COMIRNATY: Periodic safety update report assessment
19th December 2021 to 18th June 2022

This document consists of:

1. The PRAC assessment report of the Comirnaty periodic safety update report (PSUR) covering the period 19th December 2021 to 18th June 2022, and;

2. The Comirnaty PSUR itself.

The PSUR is a pharmacovigilance document intended to provide an evaluation of the risk-benefit balance of the medicinal product during the reference period mentioned above.

The objective of the PSUR is to present a comprehensive and critical analysis of the risk-benefit balance of the product, taking into account new or emerging safety information in the context of cumulative information on risk and benefits. The marketing authorisation holder is legally required to submit PSURs at defined time points after the authorisation of a medicinal product.

EMA’s safety committee, the PRAC, assesses information in the PSUR to determine if there are new risks identified for a medicine and/or if its risk-benefit balance has changed. The outcome of this assessment is summarised in the PRAC assessment report of the PSUR.

The PSUR and the PRAC assessment report of the PSUR include information about suspected side effects, i.e. medical events that have been observed following the use of the vaccine, but which are not necessarily related to or caused by the vaccine itself. Information on suspected side effects should not be interpreted as meaning that the vaccine or the active substance causes the observed event or is unsafe to use.

Only a detailed evaluation and scientific assessment of all available data, as described in the PRAC assessment report of the PSUR, can determine the impact of new data on the benefits and risks of a medicine.

Further information on the safety of COVID-19 vaccines and on PSUR submission and assessment is available on the EMA website.

This document has been redacted for commercially confidential information (CCI) and protected personal data (PPD) in accordance with applicable legislation and guidance.
PRAC PSUR assessment report

Procedure no.: EMEA/H/C/PSUSA/00010898/202206

Active substance(s): tozinameran (COMIRNATY)

Period covered by the PSUR: 19/12/2021 To: 18/06/2022

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<th>Centrally authorised Medicinal product(s):</th>
<th>Marketing Authorisation Holder</th>
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<tr>
<td>For presentations: See Annex A</td>
<td>BioNTech Manufacturing GmbH</td>
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Procedure resources

PRAC Rapporteur: Name: Menno van der Elst
<table>
<thead>
<tr>
<th>Procedure resources</th>
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<tr>
<td>Contact person - PRAC Rapporteur</td>
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<td></td>
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<td>EMA Procedure Lead</td>
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# Table of contents

1. Background information on the procedure ................................................. 4
2. Assessment conclusions and actions ............................................................. 4
3. Recommendations .......................................................................................... 5
4. Issues to be addressed in the next PSUR or as a post-authorisation measure (PAM) ........................................................................................................ 5
5 PSUR frequency .................................................................................................. 6
1. **Background information on the procedure**

This is the assessment of PSUR(s) submitted in accordance with the requirements set out in the list of Union reference dates (EURD list) for tozinamern (COMIRNATY).

2. **Assessment conclusions and actions**

The MAH submitted the 3rd EU Periodic Safety Update Report (PSUR) for Comirnaty (dated 18 Aug 2022) covering the period 19 Dec 2021 to 18 Jun 2022.

During the reporting interval, the posology recommendations for the booster use were updated to "Individuals 12 years of age and older", further details on heterologous boosting were provided and the boosting interval was shortened to at least 3 months after completion of the primary series (EMEA/H/C/005735/II/0093, EMEA/H/C/005735/II/0104 and EMEA/H/C/005735/II/0111).

Comirnaty (tozinamern) is indicated for active immunisation to prevent COVID-19 caused by SARS-CoV-2, in individuals aged 06 months and older.

During the reporting interval, an estimated 843,724,061 doses of Comirnaty were administered. Cumulatively, an estimated 2,693,922,584 doses of Comirnaty were administered.

There were no marketing authorisation withdrawals for safety reasons during the reporting interval.

During the reporting interval, the following signals were evaluated, not to be determined risks, and no new important safety issue was identified based on the data provided in the PSUR:

- Appendicitis, Hemolytic anemia, Uveitis, Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders, Capillary leak syndrome, Corneal graft rejection, Vasculitis, Cerebral venous sinus thrombosis, Lymphocytic colitis, Chronic urticaria, Polymyalgia rheumatica, Subacute thyroiditis, Cerebrovascular accident/Stroke, Amenorrhea, Loss of/altered taste and smell, and Irritability.

The following were ongoing signals during the reporting interval:

- Heavy menstrual bleeding - Please refer to the separate signal procedure heavy menstrual bleeding (EMEA/H/C/005735/SDA/053- EPITT 19783).

- Hearing loss and tinnitus - Through 18 Jun 2022, 212 HCP confirmed hearing loss and tinnitus cases (27 cases possible related to Comirnaty exposure, 110 cases unlikely, 75 cases unassessable), 295 HCP confirmed tinnitus cases (2 cases possible related, 161 cases unlikely, 132 cases unassessable), 352 HCP confirmed hearing loss cases (6 cases possible, 153 cases unlikely, 193 cases unassessable), and four relevant articles were retrieved. Despite the 27 cases reporting hearing loss and tinnitus and the 2 cases reporting tinnitus and the 6 cases reporting hearing loss that were considered possible related to Comirnaty exposure, there seems to be no causal association between Comirnaty exposure and occurrence of hearing loss and/or tinnitus in the post-marketing cases. In the observed to expected analyses all O/E ratios were below 1. In the clinical trial, there were more reports of hearing loss and of tinnitus in the placebo group compared to the Comirnaty group. The MAH’s conclusion is endorsed that a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time.

During the reporting interval, there were post-approval regulatory requests for the following topics for which no safety signal was identified based on the information provided in the PSUR:

- Acquired haemophilia, Autoimmune hepatitis, IgA nephropathy.
During the reporting interval, there were no changes in the important risks and missing information for the Comirnaty European Risk Management Plan (EU-RMP) version 4.0.

Based on the evaluation of the interval data provided, new important safety information for Comirnaty has emerged during the reporting period: Dizziness (with frequency Unknown) should be added as an ADR in section 4.8 of the Comirnaty SmPC. However, as the MAH already submitted a variation to amend the Comirnaty product information accordingly (procedure EMEA/H/C/005735/II/0152), this PSUSA procedure can be concluded with maintenance of the marketing authorisation(s).

The benefit-risk balance for the use of Comirnaty in its authorised indication remains unchanged.

3. Recommendations

Based on the PRAC Rapporteur review of data on safety and efficacy, the PRAC considers that the risk-benefit balance of medicinal products containing tozinameran (COMIRNATY) remains unchanged and therefore recommends the maintenance of the marketing authorisation(s).

4. Issues to be addressed in the next PSUR or as a post-authorisation measure (PAM)

The MAH should also address the following issues in the next PSUR:

1. In future PSURs, the MAH should only report on the number of processed cases downloaded from EudraVigilance if the percentage processed cases of the total downloaded cases decreases below 99% as presented in the current 3rd PSUR.
2. The MAH should continue to closely monitor MIS-C/-A as outlined in PRAC's signal recommendation (EPITIT 19732) and all new cases of MIS-C/-A should be reported in the future PSURs.
3. The MAH should focus on the analysis of myocarditis/pericarditis cases on aspects of these ADRs not fully known or addressed in the Comirnaty product information (myocarditis/pericarditis is already an ADR stated in section 4.8), and if the warning in section 4.4 regarding myocarditis/pericarditis is still in line with current knowledge. Therefore, the myocarditis/pericarditis analysis should focus more on information concerning the course, subsequent dosing, outcome and possible risk factors (such as age of the participant) of the myocarditis/pericarditis cases following Comirnaty exposure.
4. For future PSURs in the section 'Evaluation of AESIs', the cardiovascular AESIs, haematological AESIs, dermatological AESIs, facial paralysis, hepatic AESIs, musculoskeletal AESIs, other AESIs, respiratory AESIs, vasculitic events, local adverse reactions, and/or severe reactogenicity should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal.
5. The vaccination stress/anxiety related ADRs are considered well documented and adequately managed in clinical practice, and therefore could be removed from section 'Evaluation of other risks (not categorised as important)' in future PSURs.
6. For future PSURs in the section 'Evaluation of special situations', the overdose, abuse, misuse, and drug dependency, occupational exposure, off-label use, and/or unexpected therapeutic effect should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal.
7. For future PSURs in the section 'Update on special patient populations', the use in frail patients with co-morbidities, and/or interactions with other vaccines should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal.
8. After the DLP of the 3rd PSUR, Comirnaty has been variant updated (bivalent vaccines), approved and is currently being used in the EU. In light of this, in the next PSUR, the MAH is requested to ensure a transparent presentation of cases reported from the newly approved variant vaccines, as well as a continuous comparison of the safety issues identified with the evidence available for the original vaccine. Any differences in occurrence of safety issues, when comparing the variant updated vaccines with the original monovalent Comirnaty or, indeed, when comparing the two different variant updated bivalent Comirnaty vaccines, should be discussed.

9. The MAH should present cumulative analyses of dyspnoea, palpitations and tachycardia/heart rate increase, with special focus on the duration of the events not considered stress/anxiety-related reactions. The MAH should evaluate whether these events should be added in the section 4.8 of the Comirnaty SmPC.

10. Concerning hearing loss, the MAH is requested in future reviews of cases reporting hearing loss to conduct the analysis using the Brighton Collaboration Criteria for sensorineural hearing loss, if applicable.

11. Concerning post orthostatic tachycardia syndrome, the MAH is requested to discuss the publication of "Kwan, A.C., Ebinger, J.E., Wei, J. et al. Apparent risks of postural orthostatic tachycardia syndrome diagnoses after COVID-19 vaccination and SARS-Cov-2 Infection. Nat Cardiovasc Res (2022). https://doi.org/10.1038/s44161-022-00177-8" concerning post orthostatic tachycardia syndrome and Comirnaty exposure and, if applicable, to perform a cumulative review on the association between Comirnaty and post orthostatic tachycardia syndrome. This should include, but not be limited to, information from post-marketing cases, clinical trials, mechanistic studies and literature. The MAH should consider the need for an update of the product information and/or RMP.

5 PSUR frequency

☒ No changes to the PSUR frequency.

The current 6-month frequency for the submission of PSURs should remain unchanged.
Annex: PRAC Rapporteur assessment comments on PSUR
1. PSUR Data

1.1. Introduction

The MAH submitted the 3rd PSUR for BNT162b2 (Comirnaty) covering the period 19 Dec 2021 to 18 Jun 2022, which is assessed in this report.

The active substance of BNT162b2 is highly purified single-stranded, 5’-capped mRNA produced using a cell-free in vitro transcription from the corresponding DNA template, encoding the viral spike protein of SARS-CoV-2. The nucleoside-modified mRNA is formulated in LNPs, which enable delivery of the RNA into host cells to allow expression of the SARS-CoV-2 S antigen. The vaccine elicits both neutralizing antibody and cellular immune responses to the spike antigen, which may contribute to protection against COVID-19.

BNT162b2 was approved in the EU through a centralised procedure (conditional approval) on 21 December 2020 and is currently indicated for active immunization to prevent COVID-19 caused by SARS-CoV-2 virus, in people aged 5 years and older.

The vaccine is a white to off white frozen solution, is administered intramuscularly in the deltoid muscle and is available in 3 presentations:

<table>
<thead>
<tr>
<th>Purple cap (for 12 years of age and older)</th>
<th>Grey cap (for 12 years of age and older)</th>
<th>Orange cap (for age 5 years to &lt;12 years)</th>
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<tr>
<td>Concentrate for dispersion for Injection</td>
<td>Dispersion for Injection</td>
<td>Concentrate for dispersion for Injection</td>
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<td>30 micrograms/dose</td>
<td>30 micrograms/dose</td>
<td>10 micrograms/dose</td>
</tr>
<tr>
<td>Requires dilution</td>
<td>Do not dilute</td>
<td>Requires dilution</td>
</tr>
<tr>
<td>PBS/Sucrose presentation</td>
<td>Tris/Sucrose presentation</td>
<td>Tris/Sucrose presentation</td>
</tr>
</tbody>
</table>

**Individuals aged 12 years and older**

The 2 formulations (purple cap and grey cap) are administered as 30 μg/dose as a primary series of 2 doses (0.3 mL each) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose (third dose) may be administered approximately 6 months after the second dose in individuals 16 years of age and older.

**Individuals aged 5 through 11 years**

The Tris/Sucrose formulation (orange cap) is administered after dilution as a primary series of 2 doses (0.2 mL) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose may be administered at least 6 months after the second dose.

On 17 June 2022, an additional formulation was approved first in the United States (US): the paediatric Tris/Sucrose presentation - maroon cap (3 micrograms/dose) for individuals aged between 6 months and 4 years. This is a concentrate for dispersion for injection, to be administered after dilution intramuscularly in the anterolateral aspect of the thigh (or in the deltoid muscle in individuals 1 year of age and older) as a primary series of 3 doses (0.2 mL). The initial 2 doses are administered 3 weeks apart followed by a third dose administered at least 8 weeks after the second dose.

No changes to the product information were proposed as part of the submission of the PSUR.
Rapporteur assessment comment:

Of note, after DLP of the current PSUR: Comirnaty is currently also available as two adapted vaccines (only to be used in people aged 12 years and older who have received at least a primary vaccination course against COVID-19):

- Comirnaty Original/Omicron BA.1 contains tozinameran and riltozinameran, another mRNA molecule with instructions for producing a protein from the Omicron BA.1 subvariant of SARS-CoV-2. (procedure EMEA/H/C/005735/II/0140)

- Comirnaty Original/Omicron BA.4-5 contains tozinameran and famtozinameran, another mRNA molecule with Instructions for producing a protein from the Omicron BA.4 and BA.5 subvariants of SARS-CoV-2. (procedure EMEA/H/C/005735/II/0143)

The Comirnaty indication was extended to children 6 months - 4 years old (Tris/Sucrose presentation 3 micrograms/dose). (procedure EMEA/H/C/005735/X/0138)

An EU procedure is ongoing concerning the extension application to add a new strength of 5/5 µg (tozinameran, famtozinameran) for children between 5 to 11 years of age. (procedure EMEA/H/C/005735/X/0147)

1.2. Worldwide marketing authorisation status

BNT162b2 received first temporary authorisation for emergency supply under Regulation 174 in the UK on 01 December 2020.

BNT162b2 received first regulatory conditional marketing authorisation approval for use in individuals 16 years and older in Switzerland on 19 December 2020. In the European Union, conditional marketing authorisation was granted on 21 December 2021.

Overall BNT162b2 is authorised in 104 countries/regions. BNT162b2 is authorised for the following formulations:

- PBS/Sucrose – Purple cap 30 µg formulation:
  - in individuals aged 16 years and older in 103 countries including full (5), conditional (49), EUA and other type of approvals (52).
  - in individuals aged between 12 and 15 years in 81 countries including full (2), conditional (46), EUA and other type of approvals (34).

- Tris/Sucrose formulation:
  - Grey cap: at the dosage of 30 µg formulation in individuals aged 12 years and older in 73 countries including full (3), conditional (344), EUA and other type of approvals (28).
  - Orange cap: at the dosage of 10 µg formulation in individuals aged between 5 and 11 years in 79 countries including full (2), conditional (43), EUA and other type of approvals (35).
  - Maroon cap: at the dosage of 3 µg formulation in individuals aged between 6 months and 4 years in the US with EUA.
  - The booster dose has received approvals in 83 countries including full (3), conditional (46), EUA and other type of approvals

Rapporteur assessment comment:

The provided information regarding the worldwide marketing authorisation status is noted.
1.3. Overview of exposure and safety data

1.3.1. Actions taken in the reporting interval for safety reasons

During the reporting period, no actions have been taken with respect to BNT162b2 for safety reasons, either by a HA or by the MAH.

Rapporteur assessment comment:
The provided information is noted.

1.3.2. Changes to reference safety information

The reference safety information (RSI) for this PSUR is the Core Data Sheet (CDS) version 13.0 dated 10 May 2022, which is located in Appendix 1 of the PSUR. The 4 previous CDS versions (version 9.0 dated 02 Dec 2021, version 10.0 dated 21 Dec 2021, version 11.0 dated 14 Jan 2022 and version 12.0 dated 23 Mar 2022) were also in effect during the reporting interval.

Safety-related changes to the RSI are presented in Appendix 1.1 of the PSUR.

Rapporteur assessment comment:
The EU SmPC of Comirnaty (version 10 Aug 2022 which is after the PSUR DLP of 18 June 2022) is in line with the CDS.

1.3.3. Estimated exposure and use patterns

Clinical trials

Cumulatively, 66,656 participants have participated in the BNT162b2 clinical development program comprising several clinical candidates:

- BNT162b2: 59,260 participants of which 33,096 had received BNT162b2; 25,205 had received BNT162b2 post-unblinding and had received placebo before; 959 had received BNT162b2/placebo.
- Variant vaccines based on BNT162b2: 1836 participants of which 747 had received BNT162b2 (B.1.351); 372 had received BNT162b2 (B.1.617.2); 697 had received BNT162b2 (B.1.1.7 + B.1.617.2); 20 had received BNT162b2 (B.1.1.7).
- Early development candidates: 633 participants of which 30 had received BNT162a1; 411 had received BNT162b1; 96 had received BNT162b3; 96 had received BNT162c2.
- Blinded therapy: 7044 participants.
- Placebo: 5871 participants.

Of note, BNT162b2 is also being utilised in another Pfizer clinical development program (B747): 372 participants received BNT162b2 as a study vaccine or as a comparator in the clinical study B7471026.

Post-marketing exposure

In the current PSUR, the following regulatory requests about the exposure and number of third doses administered are addressed:
- EMEA/H/C/005735/MEA/002.8 (9th SSR), "The MAH should provide an estimate of the exposure of "third doses" in future PSURs separately (reporting period and cumulatively), if applicable."

- EMEA/H/C/005735/MEA/002.10 (11th SSR), "The MAH is requested to report the total number of administered Comirnaty dose 3 in the EU/EEA, per country, and by age group."

**MAH's response:** It is not possible to determine with certainty the number of subjects who received BNT162b2 during the period of this review, and this applies also to the "third doses".

The total number of the BNT162b2 third doses administered, downloaded from the HA’s websites (EMA, PMDA and FDA) is provided in Table 9 through Table 13 of the PSUR (only table 9 and 13 are reproduced here). Details for the cumulative number of third doses administered by age group and during the interval period in the EU/EEA countries are shown in Table 9 and in Table 13:

### Table 9. EU/EEA – Cumulative Number of BNT162b2 Administered Doses by Age Group and Dose Number

<table>
<thead>
<tr>
<th>Age Group</th>
<th>1st Dose</th>
<th>2nd Dose</th>
<th>Dose Unknown</th>
<th>3rd Dose a</th>
<th>4th Dose b</th>
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<tr>
<td>&lt; 18 years c</td>
<td>13638393</td>
<td>11718336</td>
<td>982</td>
<td>1885318</td>
<td>1839</td>
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<tr>
<td>0 – 4 years d</td>
<td>6570</td>
<td>5512</td>
<td>0</td>
<td>123</td>
<td>0</td>
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<tr>
<td>5 – 9 years</td>
<td>2400265</td>
<td>1552004</td>
<td>101</td>
<td>1101</td>
<td>0</td>
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<td>10 – 14 years</td>
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<td>4075502</td>
<td>420</td>
<td>199566</td>
<td>107</td>
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<td>15 – 17 years</td>
<td>3551465</td>
<td>3307482</td>
<td>704</td>
<td>410503</td>
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<td>18 – 24 years</td>
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<td>10563008</td>
<td>4035</td>
<td>5272098</td>
<td>13263</td>
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<tr>
<td>25 – 49 years</td>
<td>51444284</td>
<td>49059427</td>
<td>36983</td>
<td>25414999</td>
<td>112807</td>
</tr>
<tr>
<td>50 – 59 years</td>
<td>23719359</td>
<td>23084094</td>
<td>25646</td>
<td>14917669</td>
<td>115305</td>
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<tr>
<td>60 – 69 years</td>
<td>16347236</td>
<td>16155340</td>
<td>28333</td>
<td>16472372</td>
<td>508401</td>
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<td>70 – 79 years</td>
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<td>15485654</td>
<td>21790</td>
<td>15020989</td>
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<td>&gt; 80 years</td>
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<td>11939294</td>
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<td>935314</td>
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<td>223231140</td>
<td>126250</td>
<td>151603079</td>
<td>9331517</td>
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a. Indicates Dose Additional 1 in the ECDC webpage.
b. Indicates Dose Additional 2 in the ECDC webpage.
c. Data from 19 countries.
d. Data from 13 countries.
e. Data from 17 countries.
f. Data from 18 countries.
g. Data from 27 countries.
h. Data from 30 countries.

Cumulative period up to 2022 week 24 (up to 19 June 2022) – Downloaded on 15 June 2022
Table 13. EU/EEA – Interval Number of BNT162b2 Administered Doses by Age Group and Dose Number

<table>
<thead>
<tr>
<th>Age Group</th>
<th>1st Dose</th>
<th>2nd Dose</th>
<th>Dose Unknown</th>
<th>3rd Dose&lt;sup&gt;a&lt;/sup&gt;</th>
<th>4th Dose&lt;sup&gt;b&lt;/sup&gt;</th>
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<tr>
<td>&lt; 18 years</td>
<td>4421971</td>
<td>423201</td>
<td>611</td>
<td>1833167</td>
<td>1821</td>
</tr>
<tr>
<td>0 – 4 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5374</td>
<td>5605</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>5 – 9 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>203874</td>
<td>153479</td>
<td>101</td>
<td>1055</td>
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<td>10 – 14 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>126211</td>
<td>1663408</td>
<td>257</td>
<td>192672</td>
<td>101</td>
</tr>
<tr>
<td>15 – 17 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>160024</td>
<td>272676</td>
<td>290</td>
<td>383121</td>
<td>254</td>
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<tr>
<td>18 – 24 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>366151</td>
<td>598222</td>
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<td>4410546</td>
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<td>25 – 49 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1067139</td>
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<td>8151219</td>
<td>113819</td>
</tr>
<tr>
<td>60 – 69 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>262799</td>
<td>474033</td>
<td>1789</td>
<td>5777707</td>
<td>506630</td>
</tr>
<tr>
<td>70 – 79 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>163055</td>
<td>283448</td>
<td>993</td>
<td>2887858</td>
<td>842320</td>
</tr>
<tr>
<td>&gt; 80 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td>146027</td>
<td>229916</td>
<td>661</td>
<td>1652184</td>
<td>934068</td>
</tr>
<tr>
<td>Age Unknown&lt;sup&gt;c&lt;/sup&gt;</td>
<td>18890</td>
<td>12175</td>
<td>11</td>
<td>14127</td>
<td>11</td>
</tr>
<tr>
<td>EEA – All&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5369310</td>
<td>10625902</td>
<td>9845</td>
<td>69607125</td>
<td>9310883</td>
</tr>
</tbody>
</table>

<sup>a</sup> Indicated as Dose Additional 1 in the ECDC webpage.
<sup>b</sup> Indicated as Dose Additional 2 in the ECDC webpage.
<sup>c</sup> Data from 19 countries.

Interval reporting period including 2021 week 51 through 2022 week 24 (up to 19 June 2022) – Downloaded on 18 June 2022.


Rapporteur assessment comment:

The MAH reported as requested the total number of administered third and fourth doses of Comirnaty, cumulatively and during the interval period, for EU-EEA, US and Japan.

Cumulatively, in the EU-EEA an estimated:

- total of 151,603,079 third doses of Comirnaty were administered and during the interval period 69,607,125 third doses of Comirnaty.
- total of 9,331,517 fourth doses of Comirnaty were administered and during the interval period 9,310,883 fourth doses of Comirnaty.

The MAH should continue to report on the administered 1<sup>st</sup>, 2<sup>nd</sup>, 3<sup>rd</sup>, 4<sup>th</sup>, etc. doses of Comirnaty as presented above in future PSURs.

Issue solved

Worldwide exposure:

- approximately 3,555,998,805 doses of BNT162b2 were shipped worldwide from the receipt of the first temporary authorisation for emergency supply on 01 Dec 2020 through 18 Jun 2022, corresponding to 2,693,922,584 estimated administered doses.
- approximately 1,115,282,160 doses of BNT162b2 were shipped worldwide during the current reporting interval from 19 Dec 2021 through 18 Jun 2022, corresponding to 843,724,061 estimated administered doses.
• overall, through 18 Jun 2022, a total of 143,844,450 adult Tris/Sucrose doses were shipped worldwide.
• overall, through 18 Jun 2022, a total of 229,269,400 paediatric Tris/Sucrose doses were shipped worldwide.

**Rapporteur assessment comment:**
Cumulatively, worldwide a total of 2,693,922,584 doses of Comirnaty were administered.
During the reporting period, in the EU-EEA a total of 202,094,207 doses of Comirnaty were administered and cumulatively 647,275,250 doses.

### 1.3.4. Data in summary tabulations

Response to the PRAC request 1 from the 2nd PSUR (EMEA/H/C/PSUSA/00010898/202112):

*The MAH is requested to continue to report on the number of processed cases downloaded from EudraVigilance in the PSUR.*

**MAH's response:** During the reporting period, 311,460 cases were downloaded from EudraVigilance and 309,455 cases (99.4% of the total downloaded cases, 93,394 serious, 216,060 non serious and 1 unknown) were included in the data tabulations presented in the PSUR.

The remaining 2005 cases downloaded from EudraVigilance in the reporting period are not included in the data tabulations of this PSUR as they have not yet completed case processing; these include reports downloaded immediately prior to the PSUR data lock point. These reports will be included in the subsequent PSURs as Pfizer applies a late condition process that retrieves from the global safety database cases not included in the previous PSURs.

Of the 2005 case reports from EudraVigilance not included in this PSUR, 940 were serious and 1065 were non serious.

The table below provides updates on the corrective actions that have been or are being initiated with progress update from the data lock point (DLP) of the PSUR # 2 (18 December 2021) through the DLP of the PSUR # 3 (18 June 2022) to manage the volume of adverse event cases received.
<table>
<thead>
<tr>
<th>Description of Action</th>
<th>Case Type(s) within scope of action</th>
<th>Action status (proposed / completed / ongoing)</th>
<th>Completion Date/Due Date</th>
<th>Responsible party (internal Pfizer group and/or third-party)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Route to Distribution (RTD) Phase 2 BOT enabled data entry that assigns a case to a single user and subsequent routine to distribution</td>
<td>Initial and follow-up Health Authority (HA) Non-serious post-marketing (PM) cases for COVID-19 vaccine</td>
<td>Completed</td>
<td>03 March 2022</td>
<td>Pfizer</td>
</tr>
<tr>
<td>Further expand the in-scope case type for RTD</td>
<td>Initial and follow-up HA non-serious non-interventional cases where reporter causality is related or unknown</td>
<td>Ongoing</td>
<td>11 April 2022</td>
<td>Pfizer</td>
</tr>
<tr>
<td>Pfizer continuously assess resource needs. Following implementation of technology and process efficiency and through continuous monitoring of the adverse event (AE) reporting rate for COVID-19 vaccine, continue to assess the need for and onboard additional resources to help manage the increased volume of AE reports received associated with the COVID-19 vaccine</td>
<td>N/A</td>
<td>Ongoing: As of 31 December, &gt; 2700 contractor and vendor resources have been onboarded. Resources onboarded are in production as of 07 March 2022</td>
<td>N/A</td>
<td>Pfizer</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment:**

The MAH stated that the number of processed cases downloaded from Eudravigilance in the current 3rd PSUR is 309,455 cases (99.4% of the total downloaded cases, 93,394 serious, 216,060 non serious and 1 unknown). This is considered an improvement compared to the previous update provided in the previous second PSUR (96.7% of the total downloaded cases).

After DLP, on 15 Jul 2022 the MAH responded to the inspection status update regarding deprioritised non-serious cases for Comirnaty and provided the second quarter 2022 (01 Apr – 30 Jun) update on the status of COVID-19 vaccine deprioritised cases which demonstrated completed processing of all deprioritised cases according to the committed timelines:
Figure 2: Number of Non-Serious COVID-19 Vaccine Valid Cases with AESI Open In Workflow beyond Day 30

The data reflects weekly view of the number of non-serious valid COVID-19 Vaccine cases with Adverse Event of Special Interest (AESI) over day 30 which are not yet completed starting from implementation of AESI monitoring. As of 30Jun2022, there are no COVID-19 Vaccine Valid Cases with AESI beyond day 60 open in workflow.

Figure 3: Number of Serious COVID Vaccine Cases beyond day 15 Open In Workflow

The data reflects weekly view of the number of serious valid COVID Vaccine cases over day 15 not yet completed for which the reportability assessment was in progress. Timeliness with expedited reporting compliance is continually monitored and remains stable. The timeline shows volume pre-CAPA implementation to current status as of 30Jun2022.

In future PSURs, the MAH should only report on the number of processed cases downloaded from EudraVigilance if the percentage processed cases of the total downloaded cases decreases below 99% as presented in the current PSUR. **Request for next PSUR**

**Issue solved**

During the reporting period, a total of 508,351 case reports (668 from CT and 507,683 from PM) containing 1,597,673 events were retrieved, compared to 658,249 case reports in the previous second PSUR.

**Clinical trial data**

During the reporting period, in the CT dataset, the number of male participants was slightly higher than female (53.9% vs 45.5%); the number of SAEs experienced by male participants is slightly higher than female (482 vs 391); in the 18 - 30 years and the 31 - 50 years age groups, the number of SAEs reported in females was higher than in males, while in the paediatric population, in 51-64 years and in the elderly (≥ 65 years) age groups, the SAEs reported in male participants was higher than in females.
A total of 879 SAEs were reported in 668 cases.

The overall safety evaluation includes a review of the most frequently reported serious events by SOC and PT for events reported in ≥2% of all clinical trial cases during the reporting interval as compared to the cumulative period through 18 June 2022 (Table 16).

Table 16. Clinical Trial Data: Serious Events Reported in ≥2% Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>MedDRA PT</th>
<th>Reporting Period</th>
<th>Cumulatively through 18 Jun 2022</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>19 Dec 2021 - 18 Jun 2022</td>
<td></td>
</tr>
<tr>
<td></td>
<td>All Cases*</td>
<td>BNT162b2 / b2x01 / BT Cases</td>
<td>All Cases*</td>
</tr>
<tr>
<td></td>
<td>(N=879)</td>
<td>(N=668)</td>
<td>(N=2426)</td>
</tr>
<tr>
<td></td>
<td>AE(56%)</td>
<td>AE(56%)</td>
<td>AE(45%)</td>
</tr>
<tr>
<td></td>
<td>(n=504)</td>
<td>(n=557)</td>
<td>(n=3191)</td>
</tr>
<tr>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
</tr>
</tbody>
</table>

Injury, poisoning and procedural complications:
- Maternal exposure during pregnancy:
  - Condition aggravated: 25 (3.7) 25 (3.8) 121 (5.0) 113 (5.0)

General disorders and administration site conditions:
- Condition aggravated: 24 (3.6) 23 (3.5) 79 (3.3) 72 (3.2)

Infections and infestations:
- Pneumonia: 17 (2.5) 17 (2.6) 56 (2.3) 54 (2.4)
- Gastroenteritis: 15 (2.3) 15 (2.3) 22 (0.9) 21 (0.9)
- Appendicitis: 14 (2.1) 13 (2.0) 38 (2.4) 53 (2.3)

Cardiac disorders:
- Atrial fibrillation: 16 (2.4) 16 (2.4) 47 (1.9) 46 (2.0)

Nervous system disorders:
- Cerebrovascular accident: 13 (2.0) 13 (2.0) 40 (1.6) 39 (1.7)

Neoplasms: benign, malignant and unspecified (incl cysts and polyps):
- Prostate cancer: 13 (2.0) 13 (2.0) 32 (1.3) 32 (1.4)

a. Includes BNT162b2 (b2), BNT162b2x01 (b2x01), BT, and Placebo.
b. Includes BNT162b1, b2, b2x01, b3, BNT162c2 (c2), BT and Placebo.
c. The varicella vaccines b1 and c2 are study drugs in study BNT162-01, b2x01 in Study BNT162-14 and b3 in Study BNT162-04, respectively. Please refer to Section 7 for details on these studies.
d. Reporting proportion calculated as n/N (% of cases) in the current reporting period or cumulatively.
e. Reported as serious occurrence as associated to SAEs. This PT is coded in maternal cases, and in foetal cases when a foetal AE is reported. For associated SAEs, refer to Section 16.3.5.3, Use in Pregnant/Lactating Women.

AE = Adverse Event; AERP = Adverse Event Reporting Proportion; BT = Blinded Therapy; MedDRA = Medical Dictionary for Regulatory Activities; N = Number of cases; n = Number of events; PT = Preferred Term; SOC = Summary Organ Class

There were 2 SAEs assessed as related to BNT162b2 during the reporting interval:
- Dehydration was assessed as related by both the Investigator and the Sponsor.
- Abortion spontaneous was assessed as related by the Investigator and unrelated by the Sponsor.

MAH's conclusion: Based on the review of the CT cases, no new safety issues were identified.

Rapporteur assessment comment:

MAH's conclusion is endorsed that no new important safety information could be identified from the clinical trial data.
Post-authorisation data

During the reporting period, in the post-marketing dataset the number of female subjects was 2.2 times the number of male subjects (63.8% vs 29.4%); across the different age groups the ratio of female/male cases ranged between 1.1 in the less than or equal to 17 years to 2.7 in the 31-50 years group.

A total of 1,596,793 AEs (of which 439,443 were serious and 1,158,240 non-serious ) were reported in 507,683 PM cases.

The MedDRA SOCs containing the greatest number of events (≥2%) were General disorders and administration site conditions (459,731), Nervous system disorders (204,185), Musculoskeletal and connective tissue disorders (148,849), Injury, poisoning and procedural complications (130,333), Infections and infestations (82,131), Gastrointestinal disorders (81,816), Reproductive system and breast disorders (77,917), Skin and subcutaneous tissue disorders (62,405), Respiratory, thoracic and mediastinal disorders (56,663), Cardiac disorders (54,208), Surgical and medical procedures (52,531), and Blood and lymphatic system disorders (38,366).

The overall safety evaluation includes a review of the most frequently reported events by SOC and by PT for events reported in ≥2% of all post-marketing cases during the interval period as compared to the cumulative period through 18 June 2022 (table 18).
<table>
<thead>
<tr>
<th>Table 18. Post-Authorisation Data: Events Reported in ≥2% Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MedRA S&amp;O</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Nervous system disorders</td>
</tr>
<tr>
<td>Headache</td>
</tr>
<tr>
<td>Dizziness</td>
</tr>
<tr>
<td>Paraesthesia</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
</tr>
<tr>
<td>Fatigue</td>
</tr>
<tr>
<td>Pyrexia</td>
</tr>
<tr>
<td>Vaccination site pain</td>
</tr>
<tr>
<td>Chills</td>
</tr>
<tr>
<td>Malaise</td>
</tr>
<tr>
<td>Drug ineffectiveness</td>
</tr>
<tr>
<td>Vaccination site reaction</td>
</tr>
<tr>
<td>Chest pain</td>
</tr>
<tr>
<td>Pain</td>
</tr>
<tr>
<td>Arthralgia</td>
</tr>
<tr>
<td>Vaccination site swelling</td>
</tr>
<tr>
<td>Infections and infestations</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
</tr>
<tr>
<td>Myalgia</td>
</tr>
<tr>
<td>Arthralgia</td>
</tr>
<tr>
<td>Fatigue and asthenia</td>
</tr>
<tr>
<td>Limb discomfort</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
</tr>
<tr>
<td>Impromptu schedule of product administration</td>
</tr>
<tr>
<td>Self-limitedevent</td>
</tr>
<tr>
<td>Poor quality product administered</td>
</tr>
<tr>
<td>Blood and lymphatic system disorders</td>
</tr>
<tr>
<td>Lymphadenopathy</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
</tr>
<tr>
<td>Nausea</td>
</tr>
<tr>
<td>Vomiting</td>
</tr>
<tr>
<td>Diarrhoea</td>
</tr>
<tr>
<td>Surgical and medical procedures</td>
</tr>
<tr>
<td>Immunocontraception</td>
</tr>
<tr>
<td>Interaction of vaccine components</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
</tr>
<tr>
<td>Dyspnoea</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
</tr>
<tr>
<td>Rash</td>
</tr>
<tr>
<td>Cardiac disorders</td>
</tr>
<tr>
<td>Palpitations</td>
</tr>
<tr>
<td>Tachycardia</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
</tr>
<tr>
<td>Heavy menstrual bleeding</td>
</tr>
<tr>
<td>Menstrual disorder</td>
</tr>
</tbody>
</table>

a. Non-serious events are not included.
b. Listed or consistent with listed AEs in current RSI.c. Listed per case processing conventions, except for fatal cases.d. Listed per case processing conventions.e. Pts selected per case processing conventions to indicate cases reporting third/booster doses.f. Reporting proportions calculated as n/P (95% CI) of all incremental cases, incremental serious cases and all cumulative cases.g. Parensis indicate proportions were included as ADRIs in the EU-SMaP Section 4.8 as per PRAC recommendation (Procedure number EMEA/SC/0053/01/0080).h. Drug ineffectiveness represents efficacy-related conditions.i. Unlisted in the current RSI.j. Follow the listing of the associated AE.N: Number of cases; n: Number of events; MedRA=Medical Dictionary for Regulatory Activities; S&O=System Organ Class; PT=Preferred Term; AE=Adverse Event; AERP=Adverse Event Reporting Proportion; RSI=Reference Safety Information

MAH’s conclusion: Overall, during the reporting period, the serious cases represented 29.8% of the total PM; fatal outcomes occurred in less than 1% of the cases. About two-thirds of the cases occurred in female subjects and the age group 31-50 years was the group most frequently reporting AEs. The most frequently reported (≥2%) AEs (listed in the current RSI) are in majority non serious. Based on the review of the PM cases, no new safety issues were identified.
Rapporteur assessment comment:

During the reporting period, the safety signal procedures concerning **Capillary leak syndrome** (EMEA/H/C/005735/SDA/051- EPITT 19743; with outcome continue closely monitoring through routine pharmacovigilance), **Autoimmune hepatitis** (EMEA/H/C/005735/SDA/042- EPITT 19749; with outcome continue closely monitoring through routine pharmacovigilance), and **Amenorrhea** (EMEA/H/C/005735/SDA/052-EPITT 19784; with outcome follow-up requested in the PSUR) were closed.

After DLP of the current PSUR, the safety signal procedures concerning **Heavy menstrual bleeding** (EMEA/H/C/005735/SDA/053-EPITT 19783) and **Vulval ulceration** (EPITT 19840) were ongoing and the signal procedures concerning **Corneal graft rejection** (EMEA/H/C/005735/SDA/055-EPITT 19789; with outcome continue closely monitoring through routine pharmacovigilance), and **Histiocytic necrotizing lymphadenitis** (EPITT 19835; with outcome cumulative review in next PSUR) were closed.

MAH’s conclusion is endorsed that no new important safety information could be identified from the post-authorisation data.

Analysis by doses

Rapporteur assessment comment:

Please refer to the assessment of Local adverse reactions and Systemic adverse Reaction in section 2.3 'Evaluation of risks and new information' below.

Tris/Sucrose Formulation

The currently authorised presentations of BNT162b2 that use tromethamine (Tris) buffer are the following:

- Grey cap: multidose vial, formulated to provide, without need for dilution, 6 doses (each 0.3 mL dose containing 30 µg modRNA) for individuals 12 years of age and older. This presentation was approved first in the US on 29 October 2021 and then in other countries worldwide.
- Orange cap: multidose vial, formulated to provide, after dilution, 10 doses (each 0.2 mL dose containing 10 µg modRNA) for individuals 5 through 11 years of age. This paediatric presentation was approved first in the US on 29 October 2021 and then in other countries worldwide.
- Maroon cap: multidose vial, formulated to provide, after dilution, 10 doses (each 0.2 mL dose containing 3 µg modRNA) for individuals 6 months through 4 years of age. This paediatric presentation was approved first in the US on 17 June 2022.

A total of 19,789 case reports with Tris/Sucrose formulation containing 38,950 events (3.9% of the total PM dataset) fulfilled criteria for inclusion in this PSUR reporting period. Data presented in Error! Reference source not found. through Error! Reference source not found. (not reproduced here) refer to the paediatric 5-11 years old orange cap and ≥ 12 years grey cap presentations. Most cases (9055 cases, 45.8%) were reported in paediatric subjects (aged ≤ 17 years). There were no significant differences in the demographic data between paediatric subjects receiving Tris/Sucrose formulation and those receiving phosphate buffered saline (PBS)/Sucrose.
A higher percentage of medication error cases was reported in the Tris/Sucrose paediatric group and this may reflect initial difficulties in managing the new formulation. Routine pharmacovigilance activities to mitigate these medication errors, including label information (vial differentiation, instructions for reconstitution and administration, vaccination scheme, storage conditions for each formulation and available dosage), educational materials for healthcare providers, medical information call centers and traceability are listed in the approved version 5.0 of the EU-RMP adopted on 10 March 2022. The approved BLA US-PVP version 1.4.1 dated 29 April 2022 includes as routine pharmacovigilance activities label information on vial differentiation.

With regard to the reported medical events, the majority were reported in lower proportion in the Tris/Sucrose group compared to the PBS/Sucrose group although there were 7 events (Vaccination site pain, Vomiting, Abdominal pain, Diarrhoea, Rash, Urticaria, Pruritus) with a higher adverse event reporting proportion (AERP) (11.4%, 7.5%, 3.8%, 2.6%, 5.5%, 2.8%, and 2.6%, respectively) in the Tris/Sucrose paediatric group.

On review, few occurrences were serious (as important medical events – 97 for PT Vomiting, 49 for Abdominal pain, 42 for Rash, 41 for Urticaria, 24 for Diarrhoea, 20 for Pruritus, and 15 for Vaccination site pain). The clinical outcome of the serious occurrences was resolved/resolving (188), resolved with sequelae (5), not resolved (39), unknown (51), and fatal (5) at the time of reporting. In the 4 cases recording Abdominal pain, Vomiting (2 each), and Diarrhoea (1) as the fatal events, limited information was provided in 4 paediatric subjects. In these 4 cases, it is not clear whether the subjects had any underlying diseases or conditions, and date of death was unknown.

In the paediatric PBS/Sucrose cases, these events were assessed as serious as follows: PTs Vomiting (325), Abdominal pain (103), Rash (133), Urticaria (76), Diarrhoea (93), PT Pruritus (76), and PT Vaccination site pain (101). These serious events had the report proportion ≤ 0.5% of the total number of events among all paediatric PBS/Sucrose cases.

Rapporteur assessment comment:

Regarding the Tris/Sucrose formulation, 38,950 AEs from 19,789 cases were retrieved during the reporting period. 46% of the cases were from paediatric subjects aged ≤17 years and 288 AEs were considered SAEs (PT Vomiting N=97, Abdominal pain N=49, Rash N=42, Urticaria N=41, Diarrhoea N=24, Pruritus N=20, and Vaccination site pain N=15). No new important safety information could be identified.

Concerning medication errors reported in the Tris/Sucrose paediatric group, these cases have been assessed in the 13th SSR (reporting period 16 Dec 2021 – 15 Feb 2022) and 14th SSR (reporting period 16 Feb 2022 – 15 Apr 2022), and no new important safety information was identified regarding medication errors.

Booster doses (third and fourth doses)

First booster is indeed the third dose after completing a 2-dose primary series of BNT162b2 (as a homologous booster dose), or the first booster following completion of primary vaccination with another authorised COVID-19 vaccine (as a heterologous booster dose).

Second booster is indeed the fourth dose after completing a 2-dose primary series and the first booster dose with BNT162b2 (as a homologous booster dose) or the second booster dose following completion of primary vaccination and of a first booster dose with any authorised COVID-19 vaccine (as a heterologous booster dose).
Of the relevant 490 CT cases, all participants received homologous doses schedule (primary series and booster with BNT162b2). While among the relevant 117,750 PM cases, 47,759 cases received homologous doses schedule, 23,252 cases received heterologous doses schedule (primary series with a specified non-BNT162b2 COVID-19 vaccine and booster with BNT162b2), and 46,739 cases received booster doses of BNT162b2 administered after an unspecified primary COVID-19 vaccination series. The details of these cases are as follows:

**Homologous doses schedule (primary series and booster with BNT162b2)**

**Clinical trial data**

- Number of cases: 490 (BNT162b2 [441], blinded therapy [46] and placebo [3]) (73.4% of 668 cases, the total CT dataset).

**Post-authorization data**

- Number of cases: 47,759 (9.4% of 507,683 cases, the total PM dataset; 40.6% of the PM booster dataset).
- MC cases (13,848), NMC cases (33,911).
- Country of incidence (≥2%): Netherlands (15,076), UK (6998), US (4904), Austria (3222), Germany (2855), France (2377), Japan (1930), Spain (1131), Italy (1030), and Belgium (1008).
- Subjects’ gender: female (33,157), male (13,463) and unknown (1139).
- Subjects’ age in years (n=43,778), range: 0.5–120.0, mean: 45.2, median: 41.0.
- Case outcome: fatal (550), resolved/resolving (15,853), resolved with sequelae (480), not resolved (21,330), and unknown (9546).

- In 550 cases (reporting 1604 events with a fatal outcome), the reported causes of death (≥20 cases) were coded to the PTs COVID-19 (86), Vaccination failure (62), Cardiac arrest (52), COVID-19 pneumonia (46), Sudden death (31), Cardiorespiratory arrest (27), Cardiac failure, Myocardial infarction (21 each), Cerebral haemorrhage and Pulmonary embolism (20 each). Of note, in 99 cases limited information regarding the cause of death was provided (PT Death).
- Medical history (n=16,928): the most frequently (≥2% of homologous doses schedule PM cases) reported medical conditions included Disease risk factor (2101), Hypertension (2081), Asthma (1015), Drug hypersensitivity (827), Hypothyroidism (526), Seasonal allergy (509), Food allergy (483), Diabetes mellitus (480), Hypersensitivity (478), Depression (418), and Immunodeficiency (336).
- COVID-19 Medical history (n=3615): COVID-19 (2165), Suspected COVID-19 (1437), Post-acute COVID-19 syndrome (33), Exposure to SARS-CoV-2 (20), SARS-CoV-2 test positive (11), COVID-19 pneumonia (7), Asymptomatic COVID-19, Coronavirus infection (4 each), and Occupational exposure to SARS-CoV-2 (3).
- Number of events: 190,262.
- Event seriousness: serious (63,265), non-serious (127,091).
- The most reported (≥2% of homologous doses schedule PM cases) PTs were Headache (10,390), Immunisation (9993), Fatigue (9945), Malaise (8187), Myalgia (7932), COVID-19 (7123), Pyrexia (6602), Vaccination site pain (6486), Chills (6360), Lymphadenopathy (6287), Arthralgia (5138), Vaccination failure (4891), Nausea (4672), Drug ineffective (3064),
Vaccination site swelling (2813), Pain in extremity (2459), Vaccination site inflammation (2312), Vaccination site lymphadenopathy (2099), Vaccination site warmth (1900), Pain (1887), Dyspnoea (1815), Dizziness (1794), Vaccination site erythema (1774), Chest pain (1616), Axillary pain (1385), Off-label use (1171), Palpitations (1112), and Heavy menstrual bleeding (1034).

**Heterologous doses schedule (primary series with a specified non-BNT162b2 COVID-19 vaccine and booster with BNT162b2)**

**Post-authorization data**

- Number of cases: 23,252 (4.6% of 507,683 cases, the total PM dataset; 19.7% of the PM booster dataset).
- MC cases (3665), NMC cases (19,587).
- Country of incidence (≥2%): UK (9601), Netherlands (5987), Germany (1801), France (1192), Belgium (500), and US (496).
- Subjects’ gender: female (16,361), male (6296) and unknown (595).
- Subjects’ age in years (n=20,855), range: 0.3 – 102.0, mean: 46.5, median: 45.0.
- Case outcome: fatal (162), resolved/resolving (7179), resolved with sequelae (322), not resolved (12,429), and unknown (3160).
- In 162 cases (reporting 781 events with a fatal outcome), the reported causes of death (≥5 cases) were coded to the PTs Interchange of vaccine products (21), Off label use (20), Cardiac arrest (14), Sudden death (12), Cerebrovascular accident, Dyspnoea, Immunisation (11 each), Pulmonary embolism (9), Malaise, Myocardial infarction (7 each), Cerebral haemorrhage, COVID-19, Drug ineffective, Myocardial ischaemia, Pneumonia, Thrombosis (6 each), Myocarditis, Oxygen saturation decreased, and Septic shock (5 each). Of note, in 32 cases limited information regarding the cause of death was provided (PT Death).
- Medical history (n=10,734): the most frequently (≥2%) reported medical conditions included Disease risk factor (1596), Hypertension (888), Interchange of vaccine products (805), Asthma (684), Immunodeficiency (502), Hypersensitivity (289), Diabetes mellitus (278), Hypothyroidism (271), Steroid therapy (253), Depression (241), Drug hypersensitivity (229) and Clinical trial participant (224).
- COVID-19 Medical history (n=2649): Suspected COVID-19 (1346), COVID-19 (1340), Post-acute COVID-19 syndrome (22), SARS CoV 2 test positive (18), COVID-19 pneumonia (6), Coronavirus infection (4), Asymptomatic COVID-19 (2) and Exposure to SARS CoV 2 (1).
- Among the 23,252 cases reporting administration of heterologous booster dose(s) of BNT162b2 following a specified non BNT162b2 COVID-19 vaccine, the previous vaccine series consisted of:
  - 9651 subjects immunised with AstraZeneca vaccine;
  - 5334 subjects immunised with Moderna vaccine;
  - 5214 subjects immunised with unknown non Pfizer-BioNTech COVID-19 vaccine;
  - 2427 subjects immunised with Johnson and Johnson vaccine;
  - 417 subjects immunised with Coronavac (Sinovac) vaccine;
- 88 subjects immunised with SinoPharm vaccine;
- 76 subjects immunised with Sputnik vaccine;
- 29 subjects immunised with Novavax vaccine;
- 9 subjects immunised with Fiocruz vaccine;
- 2 subjects each immunised with Medicago-Clinical study and Medigen vaccine;
- 1 subject each immunised with Cansino vaccine, Covaxin vaccine, and Valneva vaccine.

- Number of events: 140,835.
- The most reported (>2% of heterologous dose schedule PM cases) PTs were Off label use (20,437), Interchange of vaccine products (20,376), Immunisation (9982), Headache (5229), Fatigue (4854), Myalgia (3412), Malaise (3362), Pyrexia (3144), Lymphadenopathy (3139), Vaccination site pain (2926), Chills (2918), Arthralgia (2578), Nausea (2382), Pain in extremity (1848), Pain (1362), Drug Ineffective (1223), COVID-19 (1192), Dizziness (1161), Vaccination site swelling (1140), Dyspnoea (1091), Chest pain (991), Axillary pain (948), Vaccination site inflammation (942), Palpitations (850), Vaccination site warmth (830), Vaccination site lymphadenopathy (815), Vaccination site erythema (751), Pruritus (630), Rash (617), Swelling (610), Heavy menstrual bleeding (608), Asthenia, Diarrhoea (565 each), Peripheral swelling (557), Paraesthesia (548), Vomiting (516), and Tachycardia (462).

**Booster doses of BNT162b2 administered after an unspecified primary COVID-19 vaccination series**

**Post-authorization data**

- Number of cases: 46,739 (9.2% of 507,683 cases, the total PM dataset; 39.7% of the PM booster dataset).
- MC cases (11,182), NMC cases (35,557).
- Country of incidence (>2%): Germany (20,876), France (6716), Japan (2922), Austria (2833), US (2553), and UK (1924).
- Subjects’ gender: female (31,045), male (13,826) and unknown (1868).
- Subjects’ age in years (n=43,405), range: 1.0 – 104.0, mean: 45.3, median: 43.0.
- Case outcome: fatal (513), resolved/resolving (18,572), resolved with sequelae (1003), not resolved (18,155), and unknown (8496).
- In 513 cases (reporting 1318 events with a fatal outcome), the reported causes of death (≥15 cases) were coded to the PTs Cardiac arrest (38), Cardiac respiratory arrest, Myocardial infarction (35 each), Sudden death (25), Pulmonary embolism (24), Cardiac failure (22), Dyspnoea (19), Cerebral haemorrhage (17), and Acute myocardial infarction (15). Of note, in 150 cases limited information regarding the cause of death was provided (PT Death).
- Medical history (n=11,782): the most frequently (>2%) reported medical conditions included Hypertension (1856), Asthma (836), Drug hypersensitivity (642), Seasonal allergy (625), Hypersensitivity (456), Diabetes mellitus (426), Hypothyroidism (384), Obesity (336), Food allergy (314), Type 2 diabetes mellitus (295), Atrial fibrillation and Depression (241 each).
Coronavirus infection, SARS CoV 2 test positive (4 each), Asymptomatic COVID-19 and Breakthrough COVID-19 (1 each).

- Number of events: 153,862.
- Event seriousness: serious (35,762), non-serious (118,147).
- The most reported (22%) PTs were Headache (8533), Lymphadenopathy (7016), Pyrexia (6893), Fatigue (6751), Vaccination site pain (5985), Immunisation (5675), Chills (4558), Myalgia (3979), Dizziness (3359), Malaise (3296), Nausea (3001), Limb discomfort (2798), Arthralgia (2615), Pain in extremity (2485), Dyspnoea (2173), Influenza (1877), Rash (1738), Drug ineffective (1563), Tachycardia (1557), Asthenia (1512), Pain (1499), Paraesthesia (1416), COVID-19 (1386), Chest pain (1381), Vaccination site swelling (1336), Vomiting (1325), Off label use (1286), Herpes zoster (1203), Menstrual disorder (1101), Diarrhoea (1062), Poor quality product administered (1061), Feeling hot (1017), Immunisation reaction (1016), Influenza like illness (1006), and Palpitations (985).

**Analysis booster doses versus primary vaccination series**

- There were 117,750 PM cases of subjects who received at least one booster dose of BNT162b2. Among the 117,750 PM cases,
  - 106,889 PM cases involved subjects who received single booster dose of BNT162b2
  - 3427 PM cases involved subjects who received >1 booster doses of COVID-19 vaccine (the latest booster dose was BNT162b2) and
  - 7434 cases involved subjects who received unknown booster dose(s) of BNT162b2.
- The most frequently (≥2%) reported clinical AEs in PM cases of subjects who received the booster dose(s) of BNT162b2 are largely reflective of reactogenicity and events associated with the immunisation process.
- The most frequently (≥2%) reported clinical AEs in PM cases of subjects who received booster dose(s) of BNT162b2 were consistent with those reported in subjects receiving primary vaccination series, as shown in Table 25.
  - A higher AERP rate was observed for 9 PTs (Lymphadenopathy [14.0% vs 3.8%], Malaise [12.6% vs 4.6%], Chills [11.8% vs 5.1%], Vaccination site swelling [4.5% vs 1.4%], Vaccination site erythema [2.9% vs 1.0%], Vaccination site lymphadenopathy [2.9% vs 0.2%], Vaccination site inflammation [2.8% vs 0.3%], Axillary pain [2.7% vs 0.5%], and Vaccination site warmth [2.4% vs 0.3%]) was observed in subjects who received the booster dose(s) of BNT162b2 compared to subjects receiving the primary vaccination series. This is consistent with the known BNT162b2 safety profile (as per the RSI), where higher rates of lymphadenopathy and reactogenicity reactions in booster doses versus primary doses were observed in interventional clinical studies.
  - No clinically significant differences were noted in the other events.

**MAH's conclusion:** Based on the review of the cases reported with the booster dose(s), no new safety issues were identified.

**Rapporteur assessment comment:**

The MAH reported on the safety profile of Comirnaty in a homologous doses schedule (primary series and booster with BNT162b2), in a heterologous doses schedule (primary series with a specified non-BNT162b2 COVID-19 vaccine and a booster with BNT162b2) and in a schedule when the third/booster
doses of BNT162b2 were administered after an unspecified primary COVID-19 vaccination series. 

MAH’s conclusion is endorsed, that at the moment no new important safety information was identified in the cases reported with the booster dose(s).

The number of persons receiving multiple booster doses (i.e., homologous, heterologous, different strains) is increasing whereas the impact on safety and efficacy remains uncertain. In addition the impact (including long-term) of repeatedly (e.g., yearly) receiving booster doses (with or without strain updates) also remains unknown. The MAH is requested to discuss whether ‘safety following multiple repeated booster doses (i.e., homologous, heterologous, different strains)’ should be considered as **Missing information** in the RMP, and proposals should be provided how to address this knowledge gap in ongoing or newly proposed PASSs, as applicable. As boosting schemes could involve multiple vaccine brands a joined approach between different MAHs would be welcomed. **Request for supplementary information**

### 1.3.5. Findings from clinical trials and other sources

#### 1.3.5.1. Clinical trials

**Completed clinical trials**

- Safety trials: During the reporting period, no interventional safety studies were completed with a final CSR.

- Other trials: During the reporting interval, there was a completed clinical trial (C4591017) with a final CSR (available upon request). No clinically important new information has emerged from this clinical trial.

**Ongoing clinical trials**

During the reporting period, there were 14 ongoing sponsor-initiated clinical trials.

**Safety trials:**

- PASS C4591015 [A phase 2/3, placebo-controlled, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate (BNT162b2) against COVID-19 in healthy pregnant women 18 years of age and older] is an ongoing PASS. No clinically important information has emerged from this ongoing PASS.

- PASS C4591024 [A phase 2b, open-label study to evaluate the safety, tolerability, and immunogenicity of vaccine candidate BNT162b2 in immunocompromised participants ≥2 years of age] is an ongoing PASS. No clinically important information has emerged from this ongoing PASS.

- Other trials where the primary aim of the trial was to identify, characterise or quantify a safety hazard or confirm the safety profile of the medicinal product: None.

- Other Trials that reported new significant efficacy information, 8 ongoing clinical trials:

  - C4591001, A phase 1/2/3, placebo-controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.
○ C4591007, A phase 1, open-label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer-blinded safety, tolerability, and Immunogenicity study of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy children <12 years of age.

○ C4591031, A phase 3 master protocol to evaluate additional dose(s) of BNT162b2 in healthy individuals previously vaccinated with BNT162b2.

○ BNT162-01, A multi-site, phase I/II, 2-Part, dose-escalation trial investigating the safety and immunogenicity of four prophylactic SARS-CoV-2 RNA vaccines against COVID-19 using different dosing regimens in healthy and immunocompromised adults.


○ BNT162-04, A multi-site, phase I/II, 2-part, dose escalation trial investigating the safety and Immunogenicity of a prophylactic SARS-CoV-2 RNA vaccine (BNT162b3) against COVID-19 using different dosing regimens in healthy adults.


○ BNT162-14, A Phase II, open-label rollover trial to evaluate the safety and immunogenicity of one or two boosting doses of Comirnaty or one dose of BNT162b2s01 in BNT162-01 trial subjects, or two boosting doses of Comirnaty in BNT162-04 trial subjects.

No clinically important information has emerged from ongoing clinical trials.

Remaining Trials, 4 ongoing clinical trials:

• C4591005, A phase 1/2, placebo-controlled, randomized, and observer-blind study to evaluate the safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy Japanese adults.

• C4591020, A phase 3, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of multiple formulations of the vaccine candidate BNT162b2 against COVID-19 in healthy adults 18 through 55 years of age.

• C4591030, A phase 3, randomized, observer-blind trial to evaluate the safety and immunogenicity of BNT162b2 when co-administered with seasonal inactivated influenza vaccine (SIIV) in adults 18 through 64 years of age.

• BNT162-17, A Phase II trial to evaluate the safety and immunogenicity of a SARS-CoV-2 multivalent RNA vaccine in healthy subjects.

No clinically important safety information has emerged from these ongoing clinical trials.

Long-term follow-up

There is no new safety information with regards to long-term follow-up of clinical trial participants for this reporting period.

Other therapeutic use of medicinal product
BNT162b2 was also utilised in another Pfizer-sponsored clinical development program (B747). The study B7471026 "A phase 3, randomized, double blind trial to describe the safety and immunogenicity of 20 valent pneumococcal conjugate vaccine when co-administered with a booster dose of BNT162b2 in adults 65 years of age and older" was completed during the reporting period.

There was no new clinically important safety information identified for this reporting period.

**New safety data related to fixed combination therapies**

BNT162b2 is not used in fixed or multi-drug combination with other compounds.

**Rapporteur assessment comment:**

No new important safety information was identified by the MAH from the clinical (safety) trials concerning long-term follow-up, other therapeutic use of the product, or related to fixed combination therapies.

### 1.3.5.2. Findings from non-interventional studies

During the reporting period, there were 11 ongoing sponsor-initiated non-interventional studies and one non-interventional study (C4591035) was completed.

#### Completed non-interventional study

**Other study**

- Study C4591035 titled 'Coronavirus Disease 2019 (COVID-19) Vaccination and Breakthrough Infections Among Persons with Immunocompromising Conditions in the US' was completed. No new safety information emerged from this non-interventional study.

#### Ongoing non-interventional studies

**Safety Studies:**

- PASS: The non-interventional studies C4591008, C4591010, C4591012, C4591021 and C4591022 are PASS. No clinically important information has emerged from PASS.

**Other Studies, 5 ongoing non-interventional studies:**

- C4591006, General Investigation of COMIRNATY intramuscular injection (follow-up study for subjects [healthcare professionals] who are vaccinated at an early post-approval stage).
- C4591014, Pfizer-BioNTech COVID-19 BNT162b2 Vaccine Effectiveness Study - Kaiser Permanente Southern California.
- C4591019, Special investigation in the population with underlying diseases considered to increase the risk of severe illness of COVID-19.
- C4591025, A prospective, single-arm, open-label, non-interventional, multicenter to assess the safety of BNT162b2 in domestic post-marketing surveillance.
- C4591034, Patient-reported health-related quality of life associated with COVID-19: A prospective survey study on symptomatic adults confirmed with RT-PCR from outpatient settings in the US.
During the reporting period, no new significant safety information from non-interventional studies was reported.

*Rapporteur assessment comment:*

No new important safety information was identified by the MAH from non-interventional studies.

### 1.3.5.3. Information from other clinical trials and sources

*Other clinical trials*

During the reporting interval, there were 11 relevant cases that originated from non-Pfizer clinical trials. In 7 of these cases, BNT162b2 was a study drug, while in the other 4 cases the administration of BNT162b2 was concomitant.

The 6 cases originated from non-Pfizer and non-BNT trials and reported the SAEs:

- COVID-19, Drug ineffective;
- Thrombophlebitis;
- Deep vein thrombosis, Pulmonary embolism;
- Myocarditis;

The SAEs reported in these 4 cases were assessed as related to the BNT162b2 by the investigators and the MAH agreed except for the case reporting Deep vein thrombosis and Pulmonary embolism, where it was considered that there was not a reasonable possibility that the events were related to vaccine administration, based on the absence of a plausible pathophysiological mechanism.

- Immunisation, Overdose (0.5 ml of Pfizer-BioNTech COVID-19 vaccine), Ventricular tachycardia;
- Condition aggravated, Endometrial thickening.

The investigator’s assessment for Ventricular tachycardia was not provided; Endometrial thickening was considered unrelated to BNT162b2 by the investigator; in both cases the MAH considered the SAEs as unrelated to the vaccine.

During this reporting period, there was no new significant safety information reported from other non-Pfizer, non-BNT sponsored clinical trials/studies

*Rapporteur assessment comment:*

No new important safety information was identified by the MAH from other clinical trials.

*Medication errors*

*Clinical trial data*

During the reporting period, there were 2 serious cases (0.3% of 668 cases, the total CT dataset) indicative of medication errors (PTs: Accidental overdose and Inappropriate schedule of product administration). In the first case, the accidental overdose referred to paracetamol and not to BNT162b2 and in the remaining case reporting inappropriate schedule of product administration, the
investigator assessed the event as not related to BNT162b2. There was 1 serious case retrieved during the reporting period of the PSUR #2.

Post-authorisation data

The potentially relevant medication error cases during the reporting period were 66,764 (13.1%) reporting 87,307 events, compared to 33,834 relevant cases (5.1%) analysed in the PSUR #2.

Out of the 66,764 relevant medication error cases, there were:

- 9426 cases reporting events indicative of medication errors related to Tris/Sucrose paediatric formulation, and
- 2750 cases reporting events indicative of medication errors related to Tris/Sucrose Grey cap presentation (adult/adolescent formulation).

These numbers represent a good index of the effectiveness of the routine pharmacovigilance activities implemented, considering that in the reporting period 182,231,200 paediatric Tris/Sucrose doses and 143,844,450 adult Tris/Sucrose doses were shipped worldwide.

Overall, among the 66,764 relevant medication error PM cases, 1326 cases (0.3% of the total interval cases, 2.0% of total relevant medication error cases) were considered harmful, 70 of which (0.1% of total relevant cases) were serious and most of them originated from vaccine administration issues (50 cases of 70 serious cases with harm).

The potential for medication errors with the new presentations are mitigated through the information in the vaccine labelling and available educational materials for the HCPs administering the vaccine.

Some medication errors are expected to occur despite written instructions for handling the vaccine (thawing, dilution, preparation) and educational activities for the HCPs or due to national or regional recommendations regarding dosing schedules that may differ from recommendations in the product labelling. The number and seriousness of the reported medication errors events do not indicate any trend and potential needs for any additional mitigation activity.

**Rapporteur assessment comment:**

Two serious clinical trial cases indicative of a medication error were reported and considered not related to or did not concern Comirnaty exposure.

During the reporting period, an increased number of medication errors (N=66,764 cases with an estimated 843,724,061 administered doses) was reported compared to the previous 2nd PSUR (N=10,776 cases with an estimated 635,763,682 administered doses). 9,426 of the 66,764 cases were indicative of medication errors related to Tris/Sucrose paediatric formulation, and 2,750 of the 66,764 cases to the Tris/Sucrose Grey cap presentation (adult/adolescent formulation). However, no specific trend or pattern was observed.

No new important safety information could be identified regarding reported medication errors.

**1.3.5.4. Non-clinical data**

During the reporting period, no new non-clinical safety findings were identified.
1.3.5.5. Literature

Nonclinical (Published)

A search of the Medline and Embase databases identified no nonclinical studies that presented important new safety findings for BNT162b2.

Clinical (Published)

A search of the Medline and Embase databases identified 8 clinical trials that presented important new safety findings for BNT162b2 (table 31).
Table 31. Clinical Literature Articles that Presented New Safety Information in the Reporting Interval

<table>
<thead>
<tr>
<th>Citation/Comment</th>
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<tbody>
<tr>
<td><strong>At Risk patients</strong></td>
</tr>
<tr>
<td>This article described a reduced immune response to BNT162b2 in patients treated with immunosuppressants. Section 4.4. Special warnings and precautions for use (Immunocompromised) individuals of the EU SmPC includes a warning regarding vaccination in immunocompromised patients, as follows: <em>The efficacy and safety of the vaccine has not been assessed in immunocompromised individuals, including those receiving immunosuppressant therapy. The efficacy of Comirnaty may be lower in immunocompromised individuals.</em></td>
</tr>
<tr>
<td>Use in immunocompromised patients is considered missing information for BNT162b2; please refer to Section 16.4.2 Description of Missing Information for the description of this topic and to Section 16.3.5.5 Use in Immunocompromised Patients for the summary of the cases received during the reporting period.</td>
</tr>
<tr>
<td><strong>Special Patients Population (Pregnancy)</strong></td>
</tr>
<tr>
<td>This article contributes to the growing evidence that risk of spontaneous abortion after COVID-19 vaccine immunisation during the first trimester of pregnancy is commensurate with the predicted risk in non-vaccinated pregnant women.</td>
</tr>
<tr>
<td>Use in pregnancy and while breastfeeding patients is considered missing information for BNT162b2; please refer to Section 16.4.2 Description of Missing Information for the description of this topic and to Section 16.3.5.3 Use in Pregnant/Lactating Women for the summary of the cases received during the reporting period.</td>
</tr>
<tr>
<td><strong>Efficacy and Effectiveness</strong></td>
</tr>
<tr>
<td>Please refer to Section 17.2 Newly Identified Information on Efficacy and Effectiveness for the comments on these articles and to Section 16.3.4.5 Lack of Therapeutic Efficacy for the review of the cases indicative of LOE reported in the current interval period.</td>
</tr>
<tr>
<td><strong>Other Safety Information</strong></td>
</tr>
<tr>
<td>This study suggests that the COVID-19 vaccine might be associated with increased risk of Sudden Sensorineural Hearing Loss; however, the effect size is very small. The study had various limitations and no causality assessment has been conducted. The MAH will continue to monitor using routine pharmacovigilance. Please refer to Appendix 6A.3 for further discussion of this article and for cumulative review of cases indicative of hearing loss.</td>
</tr>
</tbody>
</table>
In this study, BNT162b2 was associated with an immediate negative effect on anticoagulation control in patients treated with vitamin K antagonists. The author though cannot exclude the possibility that the effect on anticoagulation control was due to dose adjustments to avoid complications and patients themselves could have decided to decrease the dosage in the days following COVID-19 vaccination as they might be afraid for bleeding complications after intramuscular injection. This could result in a higher percentage of subtherapeutic INRs after vaccination. In addition, the authors use a surrogate variable for bleeding complications (INR >5).

The possible effects of vaccines on anticoagulation control remain debated even though several prospective studies have been performed (mostly on the effect of the influenza vaccine on anticoagulation control), but overall results were conflicting. As of now, there is no biological or pharmacological plausibility for a vaccine – drug interaction. The MAH will continue to monitor using routine pharmacovigilance. Please refer to Section 16.3.3.1.19 Thromboembolic AEsIs for the summary of cases indicative of coagulopathy received in the reporting period.

Rapporteur assessment comment:

The MAH identified 8 clinical trials that presented important new safety findings after Comirnaty exposure and provided an overview of the retrieved 8 studies grouped as a) At risk patients (N=1); b) Special patient population/Pregnancy (N=1); c) Efficacy and effectiveness (N=4) and d) Other safety information (N=2). No new important safety information could be identified regarding literature. Please refer regarding the study of Yanir et al. to the assessment in section 2 2.1.1. Hearing loss of this AR.

All Other Published Sources

In the final AR for PAM-MEA-002.11 - 12. SMSR (1st SBSR) received on 09 February 2022 (EMEA/H/C/005735/MEA/002.11), the MAH was requested to include in the 2nd SBSR the following article, published in the reporting interval of the PSUR # 3:


The above article was included and discussed in the SBSR no. 2 dated 04 March 2022.

In the final AR for PAM-MEA-002.12 13th SMSR (2nd SBSR) dated 05 April 2022 (EMA/PRAC/202255/2022), the MAH was requested to discuss the following publication regarding SSNHL in association with COVID-19 vaccination:


The abstract of the above article and the discussion are available in Appendix 6A.3.

Rapporteur assessment comment:

The study of Ouldali et al. was discussed in the 13th (2nd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.12). No new safety concern was identified.

Please refer regarding the study of Formeister et al. to the assessment in section 2 2.1.1. Hearing loss of this AR.
Unpublished manuscripts

In the final assessment report for PAM-MEA-002.12 13th SSR (2nd SBSR) dated 05 April 2022 (EMA/PRAC/202255/2022), the MAH was requested to include in the 3rd SBSR the following ACIP presentation, presented in the reporting interval of the PSUR # 3:


The above presentation on myocarditis outcomes following mRNA COVID-19 vaccination was included and discussed in the SBSR no. 3 dated 06 May 2022.

**Rapporteur assessment comment:**

The ACIP presentation was assessed in the 14th (3rd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.13) and resulted in a request for a variation to update sections 4.4 and 4.8 of the SmPC in order to update the occurrence of myocarditis because more information is available in the age group 5-11 years; and to update the statement in the SmPC section 4.4 regarding the risk of myocarditis after a third dose of Comirnaty based on real-world evidence, which was assessed in procedure EMEA/H/C/005735/II/0141.

### 1.3.5.6. Other periodic reports

During the reporting period, the MAH did not submit another PSUR for BNT162b2. However, the MAH was requested to prepare Summary Monthly Safety Update Reports (SMSRs), in accordance with the EMA coreRMP19 Guidance (EMA/544966/2020) and, as applicable, with ICH Guideline E2C (R2) Periodic Benefit-Risk Evaluation Report [Step 5, January 2013] considering the information for the evidence from post-EUA/conditional marketing authorisation approval data sources.

Following the proposal of discontinuation of SSR submission by PRAC included in the final PRAC AR of the 3rd SBSR (Report EMA/PRAC/577594/2022 dated 08 June 2022), the preparation of the SBSR was discontinued.

During the reporting period, no significant findings were identified for BNT162b2 in other periodic reports prepared by the MAH.

**Rapporteur assessment comment:**

After the 14th (3rd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.13), the submission of SSRs was discontinued.

### 1.3.5.7. Lack of efficacy in controlled clinical trials

Study C4591007 is the ongoing, randomised, placebo-controlled, Phase 1/2/3 paediatric study in healthy children from 6 months to <12 years of age. The Phase 2/3 primary immunogenicity objective in children from 6 months to <5 years of age was immunobridging the immune responses against SARS-CoV-2 wild-type strain from children 2 to <5 years and 6 months to <2 years of age in Study C4591007 compared to young adults 16 to 25 years of age in the Phase 3 efficacy Study C4591001.

Immunobridging data after Dose 2 met success criteria for the 6 months to <2 years group and did not meet geometric mean ratio (GMR) success criteria (but met seroresponse criteria) for the 2 to <5 years of group, compared to young adults 16 to 25 years of age. Given emerging real-world data in
the Omicron wave that two-dose protection against symptomatic infection was only modest, a third
dose was evaluated for children <5 years of age. Immunobridging data after Dose 3 met success
criteria for the 6 months to <5 years age group, compared to young adults 16 to 25 years of age.

The observed vaccine efficacy (VE) from at least 7 days after Dose 2 to before Dose 3 for BNT162b2 3-
µg administered to children 6 months to <5 years of age without prior evidence of SARS-CoV-2
infection before or during the vaccination regimen was 28.3% (2-sided 95% CI: 8.0%, 43.9%) based
on 163 cases in the BNT162b2 group and 113 cases in the placebo group, adjusted for surveillance
time (noting 2:1 randomisation of vaccine:placebo). In this population, observed VE against Delta and
Omicron was 70.2% (2-sided 95% CI: 27.2%, 88.5%) and 21.8% (2-sided 95% CI: -1.7%, 39.7%),
respectively. Note that most of the cases across this age population that were confirmed post-Dose 2
to before Dose 3 were reported in January 2022.

The observed VE from at least 7 days after Dose 3 to the cutoff date (29 April 2022) across the total
population of children 6 months to <5 years of age was 80.3% (2-sided 95% CI: 13.9%, 96.7%) based
on 3 cases in the BNT162b2 group and 7 cases in the placebo group, adjusted for surveillance
time (noting 2:1 randomization of vaccine:placebo). Note that all post-Dose 3 cases were reported in
February through April 2022.

The observed VE after 3 doses in children 6 months to <5 years of age in Study C4591007 is
consistent with real-world effectiveness data for older age groups, which indicate that in adolescents
(12 to 17 years of age) and adults (18 years of age and older), three doses of BNT162b2 are needed
to provide a high level of protection against symptomatic disease due to Omicron.

**Rapporteur assessment comment:**

MAH’s information regarding lack of efficacy is noted.

Please refer to procedure EMEA/H/C/005735/X/0138 for the line extension of Comirnaty 3 µg
concentrate for dispersion for injection for infants and children between 6 months to 4 years of age
(procedure EMEA/H/C/005735/X/0138).

### 1.3.5.8. Late-breaking information

On 08 July 2022 (after the DLP of this PSUR), the MAH submitted the EU RMP version 5.1 to remove
the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP
recommendation received in March 2022 with the positive opinion for the Type II Variation 87
(EMEA/H/C/005735/II/0087).

**Rapporteur assessment comment:**

Please refer to section 2.1 Summary of safety concerns of this AR below.
2. Signal and risk evaluation

2.1. Summary of safety concerns

The important risks and missing information for BNT162b2 at the beginning of the reporting interval, as per EU RMP v 4.0 adopted 26 Nov 2021:

<table>
<thead>
<tr>
<th>Important identified risks</th>
<th>Anaphylaxis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Myocarditis and Pericarditis</td>
</tr>
<tr>
<td>Important potential risks</td>
<td>Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)</td>
</tr>
<tr>
<td>Missing information</td>
<td>Use in pregnancy and while breast feeding</td>
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<td></td>
<td>Use in immunocompromised patients</td>
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<tr>
<td></td>
<td>Use in frail patients with co-morbidities (e.g. chronic obstructive pulmonary disease [COPD], diabetes, chronic neurological disease, cardiovascular disorders)</td>
</tr>
<tr>
<td></td>
<td>Use in patients with autoimmune or inflammatory disorders</td>
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<tr>
<td></td>
<td>Interaction with other vaccines</td>
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<tr>
<td></td>
<td>Long-term safety data</td>
</tr>
</tbody>
</table>

There were no changes to the safety concerns during the reporting period.

**Rapporteur assessment comment:**

During the reporting period there were no changes to the list of safety concerns in the Comirnaty RMP.

Of note, after the DLP of this PSUR, the MAH submitted additional Comirnaty RMPs:

1. **RMP Version 5.1** - to include the 6 months to <2 years and 2 years to <5 years phase 1 and phase 2/3 data from interventional clinical study C4591007 for the line extension of Comirnaty 3 μg concentrate for dispersion for injection for infants and children between 6 months to 4 years of age; and to remove the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP recommendation in procedure EMEA/H/C/005735/II/0087. (Procedure EMEA/H/C/005735/X/0138)

2. **RMP version 6.0** - to support the extension of the Indication to ≥12 years of age to receive an additional booster (fourth) dose of bivalent Omicron-(BA.1) modified vaccine. (procedure EMEA/H/C/005735/II/0140)

3. **RMP version 7.0** - to support the extension of the Indication to introduce a first or second booster dose of a bivalent Omicron variant-modified vaccine, (BNT162b2 15 μg + BNT162b2 OMI BA.4/5 15 μg, total 30 μg), given ≥3 months after the primary series or ≥4 months after the third dose in individuals ≥12 years of age. (procedure EMEA/H/C/005735/II/0143)

4. **RMP version 7.2** - to support the extension application to add a new strength of 5/5 μg (tozinameran, famtozinameran) for children between 5 to 11 years of age. (procedure EMEA/H/C/005735/X/0147)
### 2.2. Signal evaluation

- Tabular overview of signals: new, ongoing or closed during the reporting interval 19-12-2021 to 18-06-2022.

<table>
<thead>
<tr>
<th>Signal term</th>
<th>Date detected</th>
<th>Status (new, ongoing or closed)</th>
<th>Data closed (for closed signals)</th>
<th>Source or trigger of signal</th>
<th>Reason summary</th>
<th>Method of signal evaluation</th>
<th>Outcome, if closed</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hearing loss</strong></td>
<td>31May22</td>
<td>Ongoing</td>
<td></td>
<td>Enquiry from a competent authority (EMA PRAC, Health Canada).</td>
<td>Signal re-opened during the current reporting period due to Health Canada (HC) request and EMA PRAC request (per 14th Summary Safety Report [3rd bimonthly] Assessment Report). HC request was to provide a cumulative review of all cases of tinnitus and hearing loss. EMA PRAC requested MAH to perform a cumulative review on the association between sudden sensorineural hearing loss and Comirnaty exposure, including a review of the relevant new literature published after the reporting period of the 14th SSR, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Evaluation ongoing at datalock date (18 Jun 2022).</td>
</tr>
<tr>
<td><strong>Myocarditis and Pericarditis</strong></td>
<td>18May22</td>
<td>Closed</td>
<td>18May22</td>
<td>Other: Internal Reviews</td>
<td>The safety management team endorsed myocarditis and pericarditis being added as an ADR to the Core Data Sheet as data from multiple sources has consistently shown the association and increase from background, particularly in young males after the 2nd dose. The ongoing reviews of the post-authorization safety data, clinical trial data, pre-clinical trial, and O/E analysis data support the update to Section 4.8 of the CDS.</td>
<td>Postauthorization safety data, clinical study safety data, and preclinical data review, Literature review, and O/E analysis.</td>
<td>Important Identified Risk. Myocarditis and pericarditis are currently discussed in the Warnings/Precautions section 4.4 of the CDS. Now, based on the accumulating data and supported by signal management activities, myocarditis and pericarditis will be included in Section 4.8 (Undesirable effects) of the CDS.</td>
</tr>
<tr>
<td>Signal term</td>
<td>Date detected</td>
<td>Status</td>
<td>Data closed</td>
<td>Source or trigger of signal</td>
<td>Reason summary</td>
<td>Method of signal evaluation</td>
<td>Outcome, if closed</td>
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<tr>
<td>Irritability</td>
<td>31Jan22</td>
<td>Closed</td>
<td>31Jan22</td>
<td>Clinical Trial C4591007 unblinded review of data in 6 months to &lt;5-year-olds (Pfizer).</td>
<td>Unblinded review of the pediatric clinical trial reactogenicity data for the C4591007 6 months to &lt;5-year-old cohort. It was determined that &quot;irritability&quot; will be considered an ADR specific to the 6 months to &lt;2-year-old population.</td>
<td>Review of unblinded systemic reactogenicity events for doses 1 and 2 in 6 months to &lt;2-year-old recipients of BNT162b2 (compared to placebo).</td>
<td>Identified risk – considered NOT important for purposes of risk management. &quot;Irritability&quot; will be considered an ADR specific to the 6 months to &lt;2-year-old population. The CDS (and, subsequently, local labelling documents as appropriate) will be updated to include &quot;irritability&quot; as an ADR for this population.</td>
</tr>
<tr>
<td>Appendicitis</td>
<td>06Apr22</td>
<td>Closed</td>
<td>31May22</td>
<td>Enquiry from a competent authority (Singapore BoH).</td>
<td>Signal re-opened during the current reporting period in response to Singapore BoH request to provide a cumulative safety analysis on the association of appendicitis with Comirnaty, in particular in adolescents aged 12-17 years old in SBSR.#4. Singapore noted 18 local reports of appendicitis post vaccination with Comirnaty, with 15 assessed as possibly related.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Appendicitis. An update to product labeling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Hemolytic anemia</td>
<td>03Jan22</td>
<td>Closed</td>
<td>26Jan22</td>
<td>Enquiry from a competent authority (Saudi Arabia SFDA).</td>
<td>SFDA requested a comprehensive evaluation report of potential risks of hemolytic anemia with the use of Pfizer/BioNTech COVID-19 vaccine.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Hemolytic anemia. An update to product labeling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Signal term</td>
<td>Date detected</td>
<td>Status (new, ongoing or closed)</td>
<td>Data closed (for closed signals)</td>
<td>Source or trigger of signal</td>
<td>Reason summary</td>
<td>Method of signal evaluation</td>
<td>Outcome, if closed</td>
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<tr>
<td>Uveitis</td>
<td>31Mar22</td>
<td>Closed</td>
<td>27Apr22</td>
<td>Enquiry from a competent authority (Health Canada).</td>
<td>Request from Health Canada due to a WHO publication regarding disproportionate reporting of cases of anterior uveitis following administration of COVID-19 vaccines; request was for a cumulative review of all cases of anterior uveitis including analyses with the preferred terms uveitis, iritis, and iridocyclitis.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Uveitis. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders</td>
<td>11Jan22</td>
<td>Closed</td>
<td>02Feb22</td>
<td>Enquiry from a competent authority (EMA PRAC).</td>
<td>EMA PRAC request to review autoimmune/inflammatory disorder exacerbations reported following vaccination with COMIRNATY including data that have become available since the data-lock period of the 10th SMSR.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Capillary Leak Syndrome (CLS)</td>
<td>13Jan22</td>
<td>Closed</td>
<td>02Feb22</td>
<td>Enquiry from a competent authority (EMA PRAC).</td>
<td>EMA PRAC request based on spontaneous case reports of CLS in individuals vaccinated with COVID-19 mRNA Vaccine (nucleosidemodified) Spikevax, including in patients with medical history of CLS and a latebreaking publication from the EuréClark StudyGroup. The PRAC has agreed that the MAHs for COVID-19 mRNA vaccines, Comirnaty (BioNTech Manufacturing GmbH) and Spikevax (Moderna Biotech Spain, S.L.) should comment on the evidence provided in the publication.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and CLS. An update to product labeling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Signal term</td>
<td>Date detected</td>
<td>Status (new, ongoing or closed)</td>
<td>Data closed (for closed signals)</td>
<td>Reason summary</td>
<td>Method of signal evaluation</td>
<td>Outcome, if closed</td>
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<tr>
<td>Corneal Graft Rejection</td>
<td>01Apr22</td>
<td>Closed</td>
<td>17Jun22</td>
<td>Enquiry from a competent authority (EMA PRAC).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Corneal Graft Rejection. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
<td></td>
</tr>
<tr>
<td>Vasculitis</td>
<td>22Nov21</td>
<td>Closed</td>
<td>02Feb22</td>
<td>Notification from a competent authority (Netherlands Loreb).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Vasculitis. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
<td></td>
</tr>
<tr>
<td>Cerebral venous sinus thrombosis (CVST)</td>
<td>30Nov21</td>
<td>Closed</td>
<td>12Jan22</td>
<td>Enquiry from a competent authority (Switzerland Swissmedic).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and CVST. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
<td></td>
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<tr>
<td>Signal term</td>
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<tr>
<td>Lymphocytic colitis</td>
<td>23Dec21</td>
<td>Closed</td>
<td>10Feb22</td>
<td>Scientific Literature.</td>
<td>Signal identified from new information identified in the published literature, titled &quot;Lymphocytic colitis following mRNA vaccination for SARS-CoV2&quot;, published in the American Journal of Gastroenterology. Conference: Annual Scientific Meeting of the American College of Gastroenterology, ACG 2021. Las Vegas, NV United States. 116(SUPPL) (pp S847), 2021. (Date of Publication: October 2021. Authors Chey S.W.; Westerhoff M.; Chey W.D.).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Lymphocytic colitis. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Chronic Urticaria</td>
<td>24May22</td>
<td>Closed</td>
<td>01Jun22</td>
<td>Enquiry from a competent authority (EMA PRAC)</td>
<td>EMA PRAC preliminary Assessment Report for PSUR #2 contained a request to analyze new onset chronic urticaria following vaccination and respond as a Request for Supplemental Information. Assessment of relapse of chronic urticaria had been included in the PSUR (#3).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Chronic Urticaria, new onset or flares. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Polymyalgia Rheumatica (PMR)</td>
<td>24Jan22</td>
<td>Closed</td>
<td>10Feb22</td>
<td>Enquiry from a competent authority (EMA PRAC).</td>
<td>EMA PRAC Assessment Report to PSUR #1 requested MAH to perform a cumulative review on the association between Comirnaty and Polymyalgia Rheumatica exacerbation.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and PMR. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Signal term</td>
<td>Date detected</td>
<td>Status (new, ongoing or closed)</td>
<td>Data closed (for closed signals)</td>
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<tr>
<td>Subacute Thyroiditis (SAT)</td>
<td>18Jan22</td>
<td>Closed</td>
<td>16Feb22</td>
<td>Enquiry from a competent authority (EMA PRA®).</td>
<td>EMA PRAC Assessment Report to PSUR #1 requested MAH to perform a cumulative review on the association between Comirnaty and subacute thyroiditis.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and SAT. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Cerebrovascular Accident (CVA)/Stroke</td>
<td>27Jan22</td>
<td>Closed</td>
<td>02Mar22</td>
<td>Enquiry from a competent authority (Australia TGA).</td>
<td>Australia TGA’s Medicines and Vaccines Investigation and Surveillance (MaVIS) Section is reviewing the potential safety risk of cerebrovascular accident (CVA)/stroke with COVID-19 vaccines.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Cerebrovascular Accident/Stroke. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
<tr>
<td>Amenorrhea</td>
<td>14Feb22</td>
<td>Closed</td>
<td>16Mar22</td>
<td>Enquiry from a competent authority (EMA PRA®).</td>
<td>Having considered the available evidence from national reviews (post marketing and published studies), the EMA PRAC has requested that the MAH for COVID-19 mRNA Vaccine Comirnaty should perform a cumulative review of all cases of amenorrhea from all sources.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Amenorrhea. An update to product labelling is not warranted at this time. Routine monitoring will continue. Of note: on 13 June 2022, PRAC recommended that no update to the PI was required as the current evidence was insufficient to warrant an update at this time. However, PRAC requested that an updated analysis of amenorrhoea be included in the next PSUR with DLP 18Dec2022.</td>
</tr>
<tr>
<td>Signal term</td>
<td>Date detected</td>
<td>Status (new, ongoing or closed)</td>
<td>Data closed (for closed signals)</td>
<td>Source or trigger of signal</td>
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<tr>
<td>Heavy Menstrual Bleeding</td>
<td>14Feb22</td>
<td>Closed</td>
<td>16Mar22</td>
<td>Enquiry from a competent authority (EMA PRA©).</td>
<td>Having considered the available evidence from national reviews (post marketing and published studies), the PRAC has requested that the MAH for COVID-19 mRNA Vaccine Comirnaty should perform a cumulative review of all cases of all heavy menstrual bleeding from all sources.</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Heavy Menstrual Bleeding. An update to product labelling is not needed at this time. Routine monitoring will continue. Of note: on 13 June 2022, PRAC recommended that no update to the PI was required as the current evidence was insufficient to warrant an update at this time, however the PRAC also provided a List of Questions and the MAH is in the process of preparing a response by 24 Aug 2022.</td>
</tr>
<tr>
<td>Loss of/Altered Taste and Smell</td>
<td>17Mar22</td>
<td>Closed</td>
<td>13Apr22</td>
<td>Enquiry from a competent authority (Australia TGA).</td>
<td>Australia TGA requested an analysis for Comirnaty and loss of/ altered taste and smell (including anosmia, ageusia, dysosmia, dysgeusia, parosmia, phantogeusia).</td>
<td>Postauthorization safety data, clinical study safety data, medical literature, O/E analyses.</td>
<td>Based on the totality of data, there is insufficient evidence to conclude a causal association between the vaccine and Loss of/Altered Taste and Smell. An update to product labelling is not warranted at this time. Routine monitoring will continue.</td>
</tr>
</tbody>
</table>
Other safety topics not considered signals

Dizziness

PRAC request 3 from the 2nd Comirnaty PSUSA (procedure EMEA/H/C/PSUSA/00010898/202112):

The MAH is requested to provide a cumulative review of cases reporting dizziness after Comirnaty exposure outside the context of anxiety/stress-related reactions (as already labelled in the Comirnaty SmPC section 4.4) and a discussion on the need to add dizziness (including a proposal for the frequency of occurrence) to the ADR table of the Comirnaty SmPC section 4.8, as applicable.

MAH’s response (Appendix 6A.1 of the PSUR):

The Pfizer safety database was searched cumulatively through 18 June 2022 for all BNT162b2 cases of PT Dizziness (MedDRA v. 25.0).

All cases of dizziness

A total of 96,959 cases (with a total of 598,847 events) were retrieved from the global safety database using the search strategy above. The majority of cases (92,849, 96%) were spontaneous; there were 72,597 females, 22,435 males, and sex was not reported in 1927 cases. Most cases were nonserious (64,467; 67%). The most frequently reported age group was 31 to 50 years (40.6%).

When provided, the most frequently reported time to onset (latency) for dizziness in the 96,959 cases was Day 0 (for 32,355 relevant events) and Day 1 (24,941 events) followed by Day 2 (for 6139 events) and Day 3 (for 3201 events).

The high number of dizziness events occurring on the day of vaccination are suggestive of a large proportion of the events of dizziness occurring as a stress-related response to the vaccination process (as currently labelled). In addition, there are a large number of reports with a latency of 1 day after vaccination; this latency data and the co-reported events suggest these reports of dizziness occurred simultaneously or, potentially, as a consequence of, reactogenicity-type events.

Medically-confirmed and serious cases of dizziness

Further analysis was concentrated on the 5563 medically confirmed healthcare professional (HCP) cases (with a total of 41,777 events) in which dizziness was reported as a serious event, and the time to onset (latency) from vaccination was Day 0 (same day) to Day 21 post vaccination.

The majority of this subset of cases (5328, 96%) were spontaneous; there were 4183 females, 1316 males, and sex was not reported in 64 cases. Most of the cases were reported in adults 18-64 years of age. The mean and median ages were 44.7 and 43 years, respectively (n=5426).

Dizziness with Latency Day 0-Day 1

Out of the 5563 cases, 4113 (74 %) reported 4145 serious events of Dizziness (some cases reported the same event more than once) up to 1 day after administration of the vaccine. There were 3223 females, 839 males, and sex was not reported in 51 cases. The mean and median ages were 43.4 and 42 years, respectively (n=4003).

When dose sequence was provided, dizziness followed Dose 1 for 2015 relevant events, Dose 2 for 1207 events, Dose 3 for 223 events, and Dose 4 for 1 event.
The outcomes of dizziness at the time of reporting was resolved/resolving for 2527 events; not
resolved for 702 events; resolved with sequelae for 56 events; fatal for 21 events and unknown for
839 events.

Most cases (4063/4113, 99%) had co-reported events; the most frequently co-reported PTs are
presented below in Table 5. Similar to the overall dataset of 96,959 cases, these PTs are reactogenicity
events and events that are reflective of symptoms of a stress-related response (e.g., to the
vaccination process itself).

<table>
<thead>
<tr>
<th>PT</th>
<th>Number of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Headache</td>
<td>1472</td>
</tr>
<tr>
<td>Nausea</td>
<td>1227</td>
</tr>
<tr>
<td>Fatigue</td>
<td>784</td>
</tr>
<tr>
<td>Dyspnoea</td>
<td>764</td>
</tr>
<tr>
<td>Pyrexia</td>
<td>669</td>
</tr>
<tr>
<td>Malaise</td>
<td>617</td>
</tr>
<tr>
<td>Asthenia</td>
<td>587</td>
</tr>
<tr>
<td>Vomiting</td>
<td>531</td>
</tr>
<tr>
<td>Myalgia</td>
<td>481</td>
</tr>
<tr>
<td>Chills</td>
<td>459</td>
</tr>
<tr>
<td>Tachycardia</td>
<td>427</td>
</tr>
<tr>
<td>Anaphylactic reaction</td>
<td>408</td>
</tr>
</tbody>
</table>

Fifty of the 4113 cases reported dizziness as the only event. Out of these 50 cases:

- 18 described pre-existing medical conditions such as previous post-vaccination dizziness,
  cardiac disorders, vascular disorders, anxiety, diabetes, COVID-19 and migraine and/or use of
  concomitant medication such as antidepressants, antihypertensives, glucose-lowering agents,
  and prednisone which may cause or contribute to dizziness.

- 32 provided limited information about medical conditions or medications.

Twenty-one of the 4113 cases described a fatal outcome (15 with a latency of 1 day postvaccination
and 6 on the day of vaccination). The age of the patients ranged from 27 to 94, with 11 being 65 years
or older; 8 of whom had pre-existing cardiovascular disease or cancer and were on several
concomitant medications.

**Rapporteur assessment comment:**

Out of the retrieved 96,959 cases reporting dizziness through 18 Jun 2022, the MAH focused the
analysis on 4113 medically confirmed cases (4.2%) with serious dizziness and a TTO of 0-1 day after
Comirnaty exposure. This is problematic because all non-serious cases reporting dizziness (which are
considered useful for a non-serious AE like dizziness) are not included in the analysis. The MAH was
requested to perform a cumulative review of cases reporting dizziness after Comirnaty exposure
outside the context of anxiety/stress-related reactions, which seems not to be performed.

**Dizziness with Latency Day 2-Day 21**

Out of the 5563 cases, 1450 (26 %) reported 1471 serious events of dizziness (some cases reported
the same event more than once) with latency of Day 2 through Day 21 postvaccination. There were
960 females, 477 males, and sex was unknown in 13 cases. The mean and median ages were 48.2 and
48 years, respectively (n=1423).

When dose sequence was provided, it was Dose 1 for 619 relevant events, Dose 2 for 510 events,
Dose 3 for 133 events and Dose 4 for 2 events.
The outcome of Dizziness at the time of reporting was resolved/resolving for 679 events; not resolved for 454 events; resolved with sequelae for 37 events; fatal for 22 events; unknown for 279 events. The most frequently reported latency was Day 2 (376 events) followed by Day 3 (227), Day 4 (165) and Day 5 (106).

The most frequently co-reported PTs in these 1450 cases are presented below in Table 7.

<table>
<thead>
<tr>
<th>PT</th>
<th>Number of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Headache</td>
<td>548</td>
</tr>
<tr>
<td>Nausea</td>
<td>385</td>
</tr>
<tr>
<td>Fatigue</td>
<td>351</td>
</tr>
<tr>
<td>Pyrexan</td>
<td>221</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>216</td>
</tr>
<tr>
<td>Vertigo</td>
<td>216</td>
</tr>
<tr>
<td>Anemia</td>
<td>199</td>
</tr>
<tr>
<td>Chest pain</td>
<td>197</td>
</tr>
<tr>
<td>Malaise</td>
<td>188</td>
</tr>
</tbody>
</table>

Out of these 1450 cases, the majority (1423, 98%) co-reported additional events that were largely consistent with expected reactogenicity events.

The remaining minority of cases (27, 2%) out of 1450 cases reported dizziness as the only event. In these cases:

- 8 of 27 reported pre-existing medical histories (e.g. diabetes, infections, arthropathy, cardiac failure) and/or concomitant medications (e.g. zopiclone, alpha blockers, antihypertensives, immunosuppressants) that may cause or contribute to dizziness, and

- 21 cases provided limited details about medical conditions or concomitant medications.

There were 22 cases with fatal outcome. The most commonly co-reported PTs in these 22 cases were headache, fatigue, and syncope (5 each). The most commonly reported latency was Day 2 (for 5 Dizziness events) followed by Day 3 (for 4 events). The age of the patients ranged from 50 to 95 years with 14 cases in patients 65 years and older. More than half of these cases reported patients with pre-existing cardiovascular medical conditions on several concomitant medications.

**Rapporteur assessment comment:**

Out of the retrieved 96,959 cases reporting dizziness, there were 1450 medically confirmed cases (1.5%) with serious dizziness and a TTO of 2-21 days after Comirnaty exposure. Of these 1450 cases there were 1423 cases (98%) that reported co-events which are considered a symptom of a stress-related response to the vaccination process.

Clinical trial data

**C4591001**

In the placebo-controlled period of pivotal study C4591001 (DLP 13 MAR 2021), there were 166 participants who reported Dizziness; 74/23,037 (0.3%) were in the placebo group and 92/23,040 (0.4%) were in the BNT162b2 group. In the placebo group, 72 out of 74 events were non-serious; all events were non-serious in the BNT162b2 group. The most frequently reported latency in the placebo group were Day 0 (for 11 events) followed by Day 1 (for 8 events). Similar to this, the most frequently reported latency in BNT162b2 group were Day 0 (for 42 events) followed by Day 1 (for 35 events).

**C4591031**

In the placebo-controlled period of booster study C4591031 sub study A (DLP 08 FEB 2022), there were 25 participants who reported Dizziness; 6/5048 (0.1%) were in the placebo group and 19/5088...
(0.4%) were in the BNT162b2 group. In both groups the events of Dizziness were nonserious. In the placebo group the latency was reported as Day 1, Day 14, and Day 31 (for 2 events each). The most frequently reported latency in BNT162b2 group were Day 1 (for 12 events) followed by Day 0 (for 5 events).

C4591007

In the placebo-controlled period of pivotal study C4591007 for the 5 to >12 age group (DLP 06 SEP 2021), there were 2 participants who reported Dizziness (1/750 in the placebo group and 1/1518 in the BNT162b2 group). The reported latency was Day 19 for the event in the placebo group and the same day (Day 0) in BNT162b2 group.

Rapporteur assessment comment:
Pooled In the three placebo-controlled studies there were 112 of the 29,646 (0.38%) participants reporting dizziness in the Comirnaty group and 81 of the 28,835 (0.28%) participants in the placebo group.

MAH’s conclusion

In the large Pfizer-run clinical trials, dizziness was not commonly reported. It occurred more frequently in the vaccine groups than placebo groups, however the frequency was < 1% in both groups. Similar to the post-authorization data, the latency tended to be on the day of vaccination or the following day.

Approximately 6% of spontaneously reported BNT162b2 vaccine AE reports are cases reporting dizziness; it is the 11th most reported event for BNT162b2 in the global safety database. The characteristics of the overall number of dizziness cases in this review (96,959) are that they are mostly non-serious, occur on the day of vaccination or the following day, and are co-reported with events that are known to be systemic reactogenicity symptoms. Consistent with the overall BNT162b2 post-authorization database of AE reports, dizziness is reported more frequently in women than men and in adults up to 64 years of age more than any other age population.

Concentration on the most medically significant cases (serious and medically confirmed) shows a similar pattern, with most events occurring within a few days of vaccination and co-reported with events that are recognized reactogenicity events and stress-related responses to the vaccination process.

Based on this review of data, the MAH has determined that dizziness should be considered an adverse reaction to BNT162b2 and will update section 4.8 of the company core data sheet accordingly. Subsequent changes to local labels, including the SmPC will occur per Pfizer process.

Rapporteur assessment comment:
Although the non-serious cases reporting dizziness were excluded from the presented analysis, the MAH states that these cases showed a similar pattern as the serious and medically confirmed cases, occurrence on the day of vaccination or the following day, and are often co-reported with events that are known to be systemic reactogenicity symptoms. Dizziness is reported more frequently in women than men and in adults up to 64 years of age more than any other age population.

Of note, dizziness is stated as an ADR in the product information of Spikevax, Vaxzevria, Jcovden, and Valneva.

Overall MAH’s conclusion, that dizziness should be considered an adverse drug reaction to Comirnaty and should be added to the Comirnaty SmPC section 4.8., is endorsed. The PRAC Rapporteur considers a causal relationship between Comirnaty and dizziness is at least a reasonable possibility.
Therefore, in section 4.8 of the Comirnaty SmPC the adverse drug reaction, dizziness, should be added to the ADR table under the SOC Nervous system disorders with a frequency Unknown. The package leaflet should be updated accordingly. The MAH has already submitted a variation (procedure EMEA/H/C/005735/II/0152) to update the PI in this respect.

Acquired haemophilia

PRAC request 4 from the 2nd Comirnaty PSUSA (procedure EMEA/H/C/PSUSA/00010898/202112):

The MAH is requested to provide a cumulative review of cases reporting acquired haemophilia, including details of the underlying condition(s), time to onset, duration, outcome and an assessment of the causal relationship with the vaccination. The MAH should also provide a comprehensive discussion of the literature, mechanism of action and a discussion on the need to update the product information of Comirnaty, if applicable.

MAH’s response (Appendix 6A.2 of the PSUR):

The safety database was searched for all BNT162b2 AE reports with the Preferred Terms under the Higher-Level Term of Coagulation factor deficiencies received cumulatively to 18 June 2022.

A total of 68 cases were retrieved, 51 were spontaneous reports, 4 were from non-interventional studies and 13 were from literature reports. All the 68 cases were assessed as serious.

Gender was reported in 64 cases (32 males and 32 females), while 4 cases did not specify the gender. Sixty-three (63) cases reported an age, which ranged from 25 years to 97 years (mean: 74.59, median: 77). Consistent with known information on acquired hemophilia, the large majority of cases are in patients ≥65 years of age (79.3%).

France reported 23 cases, Italy reported 14 cases while Japan reported 5 cases. The top 10 countries reporting cases of acquired hemophilia are shown below (Table 2).

<table>
<thead>
<tr>
<th>Country</th>
<th>Number</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>France</td>
<td>23</td>
<td>34%</td>
</tr>
<tr>
<td>Italy</td>
<td>14</td>
<td>21%</td>
</tr>
<tr>
<td>Japan</td>
<td>5</td>
<td>7.4%</td>
</tr>
<tr>
<td>Switzerland</td>
<td>4</td>
<td>5.9%</td>
</tr>
<tr>
<td>United Kingdom</td>
<td>4</td>
<td>5.9%</td>
</tr>
<tr>
<td>Australia</td>
<td>3</td>
<td>4.4%</td>
</tr>
<tr>
<td>Finland</td>
<td>3</td>
<td>4.4%</td>
</tr>
<tr>
<td>Ireland</td>
<td>2</td>
<td>2.9%</td>
</tr>
<tr>
<td>Malaysia</td>
<td>2</td>
<td>2.9%</td>
</tr>
<tr>
<td>United states</td>
<td>2</td>
<td>2.9%</td>
</tr>
</tbody>
</table>

All 68 cases reported an event of Acquired Hemophilia. Other adverse event PTs co-reported in the cases included anemia (13), hematoma (11) spontaneous hematoma (6), hemorrhage (5), contusion (4), mouth hemorrhage (3), hematuria (3), ecchymosis (3), hemoglobin decreased (3), factor VIII deficiency (3), hemorrhagic shock (2).

Sixty-four (64) of the cases were medically confirmed while the remaining 4 were non medically confirmed.

Seventeen (17) of the 68 cases occurred after the first dose, 33 cases reported onset after the second dose, 8 cases reported onset after the third dose and the dose was not reported in the remaining 10 cases.
Case outcome was reported as follows: recovering/recovered (29), not recovered (25), fatal (6) and unknown (8).

**Rapporteur assessment comment:**
Post-marketing through 18 Jun 2022, a total of 68 cases reporting acquired hemophilia were retrieved. France and Italy reported most cases, 23 (34%) and 14 (21%) respectively.

Fatal case reports

1. A 90-year-old male received the first dose of BNT162b2 on 23 March 2021. Medical history included diabetes, stroke, obstructive arteriosclerosis of lower extremities, chronic renal failure and COVID-19. The patient’s concomitant medications were not reported. Information on subsequent doses were not reported. He experienced acquired hemophilia 88 days after the first dose. Lab findings included aPTT. He was hospitalized on an unspecified date in July 2021 for hemorrhagic syndrome. Therapy included prednisone, multiple transfusions for digestive hemorrhage and discontinuation of antiaggregant and anti-vitamin K recently Introduced for atrial fibrillation. The patient died on an unspecified date in 2021. Cause of death and autopsy were not reported.

**MAH comment:** Diabetes mellitus and chronic renal failure in this patient could be confounders for acquired hemophilia. The patient’s age and history of Covid-19 could also be contributory. The latency of 88 days following dose 1 and no documentation of a 2nd dose call into question the latency feasibility. Concomitant medications were not reported.

2. A 90-year-old female patient received the first dose of BNT162b2 on 25 February 2021. Her medical history included hypertension, aortic stenosis, heart failure, gout, and chronic renal failure. Concomitant medications included zopiclone, bisoprolol, furosemide, allopurinol, and paracetamol. After 14 days the patient experienced atrial fibrillation, decreased hemoglobin, hematoma, oral mucosa bleeding, malaise, an increased tendency to bruise, acquired hemophilia and aggravated cardiac failure. Patient was hospitalized and lab showed positive Anti-factor VIII antibody. Treatment with prednisolone was initiated. On 17 April patient experienced worsening heart failure, atrial fibrillation, and kidney problem. The patient died on 24 May 2021.

**MAH comment:** This patient’s elderly age and paracetamol use are known risk factors for Acquired Hemophilia. Multiple comorbidities in this patient could also be contributory.

**Rapporteur assessment comment:**
MAH’s comment is endorsed, other causes appear more likely.

3. An 84-year-old female received a second dose of BNT162b2 on 08 April 2021. The patient’s medical history and concomitant medications were not reported. Two days post vaccination, the patient experienced acquired hemophilia A, with spontaneous, superficial, and deep hematomas and significant anemia reported. Relevant lab tests included partial thromboplastin time of 65.3 seconds, Factor VIII at 1%, and positive lupus like anticoagulants. Treatment was with steroids, cyclophosphamide, vitamin k and blood transfusions. The patient died on 07 June 2021, and the cause of death was reported as acquired hemophilia, anemia, and spontaneous hematoma.

**MAH comment:** There is limited information on this patient’s medical history and concomitant medication that may be contributing factors to the development of AH in addition to her elderly age.
Rapporteur assessment comment:

MAH’s comment is endorsed, limited information for causality assessment.

4. A 67-year-old male received the second dose of BNT162b2 on 22 April 2021. Relevant medical history included rheumatoid arthritis, Crohn’s disease and pulmonary legionellosis. Concomitant medications included Duragesic, prednisone, Spiriva, indacaterol and pentasa. On 18 May 2021, patient experienced a fall at home, with diffuse hematomas. Three days later he was brought to the ER due to a deteriorating general condition and vomiting and was assessed to have hemorrhagic shock associated with acute renal failure and acquired hemophilia type A. One day after hospitalization the patient experienced sudden deterioration with onset of coma, and the following day he died from cardio-respiratory arrest.

MAH comment: Crohn’s disease and rheumatoid arthritis are confounders for acquired hemophilia. In addition, this patient’s traumatic fall likely contributed to his deteriorating condition.

Rapporteur assessment comment:

MAH’s comment is endorsed, other causes more likely.

5. A 77-year-old man received his second dose of BNT162b2 on 28 June 2021. He reported a medical history of relapsed bladder carcinoma. He was hospitalized for hematuria, 21 days after vaccination. Hemoglobin concentration was 66 g/L, activated partial thromboplastin time was increased (3.61 ratio) and FVIII:C (Factor VIII procoagulant activity) was 0.02 IU/mL with detectable inhibitor (6.9 Bethesda Units/mL). Treatment was with methylprednisolone, recombinant activated clotting Factor VII and rituximab. However, during the hospital stay the patient developed sepsis and died from respiratory complications seven weeks after admission.

MAH comment: Bladder carcinoma in this patient could be a confounder for acquired hemophilia seen in this patient.

Rapporteur assessment comment:

MAH’s comment is endorsed, other cause more likely.

6. A 97-year-old female patient received the third dose of BNT162b2 on 28 October 2021. Her medical history and concomitant medications were not reported. The patient had received the first dose on 22 February 2021 and the second dose on 15 March 2021. The patient experienced acquired hemophilia, extensive muscle hematoma, and severe anemia 18 days post vaccination. Laboratory tests revealed prolonged aPPT, Anti-factor VII antibody, Factor VIII deficiency, and Anti-factor VIII circulating inhibitor. Therapeutic measures included corticosteroids, rituximab, and concentrated red blood cell transfusion. The patient died on 11 January 2022, with cause of death not reported.

MAH comment: Elderly age is a risk factor in this patient. There is limited information on medical history and concomitant medication which precludes a meaningful medical assessment.

Rapporteur assessment comment:

MAH’s comment is endorsed, limited information for causality assessment.
Overall in the reported total of 6 fatal cases, there were 4 cases with other causes that more likely could have caused acquired hemophilia and 2 cases with too limited information to perform a meaningful causality assessment.

The remaining 62 cases are summarized below:

- 40 cases reported one or more relevant medical history or concomitant medications that could confound the event of acquired hemophilia in these patients,
  - 13 cases reported diabetes mellitus,
  - 11 cases reported cancer/neoplasm: breast cancer, prostate cancer, unspecified cancer, renal neoplasm, diffuse large b-cell lymphoma, laryngeal carcinoma, lung adenocarcinoma,
  - 6 cases reported autoimmune diseases: Giant cell arteritis, rheumatic fever, rheumatoid arthritis, Sjogren’s disease, granulomatosis with polyangitis, sarcoidosis,
  - 3 cases reported hypothyroidism,
  - 3 cases reported asthma or broncho-pneumopathy,
  - 2 cases reported polymyalgia rheumatica,
  - 2 cases reported previous history of Acquired hemophilia,
  - 1 case reported history of myelodysplastic syndrome,
  - 1 case reported angiodysplasia,
  - 1 case reported COVID-19,
  - 14 cases reported use of concomitant medications such as acetaminophen, olanzapine, apixaban, acetylsalicylic acid, levothyroxine, risperidone, chlorpromazine and levomepromazine.

- 18 cases had limited information on medical history, concomitant medication, time to onset, dose number and schedule, and clinical information surrounding the event precluding a meaningful medical assessment.

Rapporteur assessment comment:
The MAH reports that 18 of the 62 remaining cases had limited information. However, details of these 18 cases were not presented and hampers PRAC rapporteur’s assessment of these 18 cases.

Furthermore, there is no information if the cases were tested positive or negative for SARS-CoV-2 infection.

The remaining 4 cases are described below:

1. An 84-year-old male received his second dose of BNT162B2 on 06 May 2021. Medical history included transient ischemic attack, lumbago, arterial hypertension. The patient’s concomitant medications were not reported. Twenty-five days after the second dose, the patient experienced acquired hemophilia with multisite hemorrhagic anemia: retroperitoneal and psoas muscle hematoma. Therapy was started with eptacog-alfa, prednisone, cyclophosphamide and suspended on 30 June 2021 due to the onset of severe isolated thrombocytopenia. Outcome was not reported.
MAH comment: This patient’s age is a potential risk factor for AH. There is insufficient information on concomitant medication, confirmatory lab test and details surrounding the event.

Rapporteur assessment comment:
MAH’s comment is endorsed, limited information for causality assessment.

2. A 67-year-old man received the second dose of BNT162b2 on 16 Jun 2021. His medical history was unremarkable. He was admitted to the Emergency Room for urgent otolaryngological assessment due to a large hematoma of the tongue, extending in the cervical region. Hemoglobin concentration was 125 g/L, the aPTT ratio was 2.55, FVIII:C (factor VIII procoagulant activity) was 0.06 IU/mL with detectable anti-FVIII activity (2.5 Bethesda Units/mL). Recombinant activated clotting Factor VII, cyclophosphamide and prednisone was administered. Outcome of the event was reported as recovered.

MAH comment: Although this patient did not have a prior relevant history, there is insufficient information on the time to onset, concomitant medication and clinical details preceding the event.

Rapporteur assessment comment:
MAH’s comment is endorsed, limited information for causality assessment.

3. An 86-year-old female patient received the first dose of BNT162b2 on 22 June 2021. The patient’s medical history included coronary artery bypass. The patient reportedly took aspirin. She experienced acquired hemophilia 11 days after the first dose, which was described as multiple subcutaneous hemorrhages and bleeding tendency. The aspirin was stopped, but it is unclear if the bleeding subsided, with bleeding stoppage described as “not good”. On 13 July 2021, the patient received the second dose, and subcutaneous hemorrhage and muscle hemorrhage appeared. Fifteen days after the second dose, the patient was hospitalized and laboratory tests revealed prolonged aPTT with an inhibitor pattern in the cross-mixing test, low factor VIII activity and a diagnosis of acquired hemophilia was made. Outcome of the event was reported as not recovered.

MAH comment: Elderly age in this patient is a risk factor. Aspirin use could also be contributory to the event of Acquired hemophilia.

Rapporteur assessment comment:
Although the case reported a rechallenge of complaints after the 2nd Comirnaty dose, it is not clearly stated if there was Aspirin use. Therefore MAH’s comment is endorsed, other causes could be possible.

4. A 25-year-old female patient received second dose of BNT162b2 on 18 August 2021. The patient’s first dose was on 17 July 2021. Medical history included obesity treated with bariatric surgery in December 2020, cholecystectomy, appendectomy and active smoking. Familial history included phlebitis in a cousin, no autoimmune disease, no first-degree venous thromboembolism. The patient previously took lovasuvlo (oral contraceptive) and vitamin B12 and experienced digestive intolerance. On 28 August 2021, 10 days after vaccination the patient visited the emergency room for pain in the right leg, edema and multiples bruises, the patient has reportedly had the symptoms for one month. Lab test revealed anti-FVIII antibodies and assessment of acquired hemophilia A. Outcome was not reported.

MAH comment: Acquired hemophilia is more common in young females especially during pregnancy or post-partum period, however there is limited information on this patient’s obstetric history. In addition, time of onset of acquired hemophilia is unclear.
Rapporteur assessment comment:

MAH's comment is endorsed, limited information for meaningful causality assessment.

Rapporteur assessment comment:

Overall in the remaining 62 cases:
- there were 40 cases reported one or more relevant medical history or concomitant medications that could confound the event of acquired hemophilia;
- there were 18 cases had limited information event precluding a meaningful medical assessment, however lack of details hampered the PRAC rapporteur’s assessment;
- there were 4 cases with limited information for meaningful causality assessment.

No information was provided if the cases were tested positive or negative for SARS-CoV-2 infection. However, the MAH is requested to inform the PRAC rapporteur immediately if there are more supportive acquired hemophilia cases after Comirnaty exposure.

Clinical trials

There were no cases of acquired hemophilia reported in the pivotal Pfizer-led clinical trials.

Literature

There were no relevant literature articles other than literature case reports which are reviewed in the post marketing section.

Observed versus expected analysis

The MAH conducted unadjusted observed to expected ratio (O/E) analyses for the 68 acquired hemophilia cases reported cumulatively through 18 June 2022 globally. The overall expected case counts were estimated using background incidence rates (IR) reported by a surveillance study of patients treated for acquired hemophilia in hematology departments covering all National Health Service hospitals in the UK during 1 May 2002 – 30 April 30 2003. This study was selected because in a literature review, the MAH found that this figure of 1.5 cases per million is widely cited for acquired hemophilia incidence and is in the low end of the range of 1 to 4 cases per million reported by a review article.
Table 3. Observed to Expected (O/E) Ration for Spontaneously Reported Cases of Acquired Hemophilia Through 18 June 2022

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Observed cases</th>
<th>Time at risk (PY)</th>
<th>Background rates per 100,000 PY</th>
<th>Expected cases</th>
<th>O/E ratio</th>
<th>95% CI LL</th>
<th>95% CI UL</th>
</tr>
</thead>
<tbody>
<tr>
<td>21-day risk window</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td>0</td>
<td>1,917,834</td>
<td>0.005</td>
<td>0.09</td>
<td>0.000</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>≤11 years</td>
<td>0</td>
<td>3,324,066</td>
<td>0.017</td>
<td>0.57</td>
<td>0.008</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>12-17 years</td>
<td>0</td>
<td>4,733,892</td>
<td>0.029</td>
<td>1.37</td>
<td>0.008</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>18-24 years</td>
<td>2</td>
<td>19,546,111</td>
<td>0.029</td>
<td>5.67</td>
<td>0.353</td>
<td>0.043</td>
<td>1.275</td>
</tr>
<tr>
<td>25-49 years</td>
<td>0</td>
<td>9,025,268</td>
<td>0.029</td>
<td>2.62</td>
<td>0.000</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>50-59 years</td>
<td>2</td>
<td>7,901,305</td>
<td>0.310</td>
<td>24.49</td>
<td>0.082</td>
<td>0.010</td>
<td>0.295</td>
</tr>
<tr>
<td>60-69 years</td>
<td>18</td>
<td>10,962,290</td>
<td>1.032</td>
<td>113.08</td>
<td>0.159</td>
<td>0.004</td>
<td>0.252</td>
</tr>
<tr>
<td>70+ years</td>
<td>34</td>
<td>127,075,389</td>
<td>0.150</td>
<td>190.61</td>
<td>0.178</td>
<td>0.124</td>
<td>0.249</td>
</tr>
<tr>
<td>Overall Global</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td>0</td>
<td>2,779,982</td>
<td>0.005</td>
<td>0.13</td>
<td>0.000</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>≤11 years</td>
<td>0</td>
<td>5,127,136</td>
<td>0.017</td>
<td>0.87</td>
<td>0.000</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>12-17 years</td>
<td>0</td>
<td>7,477,185</td>
<td>0.029</td>
<td>2.17</td>
<td>0.080</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>18-24 years</td>
<td>2</td>
<td>30,991,113</td>
<td>0.029</td>
<td>8.99</td>
<td>0.223</td>
<td>0.027</td>
<td>0.804</td>
</tr>
<tr>
<td>25-49 years</td>
<td>1</td>
<td>14,461,569</td>
<td>0.029</td>
<td>4.19</td>
<td>0.238</td>
<td>0.006</td>
<td>1.329</td>
</tr>
<tr>
<td>50-59 years</td>
<td>7</td>
<td>12,856,835</td>
<td>0.310</td>
<td>39.86</td>
<td>0.176</td>
<td>0.071</td>
<td>0.362</td>
</tr>
<tr>
<td>60-69 years</td>
<td>24</td>
<td>17,896,776</td>
<td>1.032</td>
<td>184.61</td>
<td>0.130</td>
<td>0.083</td>
<td>0.193</td>
</tr>
<tr>
<td>70+ years</td>
<td>50</td>
<td>203,986,419</td>
<td>0.150</td>
<td>305.98</td>
<td>0.163</td>
<td>0.121</td>
<td>0.215</td>
</tr>
</tbody>
</table>

CI = confidence interval; EEA = European Economic Area; LL = lower limit; PY = person-years; UL = upper limit; US = United States.

Based on the selected background rates and the estimated number of exposure person-years through 18 June 2022, O/E ratios were well below one overall and by age for both risk windows of 21- and 42-days. This suggests that the number of observed cases is not higher than expected in the absence of Pfizer-BioNTech COVID-19 vaccines overall and within the queried strata.

Rapporteur assessment comment:
All O/E ratios are well below 1.

MAH's conclusion

Consistent with the rarity of the disorder, the search of the clinical trial database did not reveal any cases of acquired hemophilia. The medical literature search results were of case reports (included in the safety database analysis), but no larger, population level studies were retrieved.

The search of the safety database identified 68 cases, all reported as serious. Of the 68 reports, 40 cases included details in the medical history regarding medical conditions and diseases under treatment that could predispose to developing acquired hemophilia (cancer, rheumatoid arthritis, chronic kidney failure, nephroangiosclerosis, diabetes mellitus and hypothyroidism). Six cases reported fatal outcomes (4 of them were confounded by medical history and 2 had age-related risk factors and limited information on medical history). A further 18 cases had insufficient information on medical history and concomitant medications to conduct a thorough assessment. In the remaining 4 cases, although some clinical details were provided, 3 cases had age-related risk factors and history of aspirin use, while the remaining case, in a 67-year-old man, lacked clear information on concomitant
medication and time to onset to make a conclusive medical assessment. Fifty-four (54) (79%) of the 68 cases occurred in elderly patients greater than 65 years, who are the most predisposed to developing acquired hemophilia, with many of these patients having associated comorbidities.

The ratios of observed to expected cases were less than one overall and by age for both the 21- and 42-days risk windows.

Overall, the totality of data does not support a causal association between vaccination and acquired hemophilia and there is no need to update the product information at this time. Routine monitoring will continue.

**Rapporteur assessment comment:**

**Post-marketing**

Out of the retrieved 68 cases reporting acquired hemophilia through 18 Jun 2022:

- there were 6 fatal cases, of which 4 cases reported other causes that more likely could have caused acquired hemophilia and 2 cases with too limited information to perform a meaningful causality assessment.

- there were 40 cases reported one or more relevant medical history or concomitant medications that could confound the event of acquired hemophilia,

- there were 18 cases had limited information event precluding a meaningful medical assessment,

- there were 4 cases with limited information for meaningful causality assessment.

**Clinical trials**

There were no cases of acquired hemophilia in clinical trials. However, clinical trials are not designed to capture rare adverse events.

**Literature**

There were 13 literature cases reports, which were reviewed in the post-marketing section.

**O/E analysis**

Alle O/E ratios were well <1.

Overall MAH's conclusion is endorsed, that the data does not support a causal association between Comirnaty exposure and acquired hemophilia. However, the MAH is requested to inform the PRAC rapporteur immediately if there are more supportive acquired hemophilia cases after Comirnaty exposure.

**MIS-C/-A**

Introduction (Appendix 6A.4 of the PSUR)

In August 2021, the European Medicines Agency (EMA) issued a signal assessment report on Multisystem inflammatory syndrome in children (MIS-C) with SARS-CoV-2 vaccination and requested all Market Authorization Holders (MAH) of these vaccines perform cumulative review of MIS-C and Multisystem inflammatory syndrome in adults (MIS-A).

A cumulative review of cases reported within Pfizer's global safety database was performed with a data lock point (DLP) of 02 September 2021. Analysis of these cases, in conjunction with observed to expected analysis did not support a causal relationship between Comirnaty and MIS-C/A.
In concordance with Pfizer’s assessment the Pharmacovigilance Risk Assessment Committee (PRAC) agreed that the signal be closed and that no update to the product information is currently warranted.

PRAC requested the MAH continue to closely monitor MIS-C/A and report on new cases in the Monthly Summary Safety Report (MSSR) and Periodic safety update report (PSUR). Cases were requested to be assessed using the Brighton Collaboration (BC) case definition1 with MIS-C defined as patients age < 21 years and MIS-A those age ≥ 21 years.

Interval cases have subsequently been analyzed and discussed in the following documents:

- MSSR #11 (Interval 03 September through 26 October 2021),
- Summary bi-monthly safety report (SBSR) #1 (Interval 27 October through 15 December 2021),
- Periodic Safety Update Report #2 (Interval 19 June through 18 December 2021),
- SBSR #2 (Interval 16 December 2021 through 15 February 2022),
- SBSR #3 (Interval 16 February through 15 April 2022).

In accordance with the PRAC request, retrieved cases meeting BC level 1 (definitive), 2 (probable) and 3 (possible) case definition criteria are presented in this review. The following MedDRA version 25.0 Preferred Terms (PTs) were used: Multisystem inflammatory syndrome in children, Multisystem inflammatory syndrome in adults, Multisystem inflammatory syndrome, Multiple organ dysfunction syndrome, Systemic inflammatory response syndrome, Cytokine release syndrome, Distributive shock.

Methodology

The safety database was searched for all BNT162b2 cases reported in the interval 18 December 2021 through 18 June 2022.

Results

One hundred and ninety-nine (199) case reports for this reporting period were retrieved using the above method. Of these cases 49 occurred in patients aged less than 21-years-old and were classified in consideration of MIS-C, 142 occurred in patients aged greater than or equal to 21 years and were classified in consideration of MIS-A. In 8 cases no age was reported and 7 of which provided such limited information as to preclude a meaningful analysis of the case.

**MIS-C**

In 2 of the 49 cases considered for MIS-C limited clinical information was reported precluding an analysis of the case per the BC criteria. The remaining 47 cases were classified as per Table 1.

<table>
<thead>
<tr>
<th>BC level</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>26</td>
</tr>
<tr>
<td>5</td>
<td>3</td>
</tr>
</tbody>
</table>

*Table 1. BC classification of potential MIS-C cases*

---

**Rapporteur assessment comment:**
During the reporting period, the MAH identified a total of 47 potential new MIS-C cases. Of these, 10 were classified as BC level 1, 5 as BC level 2, 3 as BC level 3, 26 as BC level 4, and 3 as BC level 5.

Of the 10 BC level 1 MIC-C cases, 4 cases were previously assessed in the 13th SSR (16 Dec 2021-15 Feb 2022) (AER numbers [redacted]), and 6 cases were previously assessed in the 14th SSR (16 Feb 2022-15 Apr 2022) (AER numbers [redacted]).

The 8 cases that were classified as BC level 2 (probable MIS) and 3 (possible MIS) have not been reproduced in this AR. These were all either considered confounded by previous COVID-19 infection, no information was provided on previous COVID-19 infection or this information was incomplete.

Therefore the previous PRAC conclusion in the 14th SSR (procedure EMEA/H/C/005735/MEA/002.13) remains valid, i.e. that the data is currently insufficient to support regulatory action and therefore no update of the product information in relation to MIS-C is currently warranted. Although one additional MIS-C BC level 1 case (AER number [redacted], literature case from [redacted] that occurred in an 16-year old male) besides the Danish index case that was considered probably related with Comirnaty was identified, this is given the extensive exposure of Comirnaty (in children) not to be considered unexpected and does not present a new safety concern.

**MIS-A**

One hundred and forty-three (142) cases were in patients aged greater than or equal to 21 years and were classified in consideration of MIS-A. Twelve cases reported insufficient information to assess the case. Table 5 demonstrates the BC level classification of the remaining 130 cases.

<table>
<thead>
<tr>
<th>BC level</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>34</td>
</tr>
<tr>
<td>5</td>
<td>94</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment:**

During the reporting period, the MAH identified a total of 130 potential new MIS-A cases. Of these, 1 was classified as BC level 1, 1 as BC level 2, 0 as BC level 3, 34 as BC level 4, and 94 as BC level 5.

The case classified as BC level 1 (AER number [redacted]) is new and discussed below.

In the 14th SSR (16 Feb 2022-15 Apr 2022) (procedure EMEA/H/C/005735/MEA/002.13) there was reported another BC level 1 MIS-A (AER number [redacted]) which was considered confounded and therefore unlikely to be associated with Comirnaty. However, this case seems not to be retrieved in current safety database search.

The case classified as BC level 2 was previously assessed in the 13th SSR (16 Dec 2021-15 Feb 2022) (AER number [redacted]) and considered to be confounded by previous COVID-19 infection and causality is therefore considered unlikely.

BC level 1 case: AER #... 50-year-old female, case reported in the literature from...
The patient was hospitalized with fever, myalgia and shortness of breath approximately 2 weeks after dose 2 of BNT162b2 and antibiotics were administered. She was subsequently hospitalized approximately 6 weeks after dose 2 with persistent fever, widespread rash, proteinuria, and hyperferritinemia.

She developed pulmonary edema which required non-invasive mechanical ventilation. In addition to the persistent fever she developed seizures. Cerebrospinal fluid (CSF) analysis and magnetic resonance imaging of the brain were "noncontributory", and no evidence of infectious foci were observed serologically or microbiologically.

Her past medical record was significant for a childhood history of meningitis. She described no COVID-19 symptoms before presentation. Polymerase chain reaction (PCR) tests for SARS-CoV-2 were negative on multiple occasions. Physical examination showed widespread salmon-coloured rashes on her trunk and extremities.

C-reactive protein (CRP) of 286 mg/L, erythrocyte sedimentation rate (ESR) of 70 mm/h, ferritin of over 100,000 ng/ml (reference range (RR): 13-150), hemoglobin of 10.2 g/dl (RR: 11.9-14.6), aspartate aminotransferase of 594 U/L (RR: 0-33), alanine aminotransferase of 141 U/L (RR: 0-35), lactate dehydrogenase of 3,494 U/L (RR: 135-214), serum albumin of 3.0 mg/dl (RR: 3.5-5), creatinine of 1.0 mg/dl (RR: 0.4-0.98), D-dimer of 9.28 mg/l (RR: 0-0.5), fibrinogen of 289 mg/dl (RR: 170-420), proteinuria of 1.8 g/day, and serum proBNP of 10,522 pg/ml (RR: 0-125) with normal cardiac enzymes.

Echocardiography showed left ventricle systolic dysfunction and mild pericardial effusion.

Antinuclear antibody and rheumatoid factor were negative, bone marrow examination was unremarkable, and SARS-CoV-2 immunoglobulin (IgG) antibody levels were positive.

MIS - thought to be related to vaccine was considered and methylprednisolone was initiated. Persistent fever and high ferritin (>100,000) resulted in anakinra (Interleukin-1 receptor antagonist) being added to the treatment. The patient was discharged at day 21 after clinical and laboratory improvement and remains in remission on anakinra and low dose methylprednisolone.

**MAH comment:** The case is classified as BC level 1 based on the presence of persistent fever, mucocutaneous (rash) and neurological (seizures) clinical features, elevated inflammatory markers and 2 markers of disease activity (elevated proBNP and echocardiogram findings). The authors discount differential diagnoses of macrophage activation syndrome (discounted on normal bone marrow examination) and adult-onset Still’s disease (AOSD) although they feel the pulmonary oedema and cardiac failure are atypical. Many of the features described in the report are compatible with a diagnosis of AOSD; of note pleuritis and pericarditis are commonly described, and a biological hepatitis can be seen in up to 60% of patients. Myalgias are also common, particularly associated with fever spikes. The authors also consider MIS-A secondary to COVID-19 unlikely on the basis of a full vaccination history, presence of IgG antibodies, a negative SARS-CoV-2 PCR and a lack of COVID-19 symptoms preceding the illness. Of note it is unclear if the antibodies are anti-N or anti-S. Although providing sufficient clinical detail to classify the case as BC level 1 there is a notable lack of detail regarding the patient's complete blood count.

**Rapporteur assessment comment:**

Although MAH’s comment is acknowledged, the presented MIS-A case is considered a BC level 1 case and considered probably related with Comirnaty. However, a coincidental finding cannot be excluded either, given the extensive exposure of Comirnaty in adults.

**MAH’s conclusion**
In summary, 199 cases were reviewed for potential MIS for the PSUR period 19 December 2021 through 18 June 2022; 18 cases were classified as BC Level 1-3 MIS-C cases and 2 as BC level 1-3 MIS-A cases. As highlighted in the individual case commentaries, in the majority of cases there are either confounding elements or important clinical detail is missing which would facilitate causality assessment.

Considering the totality of the data, including the number of reports received in the context of the billions of doses of vaccine administered, the MAH does not consider that the currently available information supports a causal association between MIS-C/A and Comirnaty. No updates to current labelling or the Risk Management Plan are warranted at this time. Surveillance on this topic will continue.

**Rapporteur assessment comment:**

**MIS-C**

During the reporting period, the MAH identified a total of 47 potential new MIS-C cases. Of these, 10 were classified as BC level 1, 5 as BC level 2, 3 as BC level 3, 26 as BC level 4, and 3 as BC level 5.

The 10 BC level 1 MIC-C cases were previously assessed in the 13th SSR and the 14th SSR (procedures (procedure EMEA/H/C/005735/MEA/002.13).

Therefore the previous PRAC conclusion in the 14th SSR (procedure EMEA/H/C/005735/MEA/002.13) remains valid, i.e. that the data is currently insufficient to support regulatory action and therefore no update of the product information in relation to MIS-C is currently warranted. Although one additional MIS-C BC level 1 case (AER number [redacted], literature case from [redacted] that occurred in an 16-year old male) besides the Danish index case that was considered probably related with Comirnaty was identified, this is given the extensive exposure of Comirnaty (in children) not to be considered unexpected and does not present a new safety concern.

**MIS-A**

During the reporting period, the MAH identified a total of 130 potential new MIS-A cases. Of these, 1 was classified as BC level 1, 1 as BC level 2, 0 as BC level 3, 34 as BC level 4, and 94 as BC level 5.

The presented BC level 1 MIS-A case is considered probably related with Comirnaty, due to the absence of other etiologies or confounding. However, given the extensive exposure of Comirnaty in adults this is not considered unexpected and does not present a new safety concern.

Another BC level 1 MIS-A (AER number [redacted] reported by the MAH in the 14th SSR (interval period 16 Feb 2022-15 Apr 2022) was considered confounded by previous COVID-19 infection and therefore unlikely to be associated with Comirnaty. However, this case seems not to be retrieved in current safety database search. The MAH is requested to explain why the BC level 1 MIS-A case (AER number [redacted] from the 14th SSR was not included in current MIS-A overview (interval period 19 Dec 2021 to 18 Jun 2021). Request for supplementary information

No new important information could be identified concerning MIS-C/-A. The MAH should continue to closely monitor this safety issue as outlined in PRAC’s signal recommendation (EPITT 19732) and all new cases of MIS-C/-A should be reported in the future PSURs. Request for next PSUR

**Autoimmune hepatitis (AIH)**

PRAC request from the signal procedure concerning autoimmune hepatitis (EMEA/H/C/005735/SDA/042- EPITT 19749):
The MAH should provide in the next PSUR (submission date 27 August 2022) a cumulative review of all cases of autoimmune hepatitis, including any relevant new data, from all available sources. The cumulative review should include, but not be limited to, data from clinical trials, post-marketing cases and any relevant articles from literature, using a data lock-point as recent as possible.

MAH’s response (Appendix 6A.5 of the PSUR):

The safety database was searched for all BNT162b2 AE reports reporting PT: Immune-mediated hepatic disorder, Immune-mediated hepatitis, or Autoimmune hepatitis, cumulatively to 18 June 2022 using MedDRA version 25.

Results

A total of 194 cases (and 920 events) were retrieved from the global safety database with the search criteria mentioned above. There were 124 (63.9 %) females, 67 (34.5%) males and for 3 cases sex was not reported. Age was reported as ranging from 10 to 91 years (mean: 54.3, N: 187). There were 9 cases with age below 17 years, there were 111 cases within the adult category (≥18- <65 years), 67 elderly (≥65 years) and age was not reported in 7 cases. All cases were assessed as serious. Case outcome was reported as recovered, recovering, or recovered with sequelae in 93 cases; no outcome was reported at the time of reporting in 75 cases; outcome was unknown in 23 and fatal in 3 cases. The cases were reported mostly by Germany (57, 29.4%), Japan (23, 11.9%), UK (21, 10.8%) and France (21, 10.8%).

Medical history (MH) was reported in 111 out of 194 cases. Among these 111 cases, 21 described events consistent with liver disorders (of which 9 co-reported other underlying autoimmune diseases), an additional 27 cases reported underlying autoimmune diseases (other than autoimmunity hepatitis), 10 reported underlying cancer and 5 reported cholecystic disorders. A total of 169 of the 194 cases reported which vaccine dose was administered prior to the relevant adverse event (Table 2).

Table 2. Vaccine dose administered

<table>
<thead>
<tr>
<th>Dose</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>44</td>
</tr>
<tr>
<td>2</td>
<td>95</td>
</tr>
<tr>
<td>3</td>
<td>29</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>194</td>
</tr>
</tbody>
</table>

Time to onset was reported in 99 of the 194 cases and ranged between the same vaccination day up to 202 days post vaccination (mean: 22.8; median: 21; N: 99). Table 3 report details on the time to onset.

Table 3. Time to onset

<table>
<thead>
<tr>
<th>Days</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-2</td>
<td>9</td>
</tr>
<tr>
<td>3-7</td>
<td>29</td>
</tr>
<tr>
<td>8-15</td>
<td>26</td>
</tr>
<tr>
<td>16-30</td>
<td>34</td>
</tr>
<tr>
<td>&gt;30</td>
<td>57</td>
</tr>
<tr>
<td>Not reported</td>
<td>39</td>
</tr>
<tr>
<td>Total</td>
<td>194</td>
</tr>
</tbody>
</table>

A total of 60 of the 194 cases reported biopsy results, 39 cases did not report biopsy results but reported either viral markers/serum IgG or autoantibodies and 95 cases did not report any result of laboratory or diagnostic tests and therefore do not allow a proper assessment of the event.

Rapporteur assessment comment:

Post-marketing through 18 Jun 2022, 194 cases reporting autoimmune hepatitis were retrieved. A TTO of 3-30 days after Comirnaty exposure was reported in 89 of the 194 cases.
No results of laboratory or diagnostic tests was reported in 95 cases and did not allow a proper assessment of the event.

Biopsy results were reported in 60 cases and laboratory data only were reported in 39 cases.

Cases with liver biopsies and/or laboratory data

The International Scoring System for Diagnosis of Autoimmune Hepatitis and the Simplified criteria for the diagnosis of autoimmune Hepatitis were used when the data was provided.

The 99 cases reporting information (60 reporting biopsy results and 39 cases reporting laboratory examination were further analyzed based on diagnostic criteria for autoimmune hepatitis and divided in 2 groups:

1. Cases with reported liver biopsy results (n: 60)

Medical History was not reported in 43 cases, 17 did not report medical history, 11 autoimmune diseases, 8 pre-existing liver disorders, 5 ongoing cancers, 1 reported as substance use, 1 post-partum, 1 BRCA1 gene mutation, 15 did not report relevant MH to liver issues. Please note that one subject may have reported more than 1 of the MH events described above.

Most cases reported the event after dose 2 (Table 4).

<table>
<thead>
<tr>
<th>Dose</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>15</td>
</tr>
<tr>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>60</td>
</tr>
</tbody>
</table>

Time to onset ranged from the day of vaccination to 202 days after vaccination (Table 5).

<table>
<thead>
<tr>
<th>Days</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-2</td>
<td>4</td>
</tr>
<tr>
<td>3-7</td>
<td>10</td>
</tr>
<tr>
<td>8-15</td>
<td>10</td>
</tr>
<tr>
<td>16-30</td>
<td>9</td>
</tr>
<tr>
<td>&gt;30 (from 31 to 202)</td>
<td>20</td>
</tr>
<tr>
<td>Not reported</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>60</td>
</tr>
</tbody>
</table>

None of the cases provided sufficient information to use the International Autoimmune Hepatitis Group (IAIHG) revised scoring system for diagnosis of autoimmune hepatitis. However, 7 cases reported this score in the narrative. (Table 6, not reproduced here):

- These 7 cases had IAIHG-revised scores that ranged from 12 to 20. Six of the 7 cases are females, who are at increased risk of developing autoimmune disorders
such as AIH. Two of the 7 reports describe times to onset that suggest an unlikely association with vaccination (58 and 61 days). Of the remaining 5 cases, 2 had at least one other autoimmune disorder, suggesting a propensity for the development of AIH, and the remaining 3 did not provide any information about past medical history or concomitant medication use.

Although none of the remaining 53 cases with liver biopsies provided sufficient information for the IAIHG-revised scoring system, they were assessed using the IAIHG 4 simplified diagnostic criteria (1. positivity for autoantibodies, 2. elevated IgG levels, 3. Histological evidence of interface hepatitis and 4. the exclusion of viral hepatitis) by Hennes EM et al 2008. With this simplified criterion, a score of 7 is considered definite AIH and 6 is probably AIH (Table 7, not reproduced here):

- In these 9 cases assessed as having an IAIHG-simplified score of ≥6, 4 described a time to onset inconsistent with temporality to vaccination (31 days to 202 days). Three of the remaining 5 cases do not provide medical history or concomitant medication details. One of the remaining 2 cases describes the presence of EBV which is an alternative etiology for hepatitis.

The remaining 44 cases were assessed as having an IAIHG-simplified score <6 and are described below (subjects may have 1 or more of the following characteristics):

- In 15 cases, time to onset was not suggestive of a temporal relationship with vaccination (13 cases reported the time to onset between 31 to 90 days post vaccination (mean: 51 days); 11 of these 15 cases also lacked medical history or concomitant medication information, hampering a proper medical assessment,

- In 8 cases, an underlying autoimmune disorder was reported in medical history suggesting a predisposition to autoimmunity (e.g. thyroiditis, multiple sclerosis, pericarditis, coeliac disease and diabetes mellitus),

- In 6 cases, evidence of an underlying infection was described (e.g., EBV, CMV, herpes, tick bite on antibiotic treatment) suggesting an alternate etiology of hepatitis,

- In 4 cases, subjects had a pre-existing hepatic disorder (e.g. Autoimmune hepatitis, hepatitis C, chronic cholangitis),

- 4 subjects were undergoing treatment for cancer raising the possibility of drug-induced liver injury,

- In 3 cases, the reported time to onset was too close to vaccination (same vaccination day or the day after) for reasonable attribution,

- 1 case described illicit drug use which can cause hepatic dysfunction,

- 1 case was reported as post-partum raising the possibility of post-partum autoimmune hepatitis,

- 1 case was reported as hepatitis secondary to use of the immune checkpoint inhibitors.

One case reported a fatal outcome, the case is summarized below:

74-year-old female patient received BNT162b2 (COMIRNATY), dose 2 on 14 May 2021. Medical history included ongoing obesity, cholecystectomy, nephrectomy, bypass
surgery, ongoing non-alcoholic steatohepatitis. Concomitant medications included calcium, colecacalcalciferol, atorvastatin, paracetamol, oral acetylsalicylic acid. The same vaccination day the subject experienced fatigue and 3.5 months later was hospitalized for pneumonia, autoimmune hepatitis, circulatory collapse, respiratory failure, hepatotoxicity (all with onset 75 days after vaccination). The patient was found to have liver enzyme elevation. An abdominal ultrasound showed steatosis and minimal ascites. A liver biopsy showed acute and chronic inflammation with bridging necrosis and newly graci fibrosis and non-steatotic liver disease. Drug/Immunological reaction, AIH or viral hepatitis were suspected as was cirrhosis. Hepatitis B, C, E virus tests were negative; Hepatitis A antibody: IgG positive, IgM negative; Immunoglobulins: Normal; Smooth muscle antibody, Anti-cyclic citrullinated peptide antibody: Positive; Mitochondria antibody: Negative. The patient date of death was 20 Sep 2021 and was reported to be due to pneumonia, circulatory collapse, respiratory failure, autoimmune hepatitis. An autopsy was not performed.

**Rapporteur assessment comment:**

Of the 60 cases reporting biopsy results there were:

- 7 cases with an IAIHG-revised scores that ranged from 12 to 20:
  - 2 cases had a TTO > 30 days (unlikely association with vaccination),
  - 2 cases had at least one confounding by underlying conditions,
  - 3 cases had no information on medical history or concomitant medication use.

- 9 cases with an IAIHG-simplified score of ≥6:
  - 4 cases had a TTO > 30 days (unlikely association with vaccination),
  - 5 cases had no information on medical history or concomitant medication use.

- 44 cases with an IAIHG-simplified score <6:
  - 15 cases had a TTO > 30 days (unlikely association with vaccination),
  - 3 cases had a TTO < 3 days (unlikely association with vaccination),
  - 17 cases at least one confounding by underlying conditions,
  - 6 cases had evidence of an underlying infection,
  - 2 cases had drug induced autoimmune hepatitis,
  - 1 fatal case - TTO >30 days, underlying confounding conditions and concomitant medications.

The MAH did not perform a causality assessment for each of the autoimmune hepatitis cases.

Based on review of these 60 cases reporting biopsy results, the PRAC Rapporteur considers that a causal relation between Comirnaty and autoimmune hepatitis is:

- Unlikely in 52 cases due to non-plausible TTO, underlying conditions and/or concomitant confounding medication.
- Unassessable in 8 cases, due to limited information.
2. Cases where the diagnosis was based on laboratory data only (n: 39)

In 39 cases, only laboratory data was provided as a basis for the diagnosis of AIH. None of the cases had sufficient information to score ≥ 6 using the IAIHG-simplified score.

Characteristics of the 39 cases are described below (subjects may have 1 or more of the following characteristics):

- 13 cases did not report either MH or concomitant medication hampering a proper medical assessment,
- 9 cases that reported previous underlying hepatic issues and report the event as recurrence (1 case in the context of recurrent AIH in liver transplant),
- 8 cases reported a time to onset not suggestive of a temporal relationship with vaccination (ranging from 35 to 128 days post vaccination; mean: 77 days),
- 6 cases did not report time to onset precluding a proper causality assessment,
- 8 cases were concomitantly diagnosed with Infection (3 Covid-19, 2 hepatitis viral, 1 viral meningitis, 1 EBV and 1 Pneumonia), that are known to be possibly associated with liver issues,
- 4 cases were reported in the context of cancer under treatment (few cases reported that the treatment had also triggered previously other autoimmune diseases) suggesting an alternative aetiology for AIH (mostly reported as nivolumab or pembrolizumab),
- 3 subjects had concomitant autoimmunity diseases (such as thyroiditis, diabetes and Sjogren's syndrome), suggesting an increased risk for the development of AIH
- 2 cases were reported as suspected drug induced liver dysfunction,
- 1 case reported alcohol use which is a potential alternative explanation for the hepatic lab abnormalities
- 1 case unspecified illicit drug dependance (even if the drug is not specified most illicit drug are known to cause liver issues)
- 1 case was diagnosed with variable Immunodeficiency syndrome (pre-existing). This immune disorder is characterized by recurrent infections and low antibody levels,
- Autoimmune diseases are present in about 50% of the affected people and liver issues are common.

One case reported a fatal outcome and is summarized below:

A 76-year-old male patient received BNT162b2 on 04 Jan 2022 as dose 3. The vaccine for primary COVID-19 immunization was not known. The patient's relevant medical history included: Arterial hypertension, Steatosis hepatic, Coronary heart disease, Autoimmune hepatitis, Polyneuropathy. The patient's concomitant medications were not reported. Ten days post dose 3 he presented with abdominal pain and fulminant hepatitis and steroids were started. His general condition deteriorated, and he was diagnosed with hepatocellular carcinoma less than 2 months later; he received supportive care and passed away on an unspecified date.

*Rapporteur assessment comment:*
There were 39 cases reporting laboratory data only. The MAH did not perform a causality assessment for each of the autoimmune hepatitis cases.

Based on review of these 39 cases with diagnosis based on laboratory data only, the PRAC Rapporteur considers that a causal relation between Comirnaty and AIH is:

- Unlikely in 26 cases due to non-plausible TTO, underlying conditions and/or concomitant confounding medication.
- Unassessable in 13 cases, due to limited information.

Cases without liver biopsy or laboratory data (95 cases)

A total of 95 cases did not report any biopsy or laboratory data to confirm the diagnosis of AIH. These cases did not provide sufficient information to perform a medical assessment. Given the lack of information reported to assess the diagnosis the diagnostic criteria for autoimmune hepatitis could not be applied. Lack of information hampers a full assessment.

One case reported a fatal outcome and is summarized below:

[Redacted] an 88-year-old male patient received second dose of BNT162b2 20Mar2021. Patient concurrent conditions included hypertension, hypercholesterolemia, diabetes, and pruritus. Concomitant medications included insulin glargine; metformin; amlopidine; lansoprazole. The patient experienced autoimmune hepatitis and jaundice 31 days following the second vaccination and aggravation of already known pruritic condition, severe rash, hepatic cirrhosis and kidney and liver failure on an unknown date in 2021. The patient was treated with an unspecified Adrenal cortical hormone. CT scan on an unknown date in 2021 was normal. Blood test on an unknown date in 2021 showed high liver enzymes. Gastroscopy and ultrasounds scan on an unknown date in 2021 showed minor infection and minor gallstones. The patient died more than 100 days after vaccination. Reported causes of death: Autoimmune hepatitis, Hepatic cirrhosis, Liver failure and Kidney failure, Jaundice, Pruritus aggravated.

Rapporteur assessment comment:

It is agreed that causality cannot be established in these 95 cases in absence of data to support autoimmune hepatitis diagnosis.

The fatal case [Redacted] was previously assessed in the closed signal procedure concerning autoimmune hepatitis (EMEA/H/C/005735/SDA/042- EPITT 19749): The underlying confounding condition(s) e.g. diabetes mellitus and extensive concomitant mediation (amlodipine, lansoprazole, metformin, of which the SmPC already labels increased liver enzymes and liver disorders such as hepatitis and jaundice) hampers causality assessment.

Clinical trial data

There were no reports of autoimmune hepatitis in the Pfizer-run, placebo-controlled Phase 2/3 Study C4591001 in participants 16 years and older from dose 1 to 1 month after dose 2 (data cut-off date 13 March 2021); the safety population consisted of 21926 participants in the BNT162b2 group and 21921 participants in the placebo group.

Rapporteur assessment comment:

There were no reports of autoimmune hepatitis in clinical trial C4591001.
Observed versus expected analyses

The MAH conducted unadjusted observed to expected ratio (O/E) analyses for the 194 autoimmune hepatitis cases reported cumulatively through 18 June 2022 globally (Table 10). The overall expected case counts of autoimmune hepatitis were estimated using background incidence rates (IR) reported by a population-based study of the Clinical Practice Research Datalink (CPRD) in England during 1997-2015. The overall incidence rate was 2.08 (95% CI 1.94-2.22) per 100,000 population per year, with incidence peaking at approximately 70 years of age for both men and women. This background incidence rate is the middle of a range reported by a systematic review of 17 studies worldwide of 0.42 (United Kingdom, 1971-1987) to 3.00 (West Suffolk, England, 2003-2004) per 100,000 population (Jepsen P et al 2015, Tanner AR et al 1989, Whalley S et al 2007). The CPRD study was selected because it reported recent incidence rates by age, covering a large population.
Table 10. Observed to Expected (O/E) Ratio for Spontaneously Reported Cases of Autoimmune Hepatitis Through 18 June 2022

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Observed cases</th>
<th>Time at risk (PY)</th>
<th>Background rates per 100,000 PY</th>
<th>Expected cases</th>
<th>O/E ratio</th>
<th>95% CI LL</th>
<th>95% CI UL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>21-day risk window</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>3</td>
<td>1,917,834</td>
<td>0.51</td>
<td>10</td>
<td>0.307</td>
<td>0.063</td>
<td>0.896</td>
</tr>
<tr>
<td>12-17 years</td>
<td>1</td>
<td>3,324,066</td>
<td>1.06</td>
<td>35</td>
<td>0.028</td>
<td>0.001</td>
<td>0.158</td>
</tr>
<tr>
<td>18-24 years</td>
<td>3</td>
<td>4,733,892</td>
<td>0.84</td>
<td>40</td>
<td>0.075</td>
<td>0.016</td>
<td>0.220</td>
</tr>
<tr>
<td>25-49 years</td>
<td>19</td>
<td>19,546,111</td>
<td>1.24</td>
<td>242</td>
<td>0.078</td>
<td>0.047</td>
<td>0.122</td>
</tr>
<tr>
<td>50-59 years</td>
<td>14</td>
<td>9,025,268</td>
<td>3.29</td>
<td>297</td>
<td>0.047</td>
<td>0.026</td>
<td>0.129</td>
</tr>
<tr>
<td>60-69 years</td>
<td>11</td>
<td>7,901,305</td>
<td>4.19</td>
<td>331</td>
<td>0.033</td>
<td>0.017</td>
<td>0.059</td>
</tr>
<tr>
<td>70+ years</td>
<td>13</td>
<td>10,962,290</td>
<td>3.3</td>
<td>362</td>
<td>0.036</td>
<td>0.019</td>
<td>0.061</td>
</tr>
<tr>
<td>Overall, any dose</td>
<td>64</td>
<td>57,410,765</td>
<td>2.08</td>
<td>1194</td>
<td>0.054</td>
<td>0.041</td>
<td>0.068</td>
</tr>
<tr>
<td>Overall, dose 1</td>
<td>17</td>
<td>22,917,286</td>
<td>2.08</td>
<td>477</td>
<td>0.036</td>
<td>0.021</td>
<td>0.057</td>
</tr>
<tr>
<td>Overall, dose 2</td>
<td>35</td>
<td>21,280,003</td>
<td>2.08</td>
<td>443</td>
<td>0.079</td>
<td>0.055</td>
<td>0.110</td>
</tr>
<tr>
<td>Overall, dose 3</td>
<td>12</td>
<td>13,213,476</td>
<td>2.08</td>
<td>275</td>
<td>0.044</td>
<td>0.023</td>
<td>0.076</td>
</tr>
<tr>
<td>Overall Global</td>
<td>94</td>
<td>127,875,389</td>
<td>2.08</td>
<td>2643</td>
<td>0.036</td>
<td>0.029</td>
<td>0.044</td>
</tr>
</tbody>
</table>

| **42-day risk window** |                |                   |                                 |                |           |           |           |
| US/EEA               |                |                   |                                 |                |           |           |           |
| ≤11 years            | 3              | 2,779,982         | 0.51                            | 14             | 0.212     | 0.044     | 0.618     |
| 12-17 years          | 3              | 5,127,156         | 1.06                            | 54             | 0.055     | 0.011     | 0.161     |
| 18-24 years          | 4              | 7,477,186         | 0.84                            | 63             | 0.064     | 0.017     | 0.163     |
| 25-49 years          | 31             | 30,991,113        | 1.24                            | 384            | 0.081     | 0.055     | 0.115     |
| 50-59 years          | 20             | 14,461,569        | 3.29                            | 476            | 0.042     | 0.026     | 0.065     |
| 60-69 years          | 20             | 12,856,855        | 4.19                            | 539            | 0.037     | 0.023     | 0.057     |
| 70+ years            | 22             | 17,896,776        | 3.3                             | 591            | 0.037     | 0.023     | 0.056     |
| Overall, any dose    | 103            | 91,590,667        | 2.08                            | 1905           | 0.054     | 0.044     | 0.066     |
| Overall, dose 1      | 30             | 22,917,286        | 2.08                            | 477            | 0.063     | 0.042     | 0.090     |
| Overall, dose 2      | 56             | 42,494,750        | 2.08                            | 884            | 0.063     | 0.048     | 0.082     |
| Overall, dose 3      | 17             | 26,178,631        | 2.08                            | 545            | 0.031     | 0.018     | 0.050     |
| Overall Global       | 149            | 203,986,419       | 2.08                            | 4243           | 0.035     | 0.030     | 0.041     |

CI = confidence interval; EEA = European Economic Area; LL = lower limit; PY = person-years; UL = upper limit; US = United States


Based on the selected background rates and the estimated number of exposure person-years through 18 June 2022, O/E ratios were well below one overall, by age, and by dose for both risk windows of 21- and 42-days. This suggests that the number of observed cases of autoimmune hepatitis is not higher than expected in the absence of Pfizer-BioNTech COVID-19 vaccines overall and within the queried strata.

**Rapporteur assessment comment:**

All O/E ratios were well below 1.
Literature

The following database were searched: BIOSIS Previews <1969 to 2022 Week 31>, Embase <1974 to 2022 June 24>, Ovid MEDLINE(R) and Epub Ahead of Print, In-Process, In-Data-Review & Other Non-Indexed Citations and Daily <1946 to June 24, 2022>.

A total of 64 articles were retrieved. Relevant articles that are new compared to the previous review (previous PRAC request) where the literature was up to 2022 January 7th are discussed below:

Chow K W et al performed a systematic review to analyze every published case of AIH reported following COVID-19 vaccination and reviewed their characteristic findings, treatment, and outcomes. The data were retrieved using PubMed, Embase, and Web of Science from December 1, 2019, to November 1, 2021. The authors identified and analyzed a total of 32 cases. Although they concede that causality “cannot be proven” by this review, the authors refer to the cases as “vaccine-induced AIH” and hypothesize that molecular mimicry may be the mechanism leading to autoimmune tissue damage in susceptible individuals. They then conclude the article by stating that their findings should “under no circumstances deter individuals from getting vaccinated” as the benefits of vaccination outweigh the risks. (Chow KW et al 2022).

Roy A et al also performed a systematic review collating the available literature on potential Immune-mediated liver injury (ILI) following COVID-19 vaccination. Their final study selection included 23 patients with histopathological data in 13 studies. Reports included the AstraZeneca (n=11) and Pfizer/BNT COVID-19 (n=9) vaccines and time to onset ranged from 11.2-23.4 days. The authors note that 62.5% of the patients were female and ¼ of the patients had background autoimmune diseases; they describe jaundice as the most typical presenting symptom. IAIHG-simplified and IAIHG-revised scores were provided for 7 of the 23 cases and the authors calculated IAIHG-simplified scores for 11 cases (4 were probable, and 7 were definite AIH). Only one of the 23 patients reported a challenge-rechallenge. They conclude that most of the cases with ILI resembled AIH, that taking into consideration the limitations of the data, the crude incidence of ILI reported after vaccination seems to be far below the reported global incidence of AIH in the general population and that it is unclear whether COVID-19 vaccines are a trigger, causation or mere association with ILI.

Interestingly, Bril F wrote a letter stating that, while the resemblance cases reporting autoimmune hepatitis after Covid-19 vaccination suggests a potential causal link between the vaccine and AIH, this cannot be taken as proof that this link really exists. He further explains that considering an annual incidence of 1 case per 100,000 inhabitants as previously reported, and assuming an even distribution during the 12 months, we can estimate 1 monthly case per 1,200,000 inhabitants. Based on CDC data, during the first month of the US COVID-19 vaccination program, approximately 13,000,000 people received at least 1 dose of the vaccine (available at https://covid.cdc.gov/covid-data-tracker/#vaccinations). Based on AIH Incidence, we can therefore roughly estimate that ~10 people from this vaccinated cohort would have developed AIH within a month of getting the vaccine. Thus, it should not be surprising that we will all continue to see these cases as we continue with our vaccination efforts. Epidemiological studies assessing changes in AIH incidence may be able to shed some light on this uncertainty. Only if we observe a true increment of AIH incidence after COVID-19 vaccination, can we make the case for a potential relationship. Until then, primary care providers and hepatologists are encouraged to keep their eyes open and maximize adverse event reporting to the appropriate authorities (BrilF 2021). The same concept has been stated also very efficiently by other literature articles (Shroff H et al 2022, Lleo A et al 2022).

Bril F et al in another article state that while it is important to further explore the potential link between COVID-19 vaccines and autoimmunity, this should not discourage patients and physicians
from prescribing and/or receiving these vaccines. Even if confirmed in the future, cases of vaccine-induced autoimmunity appear to be uncommon as evidenced by only a handful of reports despite massive vaccination worldwide. Furthermore, it is likely that SARS-CoV-2 infection can trigger the same autoimmune processes as its vaccines do, leaving unvaccinated people not only at risk of developing the autoimmune process, but also the other complications associated with COVID-19 (Bril F et al 2021).

Overall, the literature at this time consists of case reports and case series which cannot confirm that COVID-19 vaccines cause autoimmune hepatitis due to confounding/triggering factors (as for example the post-partum state for the case reported from Bril et al, the underlying autoimmune disease (sarcoidosis) for the case described by Palla et al, and the use of liver toxic drugs in the cases described by Mc Shane et al and Clayton-Chubb et al.).

The European Association for the Study of the Liver (EASL), the American Association for the Study of Liver Diseases (AASLD), and British Association for the Study of Liver currently recommend the available SARS-CoV-2 vaccines for patients with chronic liver disease and liver transplant recipients (Nasa P et al 2021 and Sharma A et al 2021).

Rapporteur assessment comment:

MAH's overall statement regarding literature is accepted, the available case reports and case series cannot confirm that Comirnaty causes autoimmune hepatitis due to confounding/triggering factors.

MAH's conclusion

Overall, most of the potential AIH cases in the safety database do not provide sufficient information for a proper assessment, while cases that report information are mostly confounded by underlying comorbidities or concomitant drugs that represent an increased propensity for the development of AIH or potential alternative etiologies. Many cases reported the event with a time to onset that is not plausible for the vaccine to be causative (as for example the same vaccination day/1-2 days post vaccination or more than 30 days after vaccination).

The upper limit of the 95% confidence interval for the observed to expected ratio did not exceed 1; therefore, a signal was not identified.

In, C4591001, a large Pfizer-run double-blind placebo-controlled study of the vaccine there were no reports of Autoimmune hepatitis in either the vaccine or placebo groups, however, given the rarity of AIH, this is not unexpected.

Literature information consists of case reports and case series. Large population-based observational studies may eventually provide more useful information than cases series about the incidence of AIH in unvaccinated populations compared to those vaccinated with Pfizer/Biontech COVID-19 vaccine.

In addition, the pandemic has been associated with increased alcohol consumption, unhealthy eating habits, and interruptions to hepatology services, which might lead to an upward trend in liver disease incidence and severity that has been unnoticed until resumption of usual health care can return after the COVID-19 pandemic period.

Overall, given the totality of the available information, a causal association between Comirnaty and AIH cannot be concluded. Changes in the risk minimization measures or updates to the product information label are not warranted at this time. The benefit risk profile of the vaccine remains favorable. The topic will continue to be closely monitored.

Rapporteur assessment comment:
Post-marketing

Through 18 Jun 2022, 194 cases reporting autoimmune hepatitis were retrieved:

- Biopsy results were reported in 60 cases, a causal relation between Comirnaty and autoimmune hepatitis is considered Unlikely in 52 cases (due to non-plausible TTO, underlying conditions and/or concomitant confounding medication) and Unassessable in 8 cases (due to limited information).

- Laboratory data only were reported in 39 cases, a causal relation between Comirnaty and AIH is considered Unlikely in 26 cases (due to non-plausible TTO, underlying conditions and/or concomitant confounding medication) and Unassessable in 13 cases (due to limited information).

- Causality cannot be established in these 95 cases in absence of data to support autoimmune hepatitis diagnosis.

Clinical trial

No reports of autoimmune hepatitis in clinical trial C4591001.

Observed versus expected analyses

All O/E ratios for autoimmune hepatitis were well below 1.

Literature

The available case reports and case series cannot confirm that Comirnaty causes autoimmune hepatitis due to confounding/trIGGERING factors.

Therefore, MAH’s conclusion is accepted that a causal association between Comirnaty and autoimmune hepatitis cannot be concluded based on the available information. No new important information could be identified concerning autoimmune hepatitis. The MAH should closely monitor any new cases, patterns, or trends of reporting autoimmune hepatitis through routine pharmacovigilance.

Issue solved

Glomerulonephritis/nephrotic syndrome

Rapporteur assessment comment:

Here we refer to section ‘Evaluation of Other Risks (not categorised as important), Adverse events of special interest (AESIs)’ below in this AR and the overview of immunoglobulin A (IgA) nephropathy hereafter.

IgA nephropathy (IGAN)

Response to the PRAC request 5 from the 2nd PSUR (procedure EMEA/H/C/PSUSA/00010898/202112):

The MAH is requested to provide a cumulative review of cases reporting IgA nephropathy, including details of the underlying condition(s), time to onset, duration, outcome and an assessment of the causal relationship with the vaccination. The MAH should also provide a comprehensive discussion of the literature, mechanism of action and a discussion on the need to update the product information of Comirnaty, if applicable.

MAH’s response:
Background

In August 2021, the European Medicines Agency (EMA) issued a signal assessment report on glomerulonephritis and nephrotic syndrome and Comirnaty (EPIT no:19722). A cumulative review of the association of Comirnaty with glomerulonephritis and nephrotic syndrome using all sources (cases in the Market Authorisation Holder’s (MAH) safety database, clinical trial data, literature review and observed to expected analysis) was performed. This cumulative review covered the Preferred Terms (PTs) under the MedDRA v24.0 high level term (HLT) Glomerulonephritis and Nephrotic Syndrome, encompassing a broad range of renal pathologies, including IgA nephropathy (IgAN). The EMA’s Pharmacovigilance Risk Assessment Committee (PRAC) recommended that the MAH closely monitor the issue of ‘glomerulonephritis/nephrotic syndrome’ and present a cumulative review of cases from all sources and relevant literature in upcoming periodic safety update report (PSUR) submissions. PRAC subsequently requested a focused review of IgAN.

IgA nephropathy

IgAN is the most prevalent primary idiopathic glomerulonephritis globally, with an overall incidence of at least 2.5 per 100,000. Significant geographical variations are noted with lowest incidences in North America and Europe and highest in East Asian countries.

It has been proposed that geographical variations apparent in disease prevalence may represent regional differences in screening for renal disease and threshold for renal biopsy. Notably in Japan, all primary and secondary school students undergo urinalysis annually as part of a routine health check program under the supervision of the national government. Further testing of university students and routine health checks by employers mean that increased incidence noted in this population.

IgAN is characterised by persistent microscopic haematuria and/or proteinuria or recurrent episode of macroscopic haematuria concurrent with upper respiratory tract infections. It is often asymptomatic, identified on incidental urine analysis and diagnosed by renal biopsy with demonstration of diffuse mesangial IgA deposits in glomeruli.

Although patients can present at any age there is a peak incidence in the second and third decades of life. This is particularly pertinent given 20-40% of patients develop kidney failure within 10-20 years of diagnosis. A 2:1 male preponderance exists in North American and Western European populations, but IgAN is equally distributed between the sexes in Asian populations.

The pathogenesis of IgAN is widely accepted to follow a “multi-hit hypothesis”, involving confluence of genetic and environmental influences that modulate immune function and lead to IgAN.

Patients with IgAN have increased circulating levels of galactose-deficient IgA1 (Gd-IgA1), although this disordered glycosylation is heritable it is not sufficient in itself to cause disease. The triggers for the formation of Gd-IgA1 are not entirely understood, however exposure to environmental and infective factors may play a role. The abnormal IgA1 acts as an epitope for the formation of anti-glycan IgG antibodies. Circulating IgA immune complexes are identified in patients with IgAN, although IgA does not often activate complement the complexes contain Gd-IgA1, anti-glycan autoantibodies and complement C3. The circulating Gd-IgA1-anti-glycan IgG immune complexes deposit in the mesangium activating inflammatory and cellular proliferative signaling cascades.

The resultant local inflammation, mesangial matrix production and mesangial cell proliferation can lead to glomerular and interstitial fibrosis.

The pathognomonic finding of IgAN is the presence of mesangial IgA deposits identified by immunofluorescence-microscopy.
Clinical presentation varies from asymptomatic microscopic haematuria and or proteinuria to acute rapidly progressive glomerulonephritis. Approximately 40-50% present with gross haematuria, sometimes provoked by bacterial tonsillitis or viral upper respiratory tract infections. In these patients this first presentation is presumed to be the onset of disease although this is not proven. The concurrent "synpharyngitic" presentation differentiates this from post-streptococcal glomerulonephritis which has a usual 2-3-week latency from infection.

30-40% of cases are identified due to microscopic haematuria or proteinuria on incidental testing or during the investigation of chronic kidney disease; in these patients the disease duration is unknown.

Less than 10% of patients present with nephrotic syndrome or rapidly progressive glomerulonephritis; it is often presumed in these patients that the disease has been present for longer but not previously identified on routine urinalysis/or they have not had any prior episodes of macroscopic haematuria.

Clinical management focusses on supportive therapy; blood pressure control, reducing proteinuria with renin-angiotensin system blockage and lifestyle modifications to slow the rate of renal function decline. Immunosuppressive therapies are reserved for the highest risk patients.

IgAN can also be associated with other systemic conditions; chronic liver disease, coeliac disease, inflammatory bowel disease, Henoch-Schoenlein purpura and ankylosing spondylitis often termed "secondary IgAN".

IgA nephropathy, COVID-19 & Vaccines

Faroog et al systematically reviewed literature reports of IgAN and IgA vasculitis following COVID-19. They describe early seroconversion to IgA in COVID-19 patients and that mucosal infections are believed to enhance IL-6 production which stimulates poor galactosylation of IgA1, forming Gd-IgA1 and contributing to the disease process suggesting COVID-19 infection may trigger IgAN or flare via such mechanisms.

IgAN has not frequently been reported following preventative vaccinations (Wu et al.). Excessive production of IgA1 monomers in IgAN patients in response to Influenza vaccine has been previously seen although the clinical significance is not reported (van den Wall Bake eta al.).

Literature

A search of OVID MEDLINE (R) (1946)-present, OVID MEDLINE (R) In-Process & Epub Ahead of Print and Embase (1974 to 14 July 2022) was conducted with the following search strategy: BNT162B2 or Tozinameran or Comirnaty and the PT "IgA nephropathy" through 30 June 2022.

Of the 49 results retrieved, 28 reflected case reports or case series reported in association with COVID-19 vaccinations, of these, 22 reported cases relating to IgA nephropathy. 5 were not deemed valid individual case safety reports in the MAH safety database.

Eight (8) cases are represented in the cumulative dataset described below.

Nine (9) additional literature case reports are registered in the MAH safety database, Identified after the DLP. They do not provide sufficient clinical detail, consistency, specificity or strength of association to alter the conclusion on IgAN and Comirnaty after consideration of the totality of data.

Wu et al conducted a qualitative systematic review of new-onset and relapsed kidney histopathology cases reported in association with COVID-19 vaccination. The review includes 46 cases from 36 articles. 14 cases of IgAN were included; 6 new-onset, 6 previously known cases and 2 patients who
presented with macroscopic haematuria who did not have kidney biopsy and were presumed to have IgAN. 5 new onset cases were reported with a median time to onset of just one day (interquartile range (IQR) 1-2 days); 5 reported following Dose 2 of Moderna vaccine and 1 following Dose 2 of Comirnaty. Macroscopic haematuria was also reported in 6 patients with a previous histopathological diagnosis of IgAN with a similarly short median time to onset of 2 days (IQR 1-2 days). Onset was after Comirnaty in 3 cases (1 after dose 1, 2 after dose 2) and 3 after Dose 2 of Moderna COVID-19 vaccine. Further, 2 patients were included with presumed IgAN, given their presentation of macroscopic haematuria and mild proteinuria; both following Moderna COVID-19 Dose 2 vaccination. 1 of the patients had previously received episodic steroid therapy for IgA vasculitis as a child. The authors suggest that scenarios where subclinical IgAN becomes clinically apparent following COVID-19 vaccination are a possibility. They also suggest that the short latency from vaccination is similar to the concurrent onset of haematuria seen in sympharyngitic IgAN in clinical practice. (Wu HHL, Kaira PA, Chinnadurai R. New-Onset and Relapsed Kidney Histopathology Following COVID-19 Vaccination: A Systematic Review. Vaccines (Basel). 2021;9(11):1252.)

Racens et al investigated the impact of COVID-19 vaccination on the clinical course of IgAN in an adult Latvian patient population. 54 adult patients with a morphological diagnosis of primary IgAN at a single centre were enrolled. Clinical and laboratory parameters were evaluated at an inclusion visit and at a second visit 6 months later. Thirty-six patients were unvaccinated and 18 vaccinated; baseline proteinuria was the only significantly different parameter at baseline (lower 24-hour proteinuria in the vaccinated population). Fourteen patients were vaccinated with mRNA vaccines; 13 with Comirnaty and 1 with Spikevax. Four patients were vaccinated with Vaxzevria vector vaccine. Baseline renal function; estimated glomerular filtration rate (eGFR), haematuria, 24-hour proteinuria and disease activity markers; IgA, C3c were compared to the results at the 6-month follow-up visit with no significant difference noted in the vaccinated group compared to the non-vaccinated group. The authors concluded that COVID-19 vaccination did not affect the clinical course of IgAN, in this small prospective study. (Rācenis K, Saulīte AJ, Popova, et al. MO213: Sars-COV-2 Vaccination DID Not Affect the Clinical Course of IGA Nephropathy in Latvian Adult Cohort, Nephrology Dialysis Transplantation 2022;37(3):1145.)

Musetti et al conducted a retrospective study at a single centre in Italy from late December 2020 to 31 December 2021. Thirty-eight patients with immune-mediated nephropathies (either on or off immunosuppressive therapies), excluding those with end-stage renal disease or kidney transplant recipients were included. Seven IgAN patients were included. Five (5) male and 2 female patients, with an average age of 40.4±12.0 years. 42.9% were on immunosuppressants at the time of vaccination and 28.6% had prior relapses and the authors calculated a relapse rate prior to vaccination of 9.6 per 100 person-years. One IgAN patient had a relapse 5 days following vaccination and a post-vaccination relapse rate calculated at 34.8 per 100 person-years. The authors report a slightly higher rate of relapse post vaccination however conclude that relapses of immune-mediated nephropathies are uncommon after COVID-19 vaccination. (Musetti C, Fornara L, Cantaluppi V. MO239: Clinical Evaluation of Immunological and Clinical Recurrence of Immune-Mediated Nephropathies after SARS-COV-2 Vaccine, Nephrology Dialysis Transplantation, 2022;37(3).)

The MAH states that in summary there is a paucity of large-scale epidemiological data in the literature, however a single-centre prospective study reported that SARS-CoV-2 vaccination did not affect the clinical course of IgAN. Limited data in the literature case reports precludes meaningful assessments of case causality.

Rapporteur assessment comment:
Of the 49 results retrieved from the literature search through 30 Jun 2022, there were 22 reported cases relating to IgA nephropathy. 8 of the 22 case reports are presented in the cumulative safety
database case review below. Five cases reports were not considered valid reports by the MAH. Nine cases were retrieved after the DLP of current PSUR, however did not provide sufficient information to alter the conclusion on IgA nephropathy and Comirnaty.

In the retrieved literature, no new important information could be identified concerning IgA nephropathy.

Clinical trial data

In the blinded, placebo-controlled period of pivotal study C4591001 (data cut-off date 13 March 2021) 1131 adolescents 12-15 years of age had received primary vaccination with BNT162b2 and 1129 had received placebo. The median duration of follow-up for adolescents 12-15 years of age was ≥2 months after Dose 2. No participants reported IgA nephropathy. Of note, there were no participants in the BNT162b2 or placebo groups who had a medical history of IgA nephropathy.

Of participants 16 years of age and older, 22,026 participants had received primary vaccination with BNT162b2 and 22,021 participants had received placebo. During the blinded placebo-controlled follow-up period (up to the earlier of the time of unblinding or data cut off), 51.1% of participants in the BNT162b2 group and 51.4% of participants in the placebo group had a follow-up time between ≥4 months to <6 months after Dose 2. No participants 16 and older were reported to have IgA nephropathy. Of 22,026 participants 16 years of age and older who received primary BNT162b2, 1 had a medical history of IgA nephropathy; of the 22,021 placebo recipients, 1 had a history of IgA nephropathy.

There were no AE reports of IgA nephropathy in Study C4591007 in participants 5 to < 12 years of age (data cut-off 06 Sep 2021). Of the 1518 participants who received BNT162b2 10 mcg as a primary vaccine and of the 750 participants who received placebo, none had medical histories of IgA nephropathy.

Rapporteur assessment comment:

There were no reports of IgA nephropathy in the clinical trials.

Cumulative safety database case review

The safety database was searched for all cases of BNT162b2 reporting the MedDRA v 25.0 PT IgA nephropathy, and for cases with a medical history of IgA nephropathy coded with the PT Condition aggravated through 30 June 2022.

A total of 103 cases were retrieved. A majority (80) of the cases were spontaneous, 21 were from literature non-study while 2 were from clinical studies. A total of 100 cases were serious while 3 were non-serious. Table 1 provides the distribution of cases by age. The mean age was 33.5 years (103 cases).

<table>
<thead>
<tr>
<th>Age Range</th>
<th>Number of Cases</th>
<th>Percentage of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than or equal to 17 years</td>
<td>19</td>
<td>18.4 %</td>
</tr>
<tr>
<td>18 - 30 years</td>
<td>26</td>
<td>25.2 %</td>
</tr>
<tr>
<td>31 - 50 years</td>
<td>35</td>
<td>34.0 %</td>
</tr>
<tr>
<td>51 - 64 years</td>
<td>12</td>
<td>11.7 %</td>
</tr>
<tr>
<td>65 - 74 years</td>
<td>7</td>
<td>1.9 %</td>
</tr>
<tr>
<td>Greater than or equal to 75 years</td>
<td>1</td>
<td>1.0 %</td>
</tr>
<tr>
<td>Unknown</td>
<td>8</td>
<td>7.8 %</td>
</tr>
</tbody>
</table>
Regarding sex distribution, a majority (64) of cases were reported in female patients while 36 were in males and gender data was not provided for 3 remaining cases.

Concerning the countries of origin, of the 103 cases, most cases were received from Japan (36) and France (18). The top reporting countries are shown below in Table 2.

**Table 2. Reporting Country (n=103)**

<table>
<thead>
<tr>
<th>Country</th>
<th>Number of Cases</th>
<th>Percentage of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Japan</td>
<td>36</td>
<td>35.0%</td>
</tr>
<tr>
<td>France</td>
<td>18</td>
<td>17.5%</td>
</tr>
<tr>
<td>United States</td>
<td>10</td>
<td>9.7%</td>
</tr>
<tr>
<td>Australia</td>
<td>7</td>
<td>6.8%</td>
</tr>
<tr>
<td>Germany</td>
<td>7</td>
<td>6.8%</td>
</tr>
<tr>
<td>United Kingdom</td>
<td>5</td>
<td>4.9%</td>
</tr>
<tr>
<td>Netherlands</td>
<td>4</td>
<td>3.9%</td>
</tr>
<tr>
<td>Spain</td>
<td>3</td>
<td>2.9%</td>
</tr>
<tr>
<td>Italy</td>
<td>2</td>
<td>1.9%</td>
</tr>
</tbody>
</table>

The case outcome was recovered in 17 cases, recovering in 22 cases, recovered with sequel in 3 cases and not recovered in 42 cases while outcome was unknown in 19 cases. None of the cases had a fatal outcome.

Of these 103 cases, 2 were excluded from further analysis as PT “condition aggravation” in these cases was not in context of IgA nephropathy.

Of the remaining 101 cases, 75 were medically confirmed while 26 were non-medically confirmed cases. A total of 37 cases reported an aggravation of IgA nephropathy in context of medical history of the same while 64 reported new onset IgA nephropathy. However, 6 of these 64 cases reported a history of haematuria which could indicate that renal disease was present at baseline.

Latency from last dose was provided in 80/101 cases. Latency varied from 0 to 263 days. The latency from time of the vaccination until the development of the relevant event was reported as: the same day of vaccination to 3 days post vaccination for 53 relevant events, from day 4 to day 10 for 12 events, from day 11 to day 24 for 4 events, and post day 24 for 11 events.

Dosing information was provided in 95 of the 101 cases. Of these 95 cases, 62 reported IgA nephropathy following dose 2, 23 reported following dose 1 while 10 reported events following dose 3.

Of the 101 cases, 3 were excluded from further analysis as in these cases COVID-19 vaccine manufacturer was unknown.

Of the remaining 98 cases, 3 cases reported an implausible latency of 170-263 days. 52 of the remaining 95 cases had insufficient information regarding either latency, medical history, concomitant medications, relevant investigations, clinical course and/or event details thus, precluding a meaningful assessment of these cases both in terms of confirming the diagnosis of IgA nephropathy and/or assessing a relationship to vaccination. Of these 52 cases, 25 were medically unconfirmed cases.

The remaining 43 cases are presented below based on new onset (Table 3, N=24) or condition aggravated in cases with pre-existing history of IgA nephropathy (Table 4, N=19, not reproduced here).

**Table 3: Cases pertaining to IgA Nephropathy, new onset (n=24)**

<p>| AER Number | Age/Gender | Country | Medical history | Relevant PT(s) | Case summary | MAH comment |
|------------|------------|---------|----------------|---------------|--------------|-------------|-------------|
|            |            |         | Co-suspectif concomitant | Other PTs Latency (from) |               |             |             |</p>
<table>
<thead>
<tr>
<th>Dose Source Medically confirmed (Y/N)</th>
<th>medications</th>
<th>last dose, for IgA nephropathy</th>
<th>Seriousness Outcome</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>41 Y/F</td>
<td>Gestational diabetes</td>
<td>IgA nephropathy</td>
<td>Haematuria</td>
<td>The patient presented with subnephrotic range proteinuria, hypertension, and elevated serum creatinine 1 day after Dose 2 of Comirnaty. The patient received Dose 1 of Comirnaty 23 days before. A renal biopsy showed IgA nephropathy with fibrocellular and fibrous crescents. The chronic features on histopathology were reported to be suggestive of pre-existing undiagnosed IgA nephropathy that might have been unmasked after vaccination. Relevant investigations: Serum creatinine 153 μmol/L (normal range unknown), Urine dysmorphic red blood cells/microliter: above 200, Urine protein-to-creatinine ratio 2.03 g/g (normal range not provided), Serum IgG 12.9 g/L (normal range: 5.49-17.11 g/L), Serum IgA 6.4 g/L (normal range: 0.47-3.59 g/L), Serum IgM, 1.1 g/L (normal range: 0.15-2.59 g/L), Complement C3 0.83 g/L (normal range: 0.90-1.80 g/L), C4 0.2 g/L (normal range: 0.10-0.40 g/L), Anti-nuclear antibody: 1:320, homogeneous, anti-GBM antibody (Enzyme-linked immunosorbent assay [ELISA]): below 1.5. Patient treated with pulse methylprednisolone, followed by oral prednisolone and IV cyclophosphamide.</td>
</tr>
<tr>
<td>Dose 2 Literature Y</td>
<td>Not reported</td>
<td></td>
<td>1 day</td>
<td>Personal or family history of IgA nephropathy is not reported nor is a precedent infection or autoimmune disorder. The biopsy report indicates that the nephropathy is chronic raising the possibility that the onset may have preceded any vaccination. The patient also suffered from gestational diabetes.</td>
</tr>
</tbody>
</table>

*Rapporteur assessment comment (1):* Medically confirmed literature case reporting biopsy with pre-existing undiagnosed IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
| 13 Y/F | Salmonella test positive/ Tobacco user/ Viral pharyngitis | IgA nephropathy Acute kidney injury, Anuria, Pyrexia, Haematuria, Influenza, Proteinuria | Patient developed de novo IgA nephropathy within 24 hours following Dose 1 of Comirnaty manifesting as fever, asthenia, muscle pain and macroscopic haematuria. On examination, she had mild streptococcus-negative pharyngitis. 3 days later, a renal biopsy showed IgA nephropathy, Oxford score M1E1S1T0. Other labs: Blood creatinine: 5 mg/dl, blood urea nitrogen 96 mg/dl (normal ranges not reported), infectious testing negative, immunological test negative, urine protein 3.88 g/l, SARS-CoV-2 Serology negative. Kidney function rapidly deteriorated, and the patient became oliguric; subsequently, haemodialysis was started. Treatment consisted of IV methylprednisolone puises, followed by oral prednisone. Kidney function improved progressively; at 30 days postvaccination, serum creatinine had returned to almost normal values. | The reported co-infection is a potential and more likely aetiology of the nephropathy in this case; the latency of <24 hours after the first exposure to vaccination is less biologically plausible. |

**Rapporteur assessment comment (2):**
Medically confirmed literature case reporting a TTO of 1 day after dose 1 and a co-infection. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
| Atrial fibrillation/Urinary occult blood | IgA nephropathy  
Glomerulonephritis rapidly progressive, Haematuria, Headache, Oedema peripheral, Urinary occult blood positive, Decreased appetite, Malaise | On 04 Jul 2021, 4 days after Dose 2 of Comirnaty, the patient experienced haematuria aggravated, general malaise, lower leg oedema and inappetence. Patient received Dose 1 of Comirnaty 3 weeks before. 9 days later, the patient was diagnosed with IgA nephropathy based on a renal biopsy which showed IgA nephropathy with crescents. Labs: creatinine 3.51 mg/dl (normal range not reported), Protein urine 1.91 g/gCr (3+), urinary occult blood (3+), antinuclear antibody and ANCA were negative. In Dec 2020, creatinine 1.09 mg/dl, Protein urine (negative), urinary occult blood (2+). The patient was treated with steroids. The reporter stated, “The base was probably undiagnosed stable IgA nephropathy. It was suspected to be exacerbated by immune activation because of COVID-19 vaccination”. |

| 74 Y/M  
Dose 2  
Spontaneous | Edoxaban | The past medical history of occult blood in the urine and the reporter note that this may be an exacerbation raises the likelihood of underlying nephrology that evolved in accordance with the disease natural progression. |

### Rapporteur assessment comment (3):
Medically confirmed literature case reporting pre-existing undiagnosed IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

| Dermatitis atopic  
Duplimab | IgA nephropathy  
2 days  
Serious  
Not recovered | The patient was diagnosed with mesangial IgA nephropathy based on renal biopsy 2 days after receiving Dose 2 of Comirnaty. No information provided regarding first dose of COVID-19 vaccine. Evidence of isolated C3 consumption associated with a low positive level of anti-deoxyribonucleic acid antibodies at 33 IU. No additional information. |

| 40 Y/F  
Dose 2  
Spontaneous | 
| Limited information has been provided including why the patient had a renal biopsy. |

### Rapporteur assessment comment (4):
Spontaneous case with limited information which is considered unassessable.
<table>
<thead>
<tr>
<th><strong>20 Y/F</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dose 2</strong></td>
</tr>
<tr>
<td><strong>Spontaneous</strong></td>
</tr>
<tr>
<td>N</td>
</tr>
<tr>
<td>Not reported</td>
</tr>
<tr>
<td>IgA nephropathy, Haematuria</td>
</tr>
<tr>
<td>1 day</td>
</tr>
<tr>
<td>Serious</td>
</tr>
<tr>
<td>Unknown</td>
</tr>
<tr>
<td>1 day after receiving Dose 2 of Comirnaty, the patient experienced massive macrohaematuria. No information on first dose of COVID-19 vaccine. IgA nephropathy diagnosed based on renal biopsy on an unknown date (details unspecified). Reportedly, the patient already had persistent microhaematuria (onset unknown). No additional information.</td>
</tr>
<tr>
<td>Limited information regarding timing between 2 doses of COVID-19 vaccine. Details are lacking regarding labs, clinical course, concomitant medications and medical history precludes an adequate assessment.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (5):**

Spontaneous case with limited information which is considered unassessable.

<table>
<thead>
<tr>
<th><strong>40 Y/F</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dose Unknown</strong></td>
</tr>
<tr>
<td><strong>Spontaneous</strong></td>
</tr>
<tr>
<td>Y</td>
</tr>
<tr>
<td>None</td>
</tr>
<tr>
<td>Not reported</td>
</tr>
<tr>
<td>IgA nephropathy, Nephrotic syndrome, Proteinuria, Hypoproteinaemia, Hyperlipidaemia</td>
</tr>
<tr>
<td>7 days</td>
</tr>
<tr>
<td>Serious</td>
</tr>
<tr>
<td>Unknown</td>
</tr>
<tr>
<td>1 week after receiving Comirnaty (dose unknown), the patient experienced generalized oedema and weight increased. Examination revealed marked proteinuria (14 g/day), hypoproteinaemia, and hyperlipidaemia (values unknown) and was diagnosed with nephrotic syndrome. A renal biopsy showed mild mesangial proliferation and adhesion. On immunofluorescence, IgA, IgM, IgG, and C3c presented with the positive image of mesangial pattern and electron microscopy showed disappearing image of foot processes. Steroid pulse therapy led to complete remission within 10 days. Based on renal biopsy, the possibility of IgA nephropathy was considered. However, a possibility of minimal lesion nephrotic syndrome was also considered as there was no haematuria and immediate remission was observed.</td>
</tr>
<tr>
<td>No information on dose number of covid vaccine, medical history and concomitant medications precludes an adequate assessment. Further, the clinical presentation (no haematuria and immediate remission) raised the possibility that another nephropathy was present.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (6):**

Spontaneous medically confirmed case with limited information which is considered unassessable.
<table>
<thead>
<tr>
<th><strong>12 Y/M</strong></th>
<th>None</th>
<th>IgA nephropathy</th>
<th>The same day as receiving Dose 2 of Comirnaty, the patient experienced frank haematuria (3+), pyrexia and proteinuria (2+). History of frank haematuria an unspecified duration after receiving first dose of Comirnaty 23 days earlier. After that, only urinary occult blood continued. No other details provided.</th>
<th>Although rechallenge positive for haematuria, no information regarding concomitant medications, confirmatory diagnostic tests including biopsy and labs precluded an adequate assessment.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Spontaneous</strong></td>
<td>Not reported</td>
<td>Haematuria, Pyrexia, Proteinuria</td>
<td>0 day</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (7):**
Spontaneous medically confirmed case with limited information which is considered unassessable.

| **63 Y/F** | Hypertension / Psoriatic arthropathy | IgA nephropathy | The patient presented with gross haematuria for 6 weeks, 3 days after the Dose 2 of Comirnaty. Labs: creatinine 10 mg/dl (baseline 0.5; normal range unknown) and urine protein: creatinine ratio of 7.3 gm/gm (normal range unknown). Renal imaging including CT urogram was normal. Renal biopsy showed IgA nephropathy with a fibrocellular crescent and acute tubular necrosis likely secondary to lysed red cells in setting of multiple RBC casts in the tubules. The patient was treated with steroids and creatinine decreased to 4.5 in 15 days. As per the authors "SARS-COV2 vaccines use nucleoside modified purified mRNA which does elicit higher neutralizing antibody titer and strong cluster of differentiation response leading to production of several proinflammatory cytokines. Thus, there is a concern that vaccines might exacerbate immune mediated glomerular diseases. IgA1 is involved in the pathogenesis of IgA nephropathy and patients |
| --- | Not reported | 3 days | Serious | Unknown |

**History of autoimmune disease (psoriatic arthritis) is a possible risk factor for IgA nephropathy in this case. No information on concomitant medications and first dose of covid vaccine.**
with IgA nephropathy have higher than normal IgA1 response to other vaccines like influenza. Also, while studying the antibody response to covid-19 illness, patients with IgA nephropathy are known to express higher IgA response compared to IgG and IgM along with reports of concurrent worsening of the glomerulonephritis.

**Rapporteur assessment comment (8):**
Spontaneous medically confirmed case reporting a history of psoriatic arthritis. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

<table>
<thead>
<tr>
<th>50 Y/F</th>
<th>Appendicitis / Obesity / Peritonitis / Pregnancy</th>
<th>IgA nephropathy, Acute kidney injury, Haematuria, Pyrexia, General physical health deterioration</th>
<th>1 day after receiving Dose 2 of Comirnaty, the patient presented with macroscopic haematuria. No details provided regarding the first dose of covid vaccine. Labs: creatinine: 220 μmol/l (normal range unknown); blood Immunoglobulin A: 2.82; GFR 27, blood urea 11.5 (unit and normal range unknown). A biopsy 9 days after vaccination revealed glomerulonephritis with mesangial IgA deposits. Extensive vascular lesions were noted. Protein urine: (unspecified date) 0.9 g/l, rheumatoid factor: (unspecified date) 5.3 IU/ml (normal range unknown).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose 2</td>
<td>Spontaneous Y</td>
<td>Recovering</td>
<td>No information regarding timing between two doses of covid vaccines and concomitant medications.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (9):**
Spontaneous medically confirmed case reporting no information regarding concomitant medications and timing of dose 1, and therefore considered unassessable.
<table>
<thead>
<tr>
<th>Rapporteur assessment comment (10):</th>
</tr>
</thead>
<tbody>
<tr>
<td>A medically confirmed case reporting confounding factors for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.</td>
</tr>
</tbody>
</table>

| Abortion/ Anti-antibody/ COVID-19/ Cystitis / Hypothyroidism/ Lymphoma/ Suicide attempt/ Uterine cancer | IgA nephropathy, Acute kidney injury, Nephritis, Aethenia, Oedema peripheral, Myalgia, Decreased appetite, Weight decreased, Oropharyngeal candidiasis, Rash erythematous, Arthralgia, Cystitis, Oral candidiasis, Normocytic Anaemia | 1 week after Dose 2 of Comirnaty, the patient developed asthenia, oedema of the lower limbs, myalgia then secondarily anorexia and weight loss, oropharyngeal candidiasis. A blood test 2 weeks later revealed acute renal failure with creatinine at 18 mg/l (baseline 7.8 mg/l; normal range not reported). A urine strip test showed proteinuria 3+ and haematuria 3+. The urine test revealed proteinuria at 0.85 g/24 hours, including 50% albumin, albuminuria at 0.4 g/day; associated microscopic haematuria at more than 1000/mm3, associated hypoalbuminemia at 32 g/l (normal range unknown). | Antithyroid antibodies are suggestive of an underlying autoimmune disorder. No information provided regarding concomitant medications. |
Serum protein electrophoresis showed hypo-serum albuminemia, hyper alpha-1, hyper alpha-2, restriction of heterogeneity in the area of beta globulins and gamma globulins, immunofixation qualitatively normal. A renal biopsy showed nephropathy with mesangial deposits of Immunoglobulin. Immunological assessment: Negative anti-neutrophil cytoplasm antibodies 1/20 of homogeneous appearance, myeloperoxidase or proteinase 3 negative, rheumatoid factor negative, positive anti-nuclear antibodies of homogeneous appearance 1/320 (threshold 1/80), antideoxyribonucleic acid negative, no specificity identified, viral serology was negative for Human Immunodeficiency Virus (HIV), hepatitis B virus (HBV), hepatitis C virus (HCV).

**Rapporteur assessment comment (11):**  
Spontaneous medically confirmed case reporting confounding factors for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>Dose</th>
<th>Spontaneous</th>
<th>None</th>
<th>IgA nephropathy, Urinary retention, Haematuria, Poliakuria, Immunisation 1 day Serious Not recovered The patient experienced poliakuria, feeling of residual urine, and haematuria 1 day after receiving Dose 3 of Comirnaty. No information on primary immunization for COVID-19. A urine analysis showed protein urine 1+, occult blood 3+, RBC more than 100/HPF, WBC more than 100/HPF. A latency of 1 day after dose 3 but no Information regarding supporting labs/investigations (including biopsy), medical history and concomitant medications.</th>
</tr>
</thead>
<tbody>
<tr>
<td>46 Y/F</td>
<td></td>
<td>Dose 3</td>
<td>Spontaneous Y</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (12):**  
Spontaneous medically confirmed case with limited information which is considered unassessable.
| COVID-19 / Haematuria / Mite allergy / Seasonal allergy | IgA nephropathy, Off label use, Interchange of vaccine products, Renal failure, Fatigue | 1 week after receiving Dose 2 of Comirnaty, the patient was diagnosed with IgA nephropathy. The patient received first dose of other manufacturer's Covid vaccine 6 months earlier. A kidney biopsy showed IgA nephropathy with crescentic glomerulonephritis, GFR 17 ml/min (normal range unknown). Family history of kidney transplant (unspecified diagnosis). No additional information. | Interchange of vaccines reported. The patient had a family history of kidney transplant (unspecified diagnosis). Additionally, the patient had a history of macroscopic haematuria once, a few years ago suggesting a pre-existing condition, possibly genetic. Baseline labs not reported. |
| Dose 2 Spontaneous Y | Not reported | |

**Rapporteur assessment comment (13):**

Spontaneous medically confirmed case reporting confounding information in the medical history for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

| COVID-19 pneumonia / Cholangitis / Colitis / Endoscopic retrograde cholangiopancreatography / Pregnancy / Tooth extraction | IgA nephropathy, Nephropathy, Urine odour abnormal, Nephroangiosclerosis, Tubulointerstitial nephritis, Glomerulonephritis, Urine abnormality, Proteinuria, Haematuria, Creative protein increased, Inflammation | Approximately 3 months after the Dose 1 of Comirnaty, the patient was diagnosed with IgA nephropathy based on a kidney biopsy. Immunofluorescence results were suggestive of active subacute and chronic mesangial nephropathy. Serum creatinine 86 μmol/l (normal range not reported), haematuria at 1,000,000/ml. The patient also experienced an inflammatory syndrome with CRP at 14 mg/L and a sedimentation rate of 51 mm; normal ranges unspecified. Spontaneously favourable outcome. | Long latency of 3 months suggests a biologically less plausible causal association. In addition, immunofluorescence results were suggestive of active subacute and chronic mesangial nephropathy thus, indicating a pre-existing disease. Besides, an inflammatory syndrome is suggestive of concurrent infection potentially having a role in the nephropathy. |
| 34 Y/F | Metronidazole, Spiramycin | |
| Dose 1 Spontaneous Y | |

**Rapporteur assessment comment (14):**

Spontaneous medically confirmed case reporting a TTO of 3 months after 1st dose and confounding factors for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
<table>
<thead>
<tr>
<th>16 Y/M</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose 2</td>
</tr>
<tr>
<td>Spontaneous</td>
</tr>
<tr>
<td>Y</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Haematuria</th>
<th>Renal impairment, IgA nephropathy, Haematuria, Pyrexia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not reported</td>
<td>1 day</td>
</tr>
<tr>
<td></td>
<td>Serious</td>
</tr>
<tr>
<td></td>
<td>Not recovered</td>
</tr>
</tbody>
</table>

The patient presented with pyrexia and frank haematuria the following day of receiving Dose 2 of Comirnaty. Family history (mother) of IgA nephropathy. Details of first dose of covid vaccine unknown. Serum creatinine level increased from 0.87 to 1.26 mg/dL and urine protein to creatine ratio was 0.35 g/gCr; normal ranges unknown. On the 24th day of vaccination, a renal biopsy revealed crescent formation and mild mesangium proliferation, leading to a diagnosis of IgA nephropathy. The patient was treated with steroids and immunosuppressants.

Family history of IgA nephropathy confounds the assessment as does pre-existing haematuria. No information on first dose of covid vaccine and concomitant medications.

*Rapporteur assessment comment (15):*
Spontaneous medically confirmed case reporting confounding factors for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
<table>
<thead>
<tr>
<th>Age/Gender</th>
<th>Symptoms</th>
<th>Diagnosis</th>
<th>Description</th>
<th>Inference</th>
</tr>
</thead>
<tbody>
<tr>
<td>34 Y/F</td>
<td>Arthralgia, Gestational diabetes, Migraine, Psoriasis, Synovial cyst removal</td>
<td>IgA nephropathy, Maternal exposure during pregnancy</td>
<td>The patient received 2 doses of Comirnaty at an interval of 22 days. 47 days after second dose, proteinuria and haematuria was detected on urine strip test. Approximately 2 months later, the patient had her first nephrology consultation: renal function was normal with creatinine at 49 µmol/L, urea at 4 mmol/L (normal ranges not reported), micro haematuria at 386/mm3, proteinuria at 1.07 g/L, HBV positive. A renal biopsy revealed 36 permeable glomeruli optically with a sclerohyalin glomerulus, permeable glomeruli represented the mesangial axes that were frequently enlarged with an increase in cellularity. Discrete lesions with inability to make a formal diagnosis in the absence of a direct immunofluorescence study.</td>
<td>History of autoimmune disease (psoriasis) is a risk factor for IgA nephropathy as is chronic liver disease from HBV although it is not clear if the patient has active liver disease. In addition, the long latency from vaccination is suggestive of a less likely biological association to vaccination. Finally, renal parameters were normal, and biopsy was not confirmatory, lacking immunofluorescence to confirm the deposition of IgA.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (16):**
Spontaneous medically confirmed case reporting a TTO of 47 days and confounding factors in medical history for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

<table>
<thead>
<tr>
<th>Age/Gender</th>
<th>Symptoms</th>
<th>Diagnosis</th>
<th>Description</th>
<th>Inference</th>
</tr>
</thead>
<tbody>
<tr>
<td>15 Y/M</td>
<td>Haematuria, Proteinuria present, Urinary occult blood positive</td>
<td>IgA nephropathy, Mesangioluproliferative Glomerulonephritis, Pyrexia</td>
<td>The following day of receiving Dose 2 of Comirnaty, the patient experienced pyrexia and frank haematuria. 1 week later, a renal biopsy showed mild mesangioluproliferative nephritis. In one glomerulus, cellular crescents were observed. Fluorescence antibody technique revealed IgA deposition in the mesangial region leading to a diagnosis of IgA nephropathy. No information regarding first dose of covid vaccine.</td>
<td>No information on first dose of covid vaccine and concomitant medications. The patient already had history of occult blood in urine and haematuria suggesting disease at baseline with natural progression should be considered.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (17):**
Medically confirmed literature case reporting confounding factors in medical history for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>Allergy</th>
<th>Dose</th>
<th>Dates</th>
<th>Diagnosis</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>49 Y/F</td>
<td></td>
<td>allergic</td>
<td>Dose 2</td>
<td>Spontaneous</td>
<td>IgA nephropathy, Haematuria, Proteinuria</td>
<td>The patient received Dose 2 of Comirnaty on 21 May 2021. On an unspecified date in June 21, the patient developed haematuria, proteinuria and IgA nephropathy. Vaccination history included: Comirnaty (1st dose) and DPT (dates unspecified). The patient also had a history of urinary occult blood 2+ 3 years before. She was hospitalized 6 months later, and a kidney biopsy confirmed the diagnosis of IgA nephropathy. The patient was treated with steroids.</td>
</tr>
<tr>
<td>12 Y/M</td>
<td></td>
<td>None</td>
<td>Dose 1</td>
<td>Literature spontaneous</td>
<td>IgA nephropathy 1 day Serious Recovering</td>
<td>The patient presented with new-onset gross haematuria, proteinuria, and acute kidney injury &lt; 24 h following the Dose 1 of Comirnaty. The patient had no family history of autoimmune diseases. A kidney biopsy was consistent with IgA nephropathy. Microscopic examination of the biopsy sections revealed 11 glomeruli in the submitted cores, all of which showed mild increase of mesangial cells and matrix; no thickening of capillary loops, segmental sclerosis, crescent formation or necrosis were seen. Many of the tubules showed red cell casts with mild tubular injury and flattening of epithelial cells. The immunofluorescence studies showed 5 glomeruli revealing granular mesangial deposits of IgA (+1) and C3 (+1), and absence of IgG, IgM, C4 and fibrinogen. Other labs: Serum Creatinine 1.77 mg/dl (normal range: 0.53-0.79 mg/dl), Serum Urea 61 mg/dl (normal range: 15-36 mg/dl).</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (18):**
Spontaneous medically confirmed case with limited information which is considered unassessable.
### Rapporteur assessment comment (19):

Medically confirmed case with limited information which is considered unassessable.

<table>
<thead>
<tr>
<th>18 Y/F</th>
<th></th>
<th></th>
<th>complement factor C3: 137 mg/dl, C4: 21 mg/dl, CRP 51.23 mg/dl (0.1-2.8). Urine analysis: proteinuria at 1.7 g/l, RBCs 1920/ul. The event improved after treatment with methylprednisolone.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose 3</td>
<td>Seasonal allergy</td>
<td>IgA nephropathy, Haematuria, Cystitis, Urinary occult blood, Protein urine, Pyrexia, Fatigue, Malaise</td>
<td>2 days after receiving Dose 3 of Comirnaty, the patient presented with proteinuria (2+) and haematuria (50–99, urinary occult blood 3+). Primary immunization was completed with Comirnaty 8 months earlier. A urine culture was positive for Escherichia coli and Klebsiella pneumoniae. Suspected IgA nephropathy.</td>
</tr>
<tr>
<td>Spontaneous</td>
<td>Bepotastine Besilate</td>
<td></td>
<td>The diagnosis of IgA nephropathy was not confirmed. No confirmatory diagnostics (including biopsy) provided. Concurrent urinary tract infection confounds the assessment.</td>
</tr>
</tbody>
</table>

### Rapporteur assessment comment (20):

Spontaneous case with no confirmed diagnosis of IgA nephropathy which is considered unassessable.

<table>
<thead>
<tr>
<th>24 Y/M</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose 2</td>
<td>Allergy to animal / Asthma / Colour blindness / Deafness unilateral / Dust allergy / Mite allergy</td>
<td>IgA nephropathy, Acute kidney injury, Pyrexia, Chills, Odynophagia, Haematuria, Pharyngitis streptococcal</td>
<td>The patient presented with fever, chills, odynophagia, haematuria and streptococcal pharyngitis approximately 2.5 months after receiving Dose 2 of Comirnaty. No information regarding first dose of covid vaccine. A kidney biopsy confirmed IgA nephropathy. Serum creatine 102 μmol/l.</td>
</tr>
<tr>
<td>Spontaneous</td>
<td>Not reported</td>
<td>79 days</td>
<td>Long latency of 79 days suggests a biologically implausible latency. Additionally, concurrent streptococcal pharyngitis is a potential and more likely etiology of nephropathy.</td>
</tr>
</tbody>
</table>

### Rapporteur assessment comment (21):

Spontaneous medically confirmed case reporting a TTO of 79 days and a co-infection for the cause of IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.
| 42 Y/F | Not reported | IgA nephropathy, Haematuria, Protein urine, Renal disorder | The same day as receiving the second dose of Comirnaty, the patient developed haematuria and proteinuria and was diagnosed with IgA nephropathy. The patient received first dose of Comirnaty on an unspecified date. A renal biopsy was consistent with IgA nephropathy (details unspecified). The patient was treated with methylprednisolone. | Medical history and timing between 2 doses of Comirnaty unknown. Further, concomitant influenza vaccine (date of administration unknown) confounds the assessment. |
| Dose 2 | Influenza vaccine | 2 days | Serious |
| Spontaneous | Not recovered | |

**Rapporteur assessment comment (22):**
Spontaneous medically confirmed case with limited information which is considered unassessable.

| 14 Y/F | Haematuria / Urine analysis abnormal | IgA Nephropathy | On the following day of the Dose 2 of Comirnaty, the patient developed high fever and frank haematuria. The patient received Dose 1 of Comirnaty on an unspecified date. On the 7th day, urine albumin/Cr ratio increased to 1.99 g/gCr. On the 12th day, the serum albumin level decreased to 3.6 g/dL (serum creatinine: 0.54 mg/dL); normal ranges not reported. Based on the renal biopsy results, a diagnosis of IgA nephropathy was made. Reportedly, it was considered that latent IgA nephropathy became prominent because there was a strong temporal relationship with vaccine. | History of haematuria and abnormal urine analysis however, no diagnostic details are provided. Besides, details of biopsy results and concomitant medications are not provided. |
| Dose 2 | Not reported | 1 day | Serious |
| Literature spontaneous | IgA Nephropathy | Unknown | |

**Rapporteur assessment comment (23):**
Medically confirmed case with limited information which is considered unassessable.
<table>
<thead>
<tr>
<th>Age</th>
<th>Gender</th>
<th>Dose</th>
<th>Reactivity</th>
<th>Diagnosis</th>
<th>Concomitant Conditions</th>
<th>Duration</th>
<th>Clinical Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>13 Y/F</td>
<td>Not reported</td>
<td>Not reported</td>
<td>IgA nephropathy, Cystitis, Pyrexia, Haematuria, Proteinuria</td>
<td>2 days</td>
<td>Serious Recovering</td>
<td>2 days after receiving Dose 3 of Comirnaty, the patient developed pyrexia, frank haematuria and proteinuria. Primary immunization completed with Comirnaty 7 months earlier. Cystitis was suspected. A blood test demonstrated that an inflammatory response was alleviated, however haematuria and proteinuria persisted and deformed erythrocytes were observed in the urine. Reportedly, the patient had undiagnosed IgA nephropathy, which was possibly &quot;exteriorized&quot; owing to pyrexia (Infection). Renal biopsy was not performed.</td>
<td>The reporter indicated suspected undiagnosed nephropathy possibly unmasked due to concurrent infection raises the likelihood of underlying nephropathy. Further, no confirmatory biopsy was performed.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment (24):**

Spontaneous confirmed literature case reporting pre-existing undiagnosed IgA nephropathy. Therefore the cause of IgA nephropathy in this case is considered unlikely related to Comirnaty exposure.

**MAH’s summary of the 43 cases**

A total of 7/43 cases reported IgA nephropathy/condition aggravated after dose 1 of Pfizer/BioNTech COVID-19 vaccine. Four of these cases reported a short latency of 1-2 days thus, making a biological association less likely. Besides, 2 of these 4 cases had additional confounders including concurrent infection and concomitant medications including mycophenolate mofetil and tacrolimus. The remaining 2/4 cases had limited information regarding medical history, concomitant medications and/or relevant investigations thus, precluding an adequate assessment. The remaining 3/7 cases were confounded by a long latency of 3 months, underlying autoimmune disease, suspected concurrent infection and chronic nephropathy.

Of the remaining 36 cases, 32 reported IgA nephropathy/condition aggravated after dose 2. Of these 32 cases, 23 reported 1 or more confounders like concurrent infection (e.g.: influenza), concomitant medications (enalapril, lisinopril), pre-existing haematuria, underlying autoimmune conditions, already aggravated renal dysfunction or poorly controlled IgA nephropathy, probable natural disease progression, or the biopsy was not confirmatory, long latency of more than 6 weeks and/or family history.

The remaining 9 cases had insufficient information regarding medical history, concomitant medications, baseline labs or relevant investigations thus, precluding an adequate assessment. Of the remaining 4 cases, 3 reported IgA nephropathy after dose 3. All these 3 cases lacked information regarding medical history, concomitant medications and/or relevant investigations thus, precluding an adequate assessment.
In the remaining case, dosing information was not provided. In this case, medical history and concomitant medications were not reported. Further, the clinical presentation raised the possibility that another nephropathy was present.

Biopsy at the time of event occurrence was reported in 28/43 cases, biopsy was not done in 4 cases while 11 cases did not have information regarding biopsy. Outcome was recovered/recovering in 20/43 cases, not recovered in 12 cases while outcome was not reported in 11 cases. The patients received treatment with steroids/immunosuppressants in 12/43 cases. No treatment was provided in 6 cases of which 5 were recovering/recovered. Treatment information was not provided in the 25 remaining cases.

**Rapporteur assessment comment:**

A total of 103 cases of IgA nephropathy were retrieved from MAH's safety database, 8 cases were excluded because no IgA nephropathy, unknown COVID-19 vaccine, or implausible latency, and 52 cases reported limited information precluding a meaningful assessment. Of the remaining 43 cases:

- 24 cases reported new onset of IgA nephropathy, of which the PRAC Rapporteur considered that in 13 cases confounding factors seem to provide a more likely explanation for the event and that causality with Comirnaty cannot be established in these cases and that 11 cases are unassessable for a meaningful causality assessment.

- 19 cases reported condition aggravated in pre-existing history of IgA nephropathy, of which the PRAC Rapporteur considered 15 cases unassessable and 4 cases reported confounding factors seem to provide a more likely explanation for the event and that causality with Comirnaty cannot be established in these cases (AER numbers [redacted] and [redacted]).

In conclusion, based on the information provided in the cases reporting new onset of IgA nephropathy or condition aggravated in pre-existing history of IgA nephropathy, there is no a causal association between Comirnaty exposure and occurrence of IgA nephropathy or exacerbation of IgA nephropathy.

**Observed versus expected analyses**

The MAH conducted unadjusted observed to expected ratio (O/E) analyses for the 101 Immunoglobulin A (IgA) nephropathy cases reported cumulatively through 18 June 2022 globally (table 5).

The overall expected case counts of IgA nephropathy were estimated using background incidence rates (IR) reported by a retrospective cohort study in population older than 15 years of age residing in a north-western region of Italy. Between January 1, 1970, and December 31, 1994, 1,926 cases of biopsy-proven primary glomerulonephritis were diagnosed, and IgA nephropathy was the most frequent type with an overall incidence rate 1.47 per 100,000 population per year and a predominance in males (males, 2.27 versus females, 0.67 per 100,000 population per year). These rates provided a low range of background rates for IgA nephropathy, and therefore, low expected case counts to conservatively estimate O/E results. 2 reviews reported overall incidence rate of IgA nephropathy ranged from 2.5 to 4.5 per 100,000 population per year with higher rates in Asia.
Table 5. Observed to Expected (O/E) Ratio for Spontaneously Reported Cases of IgA Nephropathy Through 18 June 2022

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Observed cases</th>
<th>Time at risk (PY)</th>
<th>Background rates per 10,000 PY</th>
<th>Expected cases</th>
<th>O/E ratio</th>
<th>95% CI LL</th>
<th>95% CI UL</th>
</tr>
</thead>
<tbody>
<tr>
<td>7-day risk window</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td>Males</td>
<td>12</td>
<td>9,810,452</td>
<td>2.27</td>
<td>204.54</td>
<td>0.059</td>
<td>0.030</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>22</td>
<td>10,160,733</td>
<td>0.67</td>
<td>58.08</td>
<td>0.323</td>
<td>0.203</td>
</tr>
<tr>
<td>Overall</td>
<td>Any dose</td>
<td>34</td>
<td>19,171,195</td>
<td>1.47</td>
<td>281.82</td>
<td>0.121</td>
<td>0.084</td>
</tr>
<tr>
<td></td>
<td>Dose 1</td>
<td>7</td>
<td>7,645,271</td>
<td>1.47</td>
<td>112.39</td>
<td>0.062</td>
<td>0.025</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>21</td>
<td>7,899,445</td>
<td>1.47</td>
<td>194.36</td>
<td>0.291</td>
<td>0.125</td>
</tr>
<tr>
<td></td>
<td>Additional/boost er dose(s)*</td>
<td>6</td>
<td>4,426,479</td>
<td>1.47</td>
<td>65.07</td>
<td>0.952</td>
<td>0.854</td>
</tr>
<tr>
<td>Global overall</td>
<td>77</td>
<td>42,516,664</td>
<td>1.47</td>
<td>624.99</td>
<td>0.123</td>
<td>0.097</td>
<td>0.154</td>
</tr>
<tr>
<td>14-day risk window</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td>Males</td>
<td>12</td>
<td>18,090,074</td>
<td>2.27</td>
<td>408.74</td>
<td>0.029</td>
<td>0.015</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>23</td>
<td>20,304,722</td>
<td>0.67</td>
<td>136.04</td>
<td>0.169</td>
<td>0.107</td>
</tr>
<tr>
<td>Overall</td>
<td>Any dose</td>
<td>35</td>
<td>38,310,796</td>
<td>1.47</td>
<td>563.17</td>
<td>0.062</td>
<td>0.043</td>
</tr>
<tr>
<td></td>
<td>Dose 1</td>
<td>7</td>
<td>15,284,506</td>
<td>1.47</td>
<td>224.68</td>
<td>0.031</td>
<td>0.013</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>22</td>
<td>14,192,990</td>
<td>1.47</td>
<td>208.64</td>
<td>0.105</td>
<td>0.066</td>
</tr>
<tr>
<td></td>
<td>Additional/boost er dose(s)*</td>
<td>6</td>
<td>8,833,580</td>
<td>1.47</td>
<td>129.83</td>
<td>0.040</td>
<td>0.017</td>
</tr>
<tr>
<td>Global overall</td>
<td>83</td>
<td>84,874,815</td>
<td>1.47</td>
<td>1247.66</td>
<td>0.067</td>
<td>0.053</td>
<td>0.082</td>
</tr>
<tr>
<td>21-day risk window</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>US/EEA</td>
<td>Males</td>
<td>12</td>
<td>26,983,860</td>
<td>2.27</td>
<td>612.52</td>
<td>0.029</td>
<td>0.019</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>25</td>
<td>30,427,706</td>
<td>0.67</td>
<td>203.87</td>
<td>0.123</td>
<td>0.079</td>
</tr>
<tr>
<td>Overall</td>
<td>Any dose</td>
<td>37</td>
<td>57,410,765</td>
<td>1.47</td>
<td>843.94</td>
<td>0.044</td>
<td>0.031</td>
</tr>
<tr>
<td></td>
<td>Dose 1</td>
<td>8</td>
<td>22,917,286</td>
<td>1.47</td>
<td>336.88</td>
<td>0.024</td>
<td>0.010</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>22</td>
<td>21,280,003</td>
<td>1.47</td>
<td>312.82</td>
<td>0.078</td>
<td>0.044</td>
</tr>
<tr>
<td></td>
<td>Additional/boost er dose(s)*</td>
<td>7</td>
<td>13,213,476</td>
<td>1.47</td>
<td>194.24</td>
<td>0.036</td>
<td>0.014</td>
</tr>
<tr>
<td>Global overall</td>
<td>86</td>
<td>127,075,389</td>
<td>1.47</td>
<td>1868.01</td>
<td>0.046</td>
<td>0.037</td>
<td>0.057</td>
</tr>
</tbody>
</table>

CI = confidence interval; EEA=European Economic Area; LL = lower limit; PY = person-years; UL = upper limit; US=United State.
*Additional/booster dose(s) denote first booster (or additional) or second booster dose from data sources that supply vaccine doses administered for this analysis.

Rapporteur assessment comment:
All O/E ratios for were well below 1.

MAH's conclusion
The 103 cases of IgAN retrieved from the global safety database is a relatively small number of cases when considering the number of doses (estimated 1-2 billion) administered worldwide since authorization and the number of AE reports (>1.6 million) for BNT162b2 in the safety database. Concentration on cases that appeared to have sufficient information still revealed some with scant detail, lack of support of IgAN diagnosis, or plausible alternative explanations for IgAN. Cases without confounders have temporality as the main link to vaccination. Trends observed were that most cases occurred following dose 2 and many cases were from Japan. The higher number of cases describing females versus males and the fact that most cases were reported in adults 18-50 years of age is consistent with the overall dataset of spontaneous AE reports for the vaccine. It is notable that Japan was the highest reporting country (35% of cases of IgAN) because the Japanese healthcare system is recognized as being proactive in renal disorder screening.

No adverse events of IgA nephropathy were reported in the pivotal clinical trials for BNT162b2 and there were a very small number of participants with a medical history of IgAN.

PRAC PSUR assessment report
EMA/PRAC/849/2023 Page 91/272
There has been no statistical signal of disproportionate reporting in the MAH safety database for the PT IgA nephropathy.

The published literature mostly comprises clinical case reports and a systematic review qualitatively analysing 14 such cases. Although small, a prospective study of COVID-19 vaccination in patients with existing IgAN suggested no impact of vaccination on the clinical course of the disease.

O/E ratios were below one suggesting that the number of observed cases of IgA nephropathy is not higher than expected in the absence of Pfizer/BioNTech COVID-19 vaccines overall and within the queried strata.

Overall, a causal association of Comirnaty with IgA nephropathy cannot not be concluded based on this review of available clinical, literature and post-marketing data. The MAH will continue to monitor the subject using routine Pharmacovigilance.

Rapporteur assessment comment:

Literature

Of the 49 results retrieved from the literature search through 30 Jun 2022, no new important information could be identified concerning IgA nephropathy.

Clinical trial data

There were no reports of IgA nephropathy in the clinical trials.

Post-marketing

A total of 95 valid cases of IgA nephropathy were retrieved from MAH’s safety database, of which 24 cases reported new onset of IgA nephropathy (13 cases reported confounding factors seem to provide a more likely explanation for the event and that causality with Comirnaty cannot be established in these cases and 11 cases are considered unassessable) and 19 cases reported condition aggravated in pre-existing history of IgA nephropathy (15 cases are considered unassessable and 4 cases reported confounding factors seem to provide a more likely explanation for the event and that causality with Comirnaty cannot be established in these cases). The remaining 52 cases reported limited information preduding a meaningful assessment.

In conclusion, based on the information provided in the cases reporting new onset of IgA nephropathy or condition aggravated in pre-existing history of IgA nephropathy, there is not a causal association between Comirnaty exposure and occurrence of IgA nephropathy or exacerbation of IgA nephropathy.

Observed versus expected analyses

All O/E ratios for were well below 1.

Overall, MAH’s conclusion is endorsed that based on provided data no causal association of Comirnaty with IgA nephropathy can be concluded. No new important information could be identified concerning IgA nephropathy. The MAH should closely monitor any new cases, patterns, or trends of reporting IgA nephropathy through routine pharmacovigilance.

Issue solved
2.2.1. Post-approval regulatory requests

2.2.1.1. Hearing loss

Response to the PRAC request 2 from the 14th (3rd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.13):

The MAH is requested to perform a cumulative review on the association between sudden sensorineural hearing loss and Comirnaty exposure, including a review of the relevant new literature published after the reporting period of the 14th SSR, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

MAH's response (Appendix 6A.3. of the PSUR):

Reference is also made to the 08 Jun 2022 request from Health Canada for a cumulative review of cases of tinnitus and hearing loss and provision of an observed to expected analysis.

The MAH provided a cumulative review of hearing loss and tinnitus.

Literature

A cumulative search of literature was conducted through 18 June 2022, retrieved four relevant literature articles:


This cross-sectional study and case series involved an up-to-date population-based analysis of 555 incident reports of probable Sensorineural Hearing Loss (SSNHL) in the US Centers for Disease Control and Prevention Vaccine Adverse Events Reporting System (VAERS) over the first 7 months of the US vaccination campaign (December 14, 2020, through July 16, 2021). In addition, data from a multi-institutional retrospective case series of 21 patients who developed SSNHL after COVID-19 vaccination were analyzed. The study included all adults experiencing SSNHL within 3 weeks of COVID-19 vaccination who submitted reports to VAERS and consecutive adult patients presenting to 2 tertiary care centers and 1 community practice in the US who were diagnosed with SSNHL within 3 weeks of COVID-19 vaccination. Results: A total of 555 incident reports in VAERS (mean patient age, 54 years [range, 15-93 years]; 305 women [55.0%]; data on race and ethnicity not available in VAERS) met the definition of probable SSNHL (mean time to onset, 6 days [range, 0-21 days]) over the period investigated, representing an annualized incidence estimate of 0.6 to 28.0 cases of SSNHL per 100 000 people per year. The rate of incident reports of SSNHL was similar across all 3 vaccine manufacturers (0.16 cases per 100 000 doses for both Pfizer-BioNTech and Moderna vaccines, and 0.22 cases per 100 000 doses for Janssen/Johnson & Johnson vaccine). The case series included 21 patients (mean age, 61 years [range, 23-92 years]; 13 women [61.9%]) with SSNHL, with a mean time to onset of 6 days (range, 0-15 days). Patients were heterogeneous with respect to clinical and demographic characteristics. Preexisting autoimmune disease was present in 6 patients (28.6%). Of the 14 patients with posttreatment audiometric data, 8 (57.1%) experienced improvement after receiving treatment. One patient experienced SSNHL 14 days after receiving each dose of the Pfizer-BioNTech vaccine. Study Conclusions: In this cross-sectional study, findings from an updated analysis of VAERS data and a case series of patients who experienced SSNHL after COVID-19 vaccination did not suggest an association
between COVID-19 vaccination or an increased incidence of hearing loss compared with the expected incidence in the general population (the annual incidence of idiopathic SSNHL was estimated to be 11 to 77 cases per 100 000 people per year).

MAH comment: The MAH agrees with the study authors that the reporting rate of hearing loss compared with the expected incidence in the general population suggests no association with COVID-19 vaccination. Strengths of the study include population-based setting with large sample size. One of the major limitations in the study is a lack of concurrent comparison group. Formelster et al compared the incidence of SSNHL after vaccination to a historical incidence of idiopathic SSNHL that was reported during 2006-2007 in the US, therefore changes in the demographics and in the diagnostic criteria of SSNHL and the fact that the historical rate was estimated from claims data (and not self-reported data) may influence the comparison.


Yanir performed a retrospective, observational study on data from the Clalit Health Services (CHS), which provides inclusive healthcare for more than half of the Israeli population. This database, designed for administrative and clinical management is available for clinical studies. They identified cases of SSNHL by using the (International Classification of Diseases, Ninth Revision) ICD-9 diagnosis codes for sensorineural hearing loss (SNHL) and included only those patients with concurrent prednisone treatment (defined as purchasing a prescription from start date of diagnosis of SNHL up to 30 days after). Observed cases of SSNHL appearing after COVID-19 vaccination were compared with the expected cases of SSNHL based on experience of the CHS population in 2018 and 2019 (before the COVID-19 pandemic and vaccine introduction in Israel). CHS members aged > 16 years who received the first Pfizer-BioNTech vaccine dose from 20 December 2020 (the start date of mass COVID-19 vaccination in Israel) until 30 April 2021 were included. Those not diagnosed with SSNHL after dose 1, who received dose 2 before 30 April 2021 made the population for estimation of the standardized incidence ratios (SIR) after second vaccine dose. Retrospective follow-up of cases through 31 May 2021 was performed. Observed cases occurring within 21 days of the first and second Pfizer-BioNTech vaccine dose were compared with expected cases estimated from the historic comparator group. During the data collection period, 2.6 million CHS members received first Pfizer-BioNTech vaccine dose; of these 2.4 million received the second dose before 30 April 2021. SSNHL was detected within 21 days in 91 patients after the first vaccine dose and in 79 patients after the second vaccine dose. When using the 2018 comparator group for reference, the age- and sex-weighted SIRs were 1.35 (95% CI, 1.09-1.65) for the first vaccine dose and 1.23 (95% CI, 0.98-1.53) for the second dose. Higher SIRs were observed in female patients aged 16 to 44 years (SIR, 1.92; 95% CI, 0.98-3.43) and patients 65 years old (SIR, 1.68; 95% CI, 1.15–2.37) after the first dose and in male patients 16 to 44 years (SIR, 2.45; 95% CI, 1.36–4.07) after the second dose. The authors concluded that the Pfizer-BioNTech vaccine might be associated with an increased risk of SSNHL; however, the effect size is very small and the benefit of COVID-19 vaccines outweighs its potential association with SSNHL.

MAH comment: Strengths of the study include population-based setting with large sample size. One of the limitations in the study is a lack of concurrent comparison group. The study utilized recent historical data (2018 and 2019) before the COVID-19 pandemic; therefore, no major temporal differences are expected. However, confounding and detection bias could have been introduced. Although Yanir et al have controlled for age and sex through standardization, age
was controlled in wide age ranges (i.e., 16-44, 45-64, and ≥65 years). Given that age is a strong risk factor for SSNHL and vaccinated people were older (and may be sicker), especially during the early phase vaccination program, there could be residual confounding by age which contributed to the increased occurrence of SSNHL. There could also be confounding by other unmeasured factors (e.g., cardiovascular and coagulation disorders, which are risk factors for SSNHL) that might differ between the vaccinated group and the general population. Last but not least, the observed association could be susceptible to detection bias during the COVID-19 pandemic period when patients may be more likely to be alert to any adverse events after vaccination and seek medical care for hearing loss (or other adverse events) after vaccination and thus have SSNHL diagnosed. The weak association in the study could be an artifact of confounding and detection bias.


The authors conducted a retrospective chart review in recently vaccinated patients over a 30-day time frame in a tertiary otology ambulatory practice. Results: Within the same 30-day time period in 2019, 2020, and 2021, 1.6, 2.4, and 3.8% respectively, of all office visits were for patients with the diagnosis of new onset idiopathic sensorineural hearing loss (SSNHL) without other underlying otologic diagnoses. In this time frame in 2021, 30 patients out of the 1,325 clinical visits had new or significantly exacerbated otologic symptoms that began shortly after COVID-19 vaccination. Specifically, 18 patients received Moderna and 12 patients received Pfizer vaccine. Their mean age was 60.9±13.8 years old; 11 were women and 19 men. The mean onset of symptoms was 10.18 ± 9 days post-vaccination. Symptoms included 25 patients (83.3%) with hearing loss, 15 (50%) with tinnitus, eight (26.7%) with dizziness, and five (16.7%) with vertigo. Eleven patients had previous otologic diagnoses, including six patients with Menière's disease, two with autoimmune inner ear disease (AIED), and three having both. Authors' Conclusions: There are no definite correlations to the COVID-19 pandemic or vaccination and new or worsened otologic symptoms. Vaccinated patients with new or exacerbated otologic symptoms should be promptly referred for evaluation.

MAH comment: The MAH agrees with the authors that no definite correlation between vaccination and new or worsened otologic symptoms could be concluded. The sample size of this retrospective chart review is very small and the clinical data regarding vaccination is limited to the intake questionnaire and details that were documented during a clinic visit.


This non peer-reviewed article in preprint investigates whether U.S. Centers for Disease Control and Prevention Vaccine Adverse Events Reporting System (VAERS) data suggest an association between vertigo, tinnitus, hearing loss, Bell's palsy and the COVID-19 vaccines administered in the United States. VAERS reports of these events were compared with published rates for the general population. Of note, the author Ramsi A. Woodcock suffers from otologic symptoms that he believes to have been caused by COVID-19 vaccination; he is the patient of the 2nd author, Loren J. Bartels. Results: The COVID-19 vaccines were associated with statistically significant increases in the reporting of vertigo, tinnitus, hearing loss, and Bell's palsy (1877, 50, 12, and 14 cases per 100,000, respectively). In relation to the mRNA-1273 or BNT162b2 vaccines, the Ad26.COV2.S vaccine was associated with a statistically significant excess incidence of vertigo, tinnitus, and hearing loss of at least 723, 57, and 55
cases per 100,000, respectively. Authors' Conclusion: These results suggest an association between the COVID-19 vaccines and vertigo, tinnitus, hearing loss, and Bell's palsy. They also suggest that the association is relatively strong for the Ad26.COV2.S vaccine.

MAH comment: This article is not weighed heavily in consideration of this signal for these reasons: it is a pre-published and non-peer reviewed article, the author states a clear bias (experienced an AE they believe was caused by COVID-19 vaccination) and the author, while using a well-known database, does not explain their methodology for obtaining or assessing the VAERS data or the background rates. Their use of the term "incidence" as opposed to "reporting rates" with respect to the VAERS data and focus on the initial part of the observation period as a way to account for underreporting is questionable. Further, all COVID-19 vaccines were included in the assessment.

MAH's overall conclusion: The literature articles retrieved in the search do not present any new significant safety information or conclude a causal relationship.

Rapporteur assessment comment:

The study of Formeister et al. and the study of Yanir et al. were already assessed in the 14th (3rd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.13):

- Formeister et al. did not suggest an association between COVID-19 vaccination and an increased incidence of SSNHL compared with the expected incidence in the general population, based on the cross-sectional analysis of VAERS data and review of SSNHL cases.

- Yanir et al. concluded that the Pfizer-BioNTech vaccine might be associated with an increased risk of SSNHL; however, the effect size is very small. However, the strength of the study includes population-based setting with a large sample size of which the results could be considered a signal for a potential association between SSNHL and Comirnaty exposure.

As a result the MAH was requested to perform a cumulative review on the association between sudden sensorineural hearing loss and Comirnaty exposure.

In the two additional retrieved articles:

- Wichova et al. stated that no definite correlation between vaccination and new or worsened otologic symptoms could be concluded.

- The study of Woodcock et al., a publication not-peer reviewed and published on medRxiv, is not considered relevant due to missing information regarding used methodology for the comparison of the rates of vertigo, tinnitus, hearing loss, Bell's palsy between the VAERS database (after COVID-19 vaccination) and the published rates for the general population.

No new important safety information could be identified from literature concerning hearing loss and tinnitus.

Safety database review

The Pfizer safety database was searched cumulatively through 18 June 2022 for all BNT162b2; BNT162b2S01 cases reported using MedDRA v 25.0 PTs of Conductive deafness, Deafness, Deafness bilateral, Deafness neurosensory, Deafness occupational, Deafness permanent, Deafness transitory, Deafness unilateral, Hypoacusis, Mixed deafness, Neurosensory hypoacusis, Sudden hearing loss, Tinnitus, and/or Tinnitus retraining therapy.
Result

Cases with both Hearing loss and Tinnitus events

A total of 2009 cases (with a total of 10,592 events) reporting both tinnitus and hearing loss events were retrieved.

The majority of cases (1998) were spontaneous; 1601 cases were serious, 408 were nonserious. There were 1177 females, 795 males, and sex was not reported in 37 cases. When provided, the ages ranged as shown in Table 2 below. The mean and median ages were 49 years (n=1921).

<table>
<thead>
<tr>
<th>Table 2. Reported Age in 2009 Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age Range</td>
</tr>
<tr>
<td>---------------------------------</td>
</tr>
<tr>
<td>Less than or equal to 17 years</td>
</tr>
<tr>
<td>18 - 30 years</td>
</tr>
<tr>
<td>31 - 50 years</td>
</tr>
<tr>
<td>51 - 64 years</td>
</tr>
<tr>
<td>65 - 74 years</td>
</tr>
<tr>
<td>Greater than or equal to 75 years</td>
</tr>
<tr>
<td>Unknown</td>
</tr>
</tbody>
</table>

Further analysis was concentrated on 1491 cases reporting relevant events with time to onset (latency) of 0 to 21 days post vaccination.

Out of these 1491 cases, 431 cases reported a pre-existing medical condition that represents a reasonable alternative cause of the relevant events. Examples of the conditions include: deafness, tinnitus, COVID-19, autoimmune disorders, anxiety and panic disorders, balance and vestibular disorders, diabetes, chronic sinusitis, and auditory conditions. Of these 431 cases, 166 reported pre-existing deafness and/or tinnitus, 22 of which reported an aggravation or exacerbation/worsening of the condition after vaccination.

An additional 72 cases reported use of co-suspect or concomitant medications that represent a potential alternative cause of the events. Examples of the medications include: Angiotensin Converting Enzyme (ACE) inhibitors, Angiotensin Receptor Blockers (ARBs), beta blockers, tricyclic antidepressants, aspirin, Non-Steroidal Anti-Inflammatory Drugs (NSAIDs), benzodiazepines, loop diuretics, isotretinoin, immunosuppressants and other vaccines.

Out of the remaining 988 cases in this dataset, 27 cases lacked details on medical history, concomitant medications, outcome, and dose number precluding a thorough assessment.

Of the remaining 961 cases, further analysis was concentrated on the 212 reports that were healthcare professional confirmed (HCP) cases. Of these 212 cases (459 serious relevant events), 104 cases co-reported PTs unlikely to be caused by vaccination that suggested the possibility of concomitant conditions (e.g., ear infection, vestibular disorder, vertigo, syncope, tympanic membrane disorder, COVID-19, nasal congestion, middle ear effusion, ear hemorrhage, ear canal stenosis, inner ear infarction, etc.) contributing to the cause of the hearing loss/tinnitus events.

In the remaining 108 HCP cases (159 relevant events), when dose sequence was provided, it was Dose 1 for 60 relevant events, Dose 2 for 63 events and Dose 3 for 11 events (there were no events reported post Dose 4). The outcome of the relevant events at the time of reporting was resolved/resolving for 54 events; not resolved for 84 events; resolved with sequelae for 6 events; unknown for 15 events. The most frequently reported latency was Day 1 (for 24 events) followed by Day 0 (for 17 events), Day 5 (for 16 events), and Day 2 (for 15 events).
Medically confirmed cases with no co-reported events

Out of these 108 cases, 71 cases reported relevant events of hearing loss and/or tinnitus only (no other events were reported in these cases).

In 50 of these 71 cases, no information on diagnostic procedures or tests supporting the diagnosis were provided, therefore the causality assessment for these 50 cases is Unassessable per the WHO-UMC causality assessment categories8.

In 21 of the 71 cases, diagnostic tests and/or procedures such as Acoustic stimulation test, Audiogram, Ear, nose and throat examination, and Otic examination were reported. Of note, none of these cases reported confirmatory Computerized Tomography (CT) scan and/or Magnetic Resonance Imaging (MRI) results. Twenty (20) out of 21 cases occurred in adults 24 to 67 years old. One of the cases reported a patient with pre-existing alopecia areata and psoriasis. Two other cases reported pre-existing medical history and concomitant medications for hypothyroidism and benign prostatic hyperplasia, respectively, and another 17 remaining adult cases did not report any additional details. The last of these cases describes a possible rechallenge in a child:

- A 14-year-old female with no relevant medical or family history and no concomitant medications or vaccines experienced tinnitus and deafness (the patient had difficulty in talking with family members) 5 days following dose 1 of Comirnaty. There were no complaints of giddiness. The following day, the patient's tinnitus was slightly alleviated compared to the previous day, however she sought medical attention and her audiometry test showed worsening compared to testing in May (reason for pre-vaccination hearing test was not explained). A head MRI approximately 1 month later showed "no significant change." The reason for a previous MRI of the head was not explained. The patient had dose 2 of Comirnaty and reported that her tinnitus was aggravated, she underwent another head MRI (no change) and further audiometry testing. The audiometry tests show progressive worsening of hearing loss from May to 10 Dec 2021.

<table>
<thead>
<tr>
<th>Date</th>
<th>Test or Vaccine</th>
</tr>
</thead>
<tbody>
<tr>
<td>May 2021</td>
<td>Pure tone audiometry: 21.3 dB R, 17.5 dB L</td>
</tr>
<tr>
<td>26 Aug 2021</td>
<td>Dose 1 vaccine</td>
</tr>
<tr>
<td>03 Sep 2021</td>
<td>Pure tone audiometry: 26.3 dB R, 27.5 dB L</td>
</tr>
<tr>
<td>03 Oct 2021</td>
<td>Head MRI &quot;no significant change&quot;</td>
</tr>
<tr>
<td>18 Sep 2021</td>
<td>Dose 2 vaccine</td>
</tr>
<tr>
<td>03 Oct 2021</td>
<td>Head MRI &quot;no significant change&quot;</td>
</tr>
<tr>
<td>12 Nov 2021</td>
<td>Pure tone audiometry: 37.5 dB R, 42.5 dB L</td>
</tr>
<tr>
<td>10 Dec 2021</td>
<td>Pure tone audiometry: 45.0 dB R, 36.3 dB L</td>
</tr>
</tbody>
</table>

The causality assessment of these 21 cases is Possible.

Medically confirmed cases with co-reported adverse events

Out of the remaining 108 cases, 37 had co-reported events along with the hearing and tinnitus events. In 25 of these 37 cases, either no diagnostic procedures or tests supporting the diagnosis were reported, test results were reported as normal, or test results were not available at the time of reporting. The causality assessment for these 25 cases is Unassessable per the WHO-UMC causality assessment categories.

In 12 of the 37 cases diagnostic procedures/tests such as Acoustic Stimulation, Audiogram, ENT exam, Investigation, Otic exam, and/or Weber tuning were reported, supporting the diagnosis. Upon detailed review of the narratives of these 12 cases, the following factors were noted in 6 of the cases making an individual causal association Unlikely:
• One report was in a 54-year-old woman and was associated with left-sided facial numbness (head CT normal),

• One report in a 34-year-old man was associated with actual swelling of the ear on the affected side, suggesting the possibility of a contributing outer-ear etiology,

• Two cases reported a pre-existing medical history of malignancy introducing the possibility of exposure to ototoxic chemotherapeutics,

• One report in a 34-year-old woman with 2 years of seasonal allergies and intranasal steroid use introducing the possibility of eustachian tube dysfunction,

• One report in a 15-year-old girl who underwent a tympanoscopy and was found to have a probable polylymphatic fistula.

The remaining 6 reports, while lacking details, did not provide alternative explanations for the tinnitus or deafness making a causal association possible.

Rapporteur Assessment Comment:

Through 18 Jun 2022, there were a total of 2,009 cases reporting hearing loss and tinnitus. Of these 2,009 cases a TTO of >21 days was reported in 518 cases, pre-existing medical condition in 431 cases, concomitant medications in 72 cases and lack of information in 27 cases. Of the remaining 961 cases further analysis was concentrated on the 212 cases (22%) that were HCP confirmed:

- 104 cases co-reported PTs unlikely to be caused by vaccination.

- 71 cases had hearing loss and/or tinnitus only, of which 50 cases were considered unassessable and 21 cases were considered possibly related to Comirnaty exposure (including a possible rechallenge in a 14 year old female).

- 37 cases had co-reported events, of which 25 cases were considered unassessable, 6 cases were considered unlikely related to Comirnaty exposure, and 6 cases were considered possible related to Comirnaty exposure.

Cases of Tinnitus Events Only

A total of 13943 cases (with a total of 66922 events) reporting tinnitus were retrieved.

Similar to the dataset described above, the majority of cases (13781) were spontaneous; 4581 cases were serious, 9362 were nonserious. There were 8606 females, 4581 males, and sex was not reported in 356 cases. When provided, the ages ranged as shown in Table 4 below. The mean and median ages were 47.7 and 48 years, respectively (n=12756).

<table>
<thead>
<tr>
<th>Age Range</th>
<th>Number of Cases</th>
<th>% of Total Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than or equal to 17 years</td>
<td>162</td>
<td>1.20%</td>
</tr>
<tr>
<td>18 - 30 years</td>
<td>1423</td>
<td>10.20%</td>
</tr>
<tr>
<td>31 - 50 years</td>
<td>5712</td>
<td>41.00%</td>
</tr>
<tr>
<td>51 - 64 years</td>
<td>3783</td>
<td>27.10%</td>
</tr>
<tr>
<td>65 - 74 years</td>
<td>1335</td>
<td>9.60%</td>
</tr>
<tr>
<td>Greater than or equal to 75 years</td>
<td>399</td>
<td>2.90%</td>
</tr>
<tr>
<td>Unknown</td>
<td>1129</td>
<td>8.10%</td>
</tr>
</tbody>
</table>

Further analysis was concentrated on 10063 cases reporting tinnitus with latency of 0 to 21 days post vaccination,

Out of these 10063 cases, 2261 cases reported a pre-existing medical condition suggesting a reasonable alternative explanation for tinnitus. Examples of the conditions included deafness, tinnitus,
COVID-19, autoimmune disorders, and chronic sinusitis. Of these 2261 cases, 738 reported pre-existing tinnitus; 128 of these 738 cases reported "aggravation, exacerbation, or worsening of the pre-existing tinnitus. An additional 399 cases reported use of co-suspect/concomitant medication representing a reasonable alternative cause of the relevant event. Examples included ACEIs, ARBs, tricyclic antidepressants, aspirin, NSAIDs, benzodiazepines, loop diuretics, immunosuppressants, and other vaccines.

Out of the remaining 7403 of the 10063 cases, further analysis was concentrated on 295 HCP-reported cases (containing 295 serious tinnitus events). In these 295 cases, when dose sequence was provided, it was Dose 1 for 151 tinnitus events, Dose 2 for 85 events, Dose 3 for 26 events, and Dose 4 for 1 event. Outcome of tinnitus at the time of reporting was resolved/resolving for 90 events; not resolved for 165 events; resolved with sequelae for 6 events; and unknown for 34 events. The most frequently reported latency was Days 0 and 1 for 81 events each followed by Day 2 for 34 events and Day 3 for 22 events.

Medically confirmed cases with no co-reported events

Out of 7403 cases, 295 were HCP confirmed. Of these 295 cases, 55 cases reported tinnitus only (no other events were reported in these cases). In 53 of these 55 cases no information on diagnostic procedures or tests supporting the diagnosis was provided; therefore, causality assessment for these 53 cases is Unassessable.

Of the remaining 2 cases, a 53-year-old man with an unreported medical history had a Neurology visit and negative MRI (with and without contrast), and a 30-year-old woman with an unreported medical history was seen by a specialist who found no abnormalities on clinical examination and diagnosed probable vestibular neuritis (unilateral). The causality assessment in these 2 cases is Unlikely.

Medically confirmed cases with co-reported adverse events

Of these 295 cases, 240 cases reported tinnitus and other events. Of these 240 cases, 159 co-reported PTs suggesting the possibility of other conditions (e.g., syncope, vertigo, anaphylactic reaction, hypertension, gait disturbance, vestibular disorder, amaurosis, autoimmune disorder and panic attack)

Out of the remaining 81 cases, 79 did not provide information on diagnostic procedures or tests supporting the diagnosis, therefore causality assessment for these 79 cases is Unassessable. Two remaining cases described:

- A 62-year-old male (medical history and concomitant medications were not reported) experienced malaise and pyrexia on the same day of administration of Dose 2 of the vaccine (these symptoms disappeared 3 days later) and bilateral amaurosis and tinnitus on Day 3 post vaccination. The Audiometry test suggested mild to moderate grade high frequency sensorineural hearing loss (this PT was not captured as AE); case is assessed as Possible.

- A 43-year-old male physician who complained of squeaks in the left ear following dose 2 without ENT abnormalities and with an unspecified hearing test that confirmed the "abnormalities about squeaks;" case is assessed as Possible.

Rapporteur assessment comment:

Through 18 Jun 2022, there were a total of 13,943 cases reporting tinnitus. Of these 13,943 cases a TTO of >21 days was reported in 3,880 cases, pre-existing medical condition in 2,261 cases, and concomitant medications in 399 cases. Of the remaining 7,403 cases further analysis was concentrated on the 295 cases (4%) that were HCP confirmed:

- 55 cases had tinnitus only, of which 53 cases were considered unassessable and 2 cases were considered unlikely related to Comirnaty.
- 240 cases had co-reported events, of which 159 cases co-reported PTs suggesting the possibility of other conditions causing tinnitus, 79 cases were considered unassessable, and 2 cases were considered possible related to Comirnaty exposure.

**Cases of Hearing loss events only**

A total of 3177 cases (with a total of 15668 events) were retrieved.

Similar to the datasets described above, the majority of cases (3117) were spontaneously reported; 2542 cases were serious, 635 were nonserious. There were 1922 females, 1147 males, and sex was not reported in 108 cases. When provided, the ages ranged as shown in Table 7 below. The mean and median ages were 50.2 and 50 years, respectively (n=2915).

<table>
<thead>
<tr>
<th>Age Range</th>
<th>Number of Cases</th>
<th>% of Total Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than or equal to 17 years</td>
<td>97</td>
<td>3.10%</td>
</tr>
<tr>
<td>18-30 years</td>
<td>408</td>
<td>12.80%</td>
</tr>
<tr>
<td>31-50 years</td>
<td>980</td>
<td>30.80%</td>
</tr>
<tr>
<td>51-64 years</td>
<td>720</td>
<td>22.70%</td>
</tr>
<tr>
<td>65-74 years</td>
<td>363</td>
<td>11.40%</td>
</tr>
<tr>
<td>Greater than or equal to 75 years</td>
<td>355</td>
<td>11.20%</td>
</tr>
<tr>
<td>Unknown</td>
<td>254</td>
<td>8.00%</td>
</tr>
</tbody>
</table>

Further analysis was concentrated on 2039 cases reporting hearing loss events with a latency of 0 to 21 days post vaccination. Out of these 2039 cases, 433 cases reported pre-existing medical conditions (e.g., deafness, COVID-19, anxiety, hearing aid use, autoimmune disorders, diabetes, Meniere's disease, acoustic neuroma, cerebrovascular disorders) representing potential alternative causes of the events; 76 of the 433 cases reported pre-existing hearing loss and 13 of these 76 cases reported "aggravation, exacerbation, or worsening of" the pre-existing relevant condition. Additional 88 of the 2039 cases reported the use of co-suspect/concomitant medications (e.g., ACE-I, ARB, beta blockers, tricyclic antidepressants, loop diuretics, immunosuppressants, and other vaccines) potentially accounting for the hearing loss events.

Out of the remaining 1518 of the 2039 cases with a latency of 0 to 21 days, further analysis was concentrated on 352 HCP-reported cases (containing 387 serious hearing loss events). In these 352 cases, when dose sequence was provided, it was Dose 1 for 175 relevant events, Dose 2 for 131 events, Dose 3 for 33 events, and Dose 4 for 2 events. The outcome of the relevant events at the time of reporting was resolved/resolving for 148 events; not resolved for 143 events; resolved with sequelae for 21 events; and outcome was unknown for 77 events. The most frequently reported latencies were Day 1 for 96 events, Day 0 for 66 events followed by Day 2 for 52 events and Day 3 for 24 events.

Medically confirmed cases with no co-reported events

Of the 352 HCP cases, 140 reported events of hearing loss only (no other events were reported in these cases). Out of these 140 cases, 116 cases did not report Information on diagnostic procedures or tests to support the diagnoses; therefore, the causality assessment of these 116 cases is Unassessable.

Twenty-four (24) out of the 140 cases reported diagnostic tests or procedures such as acoustic stimulation test, audiogram, ENT examination, Investigation, MRI, otoacoustic emissions test, and otoscopy. All 24 cases reported adults, with 5 being patients 74 to 90 years old. Of these 5 reports of elderly patients, 1 reported a pre-existing hearing disability making the case Unassessable while the remaining 4 had limited information reported including any information that posed a possible
alternative cause of the hearing loss (apart from age); therefore, the causality assessment in these 4 cases reporting elderly patients is Possible.

Of the remaining 19 cases, 6 of the narratives described pre-existing medical histories confounding causality assessment (allergic rhinitis, alcoholism and hypersensitivity, otitis externa, traumatic deafness, and throat cancer) leading to their assessment of Unlikely.

The last 13 out of 24 cases did not report any additional details (e.g., medical history, concomitant medications, additional tests) and based on the lack of information on alternative contributions, the causality assessment in these 13 cases is Unassessable.

Medically confirmed cases with co-reported adverse events

Of the 352 HCP cases, 212 reported hearing loss and other events; 146 of these 112 cases co-reported PTs suggesting the possibility of an alternative explanation for the hearing loss (e.g., ear infection, meningitis, cerebellar condition, syncope, loss of consciousness, chronic rhinitis/sinusitis, Meniere’s disease, etc.) contributing to hearing loss and making causality assessment Unlikely.

Out of the remaining 66 cases, in 63 cases the diagnostic procedures/tests were not reported, results of these tests were reported as normal, or results were not available at the time of reporting, therefore the causality assessment for these 63 cases is Unassessable. The last 3 cases reported diagnostic procedures/tests such as acoustic stimulation and audiogram. One case described ear edema (assessment is Unlikely) and assessment for the other 2 cases was Possible.

Rapporteur assessment comment:

Through 18 Jun 2022, there were a total of 3,177 cases reporting hearing loss. Of these 3,177 cases a TTO of >21 days was reported in 1,183 cases, pre-existing medical condition in 433 cases, and concomitant medications in 88 cases. Of the remaining 1,518 cases further analysis was concentrated on the 352 cases (23%) that were HCP confirmed:

- 140 cases had hearing loss only, of which 130 cases were considered unassessable. 6 cases were considered unlikely related to Comirnaty, and 4 cases were considered possible related to Comirnaty exposure.

- 212 cases had co-reported events, of which 146 cases co-reported PTs suggesting the possibility of other conditions causing hearing loss, 63 cases were considered unassessable, 1 case was considered unlikely related to Comirnaty, and 2 cases were considered possibly related to Comirnaty exposure.

Observed to expected analyses

Hearing loss

The MAH conducted unadjusted observed to expected ratio (O/E) analyses for 5805 hearing loss cases reported cumulatively through 18 June 2022 globally (Table 9).

The overall expected case counts of hearing loss were estimated using background incidence rates (IR) reported by a population-based cross-sectional study in the US IMS Lifelink Health Plan Claims Database, as requested by the Pharmacovigilance Risk Assessment Committee (PRAC). This study included on average 66,594 new sudden sensorineural hearing loss (SSHL) cases per year identified in inpatient and outpatient healthcare settings via the International Classification of Diseases, 9th Revision (ICD-9) code (388.2) in 2006 and 2007 with an overall annual incidence of 27 per 100,000 population. Age- and sex-specific incidence rates were also reported in this study and used as the basis for the age and sex-stratified O/E analyses.
Table 9. Observed to Expected (O/E) Ratio for Spontaneously Reported Cases of Hearing Loss Through 18 June 2022

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Observed cases</th>
<th>Time at risk (PY)</th>
<th>Background rates per 100,000 PY</th>
<th>Expected cases</th>
<th>O/E ratio</th>
<th>95% CI LL</th>
<th>95% CI UL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>21-day risk window</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>US/EEA</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Males</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>2</td>
<td>901,382</td>
<td>9.0</td>
<td>81.12</td>
<td>0.025</td>
<td>0.003</td>
<td>0.089</td>
</tr>
<tr>
<td>12-17 years</td>
<td>22</td>
<td>1,562,311</td>
<td>9.0</td>
<td>140.61</td>
<td>0.156</td>
<td>0.098</td>
<td>0.237</td>
</tr>
<tr>
<td>18-24 years</td>
<td>41</td>
<td>2,224,929</td>
<td>12.0</td>
<td>266.99</td>
<td>0.154</td>
<td>0.110</td>
<td>0.208</td>
</tr>
<tr>
<td>25-49 years</td>
<td>518</td>
<td>9,186,672</td>
<td>20.3</td>
<td>1867.06</td>
<td>0.277</td>
<td>0.254</td>
<td>0.302</td>
</tr>
<tr>
<td>50-59 years</td>
<td>279</td>
<td>4,241,876</td>
<td>41.5</td>
<td>1700.38</td>
<td>0.158</td>
<td>0.140</td>
<td>0.178</td>
</tr>
<tr>
<td>60-69 years</td>
<td>182</td>
<td>3,713,614</td>
<td>67.0</td>
<td>2488.12</td>
<td>0.075</td>
<td>0.063</td>
<td>0.085</td>
</tr>
<tr>
<td>70+ years</td>
<td>195</td>
<td>5,152,276</td>
<td>81.0</td>
<td>4173.34</td>
<td>0.047</td>
<td>0.040</td>
<td>0.054</td>
</tr>
<tr>
<td><strong>Females</strong></td>
<td></td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>2</td>
<td>1,016,452</td>
<td>7.0</td>
<td>71.15</td>
<td>0.028</td>
<td>0.003</td>
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<tr>
<td>12-17 years</td>
<td>42</td>
<td>1,761,755</td>
<td>7.0</td>
<td>123.32</td>
<td>0.341</td>
<td>0.245</td>
<td>0.460</td>
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<tr>
<td>18-24 years</td>
<td>92</td>
<td>2,508,963</td>
<td>13.0</td>
<td>326.17</td>
<td>0.282</td>
<td>0.227</td>
<td>0.346</td>
</tr>
<tr>
<td>25-49 years</td>
<td>804</td>
<td>10,359,439</td>
<td>21.0</td>
<td>2175.48</td>
<td>0.370</td>
<td>0.344</td>
<td>0.396</td>
</tr>
<tr>
<td>50-59 years</td>
<td>419</td>
<td>4,783,392</td>
<td>36.5</td>
<td>1745.94</td>
<td>0.240</td>
<td>0.218</td>
<td>0.264</td>
</tr>
<tr>
<td>60-69 years</td>
<td>283</td>
<td>4,187,692</td>
<td>52.0</td>
<td>2177.60</td>
<td>0.130</td>
<td>0.115</td>
<td>0.146</td>
</tr>
<tr>
<td>70+ years</td>
<td>265</td>
<td>5,810,014</td>
<td>62.0</td>
<td>3602.21</td>
<td>0.074</td>
<td>0.065</td>
<td>0.083</td>
</tr>
<tr>
<td><strong>Overall</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any dose</td>
<td>3,146</td>
<td>57,410,765</td>
<td>27.0</td>
<td>15500.91</td>
<td>0.203</td>
<td>0.196</td>
<td>0.210</td>
</tr>
<tr>
<td>Dose 1</td>
<td>1,513</td>
<td>22,917,286</td>
<td>27.0</td>
<td>6187.67</td>
<td>0.245</td>
<td>0.252</td>
<td>0.257</td>
</tr>
<tr>
<td>Dose 2</td>
<td>1,255</td>
<td>21,280,003</td>
<td>27.0</td>
<td>5745.60</td>
<td>0.218</td>
<td>0.207</td>
<td>0.231</td>
</tr>
<tr>
<td>Dose 3</td>
<td>378</td>
<td>13,213,476</td>
<td>27.0</td>
<td>3567.64</td>
<td>0.106</td>
<td>0.096</td>
<td>0.117</td>
</tr>
<tr>
<td><strong>Overall Global</strong></td>
<td>4424</td>
<td>127,075,389</td>
<td>27.0</td>
<td>34310.35</td>
<td>0.129</td>
<td>0.125</td>
<td>0.133</td>
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<tr>
<td><strong>42-day risk window</strong></td>
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</tr>
<tr>
<td><strong>US/EEA</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Males</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>2</td>
<td>1,306,591</td>
<td>9.0</td>
<td>117.59</td>
<td>0.017</td>
<td>0.002</td>
<td>0.061</td>
</tr>
<tr>
<td>12-17 years</td>
<td>25</td>
<td>2,409,778</td>
<td>9.0</td>
<td>216.88</td>
<td>0.115</td>
<td>0.075</td>
<td>0.170</td>
</tr>
<tr>
<td>18-24 years</td>
<td>44</td>
<td>3,514,277</td>
<td>12.0</td>
<td>421.71</td>
<td>0.104</td>
<td>0.076</td>
<td>0.140</td>
</tr>
<tr>
<td>25-49 years</td>
<td>562</td>
<td>14,565,823</td>
<td>20.3</td>
<td>2061.72</td>
<td>0.190</td>
<td>0.174</td>
<td>0.206</td>
</tr>
<tr>
<td>50-59 years</td>
<td>307</td>
<td>6,796,938</td>
<td>41.5</td>
<td>2820.73</td>
<td>0.210</td>
<td>0.197</td>
<td>0.223</td>
</tr>
<tr>
<td>60-69 years</td>
<td>203</td>
<td>6,042,722</td>
<td>67.0</td>
<td>4048.62</td>
<td>0.050</td>
<td>0.043</td>
<td>0.058</td>
</tr>
<tr>
<td>70+ years</td>
<td>214</td>
<td>8,411,485</td>
<td>81.0</td>
<td>6813.30</td>
<td>0.031</td>
<td>0.027</td>
<td>0.036</td>
</tr>
<tr>
<td><strong>Females</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>2</td>
<td>1,473,390</td>
<td>7.0</td>
<td>103.14</td>
<td>0.019</td>
<td>0.002</td>
<td>0.070</td>
</tr>
<tr>
<td>12-17 years</td>
<td>45</td>
<td>2,717,409</td>
<td>7.0</td>
<td>190.22</td>
<td>0.237</td>
<td>0.173</td>
<td>0.317</td>
</tr>
<tr>
<td>18-24 years</td>
<td>98</td>
<td>3,962,908</td>
<td>13.0</td>
<td>515.18</td>
<td>0.190</td>
<td>0.154</td>
<td>0.232</td>
</tr>
<tr>
<td>25-49 years</td>
<td>864</td>
<td>16,425,290</td>
<td>21.0</td>
<td>3449.31</td>
<td>0.250</td>
<td>0.234</td>
<td>0.268</td>
</tr>
<tr>
<td>50-59 years</td>
<td>462</td>
<td>7,664,632</td>
<td>36.5</td>
<td>2797.59</td>
<td>0.165</td>
<td>0.150</td>
<td>0.181</td>
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<tr>
<td>60-69 years</td>
<td>306</td>
<td>6,814,133</td>
<td>52.0</td>
<td>3543.35</td>
<td>0.086</td>
<td>0.077</td>
<td>0.097</td>
</tr>
<tr>
<td>70+ years</td>
<td>293</td>
<td>9,485,291</td>
<td>62.0</td>
<td>5880.88</td>
<td>0.050</td>
<td>0.044</td>
<td>0.056</td>
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<tr>
<td><strong>Overall</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any dose</td>
<td>3,427</td>
<td>91,590,667</td>
<td>27.0</td>
<td>24729.48</td>
<td>0.139</td>
<td>0.134</td>
<td>0.143</td>
</tr>
<tr>
<td>Dose 1</td>
<td>1,637</td>
<td>22,917,286</td>
<td>27.0</td>
<td>6187.67</td>
<td>0.265</td>
<td>0.252</td>
<td>0.278</td>
</tr>
<tr>
<td>Dose 2</td>
<td>1,388</td>
<td>42,494,750</td>
<td>27.0</td>
<td>11473.58</td>
<td>0.121</td>
<td>0.115</td>
<td>0.128</td>
</tr>
<tr>
<td>Dose 3</td>
<td>402</td>
<td>26,178,631</td>
<td>27.0</td>
<td>7068.23</td>
<td>0.057</td>
<td>0.051</td>
<td>0.063</td>
</tr>
<tr>
<td><strong>Overall Global</strong></td>
<td>4,814</td>
<td>205,968,419</td>
<td>27.0</td>
<td>55076.33</td>
<td>0.087</td>
<td>0.085</td>
<td>0.090</td>
</tr>
</tbody>
</table>

CI = confidence interval; EEA = European Economic Area; LL = lower limit; PY = person-years; UL = upper limit; US = United States
Note: The background rate by Alexander TH et al. for age group of <18 years were used for ≤11 and 12-17 years, the average for age groups of 18-34, 35-44 and 45-54 years were used for 25-49 years, the average for age groups of 45-54 and 55-64 years were used for 50-59 years, and the average for age group 55-64 and 65+ years were used for 60-69 years, respectively.

Based on the selected background rates and the estimated number of exposure person-years through 18 June 2022, O/E ratios were well below one overall, by dose, and within strata of age groups and sex for both risk windows of 21- and 42-days. This suggests that the number of observed cases of
hearing loss is not higher than expected in the absence of Pfizer–BioNTech COVID-19 vaccines overall and within the queried strata.

**Rapporteur assessment comment:**

All O/E ratios were below 1 for hearing loss.

**Tinnitus**

The MAH conducted unadjusted observed to expected ratio (O/E) analyses for 16671 tinnitus cases reported cumulatively through 18 June 2022 globally.

The overall expected case counts of tinnitus were estimated using background incidence rates (IR) reported by a retrospective cohort study in the UK Clinical Practice Research Datalink (CPRD) with linkage to the Hospital Episode Statistics. This study included 14,303 incident clinically significant tinnitus during 01 January 2002 to 31 December 2011, defined by a discharge from the hospital with a primary diagnosis of tinnitus or a primary care recording of tinnitus with subsequent related medical follow-up within 28 days. The overall incidence rate was 54 per 100,000 person-years (95% confidence interval [CI]=53, 55). Age- and sex specific incidence rates were also reported in this study. These rates provided a low range of background rates for tinnitus, and therefore, low expected case counts to conservatively estimate O/E results. Another retrospective cohort using CPRD identified 109,783 adults with a first-time diagnosis of tinnitus between 2000 and 2016 and reported an overall incidence rate of 250 per 100,000 person years (95%CI=246, 255).
Table 10. Observed to Expected (O/E) Ratio for Spontaneously Reported Cases of Tinnitus Through 18 June 2022

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Observed cases</th>
<th>Time at risk (PY)</th>
<th>Background rates per 100,000 PY</th>
<th>Expected cases</th>
<th>O/E ratio</th>
<th>95% CI LL</th>
<th>95% CI UL</th>
</tr>
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<tr>
<td><strong>21-day risk window</strong></td>
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<tr>
<td>US-EFA</td>
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<td></td>
</tr>
<tr>
<td>Males</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>4</td>
<td>901,382</td>
<td>5.5</td>
<td>49.58</td>
<td>0.081</td>
<td>0.022</td>
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</tr>
<tr>
<td>12-17 years</td>
<td>34</td>
<td>1,562,311</td>
<td>9.0</td>
<td>140.61</td>
<td>0.242</td>
<td>0.167</td>
<td>0.338</td>
</tr>
<tr>
<td>18-24 years</td>
<td>106</td>
<td>2,224,929</td>
<td>10.0</td>
<td>355.99</td>
<td>0.298</td>
<td>0.244</td>
<td>0.360</td>
</tr>
<tr>
<td>25-49 years</td>
<td>1,872</td>
<td>9,186,672</td>
<td>42.3</td>
<td>3,889.02</td>
<td>0.481</td>
<td>0.460</td>
<td>0.504</td>
</tr>
<tr>
<td>50-69 years</td>
<td>1,002</td>
<td>4,241,876</td>
<td>106.0</td>
<td>4,496.39</td>
<td>0.223</td>
<td>0.209</td>
<td>0.237</td>
</tr>
<tr>
<td>60-69 years</td>
<td>552</td>
<td>3,713,614</td>
<td>122.0</td>
<td>4,550.61</td>
<td>0.122</td>
<td>0.112</td>
<td>0.132</td>
</tr>
<tr>
<td>70+ years</td>
<td>259</td>
<td>5,152,276</td>
<td>73.0</td>
<td>3,761.16</td>
<td>0.069</td>
<td>0.061</td>
<td>0.076</td>
</tr>
<tr>
<td>Females</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤11 years</td>
<td>3</td>
<td>1,016,452</td>
<td>4.5</td>
<td>45.74</td>
<td>0.066</td>
<td>0.014</td>
<td>0.192</td>
</tr>
<tr>
<td>12-17 years</td>
<td>58</td>
<td>1,761,155</td>
<td>7.0</td>
<td>123.32</td>
<td>0.470</td>
<td>0.357</td>
<td>0.608</td>
</tr>
<tr>
<td>18-24 years</td>
<td>235</td>
<td>2,508,963</td>
<td>13.5</td>
<td>338.71</td>
<td>0.694</td>
<td>0.608</td>
<td>0.788</td>
</tr>
<tr>
<td>25-49 years</td>
<td>3,095</td>
<td>10,599,439</td>
<td>39.3</td>
<td>4,074.71</td>
<td>0.760</td>
<td>0.733</td>
<td>0.787</td>
</tr>
<tr>
<td>50-69 years</td>
<td>1,614</td>
<td>4,783,192</td>
<td>100.0</td>
<td>4,783.39</td>
<td>0.537</td>
<td>0.521</td>
<td>0.554</td>
</tr>
<tr>
<td>60-69 years</td>
<td>935</td>
<td>4,187,692</td>
<td>107.0</td>
<td>4,480.83</td>
<td>0.209</td>
<td>0.196</td>
<td>0.222</td>
</tr>
<tr>
<td>70+ years</td>
<td>462</td>
<td>5,810,014</td>
<td>81.0</td>
<td>4,706.11</td>
<td>0.098</td>
<td>0.089</td>
<td>0.108</td>
</tr>
<tr>
<td><strong>Overall</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any dose</td>
<td>10,231</td>
<td>57,410,765</td>
<td>54.0</td>
<td>31,001.81</td>
<td>0.330</td>
<td>0.324</td>
<td>0.336</td>
</tr>
<tr>
<td>Dose 1</td>
<td>5,040</td>
<td>22,917,286</td>
<td>54.0</td>
<td>12,375.33</td>
<td>0.407</td>
<td>0.396</td>
<td>0.419</td>
</tr>
<tr>
<td>Dose 2</td>
<td>4,008</td>
<td>21,280,003</td>
<td>54.0</td>
<td>11,491.20</td>
<td>0.349</td>
<td>0.338</td>
<td>0.360</td>
</tr>
<tr>
<td>Dose 3</td>
<td>1,183</td>
<td>13,213,476</td>
<td>54.0</td>
<td>7,135.28</td>
<td>0.166</td>
<td>0.156</td>
<td>0.176</td>
</tr>
<tr>
<td><strong>Overall Global</strong></td>
<td>14,347</td>
<td>127,075,389</td>
<td>54.0</td>
<td>66,860.71</td>
<td>0.209</td>
<td>0.206</td>
<td>0.213</td>
</tr>
</tbody>
</table>

| **42-day risk window** |                |                   |                               |                |          |          |          |
| US-EFA |                |                   |                               |                |          |          |          |
| Males |                |                   |                               |                |          |          |          |
| ≤11 years | 4 | 1,306,591 | 5.5 | 71.86 | 0.056 | 0.015 | 0.143 |
| 12-17 years | 38 | 2,409,778 | 9.0 | 216.88 | 0.175 | 0.124 | 0.240 |
| 18-24 years | 113 | 3,514,277 | 16.0 | 562.28 | 0.201 | 0.166 | 0.242 |
| 25-49 years | 1,985 | 14,565,823 | 42.3 | 6166.20 | 0.322 | 0.308 | 0.336 |
| 50-69 years | 1,075 | 6,796,938 | 106.0 | 7204.75 | 0.149 | 0.140 | 0.158 |
| 60-69 years | 592 | 6,042,722 | 122.0 | 7372.12 | 0.080 | 0.074 | 0.087 |
| 70+ years | 264 | 8,411,485 | 73.0 | 6140.38 | 0.046 | 0.041 | 0.052 |
| Females |                |                   |                               |                |          |          |          |
| ≤11 years | 4 | 1,473,390 | 4.5 | 66.30 | 0.060 | 0.016 | 0.154 |
| 12-17 years | 60 | 2,717,409 | 7.0 | 190.22 | 0.315 | 0.241 | 0.406 |
| 18-24 years | 244 | 3,062,908 | 13.5 | 534.99 | 0.456 | 0.401 | 0.517 |
| 25-49 years | 2,237 | 16,425,290 | 39.3 | 6460.61 | 0.501 | 0.484 | 0.519 |
| 50-69 years | 1,728 | 7,664,632 | 100.0 | 7664.63 | 0.225 | 0.215 | 0.236 |
| 60-69 years | 991 | 6,814,133 | 107.0 | 7291.12 | 0.136 | 0.128 | 0.145 |
| 70+ years | 506 | 9,485,291 | 81.0 | 7883.09 | 0.066 | 0.060 | 0.072 |
| **Overall** |                |                   |                               |                |          |          |          |
| Any dose | 10,861 | 91,590,687 | 54.0 | 40,458.96 | 0.220 | 0.215 | 0.224 |
| Dose 1 | 5,286 | 22,917,286 | 54.0 | 12,375.33 | 0.427 | 0.416 | 0.439 |
| Dose 2 | 4,331 | 42,494,750 | 54.0 | 22947.17 | 0.189 | 0.183 | 0.194 |
| Dose 3 | 1,244 | 26,178,631 | 54.0 | 14136.46 | 0.088 | 0.083 | 0.093 |
| **Overall Global** | 15,199 | 203,986,419 | 54.0 | 110,152.67 | 0.138 | 0.136 | 0.140 |

CI = confidence interval; EEA=European Economic Area; LL = lower limit; PY = person-years; UL= upper limit; US=United State

Note: The average of background rate by Martinez C et al15: for age group of <10 and 10-19 years were used for ≤11 years, the average for age groups of 10-19 and 20-29 years were used for 12-17 years, the average for age groups of 20-29, 30-39, and 40-49 years were used for 25-49 years, and the average for age group 70-70 and 80-84 years were used for 70+ years, respectively.

Based on the select background rates and the estimated number of exposure person-years through 18 June 2022, O/E ratios were below one overall, by dose, and within strata of age groups and sex for both risk windows of 21- and 42-days. This suggests that the number of observed cases of tinnitus is not higher than expected in the absence of Pfizer-BioNTechCOVID-19 vaccines overall and within the queried strata.
Rapporteur assessment comment:
All O/E ratios were below 1 for tinnitus.

Clinical trial data

Hearing loss

In the placebo-controlled period of pivotal clinical trial C4591001 (DLP 13MAR2021), there were 12 participants who reported hearing loss events; 9/23,037 (0.04%) were in the placebo group and 3/23,040 (0.01%) were in the BNT162b2 group. In both groups the events of hearing loss were nonserious; the reported latency in the placebo group varied from Day 2 to Day 111; the latency in BNT162b2 group were Day 1, Day 3, and Day 19, respectively.

In the placebo-controlled period of clinical trial C4591031 (DLP 08FEB2022), there was 1 of 5048 participants in the placebo group who reported a hearing loss event; there were none reported in BNT162b2 group (5088 participants). The reported event was nonserious and occurred within the same day of vaccination.

In the placebo-controlled period of clinical trial C4591007 (DLP 06SEP2021), there were 3/750 participants who reported hearing loss events in the placebo group; there were none reported in the BNT162b2 group (1518 participants). The reported event were nonserious; the reported latencies were unknown (1 event), 65 days (1 event), and 135 days (1 event).

Rapporteur assessment comment:
Pooled in the three placebo-controlled studies there were 3 of the 29,646 (0.01%) participants reporting hearing loss in the Comirnaty group and 14 of the 28,835 (0.05%) participants in the placebo group.

Tinnitus

In the placebo-controlled period of pivotal clinical trial C4591001 (DLP 13MAR2021), there were 24 participants who reported tinnitus events; 23/23,037 (0.1%) were in the placebo group and 1/23,040 (0.004%) was in the BNT162b2 group. In both groups the events of tinnitus were nonserious. The most frequently reported latencies in the placebo group were Day 0 and Day 3 (for 3 events each); the latency for the event in the BNT162b2 group was 26 days.

In the placebo-controlled period of clinical trial C4591031 (DLP 08FEB2022), there was 1 of 5048 participants in the placebo group who reported an event of tinnitus; none were reported in the BNT162b2 group (5088 participants). The reported event was nonserious and occurred 42 days post vaccination.

There were no events of Tinnitus reported in the placebo-controlled period of clinical trial C4591007.

Rapporteur assessment comment:
Pooled in the three placebo-controlled studies there was 1 of the 29,646 (0.003%) participants reporting tinnitus in the Comirnaty group and 25 of the 28,835 (0.09%) participants in the placebo group.

MAH’s conclusion
Reports of recovery of SARS-CoV-2 RNA in the middle ear of individuals who died of COVID-19 and recent findings of the ability of SARS-CoV-2 to directly infect human vestibular hair and Schwann cells provide plausible biological mechanisms for COVID-19-associated hearing loss and may open avenues of investigation into immune mechanisms in the inner ear.

There is no documented mode of action of how vaccination with COVID-19 vaccines can cause hearing loss or tinnitus.

Participants in the placebo-controlled, blinded periods of the large Pfizer-run clinical trials reported very low number of tinnitus or hearing loss in either group with the placebo group having a higher number of tinnitus and hearing loss events compared to the vaccine group.

No signal of disproportionate reporting has been observed in the Pfizer safety database for any of the Preferred Terms included in the safety database search.

The spontaneously reported cases are of variable quality. While there are individual cases that provide detailed information without alternative causes of hearing loss and tinnitus, the nature of these events (their myriad etiologies) and the reports do not exclude that the events may be coincidental to vaccination.

The O/E analyses provides reassurance that the reports of hearing loss and tinnitus in the stratified populations and doses are not greater than would be expected as background occurrences.

The observational studies in the medical literature retrieved in the search do not allow a conclusion one way or the other about a causal relationship between hearing loss or tinnitus and Comirnaty at a population level. Further observational studies that include data from patients vaccinated over time (not just in the initial months of mass vaccination) will be helpful to further characterize any association.

Observational studies from healthcare networks that can combine information from vaccination records and healthcare system diagnoses may provide a more accurate picture of real-world events than passive surveillance systems (e.g., spontaneous databases). While the general expectation is that spontaneous reporting will be highest immediately following post-authorization product availability, the COVID-19 vaccines are subject to unusual reporting practices due to the global nature of the pandemic (worldwide population is affected, not a subgroup of patients), the multi-dose regimen with different mass vaccination practices in different regions, the unprecedented scrutiny on adverse events occurring in temporal relation with vaccination and the vigorous encouragement to report adverse events.

Taking into account the totality of the data available, a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time. Routine pharmacovigilance will continue.

**Rapporteur assessment comment:**

**Literature**

From the four retrieved relevant articles no new important safety information could be identified concerning hearing loss and tinnitus.

**Post-marketing**

Through 18 Jun 2022 were retrieved:

- 2009 cases reporting hearing loss and tinnitus; analysis was concentrated on the 212 HCP confirmed cases: 27 cases were considered possible related to Comirnaty exposure (including a possible
rechallenge in a 14 year old female), 110 cases were considered unlikely related to Comirnaty exposure, and 75 cases were considered unassessable.

- 13,943 cases reporting tinnitus; analysis was concentrated on the 295 HCP confirmed cases: 2 cases were considered possible related to Comirnaty exposure, 161 were considered unlikely related to Comirnaty, and 132 cases were considered unassessable.

- 3177 cases reporting hearing loss; analysis was concentrated on the 352 HCP confirmed cases: 6 cases were considered possible related to Comirnaty exposure, 153 cases were considered unlikely related to Comirnaty exposure, and 193 cases were considered unassessable.

In conclusion, despite the 27 cases reporting hearing loss and tinnitus and the 2 cases reporting tinnitus and the 6 cases reporting hearing loss that were considered possible related to Comirnaty exposure, there seems to be no causal association between Comirnaty exposure and occurrence of hearing loss and/or tinnitus in the post-marketing cases. However, a detailed MAH’s case by case assessment of the cases considered possible related to Comirnaty exposure is not presented in the PSUR, which hampers PRAC Rapporteur’s assessment.

**Observed to expected analyses**

Through 18 June 2022, for hearing loss and tinnitus, all O/E ratios were >1 overall, by dose, and within strata of age groups and sex for both risk windows of 21- and 42-days.

**Clinical trial data**

In the clinical trial there were more reports of hearing loss and of tinnitus in the placebo group compared to the Comirnaty group, 14 and 25 versus 3 and 1, respectively.

Overall, MAH’s conclusion is endorsed that a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time. No new important information could be identified concerning hearing loss or tinnitus. The MAH should continue monitor any new cases, patterns, or trends of reporting hearing loss and/or tinnitus through routine pharmacovigilance.

**Issue solved**

- the signals that were closed during the reporting period of the PSUR,
- earlier closed signals only when there are deviating trends in severity of AEs, incidence, and/or outcomes,
- previously closed signals requested by PRAC

### 2.2.2. Evaluation of closed signals

**Rapporteur assessment comment:**

General note, the MAH should present in the section ‘Evaluation of closed signals’:

- all the signals that were closed during the reporting period of the PSUR;
- earlier closed signals only when there are deviating trends in severity of AEs, incidence, and/or outcomes;
- previously closed signals as requested by PRAC.
### Signals determined to not be risks

<table>
<thead>
<tr>
<th>Condition</th>
<th>Description</th>
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<tr>
<td>Appendicitis</td>
<td>Appendicitis was identified as a signal during the reporting period based on a competent authority (Singapore BoH) inquiry following 18 local reports. The Pfizer safety database search through 01 April 2022 revealed 690 cases and those with sufficient information provided did not show any trends considered inconsistent with the underlying epidemiology and/or natural course of the condition. Placebo-controlled clinical trial data from the pivotal Pfizer-run studies did not reveal any clinically meaningful difference between the BNT162b2 and placebo groups. Of 3 large population-based studies from the US, Israel and Sweden, 2 showed no increase in appendicitis after vaccination while 1 showed a slightly increased risk ratio in the vaccinated group compared to the unvaccinated group. Observed to expected analyses conducted were well below 1. A plausible mechanism by which BNT162b2 could case appendicitis is unknown. The totality of the information was not supportive of a causal association between BNT162b2 and appendicitis signal was closed by the MAH.</td>
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<tr>
<td>Rapporteur assessment comment:</td>
<td>Through 01 Apr 2022, the MAH retrieved 690 cases reporting appendicitis and did not show any trends supportive of a causal association with Comirnaty exposure. Clinical trial data showed no clinically meaningful difference between the Comirnaty and placebo groups. O/E ratios were &lt;1. The MAH stated that there is no causal association between Comirnaty exposure and appendicitis. Therefore, the signal was closed.</td>
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<tr>
<td>Hemolytic anemia</td>
<td>Haemolytic anaemia was identified as a signal during the reporting period based on a competent authority (Saudi FDA) request for an evaluation. The Pfizer safety database search through 13 January 2022 yielded 176 cases, most of which were confounded or contained insufficient information. Among the cases with no obvious confounder or trigger, a definitive causal association could not be concluded. There were no events of haemolytic anaemia reported in the pivotal clinical trial C4591001. The medical literature yielded one case report and one prospective study of 108 patients with autoimmune cytopenias (56 with autoimmune haemolytic anaemia [AIHI]) who were vaccinated with Pfizer/BNT, Moderna or Astra-Zeneca COVID-19 vaccines. Four elderly patients with AIHI had a clinically significant haemoglobin reduction requiring treatment adjustment (2 had received Pfizer/BNT vaccine). Notably, autoimmune cytopenia recrudescences were not predictable, since they occurred in both patients on active treatment and off therapy, independently from AIHA type, after either the first or the second dose, and regardless of vaccine type. Observed to expected analyses conducted were below 1. Based on the totality of available information, a</td>
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<td><strong>Rapporteur assessment comment:</strong></td>
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<td>MAH’s response to the inquiry received from the Saudi Food and Drug Authority concerning hemolytic anemia was assessed in the 13th (2nd bi-monthly) SSR (reporting period 16 Dec 2021 – 15 Feb 2022; EMEA/H/C/005735/MEA/002.12).</td>
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<tr>
<td>MAH’s conclusion was endorsed that there is insufficient evidence to establish causality between the development of hemolytic anemia events and Comirnaty exposure. Closure of the signal hemolytic anemia was accepted.</td>
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<th><strong>Uveitis</strong></th>
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<td>Uveitis was identified as a signal during the reporting period based on a competent authority (Health Canada) request for a cumulative review. The Pfizer safety database search through 04 April 2022 yielded 538 cases, 121 of which were medically confirmed and did not report confounding factors or an implausible time to onset. Of these, 9 were determined to have a possible causality based on individual assessment. During the placebo-controlled period in the pivotal clinical trial C4591001, one case was reported in the placebo group and no cases were reported in the BNT162b2 group from Dose 1 to 1 month after Dose 2. The medical literature consisted of case reports and case series descriptions with one population-based study estimating the prevalence rates of uveitis coincident with COVID-19 vaccination as 0.9 cases per million doses or less. Observed to expected analyses conducted using both a low and high range of background rates were well below 1 overall, by dose and within age and sex strata. Based on the totality of available information, a causal association between BNT162b2 could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<th><strong>Rapporteur assessment comment:</strong></th>
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<tr>
<td>MAH’s response to the inquiry received from the Canadian Health Authority concerning uveitis was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/H/C/005735/MEA/002.13).</td>
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<tr>
<td>MAH’s conclusion was endorsed that there is insufficient evidence to establish causality between uveitis events and Comirnaty exposure. Closure of the signal uveitis was accepted.</td>
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<th><strong>Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders</strong></th>
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| A cumulative review of autoimmune and inflammatory disorder exacerbations was requested in an updated PSUR Assessment Report received from EMA/PRAC during the reporting period (30 December 2021). The search of the safety database used SMQ Immune-mediated/autoimmune disorders (narrow terms), HLGT Autoimmune disorders, HLGT Immune disorders NEC and HLT Neuromuscular junction dysfunction. There were 2223 cases describing a medical history and adverse event of an autoimmune disease (indicating a potential exacerbation). Overall, cases lacked information to ascertain baseline disease status, treatment and other factors which may affect underlying
disease activity, despite most reporting exacerbations or potential relapses within 2 days of vaccination. In the placebo-controlled portion of pivotal clinical trial C4591001, 2955/21926 BNT162b2 participants and 2977/21921 placebo participants had underlying autoimmune conditions. Of these, 7 (0.2%) and 4 (0.1%) of BNT and placebo participants, respectively, reported potential aggravations of their autoimmune disorder from Dose 1 to 1 month post Dose 2. From Dose 1 to unblinding, 8 participants in each group (IR/100 person years = 0.7) reported potential aggravations. The medical literature search yielded many studies of COVID-19 vaccination in patients with underlying autoimmune disorders. The findings consistently showed that reported post-vaccination adverse events were similar to those of healthy vaccinees. Most studies did not have control groups of participants with autoimmune disorders who did not receive COVID-19 vaccination, although those that did, did not report that vaccinees had more exacerbations than non-vaccinees. Based on the totality of the available information, a causal association between BNT162b2 and autoimmune disorder exacerbations could not be concluded, and the signal was closed by the MAH.

**Rapporteur assessment comment:**

MAH's response on the PRAC request for a cumulative review of exacerbation (flare-up) of pre-existing AI/Inflammatory disorders was assessed in the 2nd Comirnaty PSUSA (reporting period 19 Jun 2021 – 18 Dec 2021; EMEA/H/C/PSUSA/00010898/202112).

MAH's conclusion was supported and that no new important safety information could be identified. Closure of the signal was accepted.

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**Capillary leak syndrome (CLS)**

Capillary leak syndrome, or Systemic capillary leak syndrome (SCLS), was identified as a signal by PRAC on 13 January 2022. The safety database search yielded 44 cases, 2 of which were literature case reports, which occurred in individuals from 20 to 101 years of age. Four cases described a medical history of CLS. The majority of cases lacked clinical details or provided evidence of an alternative aetiology other than vaccination. There were no reported events of CLS in the placebo (21921) or BNT162b2 group (21926) in the placebo-controlled portion of C4591001 in participants 16 years and older from dose 1 to 1 month after dose 2. The medical literature has described cases of CLS occurring after COVID-19 infection and there were only individual case reports of CLS occurring after COVID-19 vaccination. Based on the totality of the available information, a causal association between BNT162b2 and CLS/SCLS could not be concluded, and the signal was closed by the MAH.

**Rapporteur assessment comment:**

Please refer to the separate signal procedure concerning capillary leak syndrome (EMEA/H/C/005735/SDA/051- EPITT 19743) in which PRAC concluded to continue closely monitoring through routine pharmacovigilance. The signal capillary leak syndrome was closed.
**Corneal graft rejection**

Corneal graft rejection was identified as a signal by PRAC on 07 April 2022. The safety database search through 14 April 2022 yielded 42 potential cases describing 40 unique individuals, all adults or elderly. There was no distinguishing trend in the cases with regard to sex, age, dose number, age of graft or time to onset. Of 12 cases with a plausible temporal relationship to vaccination, only 2 did not have reported risk factors for rejection (e.g., increased age of transplant, possible infection, graft surgery complications). Data from large clinical studies C4591001 (snapshot date of 11 April 2022), C4591031 (cut-off date of 08 February 2022) and C4591007 (cut-off dates of 08 October 2021 and 22 March 2022) were searched for PTs, corneal graft rejection and corneal graft failure. Neither of these PTs were reported in the unblinded data from the placebo-controlled portions of the studies. There were 32 clinical trial participants, all ≥16 years of age, who reported a history of corneal transplant or keratoplasty in either Study C4591001 and/or Study C4591031. There were no participants in C4591007 who reported a history of corneal transplant or keratoplasty. The medical literature consisted of case reports which were included in the safety database. There were no mechanistic studies, rather various hypotheses were theorized such as increased vascular permeability, Immune responses and immune system deregulation. Based on the totality of the available information, a causal association between BNT162b2 and corneal graft rejection could not be concluded, and the signal was closed by the MAH.

**Rapporteur assessment comment:**

Please refer to the separate signal procedure concerning corneal graft rejection (EMEA/H/C/005735/SDA/055- EPITT 19789). After DLP of the current PSUR, PRAC concluded to continue closely monitoring through routine pharmacovigilance. The signal corneal graft rejection was closed.

**Vasculitis**

During the reporting period, vasculitis was reviewed initially following a signal noted by the Lareb (Netherlands) and through 15 April at the request of PRAC in the Assessment Report for SBSR 2. Through 15 April 2022, a search of the safety database yielded 868 reports with individual ages ranging from 2 to 98 years. Reported vasculitides included vasculitis (not otherwise described), giant cell arteritis and Henoch-Schonlein purpura. The cases were generally confounded or lacked necessary details to confirm the diagnoses and/or a causal relationship. Phase 2/3 Study C4591001 placebo-controlled unblinded adverse events in participants 16 years and older from Dose 1 to 1 month after Dose 2 (data cutoff date 13 March 2021) was also reviewed for the PT Vasculitis. In the Phase 2/3 safety population, vasculitis was not reported in any of 21926 participants in the BNT162b2 (Pfizer-BioNTech COVID-19 Vaccine) group or in any of 21921 participants in the placebo group. Observed to expected analyses for the 3 most common subtypes of vasculitis (Henoch-Schonlein purpura, Giant cell arteritis, Skin manifestations of vasculitis) have repeatedly been less than one. Based on the totality of
the available information, a causal association between BNT162b2 and vasculitis could not be concluded, and the signal was closed by the MAH.

**Rapporteur assessment comment:**
An updated cumulative review of vasculitis (through 15 Apr 2022) was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/H/C/005735/MEA/002.13). PRAC concluded that based on the data provided no safety concern was identified. Closure is of the signal vasculitis was accepted.

| Cerebral venous sinus thrombosis (CVST) | Previous to the current PSUR reporting period, a cumulative review of CVST through 24 November 2021 was conducted by the MAH (SBSR 2, Appendix 3.4) in response to a request from a competent authority (Swissmedic). In the SBSR 2 PRAC Assessment Report, the MAH was requested to provide more detail on some of the cases and a further cumulative review of the topic. A search of the safety database yielded 527 cases that were reported through 24 November 2021 and 297 from 25 November 2021 to 15 April 2022. Of 37 cases in patients younger than 75 years of age with no medical history or information portending an increased risk for the development reported through 24 November 2021, only 3 were assessed as possible (the remaining were unassessable or unlikely per the WHO-UMC case causality criteria). In the interim update through 15 April 2022, the majority of the 297 cases lacked necessary detail for assessment, had implausible time to onset or described known risk factors (other than vaccination) for the development of CVST. When analyzed by age category, the cases were largely unassessable due to lack of sufficient detail for full assessment. Overall, the assessment of AE reports was that the clinical characteristics of the cases, when provided, were aligned with the known profile of CVST. There was 1 event of CVST in a clinical trial participant who received placebo in pivotal clinical trial C4S91001. Retrospective epidemiological studies of CVST after vaccination with BNT162b2 did not conclude an increased risk due to the vaccine and a retrospective analysis of 213 post-vaccine CVST cases did not demonstrate a clinically distinct profile of CVST after mRNA vaccination that differed from historical controls. Another large study showed an increased risk of CVST associated with COVID-19 compared to individuals with influenza or following COVID-19 vaccination with an mRNA vaccine. Observed to expected analyses have been conducted for CVST and when low background rates are used, the ratios are >1 in various age and sex strata. This is not seen when using higher background rates. The variation in reported background rates was notable it is possible that the delivery of healthcare, population demographics and underlying health status of the populations used for the background rate estimates differ from those in the vaccinated population. Based on the totality of the available information, a causal association between BNT162b2 and CVST could not be concluded, and the signal was closed by the MAH. |
**Rapporteur assessment comment:**

An updated cumulative review of cerebral venous sinus thrombosis (through 15 Apr 2022) was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/H/C/005735/MEA/002.13). PRAC concluded that based on the data provided no safety concern was identified. Closure is of the signal cerebral venous sinus thrombosis was accepted.

| Lymphocytic colitis | During the reporting period, this signal was identified from a published (literature) case report of a 69-year-old woman who presented for evaluation of severe abdominal pain, nausea, and diarrhea after her second vaccination with Pfizer/BNT COVID-19 vaccine. Within 24 hours of vaccination, she reported onset of diarrhea (2-3 loose to watery stools per day). Symptoms intensified over the next several days to 3-5 watery stools per day with incontinence, abdominal cramping, and nausea. GI PCR and COVID testing were negative and ondansetron and loperamide were started with minimal benefit. Two-months later, a GI consultation was obtained due to persistent symptoms. Work up demonstrated no anemia with normal CRP, celiac serologies, and GI PCR. Colonoscopy on day 98 post-onset revealed patchy erythema in the descending colon and rectosigmoid. Histologic evaluation of mucosal biopsies revealed lymphocytic colitis characterized by numerous lymphocytes infiltrating the epithelium and abundant plasma cells in the lamina propria. A previous colonoscopy performed in 2012 was unremarkable. At her most recent follow-up on day 113 post-onset, the patient reported gradual improvement of abdominal symptoms and diarrhea. This case report was recorded in the Pfizer safety database. There was no other relevant literature information on lymphocytic colitis and COVID-19 vaccination. A search of the safety database through 20 Jan 2022 yielded 40 cases for review (incl index case); in all the cases, there was either no Pfizer/BNT COVID-19 vaccine used, an unconfirmed diagnosis, lack of clinical detail, or the presence of alternative explanations or risk factors for lymphocytic colitis. Based on the totality of the available information, a causal association between BNT162b2 and lymphocytic colitis could not be concluded, and the signal was closed by the MAH. |

**Rapporteur assessment comment:**

This signal of lymphocytic colitis was assessed in the 13th (2nd bi-monthly) SSR (reporting period 16 Dec 2021 – 15 Feb 2022; EMEA/H/C/005735/MEA/002.12).

MAH's conclusion was endorsed that there is insufficient evidence to establish causality between lymphocytic colitis and Comirnaty exposure. Closure of the signal lymphocytic colitis was accepted.

| Chronic urticaria | This signal was identified following a request for a cumulative review on the subject from EMA PRAC on 09 May 2022. A search of the Pfizer safety database through 09 May 2022 yielded 244 cases; 31 of which described medical histories of chronic urticaria. Of the cases of new onset |
chronic urticaria, time to onset ranged from 0 to 90 days post vaccination, cases were reported after dose 1, dose 2 and booster doses, 26 cases described the background of underlying autoimmune disorders, 26 reported hypersensitivity conditions and 14 reported histories of COVID-19. In all, 35 cases specifically reported that the urticaria lasted more than 6 weeks (meeting criteria for chronic urticaria). Sixteen of these reported a time to onset that was reasonably temporally associated with vaccination, and among them only 7 (43.7%) of the cases provided a medical history (85.7% of which implied a predisposition to urticaria or an alternate trigger for it). During the placebo-controlled unblinded period of pivotal study C4591001 (data cut-off 15 April 2022), of participants 12 years and older, chronic urticaria was not reported in any of the 23,068 participants in the BNT162b2 group or the 23,063 participants in the placebo group from Dose 1 to data cutoff date. Observed versus expected analyses were < 1 overall, by dose and within strata of age groups. The MAH considers urticaria as an adverse reaction of BNT162b2, however, a causal association between the vaccine and chronic urticaria was not supported based on the available information.

Rapporteur assessment comment:
An updated cumulative review of chronic urticaria (through 09 May 2022) was assessed in the 2nd Comirnaty PSUSA (reporting period 19 Jun 2021 – 18 Dec 2021; EMEA/H/C/PSUSA/00010898/202112).
MAH’s conclusion was accepted that the available information did not support a causal association between chronic urticaria and Comirnaty exposure. The signal chronic urticaria was closed.

Polymyalgia rheumatica (PMR)
This signal was identified following a request for a cumulative review on the subject by EMA PRAC in the PSUR 1 Assessment Report. A search of the Pfizer safety database through 18 December 2021 yielded 628 cases, the majority of which were excluded from further consideration due to implausible time to onset, medical history or conditions confounding assessment or lack of clinical detail supporting the diagnosis of PMR. Of the reports providing laboratory data (CRP and/or ESR) supportive of the diagnosis of PMR, most were unassessable or unlikely per WHO-UMC causality criteria. The 54 cases reporting an exacerbation of PMR following vaccination were similarly hindered by lack of information. There were no reports of PMR in the pivotal Phase 2/3 Study C4591001 of individuals 16 years of age (21926 vaccine/21921 placebo) and older from Dose 1 to 1 month after Dose 2 that was placebo-controlled (data cutoff date 13 March 2021). Ten subjects reported polymyalgia rheumatica in the medical history and none of these subjects reported a flare up of the underlying disease. The medical literature search yielded several studies that did not support an association between vaccination and autoimmune disorders or flares. Based on the totality of the available information, a causal association between BNT162b2 and PMR could not be concluded, and the signal was closed by the MAH.
### Rapporteur assessment comment:

The cumulative review of polymyalgia rheumatica (through 18 Dec 2021) was assessed in the 2nd Comirnaty PSUSA (reporting period 19 Jun 2021 – 18 Dec 2021; EMEA/H/C/PSUSA/00010898/202112). PRAC concluded that the data did not suggest a causal association between Comirnaty and polymyalgia rheumatica.

| Subacute thyroiditis | This signal was identified following a request for a cumulative review on the topic by EMA PRAC on 18 January 2022 following the assessment of PSUR 2. The Pfizer safety database search yielded 498 cases through 18 December 2021. There was a similar number of cases reporting each of the PTs: Thyroiditis subacute, Autoimmune thyroiditis and Thyroiditis. The majority of reports described underlying thyroid disorder or concomitant disorders that represented confounding factors and/or did not provide a sufficient amount of information (medical history, laboratory and other diagnostic data) to allow a proper evaluation. The details of 38 cases that included laboratory work ups confirming hyperthyroid activity, did not provide enough relevant information to confirm the causal association with the vaccine. In the placebo-controlled portion of clinical trial C4591001, in the safety population of participants 16 years and older, there was 1 case of autoimmune thyroiditis reported among 21926 participants in the BNT162b2 group compared with 1 case of autoimmune thyroiditis among 21921 participants in the placebo group from dose 1 to 1 month after dose 2 (data cutoff date 13 March 2021). The medical literature consisted of case reports and case series of patients who developed thyroiditis following vaccination with various COVID-19 vaccines, including BNT162b2. Observed to expected analyses were conducted and ratios were below 1 for all age groups, doses and gender strata. Based on the totality of the available information, a causal association between BNT162b2 and subacute thyroiditis could not be concluded, and the signal was closed by the MAH. |

### Rapporteur assessment comment:

The cumulative review of subacute thyroiditis (through 18 Dec 2021) was assessed in the 2nd Comirnaty PSUSA (reporting period 19 Jun 2021 – 18 Dec 2021; EMEA/H/C/PSUSA/00010898/202112). PRAC concluded that no new safety concern was identified. Closure of the signal was accepted.

| Cerebrovascular accident (CVA)/Stroke | This signal was identified following a request from a competent authority (Australia, TGA) for an analysis on 27 January 2022. The Pfizer safety database search using a search strategy covering ischemic and hemorrhagic strokes, yielded 8934 cases through 18 December 2021. There were 4719 reports of females and 4024 of males, when sex was provided and the mean and median ages were 66.9 and 70 years of age, respectively. Cases were reviewed by age group. As would be expected |
based on the known epidemiology of stroke, the number of reported cases was highest in the oldest individuals and proportionally decreased with age; the number of ischemic strokes was greater than haemorrhagic strokes. Most of the reports described known risk factors for stroke and many of the cases in younger individuals were inconsistent with actual strokes upon individual case review. Of the 10 relevant studies in the medical literature, two studies (Shimazawa R et al [case reports] and Hippsley-Cox J et al) reported a correlative association between BNT162b2 vaccine and ischemic or haemorrhagic stroke. The study by Shimazawa R et al was based on 10 post-vaccination fatalities in Japan. Of these 10 cases, 4 females died of ICH. Insufficient details precluded further assessment. The study by Hippsley-Cox et al reported an increased risk of ischaemic stroke after a first dose of BNT162b2 but contextualized that this risk is far greater with COVID-19, emphasizing the importance of vaccination. Seven studies (Jabagi MJ et al, Simpson C.R et al, Carl L et al, Barada N et al, Koh JS et al, Klein NP et al, and Sessa M et al) did not support an association between BNT162b2 vaccine and haemorrhagic or ischemic stroke. In the remaining publication by Patone M et al, an increased risk of haemorrhagic stroke after BNT162b2 vaccination was reported in a study in England but was not replicated in a Scottish study that was somewhat smaller. Overall, the literature data does not support a clear causal association between BNT162b2 and stroke. The medical literature also did not provide a plausible mechanism for how BNT162b2 could increase the risk of haemorrhagic or ischemic strokes. All observed to expected ratios across all age, sex and dose stratifications were below 1 for haemorrhagic and ischemic strokes. Based on the totality of the available information, a causal association between BNT162b2 and hemorrhagic and ischemic stroke could not be concluded, and the signal was closed by the MAH.

Rapporteur assessment comment:

MAH's response to the Australian Therapeutic Goods Administration query concerning cerebrovascular accident/stroke data (through 18 Dec 2021) was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/C/005735/MEA/002.13) in which PRAC concluded that no new important safety information could be identified regarding cerebrovascular accident/stroke.

Amenorrhea

Amenorrhea was identified as a signal by PRAC on 14 February 2022. A search of the Pfizer safety database through 15 February 2022 yielded 9634 reports which were mostly non-serious and non-medically confirmed. Ages ranged from 11 to 66 years (mean 33.4). Of the cases without confounders and occurring in women younger than 45 years of age, causality assessments using the WHO-UMC criteria were all unlikely or unassessable. In the placebo-controlled portion of the C4591001 study, prior to treatment assignment unblinding, there were 8 events of amenorrhoea, with an equal split of 4 events after receipt of placebo vaccination and 4 events after receipt of active vaccination. Participants were followed up for a mean period of 135.8 days following the second
A heavy menstrual bleeding (HMB) was identified as a signal by PRAC on 14 February 2022. A search of the Pfizer safety database through 15 February 2022 yielded 23,659 cases of heavy menstrual bleeding. The majority are non-serious and non-medically confirmed. Of the much smaller subset of serious, medically confirmed reports that provided information about menstrual patterns, 4 were assessed as possibly related to vaccine using the WHO-UMC causality criteria as requested; one of the 4 was 1 of 2 cases that described a rechallenge. In addition, the O/E ratios do not indicate that reported events are higher than expected based on background incidence rates. In the placebo-controlled portion of the C4591001 study, prior to treatment assignment unblinding, there were 6 events of heavy menstrual bleeding; 4 of these events were after receipt of active vaccination and 2 events after receipt of placebo. The event in the C4591031 study occurred during the placebo-controlled portion of the study and was in a participant who received placebo. In the C4591001 study, participants were followed up for a mean period of 137.5 days following the second dose during the placebo-controlled follow-up period until unblinding (median 132 days; range 89 – 176 days). The participant in the C4591031 study was followed up for 96 days from blinded study vaccine (placebo) until unblinding. The medical literature on the topic reveals that menstrual abnormalities in general are very common and there have been correlations between SARS-CoV-2 pandemic stress, anxiety, and depression with menstrual cycle abnormalities. A clear pathophysiological mechanism for heavy menstrual bleeding itself is not understood. A well-designed US study of self-reported menstrual cycle data by Alison Edelman et al. did not support a significant effect of vaccination on the
number of days of menstrual bleeding. Studies are limited by their retrospective nature and self-reporting. While most menstruating women do not report menstrual changes associated with COVID-19 vaccination, it seems that variables such as age, BMI, changes in cycle over the previous year and the presence of fibroids and smoking may be playing a role. Based on the totality of the available information, a causal association between BNT162b2 and HMB could not be concluded, and the signal was closed by the MAH. On 13 June 2022, the PRAC responded with a list of questions that the MAH is in the process of preparing for submission by 24 August 2022.

Rapporteur assessment comment:
Please refer to the separate signal procedure heavy menstrual bleeding (EMEA/H/C/005735/SDA/053-EPTT 19783). PRAC concluded that heavy menstrual bleeding should be listed as ADR in the Comirnaty PI.

Loss of/ altered taste and smell
This was identified as a signal during the reporting period following a request for a competent authority (Australia, TGA) for an analysis of the topic. A search of the safety database through 01 March 2022, yielded 12,140 potentially relevant cases (1% of all AE reports for BNT162b2). There were 17 fatal cases all unrelated to vaccination but related to intercurrent diseases. To enable a focused review of the most informative cases, the MAH applied an exclusion algorithm focusing on serious and medically confirmed cases, excluding most confounding conditions and concomitant medications which could have contributed to the events. The identified 154 were further reviewed. 76 cases had insufficient information to make a thorough medical evaluation. 67 cases had alternative explanations or confounding factors which could have contributed to the events and there were 11 remaining cases, only five out of the 11 cases where judged "possible related" according to WHO causality assessment. There were no cases with a probable or definite relationship. Review of post-marketing data did not support a causal relationship between vaccination with BNT162b2 and the development of taste and smell disorders. Based on the mid-range background rates from ACCESS, O/E ratios were greater than 1 overall for all ages globally using the 7-day risk window. Using both mid- and high-range ACCESS background rates, the O/E ratios were >1 for certain age groups using both the 7- and 21-day risk windows, suggesting that the number of reported cases may be higher than expected compared to unvaccinated persons. The O/E ratios for these events may be overestimated for a few reasons. First, these pre-COVID (2017-2019) background rates from ACCESS reflect the incidence of anosmia and/or ageusia that was treated in either an inpatient or outpatient setting. Given that these symptoms may be mild in some cases, these rates might underestimate the overall incidence (and thus expected cases) of these conditions in a general, unvaccinated populations if not all cases are reported to healthcare providers. Second, they may also underestimate the incidence rate during the COVID-era since anosmia and/or ageusia are symptoms of
SARS-COV-2 infection. For example, the reported incidence rates for anosmia/ageusia were 1.4-2.3 times higher in 2020 than during 2017-2019 in both of the ACCESS sources used for background estimates. In 2020, the ES_SIDIAP_PC database rate was 35.23/100,000 persons per year, and in the ES-FISABIO database the rate was 67.5/100,000 persons per year. Third, observed cases may include cases of anosmia or ageusia due to recent or current SARS-COV-2 infection that was not documented or diagnosed during the risk window period. Fourth, due to the unprecedented attention to COVID-19 vaccination and outreach to encourage AE reporting, the long-held assumption that reported cases are an underestimation of actual cases may be incorrect. Finally, the ACCESS rates for anosmia and ageusia were defined with ICD codes that captured anosmia, parosmia, and parageusia, while the observed case definition included PTs for additional related conditions. In the pivotal clinical trial C4591001, during the placebo-controlled follow up period from Dose 1 to 1 month after Dose 2 of BNT162b2, 17 events of interest were reported in the vaccine group (N=21926) and 10 in the placebo group (N=21921) in participants ≥ 16 years of age. None of the 5 relevant PTs were reported by 12-15 year old from Dose 1 to 1 month post-Dose 2 in C4591001. None of the 5 PTs were reported by 5 to <12 year old from Dose 1 to 1 month Dose 2 in C4591007. There was a limited amount of medical literature on this topic and it was not supportive of a known relationship between vaccination and loss of taste or smell. Based on the totality of the available information, a causal association between BNT162b2 and anosmia and ageusia could not be concluded, and the signal was closed by the MAH.

**Rapporteur assessment comment:**

MAH’s response to the Australian Therapeutic Goods Administration query concerning loss of/altered taste and smell (data through 18 Dec 2021) was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/H/C/005735/MEA/002.13). PRAC concluded that the data provided did not support a causal relationship between Comirnaty exposure and the loss of/altered taste and smell.

### Important risks

**Myocarditis and Pericarditis**

*During the reporting period, myocarditis and pericarditis, which have been considered important identified risks in the US-PVP and EU-PVP, were moved from important potential risks to important identified risks in the company core list of safety concerns. After the DLP of this PSUR, they were also added as adverse reactions to the company CDS v. 14.0 dated 26 July 2022 (Section 4.8, Appendix A and Appendix B). The changes to the core list of safety concerns and CDS were made based on the summation of data that has accumulated in the surveillance of this issue, including the published incidence and reporting rates from multiple sources with consistent findings.*

**Rapporteur assessment comment:**
Please refer to the assessment of the important identified risk myocarditis and pericarditis in section 2.4.1. of this AR.

<table>
<thead>
<tr>
<th>Risks not categorized as important</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Irritability</strong></td>
</tr>
<tr>
<td>During the reporting period, placebo-controlled safety data from Clinical Trial C4591007 was unblinded for submission to regulatory authorities to support authorisation of vaccination in individuals 6 months to &lt; 5 years of age. Irritability was the most frequently reported systemic event reported within 7 days after each of the 3 doses of BNT162b2 (3 µg) for the 6 months to &lt; 2 years age group. Irritability was reported by 51.2%, 47.4% and 43.6% of participants in the BNT162b2 group and 47.2%, 40.7% and 37.6% in the placebo group, after dose 1, 2 and 3, respectively. Based on these data, irritability was determined to be an adverse reaction for the age group 6 months to &lt; 2 years.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment:**
Please refer regarding irritability to the separate ongoing line extension procedure (EMEA/H/C/005735/X/138) to add a new strength of 3 µg for individuals 6 months to 4 years of age and the RMP (version 5.1) is updated accordingly.

### 2.2.3. Signal evaluation plan for ongoing signals

#### Table 35. Signal Evaluation Plan for Ongoing Signals

<table>
<thead>
<tr>
<th>Signal</th>
<th>Evaluation Plan</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hearing loss</td>
<td>Following enquiry from a competent authority (EMA PRAC and Health Canada) this signal was reopened during the reporting period and is under evaluation at the cut-off date of this PSUR (18 June 2022). The requested cumulative review is in Appendix 6 A 3.</td>
</tr>
</tbody>
</table>

**Rapporteur assessment comment:**
Regarding hearing loss, please refer to the assessment in section 2.2.1 'Post-approval regulatory' above.

### 2.3. Evaluation of risks and new information

**Follow-up questionnaires**

Response to the PRAC request 2 from the 2nd PSUR (procedure EMEA/H/C/PSUSA/00010898/202112):

*The MAH is requested to re-assess the need for continuing the follow-up questionnaires anaphylaxis and VAED/VAERD and provide process data (e.g., response rate, extent of additional information collected) separately for cases reporting anaphylaxis and cases reporting VAED/VAERD, if applicable.*
MAH’s response (Appendix 6A of the PSUR):

As described in PSUR 1 and PSUR 2, the pursuit of additional information for specific adverse events via the use of a Data Capture Aid (DCA) remains limited in 2 regards: the number of reports that can have DCAs sent for follow-up information and the dependency on a response. As previously described, the most common reasons that a DCA is not dispatched include lack of contact details or refused contact and receipt of the report from a Health Authority. Of the adverse event reports received by Pfizer potentially meeting criteria for dispatch of a DCA for follow-up to the reporter from 01 December 2021 to 30 June 2022, 23.5% of the reports had DCAs sent in the pursuit of follow-up information. Despite the limitations, it is the opinion of the MAH that the potential for obtaining useful follow-up information justifies the continued use of the DCAs currently and as vaccine use extends to lower age populations.

Rapporteur assessment comment:

Despite that the MAH did not provide process data separately for cases reporting anaphylaxis and cases reporting VAED/VAERD, MAH’s response is endorsed that the potential for obtaining useful follow-up information justifies the continued use of the follow-up questionnaires anaphylaxis and VAED/VAERD currently and as vaccine use extends to lower age populations.

Evaluation of Important Identified risks

Anaphylaxis

Search criteria - PTs: Anaphylactic reaction; Anaphylactic shock; Anaphylactoid reaction; Anaphylactoid shock

Clinical trial data

- Number of cases: 3 (0.45% of 668 cases of the total CT dataset), compared to 2 cases (0.28%) retrieved in the PSUR #2.
- The investigator and the Sponsor reported that there was not a reasonable possibility that the events anaphylactic reactions in all cases were related to the blinded study vaccine/BNT162b2, or clinical trial procedure. In 2 cases the anaphylaxis reactions were associated with food allergies and in the remaining case anaphylaxis reaction was attributed to another product (etoricoxib).

Post-authorization data

- Number of cases: 1037 (0.20% of 507,683 cases, the total PM dataset), compared to 3507 cases (0.53%) retrieved in the PSUR #2.
- MC cases (690), NMC cases (347).
- Country of incidence (top 10): Japan (184), Germany (158), Australia (113), UK (105), US (59), Poland (48), France (47), New Zealand (41), Philippines (26), and Sweden (24); the remaining 232 cases were distributed among 34 countries.
- Subjects’ gender: female (768), male (219) and unknown (50).
- Subjects’ age in years: n = 949, range: 5 - 99, mean: 40.2, median: 40.0.
- Medical history (n = 422): the most frequently (≥ 10 occurrences) reported medical conditions Asthma (90), Food allergy (83), Drug hypersensitivity (66), Hypersensitivity (55), Hypertension (42), Seasonal allergy (38), Anaphylactic reaction (30), COVID-19 (21), Mite
allergy (20), Allergy to arthropod sting (19), Allergy to animal (14), Dermatitis contact (14), Contrast media allergy (12), Mast cell activation syndrome (12), Multiple allergies (12), Diabetes mellitus (11), Urticaria (11), Allergy to chemicals (10), Anaphylactic shock (10), Migraine (10), Obesity (10), and Rubber sensitivity (10).

- Co suspects (n = 21 cases): Relevant co-suspect vaccines/medications reported more than once were: adalimumab, herbal pollen NOS, JNJ 78436735 (2 each).
- Number of relevant events: 1073.
- Relevant event seriousness: serious (1073).
- Reported relevant PTs: Anaphylactic reaction (802), Anaphylactic shock (238), Anaphylactoid reaction (33).
- Time to event onset (n = 781), range: <24 hours to 365 days, median: 0 days.
- Duration of relevant events (n = 249 out of 1037 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 246 days, median 0 days.
- Relevant event outcome: fatal (8), resolved/resolving (647), resolved with sequelae (23), not resolved (134), unknown (263).
- Of the 433 cases reporting medical history/co-suspects, 324 cases reported relevant medical history/risk factors (e.g., asthma, drug hypersensitivity, food allergies, autoimmune disorders, hypersensitivity, prior anaphylactic reactions) and/or co-suspect (e.g., adalimumab, infliximab, influenza vaccine inact SAG 4V, herbal pollen NOS, JNJ 78436735, immunoglobulin human normal), which may have contributed to the anaphylaxis related events.

Analysis by age group
- Post-marketing:
  - Paediatric (120), Adults (751), Elderly (80) and Unknown (86). No significant difference was observed in the reporting proportion of anaphylaxis relevant PTs between paediatric, adult and elderly populations (0.38% in paediatric vs 0.21% in adults vs 0.14% in elderly).

Analysis by presence of comorbidities
- Number of subjects with comorbidities: 181 (17.5% of the cases reporting anaphylaxis). The reporting proportion of anaphylaxis related events with fatal outcome with comorbid conditions is 1.7% compared to the reporting proportion of 3.3% observed in the individuals without comorbidities. A meaningful comparison is not possible due to the low number of fatal anaphylactic related cases.

Literature
During the reporting interval, there were no new significant data received from literature sources.

O/E analysis
- O/E analysis was performed for Anaphylaxis. An O/E ratio of 2.404 (95% CI 2.353, 2.455) was observed for BNT162b2 compared to the background rate for anaphylaxis cases observed in the US. This rate has steadily declined each reporting period from the 9.47 (95% CI, 8.61,
10.40) first reported in SMSR 2 (through 31 January 2021) and has remained consistent with that reported in the most recent SBSR. The reason for the decline is unknown but could reflect decreased reporting, changes in the accuracy of the exposure estimate, or changes in population being vaccinated.

**MAH’s conclusion**

Based on the interval data, no new significant safety information was identified pertaining to the risk of anaphylaxis with BNT162b2.

This risk is communicated in the BNT162b2 CDS, Section 4.4, Special warnings and precautions for use, which includes information on appropriate action to be taken, as follows: "As with all injectable vaccines, appropriate medical treatment and supervision must always be readily available in case of a rare anaphylactic event following the administration of the vaccine.” This risk is also listed in the CDS Section 4.8, Undesirable effects, Appendix A, Appendix B.

In line with the removal of anaphylaxis from the list of safety concerns in the EU-RMP v. 5.1 submitted on 08 July 2022, the MAH proposes to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period, because anaphylaxis is a known risk of vaccines that is understood by HCPs who administer vaccines and patients and does not considerably impact the benefit/risk profile of the vaccine. Product labelling and standards of medical care during the vaccination procedure provide adequate risk mitigation.

This risk will continue to be monitored through routine pharmacovigilance.

**Rapporteur assessment comment:**

No new important safety information concerning anaphylaxis could be identified from the data in current PSUR. The current risk minimisation measures described in the product information of Comirnaty are considered adequate.

After DLP of this PSUR, the important identified risk of anaphylaxis was removed from the list of safety concerns in RMP version 5.1 (procedure EMEA/H/C/005735/X/0138). Therefore, MAH’s proposal is accepted to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period (4th PSUR), because anaphylaxis is a known risk of vaccines that is adequately being managed by HCPs who administer vaccines and the vaccinees in daily practice.

**Myocarditis and Pericarditis**

There were 8533 potentially relevant cases of Myocarditis and Pericarditis: 5423 cases reported myocarditis and 4156 cases reported pericarditis (in 1046 of these 8533 cases, the subjects developed both myocarditis and pericarditis):

**Myocarditis**

Search criteria: Autoimmune myocarditis; Carditis; Chronic myocarditis; Eosinophilic myocarditis; Giant cell myocarditis; Hypersensitivity myocarditis; Immune-mediated myocarditis; Myopericarditis; Myocarditis.

**Overall - All ages**

- Clinical Trial Data
Number of cases: 1 case of BNT162b2 (0.15% of 668 cases of the total CT dataset), compared to 2 cases (0.3% of 721 cases of the total CT dataset) retrieved in the PSUR #2.

Country of incidence: US.

Subject's gender: Male (1).

Subject's age in years: 43 years.

Medical history: PT Abstains from alcohol, Abstains from recreational drugs, Anxiety, Attention deficit hyperactivity disorder, Clinical trial participant, Dyspepsia, Gastroesophageal reflux disease, Neuralgia, Non-tobacco user, Postural orthostatic tachycardia syndrome, Prophylaxis, Seasonal allergy, Stress, Tachycardia, Thyroiditis, Vasectomy (1 each).

COVID-19 Medical history: COVID-19 (1).

Co suspets: None.

Number of relevant serious events: 1.

Reported relevant PTs: Myocarditis (not related to BNT162b2).

Relevant event outcome: Resolved (1).

Time to onset of relevant events: 98 days after dose 3.

Duration of myocarditis was reported as 2 days.

Post-Authorisation Data

Number of cases: 5422 (1.1% of 507,683 cases of the total PM dataset), compared to 6347 cases (1.0%) retrieved in the PSUR #2.

Country/region of incidence (≥10): Germany (1342), UK (1230), Australia (509), France (344), Taiwan, Province Of China (280), Canada (216), Austria (193), Japan (163), Italy (151), Sweden (119), US (118), New Zealand (107), Greece (77), Israel (64), Finland (51), Spain (45), Netherlands (44), Hong Kong (41), Poland (35), Belgium, Denmark (29 each), Switzerland (25), Norway, Portugal (24 each), Malaysia (22), Ireland (21), Czech Republic (16), Brazil (15), Romania (10). The remaining 78 cases were distributed among 25 countries.

MC (2710), NMC (2712).

Subjects' gender: female (1997), male (3307) and unknown (118).

Subjects' age in years: n = 4981, range: 6-98, mean: 35.3, median: 32.

Medical history (n = 1699): the most frequently (≥50 occurrences) reported medical conditions included Hypertension (214), Asthma (140), Seasonal allergy (130), Tobacco user (96), Drug hypersensitivity (67), Immunodeficiency (64), Obesity (62), Hypothyroidism, Non-tobacco user (59 each), and Food allergy (58).


Number of relevant events: 5458.

Relevant event seriousness: serious (5458).

Reported relevant PTs: Myocarditis (4639), Myopericarditis (697), Carditis (113), Eosinophilic myocarditis (4), Giant cell myocarditis, Hypersensitivity myocarditis (2 each), and Immune-mediated myocarditis (1).

Relevant event outcome: fatal (87), resolved/resolving (1925), resolved with sequelae (160), not resolved (1608), unknown (1682).

Of the 5422 cases, in 1108 cases (20.4% of the cases reporting myocarditis related events) the events were confounded by subject’s relevant medical history (1016 cases; e.g., COVID-19, seasonal allergy, tobacco user, drug hypersensitivity, food allergy, myocarditis, mite allergy, autoimmune thyroiditis, cardiac failure, allergy to animal, overweight, alcohol use, cardiac disorder, pericarditis, allergy to metals, breast cancer, chemotherapy, allergy to plants, dust allergy, rheumatoid arthritis, influenza, myopericarditis, systemic lupus erythematosus, mycotic allergy, radiotherapy, allergy to arthropod sting, allergy to chemicals, Epstein Barr virus infection, autoimmune disorder, Lyme disease, rheumatic disorder) and/or relevant co-suspect/concomitant medications (92 cases; e.g., influenza vaccine, isotretinoin, mesalazine, olanzapine, quetiapine, rituximab, cyclophosphamide, epirubicin, hepatitis B vaccine RHB SAG (yeast), minocycline, norepinephrine, sulfasalazine, zuclopenthixol, COVID-19 vaccine mRNA (mRNA 1273), COVID-19 vaccine NRVV AD (CHADNOX1 NCOV-19), COVID-19 vaccine, pembrolizumab, clozapine, hepatitis A vaccine, influenza vaccine INACT SAG 3V, ipilimumab, JNJ 78436735, nivolumab).

Of the 5422 cases, 236 cases involved elderly (age >70 years) subjects and 61% cases involved male subjects.

Subjects aged less than 5 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

Rapporteur assessment comment:

Comirnaty exposure in persons aged less than 5 years is considered off-label use during the interval period of the current 3rd PSUR.

Subjects aged 5 - 11 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
- Number of cases: 48 cases (0.01% of 507,683 cases of the total PM dataset; 0.6% of the 8375 subjects aged 5-11 years); 10 cases (0.002%) were retrieved in the PSUR #2.

- Country/region of incidence: Australia (15), Canada, Japan (6 each), Italy, Portugal, Spain (3 each), Greece, Taiwan, Province of China (2 each), Austria, Denmark, Finland, France, New Zealand, Philippines, UK, US (1 each).


- Medical history (n = 8): Asthma, Atioventricular block, Attention deficit hyperactivity disorder, Autoimmune thyroiditis, Cardiac failure, Cerebral palsy, Condition aggravated, Dependence on respirator, Ejection fraction decreased, Hypoxic-ischaemic encephalopathy, Intellectual disability, Motor dysfunction, Myocarditis, Neonatal asphyxia, Non-tobacco user, Obesity, Respiratory tract infection, Rhinitis allergic, Type 1 diabetes mellitus (1 each).


- Co-suspect vaccine/medications: None.

- Most frequently co-reported Pts (>5 occurrences): Chest pain (28), Dyspnoea, Pyrexia (10 each), Troponin increased (7), Chest discomfort, Electrocardiogram abnormal, Tachycardia (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 36.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>16</td>
<td>26</td>
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</tr>
<tr>
<td>No</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Relevant Pts*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>13</td>
<td>22</td>
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</tr>
<tr>
<td>Myopericarditis</td>
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<td>Cardiac</td>
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<td>0</td>
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<tr>
<td>Hospitalisation required/prolonged</td>
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<td>11</td>
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</tr>
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<td>14</td>
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<td>Relevant suspect dose</td>
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<td>Dose 2</td>
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</tr>
<tr>
<td>Dose 3</td>
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</table>

<table>
<thead>
<tr>
<th>Time to Onset n=39</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
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<tbody>
<tr>
<td>≤ 24 hours</td>
<td>3</td>
<td>3</td>
<td>0</td>
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<td>1-5 days</td>
<td>7</td>
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<td>6-13 days</td>
<td>6</td>
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<td>14-20 days</td>
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<td>5</td>
<td>4</td>
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</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female</th>
<th>Male</th>
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</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>4</td>
<td>4</td>
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</tr>
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<td>Resolved</td>
<td>4</td>
<td>12</td>
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<td>Resolving</td>
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</tr>
<tr>
<td>Unknown</td>
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<td>4</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event a n=4, median = 1 day</th>
<th>Female</th>
<th>Male</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
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</tr>
<tr>
<td>7-25 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.

b. For those cases where the event resolved.

Fatal myocarditis cases in subjects aged 5-11 years (2 cases, medically confirmed):

- A 6-year-old male subject from Portugal:
- Medical history: Autoimmune thyroiditis, Rhinitis allergic, Type 1 diabetes mellitus.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis, Cardio-respiratory arrest, COVID-19.
- Time to onset (myocarditis): 7 days after dose 1.
- Causes of death: Cardio-respiratory arrest; Myocarditis.
- Autopsy: Results awaited at the time of reporting.

- An 11-year-old female subject from Japan:
  - Co-suspect medications: None.
  - PTs with fatal outcome: Blood pressure decreased, Blood pressure immeasurable, Bradycardia, Cardio-respiratory arrest, Cyanosis, Heart rate decreased, Myocarditis, Respiratory failure.
  - Time to onset (myocarditis): 1 day after dose 2.
  - Causes of death: Blood pressure decreased; Blood pressure immeasurable; Bradycardia; Cardiac failure acute; Cardio-respiratory arrest; Cyanosis; Heart rate decreased; Myocarditis; Respiratory failure.
  - Autopsy: Pleural X-ray was performed as autopsy imaging and did not show abnormal findings.

Rapporteur assessment comment:

During the current reporting period, there were 48 cases reporting myocarditis in children aged 5-11 years compared to 10 myocarditis cases reported in the previous 2nd PSUR. Although the Comirnaty exposure in persons aged 5-11 years is considered increased based on the EU/EEA exposure (current reporting period an estimated 3,569,821 administered doses in 5-9 years versus 391,327 administered doses in 5-9 years in the previous reporting period), worldwide interval exposure in persons aged 5-11 years (or any other age category) is not presented in the PSUR and therefore the relative post-marketing reporting rate of myocarditis cases in persons aged 5-11 years is not known.

There were 2 medically confirmed fatal cases compared to no fatal cases in the previous reporting period. The MAH only briefly described the 2 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 5-11 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. Request for supplementary information.

Subjects aged 12 - 15 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.
• Post-Authorisation Data
  
  o Number of cases: 366 (0.07 % of 507,683 cases of the total PM dataset; 2.7 % of the 13,366 subjects aged 12-15 years), compared to 488 cases (0.07% of all cases in the total PM dataset) retrieved in the PSUR #2.

  o Country/region of incidence (≥10): Taiwan, Province of China (87), Germany (64), UK (30), Australia, Japan (23 each), Canada (18), France (15), Italy (14), Israel, Malaysia (11 each), Hong Kong (10). The remaining 60 cases were distributed among 20 countries.


  o Medical history (n = 72): the most frequently (≥2 occurrence) reported medical conditions included Asthma (7), Attention deficit hyperactivity disorder, Food allergy, Obesity, Seasonal allergy (4 each), Glucose-6-phosphate dehydrogenase deficiency, Hypersensitivity, Migraine, Non-tobacco user, Pericarditis, Rhinitis allergic (3 each), Anxiety, Autism spectrum disorder, Childhood asthma, Cough, Dermatitis atopic, Mite allergy, Pneumonia, Prophylaxis, Tonsillectomy (2 each).


  o Co-suspect vaccine/medications: None.

  o Most frequently co-reported PTs (>5 occurrences): Chest pain (174), Pyrexia (80), Chest discomfort (64), Dyspnoea (60), Headache (39), Palpitations (36), Pericarditis (34), Fatigue (31), Tachycardia (25), Inappropriate schedule of product administration, Troponin increased (23 each), Vomiting (20), Electrocardiogram ST segment elevation (18), Dizziness, Nausea (17 each), Malaise (16), C-reactive protein increased (15), Myalgia, Troponin I increased (14 each), Cough (13), Astenia, Off label use, Pain in extremity (11 each), Blood creatine phosphokinase MB increased, Vaccination site pain (10 each), Chills (9), Blood creatine phosphokinase increased, Diarrhoea, Pain (8 each), Decreased appetite, Immunisation, Pericardial effusion, Syncope (7 each), Arthralgia, Electrocardiogram abnormal, Heart rate increased, Multisystem inflammatory syndrome in children, Nasopharyngitis (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 37.
Table 37. Myocarditis in Subjects aged 12 – 15 Years (N=366)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>42</td>
<td>232</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>19</td>
<td>71</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Myocarditis</td>
<td>56</td>
<td>269</td>
</tr>
<tr>
<td></td>
<td>Myopericarditis</td>
<td>5</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>Extrinsic myocarditis</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Carditis</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>32</td>
<td>226</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>29</td>
<td>77</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>16</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>33</td>
<td>186</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>4</td>
<td>43</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>8</td>
<td>25</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n=274</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 24 hours</td>
<td>2</td>
<td>17</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>29</td>
<td>158</td>
<td>0</td>
</tr>
<tr>
<td>6-13 days</td>
<td>7</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>14-31 days</td>
<td>2</td>
<td>11</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>≥ 31 days</td>
<td>6</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>14</td>
<td>78</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female</th>
<th>Male</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>11</td>
<td>45</td>
<td>0</td>
</tr>
<tr>
<td>Resolved</td>
<td>21</td>
<td>88</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>21</td>
<td>104</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>7</td>
<td>63</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event* n=39, median=7 days</th>
<th>Female</th>
<th>Male</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>3</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>0</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>4</td>
<td>11</td>
<td>0</td>
</tr>
<tr>
<td>26-134 days</td>
<td>2</td>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Fatal myocarditis cases in subjects aged 12-15 years (2 cases, medically confirmed; 1 case non-medically confirmed):

- A 13-year-old male subject from Taiwan, Province of China:
  - Medical history: None.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Cardiac failure, Myocarditis.
  - Time to onset (myocarditis): 69 days after dose 2.
  - Causes of death: Cardiac failure; Myocarditis.
  - Autopsy: Not reported if autopsy was performed.

- A 13-year-old male subject from UK:
  - Medical history: Abdominal pain, Chest pain.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Anuria, Asthenia, Cardiac arrest, Compartment syndrome, Enterovirus infection, Malaise, Multi-organ disorder, Multiple organ dysfunction syndrome, Myocarditis, Pulseless electrical activity, Renal failure, Rhinovirus infection, Ventricular tachycardia.
- Time to onset (myocarditis): 5 days after dose 2.
- Causes of death: Asthenia; Cardiac arrest; Compartment syndrome; Enterovirus infection; Malaise; Multi-organ disorder; Myocarditis; Pulseless electrical activity; Renal failure; Rhinovirus infection; Ventricular tachycardia.
- Autopsy: Not reported if autopsy was performed.

- A 13-year-old female subject from Philippines:
  - Medical history: None.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Myocarditis.
  - Time to onset (myocarditis): 6 days after dose 1.
  - Causes of death: Myocarditis.
  - Autopsy: Adverse event following immunisation.

**Rapporteur assessment comment:**

During the current reporting period, there were 366 cases reporting myocarditis in persons aged 12-15 years compared to 488 myocarditis cases reported in the previous 2nd PSUR. There were 3 fatal cases compared to 3 fatal cases in the previous reporting period.

The MAH only briefly presented the 3 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 12-15 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. **Request for supplementary information**

**Subjects aged 16 - 17 years**

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 345 (0.07 % of 507,683 cases of the total PM dataset; 4.2 % of the 8313 subjects aged 16-17 years), compared to 470 cases (0.07%) retrieved in the PSUR #2.
  - Country of incidence (≥10): Germany (88), Taiwan, Province of China (55), UK (29), Australia (18), Austria (17), France, Poland (14 each), Italy, Japan (12 each), Greece (11). The remaining 75 cases were distributed among 26 countries. Subjects’ age in years: n = 345, range: 16 -17, mean: 16.6, median: 17.0.
  - Medical history (n = 71): the most frequently (>2 occurrence) reported medical conditions included Seasonal allergy (9), Asthma, Obesity (4 each), Food allergy, Mite allergy, Nasopharyngitis, Overweight (3 each).
Co suspect vaccine/medications: Clonazepam, infliximab, and levomethadone (1 each).

Most frequently co-reported PTs (>5 occurrences): Chest pain (147), Pyrexia (72), Dyspnoea (53), Chest discomfort (43), Palpitations (34), Tachycardia (33), Fatigue (29), Headache (27), Inappropriate schedule of product administration (26), Pericarditis (24), Troponin increased (23), Dizziness, Vomiting (18 each), Malaise, Nausea (17 each), Asthenia (12), Chills, Immunisation, Off label use (10 each), Arrhythmia, Blood creatine phosphokinase increased, C-reactive protein increased, Electrocardiogram ST segment elevation, Pain in extremity, Pericardial effusion, Syncope, Troponin I increased (9 each), Cough (8), Myocardial necrosis marker increased, Pain, Troponin T increased (7 each), Angina pectoris, Back pain, Blood creatine phosphokinase MB increased, Diarrhoea, Electrocardiogram abnormal, Lethargy (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 38.

### Table 38. Myocarditis in Subjects aged 16 – 17 Years (N=345)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>28</td>
<td>202</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>23</td>
<td>90</td>
<td>0</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>48</td>
<td>243</td>
<td>2</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>4</td>
<td>47</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>27</td>
<td>223</td>
<td>0</td>
</tr>
<tr>
<td>No</td>
<td>24</td>
<td>69</td>
<td>2</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>13</td>
<td>39</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>23</td>
<td>154</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>9</td>
<td>65</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>6</td>
<td>34</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤24 hours</td>
<td>7</td>
<td>18</td>
<td>1</td>
</tr>
<tr>
<td>1-5 days</td>
<td>13</td>
<td>149</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>6</td>
<td>19</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>1</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>32-90 days</td>
<td>4</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>91-150 days</td>
<td>1</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>18</td>
<td>81</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>11</td>
<td>60</td>
<td>0</td>
</tr>
<tr>
<td>Resolved</td>
<td>9</td>
<td>78</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>11</td>
<td>80</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>28</td>
<td>75</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>1</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>2</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>26-68 days</td>
<td>0</td>
<td>4</td>
<td>0</td>
</tr>
</tbody>
</table>

### Rapporteur assessment comment:

During the current reporting period, there were 345 cases reporting myocarditis in persons aged 16-17 years compared to 470 myocarditis cases reported in the previous 2nd PSUR.

There were no fatal cases compared to 2 fatal cases in the previous reporting period.
Subjects aged 18 - 24 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 968 (0.2% of 507,683 cases of the total PM dataset, 0.18% of the 382,933 subjects aged 18-24 years), compared to 1,187 cases (2.3%) retrieved in the PSUR #2.
  - Country of Incidence (≥10): Germany (289), UK (114), France (105), Australia (99), Taiwan, Province of China (46), Italy (43), Austria (38), Sweden (32), Japan (25), US (18), Greece, New Zealand (16 each), Israel (14), Canada, Spain (13 each), Finland (11), Denmark (10). The remaining 66 cases were distributed among 18 countries.
  - Medical history (n = 237): the most frequently (>2 occurrence) reported medical conditions included PT Tobacco user (28), Asthma (23), Seasonal allergy (16), Myocarditis (12), Hypertension, Non-tobacco user (11 each), Immunodeficiency, Obesity (9 each), Attention deficit hyperactivity disorder (8), Hypersensitivity, Nicotine dependence (7 each), Alcohol use, Contraception, Mite allergy (6 each), Acne, Crohn's disease, Drug hypersensitivity, Migraine, Overweight, Substance use (5 each), Anaemia, Autism spectrum disorder, Epstein-Barr virus infection (4 each), Appendicectomy, Chest pain, Food allergy, Hypothyroidism, Oral contraception, Pericarditis, Pharyngitis, Psoriasis, Syncope, Wisdom teeth removal (3 each).
  - COVID-19 Medical history (n = 52): COVID-19 (26), Suspected COVID-19 (22), SARS-CoV-2 test positive (3), and Asymptomatic COVID-19 (1).
  - Co suspect vaccine/medications: Drug COVID-19 vaccine, COVID-19 vaccine MRNA (MRNA 1273), COVID-19 vaccine NVRV AD (CHADOX1 NCOV-19), insulin, levothyroxine, and zuclopenthixol (1 each).
  - Most frequently co-reported PTs (>5 occurrences): Chest pain (354), Dyspnoea (168), Pyrexia (134), Pericarditis (116), Fatigue (115), Palpitations (112), Chest discomfort (104), Troponin increased (75), Tachycardia (74), Headache (61), Inappropriate schedule of product administration (59), Off label use (48), Dizziness, Immunisation (46 each), Interchange of vaccine products (38), Asthenia (33), Chills (32), Malaise, Myalgia (29 each), Angina pectoris (28), Pain, Syncope (25 each), Nausea (23), Arrhythmia, Dyspnoea exertional (21 each), Pericardial effusion (20), Influenza like illness (19), Cough, Vomiting (18 each), Pain in extremity (17), C-reactive protein increased, Heart rate increased (15 each), Electrocardiogram ST segment elevation, Hyperhidrosis, Lethargy (14 each), Electrocardiogram abnormal (13), Diarrhoea (12), Oropharyngeal pain (11), Arthralgia, Blood creatine phosphokinase Increased, COVID-19, Myocardial infarction, Myocardial necrosis marker increased, Troponin I increased, Troponin T increased (10 each), Acute myocardial infarction, Paraesthesia (9 each), Abdominal pain, Abdominal pain upper, Back pain, Cardiac failure, Drug ineffective, Feeling abnormal, Inflammation, Sinus tachycardia (8 each), Hypertension, Hypoesthesia, Limb discomfort, Lymphadenopathy, Night sweats, Pulmonary embolism, Vaccination site pain, Ventricular hypokinesia (7 each), Costochondritis,
Feeling hot, Incorrect route of product administration, Insomnia, Left ventricular dysfunction, Loss of consciousness, Somnolence (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 39.

Table 39. Myocarditis in Subjects aged 18 – 24 Years (N=68)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>110</td>
<td>471</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>97</td>
<td>283</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td></td>
<td>179</td>
<td>600</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td></td>
<td>23</td>
<td>147</td>
</tr>
<tr>
<td>Carditis</td>
<td></td>
<td>5</td>
<td>14</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td></td>
<td>108</td>
<td>484</td>
</tr>
<tr>
<td>No</td>
<td></td>
<td>99</td>
<td>271</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Dose 1</td>
<td></td>
<td>48</td>
<td>145</td>
</tr>
<tr>
<td>Dose 2</td>
<td></td>
<td>84</td>
<td>316</td>
</tr>
<tr>
<td>Dose 3</td>
<td></td>
<td>55</td>
<td>235</td>
</tr>
<tr>
<td>Unknown</td>
<td></td>
<td>22</td>
<td>60</td>
</tr>
</tbody>
</table>

| Time to Onset: %724          |                     |                   |                      |                       |
| <24 hours                    |                     | 18                | 39                   | 1                     |
| 1-5 days                     |                     | 73                | 362                  | 2                     |
| 6-13 days                    |                     | 15                | 57                   | 1                     |
| 14-21 days                   |                     | 9                 | 43                   | 0                     |
| 22-31 days                   |                     | 4                 | 12                   | 0                     |
| 32-60 days                   |                     | 6                 | 35                   | 0                     |
| 61-220 days                  |                     | 13                | 34                   | 0                     |
| Unknown                      |                     | 69                | 179                  | 3                     |

| Event Outcome                |                     |                   |                      |                       |
| Fatal                        |                     | 0                 | 4                    | 0                     |
| Not resolved                 |                     | 66                | 234                  | 1                     |
| Resolved                     |                     | 25                | 139                  | 1                     |
| Resolved with sequelae       |                     | 10                | 27                   | 1                     |
| Resolving                    |                     | 52                | 195                  | 1                     |
| Unknown                      |                     | 54                | 162                  | 3                     |

| Duration of event* n=71, median=7 days |                     |                   |                      |                       |
| Up to 3 days                 |                     | 4                 | 14                   | 0                     |
| 4-6 days                     |                     | 4                 | 9                    | 0                     |
| 7-25 days                    |                     | 1                 | 18                   | 0                     |
| 26-195 days                  |                     | 2                 | 18                   | 1                     |

a. All serious occurrences.
b. For those cases where the event resolved

Fatal myocarditis cases in subjects aged 18-24 years (4 cases, medically confirmed)

- A 23-year-old male subject from Estonia:
  - Medical history: Non-tobacco user.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Circulatory collapse, Endocarditis, Myocarditis, Sudden cardiac death.
  - Time to onset (myocarditis): unknown days after dose 2.
  - Causes of death: Circulatory collapse; Endocarditis; Myocarditis; Sudden cardiac death.
  - Autopsy: Autopsy was performed, results were not provided at the time of reporting.

- A 20-year-old male subject from Taiwan, Province of China:
  - Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): 20 days after dose 1.
- Cause of death: Myocarditis.
- Autopsy: Autopsy results showed cause of death as myocarditis.

- A 19-year-old male subject from Japan:
  - Co-suspect medications: None.
  - PTs with fatal outcome: Arrhythmia, Hernia, Hypoxia, Loss of consciousness, Myocardial necrosis, Myocardial necrosis marker increased, Myocarditis, Sudden death, Ventricular hypokinesia.
  - Time to onset (myocarditis): 3 days after dose 3.
  - Causes of death: Arrhythmia; Hernia; Hypoxia; Loss of consciousness; Myocardial necrosis; Myocardial necrosis marker increased; Myocarditis; Sudden death; Ventricular hypokinesia.
  - Autopsy: The autopsy revealed extensive necrosis of the left ventricular myocardium (myocardial necrosis); myocarditis/fulminant myocarditis.

- A 23-year-old male subject from Germany:
  - Medical history: Hypertension, Obesity.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Death, Myocarditis.
  - Time to onset (myocarditis): 16 days after dose 3.
  - Cause of death: Myocarditis.
  - Autopsy: Information not available.

Rapporteur assessment comment:

During the current reporting period, there were 968 cases reporting myocarditis in persons aged 18-24 years compared to 1,187 myocarditis cases reported in the previous 2nd PSUR. There were 4 fatal cases compared to 2 fatal cases in the previous reporting period.

The MAH only briefly presented the 4 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 18-24 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. Request for supplementary information

Subjects aged 25 - 29 years
• Clinical Trial Data
  o Number of cases: none. No cases were retrieved in the PSUR #2.

• Post-Authorisation Data
  o Number of cases: 519 (0.1% of 507,683 cases of the total PM dataset, 1.2 % of the 43518 subjects aged 25-29 years), compared to 589 cases (0.09%) retrieved in the PSUR #2.
  o Country of incidence (≥10): Germany (150), UK (113), Australia (54), France (39), Austria (27), Sweden, Taiwan, Province of China (17 each), Japan (14), Italy (13), New Zealand (11). The remaining 64 cases were distributed among 25 countries.
  o Subjects’ age in years: n = 519, range: 25-29, mean: 27.1, median: 27.
  o Medical history (n = 141): the most frequently (>2 occurrence) reported medical conditions included Seasonal allergy (20), Asthma (14), Tobacco user (13), Food allergy (10), Hypertension, Mite allergy (8 each), Allergy to animal, Chest pain (7 each), Non-tobacco user (6), Hypersensitivity, Hypothyroidism (5 each), Anxiety, Depression, Drug hypersensitivity, Migraine, Myocarditis, Steroid therapy (4 each), Autoimmune thyroiditis, Contraception, Gastrooesophageal reflux disease, Polycystic ovaries (3 each).
  o COVID-19 Medical history (n = 43): COVID-19 (22), Suspected COVID-19 (21), Post-acute COVID-19 syndrome (2), and SARS-CoV-2 test positive (1).
  o Co suspect vaccine/medications (n = 8): COVID-19 vaccine mRNA (MRNA 1273) (3), adalimumab, fluticasone, influenza vaccine, levothyroxine, and methylphenidate (1 each).
  o Most frequently co-reported PTs (>5 occurrences): Chest pain (217), Dyspnoea (153), Palpitations (122), Fatigue (110), Tachycardia (98), Pericarditis (88), Chest discomfort, Pyrexia (71 each), Headache (47), Immunisation (42), Dizziness (39), Arrhythmia (31), Myalgia (25), Off label use, Pain in extremity (24 each), Interchange of vaccine products, Troponin increased (23 each), Heart rate increased, Inappropriate schedule of product administration, Malaise, Pain (22 each), Angina pectoris (21), Lymphadenopathy (18), Asthenia, Nausea (16 each), Chills, Pericardial effusion (15 each), Influenza like illness, Paraesthesia, Syncope, Vaccination site pain (14 each), Arthralgia (13), Lethargy (10), COVID-19, Dyspnoea exertional, Influenza, Migraine, Vomiting (9 each), Back pain, Cardiac flutter, Heart rate irregular, Tremor (8 each), Abdominal pain upper, Blood pressure increased, Cardiac disorder, Diarrhoea, Extrasystoles, Hyperhidrosis, Peripheral swelling (7 each), Abdominal pain, Cough, Electrocardiogram ST segment elevation, Feeling abnormal, Hypertension, Inflammation, Myocardial infarction, Rash, Sleep disorder (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 40.
### Table 40. Myocarditis in Subjects aged 25 – 29 Years (N=519)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>69</td>
<td>169</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>110</td>
<td>168</td>
</tr>
<tr>
<td>Relevant PTs*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>150</td>
<td>285</td>
<td>3</td>
</tr>
<tr>
<td>Myocardiomyopathy</td>
<td>27</td>
<td>48</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>4</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>58</td>
<td>171</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>121</td>
<td>166</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>62</td>
<td>93</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>43</td>
<td>108</td>
<td>1</td>
</tr>
<tr>
<td>Dose 3</td>
<td>32</td>
<td>103</td>
<td>1</td>
</tr>
<tr>
<td>Dose 4</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>22</td>
<td>32</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset [a]</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 24 hours</td>
<td>16</td>
<td>23</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>49</td>
<td>131</td>
<td>0</td>
</tr>
<tr>
<td>6-13 days</td>
<td>12</td>
<td>27</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>15</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>3</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>5</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>61-360 days</td>
<td>3</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>78</td>
<td>166</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>70</td>
<td>108</td>
<td>0</td>
</tr>
<tr>
<td>Resolved</td>
<td>21</td>
<td>41</td>
<td>0</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>9</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>27</td>
<td>82</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>52</td>
<td>92</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event [b]</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 1 day</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>1</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>4</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>26-259 days</td>
<td>10</td>
<td>9</td>
<td>0</td>
</tr>
</tbody>
</table>

---

[a] All serious occurrences.
[b] For those cases where the event resolved.

Fatal myocarditis cases in subjects aged 25-29 years (2 cases, medically confirmed; 3 cases non-medically confirmed):

- A 29-year-old male subject from Japan:
  - Medical history: Hepatic steatosis.
  - Co-suspect medications: COVID-19 vaccine mRNA (MRNA 1273).
  - PTs with fatal outcome: Arrhythmia, Myocarditis.
  - Time to onset (myocarditis): unknown days after dose 2.
  - Causes of death: Arrhythmia; Myocarditis.
  - Autopsy: Autopsy revealed arrhythmia

- A 27-year-old male subject from Brazil:
  - Medical history: None.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use, Chest pain.
  - Time to onset (myocarditis): 10 days after dose 3.
- Causes of death: Myocarditis.
- Autopsy: Not reported if autopsy was performed

- A 26-year-old male subject from US:
  - Medical history: Aneurysm, Surgery, Vein of Galen aneurysmal malformation.
  - Co-suspect medications: Influenza vaccine.
  - PTs with fatal outcome: Myocarditis, Arrhythmia, Inflammation, Left ventricular dysfunction.
  - Time to onset (myocarditis): 4 days after dose 3.
  - Causes of death: Arrhythmia; Inflammation; Left ventricular dysfunction; Myocarditis.
  - Autopsy: Autopsy results showed myocarditis.

- A 26-year-old female subject from Germany:
  - Medical history: None.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use.
  - Time to onset (myocarditis): unknown days after dose 2.
  - Causes of death: Myocarditis.
  - Autopsy: Autopsy results showed myocarditis.

- A 27-year-old female subject from Germany:
  - Medical history: None.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use.
  - Time to onset (myocarditis): unknown days after dose 2.
  - Causes of death: Myocarditis.
  - Autopsy: Autopsy results showed myocarditis.

**Rapporteur assessment comment:**

During the current reporting period, there were 519 cases reporting myocarditis in persons aged 25-29 years compared to 589 myocarditis cases reported in the previous 2nd PSUR. There were 5 fatal cases compared to 7 fatal cases in the previous reporting period.

The MAH only briefly presented the 5 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 25-29 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. **Request for supplementary information**
Subjects aged 30 - 39 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 983 (0.2 % of 507,683 cases of the total PM dataset, 1.0 % of the 97870 subjects aged 30-39), compared to 995 cases (0.15%) retrieved in the PSUR #2.
  - Country of Incidence (≥10): UK (310), Germany (247), Australia (114), France (51), Austria (35), Taiwan, Province of China (33), New Zealand, Sweden (18 each), Italy, US (17 each), Finland (13), Canada, Greece (12 each), Japan (11), and Belgium (10). The remaining 65 cases were distributed among 23 countries.
  - Subjects' age in years: n = 983, range: 30-39, mean: 34.3, median: 34.
  - Medical history (n = 290): the most frequently (≥2 occurrence) reported medical conditions included Asthma (27), Seasonal allergy (26), Hypothyroidism (18), Tobacco user (17), Immunodeficiency (14), Drug hypersensitivity (13), Hypertension, Migraine, Myocarditis, Non-tobacco user (12 each), Food allergy (11), Breast feeding, Clinical trial participant (10 each), Diabetes mellitus, Dyspnoea, Obesity, Pregnancy (9 each), Autoimmune thyroiditis, Steroid therapy (6 each), Alcohol use, Colitis ulcerative, Dust allergy, Fibromyalgia, Histamine intolerance, Hyperhidrosis, Malaise, Pain, Pericarditis (5 each), Chest pain, Coeliac disease, Depression, Headache, Lymphadenopathy, Mast cell activation syndrome, Mite allergy, Pneumonia, Polycystic ovaries, Post viral fatigue syndrome (4 each), Allergy to animal, Allergy to metals, Cardiac disorder, Crohn's disease, Drug intolerance, Fatigue, Gastroesophageal reflux disease, Hypersensitivity, Hypophosphataemia, Lactose intolerance, Multiple sclerosis, Muscular weakness, Myotic allergy, Myocardial infarction, Nicotine dependence, Osteoporosis, Pancreatic failure, Postural orthostatic tachycardia syndrome, Pulmonary embolism, Small fibre neuropathy (3 each).
  - COVID-19 Medical history (n = 82): COVID-19 (41), Suspected COVID-19 (39), Post-acute COVID-19 syndrome, SARS-CoV-2 test positive (1 each).
  - Co suspect vaccine/medications: Drug COVID-19 vaccine mRNA (MRNA 1273) (3), Amoxicillin, clozapine, colchicine, COVID-19 VACCINE NRVV AD (CHADOX1 NCOV-19), ipilimumab, losartan, nivolumab, propranolol (1 each).
  - Most frequently co-reported PTs (≥5 occurrences): Chest pain (383), Dyspnoea (294), Palpitations (284), Fatigue (276), Pericarditis (240), Tachycardia (215), Pyrexia (128), Chest discomfort (110), Headache (104), Immunisation (100), Dizziness (92), Off label use (78), Inappropriate schedule of product administration (60), Interchange of vaccine products (58), Arrhythmia (54), Pain in extremity (51), Heart rate increased (48), Malaise, Myalgia (46 each), Pain (44), Asthenia (40), Syncope (39), Paraesthesia (38), COVID-19 (36), Drug ineffective (35), Angina pectoris, Troponin increased (34 each), Arthralgia, Hypoaesthesia (33 each), Chills, Nausea (32 each), Hyperhidrosis, Lymphadenopathy, Vomiting (25 each), Dyspnoea exertional (24), Cardiac flutter (23), Vaccination site pain (22), Cough, Feeling abnormal, Pericardial effusion (21 each), Exercise tolerance decreased (20), Discomfort, Influenza like illness (19 each), Anxiety, Back pain, Hypertension (18 each), Diarrhoea (16), Heavy menstrual
bleeding, Insomnia, Neck pain (15 each), Burning sensation (14), Hypotension, Loss of personal independence in daily activities, Oropharyngeal pain (13 each), Heart rate irregular, Menstruation irregular, Presyncope, Product use issue (12 each), Cardiac discomfort, Condition aggravated, Extrasystoles, Inflammation, Lethargy, Muscle twitching, Myocardial infarction, Pulmonary oedema, Rash (11 each), Cardiac disorder, Cardiomegaly, Disturbance in attention, Tinnitus (10 each), Atrial fibrillation, Cardiac failure, Electrocardiogram abnormal, Maternal exposure during pregnancy, Migraine, Muscular weakness, Panic attack, Pruritus, Somnolence, Supraventricular tachycardia, Thrombosis, Tremor (9 each), Abdominal pain upper, Fibrin D dimer increased, Influenza, Limb discomfort, Musculoskeletal stiffness, Night sweats, Pleural effusion, Vision blurred (8 each), Abdominal pain, Amenorrhea, Body temperature increased, Ejection fraction decreased, Heart rate decreased, Loss of consciousness, Muscle spasms, Sleep disorder, Suspected COVID-19, Ventricular extrasystoles (7 each), Asthma, Blood pressure increased, Cardiac arrest, Cardiovascular disorder, Congestive cardiomyopathy, Eczema, Feeling cold, Gait disturbance, Haemorrhage, Heart rate, Illness, Menstrual disorder, Nasopharyngitis, Pulmonary embolism (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 41.

Table 41. Myocarditis in Subjects aged 30 – 39 Years (N=983)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Medically Confirmed</strong></td>
<td>Yes</td>
<td>125</td>
<td>263</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>276</td>
<td>305</td>
</tr>
<tr>
<td><strong>Relevant PT</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>363</td>
<td>505</td>
<td>12</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>27</td>
<td>51</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>13</td>
<td>13</td>
<td>2</td>
</tr>
<tr>
<td>Immune-mediated myocarditis</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><strong>Hospitalisation required/prolonged</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>104</td>
<td>233</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>297</td>
<td>335</td>
<td>12</td>
</tr>
<tr>
<td><strong>Relevant suspect dose</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>132</td>
<td>178</td>
<td>7</td>
</tr>
<tr>
<td>Dose 2</td>
<td>116</td>
<td>204</td>
<td>2</td>
</tr>
<tr>
<td>Dose 3</td>
<td>111</td>
<td>143</td>
<td>3</td>
</tr>
<tr>
<td>Dose 4</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>42</td>
<td>42</td>
<td>7</td>
</tr>
<tr>
<td><strong>Time to Onset</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n=549</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;24 hours</td>
<td>23</td>
<td>31</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>78</td>
<td>156</td>
<td>2</td>
</tr>
<tr>
<td>6-13 days</td>
<td>283</td>
<td>70</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>16</td>
<td>31</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>17</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>12</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>61-449 days</td>
<td>12</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>218</td>
<td>210</td>
<td>12</td>
</tr>
<tr>
<td><strong>Event Outcome</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>4</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>123</td>
<td>209</td>
<td>4</td>
</tr>
<tr>
<td>Resolved</td>
<td>43</td>
<td>75</td>
<td>3</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>11</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>70</td>
<td>98</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>153</td>
<td>171</td>
<td>7</td>
</tr>
<tr>
<td><strong>Duration of event</strong></td>
<td>n=51, median=20 days</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Up to 3 days</td>
<td>2</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>2</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>6</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>26-128 days</td>
<td>10</td>
<td>11</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Fatal myocarditis cases in subjects aged 30-39 years (4 cases, medically confirmed; 1 case non-medically confirmed):

- A 36-year-old male subject from Japan:
Medical history: None.

Co-suspect medications: None.

PTs with fatal outcome: Cardio-respiratory arrest, Myocarditis.

Time to onset (myocarditis): 68 days after unknown dose.

Causes of death: Cardio-respiratory arrest; Myocarditis.

Autopsy: Autopsy revealed myocarditis and cardio-respiratory arrest.

• A 33-year-old female subject from Germany:

Medical history: None.

Co-suspect medications: None.

PTs with fatal outcome: Abdominal pain, Arrhythmia, Cardiac arrest, Chest pain, Circulatory collapse, Myocarditis, Resuscitation.

Time to onset (myocarditis): 20 days after dose 1.

Causes of death: Abdominal pain; Arrhythmia; Cardiac arrest; Chest pain; Circulatory collapse; Myocarditis.

Autopsy: Autopsy information was not reported.

• A 34-year-old male subject from UK:

Medical history: Dyspnoea, Malaise.

Co-suspect medications: None.

PTs with fatal outcome: Arrhythmia, Cardiac arrest, Cardiogenic shock, Circulatory collapse, Dyspnoea, Hypertension, Hypoxia, Left ventricular dysfunction, Myocarditis, Pulmonary oedema, Syncope.

Time to onset (myocarditis): Unknown days after unknown dose.

Causes of death: Arrhythmia; Cardiac arrest; Cardiogenic shock; Circulatory collapse; Dyspnoea; Hypertension; Hypoxia; Left ventricular dysfunction; Pulmonary oedema; Syncope.

Autopsy: Autopsy revealed cause of death as myocarditis.

• A 36-year-old female subject from UK:

Medical history: Depressed mood, Familial risk factor, Perinatal depression, Pregnancy, Tobacco user.

Co-suspect medications: None.

PTs with fatal outcome: Hypoesthesia, Menstruation Irregular, Myocardial injury, Myocarditis, Myopericarditis, Neck pain, Pain in extremity, Pain in jaw, Paraesthesia, Pleural effusion, Thrombosis, Vaccination site pain.

Time to onset (myocarditis and myopericarditis): Unknown duration after first dose.

Causes of death: COVID-19 Immunisation; Myocarditis.
o Autopsy: Autopsy revealed extensive and severe bilateral lung congestion but no evidence of ischemic, hypertensive or valvular heart disease. No evidence of subarachnoid haemorrhage was present. COVID-19 swabs were negative. Histology showed a single focus of myocarditis, with extensive lung congestion suggestive of sudden cardiac death and smoking related changes.

- A 38-year-old female subject from Finland:
  o Medical history: Cerebral palsy.
  o Co-suspect medications: None.
  o PTs with fatal outcome: Back pain, Diarrhoea, Dyspepsia, Myocarditis, Pain, Pain in extremity.
  o Time to onset (myocarditis): 41 days after dose 3.
  o Causes of death: Myocarditis.
  o Autopsy: Autopsy revealed cause of death as myocarditis.

**Rapporteur assessment comment:**

During the current reporting period, there were 983 cases reporting myocarditis in persons aged 30-39 years compared to 995 myocarditis cases reported in the previous 2nd PSUR. There were 4 fatal cases compared to 7 fatal cases in the previous reporting period.

The MAH only briefly presented the 4 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 30-39 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. Request for supplementary information.

**Subjects aged ≥40 years**

- Clinical Trial Data
  o Number of cases: 1 case of BNT162b2 (0.15% of 668 cases of the total CT dataset); 1 case (0.14%) was retrieved in the PSUR #2. Please see above the “Overall- All Ages” subsection for complete details.

- Post-Authorisation Data
  o Number of cases: 1752 (0.3% of 507,683 cases of the total PM dataset, 0.7% of the 236404 subjects ≥ 40 years), compared to 1876 cases (0.3%) retrieved in the PSUR #2.
  o Country of incidence (≥10): Germany (472), UK (464), Australia (168), France (116), Austria (66), Japan (58), New Zealand (53), Italy (41), Taiwan, Province of China (39), Canada (38), Sweden (37), Greece, US (25 each), Norway (16), Netherlands (15), Finland (14), Israel (12), Spain (11), Belgium, Denmark (10 each). The remaining 62 cases were distributed among 21 countries.
Medical history (n = 754): the most frequently (>5 occurrences) reported medical conditions included Hypertension (164), Seasonal allergy (53), Asthma (50), Drug hypersensitivity (35), Immunodeficiency (34), Obesity (32), Hypothyroidism, Tobacco user (30 each), Atrial fibrillation (29), Diabetes mellitus (28), Cardiac failure (26), Dyslipidaemia (25), Food allergy, Non-tobacco user, Type 2 diabetes mellitus (23 each), Hypersensitivity (21), Gastroesophageal reflux disease (18), Anxiety, Depression (17 each), Clinical trial participant (16), Autoimmune thyroiditis, Breast cancer, Chronic obstructive pulmonary disease, Hyperlipidaemia (15 each), Migraine, Myocarditis (14 each), Coronary artery disease (13), Cardiac disorder, Chemotherapy, Ex-tobacco user, Hypercholesterolaemia, Overweight (12 each), Myocardial infarction, Sleep apnoea syndrome, Thyroidectomy, Tobacco abuse (11 each), Allergy to animal, Allergy to metals, Fibromyalgia, Rubber sensitivity (10 each), Appendectomy, Arteriosclerosis, Fatigue, Interchange of vaccine products, Rheumatoid arthritis (9 each), Alcohol use, Menopause, Mite allergy, Osteoporosis, Radiotheraphy, Systemic lupus erythematosus (8 each), Blood cholesterol increased, Cardiac ablation, Dyspnoea, Gout, Hysterectomy, Mitral valve incompetence, Nasopharyngitis, Pulmonary embolism, Steroid therapy, Surgery (7 each), Abstains from alcohol, Allergy to arthropod sting, Allergy to plants, Arrhythmia, Arthritis, Blood cholesterol abnormal, Cerebrovascular accident, Cholecystectomy, Chronic kidney disease, Colitis ulcerative, Hormone replacement therapy, Inflammatory bowel disease, Influenza, Insomnia, Neoplasm, Nicotine dependence, Osteoarthritis, Supraventricular tachycardia, Tachycardia (6 each).


Co suspect vaccine/medications: Influenza vaccine (3), pembrolizumab (2), adalimumab, cisplatin, COVID-19 vaccine, COVID-19 vaccine mRNA (MRNA 1273), COVID-19 vaccine NRVV AD (CHADOX1 NC0V-19), gabapentin, glyceryl trinitrate, hepatitis A vaccine, ibuprofen, influenza vaccine INACT 3V, paracetamol, risankizumab, rivaroxaban, vinorelbine, vitamins NOS (1 each).

Most frequently co-reported PTs (>5 occurrences): Chest pain (572), Dyspnoea (509), Fatigue (493), Palpitations (463), Pericarditis (407), Tachycardia (339), Off label use (285), Interchange of vaccine products (266), Immunisation (264), Pyrexia (235), Headache (189), Chest discomfort (180), Dizziness (174), Arrhythmia (139), Asthenia (101), Malaise, Syncope (89 each), Inappropriate schedule of product administration, Pain in extremity (88 each), Nausea (86), Pain (83), Angina pectoris (73), Cardiac failure (71), Chills (67), Myalgia (66), Arthralgia (64), Pericardial effusion (61), Heart rate increased (58), Dyspnoea exertional, Troponin increased (57 each), Atrial fibrillation (53), Hyperhidrosis (52), Myocardial infarction (50), Cough (49), Back pain (45), Hypertension (43), Paraesthesia (42), Vomiting (40), Diarrhoea, Lethargy (39 each), COVID-19, Lymphadenopathy (35 each), Cardiac flutter, Vaccination site pain (33 each), Exsasystoles, Influenza like illness (32 each), Cardiac disorder, Hypoaesthesia (30 each), Decreased appetite (28), Drug ineffective, Insomnia (27 each), Thrombosis (26), Abdominal pain upper, Blood pressure increased (25 each), Cardiomyopathy, Condition aggravated, C-reactive protein increased, Exercise tolerance decreased (24 each), Anxiety, Feeling abnormal, Neck pain (23 each), Electrocardiogram abnormal, Somnolence, Vertigo (21 each), Abdominal pain, Acute myocardial infarction, Cardiac discomfort, Inflammation, Pulmonary embolism, Tremor
(20 each), Cardiac arrest, Gait disturbance, Muscular weakness, Ventricular extrasystoles (19 each), Hypotension (18), Cerebrovascular accident, Limb discomfort, Rash (17 each), Cardiomegaly, N-terminal prohormone brain natriuretic peptide increased, Peripheral swelling, Swelling, Ventricular tachycardia, Vision blurred (16 each), Breast pain, Bundle branch block left, Dyspepsia, Heart rate irregular, Impaired work ability, Musculoskeletal chest pain, Pneumonia, Presyncope (15 each), Axillary pain, Congestive cardiomyopathy, Coronary artery disease, Discomfort, Feeling hot, Influenza, Oedema peripheral, Pulmonary oedema, Tinnitus, Troponin T increased (14 each), Disturbance in attention, Ejection fraction decreased, Loss of personal independence in daily activities, Oedema, Pain in jaw, Sinus tachycardia (13 each), Acute coronary syndrome, Blood creatine phosphokinase increased, Feeling cold, Illness, Left ventricular dysfunction, Oropharyngeal pain, Performance status decreased, Suspected COVID-19, Weight decreased (12 each), Atrial flutter, Atrioventricular block, Migraine, Musculoskeletal pain, Pleural effusion (11 each), Cardiogenic shock, Fibrin D dimer increased, Heart rate abnormal, Joint swelling, Memory Impairment, Supraventricular tachycardia, Ventricular hypokinesia (10 each), Bradycardia, Confusional state, Electrocardiogram ST segment elevation, Feeling of body temperature change, Head discomfort, Lymph node pain, Mitral valve incompetence, Muscle spasms, Muscle twitching, Myositis, Orthopnoea, Sleep disorder, Stress, Throat tightness (9 each), Acute kidney injury, Blood pressure decreased, Burning sensation, Cold sweat, Cyanosis, Death, Depression, General physical health deterioration, Heavy menstrual bleeding, Hypokinesia, Left ventricular failure, Nasopharyngitis, Respiratory failure, Transient ischaemic attack, Urticaria, Wheezing (8 each), Abdominal discomfort, Amnesia, Arthritis, Brain natriuretic peptide increased, Bronchospasm, Cardiac-arrestive arrest, Disease recurrence, Ear pain, Echocardiogram abnormal, Electrocardiogram ST segment depression, Herpes zoster, Hypersensitivity, Loss of consciousness, Menstrual disorder, Myocardial necrosis marker increased, Pallor, Thrombocytopenia, Visual Impairment (7 each), Blood pressure abnormal, Body temperature increased, Bronchitis, Cardiac dysfunction, Cardiac failure acute, Depressed level of consciousness, Dysgeusia, Fall, Heart rate decreased, Mobility decreased, Night sweats, Pleuritic pain, Pruritus, Sepsis, Vaccination failure, Vaccination site swelling, Ventricular arrhythmia (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 42.
Table 42. Myocarditis in Subjects aged ≥40 Years (N=1752)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>357</td>
<td>374</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>565</td>
<td>437</td>
</tr>
<tr>
<td>Relevant PIs*</td>
<td>Myocarditis</td>
<td>791</td>
<td>716</td>
</tr>
<tr>
<td></td>
<td>Myonecrosis</td>
<td>110</td>
<td>81</td>
</tr>
<tr>
<td></td>
<td>Carditis</td>
<td>24</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>Eosinophilic myocarditis</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Giant cell myocarditis</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Hypersensitivity myocarditis</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>331</td>
<td>350</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>592</td>
<td>462</td>
</tr>
<tr>
<td>Relevant suspect case</td>
<td>Dose 1</td>
<td>213</td>
<td>172</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>273</td>
<td>250</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>346</td>
<td>308</td>
</tr>
<tr>
<td></td>
<td>Dose 4</td>
<td>8</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>83</td>
<td>77</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n=958</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤24 hours</td>
<td>60</td>
<td>36</td>
<td>0</td>
</tr>
<tr>
<td>1-3 days</td>
<td>167</td>
<td>166</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>88</td>
<td>90</td>
<td>4</td>
</tr>
<tr>
<td>14-21 days</td>
<td>58</td>
<td>56</td>
<td>1</td>
</tr>
<tr>
<td>22-31 days</td>
<td>34</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>40</td>
<td>42</td>
<td>0</td>
</tr>
<tr>
<td>61-365 days</td>
<td>45</td>
<td>43</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>435</td>
<td>359</td>
<td>12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>23</td>
<td>36</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>294</td>
<td>250</td>
<td>4</td>
</tr>
<tr>
<td>Resolved</td>
<td>97</td>
<td>107</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>33</td>
<td>32</td>
<td>2</td>
</tr>
<tr>
<td>Resolving</td>
<td>143</td>
<td>147</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>337</td>
<td>245</td>
<td>12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of Event n=89, median=34 days</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>2</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>1</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>7</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>26-170 days</td>
<td>18</td>
<td>19</td>
<td>1</td>
</tr>
<tr>
<td>171-822 days</td>
<td>10</td>
<td>5</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Fatal myocarditis cases in subjects aged ≥40 Years

There were 59 cases that reported 59 relevant events with fatal outcome in this age group. Of the 59 cases, 40 cases were medically confirmed and 19 were non-medically confirmed cases. There were 23 female and 36 male subjects. Subjects' ages ranged from 40 years to 96 years. The cases were reported from Japan (17), Germany (12), UK (7), Australia (5), Austria, France, Sweden (3 Each), New Zealand, Taiwan, Province of China (2 each), Hong Kong, Italy, Netherlands, Norway, and Switzerland (1 each).

The fatal events in these cases were coded to the PIs Abdominal pain upper, Acute coronary syndrome, Acute myocardial infarction, Amnesia, Aortic dissection, Aortic rupture, Aortitis, Arrhythmia, Arteriosclerosis coronary artery, Arteritis coronary, Arthritis, Asthenia, Atrial fibrillation, Atrioventricular block complete, Back pain, Bacteremia, Basal ganglia haemorrhage, Blood creatinine phosphokinase increased, Blood creatinine increased, Blood lactic acid, Bradycardia, Brain injury, Cardiac arrest, Cardiac disorder, Cardiac dysfunction, Cardiac failure, Cardiac failure high output, Cardiogenic shock, Cardiomegaly, Cardiomyopathy, Cardio-respiratory arrest, Cerebral haemorrhage, Chest pain, Chronic kidney disease, Circulatory collapse, Coiliis, Coma, Coronary artery stenosis, C-reactive protein increased, Cytology abnormal, Death, Dizziness, Dyspnoea, Dyspnoea exertional, Electrocardiogram ST segment depression, Embolism, Encephalitis, Encephalomalacia, Endocarditis, Eosinophilic myocarditis, Fatigue, Haemorrhage, Haemosiderosis, Hepatotoxicity, Hyperhidrosis, Hypersensitivity myocarditis, Immunisation, Infection, Inflammation, Influenza like illness, Interchange of vaccine products, Internal haemorrhage, Intracranial pressure increased, Ischaemic
cardiomyopathy, Malaise, Memory impairment, Multiple organ dysfunction syndrome, Myalgia, Myocardial fibrosis, Myocardial infarction, Myocardial necrosis, Myocarditis, Myopericarditis, Myositis, Obstruction, Off label use, Pain in extremity, Palpitations, Pericarditis, Peripheral coldness, pH body fluid, Pneumonia, Pneumonia aspiration, Pulmonary artery thrombosis, Pulmonary embolism, Pulmonary hypertension, Pulmonary oedema, Pulseless electrical activity, Pyrexia, Respiration abnormal, Respiratory failure, Right ventricular failure, Sepsis, Spinal cord haemorrhage, Sudden death, Syncope, Tachycardia, Tachypnoea, Thrombocytopenia, Thrombosis, Troponin I, Troponin increased, Vasculitis, Vasculitis necrotising, Ventricular fibrillation, Ventricular hypokinesia, Viral myocarditis, Vomiting (1 each).

Only 1 case reported a co-suspect medication (pembrolizumab). The most frequently reported (>1 occurrence) medical histories were coded to the PTs Hypertension (8), Cardiac failure, Diabetes mellitus, Obesity (4 each), Cardiac disorder (3), Cardiac failure chronic, Dyslipidaemia, and Type 2 diabetes mellitus (2 each). The most frequently reported (>2 occurrence) cause of death in these cases were coded to the PTs Myocarditis (47), Cardiac arrest (9), Death (7), Cardiac failure, Pericarditis (5 each), Cardio-respiratory arrest, Chest pain, Dyspnoea, Sudden death (4 each), Pneumonia, Syncope (3 each).

\[\text{Rapporteur assessment comment:}\]

During the current reporting period, there were 1752 cases reporting myocarditis in persons aged ≥40 years compared to 1876 myocarditis cases reported in the previous 2\textsuperscript{nd} PSUR. There were 59 fatal cases compared to 42 fatal cases in the previous reporting period.

The MAH only briefly presented the 59 fatal cases and did not provide an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged ≥40 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, and a WHO causality assessment. \textit{Request for supplementary information}

\begin{itemize}
\item \textbf{Subjects with booster dose}
\item \textbf{Clinical Trial Data}
  \begin{itemize}
  \item The case involved a 43-year-old male participant, who received homologous booster dose. Please see above the "Overall- All Ages" subsection for complete details.
  \end{itemize}
\item \textbf{Post-Authorisation Data}
  \begin{itemize}
  \item Number of cases: 1682 (0.3 \% of 507,683 cases of the total PM dataset, 1.4 \% of the 117750 subjects who received a booster dose), compared to 381 cases (0.06\%) in the PSUR #2.
  \item Country/region of incidence (≥10): UK (617), Germany (422), France (113), Austria (72), Italy (53), Japan (48), Israel (44), New Zealand (41), US (32), Greece, Sweden (21 each), Finland (19), Netherlands, Taiwan, Province of China (17 Each), Denmark (16), Australia, Hong Kong (15 each), Switzerland (13), Spain (12), Ireland (10); the remaining 64 cases were distributed among 19 countries.
  \item MC (702), NMC (980).
  \item Subjects’ gender: female (656), male (988), and unknown (38).
  \end{itemize}
\end{itemize}

- Medical history (n = 633): the medical conditions reported (>4 occurrence) included Hypertension (87), Asthma (46), Immunodeficiency (34), Tobacco user (31), Seasonal allergy (30), Hypothyroidism (28), Clinical trial participant (22), Myocarditis (21), Diabetes mellitus (20), Atrial fibrillation, Non-tobacco user (19), Obesity (18), Depression, Migraine (17 each), Steroid therapy (14), Food allergy (13), Anxiety, Dyslipidaemia (12 each), Drug hypersensitivity, Gastroesophageal reflux disease, Interchange of vaccine products (11 each), Chest pain, Mite allergy, Overweight (10 each), Rheumatoid arthritis, Type 2 diabetes mellitus (9 each), Alcohol use, Chronic obstructive pulmonary disease, Dyspnoea, Fibromyalgia, Hyperlipidaemia, Myocardial infarction (8 each), Autoimmune thyroiditis, Cardiac disorder, Coronary artery disease, Ex-tobacco user, Nasopharyngitis (7 each), Cerebrovascular accident, Colitis ulcerative, Crohn's disease, Hypersensitivity, Inflammatory bowel disease, Insomnia, Mitral valve incompetence, Neoplasm, Nephrolithiasis, Nicotine dependence, Osteoarthritis, Pain, Pericarditis, Pneumonia, Pulmonary embolism, Sleep apnoea syndrome (6 each), Abstains from alcohol, Allergy to animal, Appendicectomy, Arteriosclerosis, Attention deficit hyperactivity disorder, Coeliac disease, Congestive cardiomyopathy, Contraception, Endometriosis, Epstein-Barr virus infection, Fatigue, Gastritis, Gout, Hodgkin's disease, Hormone replacement therapy, Menopause, Myopericarditis, Osteoporosis, Palpitations, Pregnancy, Radiotherapy, Supraventricular tachycardia, Surgery, Urinary tract infection (5 each).


- Co suspect vaccines (n= 20) reported more than once: Influenza vaccine (5), pembrolizumab (2), amoxicillin, cisplatin, COVID-19 vaccine NRV AD (CHADOX1 NCOV-19), fluticasone, gabapentin, hepatitis A vaccine, infliximab, influenza vaccine INACT SAG 3V, JNO 78436735, paracetamol, propranolol, vinorelbine, zuclopenthixol (1 each).

- Number of relevant events: 1696.

- Relevant event seriousness: all serious.

- Reported relevant PTs: Myocarditis (1458), Myopericarditis (211), Carditis (25), and Eosinophilic myocarditis (2).

- Relevant event outcome: fatal (39), resolved/resolving (528); resolved with sequelae (29), not resolved (453), unknown (649).

- Most frequently co-reported PTs (>20 occurrence): Chest pain (691), Immunisation (537), Fatigue (494), Pericarditis (467), Dyspnoea (466), Palpitations (443), Off label use (442), Interchange of vaccine products (379), Tachycardia (358), Pyrexia (311), Headache (174), Chest discomfort (163), Dizziness (111), Malaise, Pain (86 each), Pain in extremity (84), Nausea (77), Syncope (73), Arrhythmia (71), Chills, Heart rate increased (70 each), Angina pectoris (66), Myalgia (65), Asthenia (59), Arthralgia (58), Troponin increased (52), Lymphadenopathy (51), Vomiting (45), Dyspnoea exertional (43), Back pain, Pericardial effusion (41 each), Hypertension (38), Diarrhoea, Influenza like illness (37), Atrial fibrillation (36), Cough (35), Cardiac flutter, Hyperhidrosis (34 each), Cardiac failure, Vaccination site pain (30), COVID-19 (27), Oropharyngeal pain
(26), C-reactive protein increased, Neck pain (24 each), Insomnia (23), Axillary pain, Hypoesthesia, Paraesthesia (22 each), Cardiac disorder, Myocardial infarction (21 each), Blood pressure increased, Extrasystoles (20 each).

The number of myocarditis cases occurred after a booster dose in each age group is reported in Table 44 below by gender.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Heterologous Booster dose No. of Cases</th>
<th>Homologous Booster dose No. of Cases</th>
<th>Unknown dose No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age group</td>
<td>F</td>
<td>M</td>
<td>U</td>
</tr>
<tr>
<td>0 to 17 years</td>
<td>1</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>18 to 24 years</td>
<td>4</td>
<td>30</td>
<td>0</td>
</tr>
<tr>
<td>25 to 29 years</td>
<td>7</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>30 to 39 years</td>
<td>25</td>
<td>24</td>
<td>0</td>
</tr>
<tr>
<td>40 years and older</td>
<td>142</td>
<td>109</td>
<td>5</td>
</tr>
<tr>
<td>Unknown</td>
<td>30</td>
<td>12</td>
<td>7</td>
</tr>
<tr>
<td>TOTAL</td>
<td>209</td>
<td>188</td>
<td>12</td>
</tr>
</tbody>
</table>

F=female, M=male, U=unknown

Rapporteur assessment comment:

During the current reporting period, there were 1,652 cases reporting myocarditis in persons who received a booster dose compared to 381 myocarditis cases reported in the previous 2nd PSUR. There were 39 fatal cases compared to 4 fatal cases in the previous reporting period.

The 39 fatal cases are assumed to be imbedded in the fatal cases stated in the age categories above, which are subject for a request for supplementary information.

During the reporting period there were 1,639 cases of medically confirmed myocarditis with a latency 21 days or less in subjects receiving booster dose. All cases were assessed as serious due to hospitalisation and/or medically significant (1002) or due to medically significant (637). In 1314 cases myocarditis occurred within 1 week post vaccine administration. In most of these cases, the insufficient description of During the current reporting period, there were 968 cases reporting myocarditis in persons aged 18-24 years compared to 1187 myocarditis cases reported in the previous 2nd PSUR. There were 4 fatal cases compared to 2 fatal cases in the previous reporting period cardiovascular and/or non-cardiovascular medical history and the lack of diagnostic tests confirming the aetiologies of myocarditis preclude a clear causality assessment on an individual case basis.

Rapporteur assessment comment:

During the current reporting period, there were 1,639 cases reporting medically confirmed myocarditis with a TTO 21 days or less compared to 2,007 medically confirmed myocarditis cases reported in the previous 2nd PSUR. The MAH stated that most of the cases had Insufficient information that precluded a clear causality assessment on an individual case basis.

Rapporteur assessment comment:

In general, the MAH should focus the analysis of myocarditis cases on aspects of these ADRs not fully known or addressed in the Comirnaty PI (myocarditis is already an ADR stated in section 4.8), and if the warning in section 4.4 regarding myocarditis is still in line with current knowledge. Therefore, the
| analysis should focus more on information concerning the course, outcome, and possible risk factors of the myocarditis cases following Comirnaty exposure. **Request for next PSUR** |
Myocarditis

Clinical trial data

During the reporting period, one myocarditis case (after homologous booster dose) was retrieved and considered not related to Comirnaty exposure.

Post-marketing

- Aged 5-11 years: There were 48 cases reporting myocarditis (2 fatal cases) compared to 10 myocarditis cases (no fatal cases) reported in the previous 2nd PSUR.
- Aged 12-15 years: There were 366 cases reporting myocarditis (3 fatal cases) compared to 488 myocarditis cases (3 fatal cases) reported in the previous 2nd PSUR.
- Aged 16-17 years: There were 345 cases reporting myocarditis (no fatal cases) compared to 470 myocarditis cases (2 fatal cases) reported in the previous 2nd PSUR.
- Aged 18-24 years: There were 968 cases reporting myocarditis (4 fatal cases) compared to 1187 myocarditis cases (2 fatal cases) reported in the previous 2nd PSUR.
- Aged 25-29 years: There were 519 cases reporting myocarditis (5 fatal cases) compared to 589 myocarditis cases (7 fatal cases) reported in the previous 2nd PSUR.
- Aged 30-39 years: There were 983 cases reporting myocarditis (4 fatal cases) compared to 995 myocarditis cases (7 fatal cases) reported in the previous 2nd PSUR.
- Aged ≥40 years: There were 1752 cases reporting myocarditis (59 fatal cases) compared to 1876 myocarditis cases (42 fatal cases) reported in the previous 2nd PSUR.

The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 5-11 years, aged 12-15, aged 18-24, aged 25-29, aged 30-39, and aged ≥40 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable. Request for supplementary information

Pericarditis

Search criteria: Autoimmune pericarditis; Pericarditis; Pericarditis adhesive; Pericarditis constrictive; Pleuropericarditis.

Overall - All ages

- Clinical Trial Data
  - Number of cases: No cases were retrieved during the current reporting period, compared to 1 case (0.14 %) retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 4156 (0.8% of 507,683 cases of the total PM dataset), compared to 5311 cases (0.8%) retrieved in the PSUR #2.
  - Country of incidence: Australia (1085), UK (903), France (580), Italy (281), Germany (271), Canada (174), New Zealand (111), Netherlands (97), Sweden (71), Japan (68). The remaining 515 cases were distributed among 44 countries.
o MC (2370), NMC (1786).

o Subjects' gender: female (2049), male (2017) and unknown (90).

o Subjects' age in years: n = 3847, range: 4-98 years, mean: 39.8, median: 37.0.

o Medical history: (n = 1292) the most frequently (≥1%) reported relevant medical history included: Hypertension (154), Asthma (109), Pericarditis (95), Seasonal allergy (60), Drug hypersensitivity, Tobacco user (58 each), Immunodeficiency (54), Hypothyroidism (51), Obesity (46), Hypersensitivity, Non-tobacco user (40 each).


o Co-suspects (n=48 cases): frequently (>3 occurrences) reported relevant co-suspect vaccines/medications were COVID-19 vaccine mRNA (mRNA 1273), Influenza vaccine (8 each), COVID-19 vaccine, Influenza vaccine INACT SAG 3V, Influenza vaccine INACT SPLIT 4V (3 each).

o Number of relevant events: 4164.

o Relevant event seriousness: serious (4164).

o Reported relevant PTs: Pericarditis (4133), Pleuroperticarditis (26), Pericarditis constrictive (5).

o Relevant event outcome: fatal (19), resolved/resolving (1311), resolved with sequelae (82), not resolved (1428), unknown (1325).

Subjects aged less than 5 years

- Clinical Trial Data
  
o Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  
o Number of cases: 1; 1 case was retrieved in the PSUR #2.
  
o Country of incidence: Australia.
  
o Subject's age in year: 4.
  
o Gender: female.
  
o Medical history: unknown.
  
o Co-suspects: none.
  
o Relevant PT: Pericarditis
  
o Medically Confirmed: yes.
  
o Hospitalisation required: no
  
o Time to onset (pericarditis): ≤24 hours after the 1st dose.
  
o Co-reported PTs: Chest discomfort, Chest pain, Dyspnoea, Fatigue, Headache, Myalgia, Pyrexia, and Product administered to patient of inappropriate age.
Rapporteur assessment comment:
Comirnaty exposure in persons aged less than 5 years is considered off-label use during the interval period of the current 3rd PSUR.

Subjects aged 5 - 11 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 30 (0.006 % of 507,683 cases of the total PM dataset, 0.4 % of the 8375 subjects aged 5-11 years); 4 cases (0.0006%) were retrieved in the PSUR #2.
  - Country of incidence: Australia (19), Canada (3), Italy, Japan (2 each), Germany, Israel, New Zealand, UK (1 each).
  - Subjects' age in year: n = 30, range: 5 -11, mean: 9.4, median: 10.0.
  - Medical history: Coeliac disease, Kawasaki's disease, Urinary tract infection viral (1 each).
  - COVID-19 Medical history: COVID-19 (1)
  - Co suspects: none.
  - Most frequently co-reported PTs (>2 occurrences): Chest pain (24), Dyspnoea (12), Electrocardiogram abnormal (7), Chest discomfort (6), Palpitations (5), Myocarditis (4), Pyrexia (3).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 45.

Table 45. Pericarditis in Subjects aged 5-11 years (N=30)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>7</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Pericarditis</td>
<td>9</td>
<td>21</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>8</td>
<td>19</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>7</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Time to Onset n=30</td>
<td>≤ 24 hours</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>1-3 days</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>14-21 days</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>22-31 days</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Event Outcome</td>
<td>Fatal</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Duration of event a, n=2, median: 18</td>
<td>4-6 days</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>11-26 days</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

a. All serious occurrences
b. For those cases where the event resolved or resolved with sequelae.
Rapporteur assessment comment:

During the current reporting period, there were 30 cases reporting pericarditis in children aged 5-11 years compared to 4 pericarditis cases reported in the previous 2nd PSUR. Although the Comirnaty exposure in persons aged 5-11 years is considered increased based on the EU/EEA exposure (current reporting period an estimated 3,569,821 administered doses in 5-9 years versus 391,327 administered doses in 5-9 years in the previous reporting period), worldwide interval exposure in persons aged 5-11 years (or any other age category) is not presented in the PSUR and therefore the relative post-marketing reporting rate of myocarditis cases in persons aged 5-11 years is not known.

There were no fatal cases compared to no fatal cases in the previous reporting period.

Subjects aged 12 - 15 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 118 (0.02 % of 507,683 cases of the total PSUR dataset, 0.9 % of the 13,366 subjects aged 12-15 years), compared to 215 cases (0.03%) retrieved in the PSUR #2.
  - Country of Incidence: Australia (31), UK (13), Taiwan, Province of China (11), France, Japan (8 each), Canada, Italy (7 each), Malaysia (6). The remaining 27 cases were distributed among 12 countries.
  - Medical history (n = 20): the medical conditions reported more than once included Adenotonsillectomy, Asthma, Glucose-6-phosphate dehydrogenase deficiency, and Hypersensitivity (2 each).
  - Co suspects: none.
  - Most frequently co-reported PTs (≥2%): Chest pain (60), Myocarditis (34), Dyspnoea (25), Palpitations (23), Pyrexia (22), Chest discomfort, Fatigue (15 each), Headache, Tachycardia (9 each), Dizziness, Malaise (7 each), Asthenia, Inappropriate schedule of product administration (6 each), Cough, Heart rate increased, Nausea, Pain, Pericardial effusion (5 each), Dyspnoea exertional, Syncope, Vomiting (4 each), Arthralgia, Chills, COVID-19, Electrocardiogram abnormal, Electrocardiogram ST segment elevation, Troponin increased (3 each), Angina pectoris, Back pain, Drug Ineffective, Electrocardiogram ambulatory abnormal, Exercise tolerance decreased, Immune system disorder, Lethargy, Musculoskeletal chest pain, Myalgia, Nasopharyngitis, Oropharyngeal pain, Pleural effusion, Pleuritic pain, and Sinus tachycardia (2 each).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 46.
Table 46. Pericarditis in Subjects aged 12-15 years (N=118)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>15</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>9</td>
<td>20</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Pericarditis</td>
<td>24</td>
<td>91</td>
</tr>
<tr>
<td></td>
<td>Pleuropericarditis</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>4</td>
<td>42</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>20</td>
<td>50</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>11</td>
<td>38</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>8</td>
<td>42</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>3</td>
<td>8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n=118</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 24 hours</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>10</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>6-13 days</td>
<td>4</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>61-180 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>6</td>
<td>33</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>11</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Resolved</td>
<td>4</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>4</td>
<td>29</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>5</td>
<td>25</td>
<td>1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event* n=6, median: 9</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-6 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>11-36 days</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>27-57 days</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

Rapporteur assessment comment:

During the current reporting period, there were 118 cases reporting pericarditis in persons aged 12-15 years compared to 215 pericarditis cases reported in the previous 2nd PSUR. There were no fatal cases compared to no fatal cases in the previous reporting period.

Subjects aged 16 - 17 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.
- Post-Approval Data
  - Number of cases: 106 (0.02 % of 507,683 cases of the total PM dataset, 1.3 % of the 8313 subjects aged 16-17 years), compared to 174 cases (0.03%) retrieved in the PSUR #2.
  - Country of Incidence: Australia (25), UK (20), France (15), Italy (11), Germany (6), Taiwan, province of China (5). The remaining 24 cases were distributed among 14 countries.
  - Subjects’ age in years: n = 106, range: 16 -17, mean: 16.5, median: 16.0.
  - Medical history (n = 17): the medical conditions reported more than once included the PTs Asthma, Food allergy, Pericarditis, Seasonal allergy (2 each).
Co suspects (n= 2 cases): COVID-19 vaccine mRNA (MRNA 1273), HPV vaccine VLP RL1 9V (yeast), Influenza vaccine INACT SPLIT 4V, Pneumococcal vaccine polysacch 23V (1 each).

Most frequently co-reported PTs (≥2%): Chest pain (56), Dyspnoea (25), Myocarditis, Pyrexia (22 each), Fatigue, Palpitations (19 each), Tachycardia (15), Chest discomfort (14), Inappropriate schedule of product administration, Nausea, Pain (7 each), Headache, Pericardial effusion, Vomiting (6 each), Electrocardiogram abnormal, Malaise, Myopericarditis (5 each), Chills, Cough, Dizziness, Troponin increased, Abdominal pain upper, Influenza like illness, Lethargy, Pain in extremity (3 each), Asthenia, Back pain, Cellulitis, Cold sweat, C-reactive protein increased, Decreased appetite, Feeling hot, Heart rate irregular, Hyperhidrosis, Interchange of vaccine products, Myalgia, Off label use, Product use issue, Syncope, Vaccination site pain (2 each).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 47.

Table 47. Pericarditis in Subjects aged 16-17 years (N=106)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>20</td>
<td>50</td>
<td>1</td>
</tr>
<tr>
<td>No</td>
<td>15</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>35</td>
<td>70</td>
<td>1</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>7</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>No</td>
<td>28</td>
<td>49</td>
<td>1</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>12</td>
<td>23</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>20</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>2</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>1</td>
<td>8</td>
<td>0</td>
</tr>
</tbody>
</table>

| Time to Onset n=106         |                     |                  |                      |
| ≤ 24 hours                 | 1                   | 6                | 0                    |
| 1-5 days                   | 9                   | 20               | 1                    |
| 6-13 days                  | 3                   | 5                | 0                    |
| 14-21 days                 | 1                   | 3                | 0                    |
| 22-31 days                 | 2                   | 3                | 0                    |
| 32-68 days                 | 5                   | 2                | 0                    |
| 61-180 days                | 2                   | 3                | 0                    |
| 181-375 days               | 0                   | 1                | 0                    |
| Unknown                    | 12                  | 27               | 0                    |

| Event Outcome              |                     |                  |                      |
| Fatal                      | 0                   | 0                | 0                    |
| Not resolved               | 11                  | 17               | 1                    |
| Resolved                   | 4                   | 19               | 0                    |
| Resolved with sequelae     | 0                   | 1                | 0                    |
| Resolving                  | 11                  | 13               | 0                    |
| Unknown                    | 9                   | 20               | 0                    |

| Duration of eventb n=7, median: 10 |                     |                  |                      |
| Up to 3 days               | 1                   | 2                | 0                    |
| 3-10 days                  | 0                   | 1                | 0                    |
| 11-26 days                 | 0                   | 1                | 0                    |
| 27-57 days                 | 0                   | 1                | 0                    |
| 58-180 days                | 0                   | 1                | 0                    |

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

Rapporteur assessment comment:

During the current reporting period, there were 106 cases reporting pericarditis in persons aged 16-17 years compared to 174 pericarditis cases reported in the previous 2nd PSUR.

There were no fatal cases compared to no fatal cases in the previous reporting period.
Subjects aged 18 - 24 years

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 479 (0.09 % of 507,683 cases of the total PM dataset, 1.3% of the 38,293 subjects aged 18-24 years), compared to 659 cases (0.10%) retrieved in the PSUR #2.
  - Country of Incidence: Australia (135), France (79), UK (73), Germany (44), Italy (33), New Zealand (19), Japan (14), Netherlands, Sweden (12 each), Norway (11), US (6). The remaining 41 cases were distributed among 16 countries.
  - Medical history (n = 120): the medical conditions reported more than twice included Asthma (21), Immunodeficiency, Pericarditis (6 each), Attention deficit hyperactivity disorder, Mite allergy, Non-tobacco user, Obesity, Overweight, Tobacco user (5 each), Food allergy, Irritable bowel syndrome (4 each), Disease risk factor, Drug hypersensitivity, Endometriosis, Hospitalisation, Hypersensitivity, Hypothyroidism, Migraine, Seasonal allergy, and Substance use (3 each).
  - Co suspects (n= 8 cases): COVID-19 vaccine MRNA (MRNA 1273) (2), COVID-19 vaccine, duplumab, Influenza vaccine INACT SPLIT 4V, Insulin, levothyroxine, salbutamol, zuclopenthixol (1 each).
  - Most frequently co-reported PTs (>2%): Dyspnoea (141), Myocarditis (112), Palpitations (95), Fatigue (83), Chest discomfort (75), Pyrexia (66), Tachycardia (65), Headache, Pericardial effusion (35 each), Dizziness (33), Inappropriate schedule of product administration (26), Electrocardiogram abnormal (24), Pain (23), Immunisation, Myalgia (20 each), Malaise, Off label use (19 each), Interchange of vaccine products, Syncope (18 each), Asthenia (17), Pain in extremity (16), Nausea (15), Angina pectoris, Chills, Vomiting (14 each), Cough (13), C-reactive protein increased, Dyspnoea exertional, Hyperhidrosis, Lethargy (12 each), Anxiety, Sinus tachycardia (10 each), Arthralgia, Heart rate increased (9 each), Back pain, Electrocardiogram ST segment elevation, Paraesthesia, Troponin increased (8 each).
  - Pericarditis events with fatal outcome (1).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 48.
Table 48. Pericarditis in Subjects aged 18-24 years (N=479)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>120</td>
<td>169</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>72</td>
<td>110</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Pericarditis</td>
<td>192</td>
<td>279</td>
</tr>
<tr>
<td>Hospitalisation</td>
<td>Yes</td>
<td>46</td>
<td>96</td>
</tr>
<tr>
<td>required/prolonged</td>
<td>No</td>
<td>146</td>
<td>183</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>79</td>
<td>102</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>52</td>
<td>96</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>49</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>Dose 4</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>11</td>
<td>19</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n=479</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 24 hours</td>
<td>27</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>66</td>
<td>93</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>22</td>
<td>34</td>
<td>2</td>
</tr>
<tr>
<td>14-21 days</td>
<td>5</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>5</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>1</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>61-180 days</td>
<td>7</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>181-375 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>59</td>
<td>86</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>74</td>
<td>97</td>
<td>2</td>
</tr>
<tr>
<td>Resolved</td>
<td>20</td>
<td>33</td>
<td>2</td>
</tr>
<tr>
<td>Resolved with sequela</td>
<td>7</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>Resolving</td>
<td>47</td>
<td>65</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>44</td>
<td>76</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event a=18, median = 21</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>11-26 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>27-57 days</td>
<td>3</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>58-180 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

- All serious occurrences.
- For these cases where the event resolved or resolved with sequela.

Fatal pericarditis cases in adult (18-24 years of age) (1 case, medically confirmed):

- A 22-year-old male subject from Israel:
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Cardiac tamponade, Multiple organ dysfunction syndrome, Pericardial effusion, Pericardial mass, Pericardial mesothelioma malignant, Pericarditis, Right ventricular dysfunction, Right ventricular failure.
  - Time to onset (pericarditis): 31 days after dose 2.
  - Causes of death: all the above events.

**Rapporteur assessment comment:**

During the current reporting period, there were 479 cases reporting pericarditis in persons aged 18-24 years compared to 659 pericarditis cases reported in the previous 2nd PSUR. There was 1 fatal case compared to no fatal cases in the previous reporting period.

The MAH only briefly presented the fatal case and did not provide an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and no WHO causality assessment for the case regarding Comi maty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal case with pericarditis in persons aged 18-24 years and perform an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and a WHO causality assessment. **Request for supplementary information**
Subjects aged 25 - 29 years

- Clinical Trial Data
  - Number of cases: none; no cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 417 (0.08 % of 507,683 cases of the total PM dataset, 1.0 % of the 43,518 subjects aged 25-29 years), compared to 614 cases (0.09%) retrieved in the PSUR #2.
  - Country of Incidence: Australia (136), UK (75), France (71), Germany (21), Italy, Netherlands (18 each), New Zealand (16), Sweden (8), Japan, Spain (7 each), Denmark (6), Canada (5). The remaining 29 cases were distributed among 17 countries.
  - Subjects' age in years: n = 417, range: 25-29, mean: 27.0, median: 27.0.
  - Medical history (n = 87): the medical conditions reported more than twice included Asthma (10), Tobacco user (7), Obesity (6), Disease risk factor, Drug hypersensitivity, Non-tobacco user, Pericarditis (4 each), Abstains from alcohol, Contraception, Gastritis, Steroid therapy (3 each).
  - Co suspects (n=3 cases): COVID-19 vaccine (2), and Methylphenidate (1).
  - Most frequently co-reported PTs (≥2%): Dyspnœa (137), Palpitations (101), Fatigue (94), Myocarditis (87), Tachycardia (67), Chest discomfort (60), Pyrexia (49), Headache (40), Dizziness, Immunisation (33 each), Nausea (22), Pain (21), Off label use (20), Interchange of vaccine products (19), Malaise, Pericardial effusion (17 each), Syncope (16), Electrocardiogram abnormal, Myalgia (14 each), Angina pectoris (13), Arthralgia, Asthenia, Heart rate increased (12 each), Dyspnœa exertional, Pain in extremity, Paraesthesia, Vaccination site pain (11 each), Lethargy, Lymphadenopathy (10 each), Inappropriate schedule of product administration (9), Troponin increased (8), Cardiac flutter, Diarrhoea, Vomiting (7 each).
  - Pericarditis events with fatal outcome (1).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 49.
### Table 49. Pericarditis in Subjects aged 25-29 years (N=417)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>102</td>
<td>146</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>71</td>
<td>91</td>
</tr>
<tr>
<td>Relevant: PT*</td>
<td>Pericarditis</td>
<td>171</td>
<td>237</td>
</tr>
<tr>
<td></td>
<td>Pleuropericarditis</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>41</td>
<td>47</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>132</td>
<td>190</td>
</tr>
<tr>
<td>Relevant: suspect dose</td>
<td>Dose 1</td>
<td>68</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>40</td>
<td>61</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>50</td>
<td>58</td>
</tr>
<tr>
<td></td>
<td>Dose 4</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>15</td>
<td>17</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n=418</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 24 hours</td>
<td>15</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>42</td>
<td>76</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>23</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>15</td>
<td>20</td>
<td>1</td>
</tr>
<tr>
<td>22-31 days</td>
<td>6</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>8</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>61-180 days</td>
<td>11</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>181-375 days</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>52</td>
<td>73</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>78</td>
<td>84</td>
<td>1</td>
</tr>
<tr>
<td>Resolved</td>
<td>12</td>
<td>37</td>
<td>2</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>34</td>
<td>47</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>47</td>
<td>65</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event n=13, median: 13</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>11-26 days</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>27-57 days</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>58-180 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

- All serious occurrences
- For those cases where the event resolved or resolved with sequelae.

**Fatal pericarditis cases in adult (25-29 years of age) (1 case, medically confirmed)**

- A 29-year-old male subject from Finland:
  - Medical history: Hypoventilation, Obesity, Pulmonary fibrosis, Sleep apnoea syndrome, Still's disease
  - Co-suspect medications: None
  - PTs with fatal outcome: Acute kidney injury, Aortic dissection, Chest pain, Hypoventilation, Inflammatory marker increased, Multiple organ dysfunction syndrome, Pericardial disease, Pericarditis, Respiratory failure, Sepsis.
  - Time to onset (pericarditis): 6 days after dose 3.

**Causes of death:** Multiple organ dysfunction syndrome; Sepsis; Still's disease.

**Rapporteur assessment comment:**

During the current reporting period, there were 417 cases reporting pericarditis in persons aged 25-29 years compared to 614 pericarditis cases reported in the previous 2nd PSUR. There was 1 fatal case compared to no fatal cases in the previous reporting period.

The MAH only briefly presented the fatal case and did not provide an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and no WHO causality.
assessment for the case regarding Community exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal case with pericarditis in persons aged 18-24 years and perform an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and a WHO causality assessment. **Request for supplementary information**

**Subjects aged 30 – 39 years**

- Clinical Trial Data
  - Number of cases: none. No cases were retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 940 (0.2 % of 507,683 cases of the total PM dataset; 1.0 % of the 97,870 subjects aged 30-39), compared to 1222 cases (0.2%) retrieved in the PSUR #2.
  - Country/region of incidence: Australia (356), UK (217), France (114), Germany (46), Italy (39), New Zealand (26), Netherlands (21), Canada (18), Norway (16), Sweden (13), Belgium (9), Greece, US (8 each), Denmark, Japan (6 each), Austria, and Hong Kong (5 each). The remaining 27 cases were distributed among 17 different countries.
  - Subjects’ age in years: n = 940, range: 30 -39, mean: 34.3, median: 34.0.
  - Medical history (n = 217): the medical conditions reported more than 5 times included the PTs Pericarditis (27), Asthma (20), Drug hypersensitivity (18), Seasonal allergy (13), Mite allergy, Non-tobacco user, Pregnancy, Tobacco user (11 each), Migraine (10), Chest pain, Hypothyroidism (8 each), Anxiety, Clinical trial participant, Immunodeficiency (7 each), Alcohol use, Eczema, Obesity (6 each).
  - COVID-19 Medical history (n = 70): COVID-19 (43), Suspected COVID-19 (24), SARS-CoV-2 test positive (3).
  - Co suspect vaccines/medications (n=6): colchicine (2), amoxicillin, interferon Beta-1A, iron isomaltoside 1000, propranolol (1 each).
  - Most frequently co-reported PTs (≥2%): Chest pain (547), Dyspnoea (345), Palpitations (260), Fatigue (239), Myocarditis (236), Tachycardia (179), Chest discomfort (125), Pyrexia (113), Headache (93), Dizziness (75), Immunisation (74), Malaise, Pain in extremity (54 each), Nausea, Parasthesia (48 each), Arthralgia, Pain (46 each), Off label use (45), Myalgia (44), Inappropriate schedule of product administration (40), Heart rate increased, Interchange of vaccine products (39 each), Hypoaesthesia, Pericardial effusion (38 each), Asthenia (32), Hyperhidrosis, Syncope (29 each), Electrocardiogram abnormal, Influenza like illness (25 each), Cardiac flutter (24), Arrhythmia (23), Chills, Lethargy (21 each), Feeling abnormal, Vaccination site pain (20 each), Diarrhoea, Exercise tolerance decreased, Vomiting (18 each), Cough (17), Anxiety, Back pain, Dyspnoea exertional, Lymphadenopathy, Neck pain (16 each).
Pericarditis relevant data in this subgroup of subjects are summarised in below Table 50.

Table 50. Pericarditis in Subjects aged 30-39 years (N=940)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>277</td>
<td>284</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>193</td>
<td>179</td>
</tr>
<tr>
<td>Relevant PT(a)</td>
<td>Pericarditis</td>
<td>470</td>
<td>462</td>
</tr>
<tr>
<td></td>
<td>Pleuropericarditis</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>68</td>
<td>96</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>402</td>
<td>367</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dose 1</td>
<td>223</td>
<td>243</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>119</td>
<td>111</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>108</td>
<td>83</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>20</td>
<td>24</td>
</tr>
<tr>
<td>Time to Onset n=941</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>≤ 24 hours</td>
<td>37</td>
<td>35</td>
</tr>
<tr>
<td></td>
<td>1-5 days</td>
<td>105</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>47</td>
<td>52</td>
</tr>
<tr>
<td></td>
<td>14-21 days</td>
<td>27</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>22-31 days</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>32-60 days</td>
<td>17</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>61-180 days</td>
<td>21</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>181-375 days</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>194</td>
<td>187</td>
</tr>
<tr>
<td>EventOutcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fatal</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>181</td>
<td>189</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>37</td>
<td>63</td>
</tr>
<tr>
<td></td>
<td>Resolved with sequelae</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>81</td>
<td>65</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>166</td>
<td>144</td>
</tr>
<tr>
<td>Duration of event n=27, median: 15</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Up to 3 days</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>4-6 days</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>7-10 days</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>11-26 days</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>27-57 days</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>58-180 days</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

\(a\) All serious occurrences.
\(b\) For those cases where the event resolved or resolved with sequelae.

Rapporteur assessment comment:

During the current reporting period, there were 940 cases reporting pericarditis in persons aged 30-39 years compared to 1222 pericarditis cases reported in the previous 2\(^{nd}\) PSUR. There were no fatal cases compared to 1 fatal case in the previous reporting period.

Subjects aged ≥40 years

- Clinical Trial Data
  - Number of cases: none. One (1) case (0.14%) retrieved in the PSUR #2. Please see above the "Overall – All Ages" subsection.
- Post-Authorisation Data
  - Number of cases: 1756 (0.3 % of 507,683 cases of the total PM dataset, 0.7% of the 236,404 subjects ≥ 40 years), compared to 2059 cases (0.3%) retrieved in the PSUR #2.
  - Country of incidence: UK (375), Australia (333), France (288), Italy (169), Germany (137), Canada (61), New Zealand (44), Netherlands (41), Greece (40), Sweden (35),

PRAC PSUR assessment report
EMA/PRAC/849/2023
Austria, Norway (28 each), Japan (25), Denmark (20). The remaining 132 cases were distributed among 25 different countries.

- **Subjects’ age in years:** n = 1756, range: 40-98, mean: 54.6, median: 52.0.
- **Medical history** (n = 738): the medical conditions reported more than 10 times included PTs Hypertension (133), Pericarditis (47), Asthma (43), Immunodeficiency, Seasonal allergy (36 each), Hypothyroidism (34), Hypersensitivity (29), Obesity, Tobacco user, Type 2 diabetes mellitus (28 each), Diabetes mellitus (27), Drug hypersensitivity, Gastrooesophageal reflux disease (25 each), Depression (20), Atrial fibrillation (19), Dyslipidaemia, Rheumatoid arthritis (18 each), Anxiety (17), Breast cancer, Dyspnoea, Hypercholesterolaemia, Myocardial ischaemia, Non-tobacco user (16 each), Chronic kidney disease (15), Myocardial infarction (14), Chronic obstructive pulmonary disease, Food allergy, Gastritis (13 each), Autoimmune thyroiditis, Chest pain, Systemic lupus erythematosus (12 each), Overweight, Palpitations, Psoriasis, Steroid therapy (11 each).
- **Co suspect vaccines/medications** (n= 24): Influenza vaccine (7), COVID-19 vaccine mRNA (MRNA 1273), Influenza vaccine INACT SAG 3V (3 each), Adalimumab, Aiplaban, COVID-19 vaccine NRV AD (CHADOX1 NCOV-19), Etanercept, Glyceryl trinitrate, Influenza vaccine INACT SPLIT 4V, Levetiracetam, Peginterferon alfa-2A, Pembrolizumab, Rivaroxaban, Sotrovimab (1 each).
- **Most frequently co-reported PTs (≥2%):** Chest pain (787), Dyspnoea (567), Fatigue (455), Myocarditis (396), Palpitations (372), Tachycardia (286), Off label use (272), Intercambio of vaccine products (246), Immunisation (243), Pyrexia (223), Chest discomfort (199), Pericardial effusion (167), Headache (157), Dizziness (131), Malaise (87), Pain in extremity (83), Astenia (82), Arthralgia (74), Nausea (73), Pain (71), Inappropriate schedule of product administration (70), Syncope (69), Myalgia (67), Angina pectoris, Paraesthesia (60 each), Arrhythmia, Cough (57 each), Lymphadenopathy (50), Heart rate increased (49), Chills (46), Hyperhidrosis (44), Electrocardiogram abnormal (43), Back pain (42), Hypertension (41), Lethargy (40), Pleural effusion, Vaccination site pain (39 each), Atrial fibrillation, Diarrhoea (38 each), Dyspnoea exertional (37), Influenza like illness (36), Myocardial infarction (33), Neck pain (32), Cardiac flutter, Condition aggravated (31 each), C-reactive protein increased, Vomiting (29 each).
- **Pericarditis events with fatal outcome** (17) occurred in subjects aged ≥40 years (n=17, ranged between 41 to 92 years of age).

Pericarditis relevant data in this subgroup of subjects are summarised in Table 51 below.
Table 51. Pericarditis in Subjects aged ≥ 40 years (N=1756)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>538</td>
<td>388</td>
<td>8</td>
</tr>
<tr>
<td>No</td>
<td>474</td>
<td>338</td>
<td>10</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>1001</td>
<td>720</td>
<td>17</td>
</tr>
<tr>
<td>Pericarditis constrictive</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Pleurapericarditis</td>
<td>15</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>225</td>
<td>264</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>789</td>
<td>462</td>
<td>16</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>296</td>
<td>224</td>
<td>5</td>
</tr>
<tr>
<td>Dose 2</td>
<td>295</td>
<td>210</td>
<td>4</td>
</tr>
<tr>
<td>Dose 3</td>
<td>348</td>
<td>241</td>
<td>6</td>
</tr>
<tr>
<td>Dose 4</td>
<td>15</td>
<td>8</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>58</td>
<td>43</td>
<td>2</td>
</tr>
</tbody>
</table>

| Time to Onset n=1763                   |                     |                   |                      |
| ≤ 24 hours                             | 53                  | 33                | 0                    |
| 1-5 days                               | 190                 | 122               | 2                    |
| 6-13 days                              | 119                 | 99                | 0                    |
| 14-21 days                             | 77                  | 74                | 0                    |
| 22-31 days                             | 57                  | 45                | 3                    |
| 32-60 days                             | 64                  | 44                | 0                    |
| 61-180 days                            | 67                  | 48                | 0                    |
| 181-375 days                           | 17                  | 5                 | 0                    |
| Unknown                                | 374                 | 257               | 13                   |

| Event Outcomeb                         |                     |                   |                      |
| Fatal                                  | 5                   | 12                | 0                    |
| Not resolved                           | 355                 | 207               | 2                    |
| Resolved                               | 114                 | 121               | 2                    |
| Resolved with sequelae                 | 26                  | 21                | 1                    |
| Resolving                              | 215                 | 148               | 0                    |
| Unknown                                | 304                 | 218               | 13                   |

| Duration of eventc n=87, median: 28    |                     |                   |                      |
| Up to 3 days                           | 1                   | 7                 | 0                    |
| 4-6 days                               | 6                   | 2                 | 0                    |
| 7-10 days                              | 2                   | 2                 | 0                    |
| 11-26 days                             | 11                  | 11                | 0                    |
| 27-57 days                             | 6                   | 12                | 0                    |
| 58-180 days                            | 16                  | 8                 | 0                    |
| 181-265 days                           | 3                   | 0                 | 0                    |

a. All serious occurrences.
b. Multiple episodes of the same PT event were reported with a different clinical outcome in one case hence the sum of the events outcome exceeds the total number of PT events.
c. For those cases where the event resolved or resolved with sequelae.

Fatal Pericarditis cases in adult (40-50 years of age) (4 cases; 2 cases medically confirmed and 2 non-medically confirmed):

- 2 cases medically confirmed:
  - A 43-year-old male subject from Japan.
    - Medical history: Diabetes mellitus, Obesity.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Myocarditis, Pericarditis, Sudden death.
    - Time to onset (pericarditis and myocarditis): On the same day of receiving dose 3, the patient died.
    - Cause of death: Myocarditis, Pericarditis, Sudden death.
  - A 48-year-old male subject from Hong Kong.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Brain stem haemorrhage, Pericarditis.
- Time to onset (pericarditis): 17 days after dose 2.
- Cause of death: Both the above events.

- 2 cases non-medically confirmed:
  - A 41-year-old male subject from UK.
    - Medical history: Congestive cardiomyopathy, Huntington's disease, Positive airway pressure therapy, Sleep apnoea syndrome, Type 2 diabetes mellitus.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Interchange of vaccine products, Myocarditis, Off label use, Pericarditis, Sudden death.
    - Time to onset (pericarditis and myocarditis): 11.5 hours after dose 3, the patient died.
    - Cause of death: Myocarditis; Pericarditis; Sudden death.
  - A 49-year-old male subject from UK.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Aortic rupture, Back pain, Cardiomegaly, Internal haemorrhage, Myocarditis, Pericarditis, Pyrexia, Syncope, Vomiting.
    - Time to onset (pericarditis): ~50 days after dose 1, the patient died due to the above events.
    - Cause of death: Cardiomegaly.

Fatal Pericarditis cases in adult (51-64 years of age) (7 cases; 5 cases medically confirmed and 2 non-medically confirmed)

- 5 cases medically confirmed:
  - A 56-year-old male subject from Australia.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Malaise, Pericarditis.
    - Time to onset (pericarditis): On the same day of receiving dose 1.
    - Cause of death: Both the above events.
  - A 57-year-old female subject from Austria.
    - Medical history: Thyroid cancer.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Pericarditis.
    - Time to onset (pericarditis): Unspecified days after dose 3.
    - Cause of death: Pericarditis.
  - A 59-year-old female subject from Australia.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Atrial fibrillation, Atrioventricular block complete, Cardiac arrest, Chest pain, Electrocardiogram ST segment depression, Myocarditis, Pericarditis, Troponin increased.
    - Time to onset (pericarditis): 67 days after dose 3.
    - Cause of death: All the above events.
  - A 61-year-old female subject from Japan.
    - Medical history: Cerebrovascular accident, Syncope, Thymic carcinoma, Thymoma.
    - Co-suspect medications: None.
- PTs with fatal outcome: Cardio-respiratory arrest, Coronary artery stenosis, Endocarditis, Myocarditis, Pericarditis, Right ventricular failure, Sudden death.
- Time to onset (pericarditis): 11 days after dose 3.
- Cause of death: All the above events.
  - A 62-year-old male subject from Austria.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Cardiac failure, Pericarditis.
    - Time to onset (pericarditis): 14 days after dose 2.
    - Cause of death: Both the above events.

- 2 cases non-medically confirmed:
  - A 53-year-old male subject from Italy.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Pericarditis.
    - Time to onset (pericarditis): Within 7 days after dose 1.
    - Cause of death: Pericarditis.
  - A 62-year-old male subject from UK.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Abdominal pain upper, Cardiac arrest, Chest pain, Dizziness, Dyspnoea, Fatigue, Immunisation, Myocarditis, Pain in extremity, Palpitations, Pericarditis, Thrombosis.
    - Time to onset (pericarditis): 6 days after dose 3.
    - Cause of death: All the above clinical events.

Fatal Pericarditis cases in elderly (65-74 years of age) (2 cases, both non-medically confirmed)

- A 69-year-old female subject from UK.
  - Medical history: Arthralgia, Brain neoplasm, Hypertension.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Amnesia, Death, Interchange of vaccine products, Memory impairment, Myocarditis, Off label use, Pericarditis.
  - Time to onset (pericarditis and myocarditis): Unspecified days after the dose 3.
  - Causes of death: Brain neoplasm.

- A 71-year-old male subject from UK.
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Chest pain, Death, Dyspnoea, Fatigue, Myocarditis, Palpitations, Pericarditis, Pulmonary embolism, Thrombosis.
  - Time to onset (pericarditis and myocarditis): Unspecified days after the dose 2.
  - Causes of death: Death; Thrombosis.

Fatal Pericarditis cases in elderly (> 75 years of age -4 cases; 2 cases medically confirmed and 2 cases non-medically confirmed)

- 2 cases medically confirmed:
  - An 89-year-old female subject from Spain.
    - Medical history: Anaemia megaloblastic, Aphasia, Arthropathy, Atrial fibrillation, Benign tumour excision, Cardiac assistance device user, Cerebrovascular accident, Chronic gastritis, Cognitive disorder, Diverticulum,
Dyslipidaemia, Hypertension, Neoplasm, Oropharyngeal surgery, Parotitis, Salivary gland neoplasm, Sinus node dysfunction, Type 2 diabetes mellitus.

- Co-suspect medications: Influenza vaccine INACT SAG 3V.
- PTs with fatal outcome: Pericarditis.
- Time to onset (pericarditis): 2 days after dose 3.
- Causes of death: Pericarditis
  - A 92-year-old male subject from Japan.
    - Medical history: Unknown.
    - Co-suspect medications: None.
    - PTs with fatal outcome: Aortic dissection, Cardiac failure, Cardiac tamponade, Pericarditis.
    - Time to onset (pericarditis): Unspecified days after dose 3.
    - Causes of death: All the above events.
  - 2 cases not medically confirmed:
    - An 81-year-old male subject, from Italy.
      - Medical history: Unknown.
      - Co-suspect medications: None.
      - PTs with fatal outcome: Pericarditis.
      - Time to onset (pericarditis): 60 days after dose 3.
    - A 78-year-old male subject from Slovakia.
      - Medical history: Unknown.
      - Co-suspect medications: None.
      - PTs with fatal outcome: Multiple organ dysfunction syndrome, Myocardial infarction, Pericarditis.
      - Time to onset (pericarditis): 8 days after dose 3.
      - Causes of death: Due to all the above events.

**Rapporteur assessment comment:**

During the current reporting period, there were 1756 cases reporting pericarditis in persons aged ≥40 years compared to 2059 pericarditis cases reported in the previous 2\textsuperscript{nd} PSUR. There were 17 fatal cases compared to 8 fatal cases in the previous reporting period.

The MAH only briefly presented the 17 fatal cases and did not provide an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and no WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases with pericarditis in persons aged ≥40 years and perform per case an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, and a WHO causality assessment. **Request for supplementary information**

**Subjects with booster dose**

- Clinical Trial Data
  - Number of cases: none; one (1) case was retrieved in the PSUR #2.

- Post-Authorisation Data
  - Number of cases: 1216 (0.2% of 507,683 cases of the total PM dataset, 1.0% of the 117,750 subjects who received a booster dose), compared to 283 cases (0.04%) in the PSUR #2.
• Country of incidence: UK (474), France (202), Germany (94), Italy (93), Netherlands (46), New Zealand (38), Norway (32), Japan (30), Israel, Sweden (21 each); the remaining 165 cases were distributed among 24 countries.

• MC (500), NMC (716).

• Subjects’ gender: female (661), male (531), and unknown (24).

• Subjects’ age in year: n = 1130, range: 13 -93, mean: 45.1, median: 44.0

• Medical history (n = 566): the medical conditions reported more or equal to 10 times included the PTs Hypertension (79), Pericarditis (40), Asthma (38), Immunodeficiency (32), Hypothyroidism (29), Obesity (22), Diabetes mellitus, Drug hypersensitivity (19 each), Seasonal allergy (18), Depression, Steroid therapy, Tobacco user (15 each), Atrial fibrillation, Non-tobacco user, Type 2 diabetes mellitus (14 each), Anxiety, Dyslipidaemia, Gastroesophageal reflux disease (13 each), Clinical trial participant, Disease risk factor, Migraine (12 each), Myocardial infarction, Rheumatoid arthritis (11 each), Chronic kidney disease, Food allergy (10 each).

• COVID-19 Medical history (n = 114): Suspected COVID-19 (60), COVID-19 (55), Post acute COVID-19 syndrome (3), Exposure to SARS-CoV-2, SARS-CoV-2 test positive (1 each).

• Co suspects (n=20 cases): Influenza vaccine (6), Influenza vaccine INACT SAG 3V (3), Adalimumab, amoxicillin, Apixaban, Colchicine, COVID-19 vaccine, COVID-19 vaccine mRNA (MRNA 1273), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19), Etanercept, Propranolol, Saliutamol, Zuclopihexol (1 each).

• Number of relevant events: 1220.

• Relevant event seriousness: all serious.

• Reported relevant PTs: Pericarditis (1212), Pleuropericarditis (8)

• Relevant event outcome: fatal (12), resolved/resolving (414), resolved with sequelae (21), not resolved (296), unknown (478).

• Most frequently co-reported PTs (>3%): Chest pain (620), Myocarditis (461), Dyspnoea (448), Fatigue (427), Immunisation (418), Off label use (365), Palpitations (363), Interchange of vaccine products (332), Tachycardia (291), Pyrexia (218), Chest discomfort (151), Headache (124), Pericardial effusion (88), Malaise (87), Pain (82), Dizziness (74), Pain in extremity (66), Syncope (61), Arthralgia (58), Heart rate increased (57), Angina pectoris, Nausea (50 each), Asthenia (48), Arrhythmia (41), Chills, Lymphadenopathy, Myalgia (40 each), Back pain, Vaccination site pain (34 each), Cough (33).

The number of pericarditis cases occurred after a booster dose in each age group is reported in the below Table 53 by gender.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Heterologous Booster dose</th>
<th>Homologous Booster dose</th>
<th>Unknown dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. of Cases</td>
<td>No. of Cases</td>
<td>No. of cases</td>
</tr>
<tr>
<td><strong>Age group</strong></td>
<td>F  M  U</td>
<td>F  M  U</td>
<td>F  M  U</td>
</tr>
<tr>
<td>0 to 17 years</td>
<td>1  0  0</td>
<td>2  9  0</td>
<td>2  9  0</td>
</tr>
<tr>
<td>18 to 24 years</td>
<td>4  13 0</td>
<td>24 32 0</td>
<td>25 22 0</td>
</tr>
<tr>
<td>25 to 29 years</td>
<td>8  10 0</td>
<td>25 35 0</td>
<td>22 20 2</td>
</tr>
<tr>
<td>30 to 39 years</td>
<td>23 15 0</td>
<td>62 50 2</td>
<td>32 24 0</td>
</tr>
<tr>
<td>40 years and older</td>
<td>140 94 4</td>
<td>143 85 2</td>
<td>100 87 2</td>
</tr>
<tr>
<td>Unknown</td>
<td>26 8 5</td>
<td>16 12 5</td>
<td>6 6 2</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>202 140 9</td>
<td>272 223 9</td>
<td>187 168 6</td>
</tr>
</tbody>
</table>

F=female; M=male; U=unknown
Rapporteur assessment comment:

During the current reporting period, there were 1216 cases reporting pericarditis in persons who received a booster dose compared to 283 pericarditis cases reported in the previous 2nd PSUR. There were 12 fatal cases compared to 1 fatal case in the previous reporting period.

The 12 fatal cases are assumed to be imbedded in the fatal cases stated in the age categories above, which are subject for a request for supplementary information.

During the reporting period, of the 4156 cases reported, there were 1319 cases of medically confirmed pericarditis with a latency 21 days or less, of which in 975 cases pericarditis occurred within 1 week post vaccine administration. The majority (1255) of the cases were assessed as serious due to hospitalisation and/or medically significant. In 58 other cases, the seriousness criterion was reported as disability or life threatening, and in 6 cases, a fatal outcome was reported, which are reviewed above in the age-stratified sections.

Rapporteur assessment comment:

During the current reporting period, there were 1,319 cases reporting medically confirmed pericarditis with a TTO 21 days or less compared to 1,461 medically confirmed pericarditis cases reported in the previous 2nd PSUR.

Observed versus Expected analyses

The MAH conducted analyses for myocarditis and myocarditis/pericarditis stratified by age and sex, and by dose. Since background incidence rates stratified by age and sex were not available for pericarditis alone, analyses performed for a combined myocarditis/pericarditis (table 17 and 18).

Spontaneously reported cases of myocarditis and myocarditis/pericarditis were limited to those occurring in EEA countries or the United States. Cases were also limited to those with time to onsets occurring within the 14-day (results not reproduced here) or 21-day risk windows (table 15 and 16) to increase the sensitivity for signal detection for these events.
<table>
<thead>
<tr>
<th>Stratification</th>
<th>Person Years</th>
<th>Obs Cases</th>
<th>Background rate at 100,000</th>
<th>Exp Cases</th>
<th>O/E Ratio</th>
<th>95% CI</th>
<th>Background rate at 100,000</th>
<th>Exp Cases</th>
<th>O/E Ratio</th>
<th>95% CI</th>
<th>Background rate at 100,000</th>
<th>Exp Cases</th>
<th>O/E Ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Males &lt;51 years</td>
<td>513,444</td>
<td>14</td>
<td>0.48</td>
<td>2.5</td>
<td>5.681</td>
<td>9.531</td>
<td>3.186</td>
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<td>4.40</td>
<td>22.6</td>
<td>0.630</td>
<td>1.040</td>
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<tr>
<td>Males 51-64 years</td>
<td>918,902</td>
<td>604</td>
<td>0.48</td>
<td>4.4</td>
<td>136.939</td>
<td>148.30</td>
<td>12.734</td>
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<td>4.40</td>
<td>40.4</td>
<td>14.939</td>
<td>16.179</td>
</tr>
<tr>
<td>Males 65-74 years</td>
<td>1,346,972</td>
<td>990</td>
<td>2.77</td>
<td>1.37</td>
<td>26.373</td>
<td>28.281</td>
<td>19.943</td>
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<td></td>
<td></td>
<td>4.40</td>
<td>59.2</td>
<td>16.729</td>
<td>17.804</td>
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<tr>
<td>Males 75-84 years</td>
<td>5,884,458</td>
<td>1,271</td>
<td>2.48</td>
<td>1.45</td>
<td>9.799</td>
<td>10.382</td>
<td>8.237</td>
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<td></td>
<td></td>
<td>4.40</td>
<td>258.9</td>
<td>4.909</td>
<td>5.186</td>
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<tr>
<td>Males 85-94 years</td>
<td>2,797,358</td>
<td>729</td>
<td>2.66</td>
<td>4.16</td>
<td>6.008</td>
<td>6.756</td>
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<td>4.40</td>
<td>123.1</td>
<td>2.267</td>
<td>2.549</td>
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<tr>
<td>Males 95+ years</td>
<td>2,371,600</td>
<td>137</td>
<td>2.37</td>
<td>3.52</td>
<td>2.437</td>
<td>2.581</td>
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<td>4.40</td>
<td>104.4</td>
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<tr>
<td>Females &lt;51 years</td>
<td>578,990</td>
<td>4</td>
<td>0.08</td>
<td>0.05</td>
<td>8.636</td>
<td>9.211</td>
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<td>4.40</td>
<td>20.5</td>
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<tr>
<td>Females 51-64 years</td>
<td>1,936,208</td>
<td>92</td>
<td>0.08</td>
<td>0.8</td>
<td>10.982</td>
<td>13.109</td>
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<td>4.40</td>
<td>45.6</td>
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<tr>
<td>Females 65-74 years</td>
<td>1,316,571</td>
<td>202</td>
<td>0.47</td>
<td>1.12</td>
<td>17.998</td>
<td>20.559</td>
<td>13.602</td>
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<td></td>
<td>4.40</td>
<td>66.7</td>
<td>3.027</td>
<td>3.474</td>
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<tr>
<td>Females 75-84 years</td>
<td>6,635,676</td>
<td>694</td>
<td>0.72</td>
<td>0.47</td>
<td>14.526</td>
<td>15.638</td>
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<td>4.40</td>
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<tr>
<td>Females 85-94 years</td>
<td>3,154,468</td>
<td>247</td>
<td>0.87</td>
<td>0.36</td>
<td>8.072</td>
<td>9.144</td>
<td>1.565</td>
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<td>4.40</td>
<td>138.8</td>
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<tr>
<td>Females 95+ years</td>
<td>2,672,357</td>
<td>108</td>
<td>1.48</td>
<td>3.96</td>
<td>2.729</td>
<td>3.294</td>
<td>2.715</td>
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<td>4.40</td>
<td>117.7</td>
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<td>1.108</td>
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<tr>
<td>Overall, &lt;51 years</td>
<td>14,052,364</td>
<td>1,193</td>
<td>1.34</td>
<td>1.83</td>
<td>6.867</td>
<td>7.251</td>
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<td>4.40</td>
<td>618.3</td>
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<tr>
<td>Overall, 51-64 years</td>
<td>13,858,666</td>
<td>6</td>
<td>2.66</td>
<td>1.85</td>
<td>14.382</td>
<td>13.814</td>
<td>13.841</td>
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<td>4.40</td>
<td>608.9</td>
<td>4.380</td>
<td>4.549</td>
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<tr>
<td>Overall, 65-74 years</td>
<td>9,287,821</td>
<td>643</td>
<td>1.34</td>
<td>1.25</td>
<td>7.099</td>
<td>7.580</td>
<td>6.641</td>
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<td>4.40</td>
<td>413.5</td>
<td>2.162</td>
<td>2.308</td>
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</table>

a. Background rate per 100,000 person years
<table>
<thead>
<tr>
<th>Stratification</th>
<th>Person Years</th>
<th>Obs Cases</th>
<th>Low Bkgd rate&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Exp Case</th>
<th>O/E Ratio</th>
<th>95% CI</th>
<th>Mid Bkgd rate&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Exp Case</th>
<th>O/E Ratio</th>
<th>95% CI</th>
<th>High Bkgd rate&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Exp Case</th>
<th>O/E Ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Males &lt;11 years</td>
<td>387,938</td>
<td>2</td>
<td>0.48 1.9 1.074 3.888&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 171</td>
<td>1.117</td>
<td>0.014</td>
<td>8.26 32.0 0.062 0.225&lt;sup&gt;b&lt;/sup&gt;</td>
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</tr>
<tr>
<td>Males 12-17 years</td>
<td>643,410</td>
<td>63</td>
<td>0.48 3.1 20.399 26.099&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 283</td>
<td>2.225</td>
<td>0.116</td>
<td>8.26 53.1 1.185 1.517&lt;sup&gt;b&lt;/sup&gt;</td>
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</tr>
<tr>
<td>Males 18-24 years</td>
<td>879,957</td>
<td>80</td>
<td>2.77 24.4 3.782 4.085&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 387</td>
<td>2.066</td>
<td>0.557</td>
<td>20.20 177.8 0.450 0.560&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Males 25-49 years</td>
<td>3,302,205</td>
<td>83</td>
<td>2.48 81.9 1.013 1.156&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 145.3</td>
<td>0.571 0.708&lt;sup&gt;b&lt;/sup&gt;</td>
<td>14.97 494.3 0.168 0.268&lt;sup&gt;b&lt;/sup&gt;</td>
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</tr>
<tr>
<td>Males 50-59 years</td>
<td>1,444,518</td>
<td>13</td>
<td>1.66 24.0 0.542 0.977&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 63.6</td>
<td>2.050 0.358&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8.48 122.5 0.306 0.181&lt;sup&gt;b&lt;/sup&gt;</td>
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</tr>
<tr>
<td>Males 60-69 years</td>
<td>1,342,014</td>
<td>11</td>
<td>2.37 31.8 0.346 0.619&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 59.0</td>
<td>2.086 0.333&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8.40 64.4 0.171 0.360&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Males 70+ years</td>
<td>1,461,865</td>
<td>17</td>
<td>2.47 36.1 0.471 0.754&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 64.3</td>
<td>2.064 0.423&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.31 63.0 0.270 0.432&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females &lt;11 years</td>
<td>437,492</td>
<td>2</td>
<td>0.08 0.3 5.715 28.644&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 19.2</td>
<td>0.884 0.375&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.42 6.2 0.322 1.163&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 12-17 years</td>
<td>725,547</td>
<td>14</td>
<td>0.08 0.6 24.120 48.469&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 31.9</td>
<td>0.639 0.736&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.42 10.3 1.359 2.980&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 18-24 years</td>
<td>992,292</td>
<td>8</td>
<td>0.74 7.3 1.089 2.147&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 43.7</td>
<td>0.183 0.361&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.55 45.1 0.177 0.349&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 25-49 years</td>
<td>3,723,763</td>
<td>47</td>
<td>0.72 26.8 1.753 2.331&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 161.8</td>
<td>0.287 0.381&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.97 147.8 0.313 0.423&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 50-59 years</td>
<td>1,628,924</td>
<td>19</td>
<td>0.17 15.8 1.262 1.878&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 71.7</td>
<td>0.265 0.414&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.46 56.4 0.337 0.526&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 60-69 years</td>
<td>1,513,335</td>
<td>13</td>
<td>1.48 22.4 0.589 0.993&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 66.6</td>
<td>0.195 0.334&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.20 63.6 0.205 0.350&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Females 70+ years</td>
<td>1,648,486</td>
<td>13</td>
<td>0.76 12.5 1.088 1.774&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 72.5</td>
<td>0.179 0.306&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.84 63.3 0.205 0.351&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Overall, female</td>
<td>8,864,722</td>
<td>121</td>
<td>1.34 8.0 1.019 1.217&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 300.0</td>
<td>0.310 0.371&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.27 555.8 0.218 0.260&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Overall, male</td>
<td>5,491,992</td>
<td>552</td>
<td>0.552 0.952 1.088 1.774&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 72.5</td>
<td>0.179 0.306&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.84 63.3 0.205 0.351&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td>Overall, both</td>
<td>14,356,714</td>
<td>1,345</td>
<td>1.34 8.9 1.019 1.217&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4.40 300.0</td>
<td>0.310 0.371&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.27 555.8 0.218 0.260&lt;sup&gt;b&lt;/sup&gt;</td>
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a. Background rate per 100,000 person-years

PRAC PSUR assessment report
EMA/PRAC/849/2023 Page 171/272
Table 17. Observed to Expected (O/E) Analysis of Myocarditis/Pericarditis in European Economic Area Countries, Cumulative Period

<table>
<thead>
<tr>
<th>Stratification</th>
<th>14-Day Risk Window</th>
<th>21-Day Risk Window</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Bkgd Cases</td>
<td>Exp Cases</td>
</tr>
<tr>
<td>Males ≤11 years</td>
<td>16.77</td>
<td>3</td>
</tr>
<tr>
<td>Males 12-17 years</td>
<td>16.77</td>
<td>3</td>
</tr>
<tr>
<td>Males 18-24 years</td>
<td>16.77</td>
<td>3</td>
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<tr>
<td>Males 25-49 years</td>
<td>16.77</td>
<td>3</td>
</tr>
<tr>
<td>Males 50-69 years</td>
<td>16.77</td>
<td>3</td>
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<tr>
<td>Males 70+ years</td>
<td>16.77</td>
<td>3</td>
</tr>
<tr>
<td>Females ≤11 years</td>
<td>1.29</td>
<td>4</td>
</tr>
<tr>
<td>Females 12-17 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 18-24 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 25-49 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 50-69 years</td>
<td>1.30</td>
<td>4</td>
</tr>
<tr>
<td>Females 70+ years</td>
<td>1.30</td>
<td>4</td>
</tr>
<tr>
<td>Overall, dose 1</td>
<td>1.88</td>
<td>3</td>
</tr>
<tr>
<td>Overall, dose 2</td>
<td>1.88</td>
<td>3</td>
</tr>
<tr>
<td>Overall, dose 3</td>
<td>1.88</td>
<td>3</td>
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</tbody>
</table>

Table 18. Observed to Expected (O/E) Analysis of Myocarditis/Pericarditis in the United States, Cumulative Period

<table>
<thead>
<tr>
<th>Stratification</th>
<th>14-Day Risk Window</th>
<th>21-Day Risk Window</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Bkgd Cases</td>
<td>Exp Cases</td>
</tr>
<tr>
<td>Males ≤11 years</td>
<td>16.77</td>
<td>3</td>
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<tr>
<td>Males 12-17 years</td>
<td>16.77</td>
<td>3</td>
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<tr>
<td>Males 18-24 years</td>
<td>16.77</td>
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<tr>
<td>Males 25-49 years</td>
<td>16.77</td>
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<tr>
<td>Males 50-69 years</td>
<td>16.77</td>
<td>3</td>
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<tr>
<td>Males 70+ years</td>
<td>16.77</td>
<td>3</td>
</tr>
<tr>
<td>Females ≤11 years</td>
<td>1.30</td>
<td>4</td>
</tr>
<tr>
<td>Females 12-17 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 18-24 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 25-49 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 50-69 years</td>
<td>1.30</td>
<td>4</td>
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<tr>
<td>Females 70+ years</td>
<td>1.30</td>
<td>4</td>
</tr>
<tr>
<td>Overall, dose 1</td>
<td>1.88</td>
<td>3</td>
</tr>
<tr>
<td>Overall, dose 2</td>
<td>1.88</td>
<td>3</td>
</tr>
<tr>
<td>Overall, dose 3</td>
<td>1.88</td>
<td>3</td>
</tr>
</tbody>
</table>

Consistent with the analyses in the most recent SBSR, for myocarditis in the EEA, all O/E ratios were above 1 across age groups, genders, and doses, using the low background rate. This was also true for most age groups other than the youngest and oldest in both genders using the mid and high background rates.
For myocarditis in the US, O/E ratios were above 1 for all stratifications except males 50+, females 60-69 years, and dose 3 using the low background rate, for males 12-24 years using the mid background rate, and for males and females 12-17 years using the high background rate.

Recent increases in O/E ratios for the younger age groups may have been influenced by increased reporting of cases after the release of a Dear Healthcare Provider letter in late July 2021.

For myocarditis/pericarditis, the O/E ratios were above 1 for the 12-24 years age groups in males, the 12-59 years age groups in females, and dose 1, 2, and 3 in the EEA. All O/E ratios were below 1 for myocarditis/pericarditis in the US except for females 12-17 years.

**Rapporteur assessment comment:**

**Myocarditis in EEA**
Cumulatively using 21-days risk window and the low background rate, all O/E ratios were above 1 across age groups, genders, and doses.

**Myocarditis/pericarditis in EEA**
Cumulatively using 14-days or 21-days or risk window, the O/E ratios were above 1 for males aged 12-24 years, females aged 12-59 years, and dose 1, 2, and 3.

**Rapporteur assessment comment:**

In general, the MAH should focus the analysis of pericarditis cases on aspects of these ADRs not fully known or addressed in the Conimly PI (pericarditis is already an ADR stated in section 4.8), and if the warning in section 4.4 regarding pericarditis is still in line with current knowledge. Therefore, the analysis should focus more on information concerning the course, outcome, and possible risk factors of the pericarditis cases following Conimly exposure. **Request for next PSUR**

**Pericarditis**

**Clinical trial data**

During the reporting period, no cases were retrieved.

**Post-marketing**

- Aged 5-11 years: There were 30 cases reporting pericarditis (no fatal cases) compared to 4 pericarditis cases (no fatal cases) reported in the previous 2nd PSUR.

- Aged 12-15 years: There were 118 cases reporting pericarditis (no fatal cases) compared to 215 pericarditis cases (no fatal cases) reported in the previous 2nd PSUR.

- Aged 16-17 years: There were 106 cases reporting pericarditis (no fatal cases) compared to 174 pericarditis cases (no fatal cases) reported in the previous 2nd PSUR.

- Aged 18-24 years: There were 479 cases reporting pericarditis (1 fatal case) compared to 659 pericarditis cases (no fatal cases) reported in the previous 2nd PSUR.

- Aged 25-29 years: There were 417 cases reporting pericarditis (1 fatal case) compared to 614 pericarditis cases (no fatal cases) reported in the previous 2nd PSUR.

- Aged 30-39 years: There were 940 cases reporting pericarditis (no fatal cases) compared to 1222 pericarditis cases (1 fatal case) reported in the previous 2nd PSUR.
- Aged ≥40 years: There were 1756 cases reporting pericarditis (17 fatal cases) compared to 2059 pericarditis cases (8 fatal cases) reported in the previous 2nd PSUR.

The MAH is requested to provide detailed information concerning the fatal cases with pericarditis in persons aged 18–24 years, 25–29 years, and ≥40 years and perform per case an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable. Request for supplementary information

Rapporteur assessment comment:

In spite of the recent risk communication and routine risk minimisation measures concerning myocarditis and pericarditis fatal cases continue to be reported, which is worrisome. Upon analysis of the requested details of the fatal myocarditis/pericarditis cases the MAH is expected to discuss whether the current risk minimisation measures is still adequate. Request for supplementary information

Evaluation of important potential risks

**Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)**

Search criteria:

1 - PTs Vaccine associated enhanced respiratory disease OR Vaccine associated enhanced disease

OR

2 - Standard Decreased Therapeutic Response Search (Drug Ineffective OR Vaccination failure) AND 1 among the following PTs: Dyspnoea; Tachypnoea; Hypoxia; COVID 19 pneumonia; Respiratory failure; Acute respiratory distress syndrome; Cardiac failure; Cardiogenic shock; Acute myocardial infarction; Arrhythmia; Myocarditis; Vomiting; Diarrhoea; Abdominal pain; Jaundice; Acute hepatic failure; Deep vein thrombosis; Pulmonary embolism; Peripheral ischaemia; Vasculitis; Shock; Acute kidney Injury; Renal failure; Altered state of consciousness; Seizure; Encephalopathy; Meningitis; Cerebrovascular accident; Thrombocytopenia; Disseminated intravascular coagulation; Chilblains; Erythema multiforme; Multiple organ dysfunction syndrome; Multisystem inflammatory syndrome in children.

Clinical trial data

- There were no cases reporting COVID-19 infection associated to one of the PTs utilised to identify potential severe or atypical cases of COVID-19.

Post-Authorisation Data

- Number of cases: 1268 (0.2% of 507,683 cases, the total PM dataset), compared to 1490 (0.2%) retrieved in the PSUR # 2. All cases are serious.
- MC cases (878), NMC cases (390).
- Country of incidence: France (346), Spain (142), UK (139), US (117), Italy (105), Estonia (94), Germany (66), Philippines (45), Australia, Canada (19 each), Switzerland (18), Portugal,
(17), Netherlands (14), Austria (10); the remaining 117 cases originated from 117 different countries.

- Gender: female (636), male (604), and unknown (28).
- Age in years (n = 1215), range: 5 – 102, mean: 61.4, median: 65.0.
- Relevant event seriousness: 1295 serious, 406 non-serious.
- Reported relevant PTs by organ system:
  
  o Respiratory system PTs (1631): COVID-19 pneumonia (524), Dyspnoea (398), Respiratory failure (48), Acute respiratory distress syndrome (42), Pulmonary embolism (40), Hypoxia (28), and Tachypnoea (27).
  
  o Gastrointestinal/Hepatic system PTs (288): Diarrhoea (139), Vomiting (88), Abdominal pain (54), and Jaundice (7).
  
  o Cardiovascular system PTs (143): Myocarditis (85), Arrhythmia (32), Cardiac failure (18), Acute myocardial infarction (6), and Cardiogenic shock (2).
  
  o Renal and urinary system PTs (39): Acute kidney injury (27), and Renal failure (12).
  
  o Nervous system PTs (47): Seizure (21), Cerebrovascular accident (18), Encephalopathy (6), and Altered state of consciousness (2).
  
  o Vascular system PTs (23): Deep vein thrombosis (12), Shock (6), Vasculitis (3), and Peripheral ischaemia (2).
  
  o Blood and lymphatic system PTs (14): Thrombocytopenia (12), and Disseminated Intravascular coagulation (2).
  
  o Immune system PTs (30): Vaccine associated enhanced disease (12), and Multisystem inflammatory syndrome in children (9), and Vaccine associated enhanced respiratory disease (9 each).
  
  o Other PTs (24): Multiple organ dysfunction syndrome (13), Chilblains (5), Meningitis (4), and Erythema multiforme (2).

- Case outcome: fatal (184), not resolved (329), resolved/resolving (582), resolved with sequelae (31), and unknown (142).

MAH’s conclusion

The nature of spontaneously reported data provides a challenge because of the lack of a comparison group. Further, the background rate of VAED is not known. Considering the limitations of the review, VAED/VAERD remains a theoretical risk for the vaccine. Surveillance will continue.

**Rapporteur assessment comment:**

No new important safety concern could be identified regarding VAED/VAERD.

**Evaluation of Other Risks (not categorised as important)**

*Adverse events of special interest (AESIs)*

**Anaphylactic AESIs**

**Rapporteur assessment comment:**
Please refer to the section 2.3 ‘Evaluation of important identified risks’ of this assessment report.

**Cardiovascular AESIs**

- **Search criteria:** PTs Acute myocardial infarction; Arrhythmia; Cardiac failure; Cardiac failure acute; Cardiogenic shock; Coronary artery disease; Myocardial infarction; Postural orthostatic tachycardia syndrome; Stress cardiomyopathy; Tachycardia

**Clinical trial data**

- Number of cases: 27 (blinded therapy [6], and BNT162b2 [21]) (4.0 % of 668 cases, the total CT dataset) compared to 35 cases (4.9%) retrieved in the PSUR #2. None of the events were related BNT162b2 or blinded therapy.

**Post-authorization data**

- Number of cases: 32,712 (6.4 % of 507,683 cases, the total PM dataset), compared to 29,486 (4.5%) cases retrieved in the PSUR #2.
- MC cases (11,952), NMC cases (20760).
- Country of incidence (>16 occurrences): Germany (11,180), Australia (4456), UK (3049), France (2612), Taiwan Province of China (1393), Italy (1334), Netherlands (810), Austria (677), Malaysia (591), Philippines (475), US (422), Japan (397), New Zealand (396), Norway (369), Finland (338), Sweden (333), Belgium (328), Canada (317), Poland (307), Iraq (298), Greece (275), Spain (271), Ireland (237), Romania (217), Czech Republic (198), Denmark (125), Switzerland (119), Brazil (118), Lithuania (110), Croatia (92), Estonia (88), Portugal (82), Egypt (77), Israel (74), Slovenia (70), Mexico (62), Iceland (52), Slovakia (51), Hungary (50), Singapore (35), South Africa (30), Georgia (24), Latvia (24), Luxembourg (22), Turkey (20), Bulgaria (18), Cyprus (17); the remaining 72 cases were distributed among 31 countries.
- Subjects' gender: female (19,730), male (12,424) and unknown (558).
- Subjects' age in years (n = 31,124), range: 2 months-99, mean: 40.3, median: 39.
- Medical history (n = 9348): the most frequently (>2%) reported relevant medical conditions included Hypertension (1349), Seasonal allergy (1121), Asthma (839), Drug hypersensitivity (758), Hypersensitivity (511), Food allergy (502), Mite allergy (387), Hypothyroidism (365), Tobacco user (290), Allergy to animal (285), Autoimmune thyroiditis (280), Diabetes mellitus (274), Obesity (253), Atrial fibrillation (232), Non-tobacco user (232), Arrhythmia (220), Allergy to metals (216), Migraine (214).
- COVID-19 Medical history (n = 1546): the medical conditions reported included COVID-19 (1021), Suspected COVID-19 (492), Post-acute COVID-19 syndrome (40), COVID-19 pneumonia (16), SARS-CoV-2 test positive (14), Coronavirus infection (7), Asymptomatic COVID-19 (6), Exposure to SARS-CoV-2 (4), and Coronavirus pneumonia (1).
- Co suspects (n = 295 cases): the frequently (>12 occurrences) reported relevant co-suspect medications were COVID-19 vaccine mRNA (MRNA 1273) (66), COVID-19 vaccine (34), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (20), INFLUENZA VACCINE (19), adalimumab (13).
- Number of relevant events: 36,790.
- Relevant event seriousness: serious (16,539), non-serious (20,268).
- Relevant PTs: Chest pain (17,945), Tachycardia (10,914), Arrhythmia (5627), Myocardial infarction (921), Cardiac failure (583), Acute myocardial infarction (364), Postural orthostatic tachycardia syndrome (149), Coronary artery disease (114), Cardiogenic shock (72), Cardiac failure acute (57), Stress cardiomyopathy (44).

- Time to event onset (n = 26,744 occurrences), range: <24 hours to 382 days, median: 1 day.

- Duration of relevant events (n = 8262 out of 8906 occurrences with outcome of resolved and resolved with sequelae), range: <24 hours to 430 days, median: 4 days.

- Relevant event outcome: fatal (496), resolved/resolving (13,937), resolved with sequelae (1321), not resolved (12,839), unknown (8437).
  - In 449 cases (reporting 496 relevant events with fatal outcome), the reported causes of death (>10 occurrences) were coded to the PTs Myocardial Infarction (147), Cardiac failure (94), Chest pain (55), Acute myocardial infarction (54), Dyspnoea (43), Cardiac arrest (42), Arrhythmia (34), Cardiac failure acute (26), Myocarditis (22), Cardiogenic shock, Cardio-respiratory arrest (20 each), Thrombosis (15), Malaise (13), Loss of consciousness, Pulmonary embolism, Pulmonary oedema, Tachycardia (12 each), Pneumonia, Respiratory failure (11 each). Of note, in 16 cases limited information regarding the cause of death was provided (PT Death [11]; PT Sudden death [1]; Unknown [4]). Most (250 of 449 cases) of the fatal cases involved elderly subjects. When the medical history was provided (253 cases), the most frequently (≥ 9 occurrences) relevant medical conditions included events coded to the PTs Hypertension (91), Diabetes mellitus (32), Atrial fibrillation (28), Obesity (24), Cardiac failure (23), Type 2 diabetes mellitus (16), Coronary artery disease (15), Dyslipidaemia, Myocardial Infarction (13 each), Chronic kidney disease, Chronic obstructive pulmonary disease (12 each), Cardiac disorder, Tobacco user (11 each), Cardiac failure chronic, Hyperlipidaemia (10 each), Arteriosclerosis, Asthma, Cerebral infarction, Myocardial ischaemia (9 each).

**Analysis by age group**

- Clinical trials: Paediatric (1), Adults (14), and Elderly (12). A meaningful comparison between the different age groups is not possible due to the low number of cases.

- Post-marketing: Paediatric (2808), Adults (25850), Elderly (2996) and Unknown (1058).
  - Higher reporting proportion of events coded to the PTs Arrhythmia [4.6% in paediatrics vs 18.2% in adults vs 24.3% in elderly], Cardiac failure [0.5% in paediatrics vs 0.8% in adults vs 11.5% in elderly], Myocardial infarction [0.3% in paediatrics vs 2.2% in adults vs 9.0% in elderly], Cardiogenic shock [0.1% in paediatrics vs 0.2% in adults vs 0.8% in elderly], Acute myocardial infarction [0.1% in paediatrics vs 0.8% in adults vs 4.8% in elderly], Cardiac failure acute [0.1% in paediatrics vs 0.1% in adults vs 0.8% in elderly], Stress cardiomyopathy [0.04% in paediatrics vs 0.1% in adults vs 0.7% in elderly], and Coronary artery disease [0% in paediatrics vs 0.3% in adults vs 1.5% in elderly] and was reported in elderly population when compared to adult and paediatric population.

Higher reporting proportion of PT Chest pain [81.6% in paediatrics vs 53.5% in adults vs 36.3% in elderly] was reported in paediatrics compared to adults and elderly subjects.
Higher reporting proportion of PT Tachycardia [19.6% in paediatrics vs 36.5% in adults vs 22.7% in elderly] was reported in adults compared to paediatrics and elderly subjects.

The PT Postural orthostatic tachycardia syndrome was reported among the paediatric and adult subjects only (0.4% each).

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 3726 (0.7% of 507,683 cases, the total dataset).
- No significant difference was observed in the reporting proportion of cardiovascular AESIs with fatal outcome in individuals with comorbid conditions (0.4% of events with fatal outcome) when compared to the reporting proportion observed in the individuals without comorbidities (0.9 % of events with fatal outcome).

**O/E analysis**

- O/E analysis was performed for Acute myocardial infarction/Myocardial infarction; Arrhythmia; Coronary artery disease; Heart failure; Postural orthostatic tachycardia syndrome; Stress cardiomyopathy. All O/E ratios were <1.

**MAH’s conclusion**

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

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**Rapporteur assessment comment:**

During the interval period, post-marketing there were retrieved 32,712 cases (6.4 % of 507,683 cases, the total PM dataset) reporting cardiovascular AESIs which is an increase compared to 29,486 (4.5%) cases retrieved in the previous 2nd PSUR. Of note, during the current interval period the overall (all ages) exposure decreased compared to the previous interval period: 843 Mio doses versus 1.4 billion doses. However, the paediatric exposure increased from 47 Mio paediatric Tris/Sucrose doses shipped worldwide in the previous interval period to 182 Mio paediatric Tris/Sucrose doses in current interval period.

The O/E analysis showed that all O/E ratios were below 1 and age stratified O/E ratios were also <1, when available.

No new important safety concern could be identified for cardiovascular AESIs. For future PSURs in the section 'Evaluation of AESIs', the cardiovascular AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

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**Haematological AESIs**

Search Criteria: *Leukopenias NEC; Neutropenias OR SMQ Haemorrhage terms (excl laboratory terms) OR PT Acquired haemophilia.*

**Clinical trial data**

- Number of cases: 15 (BNT162b2 [12] and blinded therapy [3]) (2.2 % of 668 cases, the total CT dataset) compared to 19 cases (2.4%) retrieved in the PSUR #2. None of these SAEs were assessed as related to BNT162b2/blinded therapy.

**Post-authorisation data**
- Number of cases: 30,302 (5.9% of 507,683 cases, the total PM dataset), compared to 37,327 cases (5.7%) retrieved in the PSUR #2.

- MC cases (4952), NMC cases (25,350).

- Country of incidence: Germany (7802), Netherlands (6166), UK (3266), Norway (2930), France (2905), Australia (1010), Spain (626), Italy (579), Sweden (557), Belgium (438); the remaining 4023 cases were distributed among 63 countries.

- Subjects' age in years (n = 28,488), range: 5 months-100 years, mean: 38.9, median: 37.0.

- Medical history (n = 10,294): the most frequently (≥200 occurrences) reported relevant medical conditions were coded to the PTs Disease risk factor (957), Hypertension (648), Menopause (620), Asthma (508), Seasonal allergy (427), Drug hypersensitivity (419), Amenorrhoea (404), Hypersensitivity (379), Hypothyroidism (295), Endometriosis (225), Food allergy (223), Migraine (213), and Contraception (200).

- COVID-19 Medical history (n = 2397): Medical conditions reported more than once were coded to the PTs COVID-19 (1636), Suspected COVID-19 (730), Post-acute COVID-19 syndrome (12), SARS CoV 2 test positive (8), COVID-19 pneumonia (5), Coronavirus Infection, Exposure to SARS CoV 2 (2 each), Asymptomatic COVID-19, and Coronavirus test positive (1 each).

- Co-suspects: the most frequently (≥10 occurrences) reported relevant co-suspect vaccines/medications were COVID-19 vaccine mRNA (97), adalimumab (43), Influenza vaccine (36), COVID-19 vaccine NRVV (31), levonorgestrel (24), and COVID-19 vaccine (19).

- Number of relevant events: 33,677.

- Relevant event seriousness: serious (8090) and non-serious (25,587).

- Most frequently reported relevant PTs (≥2%): Heavy menstrual bleeding (12,905), Intermenstrual bleeding (6088), Vaginal haemorrhage (1759), Epistaxis (1645), Contusion (1450), Vaccination site haematoma (1137), Postmenopausal haemorrhage (1137), Haematoma (944), and Haemorrhage (677).

- Time to event onset (n = 24,005 events), range: <24 hours to 7337 days, median: 3 days.

- Duration of relevant events (n = 240 out of 572 occurrences with outcome of resolved/resolved with sequela), range: 1 day to 21,170 days.

- Relevant event outcome: fatal (146), resolved/resolving (11,605), resolved with sequela (571), not resolved (13,999), and unknown (7480).

  - In the 174 fatal cases (including 146 relevant events with fatal outcome, reported in 114 cases), the reported causes of death (>8 occurrences) were coded to the PTs Haemorrhage (12), Gastrointestinal haemorrhage, Haematemesis, and Pericardial haemorrhage (9 each). Of note, in 19 cases limited information regarding the cause of death was provided (PT Death). Most (122 of 174 cases) of the fatal cases involved elderly subjects. When the medical history was provided (114 cases), the most frequently (≥10 occurrences) relevant medical conditions included the PTs Hypertension (44), Cardiac arrest (16), Myocardial infarction (14), Cardiac failure, Cardio-respiratory arrest, Haemorrhage, and Myocardial ischaemia (10 each).

Analysis by age group

- Clinical trials: Adults (9) and Elderly (5). A meaningful comparison between the different age groups is not possible due to the low number of cases.
Post-marketing: Paediatric (1044), Adults (26,592), Elderly (1731) and Unknown (935).

- A significantly higher reporting proportion of events coded to the PTs Heavy menstrual bleeding and Intermenstrual bleeding was observed in paediatric and adult population when compared to elderly population (Heavy menstrual bleeding [33.9% in paediatrics vs 45.7% in adults vs 0.2% in elderly] and Intermenstrual bleeding [8.3% in paediatrics vs 22.1% in adults vs 1.2% in elderly]).

The reporting proportion of the PT Epistaxis was significantly higher in paediatric and elderly population when compared to adult population (21.3% in paediatrics vs 14.3% in elderly vs 4.2% in adults).

The reporting proportion of PT Haematoma was higher in elderly population (12.1%) when compared to paediatrics (1.2%) and adult (2.0%) population.

The comparative differences in reporting proportions are not unexpected given the generally expected medical issues affecting each age group (paediatrics, adults, elderly).

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 2542 (8.4% of the CT and PM cases reporting haematological AESIs).

- The reporting proportion of haematological AESIs with fatal outcome (1.6%) is higher in individuals with comorbid conditions when compared to the reporting proportion observed in the individuals without comorbidities (0.3%).

O/E analysis

- O/E analysis was performed for Acquired haemophilia and Haemorrhage. All O/E ratios below 1.

MAH's conclusion

Acquired haemophilia was evaluated during this reporting period. No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:

Please refer regarding Heavy menstrual bleeding to the separate signal procedure (EMEA/H/C/005735/SDA/053- EPITT 19783), in which PRAC concluded that heavy menstrual bleeding should be listed as an ADR in the Comirnaty PI.

Please refer regarding the assessment of acquired haemophilia to section 2.2 Signal evaluation of this AR.

Haematological AESIs were continuously monitored in the 13th and 14th SSRs (through 15 Apr 2022) for Comirnaty during the reporting period.

No new important safety concern could be identified for haematological AESIs. For future PSURs in the section 'Evaluation of AESIs', the haematological AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

COVID-19 AESIs
Search criteria: SMQ COVID-19 (Narrow and Broad) OR PTs Ageusia; Anosmia; Vaccine associated enhanced disease; Vaccine associated enhanced respiratory disease.

Clinical trial data

- Number of cases: 7 (blinded therapy [3] and BNT162b2 [4]) (1.0% of 668 cases, the total CT dataset) compared to 3 cases (0.4%) retrieved in the PSUR #2. None of the events were related to BNT162b2 or blinded therapy.

Post-authorisation data

- Number of relevant cases: 54,335 (10.7% of 507,683 cases, the total PM dataset), compared to 25,453 cases (3.9%) retrieved in the PSUR #2. The increase in the number of cases reported during the current PSUR is attributed to the increase in cases reported from Austria (9068 cases in the PSUR #2 vs 31,769 cases in the current PSUR #3) due to active solicitation of LOE cases from the Austrian BoH.

- MC cases (40,416); NMC cases (13,919).

- Country of Incidence (≥2%): Austria (31,769), US (4874), UK (2725), Germany (2386), France (1934), Netherlands (1495); the remaining 9152 cases were distributed among 77 countries.

- Subjects’ gender: female (29,370), male (22,867) and unknown (2098).

- Subjects’ age in years: (n = 51,267), range: 6 months – 107 years, mean: 47.1, median: 46.0.

- Medical history (n = 8328): the most frequently (≥2%) reported relevant medical conditions included Hypertension (1429), Asthma (766), Drug hypersensitivity (617).

- COVID-19 Medical history: COVID-19 (1018), Suspected COVID-19 (361), Exposure to SARS-CoV-2 (49), Post-acute COVID-19 syndrome (48), COVID-19 pneumonia (10), SARS-CoV-2 test positive (9), Asymptomatic COVID-19, Coronavirus Infection (3 each), Coronavirus test positive, Occupational exposure to SARS-CoV-2 (2 each).

- Co-suspects (n = 3995 cases): the most frequently (≥10) reported relevant co-suspect vaccines/medications were COVID-19 vaccine (1861), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (798), COVID-19 vaccine mRNA (MRNA 1273) (768), Adalimumab (256), JNJ 78436735 (100), Ocrelizumab (35), Influenza vaccine (34), Upadacitinib (31), COVID-19 vaccine INACT (VERO) CZ02 (25), Risankizumab (21), Prednisone (18), Casirivimab/Imdevimab, Rituximab (13 each), Mycophenolate (10).

- Number of relevant events: 55,437.

- Relevant event seriousness: serious (52,185), non-serious (3254).

- Most frequently reported relevant PTs (≥2%): COVID-19 (47,981) Suspected COVID-19 (3002), and Ageusia (1094).

- Time to event onset (n = 46,269), range: <24 hours to 564 days, median: 117 days.

- Duration of relevant events (n = 1968 out of 4800 occurrences with outcome of resolved/resolved with sequelae), range: 24 hours to 373 days, median: 9 days.

- Relevant event outcome: fatal (506), resolved/resolving (7289), resolved with sequelae (296), not resolved (3281), unknown (44071).
In 493 cases (reporting 543 relevant events of which 506 relevant events reported a fatal outcome), the reported causes of death (>20 occurrences) were coded to the Pts COVID-19 (297), Vaccination failure (143), Drug ineffective (131), COVID-19 pneumonia (127), Death (46), Dyspnoea (21). Of note, in 39 cases limited information regarding the cause of death was provided (PT Death [38] and Sudden death [1]). Most (406 of 493 cases) of the fatal cases involved elderly subjects. When the medical history was provided (272 cases), the most frequently (>20 occurrences) relevant medical conditions included the Pts Hypertension (117), Atrial fibrillation (53), Chronic kidney disease, Dyslipidaemia (33 each), Type 2 diabetes mellitus (29), Myocardial ischaemia (25), Cardiac failure (24), COVID-19 (22), and Diabetes mellitus (21).

Analysis by age group

- Clinical trials: Paediatric (1), Adults (5), Elderly (1). Due to low volume of paediatric cases, a meaningful comparison of the same with the other age groups is not possible.
- Post-marketing: Paediatric (2158), Adults (39,726), Elderly (9566). No significant difference was observed in the reporting proportion of the most frequently reported COVID-19 AEs (>2%) between adult, elderly and paediatric population.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 3846 (0.8% of 507,683 cases, the total dataset).
- The reporting proportion of COVID-19 AEs with fatal outcome (5.6% [230 of 4093 events]) is higher in subjects with comorbid conditions, compared to the reporting proportion observed in the individuals without comorbidities (0.5% [276 of 51,344 cases] of fatal events).

Long COVID

Search criteria: PT Post-acute COVID-19 syndrome.

Clinical trial data

- No cases.

Post-authorization data

- Number of relevant cases: 200 (0.04% of 507,683 cases, the total PM dataset), compared to 72 cases (0.3% of 25,453 cases) retrieved in the PSUR #2.
- MC cases (62); NMC cases (138).
- Country of incidence: Germany (106), France (15), UK (14), Austria (13), Sweden (9), Australia, Finland (8 each), Netherlands (6), Italy (4), Ireland, US (3 each), Belgium, Hungary, New Zealand, Spain (2 each), Brazil, Greece and Luxembourg (1 each).
- Subjects' gender: female (151), male (46) and unknown (3).
- Subjects' age in years: (n = 174), range: 9 – 85 years, mean: 43.6, median: 45.0. Of these 174 subjects, there were 10 paediatric, 156 adults, and 8 elderly subjects.
- Medical history (n = 104): the most frequently (>2%) reported medical conditions included Asthma, Drug hypersensitivity (8 each), and Seasonal allergy (7).

Q/E analysis
- O/E analysis was performed for Ageusia/anosmia: all O/E ratios <1.

**MAH’s conclusion**

Loss of/Altered Taste and Smell was evaluated as signal during the reporting period and determined not to be a risk.

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

**Rapporteur assessment comment:**

During the interval period, there was an increased number of cases reporting COVID-19 AESIs (54,335 [10.7% of the total PM dataset] compared to 25,453 cases (3.9%) retrieved in the 2nd PSUR which is attributed to the increase in cases reported from Austria (9,068 cases in the 2nd PSUR versus 31,769 cases in the current 3rd PSUR) due to active solicitation of lack of efficacy cases from the Austrian board of health.

Please refer regarding the closed signal ‘Loss of/altered taste and smell’ to section 2.2.2. Evaluation of closed signals of this AR (PRAC concluded that the data provided did not support a causal relationship between Comirnaty exposure and the loss of/altered taste and smell).

COVID-19 AESIs were continuously monitored in the 13th and 14th SSRs (through 15 Apr 2022) for Comirnaty during the reporting period.

No new important safety concern could be identified for COVID-19 AESIs.

**Dermatological AESIs**

Search criteria: *Pts Chillblains; Erythema multiforme.*

**Clinical trial data**

- No cases, and no cases were retrieved in the PSUR #2.

**Post-authorization data**

- Number of cases: 284 (0.06% of 507,683 cases, the total PM dataset), compared to 339 (0.05%) cases retrieved in the PSUR #2.
- MC cases (158), NMC cases (126).
- Country of incidence: France (72), Germany (38), UK (25), Italy (24), Singapore (18), Japan, Poland (11 each), the Netherlands, US (9 each), Australia (8), Belgium (7), Canada, New Zealand, Spain (6 each); the remaining 34 cases were distributed among 18 countries.
- Subjects’ gender: female (182), male (93) and unknown (9).
- Subjects’ age in years: (n = 269), range: 7-89, mean: 46.4, median: 46.
- Medical history (n = 102): the most frequently (≥ 4 occurrences) reported relevant medical conditions included Hypertension (16), COVID-19 (12), Asthma (8), Drug hypersensitivity, Suspected COVID-19 (6 each), Diabetes mellitus (5), Cerebrovascular accident, Food allergy, Herpes simplex, Hypothyroidism, Type 2 diabetes mellitus (4 each).
- COVID-19 Medical history (n = 17): COVID-19 (12), Suspected COVID-19 (6), and Post-acute COVID-19 syndrome (1).
• Co-suspects (n = 4 cases): Acetylcysteine/benzalkoniumchloride/taurinoheptane sulfate, Albendazole, Dextromethorphan, Ibuprofen, Ketoprofen, Ocrelizumab, Prednisolone metasulfobenzoate sodium, Sulfasalazine (1 each).

• Number of events: 284.

• Relevant event seriousness: serious (206), non-serious (78).

• Reported relevant PTs: Erythema multiforme (181), Chillblains (103).

• Time to event onset (n = 72), range: <24 hours to 262 days, median: 4 days.

• Duration of relevant events (n = 14 out of 53 occurrences with outcome of resolved/resolved with sequelae), range: 0 days to 67 days, median: 20.5 days:

• Relevant event outcome: resolved/resolving (108), resolved with sequelae (8), not resolved (118), unknown (50). No fatal events were reported.

Analysis by age group

• Post-marketing: Paediatric (31), Adults (183), Elderly (60) and Unknown (10).
  o Due to low volume of paediatric cases a meaningful comparison of the same with the other age groups is not possible. No significant difference observed in the reporting proportion of events chillblains and erythema multiforme between adult and elderly population.

Analysis by presence of comorbidities

• Number of subjects reporting comorbidities: 53 (18.7 % of the cases reporting dermatological AESIs). A higher reporting proportion of dermatological AESIs was reported in subjects without significant comorbidities (81.3 %) when compared to subjects with significant comorbidities.

O/E analysis

• O/E analysis was performed for Chillblains and Erythema multiforme: all O/E ratios <1.

MAH's conclusion

No other safety signals have emerged based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:

No new important safety concern could be identified for dermatological AESIs. For future PSURs in the section 'Evaluation of AESIs', the dermatological AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Please refer regarding Pemphigus and Pemphigoid to the separate ongoing signal procedure (EPITT 19859).

Facial paralysis

Search Criteria: PTs Bell's palsy, Facial paralysis, Facial paresis, Oculo facial paralysis.

Clinical trial data

• Number of cases: 1 (BNT162b2 [1]) (0.1% of 668 cases, the total CT dataset) compared to no cases retrieved in the PSUR #2. Not related to BNT162b2.
Post-authorization data

- Number of cases: 2589 (0.5% of 507,683 cases, the total PM dataset), compared to 4515 cases (0.7%) retrieved in the PSUR #2.
- MC cases (1105), NMC cases (1484).
- Country/region of incidence: Germany (714), France (387), UK (229), Australia (184), Italy (112), Austria (97), Sweden (87), Hong Kong (70), Taiwan, Province of China (69), US (54); the remaining 586 cases were distributed among 40 countries.
- Subjects’ gender: female (1487), male (1060), and unknown (42).
- Subjects’ age in years: (n = 2473), range: 1.42 – 99, mean 47.3, median 47.0.
- Medical history (n = 934): the most frequently (>2%) reported relevant medical conditions were coded to the PTs Hypertension (185), Asthma (89), Seasonal allergy (85), Drug hypersensitivity (63), Hypersensitivity (57), Diabetes mellitus (55), Type 2 diabetes mellitus (40), Obesity (35), Food allergy (34), Hypothyroidism (32), Facial paralysis, Mite allergy (28 each), Allergy to animal (26), Bell’s palsy (25), Hypercholesterolaemia (24), Chronic obstructive pulmonary disease, Tobacco user (23 each), Migraine (20), and Coronary artery disease (19).
- COVID-19 Medical history (n = 133): reported medical conditions were coded to the PTs COVID-19 (100), Suspected COVID-19 (31), Post-acute COVID-19 syndrome (4), SARS CoV 2 test positive (2), Asymptomatic COVID-19, Coronavirus infection, and COVID-19 pneumonia (1 each).
- Co-suspects (n = 33): the relevant co-suspect vaccines/medications were diphtheria vaccine toxoid/ polio vaccine inact 3V (vero)/ tetanus vaccine toxoid and meningococcal group C tetanus toxoid conjugate vaccine (1 each).
- Number of relevant events: 2706.
- Relevant event seriousness: serious (2431) and non-serious (543).
- Reported relevant PTs: Facial paralysis (1428), Bell’s palsy (733), Facial paresis (543), and Oculofacial paralysis (2).
- Time to event onset (n = 2152 events), range: <24 hours to 389 days, median 7 days.
- Duration of relevant events (n = 286 out of 613 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 246 days, median: 6 days.
- Relevant event outcome: fatal (6), resolved/resolving (999), resolved with sequelae (111), not resolved at the time of reporting (1063), and unknown (534).
  - In 6 cases (reporting 6 relevant events with a fatal outcome), the causes of death (≥2 occurrences) were coded to the PTs Facial paralysis (3), Cerebrovascular accident and Death (2 each). Of note, in 2 cases limited information regarding the cause of death was provided (PT Death). All of the patients were >60 years of age (range 61 to 99 years). When the medical history was provided (4 cases), significant medial conditions reported Arthralgia, Cerebral infarction, Diverticulitis, Lung adenocarcinoma, Neoplasism malignant, Pemphigoid, and Pulmonary embolism (1 each).

Analysis by age group

- Paediatric (146), Adults (1914), Elderly (420), and Unknown (109).
There was no significant difference observed in the reporting proportion of facial paralysis events between age groups.

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 407 (15.7% of the CT and PM cases reporting facial paralysis).
  - The reporting proportion of cases reporting a facial paralysis event with a fatal outcome is higher in subjects with comorbid conditions (0.74%) when compared to the reporting proportion observed in the subjects without comorbidities (0.14%).

**O/E analysis**

- O/E analysis was performed for Bell’s palsy (PTs: Bell’s palsy, Facial paralysis, Facial paresis, Oculofacial paralysis): all O/E ratios <1.

**MAH’s conclusion**

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

**Rapporteur assessment comment:**

No new important safety concern could be identified for facial paralysis. For future PSURs in the section ‘Evaluation of AESIs’, the facial paralysis should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

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**Hepatic AESIs**

Search Criteria: *SMQ Liver related investigations, signs and symptoms (Narrow and Broad) OR PTs Autoimmune hepatitis, Liver Injury.*

**Clinical trial data**

- No cases compared to 2 cases (0.28%) retrieved in the PSUR #2.

**Post-authorization data**

- Number of relevant cases: 1091 (0.2% of 507,683 cases, the total PM dataset), compared to 1393 cases (0.2%) retrieved in the PSUR #2.
- MC cases (560), NMC cases (531).
- Country of incidence: Germany (276), Japan (157), France (152), UK (71), Australia (66), US (55), Italy (44), Austria (35), Spain (29), Taiwan, province of China (26), Netherlands (17), Sweden (15), Belgium, Finland (14 each), New Zealand (13), Greece (11), Canada (10), Denmark (9), Czech Republic (8), Norway, Poland (6 each), Croatia, Ireland, Romania (5 each), Switzerland (4), Brazil, Latvia, Philippines, Portugal, Slovakia, Slovenia (3 each), Estonia, Hungary, Lithuania, Malaysia, Mexico (2 each); the remaining 10 cases were distributed among 10 countries.
- Subjects’ gender: female (661), male (406) and unknown (24).
- Subjects’ age in years (n = 1017), range: 5 - 94, mean: 49.3, median: 51.0.
- Medical history (n = 518): the most frequently reported relevant medical conditions (≥5 occurrences) included Hypertension (80), Drug hypersensitivity (38), Hypothyroidism,
Seasonal allergy (33), Asthma (32), Food allergy (23), Autoimmune thyroiditis, Type 2 diabetes mellitus (21 each), Allergy to animal (20), Allergy to metals, Dyslipidaemia (19 each), Mite allergy (17), Atrial fibrillation (16), Diabetes mellitus, Hepatic steatosis, Tobacco user (15 each), Obesity (14), Hypercholesterolaemia, Hypersensitivity (13 each), Gastro-oesophageal reflux disease, Rheumatoid arthritis (12 each), Breast cancer, Mycotic allergy, Non-tobacco user, Tonsilloctomy (11 each), Autoimmune hepatitis, Depression, Ovarian cystectomy (10 each), Interchange of vaccine products, Osteoporosis, Salivary gland operation (9 each), Cardiac failure, Colecystectomy, Liver disorder, Migraine (8 each), Abstains from alcohol, Alcohol use, Allergy to plants, Anxiety, Arrhythmia, Epstein-Barr virus infection, Headache, Hysterectomy, Pyrexia, Type 1 diabetes mellitus (7 each), Coeliac disease, Disease risk factor, Hepatic cirrhosis, Hyperuricaemia, Thyroid disorder, Weight decreased (6 each), Colon cancer, Dermatitis contact, Diverticulum intestinal, Epilepsy, Hepatic function abnormal, Hepatitis, Hyperlipidaemia, Immunodeficiency, Insomnia, Nephrolithiasis, Neuropathy peripheral, Pericarditis, Primary biliary cholangitis, Sinus operation, Sjogren’s syndrome, Sleep apnoea syndrome, and Thyroid cancer (5 each).

- COVID-19 Medical history (n = 46): the medical conditions reported included COVID-19 (34), Post-acute COVID-19 syndrome, Suspected COVID-19, (5 each), Asymptomatic COVID-19, and COVID-19 pneumonia (1 each).

- Co-suspects (n = 58): the relevant co-suspect medications reported were adalimumab (10), upadacitinib (3), atorvastatin, hepatitis A vaccine, methotrexate, paracetamol (2 each), amiodipine, amoxicillin, cabozantinib, cefuroxime, ceftriaxone, clopidogrel, clozapine, colchicine, dospirenone ethinylestradiol, ebastine, ethinylestradiol gestodene, exemestane, fingolimod, ibuprofen, lipilumab, lanreotide, nitrofurantoin, nivolumab, paclitaxel, ribociclib, rosuvastatin, sorafenib, spironolactone, teriflunomide, torasemide, and valsartan (1 each).

- Number of relevant events: 1422.

- Relevant event seriousness: serious (676) and non-serious (746).

- Most frequently reported relevant PTs (≥50 occurrences): Hepatic enzyme increased (131), Alanine aminotransferase increased (126), Hepatic function abnormal (124), Liver function test abnormal (119), Aspartate aminotransferase increased (110), Autoimmune hepatitis (99), Hepatic pain (98), Gamma-glutamyltransferase increased (86), Liver function test increased (79), Transaminases increased (72), Ascites (60).

- Time to event onset (n = 876 events), range: <24 hours to 177 days, median: 7 days.

- Duration of relevant events (n = 120 out of 1425 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 210 days, median: 23 days.

- Relevant event outcome: fatal (23), resolved/resolving (426), resolved with sequelae (46), not resolved at the time of reporting (343), and unknown (586).
  - In 22 cases with fatal outcome (reporting 23 relevant events with fatal outcome), the reported causes of death were coded to Ascites (5), Congestive hepatopathy, Hepatic function abnormal, Hepatic pain, Hypertransaminasaemia (2 each), Alanine aminotransferase increased, Autoimmune hepatitis, Blood bilirubin increased, Hepatic enzyme increased, Hepatic mass, Hepatomegaly, Hepatosplenomegaly, Hypoalbuminaemia, Liver function test abnormal, and Liver injury (1 each). Of note, in 6 cases limited information regarding the cause of death was provided (Alanine aminotransferase increased, Ascites, Hepatic mass, Hepatic pain,
Hypertransaminasaemia, Liver injury (1 each). Most (13 of 22 cases) of the fatal cases involved subjects who were ≥60 years of age.

When the medical history was provided (13 cases), the relevant medical conditions included Hepatic steatosis, Type 2 diabetes mellitus (3 each), Diabetes mellitus (2), Autoimmune hepatitis, and Hepatic function abnormal (1 each).

**Analysis by age group**

- Post-marketing: Paediatric (66), Adults (712), Elderly (243) and No data (70).
  - Among the frequently (≥2%) reported relevant hepatic events, Hepatic pain was reported significantly higher in the adult population when compared to elderly population (25.5% in adult vs 6.3% in elderly). Upon further review, the majority of the events of hepatic pain were assessed as non-serious in the adult population (63 of 84 events).

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 265 (24.3% of the CT and PM cases reporting hepatic AESIs).
  - The reporting proportion of hepatic AESIs with fatal outcome (2.1%) is slightly higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subjects without comorbidities (1.4%).

**O/E analysis**

- O/E analysis was performed for Acute liver injury/Liver injury and Autoimmune hepatitis. All O/E ratios were <1.

**MAH’s conclusion**

A cumulative review of Autoimmune hepatitis has been performed. No other safety signals have emerged based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

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**Rapporteur assessment comment:**

Please refer regarding autoimmune hepatitis to section 2.2 Signal evaluation of this AR (a causal association between Comirnaty and autoimmune hepatitis cannot be concluded based on the available information).

No new important safety concern could be identified for hepatic AESIs. For future PSURs in the section ‘Evaluation of AESIs’, the hepatic AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

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**Immune-mediated/autoimmune AESIs**

Search criteria: *SMQ* Immune-mediated/autoimmune disorders (Narrow and Broad) OR HLG (All Path) Autoimmune disorders OR PTs Cytokine storm; Hypersensitivity; Thyroiditis subacute; Thrombocytopenia.

**Clinical Trial Data**

- Number of cases: 19 (BNT162b2 [17] and blinded therapy [2] (2.8% of 668 cases, the total CT dataset) compared to 20 cases (2.8%) retrieved in the PSUR #2. All SAEs were assessed as not related to BNT162b2 or blinded therapy.
Post-authorization data

- Number of cases: 11,726 (2.3% of 507,683 cases of the total PM dataset), compared to 21,994 cases (3.3%) retrieved in the PSUR #2.

- MC cases (4822), NMC cases (6904).

- Country of incidence: Germany (3094), France (1474), UK (1038), US (718), Italy (582), Japan (490), Australia (461), Netherlands (370), Austria (362), Sweden (250), Belgium (237), Norway (230), Greece (229), Finland (216), Poland, Spain (184 each), Taiwan, Province of China (156), Canada (145), New Zealand (125); the remaining 1181 cases were distributed among 64 countries.

- Subjects’ gender: female (7678), male (3661), and unknown (387).

- Subjects’ age in years (n = 10,827), range: 5 – 98, mean: 47.5, median: 47.0.

- Medical history (n = 4887): the most frequently (≥150 occurrences) reported relevant medical conditions were coded to the PTs Seasonal allergy (400), Asthma (378), Drug hypersensitivity (317), Hypersensitivity (306), Psoriasis (269), Hypothyroidism (252), Autoimmune thyroiditis (237), Food allergy (235), Diabetes mellitus (186), and Colitis ulcerative (158).

- COVID-19 Medical history (n = 507): the reported medical conditions were coded to the PTs COVID-19 (382), Suspected COVID-19 (119), COVID-19 pneumonia (11), Post-acute COVID-19 syndrome (5), Coronavirus infection (4), SARS-CoV-2 test positive (3), and Asymptomatic COVID-19 (2).

- Co-suspects (n = 460): the most frequently (≥10 occurrences) reported relevant co suspects were adalimumab (168), COVID-19 vaccine MRNA (MRNA 1273) (36), Influenza vaccine (26), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (19), influenza vaccine inact split 4V, and Risankizumab (13 each).

- Number of relevant events: 12,795.

- Relevant event seriousness: serious (8445) and non-serious (4356).

- Most frequently reported relevant PTs (≥2%): Hypersensitivity (2393), Psoriasis (660), Thrombocytopenia (487), Polymyalgia rheumatica (431), Dermatitis (305), Rheumatic disorder (286), and Alopecia areata (281).

- Time to event onset (n = 7591), range: □24 hours to 499 days, median: 6 days.

- Duration of relevant events (n = 969 out of 2334 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 419 days, median 14 days.

- Relevant event outcome: fatal (133), resolved/resolving (3786), resolved with sequelae (664), not resolved at the time of reporting (4934), and unknown (3304).

  - In 112 cases (reporting 133 relevant events with a fatal outcome), the reported causes of death (≥5 occurrences) were coded to the PTs Thrombocytopenia (19), Death, Interstitial lung disease (13 each), Haemophagocytic lymphohistiocytosis, Immune thrombocytopenia (8 each), Cerebral haemorrhage, Encephalitis, Multiple organ dysfunction syndrome, Renal failure (7 each), Pneumonia, Respiratory failure (6 each), and Pulmonary embolism (5). Of note, in 13 cases limited information regarding the cause of death was provided (PT Death). Most (78 of 104 cases that provided age) of the fatal cases involved elderly subjects. When the medical history was provided (67 cases), significant medical conditions reported in more than 3 cases included
Hypertension (23), Atrial fibrillation (9), Osteoporosis (7), Dyslipidaemia (6), Diabetes mellitus, Hyperlipidemia, Type 2 diabetes mellitus (5 each), Hypothyroidism, Myocardial Infarction, Radiotherapy, and Thrombocytopenia (4 each).

Analysis by age group

- Clinical trial: Paediatric (5), Adults (11), and Elderly (3). A meaningful comparison between the different age groups is not possible due to the low number of cases.
- Post-marketing: Paediatric (591), Adults (8319), Elderly (2125) and Unknown (691).
  - Among the frequently (>2%) reported immune mediated/autoimmune AESIs, a higher reporting proportion of events coded to the PT Polymyalgia rheumatica were observed in the elderly population when compared to paediatric and adult populations (none in paediatrics vs 1.7% in adults vs 12.9% in elderly).
  - A higher reporting proportion of events coded to the PTs Hypersensitivity and Alopecia areata were observed in the paediatric and adult populations when compared to the elderly population (Hypersensitivity [24.2% in paediatrics vs 20.8% in adults vs 11.8% in elderly], Alopecia areata [3.6% in paediatrics vs 2.7% in adults vs 0.9% in elderly]).
  - A higher reporting proportion of events coded to the PTs Psoriasis and Rheumatic disorder were observed in the adult and elderly populations when compared to the paediatric population (Psoriasis [2.2% in paediatrics vs 5.8% in adults vs 5.9% in elderly], Rheumatic disorder [0.5% in paediatrics vs 2.3% in adults vs 3.4% in elderly]).
  - A higher reporting proportion of events coded to the PT Thrombocytopenia were observed in the paediatric and elderly populations when compared to the adult population (8.8% in paediatrics vs 3.3% in adults vs 6.3% in elderly).

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 3199 (27.2% of the CT and PM cases reporting immune mediated/autoimmune AESIs).
  - The reporting proportion of immune mediated/autoimmune AESIs with a fatal outcome (2.6%) is higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subjects without comorbidities (0.6% of events with fatal outcome).

O/E analysis

- O/E analysis was performed for Acute disseminated encephalomyelitis (ADEM), ADEM and encephalitis, Autoimmune thyroiditis, Myasthenia gravis, Polymyalgia rheumatica, and Type 1 diabetes mellitus. All O/E ratios were <1.

MAH's conclusion

Polymyalgia rheumatica, Uveitis and Subacute Thyroiditis (SAT) were evaluated as signals in the reporting period and determined not to be risks. No new safety signals have emerged based on a review of the remaining events and on O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:

Please refer regarding polymyalgia rheumatica, uveitis and subacute thyroiditis to section 2.2 Signal evaluation of this AR.
Immune-mediated/autoimmune AESIs were continuously monitored in the 13th and 14th SSRs (through 15 Apr 2022) for Comirnaty during the reporting period.

No new important safety concern could be identified for immune-mediated/autoimmune AESIs.

Multisystem Inflammatory Syndrome in Children / Adults

Search criteria: Pts Cytokine release syndrome; Distributive shock; Multiple organ dysfunction syndrome; Multisystem inflammatory syndrome; Multisystem inflammatory syndrome in adults; Multisystem inflammatory syndrome in children; Systemic inflammatory response syndrome.

Rapporteur assessment comment:
The amended the search strategy since the initial signal evaluation was considered acceptable by PRAC (13th SSR; EMEA/H/C/005735/MEA/002.12), because no additional MIS-C/A cases were identified using the extended search strategy.

Clinical Trial Data

- No serious cases from the CT dataset were reported. For comparison, 2 cases (0.3%) were retrieved in the PSUR #2.

Post-authorization data

- Number of relevant cases: 207 (0.04% of 507,683 cases in the total PM dataset), compared to 438 (0.07%) retrieved in PSUR #2.
- MC cases (170), NMC cases (37).
- Country of incidence (≥5 occurrences): France (55), Germany (27), UK (18), Australia (15), US (14), Japan (12), Norway (6), Spain (5); the remaining 55 cases were distributed among 30 countries.
- Subjects’ gender: female (92), male (109), unknown (6).
- Subjects’ age in years (n = 196), range: 3 – 95, mean: 46.6, median: 50.
- Medical history (n = 132): the most frequently (≥5 occurrences) reported medical conditions included the PTs Hypertension (40), Obesity (11), Diabetes mellitus (9), Ex-tobacco user, Hypothyroidism (8 each), Atrial fibrillation, Tobacco user, Type 2 diabetes mellitus (7 each), Alcohol use, Osteoporosis, Pyrexia, Sleep apnoea syndrome (6 each), Asthma, Non-tobacco user, Prostate cancer, and Rheumatoid arthritis (5 each).
- Co-suspects (n = 16 cases): COVID-19 vaccine mRNA (mRNA 1273) (2), carboplatin, cefotaxime, ciclosporin, colchicine, COVID-19 vaccine, etrombopag, enoxaparin, everolimus, mesalazine, methotrexate, pembrolizumab, pemetrexed, rituximab, treprostinil (1 each).
- Number of relevant events: 210.
- Relevant event seriousness: serious (210).
- Relevant PTs: Multiple organ dysfunction syndrome (82), Multisystem inflammatory syndrome (43), Multisystem inflammatory syndrome in children (38), Systemic inflammatory response
syndrome (32), Multisystem inflammatory syndrome in adults (10), Cytokine release syndrome (5).

- Time to event onset (n = 115), range: <24 hours to 234 days, median: 15 days.
- Duration of relevant events (n = 12 out of 39 occurrences with outcome of resolved or resolved with sequelae), range: 3 days to 57 days, median: 16 days.
- Relevant event outcome: fatal (57), resolved/resolving (61), resolved with sequelae (3), not resolved (20), unknown (72).
  - In 56 cases (reporting 57 relevant events with fatal outcome), the reported causes of death (≥5 occurrences) were coded to Multiple organ dysfunction syndrome (55), Septic shock (10), Renal failure (9), Immunisation, Sepsis (8 each), Pneumonia (7), Acute respiratory distress syndrome (6), Acute kidney injury, Cardiac arrest, COVID-19, COVID-19 pneumonia, Drug ineffective, Hepatic failure, Respiratory failure, and Vaccination failure (5 each).

Most (35 of 56 cases) of the fatal cases involved elderly subjects. When the medical history was provided (43 cases), the most frequently (≥3 occurrences) medical conditions included Hypertension (19), Diabetes mellitus, Obesity (6 each), COVID-19 (5), Atrial fibrillation, Ex-tobacco user, Hypothyroidism, Osteoarthritis, Renal transplant, and Tobacco user (3 each).

Analysis by age group

- Post-marketing: Paediatric (46 [16 Child, 30 Adolescent]), Adult (84), Elderly (69), Unknown (8).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 73 (35.3% of the 207 cases reporting Multisystem Inflammatory Syndrome AESIs).
  - Of the PM cases that reported medical histories, the percentage of cases with a fatal Multisystem Inflammatory Syndrome AESI is higher in subjects with comorbid conditions (60.5%) when compared to the percentage of cases with a fatal Multisystem Inflammatory Syndrome AESI in subjects without comorbidities (39.5%).
  - Upon review of the relevant events in PM cases that recorded medical histories, no Multisystem Inflammatory Syndrome AESIs had a significant proportional reporting ratio of >3:1 in subjects with comorbidities compared to subjects without comorbidities.

Q/E analysis

- Q/E analysis was performed for Multisystem inflammatory syndrome:
Table 10. Observed to Expected (O/E) Analysis of Multisystem Inflammatory Syndrome in European Economic Area Countries and in the United States, Cumulative Period

<table>
<thead>
<tr>
<th>Stratification</th>
<th>Blgld. rate*10⁵</th>
<th>21-Day Risk Window</th>
<th>42-Day Risk Window</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Obs Cases</td>
<td>Exp Cases</td>
<td>O/E</td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;11 years</td>
<td>2.77</td>
<td>12</td>
<td>1,520</td>
</tr>
<tr>
<td>12-17 years</td>
<td>2.77</td>
<td>31</td>
<td>3,324,066</td>
</tr>
<tr>
<td>18-24 years</td>
<td>1.50</td>
<td>7</td>
<td>2,028,811</td>
</tr>
<tr>
<td>21-24 years</td>
<td>0.23</td>
<td>7</td>
<td>2,705,081</td>
</tr>
<tr>
<td>25-49 years</td>
<td>0.58</td>
<td>54</td>
<td>19,546,121</td>
</tr>
<tr>
<td>50-59 years</td>
<td>1.47</td>
<td>29</td>
<td>9,025,205</td>
</tr>
<tr>
<td>60-69 years</td>
<td>3.38</td>
<td>36</td>
<td>7,901,385</td>
</tr>
<tr>
<td>70+ years</td>
<td>7.42</td>
<td>107</td>
<td>10,962,290</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Males</td>
<td>2.36</td>
<td>147</td>
<td>26,983,060</td>
</tr>
<tr>
<td>Females</td>
<td>2.36</td>
<td>156</td>
<td>19,427,120</td>
</tr>
</tbody>
</table>

a. Background rate per 100,000 person years (PY). Background rates from ACCESS include Kawasaki disease codes

- As in the most recent SBSR #3, the 21-24 years age group using the 21-day risk window meets the signal criteria with an O/E ratio >1, however, the result if not statistically significant as the 95% CI includes 1.

MIS is a condition known to be associated with COVID-19. The MAH's COVID-19 vaccine requires the first two doses to be spaced 21 days apart before achieving optimal effectiveness. It is possible that COVID-19 may occur within the 21-day window between doses. Additionally, general awareness of MIS is currently increased because of COVID-19, leading to possible stimulated reporting. The ACCESS background rates used for the MIS analysis are primarily based on codes for Kawasaki disease and may mis-characterize background rates of true MIS cases in the presence of COVID-19 infections.

MAH's conclusion

Cases of potential MIS in adults (MIS-A) and children (MIS-C) reported during this interval period are assessed in Appendix 6A.4 of the PSUR.

During the reporting period, an article including important safety information on MIS-C was reviewed.

No new safety signals have emerged based on a review of these cases, literature or of the O/E analysis. The MAH will continue to monitor MIS

Rapporteur assessment comment:

Please refer regarding new cases of MIS-C/ -A to section 2.2 Signal evaluation of this AR.

During the interval period, the MAH reported post-marketing 207 relevant MIS-C/ -A cases. However, in Appendix 6A.4 of the PSUR the MAH reported 199 MIS-C/ -A cases during the interval period. The MAH is requested to explain this discrepancy in the numbers of retrieved MIS-C/ -A cases. Request for supplementary information

The study of Ouldali et al. was discussed in the 13th (2nd bi-monthly) SSR (procedure EMEA/H/C/005735/MEA/002.12). No new safety concern was identified.

In the O/E analyses for multisystem inflammatory syndrome, the O/E ratio was elevated (O/E ratio 1.1, 95% CI 0.5;2.3) for the 21-day risk window in the 21-24 years age group.

Provided MAH's response regarding the Requests for supplementary information for MIS-C/ -A, no new important safety concern could be identified for multisystem inflammatory syndrome in children / adults.
Musculoskeletal AESIs

Search Criteria: PTs Arthralgia; Arthritis; Chronic fatigue syndrome; Juvenile idiopathic arthritis; Polyarthritis; Post viral fatigue syndrome; Rhabdomyolysis; Rheumatoid arthritis.

Clinical Trial Data

- Number of cases: 6 (BNT162b2) (0.9% of 668 cases, the total CT dataset) compared to 4 cases (0.6%) retrieved in the PSUR #2. Not related to BNT162b2.

Post-authorization data

- Number of relevant cases: 31,012 (6.1% of 507,683 cases, the total PM dataset), compared to 58,250 cases (8.9%) retrieved in the PSUR #2.
- Relevant PTs: Arthralgia (29,429), Arthritis (996), Rheumatoid arthritis (660), Chronic fatigue syndrome (219), Polyarthritis (145), Post viral fatigue syndrome (92), Rhabdomyolysis (78), Juvenile Idiopathic arthritis (14).

Analysis by age group

- Clinical trial: Adult (1) and Elderly (5).
- Post-marketing: Paediatric (664), Adult (25,307), Elderly (3469), Unknown (1572).

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 3193 (10.3% of the cases reporting musculoskeletal AESIs).

O/E analysis

- O/E analysis was performed for Chronic fatigue syndrome/ME/PVFS, Rhabdomyolysis, Rheumatoid arthritis, polyarthritis, juvenile idiopathic arthritis. All O/E ratios were <1.

MAH's conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. The majority of the events reported in this category are arthralgia which is considered to be an adverse reaction for the vaccine and is labelled as such. Arthralgia will be removed from the search strategy in the next PSUR. Safety surveillance will continue.

Rapporteur assessment comment:

MAH's proposal to remove arthralgia from the search strategy for musculoskeletal AESIs in the next PSUR is endorsed because the majority of the musculoskeletal events reported are arthralgia which is labelled in the ADR table in section 4.8 of the Comirnaty SmPC (frequency very common).

Musculoskeletal AESIs were continuously monitored in the 13th and 14th Comirnaty SSRs (through 15 Apr 2022) during the reporting period.

No new important safety concern could be identified for musculoskeletal AESIs. For future PSURs in the section 'Evaluation of AESIs', the musculoskeletal AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Myocarditis and Pericarditis AESIs

Rapporteur assessment comment:
Neurological AESIs (including demyelination)

Search Criteria: SMQ Generalised convulsive seizures following immunisation (Narrow) OR SMQ Demyelination (Narrow and Broad) OR PTs Ataxia; Cataplexy; Fibromyalgia; Intracranial pressure increased; Meningitis; Meningitis aseptic; Narcolepsy; Neuropathy peripheral; Polyneuropathy.

Clinical Trial Data

- Number of cases: 15 cases (BNT162b2 [11], blinded therapy [4]; 2.2% of 668 cases in the total CT dataset) compared to 7 cases (0.97%) retrieved in the PSUR #2. None of these SAEs were assessed as related to BNT162b2/blinded therapy.

Post-authorization data

- Number of relevant cases: 5111 (1.0% of 507,683 cases in the total PM dataset), compared to 7197 cases (1.1%) retrieved in the PSUR #2.

Analysis by age group

- Clinical trial: Paediatric (Infant [2], Child [4]), Adult (6), Elderly (3).
- Post-marketing: Paediatric (523 [162 Child, 361 Adolescent]), Adult (3574), Elderly (787), Unknown (227).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 1201 (23.5% of the 5111 cases reporting Neurological AESIs).

O/E analysis

- O/E analysis was performed for Generalized convulsive, Fibromyalgia, Guillain-Barré syndrome, Meningitis, Narcolepsy, Multiple sclerosis and Polyneuropathy. All O/E ratios were <1.

MAH’s conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:

Neurological AESIs were continuously monitored in the 13th and 14th Comirnaty SSRs (through 15 Apr 2022) during the reporting period.

No new important safety concern could be identified for neurological AESIs.

Other AESIs

Search Criteria: HLT (All Path) Herpes viral infections OR PTs Adverse event following immunisation; Appendectomy; Appendicitis; Appendiceal abscess; Appendicitis perforated; Complicated appendicitis; Deafness; Deafness bilateral; Deafness neurosensory; Deafness permanent; Deafness transitory; Deafness unilateral; Hypoacusis; Inflammation; Manufacturing laboratory analytical testing issue; Manufacturing materials issue; Manufacturing production issue; MERS-CoV test; MERS-CoV test negative; MERS-CoV test positive; Occupational exposure to communicable disease; Patient isolation;
**Product availability issue; Product distribution issue; Product supply issue; Pyrexia; Quarantine; SARS-CoV-1 test; SARS-CoV-1 test negative; SARS-CoV-1 test positive; Sudden hearing loss.**

**Clinical Trial Data**

- Number of cases: 26 (BNT162b2 [22], blinded therapy [3] and placebo [1]) (3.9% of 668 cases, the total CT dataset) compared to 2 cases (0.28%) retrieved in the PSUR #2. None of the SAEs were assessed as related to BNT162b2 or blinded therapy or placebo.

**Post-authorization data**

- Number of relevant cases: 68,548 (13.5% of 507,683 cases, the total PM dataset), compared to 118,843 cases (18.1%) retrieved in the PSUR #2.
- Most frequently reported relevant PTs (≥ 50 occurrences): Pyrexia (57,474), Herpes zoster (6216), Inflammation (1585), Oral herpes (794), Hypoacusis (744), Deafness (488), Sudden hearing loss (370), Herpes virus infection (297), Appendicitis (254), Deafness unilateral, Ophthalmic herpes zoster (222 each), Herpes simplex (206), Adverse event following immunisation (188), Genital herpes (163), Herpes ophthalmic (70), Deafness neurosensory (63), Herpes zoster oticus (61), Herpes zoster reactivation, and Varicella (59 each).

**Analysis by age group**

- Clinical trial: Adult (14), Paediatric (10), Elderly (2).
- Post-marketing: Paediatric (5092), Adult (53,918), Elderly (6771), and Unknown (2767).

**Analysis by presence of comorbidities**

- Number of subjects reporting comorbidities: 5215 (7.6 % of the cases reporting other AESI).

**O/E analysis**

- O/E analysis was performed on Appendicitis, Herpes zoster and Sudden hearing loss. All O/E ratios were <1.

**MAH’s conclusion**

Hearing loss and Appendicitis was evaluated as a signal during the reporting period. No other safety signals than those mentioned have emerged based on the review of these cases, or from the O/E analysis. No risks have been identified following the evaluations of appendicitis and hearing loss. Safety surveillance will continue.

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**Rapporteur assessment comment:**

Regarding hearing loss please refer to the assessment in section 2.2.1.1 Post-approval regulatory requests of this AR above.

Regarding Appendicitis please refer to the assessment in section 2.2.2.1 Signals determined not to be risks of this AR above.

Other AESIs were continuously monitored in the 13th and 14th Comirnaty SSRs (through 15 Apr 2022) during the reporting period.

No new important safety concern could be identified for other AESIs. For future PSURs in the section 'Evaluation of AESIs', the other AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**
Pregnancy related AESIs

Rapporteur assessment comment:

Please refer to the assessment in section ‘Use in pregnant/lactating women’ in this AR below.

Glomerulonephritis and Nephrotic Syndrome AESIs

Search Criteria: HLT Glomerulonephritis and nephrotic syndrome.

The PTs Acute kidney injury and Renal failure have been removed from the search criteria and replaced with a more focused search of glomerulonephritis and nephrotic syndrome based on the evolving pharmacovigilance and medical literature. An evaluation of IgA nephropathy, as requested by EMA in the PSUR 2 assessment report is ongoing and will be provided to EMA under separate cover from the PSUR.

Clinical Trial Data

- No serious cases from the CT dataset were reported. No comparison with PSUR #2 is possible due to the change in the search criteria.

Post-authorization data

- Number of cases: 276 (0.05% of 507,683 cases, the total PM dataset). No comparison with PSUR #2 is possible.
- MC cases (172), NMC cases (104).
- Country of incidence: Germany (74), Japan (50), France (29), Australia (13), Italy, UK (11 each); the remaining 88 cases were distributed among 28 countries.
- Subjects’ gender: female (150), male (124) and unknown (2).
- Subjects’ age in years (n = 270), range: 5 – 88, mean: 44.2, median: 43.0.
- Medical history (n = 148): the most frequently (≥5 occurrences) reported relevant medical conditions included Hypertension (25), Nephrotic syndrome (12), Hypercholesterolaemia (8), Dyslipidaemia, Glomerulonephritis, Haematuria, IgA nephropathy (7 each), Proteinuria (6).
- Co-suspects (n = 3 cases): the reported relevant co-suspect medications included Hepatitis A vaccine, Influenza vaccine, and Tocilizumab (1 each).
- Number of relevant events: 323.
- Relevant event seriousness: serious (318), non-serious (5).
- Most frequently reported relevant PTs: Nephrotic syndrome (99), IgA nephropathy (47), Glomerulonephritis (46), Glomerulonephritis minimal lesion (25), Granulomatosis with polyangiitis (22), Microscopic polyangiitis (14), Glomerulonephritis membranous (12), Focal segmental glomerulosclerosis, and Glomerulonephritis rapidly progressive (10 each).
- Time to event onset (n = 172), range: <24 hours to 172 days, median: 12 days.
- Duration of relevant events (n = 12 out of 323 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 137 days, median 48 days.
- Relevant event outcome: fatal (2), resolved/resolving (95), resolved with sequelae (23), not resolved (111), unknown (92).

In 2 cases (reporting 2 relevant events with fatal outcome), the reported causes of death were coded to Glomerulonephritis and Granulomatosis with polyangiitis (1 each). Both fatal cases involved elderly subjects. Medical history was provided in both cases and included Autoimmune hypothyroidism, Hypertension and Obesity (1 each).

Analysis by age group

- Post-marketing: Paediatric (33), Adult (177), Elderly (62) and Unknown (4).
- Among the frequently (>2%) reported relevant events Glomerulonephritis and Nephrotic Syndrome AESIs, the PT Renal failure was reported significantly higher in elderly population when compared to adult population (2.8% in adults vs 8.4% in elderly). A higher reporting proportion of events coded to the PTs Haematuria and IgA nephropathy were observed in the adult population when compared to the elderly population (Haematuria [10.4% in adults vs 2.1% in elderly] and IgA nephropathy [11.1% in adults vs 1.1% in elderly]). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 14 (5.1% of the cases reporting Glomerulonephritis and nephrotic syndrome AESIs).
- The reporting proportion of Glomerulonephritis and nephrotic syndrome AESIs with fatal outcome is 0.4% in subjects without comorbid conditions. There were no fatal outcomes in the subjects with comorbidities.

O/E analysis

O/E analysis was performed for Glomerulonephritis/nephrotic syndrome. All O/E ratios were <1.

MAH’s conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. The ongoing evaluation of IgA nephropathy will be submitted separately from the PSUR. Safety surveillance will continue.

Rapporteur assessment comment:

Regarding the assessment of the cumulative review IgA nephropathy please refer to section 2.2 ‘Signal evaluation’ of this AR above.

During the reporting interval, retrieved were 46 glomerulonephritis cases which is considered an increase compared with the 27 glomerulonephritis cases from the previous reporting period. Therefore, the MAH is requested to provide detailed information concerning the 46 cases with glomerulonephritis in the interval period and perform an WHO-UMC causality assessment per case, and to provide an updated review with DLP 14 Nov 2022 of cases reporting glomerulonephritis from their safety database including a WHO-UMC causality assessment per case regarding Comirnaty exposure. Request for supplementary information

Respiratory AESIs
Search Criteria: HLTs (All Path) Lower respiratory tract infections NEC; Respiratory failures (excl. neonatal); Viral lower respiratory tract infections OR PTs Acute respiratory distress syndrome; Endotracheal intubation; Hypoxia; Respiratory disorder.

Clinical trial data
- Number of cases: 33 (Blinded therapy [10], BNT162b2 [23]) (4.9 % of 668 cases, the total CT dataset) compared to 38 cases (5.3%) retrieved in the PSUR #2. All were assessed as not related to BNT162B2 or blinded therapy.

Post-authorisation data
- Number of cases: 2188 (0.4% of 507,683 cases, the total PM dataset), compared to 3356 cases (0.51%) retrieved in the PSUR #2.
- Most frequently reported relevant PTs (≥ 100 occurrences): Pneumonia (809), Respiratory disorder (325), Bronchitis (303), Respiratory failure (213), Lower respiratory tract infection (175), Cardiorespiratory arrest (140), and Hypoxia (133).

Analysis by age group
- Clinical trial: Paediatric (8), Adult (12) and Elderly (13).
- Post-marketing: Paediatric (83), Adult (1168), Elderly (836) and Unknown (101).

Analysis by presence of comorbidities
- Number of subjects reporting comorbidities: 113 (5.16 % of the cases reporting respiratory AESIs).

O/E analysis
O/E analysis was performed for Acute respiratory distress syndrome: all O/E ratios were below 1.

MAH’s conclusion
No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue. Respiratory events were originally included in the AESI list in order to capture potential cases of respiratory failure that may occur in cases of severe COVID-19. The search strategy will be amended to focus on acute respiratory distress syndrome and respiratory failure for the next PSUR.

Rapporteur assessment comment:
No new important safety concern could be identified for respiratory AESIs. For future PSURs in the section ‘Evaluation of AESIs’, the respiratory AESIs should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Stroke
Search criteria: HLT Central nervous system haemorrhages and cerebrovascular accidents (All path); Cerebrovascular venous and sinus thrombosis (Primary Path).

Clinical trial data
- Number of cases: 19 cases (BNT162b2 [18], blinded therapy [1]; 2.8% of 668 cases in the total CT dataset) compared to 19 cases (2.6%) retrieved in the PSUR #2. None of the SAEs were assessed as related to BNT162b2/blinded therapy.
Post-authorisation data

- Number of cases: 3091 (0.6% of 507,683 cases in the total PM dataset), compared to 4834 cases (0.7%) retrieved in the PSUR #2.
- Most frequently (>25 occurrences) reported relevant PTs: Cerebrovascular accident (1363), Cerebral infarction (416), Ischaemic stroke (367), Cerebral haemorrhage (306), Cerebral venous sinus thrombosis (166), Cerebral thrombosis (93), Cerebral ischaemia (76), Subarachnoid haemorrhage (72), Cerebral venous thrombosis (68), Cerebellar infarction (42), Brain stem infarction, Haemorrhage intracranial (35 each), Ischaemic cerebral infarction (33), Embolic stroke (31), Haemorrhagic stroke (29), Thalamic infarction (26).

Analysis by age group

- Clinical trial: Adult (6), Elderly (13).
- Post-marketing: Paediatric [33 [9 Child, 24 Adolescent]], Adult (1575), Elderly (1352), Unknown (131).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 719 (23.3% of the 3091 cases reporting stroke-related events).

O/E analysis

- O/E analysis was performed for Cerebral venous sinus thrombosis, Ischemic stroke and Hemorrhagic stroke. For the CVST analysis, males and females 18-24 years, males and females 25-49 years, females 60-69 years, and overall dose 2 using the low background rate meet the signal criteria with an O/E ratio greater than 1 in either the 21-day or 42-day risk windows. However, the 95% CIs for some age groups included 1, indicating that the result is not statistically significant. For all other stratifications using the low background rate, the O/E ratio is less than 1. Using the mid-range background rate, all stratifications have an O/E ratio less than 1. The O/E were similar to the most recent SBSR #3.

MAH's conclusion

Cerebral venous sinus thrombosis (CVST) and Cerebrovascular Accident (CVA)/Stroke were evaluated as signals during the reporting period and were not determined to be risks causally associated with the vaccine. No additional safety signals other than those mentioned above have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:

Regarding cerebral venous sinus thrombosis (CVST) please refer to section 2.2. ‘Evaluation of closed signals’ of this AR above.

Stroke was continuously monitored in the 13th and 14th Comirnaty SSRs (through 15 Apr 2022) during the reporting period, and concerning cerebrovascular accident (CVA)/stroke/ cerebral venous sinus thrombosis (CVST) no new important safety information could be identified.

In the age-stratified analysis, some of the O/E ratios (incl. 95% CI) were >1 (not in paediatric persons), this was only seen when applying the low background rates. Using the mid-range background rate all O/E ratios were below 1 and similar to the O/E analyses results provided in the 14th Comirnaty SSR.

No new important safety concern could be identified for stroke.
Sudden Death

Rapporteur assessment comment:

Please refer to ‘Death’ in section ‘Evaluation of special situations’ of this AR below.

Thromboembolic AESIs

Search criteria: HLGT (All path) Embolism and thrombosis (excluding PTs reviewed as Stroke AESIs) OR PT Coagulopathy.

Clinical trial data

- Number of cases: 17 (BNT162b2 [16], blinded therapy [1]; 2.5% of 668 cases in the total CT dataset) compared to 15 cases (2.1%) retrieved in the PSUR #2. None of these SAEs were assessed as related to BNT162b2/blinded therapy.

Post-authorisation data

- Number of cases: 6102 (1.2% of 507,683 cases in the total PM dataset), compared to 6507 cases (1.0%) retrieved in the PSUR #2.

- Most frequently (≥50 occurrences) reported relevant PTs: Pulmonary embolism (2068), Thrombosis (1461), Deep vein thrombosis (1321), Thrombophlebitis (285), Venous thrombosis limb (276), Superficial vein thrombosis (258), Venous thrombosis (173), Coagulopathy (164), Retinal vein occlusion (127), Embolism (103), Pulmonary thrombosis (77), Ophthalmic vein thrombosis (74), Retinal vein thrombosis (54), Retinal artery occlusion (52), Portal vein thrombosis (50).

Analysis by age group

- Clinical trial: Adults (12), Elderly (5).

- Post-marketing: Paediatric (79 [7 Child, 72 Adolescent]), Adults (3833), Elderly (1966), Unknown (224). The reporting proportion of the PT Coagulopathy was significantly higher in the paediatric population (11.4%) when compared to the adult and elderly populations (2.7% and 2.1%, respectively).

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 1356 (22.2% of the 6102 cases reporting thromboembolic AESIs).

O/E analysis

- O/E analysis was performed for Arterial thromboembolism, Deep vein thrombosis, Disseminated Intravascular coagulation, Thrombotic thrombocytopenia syndrome and Venous thromboembolism. All O/E ratios were below 1.

MAH’s conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

Rapporteur assessment comment:
Thromboembolic AESIs were continuously monitored in the 13th and 14th Conmiitary SSRs (through 15 Apr 2022) during the reporting period.

No new important safety concern could be identified for thromboembolic AESIs.

**Vasculitic events**

Search criteria: HLT (All Path) Vasculitides OR PTs Microangiopathy; Peripheral ischaemia.

**Clinical trial data**

- During the reporting period no serious cases from the CT dataset were reported; no cases were retrieved in the PSUR #2.

**Post-authorisation data**

- Number of cases: 612 (0.12% of 507,683 cases, the total PM dataset), compared to 854 cases (0.13%) retrieved in the PSUR #2.

- Most frequently reported relevant PTs (≥20 occurrences): Vasculitis (267), Giant cell arteritis (102), Henoch-Schonlein purpura (66), Peripheral ischaemia (60).

**Analysis by age group**

- Post-marketing: Paediatric (62), Adults (317), Elderly (208) and Unknown (25).

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 224 (36.6 % of the PM cases reporting vasculitic events).

**O/E analysis**

- O/E analysis was performed for Behcet’s syndrome, Giant cell arteritis, Henoch-Schonlein purpura, Limb Ischaemia, and Vasculitis.

**MAH’s conclusion**

Vasculitis was evaluated as signal during the reporting period and was determined to not be a risk. No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

**Rapporteur assessment comment:**

An updated cumulative review of vasculitis (through 15 Apr 2022) was assessed in the 14th (3rd bi-monthly) SSR (reporting period 16 Feb 2022 – 15 Apr 2022; procedure EMEA/H/C/005735/MEA/002.13). The MAH had provided an updated review of vasculitis and included WHO-UMC causality assessments for the relevant vasculitis cases. Given that the O/E ratios are <1, only two probable cases were identified and no relevant literature was available, PRAC agreed that based on the data provided no safety concern was identified.

No new important safety concern could be identified for vasculitic events. For future PSURs in the section ‘Evaluation of AESIs’, the vasculitic events should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

**AESIs in subjects with Malnutrition; HIV infection**
As part of the approval letter for the emergency use of Tozinameran - COVID-19 mRNA vaccine (nucleoside modified) - COMIRNATY, the WHO requested the MAH to provide additional data on vaccine immunogenicity, effectiveness and safety on population groups represented in Low- and Middle-Income Countries (LMICs) including individuals with conditions such as malnutrition and populations with existing co-morbidities such as tuberculosis, human immunodeficiency virus (HIV) infection and other high prevalent infectious diseases.

Search criteria - PT Decode (History): Acute HIV infection; Asymptomatic HIV infection; Congenital HIV infection; HIV infection; HIV infection CDC Group I; HIV infection CDC Group II; HIV infection CDC Group III; HIV infection CDC Group IV subgroup A; HIV infection CDC Group IV subgroup B; HIV infection CDC Group IV subgroup C1; HIV infection CDC Group IV subgroup C2; HIV infection CDC Group IV subgroup D; HIV infection CDC Group IV subgroup E; HIV infection CDC category A; HIV infection CDC category B; HIV infection CDC category C; HIV infection CDC group IV; HIV infection WHO clinical stage I; HIV infection WHO clinical stage II; HIV infection WHO clinical stage III; HIV infection WHO clinical stage IV; Malnutrition; Perinatal HIV infection; Prophylaxis against HIV infection; Tuberculosis.

Clinical trial data

- Number of cases: 11 (blinded therapy [2], BNT162b2 [9]) (1.6% of 668 cases, the total CT dataset, compared to 7 cases (1.0%) retrieved in the PSUR #2.

- Reported Pts (16): Condition aggravated, Maternal exposure during pregnancy, Mental disorder (2 each), Atrial fibrillation, Cephalo-pelvic disproportion, Constipation, Craniocerebral injury, Failed trial of labour, Headache, Intestinal obstruction, Lumbar spinal stenosis, Prostate cancer, and Spinal claudication (1 each). None of the events were related to BNT162b2 or blinded therapy.

Post-authorisation data

- Number of cases: 197 (0.04% of 507,683 cases, the total PM dataset), compared to 393 cases (0.06%) retrieved in the PSUR #2.

  Patients with pre-existing HIV Infection:

  - 107 cases (0.02% of 507,683 cases, the total PM dataset).

  Patients with pre-existing tuberculosis:

  - 67 cases (0.01% of 507,683 cases, the total PM dataset).

  Patients with pre-existing malnutrition:

  - 23 cases (<0.01% of 507,683 cases, the total PM dataset).

MAH’s conclusion

No safety signals have emerged based on the review of these cases. Safety surveillance will continue.

Rapporteur assessment comment:

Based on the data presented concerning individuals with pre-existing HIV infection, with pre-existing tuberculosis, or with pre-existing malnutrition, no new important safety concern could be identified.

Clinical reactogenicity data on individuals previously exposed or not to SARS-COV-2
Data are available from 3 analyses: children 2 to <5 years and children 6 months to <2 years receiving up to 3 primary doses of BNT162b2 3 μg, and adults 18-55 years receiving a fourth dose booster of either the current vaccine or a monovalent Omicron-modified vaccine.

Children 6 months to <2 years (from C4591007)

- Subgroups of Phase 2/3 pediatric participants 6 months to <2 years of age had similar reactogenicity, with regard to local reactions and systemic events, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution.
- There were no meaningful differences in the overall patterns of local reactions and of systemic events across these subgroups.

Children 2 to <5 years (from C4591007)

Subgroups of Phase 2/3 pediatric participants 2 to <5 years of age had similar reactogenicity, with regard to local reactions and systemic events, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution.

There were no meaningful differences in the overall patterns of local reactions and of systemic events across these subgroups.

Adults 18 through 55 years (from C4591031 Substudy D)

- Any local reactions reported within 7 days after first study (Dose 4) vaccination for Cohort 2 participants were similar in the BNT162b2 OMI (78.6%) and BNT162b2 (79.4%) groups, and most events were mild or moderate in severity. No Grade 4 local reactions were reported.
- Any systemic events reported within 7 days after first study (Dose 4) vaccination for Cohort 2 participants were similar in the BNT162b2 OMI (77.6%) and BNT162b2 (72.9%) groups, and most events were mild or moderate in severity. No Grade 4 systemic events were reported.
- The baseline positive subgroup included a limited number of participants, and their results should be interpreted with caution. Overall, numerical differences in any of the local reactions and of the systemic events by SARS-CoV-2 baseline status were not considered clinically meaningful.

Rapporteur assessment comment:

No new important safety concern could be identified based on the data presented regarding reactogenicity on individuals previously exposed or not to SARS-202.

Local adverse reactions

Search criteria: Pt's Erythema; Injection site erythema; Injection site pain; Injection site swelling; Swelling.

Clinical trial data

- No serious clinical trial cases of local reactions reported during the reporting interval; no cases were retrieved in the PSUR #2.

Post-authorisation data
- Number of cases: 8,597 (1.7% of 507,683 cases, the total PM dataset), compared to 21,240 cases (3.2%) retrieved in the PSUR #2.

- Most frequently reported relevant PTs (≥2%): Erythema (4137), Swelling (4036), Injection site pain (690), and Injection site swelling (212).

- Time to event onset (n = 5683) range: range: <24 hours to 366 days, median: 1 day.

- Duration of relevant events (n = 1328 out of 9440 occurrences with outcome of resolved/resolved with sequelae), range = <24 hours to 233 days, median 3 days.

- Relevant event outcome: fatal (6), resolved/resolving (4065), resolved with sequelae (99), not resolved (2582), unknown (2524).
  - There were 6 cases reporting fatal events of interest (Erythema [4 cases] and Swelling [2 case]) in elderly (4 cases) and adult (2 cases) patients. Time to onset of fatal events were < 24 hours (1 event), 1 day (1 event), 3 days (1 event), 4 days (1 event), and unknown days (2 events). Review of these cases identified additional fatal adverse events reported in these cases and the local adverse reactions were not the primary cause of death in these cases.

Analysis by age group

- Post-marketing: Paediatric (512), Adults (6972), Elderly (373) and Unknown (740). In general, the events of interest were similar by percentage across age group.

Analysis by presence of comorbidities

- Post-marketing:
  - Number of subjects with comorbidities: 38,787 (7.6% of 507,683 cases, the total PM dataset). Subjects with comorbidities were reported in (902/10.5%) of the Local Adverse Reactions dataset. There were no differences between the group with comorbidities and the one without comorbidities.

Analysis by dose

- Post-marketing: Number of post-authorisation vaccine doses administered at the time of the event onset: Dose 1 in 2140 cases, Dose 2 in 1874 cases, Dose 3 in 2627 cases, Dose 4 in 77 case, Dose 5 in 1 case, and the dose number was not specified in 2039 cases. The majority of post-authorisation events reported across doses were similar with the exception of injection site pain being reported more frequently in the unspecified dose group.

MAH's conclusion

Local adverse reactions were reported in 8597 relevant cases representing 1.7% of the cases in the reporting period. The majority of events (79.8%) were non-serious events with 44.9% of the events resolved, resolved with sequelae or resolving at the time of reporting. There were 9 fatal cases describing fatal local adverse reactions in 6 cases; two were in adult and 4 were elderly subjects. Three of the 9 fatal cases did not report fatal local adverse reaction events. Review of these cases indicated that there were additional fatal adverse events reported and the event of interests (Erythema, Swelling) were not the primary cause of death in these subjects. When reported, the majority onset of events occurred within to <24 hours, with durations lasting <24 hours to 7 days.

The PM data appears to differ from the clinical trial data where injection site pain is generally the most frequently reported local reactogenicity event in adults and children. However, this is considered to be an effect of coding conventions given that commonly co-reported PTs in the cases are: Pain, Pain in
extremity and Vaccination site pain. Evaluation of local adverse reaction cases did not reveal any significant new safety information. Local adverse reactions are appropriately described in the RSI. Surveillance of local adverse reactions will continue.

Rapporteur assessment comment:

During the interval period, a significant decreased number of 8,597 cases (1.7% of 507,683 cases, the total PM dataset) reporting local adverse reactions were retrieved compared to 21,240 cases (3.2% of the total PM dataset) retrieved in the previous 2\textsuperscript{nd} PSUR.

Local adverse reactions are stated in the ADR table of section 4.8 of the Comirnaty Smpc. No new important safety concern could be identified for local adverse reactions. For future PSURs in the section ‘Evaluation of AEs’, the local adverse reactions should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Systemic adverse reactions

Search criteria: PTs Arthralgia; Chills; Fatigue; Headache; Myalgia; Pyrexia.

Clinical trial data

- Number of cases: 11 (BNT162b2 [10], and blinded therapy [1]) (1.6% of 668 cases, the total CT dataset) compared to 4 cases (0.6%) retrieved in the PSUR #2.
- Relevant PTs: Pyrexia (6), Arthralgia (3), Headache, and Myalgia (1 each), none of which were assessed as related to BNT162b2 by the investigator and Sponsor.

Post-authorisation data

- Number of cases: 167,760 (33% of 507,683 cases in the total PM dataset), compared to 279,184 (42.5% retrieved in the PSUR #2).
- Relevant event seriousness: serious (36801), non-serious (273,863).
- Relevant PTs: Headache (77,970), Fatigue (67,855), Pyrexia (57,671), Myalgia (43,916), Chills (33,541), and Arthralgia (29,430).
- Time to event onset (n = 253,501 ) range: <24 hours to 3654 days, median: 1 day.
- Duration of relevant events (n = 76,627 , out of which 76,067 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 1 year, 2 months 8 days, median 2 days.
- Relevant event outcome78: fatal (300), resolved/resolving (175,756), resolved with sequelae (3756), not resolved (86,147), unknown (45,676).
  - In 233 cases, the following relevant events (300) were reported as fatal: PTs Pyrexia (119), Fatigue (63), Headache (55), Chills (29), Myalgia (20), and Arthralgia (14).

Analysis by age group

- Clinical trial: Paediatric (5, PTs Pyrexia [4], Myalgia [1]), Adults (1, PT Headache), Elderly (5, PTs Arthralgia [3], Pyrexia [2]). A meaningful comparison between the different age groups is not possible due to the low number of cases.
• Post-marketing: In the current reporting interval, the most frequent systemic adverse reactions in subjects 16 years of age and older (in order from highest to lowest frequencies) were Headache (69,392), Fatigue (61,567), Pyrexia (48,928), Myalgia (40,707), Chills (30,837) and Arthralgia (27,333); the most frequent systemic adverse reactions in adolescents 12 through 15 years of age were PTs Headache (4485), Pyrexia (4727), Fatigue (2381), Myalgia (1164), Chills (1121), Arthralgia (614). Across the age groups in the table below, the greatest number of events were reported in the adult population, followed by the elderly. In general, relevant events were more likely to be assessed as non-serious and/or associated with a resolving outcome with increasing age. Generally, there were less relevant events associated with a worse outcome (not resolved/fatal).

Analysis by presence of comorbidities

• Number of subjects with comorbidities: 13,030 (2.6% of 508,351 cases in the total dataset and 7.8% of 167,771 [11 CT and 167,760 PM] cases reporting systemic adverse reactions).

• Clinical trial: None of the CT cases reported selected comorbidities.

• Post-marketing: The total proportion of relevant events were generally evenly distributed among subjects that reported selected comorbidities and subjects that did not report selected comorbidities. In subjects with selected comorbidities, the relevant event was more likely to be assessed as non-serious and/or with a resolved or resolving event outcome. Of note, subjects that reported comorbidities were more likely to be of advanced age, polypharmacy users, report more AEs on average (e.g., concurrent conditions) and/or prone to hospitalisation; therefore, assessment of the contributory role of BNT162b2 on the seriousness and outcome of these relevant events is confounded.

Analysis by dose

• Number of vaccine doses administered: 1 dose in 47,268 cases, 2 doses in 49,553 cases; 3 doses in 44,738 cases, 4 doses in 893 cases, and in 25,515 cases the dose was either not specified or reported as others.

• Clinical trial: Vaccination dose number: 2 doses (3), 3 doses (7) and 4 doses (1). A meaningful comparison by dose is not possible due to the low number of CT cases.

• Post-marketing: In general, the total proportion of relevant events, event seriousness, and event outcome were highest in those subjects who had received three doses of the vaccine; following this, most events were reported in those who had received two doses of the vaccine.

MAH’s conclusion

Systemic adverse reactions were reported in 167,771 (11 CT and 167,760 PM) cases representing 33.0% of the cases in the total dataset for the reporting period. The majority of events (88.2%) were non-serious events with 57.8% of the events resolved, resolved with sequelae or resolving at the time of reporting. Evaluation of systemic adverse reaction cases did not reveal any significant new safety information based on analysis by age group, by presence of comorbidities or by dose. Systemic adverse reactions are appropriately described in the RSI. Surveillance of systemic adverse reactions will continue.

Rapporteur assessment comment:

During the interval period, a decreased number of 167,760, (33% of 507,683 cases in the total PM dataset) reporting systemic adverse reactions were retrieved compared to 279,184 cases (42.5% of the total PM dataset) retrieved in the previous 2nd PSUR.
Systemic adverse reactions are stated in the ADR table of section 4.8 of the Comirnaty SmPC. No new important safety concern could be identified for systemic adverse reactions. For future PSURs in the section ‘Evaluation of AESIs’, the systemic adverse reactions should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Severe reactogenicity

Search criteria: PT Extensive swelling of vaccinated limb.

Clinical trial data

- There were no serious CT cases indicative of extensive swelling of vaccinated limb; no cases were retrieved in the PSUR #2.

Post-authorisation data

- Number of cases: 1613 (0.32% of 507,683 cases, the total PM dataset), compared to 1558 cases (0.24%) retrieved in the PSUR #2.
- MC cases (196), NMC cases (1417).
- Country of incidence: Netherlands (921), Belgium (590), Iraq (26), Australia (24), UK (12), France, Germany (8 each), Philippines (5); the remaining 19 cases were distributed among 10 countries.
- Subjects’ gender: female (1310), male (300) and unknown (3).
- Subjects’ age in years (n = 1536), range: 7 – 94, mean: 38.3, median: 36.0.
- Medical history (n = 497): the relevant reported medical conditions included Drug hypersensitivity (24), Hypersensitivity (8), Allergic reaction to excipient, Allergy to vaccine, Reaction to preservatives (1 each).
- COVID-19 Medical history (n = 219): medical conditions reported included COVID-19 (162), Suspected COVID-19 (54), Post-acute COVID-19 syndrome (2), and SARS-CoV-2 test positive (1).
- Co-suspects (n= 17 cases): Influenza vaccine (6), Pneumococcal vaccine polysacchar 23V (2).
- Number of relevant events: 1613
- Relevant event seriousness: serious (202), non-serious (1,411).
- Time to event onset (n = 1450) , range: range:<24 hours to 175 days, median: 1 day.
- Duration of relevant events (n = 375 out of 1,615 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 157 days, median 4 days.
- The reported relevant PT included Extensive swelling of vaccinated limb (1613).
- Relevant event outcome: fatal (1), resolved/resolving (910), resolved with sequelae (8), not resolved (583), unknown (112).
  - During the reporting period, 1 case was received from a Health Authority, reporting the relevant PT Extensive swelling of vaccinated limb with a fatal outcome. This case described a 14-year-old male patient who received BNT162b2 Intramuscularly for COVID-19 immunisation and experienced swelling of limb. The patient also experienced...
difficulty breathing (PT Dyspnoea), cyanosis (PT Cyanosis) and oedematous lower extremities (PT Oedema), all of which were reported as non-serious events. The reported cause of death was peripheral swelling. Limited information was provided in this case precluding a meaningful medical assessment, including a lack of event onset dates, event details, test results, medical history, and concomitant medications.

- A majority of the cases did not describe the type or extent of swelling and reported (verbatim) terms such as, “extensive swelling of the arm, reaction at or around the injection site, swelling limb, or extended swelling of the arm: extensive swelling of vaccinated limb”. Many cases also reported additional events related to pain, warmth, or erythema at the injection site, with no additional relevant details. Most cases described localized redness or swelling limited to the injection site and/or reports of lymph node swelling with no evidence in the case detail regarding any additional extensive swelling. For those cases reporting details of swelling, most appeared limited to the area surrounding the injection site with little evidence of additional extensive swelling of the rest of the limb. In a majority of the cases reporting swelling associated with the injection site, it was not reported if treatment was required, and no case reported long lasting or permanent sequelae following the event.

**Analysis by age group**

- Post-marketing: Paediatric (25), Adult (1506), Elderly (65), Unknown (17). A higher reporting proportion of events coded to the PT Extensive swelling of vaccinated limb was observed in elderly versus adult population (26.5% in elderly vs 20.3% in adults). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

**Analysis by presence of comorbidities**

- 51 (3.2% of the cases reporting the event severe reactivity). A higher reporting proportion of severe reactogenicity was reported in patients without significant comorbidities (96.9%) when compared to patients with significant comorbidities. The reporting proportion of the event severe reactogenicity with the outcome of resolved/resolving (58.8%) is slightly higher in individuals with comorbid conditions when compared to the reporting proportion observed in the individuals without comorbidities (56.3% of events with resolved/resolving).

**MAH’s conclusion**

There was a total of 1613 cases in the safety database reporting the PT Extensive swelling of vaccinated limb with the use of BNT162b2 which were mostly reported from the Netherlands (921) and Belgium (590). A majority of the cases involved females (1310, 81.2%) and were reported in subjects aged 31-50 years (793, 49.2%). Two-hundred and two (202; 12.5%) of the events were assessed as serious due to meeting medically significant criteria (there were 6 hospitalisations due to reported events). There was 1 case reporting a fatal outcome. One thousand two hundred and thirty-seven (1237) cases reported time to onset of the event as the same day or the day following vaccination. The majority of cases reporting swelling associated with the injection site, did not report that treatment was required, and no case reported long lasting or permanent sequelae following the event. Injection site swelling and lymphadenopathy are listed adverse drug reactions in the RSI for BNT162b2, and based on the data reviewed, there is insufficient evidence from reported cases to date that would warrant a change to the existing product information.

**Rapporteur assessment comment:**
Age-related adverse reactions

Clinical trial data

- Number of cases: 668.
- Time to event onset (n = 793), range: <24 hours to 558 days, median: 116 days.
- Relevant event outcome: fatal (50), resolved/resolving (663), resolved with sequelae (49), not resolved (115), unknown (3).

Post-authorisation data

- Number of cases: 507,683.
- Time to event onset (n = 1,196,069), range: <24 hours to 7337 days, median: 1 day.
- Relevant event outcome: fatal (8526), resolved/resolving (595,395), resolved with sequelae (26,518), not resolved (434,513), unknown (536,733).

Analysis by age group

- Clinical trial: Paediatric (103), Adults (336), Elderly (211) and Unknown (1).
  - The 5 MedDRA SOCs with the most frequently reported events for the current reporting period for each age group is presented are Table 57, Table 58 and Table 59 (not reproduced here). Of note, 139 cases reported 151 events pertaining the Infections and infestations SOC, which was included among the SOCs of the most frequently reported AEs in all 3 age groups.
  - There were 59 cases reporting 65 events in the Cardiac disorders SOC for the adult and elderly age group. Forty-five (45) cases reported relevant medical history (e.g., coronary artery disease, atrial fibrillation, congestive cardiac failure, cardiovascular disorder), which may have contributed to the relevant events. The most frequently reported events (≥3 occurrences) in the Cardiac disorders SOC for the adult and elderly age group were Atrial fibrillation (16), Myocardial infarction (9), Cardiac failure congestive, Coronary artery disease (5 each), Acute coronary syndrome, Acute myocardial infarction (4 each), Angina pectoris and Angina unstable (3).
  - There were 96 cases reporting 98 events in the Neoplasms benign, malignant and unspecified (incl cysts and polyps) SOC for the adult and elderly age group. Twenty-eight (28) cases reported pre-existing medical history of cancer (e.g., basal cell carcinoma, neoplasm malignant, pituitary tumour benign, prostate cancer). The most frequently reported events (≥3 occurrences) in the Neoplasms benign, malignant and unspecified (incl cysts and polyps) SOC for the adult and elderly age group were Prostate cancer (13), Adenocarcinoma of colon, Breast cancer, Pancreatic carcinoma (5 each), Brain neoplasm (4), Invasive ductal breast carcinoma, and Oesophageal carcinoma (3 each). When reported, latency ranged from 1 day to 437 days with a
median of 104 days. Of the 78 events reporting latency, the majority of the relevant event latency (65 events) was reported between 1 day to 6 months.

- There were 7 cases reporting 9 events in the Psychiatric disorders SOC for the paediatric age group. The 9 events reported were Depression, Suicidal ideation, Suicide attempt (2 each), Depression suicidal, Major depression and Mental status changes (1 each). The events were assessed as unrelated to BNT162b2/Blinded therapy by the investigator and the Sponsor.

- Post-marketing: Paediatric (31,832), Adults (361,138), Elderly (56,588) and Unknown (56,647).

- The top 5 MedDRA SOCs with the most frequently reported events for the current reporting period for each age group are presented in Table 60, Table 61, and Table 62 (not reproduced here). The top 5 SOCs were generally comparable for all age groups except Reproductive system and breast disorders in the adult age group, Skin and subcutaneous tissue disorders in the paediatric age group and Infections and infestations in the elderly age group.

- In the Reproductive system and breast disorders SOC for adult age group, event seriousness was assessed as serious (8609) and non-serious (61,891). Event outcome was reported as resolved/resolving (19,328), not resolved (33,732), resolved with sequelae (1,390), unknown (16,268), and fatal (6). The most commonly reported PTs (>1000 occurrences) in Reproductive system and breast disorders for the adult age group were Heavy menstrual bleeding (11,691), Menstrual disorder (11,655), Menstruation irregular (6481), Dysmenorrhea (5824), Intermenstrual bleeding (5650), Amenorrhea (5267), Polymenorrhea (4522), Menstruation delayed (4500), Oligomenorrhea (1818), Breast pain (1816), Vaginal haemorrhage (1588), and Postmenopausal haemorrhage (1028). It is not unexpected for these events of reproductive system and breast disorders to be reported more frequently in adult subjects compared to elderly and paediatric subjects (males or females of non-puberty age).

- In the Skin and subcutaneous tissue disorders SOC for paediatric age group, event seriousness was assessed as serious (966) and non-serious (4194). Event outcome was reported as resolved/resolving (2903), not resolved (1161), resolved with sequelae (20), unknown (1085), and fatal (3). The fatal cases are reviewed in Section 16.3.4.1 Death of the PSUR. The most commonly reported PTs (>110 occurrences) in Skin and subcutaneous tissue disorders for the paediatric age group were Rash (1538), Pruritus (718), Urticaria (681), Erythema (326), Hyperhidrosis (270), Rash pruritic (198), Cold sweat (131), and Sensitive skin (110). Most of these events are listed or consistent with listed events as per the current RSI.

- In the Infections and infestations SOC for elderly age group, event seriousness was assessed as serious (11,447) and non-serious (2756). Event outcome was reported as resolved/resolving (3096), not resolved (2014), resolved with sequelae (157), unknown (8305), and fatal (649). The fatal cases are reviewed in Section 16.3.4.1 Death of the PSUR. The most commonly reported PTs (>250 occurrences) in Infections and infestations for the elderly age group were coded to the PTs COVID-19 (8394), Herpes zoster (1771), Suspected COVID-19 (462), COVID-19 pneumonia (408), Influenza (407), Pneumonia (346), and Nasopharyngitis (251). It is not unexpected for these events to be reported more frequently in elderly subjects compared to adult and paediatric subjects.

MAH's conclusion
The most frequently reported SOCs and overall PTs with the COVID-19 vaccine across the age groups were listed or consistent with listed events as per the current RSI. The analysis of age-related AEs did not identify any new significant safety information.

Rapporteur assessment comment:

No new important safety concern could be identified for age-related adverse reactions.

**Vaccination stress/anxiety related ADRs**

Search criteria: PTs Anxiety; Blood pressure decreased; Blood pressure increased; Dizziness; Dyspnoea; Hyperhidrosis; Loss of consciousness; Palpitations; Parasthesia; Parasthesia oral; Syncope; Tachycardia (reported in very close temporal proximity to vaccination, e.g., when time to event onset for the relevant PTs is same day or 1 day after vaccination).

**Clinical trial data**

- Number of cases: 2, both involving BNT162b2 (0.3 % of 668 cases in the total CT dataset) compared to no cases retrieved in the PSUR #2.
- Country of Incidence: Israel, Poland (1 each).
- Subjects’ gender: male (2).
- Subjects’ age in years (n = 2), 10 years and 73 years.
- Medical history: Hypertension, Hypercholesterolaemia, Myocardial ischaemia and Glucose-6-phosphate dehydrogenase deficiency (1 each).
- COVID-19 Medical history: none.
- Co-suspects: none.
- Reported relevant PTs: Dyspnoea and Syncope (1 each).
- Time to event onset: 1 days for both the relevant events.
- Duration of relevant events: 1 day for the event Syncope, 5 days for the event Dyspnoea.
- Relevant event outcome: resolved (2).

**Post-authorisation data**

- Number of relevant cases: 39,800 (7.8% of 507,683 cases, the total PM dataset), compared to 56,230 cases (8.6%) retrieved in the PSUR #2.
- Most frequently reported relevant PTs (≥2%): Dizziness (16,611), Dyspnoea (7875), Parasthesia (6846), Tachycardia (4757), Palpitations (4754), Blood pressure increased (2539), Hyperhidrosis (2339), Syncope (2294) and Loss of consciousness (1037).
- Relevant event outcome: fatal (81), resolved/resolving (26,704), resolved with sequelae (1170), not resolved (16,695), unknown (5789).

**Analysis by age group**

- Clinical data: Paediatric (1) and Adults (1). A meaningful comparison between the different age groups is not possible due to the low number of cases.
• Post-marketing: Paediatric (3681), Adults (31,950), Elderly (2921) and Unknown (1248). No significant difference was observed in the reporting proportion of frequently (≥2%) reported relevant events between the adult and elderly populations. A higher reporting proportion of relevant PT Syncope was observed in the paediatric population when compared to the adult or elderly population (15.2% in paediatric vs 4.6% in adult vs 6.1% in elderly subjects). This is consistent with expectations based on age-related event reports from other vaccines.

Analysis by presence of comorbidities

• Number of subjects with comorbidities: 3277 (0.8 % of the cases reporting stress/anxiety ADRs). Upon review, no significant difference in the occurrence of the most frequently reported AEs related to vaccination stress/anxiety and in relevant AEs with fatal outcome in the subjects with comorbidities compared to the population without underlying diseases was identified, apart from the event syncope that was reported with higher proportion (6.3%) in subjects with comorbidities with respect to subjects without comorbidities (0.6%). The subjects’ underlying conditions are likely to be contributory to the occurrence of syncope in these cases.

MAH’s conclusion

No new significant safety information was identified based on a review of these cases.

Rapporteur assessment comment:

Anxiety and stress-related adverse events (e.g., dizziness, paraesthesia, hypoesthesia, hyperhidrosis) are stated in section 4.4 and 4.8 of the Comirnaty SmPC. No new important safety concern could be identified for vaccination stress/anxiety related ADRs.

The vaccination stress/anxiety related ADRs are considered well documented and adequately managed in clinical practice, and therefore could be removed from section 'Evaluation of other risks (not categorised as important)' in future PSURs. Request for next PSUR

Evaluation of special situations

Death

Search criteria - Death cases are identified based on the following criteria: If the case or event outcome is "Fatal"; If the date of death field has a value; If any of the history type values is "Death" or "Autopsy"; If the death field is set to "Yes"; If the case contains one of the following terms: High Level Group Term: Fatal outcomes; Preferred Terms: Assisted suicide, Completed suicide.

Clinical trial data

• Number of cases: 34 (blinded therapy [4] and BNT162b2 [30]) (5.1 % of 668 cases, the total CT dataset) compared to 44 cases (6.1%) retrieved in the PSUR #2.

• Causes of death most frequently reported (>2 occurrences): Death (6), Disease progression (5), and Completed suicide (4). None of the fatal events are considered related to blinded therapy/BNT162b2.

Post-authorisation data

• Number of cases: 3163 (0.6% of 507,683 cases, the total PM dataset) compared to 5215 (0.8%) analysed in the PSUR #2.

• MC cases (2061), NMC cases (1102).
- Country of incidence (≥107 occurrences): Germany (655), France (304), Japan (252), Philippines (205), Austria (194), the UK (164), Malaysia (151), the US (138), Australia (122), and Italy (107).

- Subjects’ gender: female (1304), male (1722), unknown (137).

- Subjects’ age in years (n = 2901), range: 5.0 – 107.0 years, mean: 68.0 years, median: 73.0 years.

- Medical history (n = 1631): The most frequently reported (>70 occurrences) medical conditions included cardiac and vascular disorders [e.g., Hypertension (588), Atrial fibrillation (171), Cardiac failure (113), Dyslipidaemia (80), and Myocardial ischaemia (72)]. Other most frequently reported (>70 occurrences) medical conditions included Diabetes mellitus (169), Type 2 diabetes mellitus (117), Obesity (102), Chronic obstructive pulmonary disease (95), Dementia (83), and Chronic kidney disease (72).

- COVID-19 Medical history (n = 98): COVID-19 (86), Suspected COVID-19 (9), COVID-19 pneumonia (8), Coronavirus infection, Post-acute COVID-19 syndrome, and SARS-CoV-2 antibody test positive (1 each).

- Causes of death most frequently reported (>100 occurrences): Death (739), COVID-19 (301), Cardiac arrest (215), Dyspnoea (185), Myocardial infarction (154), Vaccination failure (144), Drug ineffective (131), COVID-19 pneumonia (129), Sudden death (110), Pulmonary embolism (105), Cardio-respiratory arrest (102), and Cardiac failure (101).

- Autopsy results were provided in 165 cases and the most commonly reported (≥7 occurrences) were: Pulmonary embolism (22), Myocarditis (18), Pulmonary oedema (12), Arteriosclerosis coronary artery, Myocardial infarction, Myocardial ischaemia (10 each), Acute myocardial infarction, Arteriosclerosis (9 each), Arrhythmia, Death (8 each), Cardiac failure (7).

- Co-suspect vaccines/medications (n = 144): the most frequently reported (>3 occurrences) were COVID-19 vaccine (25), Influenza vaccine (16), COVID-19 vaccine MRNA (MRNA 1273) (15), COVID-19 vaccine AstraZeneca (CHADOX1 NCOV-19) (14), Influenza vaccine INACT SPLIT 4V (8), Influenza vaccine INACT SAG 4V (6), Casirivimab/lumivimab (5), apixaban, furosemide, and lenalidomide (4 each).

- Cases with confounders and risk factors: 1726 fatal cases included one or more contributing factors, which precluded a meaningful causality assessment: co-suspect (144 cases), concomitant drugs (638 cases) and/or underlying medical history/risk factors (1652 cases).

- Events with a fatal outcome (n = 8335): The most frequently reported (>100 occurrences) fatal events were coded to the PTs: Death (652), COVID-19 (340), Immunisation (240), Cardiac arrest (222), Vaccination failure (218), Dyspnoea (217), Drug ineffective (209), Off label use (193), Myocardial infarction (155), Interchange of vaccine products (144), Sudden death (140), COVID-19 pneumonia (137), Pyrexia (119), Pulmonary embolism (116), and Cardiac failure (102).

- Time to fatal event onset (n = 5580), range: <24 hours to 365 days, median: 8 days

**Analysis by age group**

- Clinical trial: Adults (18–64) (17) and Elderly (65 years and older) (17). A meaningful comparison between age groups is not possible due to the low number of cases with a fatal outcome.
• Post-marketing: Paediatric (17 years and under) (82), Adults (18-64 years) (932), Elderly (65 years and older) (1946), and Unknown (203).
  
  o There is a significant difference observed in the reporting proportion for the majority of the frequently reported fatal events (>100 occurrences) in the elderly population when compared to the adult population due to a higher proportion of fatal cases reported in subjects over 64 years of age (61.5% vs 29.5%, respectively). There is no meaningful comparison between elderly vs paediatric population possible due to the low number of paediatric fatal cases reported (2.6% vs 61.5%, respectively).
  
  o Most of the cases reporting a fatal outcome (42.1%) were in subjects over 75 years of age. The elderly population is generally considered a priority group targeted for vaccination by many regions and countries, including Europe and the US (with various lower age cut-offs across countries), because of their higher risk of severe disease and mortality if infected with SARS-CoV-2.

Analysis by presence of comorbidities

• Number of subjects with comorbidities: 1094 (0.2 % of 508,351, the total dataset) when compared to 2090 (0.3% of 658,249 cases) in the PSUR #2. Upon review, there were no significant differences observed in the patterns of the most frequently reported fatal events (>100 occurrences) between the group with comorbidities and the one without comorbidities.

Analysis by dose

• Number of vaccine doses administered at the time of the subjects' death:
  
  o First dose (378 cases).
  
  o Second dose (934 cases). Of the 934 cases, 163 cases (17.5 %) reported a latency of same day to 3 days after vaccination. There were 2477 fatal events. The most frequently reported (>100 occurrences) fatal events were coded to the PTs COVID-19 (178), Death (154), Drug ineffective (113), Vaccination failure (111).
  
  o Third dose (1084 cases). Majority of these cases (>50 occurrences) originated from Germany (240), Japan (151), France (139), the UK (78), and Austria (56). There were 3267 fatal events. The most frequently reported (>100 occurrences) fatal events were coded to the PTs Death (206) Immunisation (188), Off label use (117), COVID-19 (112), Interchange of vaccine products (107), and Vaccination failure (101).
  
  o Fourth dose (71 cases). Majority of these cases (>10 occurrences) originated from Germany (23), France, and the UK (11 each). There were 254 fatal events. The most frequently reported (>20 occurrences) fatal events were coded to the PTs Off label use (42) Immunisation43 (39), and Death (22).
  
  o Fifth dose (1 case). This is a spontaneous case reported by a consumer. In this case, a 66-year-old male subject received BNT162b2, as dose 5 (booster), for COVID-19 immunisation (Off label use). Relevant medical history included Interchange of vaccine products (first 2 doses with Coronavac; third and fourth doses with BNT162b2) and hospitalisation for the drop in oxygen saturation. The subject's condition worsened after receiving the fifth dose and he experienced immunisation reaction such as low oxygen saturation, lung oedema, abnormal lung function and shortness of breath, and he died 3 days later. Oxygen deficiency and failure of the lungs to function were cited as the cause of death. It was unknown if an autopsy was performed.
In the remaining cases (695), dose number was not specified at the time of the subject's death.

**Literature**

- Review of the literature did not identify any significant new information regarding the use of BNT162b2 and death.

**Observed versus expected analysis**

O/E analysis was performed for events with a fatal outcome. All O/E ratios were below 1.

**MAH’s conclusion**

No new risks were identified following review of fatal cases.

**Rapporteur assessment comment:**

No new important safety concern could be identified for cases reporting fatal outcome.

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**Overdose**

Search criteria: HLT Overdoses NEC OR PT Accidental overdose.

**Clinical trial data**

- No serious clinical trial cases of overdose of the vaccine reported during the current interval period, similar to no cases in the PSUR #2.

**Post-authorisation data**

- Number of cases: 1595 (0.3% of 507,683 cases, the total PM dataset), compared to 1985 cases (0.3%) retrieved in the PSUR #2.
- Relevant PTs: Overdose (1510), Accidental overdose (81), and Intentional overdose (4).
- Relevant event outcome: resolved/resolving (68), not resolved (12), fatal (3), resolved with sequelae (2), unknown (1510).

**Analysis by age group**

- Paediatric (630), Adults (420), Elderly (89) and Unknown (456). Upon review, no significant differences in the reporting proportion of the most frequently co-reported AEs were noted between the different age groups.

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 63 (4.0% of the total cases reporting overdose). Upon review, no significant differences in the occurrence of the most frequently co-reported AEs in the subjects with comorbidities compared to the population without underlying diseases was identified.

**Literature**

- Review of the literature did not identify any significant new information regarding overdoses of BNT162b2.

**MAH’s conclusion**

The most frequently reported reasons (≥ 2%) for overdose were:
• administration of incorrect dose of diluted vaccine, different from the recommended 0.3 ml for the subjects aged ≥ 12 years and 0.2 ml for the paediatric subjects aged 5 through 11 years (411; 25.8% of the total cases reporting overdose);

• administration of undiluted vaccine (582; 36.5% of the total cases reporting overdose);

• dilution with a volume of sodium chloride different from the recommended 1.8 ml for the subjects aged ≥ 12 years and 1.3 ml for the paediatric subjects aged 5 through 11 years (170; 10.7% of the total cases reporting overdose);

• administration of more than 1 dose of vaccine (39; 2.4% of the total cases reporting overdose);

• incorrect vaccine formulation administered to paediatric subjects aged 5 through 11 years instead of the recommended 10 mcg dosage (71; 4.4% of the total cases reporting overdose).

In most cases, the incorrect preparation and/or administration of vaccine occurred by mistake. In 218 cases, the reason for overdose was not reported or unclear, 2 of which reported the PT Intentional overdose. In the remaining 2 cases reporting intentional overdose, an administration of 30 mcg in children (aged 9 and 10 years old) was reported.

No new significant safety information was identified based on the review of these cases. The majority of the most frequently co-reported AEs other than overdose and medication error PTs were events consistent with the known reactogenicity of the vaccine and listed in Section 4.8 Undesirable effects of the CDS.

**Rapporteur assessment comment:**

No new important safety concern could be identified for overdose. For future PSURs in the section 'Evaluation of special situations’, the overdose should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

**Abuse, misuse, and drug dependency**

Abuse Search Criteria: PTs Alcohol use disorder; Dependence; Disturbance in social behaviour; Dopamine dysregulation syndrome; Drug abuse; Drug abuser; Drug dependence; Drug dependence, antepartum; Drug dependence, postpartum; Drug detoxification; Drug diversion; Drug level above therapeutic; Drug level increased; Drug rehabilitation; Drug screen; Drug screen positive; Drug tolerance; Drug tolerance decreased; Drug tolerance increased; Drug use disorder; Drug use disorder, antepartum; Drug use disorder, postpartum; Drug withdrawal convulsions; Drug withdrawal headache; Drug withdrawal maintenance therapy; Drug withdrawal syndrome; Drug withdrawal syndrome neonatal; Maternal use of illicit drugs; Needle track marks; Neonatal complications of substance abuse; Pharmaceutical nomadism ; Substance abuse; Substance abuser; Substance dependence; Substance use; Substance use disorder; Toxicity to various agents; Withdrawal syndrome.

Misuse Search Criteria: Intentional product misuses; Intentional device use issue; Intentional dose omission; Intentional medical device removal by patient; Intentional product use issue; Intentional removal of drug delivery system by patient; Intentional overdose; Performance enhancing product use; Prescription drug used without a prescription; Treatment noncompliance.

Of the 55 cases, 44 cases were determined to be non-contributory and were not included in the discussion.

**Clinical trial data**
• No serious clinical trial cases of abuse or misuse of the vaccine reported during the reporting period; no cases were retrieved in the PSUR #2.

Post-authorisation data

• Number of cases: 11 (0.002% of 507,683 cases, the total PM dataset), compared to 45 cases (0.01%) retrieved in the PSUR #2.
• Relevant PTs: Intentional product misuse, Intentional product use issue (4 each), Intentional underdose (3).
• Relevant event outcome: fatal (1), not resolved (3), unknown (7).

In the case involving the fatal outcome, a female subject (age unknown) received three vaccines COVID-19 (BNT162b2), pneumonia vaccine (unspecified) and the flu vaccine at one time. The patient experienced a myocardial infarction and died. Onset date of myocardial infarction was not reported.

Analysis by age group

• Post-marketing: Paediatric (2), Adults (5), Elderly (2), and Unknown (2). There was no meaningful difference between different age groups.

Analysis by dose

• Post-marketing: Number of vaccine doses administered at the time of the event onset: dose 1 in 1 case, dose 2 in 1 case, dose 3 in 1 case, and number of doses was not specified in 8 cases. There are no differences between the AEs that occurred after the first, the second and the booster dose.

Literature

• Review of the literature did not identify any significant new information regarding the use of BNT162b2 and abuse, dependence or misuse.

MAH’s conclusion

Overall, there were 11 cases representing 0.002% of the overall post-marketing dataset, that reported events indicative of misuse. These cases involved either improper storage, improper dilution of vaccine, administration of vaccine to unapproved age groups or administration of vaccine at a dose lower than the recommended dose. In general, the most frequently co-reported events observed in these cases were consistent with those observed in the overall population. No safety signals have emerged that would be considered specific to this population.

Rapporteur assessment comment:

No new important safety concern could be identified for abuse, misuse, and drug dependency. For future PSURs in the section 'Evaluation of special situations', the abuse, misuse, and drug dependency should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Occupational exposure

Search criteria: PTs Exposure to contaminated device; Occupational exposure to product; Occupational exposure to radiation; Occupational exposure to toxic agent.

Clinical trial data
• No serious clinical trial cases indicative of occupational exposure during the reporting period; no cases were retrieved in the PSUR #2.

Post-authorisation data

• Number of cases: 20 (0.004% of 507,683 cases, the total PM dataset), compared to 41 cases (0.01%) retrieved in the PSUR #2.
• Relevant PTs: Occupational exposure to product (20).
• Relevant event outcome: resolved/resolving (2), resolved with sequelae (1), unknown (17).

Analysis by age group

• Post-marketing: Paediatric (2), Adults (5), Elderly (0) and Unknown (13). A meaningful comparison between the different age groups is not possible due to the low number of cases.

Literature

• Review of the literature did not identify any significant new information regarding the use of BNT162b2 and occupational exposure.

MAH’s conclusion

Overall, there were 20 cases representing 0.004% of the overall post-marketing dataset, that reported events indicative of occupational exposure. Review of the cases did not identify any significant new information regarding the use of BNT162b2 and occupational exposure. No safety signals have emerged that would be considered specific to this population.

Rapporteur assessment comment:

No new important safety concern could be identified for occupational exposure. For future PSURs in the section ‘Evaluation of special situations’, the occupational exposure should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Lack of therapeutic efficacy

Search Criteria: PTs Drug Ineffective, Vaccination failure.

Clinical trial data

• No lack of efficacy cases in the clinical trial dataset for this reporting period or for the reporting period of PSUR #2.

Post-authorization data

• Number of cases: 51,028 (10.1% of 507,683 cases, the total PM dataset), compared to 21,457 cases (3.3%) in PSUR #2. The increase in the reporting proportion of LOE cases was multifactorial. A high number of cases were reported from Austria (31,629 cases in the current PSUR), as compared to the previous PSURs (9009 cases in PSUR #2 and 204 cases in PSUR #1) due to the active solicitation of LOE cases, including retrospective cases, by the Austrian Board of Health starting from August 2021. In addition, the epidemiology of the virus has changed since December 2021 in that the Omicron variant has become predominant in most regions. BNT162b2 efficacy against Omicron variants is less than against the previous dominant variants of concern.
• MC cases (39,368), NMC cases (11,660).
• Relevant lack of efficacy events: 51,028 (Vaccination failure [24,762] and Drug ineffective [26,266]).

• Country of incidence (≥2%): Austria (31,629), US (4734), UK (2316), Germany (1856), France (1478), Netherlands (1291); the remaining 7724 cases were distributed among 71 countries.

• Subjects’ gender: female (27,177), male (21,802) and unknown (2049).

• Subjects’ age in years (n = 48,297), range: 1.5 – 107.0 years, mean: 47.3 years, median: 47.0 years.

• Relevant event seriousness: all serious.

Confirmed vaccination failure (24,077 cases)

• Age groups: Child (40), Adolescent (1053), Adult (18,337), Elderly (4475) and Unknown (172).

• Time to event onset was known for 23,013 cases:
  o Time to onset reported after the second dose ranged from 7 – 501 days.
  o Time to onset reported after the third dose ranged from 1 – 293 days.
  o Time to onset reported after the fourth dose ranged from 1 – 213 days.


• Outcome of COVID-19 infection related events: resolved/resolving (2187), resolved with sequelae (29), not resolved (673), unknown (21,115), and fatal (221).

• Of the 24,077 subjects with confirmed vaccination failure, in 880 cases, the COVID-19 events were severe, resulting in: Hospitalisation (non-fatal/non-life threatening) (623), Disability (13), Life threatening (40), Death (204).

Suspected vaccination failure (1402 cases)

Not a vaccination failure case (25,549 cases)

SARS-CoV-2 Variants (11,901 cases)

In 11,901 of the 51,028 cases, information on SARS-CoV-2 variants was provided:

• Delta (India) variant (11,274 cases)
  o Country of Incidence (≥3 occurrences): Austria (11,164), France (84), Germany (16), and US (4).
  o Lack of efficacy events: Vaccination failure (6591) and Drug ineffective (4683).
  o Outcome of COVID-19 infection related events160: resolved/resolving (50), resolved with sequelae (2), not resolved (23), unknown (11,156), and fatal (51).

• Omicron variant (606 cases)
  o Country/region of incidence (≥2 occurrences): Hong Kong (391), France (79), Germany (40), US (39), Japan (12), Spain (6), Austria (4), Belgium, Brazil, Mexico, and Norway (3 each).
- Lack of efficacy events: Vaccination failure (404) and Drug ineffective (202).
- Outcome of COVID-19 infection related events: resolved/resolving (81), not resolved (11), unknown (503), and fatal (18).

- Alpha (UK) variant (19 cases)
  - Country of incidence: Austria, Germany (5 each), France, Italy (4 each), and Poland (1).
  - Lack of efficacy events: Vaccination failure (16) and Drug ineffective (3).
  - Outcome of COVID-19 infection related events: resolved/resolving (8), not resolved (1), unknown (10), and fatal (1).
- Others (2 cases)
  - In 2 other cases, variant was reported as Beta (South Africa162) and South African or Brazilian (as reported), respectively.

**Literature**

- Review of the literature identified significant new information with regards to the use of BNT162b2 and lack of therapeutic efficacy.

**MAH’s conclusion**

No new safety signals have emerged based on a review of these cases.

**Rapporteur assessment comment:**

During the interval period, a substantial increase of cases reporting lack of efficacy was retrieved by the MAH (51,028 [10.1% of total PM dataset] compared to 21,457 cases [3.3%] in the previous 2nd PSUR. The MAH stated as reasons for this increase that a high number of cases were reported from Austria due to the active solicitation of lack of efficacy cases (including retrospective cases), and that Comirnaty efficacy against current Omicron variants is less than against the previous dominant variants.

In view of the 843,724,061 administered Comirnaty doses during the current interval period, a total of 24,077 (0.003%) confirmed vaccination failures cases is not considered a safety signal.

No new important safety concern could be identified for lack of therapeutic efficacy.

**Off-label use**

Search criteria: PTs Contraindicated product administered; Contraindicated product prescribed; Drug effective for unapproved indication; Drug ineffective for unapproved indication; Intentional device use issue; Intentional product use issue; Intentional overdose; Off label use; Off label use of device; Prescribed overdose; Product administered to patient of inappropriate age; Product use in unapproved indication; Product use issue; Therapeutic product effective for unapproved indication; Therapeutic product ineffective for unapproved indication.

**Post-authorisation data**

- Number of cases: 29,805 (5.9% of 507,683 cases, the total PM dataset), compared to: 22,533 (3.4%) cases retrieved in the PSUR #2. A general increase in cases reporting Interchange of
vaccine products was noted (54.0% of PM cases from PSUR #2 versus 83.3% of PM cases retrieved during this reporting period).

- **MC cases (6091), NMC cases (23,714).**
- **Country of incidence (≥2%):** UK (10,172), Netherlands (6230), Germany (4368), France (1516), Poland (602)
- **Subjects' gender:** female (20,994), male (7831) and unknown (980).
- **Subjects' age in years (n = 26,283), range:** 0.01–104 years, mean: 45.8 years, median: 44.0 years.
- **Medical history (n = 12,399):** the most frequently (≥2%) reported medical conditions include PT Disease risk factor (1663), COVID-19 (1591), Suspected COVID-19 (1448), Hypertension (1089), Breast feeding (1061), Asthma (746), Immunodeficiency (581), Hypothyroidism (340), Diabetes mellitus (319), Hypersensitivity (296), Steroid therapy (293), Depression (281), Drug hypersensitivity (279), Seasonal allergy (271).
- **COVID-19 Medical history (n = 3001):** the most frequently (≥2%) reported medical conditions included COVID-19 (1591) and Suspected COVID-19 (1448).
- **Co-suspects (n = 1745 cases):** the most frequently (≥2%) reported co-suspect vaccines/medications included COVID-19 vaccine MRNA (MRNA 1273) (681), COVID-19 vaccine NRVV AD (CHADOX1 NCov-19) (420), Influenza vaccine (188), Influenza vaccine inact SAG 4V (80), Influenza vaccine inact SPLIT 4V (64), JNJ 78436735 (51), COVID-19 vaccine (50).
- **Number of events:** 174,381 (of which 32,211 were events of interest).
- **Relevant event seriousness:** 42 serious (10,382), non-serious (21,845).
- **Most frequently reported relevant PTs (≥2%):** Off label use (29,562) and Product use issue (2531). Of note, of the 29,805 cases, 696 did not report additional events. The majority of cases described off-label use as:
  - intentionally used in unapproved populations such as those mentioned below:
    - It is unknown whether the BNT162b2 vaccine is excreted in human milk.
    - Administration of the vaccine in pregnancy should be considered when potential benefits outweigh any potential risks for the mother and foetus.
    - Immunocompromised persons, including individuals receiving immunosuppressant therapy, may have a diminished immune response to the vaccine.
    - The administration of the BNT162b2 vaccine should be postponed in individuals suffering from acute severe febrile illness.
    - The safety and efficacy have not yet been established in individuals under 5 years of age. The safety and effectiveness of a booster dose of in individuals 16 through 17 years of age is based on safety and effectiveness data in adults at least 18 through 55 years of age.
  - alternative dosing or scheduling regimens (i.e., Full primary series not received, longer/shorter number of days between doses than recommended)
- The primary series of the BNT162b2 vaccine is administered as 2 doses at
greater than or equal to 21 days (preferably 3 weeks) apart. Off label is
currently considered when the 2nd dose of the vaccine is administered outside
the 19-42 day range from the 1st dose.
  - co-administration with other vaccines (i.e., influenza)
    - No interaction studies have been performed
      - administration of COVID-19 vaccines from different manufacturers and
        third/booster(extra doses).
      - administration of COVID-19 vaccine formulations indicated for a different age group.
      - usage of poor quality COVID-19 vaccines due to either preparation (i.e., dilution
technique) and/or storage issues (i.e., used after the expiry or beyond use date).

**Analysis by dose interval**

- Among these cases, 9 (all non-serious) reported administration of 3 doses of BNT162b2 with
different time intervals than the recommended posology and included the relevant PTs Off label
use (9) and Product use issue (1).

- Upon review, there were no significant differences were identified in the occurrence of the most
frequently relevant PTs and clinical co-reported AEs reported in those who received the 3 doses
of vaccine at a different time interval than the recommended posology when compared to the
population receiving BNT162b2 in unapproved conditions Clinical events reported more than
once in this population included Headache (4), Pain, Pyrexia, and Vaccination site pain (2
each).

**Literature**

- Review of the literature did not identify any significant new information with regards to the off-
label use of BNT162b2.

**MAH's conclusion**

Review of these cases did not identify new safety information related to off-label use.

**Rapporteur assessment comment:**

No new important safety concern could be identified for off-label use. For future PSURs in the section
'Evaluation of special situations', the off-label use should only be included and discussed in the PSUR if
the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

**Unexpected therapeutic effect**

Search criteria: **PTs Device effect increased; Drug effect faster than expected; Drug effective for
unapproved indication; Therapeutic product effective for unapproved indication; Therapeutic response
changed; Therapeutic response increased; Therapeutic response prolonged; Therapeutic response
unexpected; Therapeutic product effect increased; Therapeutic product effect prolonged.**

**Clinical trial data**

- No serious clinical trial cases with the above PTs reported during the reporting period; no
serious cases were retrieved in the PSUR #2.
Post-authorisation data

- Number of cases: 664 (0.1 % of 507,683 cases in the total PM dataset), compared to 844 cases (0.1%) retrieved in the PSUR #2.
- In most of the cases (when specified), the unexpected therapeutic effect included improvement in the following: pain, breathing, allergies, skin conditions (including warts and psoriasis), autoimmune/inflammatory diseases (e.g., arthritis, multiple sclerosis, ulcerative colitis), migraine/headache, infections (e.g., herpes and other viral infections, fungal infections), neoplasia (remission/regression of various cancers), movement/mobility, energy, hair growth/loss, menstruation, and general health/well-being (e.g., "felt better").

Analysis by age group

- Post-marketing: Paediatric (3 [1 Child, 2 Adolescent]), Adults (285), Elderly (109), Unknown (267). There was no meaningful difference between different age groups.

Literature

- Review of the literature did not identify any significant new information regarding the use of BNT162b2 and unexpected therapeutic effect.

MAH's conclusion

In most of the cases, the unexpected therapeutic effect included improvement in the following: pain, allergies, menstrual disorders, breathing, skin conditions (including warts and psoriasis), arthritis, migraine, headache, herpes infections, taste, smell, eyesight and cognitive skills. In the majority of the cases, the subject's experienced the unexpected therapeutic effect following the first dose. No significant new information was identified with regards the use of BNT162b2 and unexpected therapeutic effects.

Rapporteur assessment comment:

No new important safety concern could be identified for unexpected therapeutic effect. For future PSURs in the section 'Evaluation of special situations', the unexpected therapeutic effect should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. Request for next PSUR

Update on special populations

Use in elderly

Clinical trial data

- Number of cases: Number of cases: 211 (BNT162b2 [180], blinded therapy [26], placebo [4], BNT162b2S01 [1]; (31.6 % of 668 cases in the total CT dataset), compared to 233 cases (32.3%) retrieved in the PSUR #2.
- Number of events: 274.
- Most frequently (≥5 occurrences) reported PTs: Atrial fibrillation (11), Cerebrovascular accident, Osteoarthritis, Prostate cancer (9 each), Condition aggravated (8), Acute kidney injury, Acute respiratory failure, Dyspnoea (5 each).
- Of the 274 events, the only related event was for BNT162b2 and coded to the PT Dehydration (1).
Post-authorisation data

- Number of cases: 56,584 (11.1% of 507,683 cases in the total PM dataset), compared to 87,982 cases (13.4%) retrieved in the PSUR #2.

- MC cases (28,690), NMC cases (27,894).

- Country of incidence (>500 occurrences): Germany (10,884), Austria (9277), France (7504), US (3203), UK (3119), Japan (2225), Netherlands (2140), Sweden (2091), Australia (1747), Italy (1596), Spain (1187), Malaysia (1025), Denmark (992), Belgium (984), Poland (804), Philippines (689), Slovenia (614), Norway (602), Finland (592), Canada (556); the remaining 4753 cases were distributed among 62 countries.

- Subjects' gender: female (33,348), male (22,179), unknown (1057).

- Subjects' age in years (n = 54,943), range: 65 – 120, mean: 73.9, median: 72.

- Medical history (n = 18,647): the most frequently (≥1000 occurrences) reported medical conditions included the following HLGTs: Vascular hypertensive disorders (6169), Glucose metabolism disorders (incl diabetes mellitus) (2721), Allergic conditions (2365), Bronchial disorders (excl neoplasms) (1789), Cardiac arrhythmias (1712), Lipid metabolism disorders (1683), Joint disorders (1580), Thyroid gland disorders (1409), Therapeutic procedures and supportive care NEC (1392), Lifestyle issues (1342), Coronary artery disorders (1226), Central nervous system vascular disorders (1081).


- Co-suspects (n = 1956 cases) the most frequently (≥10 occurrences) reported co-suspect medications included: COVID-19 vaccine (422), COVID-19 vaccine mRNA (mRNA 1273) (274), COVID-19 vaccine NRV AD (258), influenza vaccine (154), adalimumab (151), Influenza vaccine INACT SAG 4V (100), Influenza vaccine INACT SPLIT 4V (58), pneumococcal polysaccharide vaccine 23-valent (32), apixaban (29), upadacitinib (24), Influenza vaccine INACT SAG 3V (23), N3N 78436735 (18), prednisone (17), mepolizumab (16), rivaroxaban (13), rituximab (12), casirivimab,Imdevimab, Influenza vaccine INACT SPLIT 3V (11 each), atorvastatin, ibritinib, levothyroxine, risankizumab (10 each).

- Number of events: 167,970; the most frequently (>1000 occurrences) reported events were coded to the PTs were: COVID-19 (8394), Inappropriate schedule of product administration (5063), Fatigue (4864), Headache (4712), Drug ineffective (4627), Vaccination failure (4515), Pyrexia (4261), Off label use (3847), Myalgia (3431), Immunisation (3355), Arthralgia (3137), Interchange of vaccine products (2920), Dizziness (2815), Vaccination site pain (2798), Pain in extremity (2796), Malaise (2468), Dyspnoea (2455), Nausea (2155), Chills (212), Asthenia (166), Herpes zoster (1771), Pain (1706), Rash (1677), Pruritus (1194), Vomiting (1178), Diarrhoea (1179), Paraesthesia (1096), Chest pain (1089).

- Event seriousness: serious (73,170), non-serious (94,882).

- Time to event onset (n = 119,721), range: from <24 hours to 492 days, median: 2 days.

- Event outcome: fatal (5367), resolved/resolving (52,311), resolved with sequelae (4003), not resolved (39,949), unknown (66,774).
Analysis by presence of comorbidities

- Number of elderly subjects with reported comorbidities: 10,304 (18.2% of the 56,584 cases in the total elderly dataset).

- Of the cases that reported medical histories, the percentage of cases reporting an AE with a fatal outcome is higher in subjects with comorbid conditions (72.1%) when compared to the percentage of cases involving an AE with a fatal outcome in subjects without comorbidities (27.9%).

- Upon review of the most frequently (≥200 occurrences) reported AEs in cases that recorded medical histories, the PT COVID-19 Pneumonia was the only event that had a significant proportional reporting ratio of >3:1 in the elderly population with comorbidities compared to the elderly population without comorbidities.

Literature

- Review of the literature did not identify any significant new information regarding the use of BNT162b2 in elderly patients.

MAH’s conclusion

No significant differences in the reporting proportion of the most frequently reported AEs were noted between the elderly dataset and the non-elderly dataset, apart from the PTs indicative of lack of therapeutic effect, for which the reporting proportion is higher in the elderly population: COVID-19 (14.8% versus 8.8%) and Vaccination failure (7.9% versus 4.4%). This is expected due to age-related decline in immunity that results not only in increased susceptibility to infection, but also reduces the prophylactic efficacy of vaccinations. The interval data reviewed did not identify any new safety information regarding the use of BNT162b2 in elderly patients.

**Rapporteur assessment comment:**

No important new information could be identified regarding the use of Comirnaty in the elderly.

Use in paediatric population

Paediatric subjects <5 years of age

Clinical trial data

- Number of cases: 62 (blinded therapy [43], BNT162b2 [18] and pre-randomisation [1]), originated from Protocols C4591007, C4591007-OPENLABEL and C4591024 (9.3% of 668 cases, the total CT dataset), compared to 25 cases (3.5%) retrieved in the PSUR #2.

Post-authorisation data

- Number of cases: 275 (0.5% of 507,683 cases, the total PM dataset), compared to 83 cases (0.01%) retrieved in the PSUR #2.

**Rapporteur assessment comment:**

During the reporting period, the use of Comirnaty in children <5 years old was not within the current approved indication in the EU and therefore considered off-label use.

Paediatric subjects ≥5 years and ≤11 years of age
Clinical trial data

- Number of cases: 25 (blinded therapy [6] and BNT162b2 [19]), originated from Protocols C4591007, C4591007-OPENLABEL and C4591024 (3.7% of 668 cases, the total CT dataset), compared to 18 cases (2.5%) retrieved in the PSUR #2.
- PTs (34): Gastroenteritis (3), Dermatomyositis, Intestinal obstruction, Pyrexia (2 each), Appendicitis, Asthma, Colitis, Condition aggravated, Constipation, Depression, Depression suicidal, Device related infection, Diarrhoea, Drug therapy, Febrile convulsion, Hypertension, Hyponatraemia, Influenza, Kidney transplant rejection, Large intestine benign neoplasm, Mental status changes, Myalgia, Myositis, Rhinovirus infection, Seizure, Small intestinal obstruction, Syncope, Tibia fracture, and Vomiting (1 each).
- All events were assessed as unrelated to BNT162b2 or blinded therapy.

Post-authorisation data

- Number of cases: 9605 (1.9% of 507,683 cases, the total PM dataset), compared to 1227 cases (0.2%) retrieved in the PSUR #2.
- MC cases (6573), NMC cases (3032).
- Country of incidence (≥2%): US (2503), Australia (1428), Philippines (1264), Germany (1177), Japan (859), Italy (409), and Spain (386).
- Subjects' gender: female (3925), male (4133) and unknown (1547).
- Subjects' age in years (n = 8372), range: 5 – 11,25, mean: 8.4, median: 9.0.
- Medical history (n = 846): the most frequently (≥10) reported medical conditions included Asthma (158), Food allergy (62), Seasonal allergy (59), Attention deficit hyperactivity disorder (41), Hypersensitivity (34), Autism spectrum disorder, Epilepsy (30 each), Drug hypersensitivity (28), Rhinitis allergic (27), Eczema (26), Dermatitis atopic (23), Mite allergy (21), Allergy to animal (19), Constipation, Type 1 diabetes mellitus, Urticaria (15 each), Bronchospasm, Headache, Seizure (12 each), Obesity (11), Migraine (10).
- COVID-19 Medical history (n = 136): COVID-19 (121), Suspected COVID-19 (10), Asymptomatic COVID-19, Exposure to SARS-CoV-2 (2 each), and Post-acute COVID-19 syndrome (1).
- Co-suspects (n = 44): the most frequently (>1) reported co-suspect vaccines/medications included influenza vaccine (8), adalimumab, COVID-19 vaccine (7 each), measles vaccine live (Enders-Edmonston)/ mumps vaccine live (Jeryl Lynn)/ rubella vaccine live (Wistar RA 27/3), varicella zoster vaccine live (Oka/Merck) (3 each), diphtheria vaccine toxoid/ pertussis vaccine acellular/tetanus vaccine toxoid, influenza vaccine inact split 3V, meningococcal vaccine B RFHBp, NADA, NHBA OMV, and sodium chloride (2 each).
- Number of events: 22,457.
- Event seriousness: (3735), non-serious (18,725).
- Most frequently reported PTs (>3% of cases): Product administered to patient of inappropriate age (1338), Pyrexia (1289), Vaccination site pain (1213), Poor quality product administered (1063), Headache (976), Product administration error (753), Vomiting (733), Rash (556), Overdose (516), Product preparation error (429), Fatigue (425), Nausea (410), Abdominal pain (371), Dizziness (366), Chest pain (331), COVID-19 (309), Pain in extremity (293), and Underdose (290).
• Time to event onset (n = 16,236), range: from <1 day to 385 days, median: <1 day.

• Duration of relevant events (n = 3787 out of 7329 occurrences with outcome of resolved/resolved with sequelae), range: from <1 day to 109 days, median 1 day.

• Relevant event outcome: resolved/resolving (9811), resolved with sequelae (73), not resolved (3274), fatal (58), unknown (9257).

• Fatal cases: 20
  o Age: 5 years (1), 6 years (3), 7 years (4), 8 years (2), 9 years (1), 10 years (2), 11 years (5), unknown (2).
  o MC cases (17), NMC cases (3).
  o Gender: females (9), males (9), unknown (2).
  o Country: Philippines (6), Australia (4), Germany, Spain (3 each), Albania, Japan, Portugal, UK (1 each).
  o Fatal PTs (58): the most frequently (≥ 2) reported AEs included Dyspnoea (4), Cardiac arrest, Cardiopulmonary arrest, Pyrexia (3 each), Abdominal pain, Cough, COVID-19, Death, Headache, Myocarditis, Seizure, and Vomiting (2 each).
  o Medical history (n = 7): Autoimmune thyroiditis, Asphyxiating thoracic dystrophy, Brain malformation, Bronchitis, Bronchospasm, Cerebral palsy, Cognitive disorder, COVID-19, Dependence on respirator, Developmental delay, Dysphagia, Epilepsy, Gastrostomy, Hypoxic-ischaemic encephalopathy, Immunodeficiency, Intellectual disability, Joint dislocation, Kidney transplant rejection, Motor dysfunction, Myoclonic epilepsy, Neonatal asphyxia, Obstructive sleep apnoea syndrome, Pneumonia, Renal impairment, Renal transplant, Rhinitis allergic, Scoliosis, Seizure, Severe myoclonic epilepsy of infancy, Type 1 diabetes mellitus, and Varicella zoster virus infection (1 each).
  o The 20 fatal cases are summarised below:
    ▪ In 2 cases (1 MC and 1 NMC) reporting only Death as the fatal AE, neither cause of death nor information on autopsy was provided. The limited information provided prevented any meaningful assessment.
    ▪ In 2 cases, the underlying medical conditions could have predisposed to the occurrence of the fatal AEs:
      • MC case; age: 11 years; gender: male; fatal PT: Acute respiratory failure, occurred 2 days after the 1st dose of BNT162b2; medical history: brain malformation, bronchitis, cognitive disorder, dysphagia, gastrostomy, joint dislocation, myoclonic epilepsy, obstructive sleep apnoea syndrome, pneumonia, scoliosis; autopsy: not performed.
      • MC case; age: 6 years; gender: female; fatal PTs: Renal impairment, Epilepsy, Apnoea, Varicella zoster virus infection, Seizure, Sudden death, Product administered to patient of inappropriate age, death occurred 7 days after the 1st dose of BNT162b2; medical history: developmental delay, epilepsy, Immunodeficiency, renal impairment, seizure, severe myoclonic epilepsy of infancy, varicella zoster virus infection; autopsy: unknown if performed.
In one case, the reporter concluded that the death “had nothing to do” with the administration of BNT162b2 and was due to natural causes:

- **MC case; age: 6 years; gender: male; fatal PTs: Myocarditis, Cardiorespiratory arrest, COVID-19, occurred 7 days after the 1st dose of BNT162b2; medical history: rhinitis allergic, autoimmune thyroiditis, type I diabetes mellitus); autopsy: performed, results are pending.**

In the remaining 15 cases (13 MC and 2 NMC) reporting the following fatal PTs Dyspnoea (4), Cardiac arrest, Pyrexia (3 each), Abdominal pain, Cardiorespiratory arrest, Cough, Headache, Vomiting (2 each), Abdominal pain upper, Acute respiratory distress syndrome, Adverse event following immunisation, Arteriovenous malformation, Blood pressure decreased, Blood pressure immeasurable, Bradycardia, Cardiac failure acute, Cerebral haemorrhage, COVID-19, Cyanosis, Diarrhoea, Drug ineffective, haematemesis, Heart rate decreased, Immunisation, Influenza like illness, Multisystem inflammatory syndrome, Myocarditis, Nasopharyngitis, Nausea, Off label use, Pulmonary embolism, Respiratory failure, and Seizure (1 each), no confounding factors have been identified. In most cases (9) the limited information available does not allow a medically meaningful assessment; in the remaining cases (6) a causality between the vaccination and the occurrence of the fatalities cannot be ruled out, based on the temporal relationship, although no laboratory data or autopsy results provided evidence of a causal relationship.

**Rapporteur assessment comment:**

Although the Comirnaty exposure in persons aged 5-11 years is considered increased based on the EU/EEA exposure (current reporting period an estimated 3,569,821 administered doses in 5-9 years versus 391,327 administered doses in 5-9 years in the previous reporting period), worldwide interval exposure in persons aged 5-11 years (or any other age category) is not presented in the PSUR and therefore the relative post-marketing reporting rate of fatal cases in persons aged 5-11 years is not known.

There were 17 medically confirmed fatal cases in persons aged 5-11 years compared to 1 medically confirmed fatal cases in the previous reporting period. The MAH only briefly described the 17 medically confirmed fatal cases and did not provide a WHO causality assessment per case regarding Comirnaty exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases in persons aged 5-11 years, perform per case a WHO causality assessment, and an assessment according to Brighton Collaboration case definition and Level of certainty classification, if applicable. **Request for supplementary information**

**Paediatric subjects ≥12 years of age**

**Clinical trial data**

- Number of cases: 15 (BNT162b2 [14] and blinded therapy [1]) originated from Protocol C4591001 (2), C4591001-OPEN LABEL (10), C4591007-OPEN LABEL (1), C4591024 (1), and C4591031-OPEN LABEL (1) (2.2% of 668 cases, the total CT dataset), compared to 24 cases (3.3%) retrieved in the PSUR #2.

- PTs (17): Suicidal ideation, Suicide attempt, Toxic shock syndrome (2 each), Addison’s disease, Appendicitis, Constipation, Depression, Fractured skull depressed, Herpes zoster,
Major depression, Mucocutaneous rash, Pectus excavatum, Subdural haematoma, and Syncope (1 each).

- All events were assessed as unrelated to BNT162b2 or blinded therapy.

**Post-authorisation data**

- Number of cases: 21,945 (4.3% of 507,683 cases, the total PM dataset), compared to 18,451 cases (2.8%) retrieved in the PSUR #2.
- MC cases (13,478), NMC cases (8467).
- Country of incidence (>2%): Germany (3333), Philippines (3026), Australia (2220), UK (1656), Austria (1645), Malaysia (1189), Taiwan, Province of China (1186), France (1061), US (1025), Italy (628), Netherlands (582), Japan (484), and Mexico (454).
- Subjects' gender: female (11,656), male (9813) and unknown (476).
- Subjects' age in years (n = 21,661), range: 12 - 17, mean: 14.7, median: 15.0.
- Medical history (n = 2837): the most frequently (>1%) reported medical conditions included Asthma (286), Hypersensitivity (210), Seasonal allergy (194), Food allergy (108), Attention deficit hyperactivity disorder (90), Drug hypersensitivity (74), Mite allergy (73), Epilepsy (62), Depression (51), Autism spectrum disorder (48), Allergy to animal (45), Anxiety, Migraine (40 each), Immunodeficiency, Obesity (39 each), Rhinitis allergic (37), Eczema (35), Non-tobacco user (34), Acne (33), Headache (31), and Dermatitis atopica (29).
- Co-suspects (n = 148): the most frequently (>2%) reported co-suspect vaccines/medications included COVID-19 vaccine (33), adalimumab (18), influenza vaccine (15), COVID-19 vaccine mRNA (MRNA 1273) (13), influenza vaccine inact split 4V, mestranol/norethisterone (8 each), HPV vaccine VLP RL1 9V (yeast) (7), HPV vaccine VLP RL1 2V (baculovirus) (5), HPV vaccine, infliximab (4 each), ibuprofen, and semaglutide (3 each).
- Number of events: 61,071.
- Relevant event seriousness: serious (19,558), non-serious (41,530).
- Most frequently reported PTs (>2%): Headache (3495), Pyrexia (3395), Dizziness (2376), Chest pain (1956), Fatigue (1919), Vaccination site pain (1804), Nausea (1669), COVID-19 (1600), and Dyspnoea (1267).
- Time to event onset: (n = 45,162), range: from <1 day to 476 days, median: 1 day.
- Duration of relevant events (n = 9201 out of 19,141 occurrences with outcome of resolved/resolved with sequelae), range: <1 day to 329 days, median 1 day.
- Relevant event outcome: fatal (169), resolved/resolving (28,719), not resolved (12,336), resolved with sequelae (332), unknown (19,645).
- Fatal cases (62)
- Age: 12 years (12), 13 years (13), 14 years (5), 15 years (6), 16 years (11), 17 years (9), unknown (6).
- MC cases (45), NMC cases (17).
• Gender: females (28), males (32), unknown (2).

• Country (≥ 2): Philippines (19), US (8), Malaysia, Poland (6 each), Germany (4), Austria, Brazil, Japan, Taiwan (Province of China), UK (2 each).

• Fatal PTs (169): the most frequently (≥ 3) reported AEs included Death (16), Dyspnoea (8), Pyrexia (7), Cardiac arrest (6), Myocarditis (5), Cardiac failure, Headache (4 each), Asthenia, Seizure, Shock, and Vomiting (3 each).

• Medical history (n = 13): Attention deficit hyperactivity disorder, Obesity (2 each), Abdominal pain, Agitation, Amenorrhoea, Asthma, Bedridden, Chest pain, Colloid brain cyst, Cough, Cystic fibrosis, Cyst removal, Decreased appetite, Depression, Diabetes insipidus, Dizziness, Drug hypersensitivity, Dyspnoea, Dysomnia, Exercise adequate, Fatigue, Feeling abnormal, Fracture, Headache, Hereditary cerebral degeneration, Hypertension, Kawasaki’s disease, Lipoedema, Liver disorder, Lymphoedema, Lymphostasis, Oral contraception, Osteogenesis imperfecta, Ovarian enlargement, Palpitations, Physical deconditioning, Pulmonary embolism, Pulmonary veno-occlusive disease, Seasonal allergy, Somatic symptom disorder, Substance abuser, Substance use, and Weight decreased (1 each).

• The 62 fatal cases are summarised below:

• In 15 cases (9 MC and 6 NMC) reporting only Death as the fatal AE, neither cause of death nor information on autopsy was provided. The time to fatal event onset is available in 4 cases: 5 days, 7 days, 49 days, and 144 days (1 each). The limited information provided prevented any meaningful assessment.

• In 2 cases, the subjects did not die due to illness, but due to unfortunate accidents:

• MC case; age: 17 years; gender: male; fatal PT: Fall, occurred 24 days after the vaccination; autopsy: unknown if performed.

• MC case; age: 16 years; gender: male; fatal PT: Road traffic accident, occurred approximately 110 days after the 2nd dose of BNT162b2; autopsy: unknown if performed.

• In 6 cases, the underlying medical conditions could have predisposed to the occurrence of the fatal AEs:

• MC case; age: 16 years; gender: female; fatal PT: Dyspnoea, occurred 3 days after the 1st dose of BNT162b2; medical history: bronchial asthma; autopsy: unknown if performed.

• NMC case; age: 16 years; gender: female; fatal PTs: Dyspnoea (developed 6 days after the 1st dose of BNT162b2), Brain injury, Cardiac failure acute, Hypoxia, Cardiac failure (all developed 38 days after the 1st dose of BNT162b2), Sudden death, Pulmonary veno-occlusive disease, Pulmonary arterial hypertension (all developed 41 days after the 1st dose of BNT162b2), Brain oedema, Sudden cardiac death, Brain injury, Acute kidney injury, Pneumonitis, Epistaxis, Acute respiratory failure, Cardiac failure congestive (all unknown onset date); medical history: pulmonary veno-occlusive disease, amenorrhoea, cough, dyspnoea, fatigue, fracture, hypertension, lipoedema, lymphoedema, lymphostasis, osteogenesis imperfecta, ovarian enlargement; autopsy: not performed.

• NMC case; age: 16 years; gender: female; fatal PTs: Pulmonary embolism, Cardiac arrest (all developed 2 days after the 3rd dose of BNT162b2); medical history: obesity, oral contraception, pulmonary embolism; autopsy: performed, results not provided.

• MC case; age: 17 years; gender: male; fatal PTs Pneumococcal sepsis, Cardiac failure, Pneumonia pneumococcal (all occurred 92 days after the 2nd dose of BNT162b2); medical history: [redacted]
agitation, attention deficit hyperactivity disorder, depression, dyssomnia, regular exercise. Autopsy results: the subject died after consumption of from the beginning pneumonia and the influx of germs into the bloodstream as a result of cardiovascular failure. The concentration determined in the blood and brain does not justify in itself a fatal intoxication in view of a long-term intake with a tolerance effect but may have favoured the onset of death due to a substance-typical respiratory and circulatory depressive effect, also increased in combination with the effect of. The findings obtained during the autopsy and the results of the chemical-toxicological examination can be reconciled with a protracted occurrence of death.

- MC case; age: 13 years; gender: female subject; fatal PTs: Malaise (developed 1 day after the 1st dose of BNT162b2, Lot number FJ1763), Palpitations, Chest pain (all occurred 5 days after the 1st dose of BNT162b2, Lot number FJ1763), Loss of consciousness, Pulseless electrical activity, Cardiac arrest (all occurred 64 days after the 1st dose of BNT162b2, Lot number FJ1763); medical history: Kawasaki’s disease, palpitations, weight decreased, decreased appetite, feeling abnormal. Autopsy results showed that there was no possibility of myocarditis and angina pectoris, and there was no thrombus. Since symptoms such as episodes of palpitations had appeared before the vaccination, it was assessed that the vaccination was possibly related to the death, but the possibility of being the exacerbation factor could not be ruled out.

- MC case; age: 13 years; gender: male; fatal PTs Brain death, Condition aggravated (all occurred within 1 month of unknown dose number of BNT162b2, Lot number FG9428); medical history: colloid brain cyst; autopsy: unknown if performed.

- In the remaining 39 cases (30 MC and 9 NMC) reporting the following fatal PTs Pyrexia (7), Dyspnoea (6), Myocarditis (5), Cardiac arrest, Headache (4 each), Asthenia, Seizure, Shock, Vomiting (3 each), Cardiac failure, Cardiac infection, Cardiomegaly, Depressed level of consciousness, Diarrhoea, Dizziness, Hypoaesthesia, Multiple organ dysfunction syndrome, Myocardial infarction, Myocardial Injury, Pneumonia, Toxic cardiomyopathy (2 each), Abdominal pain upper, Adverse event following immunisation, Agranulocytosis, Aneurysm ruptured, Anisocoria, Anuria, Atrioventricular block, B-cell type acute leukaemia, Brain injury, Cardiogenic shock, Cerebral haemorrhage, Chest discomfort, Chills, Coma, Compartment syndrome, Completed suicide, Contusion, Cough, COVID-19, Death, Dehydration, Diabetic ketoacidosis, Enterovirus infection, Extensive swelling of vaccinated limb, Fallot’s tetralogy, Gait inability, Haematemesis, Haemorrhage intracranial, Head banging, Hypertension, Immunisation, Loss of consciousness, Malaise, Meningitis meningococcal, Metabolic acidosis, Multi-organ disorder, Musculoskeletal stiffness, Nausea, Nervous system disorder, Off label use, Pain in extremity, Peripheral swelling, Pleural effusion, Pruritus, Pulse absent, Pulseless electrical activity, Rash, Rash pruritic, Renal failure, Respiratory arrest, Rhinovirus infection, Sepsis, Septic shock, Slow response to stimuli, Stress cardiomyopathy, Sudden death, Thrombosis, Unresponsive to stimuli, Vaccination failure, Vaccination site pain, and Ventricular tachycardia (1 each), no confounding factors have been identified. In 19 cases the limited information available does not allow a medically meaningful assessment, in the remaining 20 cases a causality between the vaccination and the occurrence of the fatalities cannot be ruled out, based on the temporal relationship, although no laboratory data or autopsy results provided evidence of a causal relationship.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 960 (3.0% of 31,927 cases, the total paediatric dataset). Upon review, there was no significant differences in the occurrence of the most frequently reported AEs in the subjects with comorbidities compared to the population without underlying diseases was identified.
Analysis of confounders and risk factors

- Among the 31,927 cases involving paediatric subjects, 4423 included one or more confounders that prevented a clear causality assessment: co-suspect and/or multiple concomitant drugs (1286 cases), underlying medical history and/or comorbidities (4037 cases) or predisposing factors (e.g., asthma, cardiac disorders, depression, diabetes, menstrual disorders, renal disease, respiratory disorders, seizures/epilepsy) (503 cases).

Literature

- Review of the literature did not identify any significant new information regarding the use of BNT162b2 in paediatric subjects.

MAH’s conclusion

No relevant differences in the occurrence of the most frequently reported events, case seriousness and case outcomes were identified among the 3 paediatric age groups presented above. Additionally, no significant differences in the reporting proportion of the most frequently reported AEs were noted between the paediatric dataset and the non-paediatric dataset, apart from the PTs Vomiting (6.1% versus 2.0%) and Product administered to patient of inappropriate age (5.8% versus 1.3%).

No new significant safety information was identified based on the review of the cases reported in the overall paediatric population. The most frequently reported AEs were consistent with the known reactogenicity of the vaccine and listed in Section 4.4 Special warnings and precautions for use and/or in Section 4.8 Undesirable effects of the CDS.

Rapporteur assessment comment:

There were 45 medically confirmed fatal cases in persons aged 12-17 years compared to 40 medically confirmed fatal cases in the previous reporting period. The MAH only briefly described the 40 medically confirmed fatal cases and did not provide a WHO causality assessment per case regarding Cominatory exposure. This is not accepted. The MAH is requested to provide detailed information concerning the fatal cases in persons aged 12-17 years, perform per case a WHO causality assessment, and an assessment according to Brighton Collaboration case definition and Level of certainty classification, if applicable. Request for supplementary information

Use in pregnant/lactating women

Search criteria: Pregnancy cases are identified as cases where:

- Patient Pregnant Flag is "Yes";
- If there is a value for Pregnancy Outcome, Birth Outcome, or Congenital Anomaly;
- If Delivery Notes are available;
- If any of the valid events on the case contains one of the following:"
  - SOC Pregnancy, puerperium and perinatal conditions, or
  - HLT Exposures associated with pregnancy, delivery and lactation; Lactation disorders, or
  - PT Exposure via body fluid.

Clinical trial data
Cumulative review (Pregnancy Cases)

- Number of pregnancy cases: 697 (28.7% of the total 2426 cases from the CT dataset). These 697 cases represent 669 unique pregnancies (2 cases [a mother case and a foetus/baby case] or 3 cases [a mother case and 2 foetuses/baby cases for twins] were created for 28 pregnancies). Cases originated from clinical studies C4591001 (155), C4591015 (120), C4591001-OPENLABEL (91), C4591031-OPENLABEL (7), C4591031 (6), C4591020 (2), C4591017 (1), BNT162-01-OPENLABEL (1), BNT162-17 (2), and C4591006 (328) and study treatment was reported as BNT162B2 (466), blinded therapy (188), placebo (42) and BNT162C2 (1).

- Country of incidence: Japan (322), US (200), Brazil (49), Argentina (46), South Africa (44), Spain (19), UK (12), Germany (3) and Turkey (2).

- Of the 597 mother cases, 431 cases reported exposure to vaccine in utero without the occurrence of any clinical event. The frequently reported pregnancy related events (>1 occurrence) were coded to the PTs Maternal exposure before pregnancy (272), Maternal exposure during pregnancy (139), Maternal exposure timing unspecified (12), Exposure during pregnancy (6), Drug exposure before pregnancy (2).

- One hundred sixty-six (166) mother cases, 139 serious and 27 non-serious, reported additional clinical events, which occurred in the vaccinated mothers:
  - The frequently reported pregnancy related events (>1 occurrence) reported in these cases were coded to the PTs Maternal exposure during pregnancy (57), Abortion spontaneous (46), Maternal exposure before pregnancy (30), Pre-eclampsia (7), Cephalo pelvic disproportion (6), Abortion missed, Foetal death, Postpartum haemorrhage, Premature separation of placenta (4 each), Abortion threatened, Delivery, Ectopic pregnancy, Gestational hypertension, Premature delivery, Premature labour (3 each), Abortion incomplete, Hyperemesis gravidarum, Maternal exposure via partner during pregnancy, Miscarriage of partner, Uterine disorder (2 each).
  - Other reported clinical events were coded to the PTs COVID-19 (9), Anaemia (2), Abdominal wall haematoma, Cholelithiasis, Dehydration, Drug eruption, Endometritis, Lower respiratory tract infection, Osteoarthritis, Pneumonia, Pruritis, Pyelonephritis, Urinary tract Infection, Urinary tract procedural complication, Vascular pseudoaneurysm, Venous thrombosis limb (1 each).
  - Of the 58 cases reporting spontaneous abortion or abortion related events, in 25 cases the mother had a medical history of spontaneous abortion, alcohol/tobacco use during pregnancy, ectopic pregnancy, obesity, diabetes mellitus, uterine disorder, depression or anembryonic gestation, which might have contributed to the event and in 33 cases there was limited information regarding the mother’s obstetric history, which precluded meaningful assessment.
  - Of the 19 cases reporting elective termination, in 10 cases, the mother had a medical history of spontaneous abortion, induced abortion, alcohol/tobacco use and in the remaining 9 cases there was limited information regarding mother’s obstetric history which precluded meaningful assessment.
  - In 3 cases reporting foetal death/stillbirth the mother had a medical history of amniotic cavity infection, HIV infection and/or spontaneous abortion, which might have contributed to the event.
- In 3 cases reporting ectopic pregnancy, in 1 case, the mother had a medical history of tobacco use which might have contributed to the event, and in the remaining 2 cases, there was limited information regarding the mother's obstetric history, which precluded meaningful assessment.

- Hundred (100) baby/foetal cases, 98 serious and 2 non serious. Cases are classified according to pregnancy outcome.
  
  o Pregnancy outcome: Live birth with congenital anomaly: Thirty-one (31) of these cases reported 39 congenital anomalies that were coded to the PTs Atrial septal defect (4), Ankyloglossia congenital, Hypoxic-ischaemic encephalopathy, Neonatal hypotension, Trisomy 21 (2 each), Cleft lip, Coma neonatal, Congenital rubella syndrome, Congenital skin dimples, Congenital skin disorder, Craniosynostosis, DiGeorge's syndrome, Gnahtoschisis, Microcephaly, Neonatal pneumothorax, Neonatal intestinal perforation, Neonatal seizure, Nervous system disorder, Newborn persistent pulmonary hypertension, Osteochondrodysplasia, Patent ductus arteriosus, Polydactyly, Pyelonephritis acute, Renal failure neonatal, Renal tubular necrosis, Sepsis neonatal, Sex chromosome abnormality, Syndactyly, Thanatophoric dwarfism, Thrombocytopenia neonatal, Ventricular septal defect, Vesicoureteric reflux (1 each). Of these 31 cases, information regarding trimester of exposure was available in 17 cases. Of these 17 cases, in 12 cases foetus was exposed during the 3rd trimester, in 4 cases foetus was exposed during the 2nd trimester, and in 1 case exposure occurred during the 1st trimester. Of these 31 cases, in 5 cases the mother of the baby was on multiple concomitant medications, alcohol use, advanced age of the mother (i.e., 43 years) and/or had a medical history of in vitro fertilization which increases the chance of gene mutation. In the remaining 26 cases, there was limited information regarding the mother's obstetric history, which precluded meaningful causality assessment.

  o Pregnancy outcome: Still birth without foetal defect: During the reporting period there was 1 case reporting stillbirth without foetal defect. The event reported in this case was coded to the PT Neonatal respiratory distress syndrome. The information regarding trimester of exposure was unknown. In this case the mother of the baby had underlying medical history of amniotic cavity infection, which might have led to the development of the reported event.

  o Pregnancy outcome: Live birth without congenital anomaly: Sixty-eight (68) cases reported live birth babies without congenital anomaly. Of these 68 cases, information regarding trimester of exposure was available in 40 cases. Of these 40 cases, in 23 cases, foetus was exposed during the 3rd trimester, in 14 cases foetus was exposed during the 2nd trimester, and in 3 cases exposure occurred during the 1st trimester. The frequently reported events (>1 occurrence) in these 68 cases were coded to PTs Jaundice neonatal (11), Foetal distress syndrome (8), Premature baby (6), Neonatal pneumonia, Neonatal respiratory distress, Bronchiolitis, Neonatal respiratory distress syndrome, Hyperbilirubinaemia neonatal (3 each), Foetal hypokinesia, Neonatal tachypnoea, Dehydration, Gastroenteritis, Patent ductus arteriosus, Anaemia neonatal, Sepsis neonatal, Hypoglycaemia neonatal, Meconium aspiration syndrome (2 each). In all these 68 cases, there was limited information regarding the mother's obstetric history, which precluded meaningful causality assessment.

- Of the 697 cases, 658 cases provided pregnancy outcomes, which are provided in Table 67 (not reproduced here).
Cumulative review (Lactation cases)

- Number of lactation cases: 141 (5.8% of the total 2426 cases from the CT dataset). All these 141 cases were non-serious. Of these 141 cases, 140 cases reported only exposure to vaccine during breastfeeding (PT Exposure via breast milk) without the occurrence of any clinical events. In the remaining case the clinical event was coded to the PT Respiratory syncytial virus infection. In this case there was limited information regarding mother’s obstetric history, which precluded meaningful causality assessment.

Incremental review (CT cases)

- Number of pregnancy cases: 41 (6.1% of the total 668 cases from the CT dataset). These 41 cases represent 37 unique pregnancies (2 cases [a mother case and a foetus/baby case] for 4 pregnancies). Cases originated from clinical studies C4591015 (24), C4591001-OPENLABEL (10), C4591001, C4591031-OPENLABEL (3 each), C4591031 (1) and study treatment was reported as blinded therapy (27), and BNT162b2 (14).

- Country of incidence: South Africa (15), Brazil (11), US (6), Argentina (5), Spain (3), UK (1).

- Twenty-three (23) serious maternal cases reported additional clinical events, which occurred in the vaccinated pregnant females:
  
  - The frequently reported pregnancy related events (>1 occurrence) reported in these cases were coded to the PTs Maternal exposure during pregnancy (8), Abortion spontaneous (7), Cephalo pelvic disproportion (3), Abortion missed, Maternal exposure before pregnancy (2 each).

  - Other reported clinical events were coded to the PTs Abdominal wall haematoma, COVID-19, Pneumonia, Urinary tract infection (1 each).

  - Of the 11 cases reporting spontaneous abortion or abortion related events, in 4 cases, the mother had a medical history of spontaneous abortion or had underlying condition of obesity, which might have contributed to the event and in 7 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

- Eighteen (18) serious baby/foetal cases are classified according to pregnancy outcome:

  - Pregnancy outcome: Live birth with congenital anomaly: Five (5) of these cases reported 5 congenital anomalies that coded to the PTs Congenital rubella syndrome, DiGeorge’s syndrome, Pyelonephritis, Syndactyly, Trisomy 21 (1 each). Of these 5 cases, information regarding trimester of exposure was available in 2 cases and in these 2 cases foetus was exposed during the 2nd trimester in 1 case and the 3rd trimester in the remaining case. Of these 5 cases, in 1 case reporting Trisomy 21, the age of the mother was 43 years and advanced maternal age is a risk factor for Trisomy 21. In the remaining 4 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

  - Pregnancy outcome: Live birth without congenital anomaly: Thirteen (13) cases reported live birth babies without congenital anomaly. Of these 13 cases, information regarding trimester of exposure was available in 7 cases. Of these 7 cases, in 5 cases, foetus was exposed during the 2nd trimester and in 2 cases foetus was exposed during the 1st and the 3rd trimester each. The frequently reported clinical events (>1 occurrence) in these 13 cases were coded to the PTs Foetal distress syndrome (3), Meconium aspiration syndrome, Gastroenteritis, Jaundice neonatal (2 each). In all
these 13 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

- Of the 41 cases, 38 cases provided pregnancy outcomes, which are provided in Table 68 (not reproduced here).

Post-authorisation data

Incremental review (Pregnancy)

- Number of pregnancy cases: 3642 (0.7% of 507,683 cases, the total PM dataset), compared to 5239 cases (0.8%) retrieved in the PSUR #2. These 3642 cases represent 3419 unique pregnancies (2 cases [a mother case and a foetus/baby case] or 3 cases [a mother case and 2 foetus/baby cases for twins] were created for 223 pregnancies).

- Country of incidence (>100 occurrences): Germany (837), UK (475), Netherlands (461), Philippines (309), France (302), Sweden (162), Australia (110).

- Of the 3320 mother cases, 535 cases reported exposure to vaccine in utero without the occurrence of any clinical event. The pregnancy exposure events were coded to the PTs Maternal exposure during pregnancy (355), Maternal exposure timing unspecified (116), Maternal exposure before pregnancy (52), Exposure during pregnancy (7), Drug exposure before pregnancy (4), Foetal exposure during pregnancy (1).

- There were 2785 mother cases of which 1479 were serious and 1306 were non-serious reporting additional clinical events, which occurred in the vaccinated pregnant females. Additional pregnancy related events reported in these cases (>50 occurrences) were coded to the PTs Abortion spontaneous (566), Labour pain (151), Vaginal haemorrhage (78), Heavy menstrual bleeding (50). Other frequently reported (>100 occurrences) clinical events were coded to the PTs Headache (410), Vaccination site pain (407), Fatigue (363), Pyrexia (206), Malaise (194), Myalgia (192), Nausea (178), Chills (156), Pain in extremity (135). The distribution of clinical events that were not pregnancy related (>100 occurrences) was similar in the pregnant mothers when compared with non-pregnant women of childbearing age.

- Three hundred twenty-two (322) baby/foetal cases, 283 serious and 39 non-serious. Cases are classified according to pregnancy outcome:
  - Pregnancy outcome: Live birth with congenital anomaly: Thirty-nine (39) of these cases reported 72 congenital anomalies that were coded to the PTs Foetal malformation (4), Atrial septal defect, Congenital anomaly, Ventricular septal defect (3 each), Congenital cystic lung, Congenital hydronephrosis, Congenital skin dimples, Exomphalos, Foetal cardiac disorder, Foetal chromosome abnormality, Foetal growth restriction, Kidney malformation, Pulmonary valve stenosis congenital (2 each), Anal atresia, Ankyloglossia congenital, Arnold-Chiari malformation, Cleft lip, Cleft palate, Cloacal exstrophy, Congenital amputation, Congenital foot malformation, Congenital haematological disorder, Congenital hand malformation, Congenital heart valve disorder, Congenital musculoskeletal disorder, Congenital musculoskeletal disorder of limbs, Congenital musculoskeletal disorder of spine, Congenital oral malformation, Cryptorchism, Double outlet right ventricle, Dysmorphism, Enlarged foetal cisterna magna, Fallot's tetralogy, Foetal arrhythmia, Foetal growth abnormality, Growth retardation, Heart disease congenital, Heart valve incompetence, Hepatic cytolysis, Hypospadias, Meningomyelocele, Neonatal deafness, Neonatal infection, Polydactyly, Pulmonary artery stenosis congenital, Pulmonary sequestration, Renal aplasia, Renal disorder, Renal dysplasia, Renal failure, Renal fusion anomaly, Renal hypertrophy,
Spina bifida, VACTERL syndrome (1 each). Of these 39 cases, information regarding trimester of exposure was available in 19 cases. Of these 19 cases, in 13 cases foetus was exposed during the 1st trimester, in 4 cases foetus was exposed during the 2nd trimester, and in 2 case exposure occurred during the 3rd trimester. Of these 39 cases, in 2 cases the mother of the baby was an asymptomatic gene carrier or had familial risk factors. In the remaining 37 cases, there was limited information regarding mother’s obstetric history, which precluded meaningful causality assessment.

- Pregnancy outcome: Spontaneous abortion: Thirty-seven (37) cases reported spontaneous abortion. Of these 37 cases, information regarding trimester of exposure was provided in 17 cases. Of these 17 cases, in 15 cases, foetus was exposed during the 1st trimester, in 2 cases foetus was exposed during the 2nd and the 3rd trimester each. The most frequently reported events (>1 occurrence) in these 37 cases other than exposure related events were coded to PTs Foetal growth restriction (18), Congenital anomaly (8), Foetal heart rate abnormal (3), Cytogenetic abnormality, Foetal vascular malperfusion (2 each). Of these 37 cases, in 4 cases mother had underlying medical history (i.e., spontaneous abortion, induced abortion and/or tobacco abuse), which might have contributed to the reported events. In the remaining 33 cases, there was limited information regarding obstetric history or co suspect medications of the mother, which precluded meaningful causality assessment.

- Pregnancy outcome: Elective termination: Twenty-three (23) cases reported elective termination of pregnancy. Of these 23 cases, 22 cases reported elective termination due to foetal defects and 1 case reported elective termination without foetal defects or unknown. Of these 23 cases, information regarding trimester of exposure was provided in 8 cases. Of these 8 cases, in 7 cases foetus was exposed during the 1st trimester, in 1 case, foetus was exposed during the 2nd trimester. The most frequently reported events (>1 occurrence) in these 23 cases other than exposure related events were coded to the PTs Heart disease congenital (4), Foetal malformation (3), Congenital central nervous system anomaly, Induced abortion (2 each). Of these 23 cases, in 5 cases mother had underlying medical history (i.e., spontaneous abortion, and/or gestational diabetes), which might have contributed to the reasons for elective termination of foetus. In the remaining 18 cases, there was limited information regarding obstetric history or co suspect medications of mother, which precluded meaningful assessment.

- Pregnancy outcome: Stillbirth: Twenty one (21) cases reported foetal death/neonatal death. Of these 21 cases, 15 cases reported stillbirth with foetal defects and remaining 6 cases reported stillbirth without foetal defect. Of these 21 cases, information regarding trimester of exposure was provided in 6 cases. Of these 6 cases, in 3 cases foetus was exposed during the 1st trimester, in the remaining 3 cases, foetus was exposed during the 2nd trimester. The most frequently reported events (>1 occurrence) in these 21 cases other than exposure related events were coded to the PTs Premature baby (7), Foetal hypokinesia (5), Foetal death, Foetal heart rate abnormal (4 each), Foetal growth restriction (3). Of these 21 cases, in 5 cases the mother had underlying medical history (i.e., spontaneous abortion, and/or obesity), which might have contributed to the reported event. In the remaining 16 cases, there was limited information regarding obstetric history or co suspect medications of mother, which precluded meaningful causality assessment.
Pregnancy outcome: Live birth without congenital anomaly: Two hundred two (202) cases reported live birth babies without congenital anomaly. Of these 202 cases, information regarding trimester of exposure was available in 58 cases. Of these 58 cases, in 26 cases, foetus was exposed during the 3rd trimester, in 20 cases foetus was exposed during the 2nd trimester, and in 12 cases exposure occurred during the 1st trimester. The frequently reported events (≥5 occurrence) in these 202 cases other than exposure related events were coded to Pts Premature baby (74), Foetal growth restriction (22), Foetal hypokinesia (12), Jaundice neonatal (9), Foetal heart rate abnormal, Congenital anomaly, Foetal distress syndrome (7 each), Immunisation (6), Neonatal respiratory distress syndrome, Breech presentation (5 each). Of these 202 cases, in 1 case reporting cerebral thrombosis and cerebral haemorrhage foetal the baby was delivered using vacuum extractor, which might have led to development of reported event. In the remaining 201 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

- Of the 3642 cases, 1898 cases provided pregnancy outcomes, which are provided in Table 69 (not reproduced here).

**Literature**

During the reporting period an article including new significant information regarding the use of BNT162b2 in pregnant/lactating women was identified: CItu IM, CItu C, Gorun F, et al. The Risk of Spontaneous Abortion Does Not Increase Following First Trimester mRNA COVID-19 Vaccination. J Clin Med. 2022; 11(6):1698. This article contributes to the growing evidence that risk of spontaneous abortion after COVID-19 vaccine immunisation during the first trimester of pregnancy is commensurate with the predicted risk in nonvaccinated pregnant women.

**MAH’s conclusion**

There were no safety signals regarding use in pregnant/lactating women that emerged from the review of these cases or the medical literature.

**Rapporteur assessment comment:**

**Clinical trial data**

Cumulatively, 58 cases (35%) out of 166 mother cases reported spontaneous abortion or abortion related events. In 25 cases the mother had a confounding medical history (spontaneous abortion, alcohol/tobacco use during pregnancy, ectopic pregnancy, obesity, diabetes mellitus, uterine disorder, depression or anembryonic gestation) and in 33 cases there was limited information which precluded meaningful assessment.

**Post-marketing data**

During the interval period, 566 cases (20%) out of 2,785 mother cases reported abortion spontaneous compared to 1040 cases (27%) out of 3,740 mother cases retrieved in the previous interval period.

Regarding pregnancy outcome, 37 cases (11%) out of 322 baby/foetal cases reported spontaneous abortion compared to 101 cases (23%) out of 443 baby/foetal cases retrieved in the previous interval period.

**Literature**

An article was published which showed that the risk of spontaneous abortion does not increase following first trimester mRNA COVID-19 vaccination (CItu et al.).
Overall, based on the information provided by the MAH in the current PSUR, it is agreed that no new safety concerns were identified for use in pregnant/lactating women. The Comirnaty product information reflects that Comirnaty can be used during pregnancy and breastfeeding.

**Use in Immunocompromised Patients**

Search criteria: Patients with Medical history of PTs included in Malignancy related conditions (SMQ Narrow); Malignancy related therapeutic and diagnostic procedures (SMQ Narrow); Malignant or unspecified tumours (SMQ Narrow); HLGT: Immunodeficiency syndromes (Primary Path); HLT: Retroviral infections (Primary Path); PTs: Allogenic bone marrow transplantation therapy; Allogenic stem cell transplantation; Autologous bone marrow transplantation therapy; Autologous haematopoietic stem cell transplant; Bone marrow transplant; Cord blood therapy; Heart transplant; Liver transplant; Lung transplant; Pancreas islet cell transplant; Renal transplant; Small intestine transplant; Stem cell transplant.

**Clinical trial data**

- Number of cases: 110 (BNT162b2 [90], blinded therapy [18], and BNT162B2S01, placebo [1 each]) (16.5% of 668 cases, the total CT dataset), compared to 110 cases (15.3%) retrieved in the PSUR #2.
- Most frequently reported clinical PTs (>2%): Condition aggravated (8), Atrial fibrillation (4), Cerebrovascular accident (4), Gastroenteritis (4), Osteoarthritis (4), Pneumonia (4), Acute kidney injury (3), Pertussis (3), Pyrexia (3).
- BNT162b2 related events coded to the PT: None of the events were assessed as related to BNT162b2 and/or blinded therapy by the Sponsor or investigator.

**Post-authorisation data**

- Number of cases: 8,815 (1.7% of 507,683 cases, the total PM dataset), compared to 14,657 cases (2.2%) retrieved in the PSUR #2.
- MC cases (3474), NMC cases (5341).
- Country of incidence: France (2200), UK (2070), Germany (1085), US (726), Italy (314), Sweden (312), Japan (212), Austria (192), Spain (158), Netherlands (156), Denmark (131), Belgium (119), Canada (116), Norway (112); the remaining 912 cases were distributed among 53 countries.
- Subjects' gender: female (5967), male (2628) and unknown (220).
- Subjects' age in years (n = 8073), range: 5 – 100, mean: 58.2, median: 60.0.
- Medical history (n = 8815). The most frequently (>200 occurrences) reported relevant medical conditions included Immunodeficiency (1647), Breast cancer (1121), Thyroidectomy (566), Neoplasm malignant (466), Hysterectomy (407), Chemotherapy (377), Prostate cancer (330), Radiotherapy (272), Chronic lymphocytic leukaemia (243), Neoplasm (239).
- COVID-19 Medical history (n = 689): COVID-19 (418), Suspected COVID-19 (249), COVID-19 pneumonia (22), Post-acute COVID-19 syndrome (15), SARS-CoV-2 test positive (4), Asymptomatic COVID-19, Exposure to SARS CoV 2 (3 each), Coronavirus infection, Coronavirus test positive (2 each).
Co-suspects (n = 608): The most frequently (≥10 cases) reported co-suspect vaccines/medications included COVID-19 vaccine NRV AD (113), COVID-19 vaccine (101), COVID-19 vaccine MRNA (95), Influenza vaccine (23), prednisone (20), mycophenolate mofetil (18), adalimumab (16), casirivimab/imdevimab, tacrolimus (13 each), Influenza vaccine Inact split 4V, JNJ 78436735, nivolumab, ocrelizumab (10 each).

- Number of events: 38,399.
- Event seriousness: serious (21,926), non-serious (16,507).
- Most frequently reported clinical PTs (≥3%): Immunisation (1248), Interchange of vaccine products (1223), Headache (1096), Fatigue (1030), Pyrexia (827), COVID-19 (740), Pain in extremity (686), Dyspnoea (605), Arthralgia (589), Myalgia (535), Dizziness (516), Pain (510), Nausea (488), Asthenia (478), Lymphadenopathy (456), Malaise (420), Chills (401), Chest pain (389), Vaccination site pain (374), Palpitations (326), Paraesthesia (313), Vomiting (292), Condition aggravated (254), Tachycardia (246).
- Time to event onset (n = 23,969 events), range: from <24 hours to ≤540 days, median: 1 day.
- Duration of event (n = 3184 of 6987 events with outcome of resolved/resolved with sequelae), range: <24 hours to 200 days, median: 3 days.
- Event outcome: fatal (1006), resolved/resolving (10,930), resolved with sequelae (821), not resolved (8997), unknown (16,862).

Analysis by age group

- Clinical trial data: Paediatric (16), Adults (39), and Elderly (55). A meaningful comparison between the different age groups is not possible due to the low number of cases.
- Post-marketing data: Paediatric (96), Adults (4828), Elderly (3198) and Unknown (693).
  - No significant difference was observed in the reporting proportion of frequently (≥3%) reported events between adult and elderly population except for the events coded to the PTs Headache, Lymphadenopathy, Palpitations and Tachycardia.
  - A higher reporting proportion of events coded to the PT Headache was observed in the adult population (16.6% [752 cases] in adults vs 8.2% [251 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Lymphadenopathy was observed in the adult population (7.4% [334 cases] in adults vs 2.6% [78 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Palpitations was observed in the adult population (5.2% [234 cases] in adults vs 2.1% [64 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Tachycardia was observed in the adult population (4.1% [185 cases] in adults vs 1.5% [46 cases] in elderly) compared to the elderly population.
  - No comparison was made to the paediatric population considering the limited number of cases.

MAH's conclusion

No new significant safety information was identified based on a review of these cases.

Rapporteur assessment comment:
No new important safety information could be identified in immunocompromised patients exposed to Comirnaty.

**Use in patients with autoimmune or inflammatory disorders**

Search criteria: Patients with Medical history PTs included in SMQ Immune-mediated/autoimmune disorders (Narrow and Broad scope); HLTs (Primary Path) Autoimmune disorders; Immune disorders NEC; HLT (Primary Path) Neuromuscular junction dysfunction.

**Clinical trial data**

- Number of cases: 102 (BNT162b2 [86], blinded therapy [14], and placebo [2]) (15.3% of 668 cases, the total CT dataset), compared to 101 cases (14.0%) retrieved in the PSUR #2.

- Of the 102 cases, the most frequently reported PTs (≥23%) included: Condition aggravated (6, 5.9%) and Atrial fibrillation (4, 3.9%).

- Event outcome: fatal (10), resolved/resolving (93), resolved with sequelae (3), and not resolved (24).

- In 6 cases (reporting 10 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Acute myeloid leukaemia, Adenocarcinoma pancreas, Cardiac failure congestive, Cardio-respiratory arrest, Completed suicide, Condition aggravated, COVID-19, Death, Pneumonia, and Sudden cardiac death (1 each). Of note, limited information regarding the cause of death was provided in 1 case (PT Death). Half (3 of 6 cases) of the fatal cases involved elderly subjects. The medical history reported included hypothyroidism, (3), colitis ulcerative, diabetes mellitus, narcolepsy, neuropathy peripheral (1 each).

- BNT162b2 related events coded to the PT Dehydration (1). Time to onset of event is 1 day and the event outcome is reported as resolved. None of the events were related to blinded therapy.

**Post-authorisation data**

- Number of cases: 21,000 (4.1% of 507,683, the total PM dataset), compared to 35,514 cases (5.4%) retrieved in the PSUR #2.

- MC cases (6424), NMC cases (14,576).

- Of the 21,000 cases, the most frequently reported clinical PTs (>3%) Included: Fatigue (3103, 14.8%), Headache (3082, 14.7%), Pyrexia (2207, 10.5%), Immunization (1750, 8.3%), Pain in extremity (1680, 8.0%), Arthralgia (1675, 8.0%), Interchange of vaccine products (1568, 7.5%), Myalgia (1535, 7.3%), Dizziness (1478, 7.0%), Dyspnoea (1385, 6.6%), Vaccination site pain (1360, 6.5%), COVID-19 (1344, 6.4%), Nausea (1308, 6.2%), Pain (1226, 5.8%), Malaise (1180, 5.6%), Chills (1174, 5.6%), Asthenia (1083, 5.2%), Chest pain (932, 4.4%), Paraesthesia (929, 4.4%), Lymphadenopathy (896, 4.3%), Condition aggravated (813, 3.9%), Palpitations (794, 3.8%), Tachycardia (646, 3.1%), and Hypoaesthesia (640, 3.1%).

- Event seriousness: serious (39,651), non-serious (43,889).

- Event outcome: fatal (1295), resolved/resolving (27,683), resolved with sequelae (2277), not resolved (25,409), unknown (27,206).

- In 409 cases (reporting 1295 relevant events with a fatal outcome), the reported causes of death (≥ 20 occurrences) were coded to the PTs Death (63), Immunisation (44), Cardiac arrest, COVID-19 (36 each), COVID-19 pneumonia (33), Dyspnoea (23), Cardio-respiratory
arrest (22), Interchange of vaccine products, Sudden death (21 each), and Cardiac failure (20). Of note, in 84 cases, limited information regarding the cause of death was provided (PTS Death and Sudden death). Immunisation and Interchange of vaccine products are discussed in the Section Off Label Use. Most (326 of 409 cases) of the fatal cases involved elderly subjects. The most frequently (>10 occurrences) reported medical history included diabetes mellitus (169), hypothyroidism (53), rheumatoid arthritis (36), type 1 diabetes mellitus (20), pulmonary fibrosis (15), rheumatic disorder (13), colitis ulcerative, psoriasis, and thyroid disorder (10 each).

- The most frequently reported clinical events observed in subjects with autoimmune or inflammatory disorders were consistent with those observed in the overall population.

**Exacerbation or flare-up**

- A focused analysis on exacerbation or flare of autoimmune or inflammatory disorders was conducted using PTs of interest (i.e., condition aggravated, disease progression), rather than all events.

- Of the 1117 cases that reported PTs indicative of exacerbation or flare, 345 cases were determined to be non-contributory and were not included in the discussion for the following reasons:
  
  - The events referred to an exacerbation/progression of an underlying condition that was not an autoimmune or inflammatory disorder (e.g., pain, arrhythmia, elevated blood pressure/hypertension, deep vein thrombosis, renal disease, migraine, fatigue/tiredness).

- Therefore, 772 cases are included in the analysis below.

- Clinical trial data
  
  - 1 case (BNT162b2) (0.1% of 668 cases, the total CT dataset), compared to 1 (0.1%) retrieved in the PSUR #2. The events were considered unrelated to BNT162b2.

- Post-authorisation data
  
  - Number of cases: 771 (0.2% of 507,683 cases, the total PM dataset), compared to 750 (0.1%) retrieved in the PSUR #2.
  
  - MC cases (274), NMC cases (497).
  
  - Country of Incidence: France (185), Germany (126), UK (118), Netherlands (54), Italy (51), US (35), Austria (23); the remaining 179 cases were distributed among 34 countries.
  
  - Subjects’ gender: female (584), male (180) and unknown (7).
  
  - Subjects’ age in years (n = 736), range: 9 – 90 years, mean: 50.7 years, median: 51 years.
  
  - Relevant medical history: the most frequently (>20 occurrences) reported medical conditions included: Autoimmune thyroiditis (79), Hypothyroidism (53), Rheumatoid arthritis (49), Psoriasis (34), Pericarditis (29), Colitis ulcerative, Diabetes mellitus, Multiple sclerosis (28 each), Autoimmune disorder, Basedow’s disease (27 each), Ankylosing spondylitis, Systemic lupus erythematosus (26 each), Immune thrombocytopenia (25), Sjogren’s syndrome (22), Crohn’s disease (21), Arthritis, and Psoriatic arthropathy (20 each).
COVID-19 Medical history (n = 61): COVID-19 (43), Suspected COVID-19 (20), Post-acute COVID-19 syndrome (5), and SARS-CoV-2 test positive (1).

- Co-suspect vaccines/medications: Influenza vaccine (5), COVID-19 Vaccine MRNA (MRNA 1273) (3), Adalimumab, COVID-19 Vaccine NRVV AD (CHADEX1 NCOV-19) (2 each), acyclovir, colchicine, Hepatitis B vaccine, hydroxychloroquine, ocrelizumab, and pneumococcal vaccine polysacchar 23V (1 each).

- Number of events: 4633 (of which 782 were events of interest i.e., exacerbation/flare AE).

- Relevant event seriousness: serious (521), non-serious (266).

- Most frequently reported relevant PTs (≥2%): Condition aggravated (548), Disease recurrence (200), and Concomitant disease aggravated (22).

- Time to event onset (n = 424), range: from 1 day to 164 days, median: 4 days.

- Duration of relevant events (n = 41 out of 112 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 160 days, median 17 days.

- Relevant event outcome: fatal (4), resolved/resolving (224), resolved with sequelae (18), not resolved (332), unknown (208).

- In 4 cases (reporting 4 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Disease recurrence (3), and Condition aggravated (1). Three of the 4 cases involved elderly subjects. The medical history reported included arthritis, autoimmune hepatitis, Miller Fisher syndrome, and thrombotic thrombocytopenic purpura.

- Analysis by age group

  - Clinical trial: Paediatric (1).

  - Post-marketing: Paediatric (19), Adults (572), Elderly (155) and Unknown (25).
    - Exacerbation and/or flare of underlying autoimmune or inflammatory disorders occurred more frequently in the adult population, which is likely due to autoimmune disorders being more common in adults and the fact that adults are the largest group of vaccinated individuals reporting adverse events.

MAH's conclusion

Overall, there were 772 cases (1 CT case and 771 PM cases [0.2% of the overall dataset]) that reported exacerbation/flare in subjects with autoimmune or inflammatory disorders following administration of BNT162b2. Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders occurred more frequently in the adult population which is likely due to the fact that the largest group of vaccinated individuals reporting AEs are in this age group. Also, the autoimmune or inflammatory disorders often occur during adulthood. The most frequently reported clinical events observed in subjects with autoimmune or inflammatory disorders were consistent with those observed in the overall population.

Rapporteur assessment comment:

No new important safety information could be identified in patients with autoimmune or inflammatory disorders.
Use in frail patients with comorbidities (e.g. COPD, Diabetes, Chronic Neurological Disease, Cardiovascular Disorders, active Tuberculosis)

Search criteria: Patients with Medical history of PTs included in HLGTs (Primary Path) Bronchial disorders (excl neoplasms), Heart failures, Mental impairment disorders, Movement disorders (incl parkinsonism), Nephropathies, Pulmonary vascular disorders, Renal disorders (excl nephropathies); HLTS (Primary Path) Autonomic nervous system disorders, Diabetes mellitus (incl subtypes), Hepatic failure and associated disorders, Hepatic fibrosis and cirrhosis, Motor neurone diseases, Muscle tone abnormal, Neuromuscular disorders NEC, Pulmonary hypertensions, Respiratory failures (excl. neonatal); patients with Medical history of ongoing Tuberculosis.

Clinical trial data

- Number of cases: 153 (BNT162b2 [125], blinded therapy [25], and placebo [3]) (22.9% of 668 cases, the total CT dataset), compared to 176 cases (24.4%) retrieved in the PSUR #2.
- Country of incidence: US (123), Argentina (11), Germany (9), Brazil (3), China, Spain (2 each); the remaining 3 cases were distributed among 3 countries.
- Subjects' gender: female (56), male (97).
- Subjects' age in years (n = 153), range: 0.83 – 87 years, mean: 59.6 years, median: 64 years.
- Medical history (n = 153): the most frequently (≥5 occurrences) reported relevant medical conditions included Type 2 diabetes mellitus (71), Asthma (33), Chronic obstructive pulmonary disease (23), Diabetes mellitus (14), Cardiac failure congestive, Chronic kidney disease (10 each), Pulmonary embolism (7), and Bronchitis chronic (5).
- Co-suspects (n = 33 cases): The reported co-suspect agents included metformin (2), amiodarone, amiodipine, aripiprazole, atenolol, diltiazem, duloxetine, furosemide, hydrochlorothiazide/triamterene, hydroxyzine, losartan, semaglutide, tamsulosin, warfarin (1 each).
- Number of events: 187.
- Most frequently reported clinical PTs (>2%): Condition aggravated, Pneumonia (6 each), Cerebrovascular accident, Dyspnoea (5 each), and Coronary artery disease (4).
- BNT162b2 related events were coded to the PT: Dehydration (1). Time to onset of event is 1 day and the event outcome is reported as resolved. None of the events were related to blinded therapy.
- Time to event onset: (n = 131), range: from 1 day to 178 days, median: 106 days.
- Duration of relevant events (n = 78 out of 103 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 78 days, median 5 days..
- Reported event outcome: fatal (13), resolved/resolving (128), resolved with sequelae (10), not resolved (36), and unknown (0).
- In 9 cases (reporting 13 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Death (2), Adenocarcinoma pancreas, Cardiac failure congestive, Cardio-respiratory arrest, Completed suicide, Condition aggravated, COVID-19, Drowning, Pneumonia, Pulmonary embolism, Respiratory failure, and Sudden cardiac death (1 each). Of
note, in 2 cases, limited information regarding the cause of death was provided (PT Death).
Most (5 of 9 cases) of the fatal cases involved elderly subjects. The most frequently (>1 occurrence) reported medical histories included type 2 diabetes mellitus (6) and Asthma (2).

Post-authorisation data

- **Number of cases:** 18,276 (3.6% of 507,663, the total PM dataset), compared to 33,889 cases (5.2%) retrieved in the PSUR #2.
- **MC cases (6964), NMC cases (11,312).**
- **Country of incidence:** France (3532), Germany (3124), UK (2189), US (1520), Sweden (1062), Japan (765), Italy (616), Austria (471), Norway (448), Spain (442), Denmark (408), Netherlands (396), Finland (305), Canada (260), Belgium (240), Czech Republic (234), Estonia (222), Iraq (220), Ireland (196), Greece (164), Taiwan, province of China (144), Portugal (143), Switzerland (136), Poland (102); the remaining 937 cases were distributed among 54 countries.
- **Subject’s gender:** female (11,576), male (6436), and unknown (264).
- **Subject’s age in years (n = 17,342), range: 3 - 107 years, mean: 54.1 years, median: 55 years.
- **Medical history (n = 18,276):** the most frequently (>75 occurrences) reported relevant medical conditions included Asthma (7896), Diabetes mellitus (3121), Type 2 diabetes mellitus (2001), Chronic obstructive pulmonary disease (1201), Type 1 diabetes mellitus (649), Cardiac failure (616), Chronic kidney disease (608), Pulmonary embolism (564), Renal failure (343), Parkinson’s disease (247), Dementia (242), Hypokinesia (168), Cognitive disorder (166), Dementia Alzheimer’s type (146), Bronchitis chronic (133), Renal disorder (117), Bronchiectasis (107), Asthma exercise Induced (100), Cardiac failure chronic (81), Cardiac failure congestive (77), Bronchospasm, IgA nephropathy (76 each), and Hepatic cirrhosis (75).
- **COVID-19 Medical history (n = 1226):** COVID-19 (912), Suspected COVID-19 (268), COVID-19 pneumonia (38), Post-acute COVID-19 syndrome (36), SARS-CoV-2 test positive (13), Coronavirus infection (8), Asymptomatic COVID-19 (5), and Exposure to SARS-CoV-2 (1).
- **Co-suspects (n = 929 cases):** The most frequently (>5 occurrences) reported co-suspect vaccines/medications included COVID-19 vaccine (250), COVID-19 vaccine MRNA (1273) (141), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (129), Influenza vaccine (58), Influenza vaccine inact split 4V (24), ocrelizumab, prednisone (19 each), JNJ 78436735, mycophenolate mofetil (15 each), apixaban (14), Influenza vaccine inact SAG 4V (13), tacrolimus (12), adalimumab (11), rituximab (9), prednisolone (8), atorvastatin, levethroxine, methotrexate (7 each), allopurinol, clopidogrel, influenza vaccine inact SAG 3V, and pregabalin (6 each).
- **Number of events:** 70,918
- **Relevant event seriousness:** serious (34,905), non-serious (36,098).
- **Most frequently reported (≥3%) clinical PTs:** Headache (2624, 15.0%), Fatigue (2570, 14.6%), Pyrexia (2012, 11.5%), Dyspnoea (1797, 10.2%), Immunisation (1533, 8.7%), COVID-19 (1446, 8.2%), Interchange of vaccine products (1383, 7.9%), Pain in extremity (1366, 7.8%), Dizziness (1255, 7.2%), Myalgia (1212, 6.9%), Arthralgia (1179, 6.7%), Vaccination site pain (1173, 6.7%), Nausea (1146, 6.5%), Malaise (1073, 6.1%), Asthenia (970, 5.5%), Chills (925, 5.3%), Pain (907, 5.2%), Chest pain (826, 4.7%), Palpitations (668, 3.8%).
Lymphadenopathy (614, 3.5%), Paraesthesia (602, 3.4%), Cough (585, 3.3%), and Vomiting (561, 3.2%).

- Time to event onset (n = 46,814), range: from 1 day to 180 days, median: 2 days.
- Duration of relevant events (n = 8391 out of 16,690 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 181 days, median 3 days.
- Relevant event outcome: fatal (2258), resolved/resolving (24,735), resolved with sequelae (1867), not resolved (19,410), unknown (23,001).
- In 801 cases (reporting 2258 relevant events with a fatal outcome), the reported cause of death (226 occurrences) was coded to the PTs Death (144), Immunisation (92), COVID-19 (91), COVID-19 pneumonia (80), Cardiac arrest (62), Cardiac failure, Dyspnoea (50 each), Interchange of vaccine products (49), Sudden death (42), Cardio-respiratory arrest (40), Pulmonary embolism (38), Pneumonia (34), Respiratory failure (29), Pyrexia (28), and Myocardial infarction (26). Of note, in 186 cases, limited information regarding the cause of death was provided (PTs Death and Sudden death). Most (689 of 801 cases) of the fatal cases involved elderly subjects. The most frequently (≥20 occurrences) reported medical history included diabetes mellitus (169), type 2 diabetes mellitus (117), cardiac failure (113), chronic obstructive pulmonary disease (95), dementia (83), chronic kidney disease (72), asthma (55), cognitive disorder, pulmonary embolism (39 each), renal failure (38), Parkinson's disease (35), dementia Alzheimer's type (31), Cardiac failure chronic (27), and type 1 diabetes mellitus (20).

Analysis by age group

- Clinical trial: Paediatric (12), Adults (67), Elderly (74). A meaningful comparison between the different age groups is not possible due to the low number of cases.
- Post-marketing: Paediatric (625), Adults (11,157), Elderly (5906) and Unknown (588).
  - No significant difference was observed in the reporting proportion of frequently (≥3%) reported events between adult and elderly population except for the event coded to PT Lymphadenopathy.
  - A higher reporting proportion of events coded to PT Lymphadenopathy was observed in the adult population (4.7% [520 cases] in adults vs 1.0% [59 cases] in elderly) compared to the elderly population.
  - No comparison was made to the paediatric population considering the limited number of cases.

MAH’s conclusion

The reporting proportion of not resolved cases (36.1%) and cases resolved with sequelae (3.1%) in frail subjects is similar to the reporting proportion observed in the overall population (31.7% for outcome of not resolved, 1.9% for outcome of resolved with sequelae). The reporting proportion of cases reporting fatal outcome (4.4%) in frail subjects is higher than the reporting proportion of cases reporting fatal outcome in the overall population (0.6%). This is expected, considering that most of the cases reporting a fatal outcome (64.4%) among the frail subjects involved subjects over 75 years of age who, due to their advanced age and underlying comorbidities, are more likely to die than younger individuals. Underlying comorbidities are likely to be contributory to their deaths.

The vaccine has been studied in individuals with stable chronic diseases (e.g., hypertension, obesity). It has not been systematically studied in frail individuals with severe comorbidities but there is much
post-authorisation data in this population as they have generally been targeted as high priority for vaccination. No safety signals have emerged that would be considered specific to this population.

**Rapporteur assessment comment:**

No important new safety information could be identified regarding use in frail patients with co-morbidities. For future PSURs in the section ‘Update on special patient populations’, the use in frail patients with co-morbidities should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

**Interactions with other vaccines**

Search criteria: **HLT Interactions**.

**Clinical trial data**

- No relevant serious clinical trial cases reported during the reporting period, as in the PSUR #2.

**Post-authorisation data**

- Number of cases: 3 (0.0006% of 507,683 cases, the total PM dataset), compared to 18 (0.003%) retrieved in the PSUR #2.

**MAH’s conclusion**

Among the overall 146 cases, 143 were considered not relevant, as a drug interaction did not occur in 1 case, the interacting agents was not specified in 32 cases, BNT162b2 was not involved in 1 case and in the remaining 110 cases, the interaction occurred with alcohol, herbal or medications rather than another vaccine.

There were 3 cases in the overall post-marketing dataset that involved a vaccine interaction. The most frequently co-reported event (>2 occurrences) other than off label use and interchange of vaccines PTs was Pyrexia, which is consistent with the known reactogenicity of the vaccine and listed in Section 4.8 Undesirable effects of the CDS. There is no indication of a safety signal noted based on the review of these cases.

**Rapporteur assessment comment:**

No important new safety information could be identified regarding interactions with other vaccines. For future PSURs in the section ‘Update on special patient populations’, the interactions with other vaccines should only be included and discussed in the PSUR if the reporting pattern changes and/or there is a safety issue/signal. **Request for next PSUR**

**2.4. Characterisation of risks**

As reported in Section 16.1 Summary of Safety Concerns of the PSUR, on 08 July 2022 (after the DLP of this PSUR), the MAH submitted the EU RMP version 5.1 to remove the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP recommendation received on 10 March 2022 with the positive opinion for the Type II Variation 87 (EMEA/H/C/005735/II/0087) and based on the accumulation of post-authorisation safety information.

In line with this update to the EU-RMP, the MAH proposes to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period, because anaphylaxis is
a known risk of vaccines that is understood by HCPs who administer vaccines and patients and does not considerably impact the benefit/risk profile of the vaccine. Product labelling and standards of medical care during the vaccination procedure provide adequate risk mitigation.

**Rapporteur assessment comment:**

Please refer regarding the important identified risk Anaphylaxis to 2.3. ‘Evaluation of risks and new information’, section ‘Evaluation of important identified risks’ of this AR.

### 2.4.1. Characterisation of important identified and potential risks

- Important Identified Risk: Anaphylaxis
- Important Identified Risk: Myocarditis and Pericarditis
- Important Potential Risk: Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)

**Rapporteur assessment comment:**

Please refer regarding the important identified risks - Anaphylaxis – and - Myocarditis and Pericarditis - to 2.3. ‘Evaluation of risks and new information’, section ‘Evaluation of important identified risks’ of this AR.

Please refer regarding the important potential risk - Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD) – to 2.3. ‘Evaluation of risks and new information’, section ‘Evaluation of important potential risks’ of this AR.

### 2.4.2. Description of missing information

Missing information:

- Use in pregnancy and while breast feeding

**Rapporteur assessment comment:**

Please refer regarding pregnancy and lactation to 2.3. ‘Evaluation of risks and new information’, section ‘Use in pregnant/lactating women’ of this AR. No important new safety information could be identified.

- Use in immunocompromised patients

**Rapporteur assessment comment:**

Please refer regarding immunocompromised patients to 2.3. ‘Evaluation of risks and new information’, section ‘Use in immunocompromised patients’ of this AR. No important new safety information could be identified.

- Use in frail patients with co-morbidities (e.g. chronic obstructive pulmonary disease (COPD), diabetes, chronic neurological disease, cardiovascular disorders)

**Rapporteur assessment comment:**

Please refer regarding frail patients with co-morbidities to 2.3. ‘Evaluation of risks and new information’, section ‘Use in frail patients with co-morbidities’ of this AR. No important new safety information could be identified.
• Use in patients with autoimmune or inflammatory disorders

Rapporteur assessment comment:
Please refer regarding patients with autoimmune or inflammatory disorders to 2.3. ‘Evaluation of risks and new information’, section ‘Use in patients with autoimmune or inflammatory disorders’ of this AR. No important new safety information could be identified.

• Interaction with other vaccines

Rapporteur assessment comment:
Please refer regarding interaction with other vaccines to 2.3. ‘Evaluation of risks and new information’, section ‘Interaction with other vaccines’ of this AR. No important new safety information could be identified.

• Long term safety data

At the time of the initial submission, 2-month post dose 2 safety data were available for approximately half of the patients who have received BNT162b2 mRNA vaccine in Study C4591001. The pivotal clinical study is ongoing and ongoing non-interventional safety studies will collect longer term post-marketing safety data.

Rapporteur assessment comment:
The information regarding long-term safety data is noted.

3. Benefit evaluation

BNT162b2 is indicated for active immunisation to prevent COVID-19 caused by SARS CoV-2 virus in individuals 5 years of age and older.

Rapporteur assessment comment:
There are no new data on efficacy that alters previous assessments, and which are described in the approved product information of Comirnaty.

Of note, after the DLP of this PSUR:
Comirnaty is currently also available as two adapted vaccines (only to be used in people aged 12 years and older who have received at least a primary vaccination course against COVID-19):
- Comirnaty Original/Omicron BA.1 contains tozinameran and riltozinameran, another mRNA molecule with instructions for producing a protein from the Omicron BA.1 subvariant of SARS-CoV-2. (procedure EMEA/H/C/005735/II/0140)
- Comirnaty Original/Omicron BA.4-5 contains tozinameran and famtozinameran, another mRNA molecule with instructions for producing a protein from the Omicron BA.4 and BA.5 subvariants of SARS-CoV-2. (procedure EMEA/H/C/005735/II/0143)

The Comirnaty indication was extended to children 6 months - 4 years old (Tris/Sucrose presentation 3 micrograms/dose). (procedure EMEA/H/C/005735/X/0138)

An EU procedure is ongoing concerning the extension application to add a new strength of 5/5 μg (tozinameran, famtozinameran) for children between 5 to 11 years of age. (procedure
4. Benefit-risk balance

During the reporting period of the PSUR, the posology recommendations for the booster use was amended from "individuals 18 years of age and older" to "individuals 12 years of age and older", provided further details on heterologous boosting and the boosting interval was shortened to at least 3 months after completion of the primary series (EMEA/H/C/005735/II/0093, EMEA/H/C/005735/II/0104 and EMEA/H/C/005735/II/0111).

There are safety issues identified, which include that dizziness should be added as an ADR to section 4.8 with frequency unknown in the Comirnaty product information. However, as the MAH already submitted a variation to amend the Comirnaty product Information accordingly (procedure EMEA/H/C/005735/II/0152), this PSUSA procedure can be concluded with maintenance of the marketing authorisation(s).

The risks have been evaluated in the context of the benefits of the vaccine. No additional changes to the Comirnaty risk minimisation measures are warranted.

Based on the PRAC Rapporteur review of the available safety and efficacy/effectiveness data from the current reporting period for the Comirnaty PSUR, the benefit-risk balance of Comirnaty (tozinameran) remains unchanged.

The MAH should continue to review the safety of Comirnaty, including all reports of adverse events and should propose an update of the product information if an evaluation of the safety data identifies important new safety information, as applicable.

There is no need for changes to the frequency of PSUR submission for Comirnaty.

5. Rapporteur request for supplementary information

1. Regarding multiple repeated booster doses, during the reporting period the number of persons receiving multiple booster doses (i.e., homologous, heterologous, different strains) is increasing whereas the impact on safety and efficacy remains uncertain. In addition the impact (including long-term) of repeatedly (e.g., yearly) receiving booster doses (with or without strain updates) also remains unknown. The MAH is requested to discuss whether ‘safety following multiple repeated booster doses (i.e., homologous, heterologous, different strains)’ should be considered as Missing information in the RMP, and proposals should be provided how to address this knowledge gap in ongoing or newly proposed PASSs, as applicable.

2. Regarding multisystem inflammatory syndrome in children and in adults (MIS-C/ -A):
   a. During the reporting period, the MAH identified a total of 130 potential new MIS-A cases. Of these, 1 was classified as BC level 1 and considered probably related with Comirnaty. However, given the extensive exposure of Comirnaty in adults this is not considered unexpected and does not present a new safety concern. Another BC level 1 MIS-A (AER number [redacted]) reported by the MAH in the 14th SSR (interval period 16 Feb 2022-15 Apr 2022) was considered confounded by previous COVID-19 infection and therefore unlikely to be associated with Comirnaty. However, this case seems not to be retrieved in current safety database search. The MAH is requested to
explain why the BC level 1 MIS-A case (AER number [redacted]) from the 14th SSR was not included in current MIS-A overview (interval period 19 Dec 2021 to 18 Jun 2021).

b. During the interval period, the MAH reported post-marketing 207 relevant MIS-C/-A cases. However, in Appendix 6A.4 of the PSUR the MAH reported 199 MIS-C/-A cases during the interval period. The MAH is requested to explain this discrepancy in the numbers of retrieved MIS-C/-A cases.

3. Regarding myocarditis,

a. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 5-11 years, aged 12-15, aged 18-24, aged 25-29, aged 30-39, and aged ≥40 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

b. Upon analysis of the requested details of the fatal myocarditis cases the MAH is expected to discuss whether the current risk minimisation measures is still adequate.

4. Regarding pericarditis:

a. The MAH is requested to provide detailed information concerning the fatal cases with pericarditis in persons aged 18-24 years, 25-29 years, and ≥40 years and perform per case an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

b. Upon analysis of the requested details of the fatal pericarditis cases the MAH is expected to discuss whether the current risk minimisation measures is still adequate.

5. Regarding glomerulonephritis, the MAH is requested to provide detailed information concerning the 46 cases with glomerulonephritis in the interval period and perform an WHO-UMC causality assessment per case, and to provide an updated review with DLP 14 Nov 2022 of cases reporting glomerulonephritis from their safety database including a WHO-UMC causality assessment per case regarding Comirnaty exposure.

6. Regarding fatal cases reported in paediatric persons, the MAH is requested to provide detailed information concerning the fatal cases in persons aged 5-11 years and 12-17 years, perform per case a WHO causality assessment, and an assessment according to Brighton Collaboration case definition and Level of certainty classification, if applicable.
6. MAH responses to request for supplementary information

1. Regarding **multiple repeated booster doses**, during the reporting period the number of persons receiving multiple booster doses (i.e., homologous, heterologous, different strains) is increasing whereas the impact on safety and efficacy remains uncertain. In addition the impact (including long-term) of repeatedly (e.g., yearly) receiving booster doses (with or without strain updates) also remains unknown. The MAH is requested to discuss whether 'safety following multiple repeated booster doses (i.e., homologous, heterologous, different strains)' should be considered as Missing information in the RMP, and proposals should be provided how to address this knowledge gap in ongoing or newly proposed PASSs, as applicable.

**MAH response**

For the monovalent and bivalent booster doses, the safety of the vaccine in persons receiving multiple booster shots (including homologous, heterologous, and different strains) is being monitored across the MAH’s portfolio of ongoing and planned PASS studies (Table 1).

<table>
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<th>PASS Study</th>
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All studies include a subset of patients with the MAH’s primary series (2 or 3 doses, as applicable per age population). Some studies also include subsets of heterologous primary series. An overview of the booster combination doses collected in each study is summarised in Table 2.

<table>
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* All studies include a subset of patients with a Pfizer primary series (2 or 3 doses, as applicable). Some studies also include subsets of heterologous primary series.

For future booster doses, the safety surveillance approach will depend on the outcome of current and future discussions with the EMA and the FDA surrounding the future of the COVID-19 vaccine. The MAH will assess the feasibility and the suitability of implementing an enhanced active safety surveillance when a new booster dose becomes available. For example, once a new booster dose becomes available, and depending on the level of completeness of ongoing studies, vaccinees from at
least 1 country could be enrolled and followed for potential safety events. Initial findings from this new active safety surveillance could be available shortly after launch of the new booster dose. This framework could be established then repeated for each subsequent booster dose. Therefore, studies of subsequent booster doses could enrol new patients but apply the same enhanced active surveillance framework for consistent safety surveillance.

The MAH does not consider that “safety following multiple repeated booster doses (i.e., homologous, heterologous, different strains)” should be considered missing information in the EU-RMP. For the purpose of the RMP, missing information refers to gaps in knowledge about the safety of the product within the approved indication. Regulatory authorities in various countries, including in the EU and US, have included in COMIRNATY product labelling, implicit and explicit information regarding the use of COMIRNATY in homologous and heterologous dosing scenarios. As such, there has been widespread use of COMIRNATY in primary and boosting mixed use scenarios without new significant safety information emerging from the experience, particularly as followed in the medical literature. In addition, routine pharmacovigilance has not uncovered new safety concerns relating to the use of different COVID-19 strain vaccines, namely bivalent vaccines against original and OMEICRON BA.1 and BA.4/BA.5 strains of SARS-CoV-2. Routine pharmacovigilance will continue, and any new significant safety information will be reported appropriately.

**Rapporteur assessment comment:**

At the moment we agree that there is no gap in knowledge about the safety concerning the primary and boosting mixed use scenarios of Comirnaty since there is no new significant information that Comirnaty would be associated with a different safety profile when administered in such situations. Therefore, MAH’s conclusion is endorsed that ‘safety following multiple repeated booster doses (i.e., homologous, heterologous, different strains)” should not be considered missing information in the Comirnaty RMP.

However, after the DLP of the 3rd PSUR, Comirnaty has been variant updated (bivalent vaccines), approved and is currently being used in the EU. In light of this, in the next PSUR, the MAH is requested to ensure a transparent presentation of cases reported from the newly approved variant vaccines, as well as a continuous comparison of the safety issues identified with the evidence available for the original vaccine. Any differences in occurrence of safety issues, when comparing the variant updated vaccines with the original monovalent Comirnaty or, indeed, when comparing the two different variant updated bivalent Comirnaty vaccines, should be discussed. **Request for next PSUR**

**Issue solved**

2. **Regarding multisystem inflammatory syndrome in children and in adults (MIS-C/ -A):**

   a. *During the reporting period, the MAH identified a total of 130 potential new MIS-A cases. Of these, 1 was classified as BC level 1 and considered probably related with Comirnaty. However, given the extensive exposure of Comirnaty in adults this is not considered unexpected and does not present a new safety concern. Another BC level 1 MIS-A (AER number [redacted]) reported by the MAH in the 14th SSR (interval period 16 Feb 2022-15 Apr 2022) was considered confounded by previous COVID-19 infection and therefore unlikely to be associated with Comirnaty. However, this case seems not to be retrieved in current safety database search. The MAH is requested to explain why the BC level 1 MIS-A case (AER number [redacted] from the 14th SSR was not included in current MIS-A overview (interval period 19 Dec 2021 to 18 Jun 2021).*
MAH response

AER numbers [redacted] and [redacted] are individual case reports which were identified from a single literature article and entered into the MAH safety database. One case (AER# [redacted]) described a 48-year-old female patient who developed multisystem inflammation following receipt of the Moderna COVID-19 vaccine. This case was made invalid during the PSUR reporting period because the suspect product was not BNT162b2. The second case (AER# [redacted]) describes a 51-year-old male patient which was reported by the MAH in the 14th SSR; in error this case was presented with the other AER number from the 2 literature cases [redacted]. This error has been corrected in the dataset and the BC level 1 case of the 51-year-old male who received BNT162b2 (AER# [redacted]) is presented below. This case is significantly confounded by a recent COVID-19 infection. As COVID-19 infection within the last 12 weeks is the current known aetiology of MIS-A, this is the most likely cause in this case and the case does not change the MAH's overall assessment of the potential association between BNT162b2 and MIS-A/C.

AER # [redacted] 51-year-old male, reported to be previously healthy.


On admission he was tachycardic (130 bpm), hypotensive (90/60 mmHg), leucocytosis 19.4 x 103/μL (92% neutrophils). Anaemia (Hb 11 g/dL), thrombocytopenic (72000/μL) and had elevated CRP (334 mg/L), brain natriuretic protein (17768 μg/ml) and troponin (0.248 μg/l). Antibody testing confirmed previous COVID-19 infection. PCR testing for SARS-CoV-2 and enteric pathogens was negative. Imaging of the chest and abdomen was initially normal.

Despite fluids he required vasopressors and overt pulmonary oedema developed. Echocardiography confirmed biventricular dilatation with EF 20%.

After empiric MIS-A treatment with steroids and 1 dose of intravenous immunoglobulins (IV Ig) the symptoms, haemodynamics and inflammatory markers rapidly improved. EF was normal (60%) on 14 and 28 June 2021 whilst patient was on prednisolone. On steroids he experienced superficial desquamation of palms of hands and soles of feet and 2 episodes of mild conjunctivitis. He remained fully recovered as of February 2022 and received no further vaccination.

References


Rapporteur assessment comment:

The BC level 1 MIS-A case (AER number [redacted]) from the 14th SSR was not included in current MIS-A overview (interval period 19 Dec 2021 to 18 Jun 2021) because in error this case was presented with the wrong AER number [redacted]. This error has been corrected in the dataset and the BC level 1 case of the 51-year-old male who received Comirnaty (AER number [redacted]) is now also included. Because this case [redacted] is confounded by a recent COVID-19 infection which could be the cause of the MIS-A. Therefore, the case is considered unlikely related to Comirnaty exposure.

In conclusion, there were 2 BC level 1 MIS-A cases with AER numbers [redacted] and [redacted] a 50-year-old female (case reported in the literature from [redacted]; not considered unexpected and does not present a new safety concern) and a 51-year-old male (case reported in the
Among the literature from [ ], considered not likely related to Comirnaty exposure), respectively.

No new important safety concern could be identified for MIS-A.

**Issue solved**

*b. During the interval period, the MAH reported post-marketing 207 relevant MIS-C/ -A cases. However, in Appendix 6A.4 of the PSUR the MAH reported 199 MIS-C/ -A cases during the interval period. The MAH is requested to explain this discrepancy in the numbers of retrieved MIS-C/ -A cases.*

**MAH response**

Eight cases were included in PSUR #3 section 16.3.3.1.9 which were erroneously not presented in Appendix 6.4, as the cases had previously been presented by the MAH in SBSR #2 (reporting period 16 December 2021 through 15 February 2022) (AER #s: [redacted]) and SBSR #3 (reporting period 16 February through 15 April 2022) (AER #s: [redacted]).

Three cases were classified as BC level 4, and 4 cases as BC level 5 (all adult patients). One case was classified as a BC level 2 (probable) case of MIS-C. The clinical details are presented below and were reported in SBSR #2.

**15-year-old male from [ ].**

Reported preferred terms: drug ineffective, covid-19, myocarditis, pericarditis, renal impairment, shock, multisystem inflammatory syndrome in children, blood creatinine increased, oropharyngeal pain, abdominal pain, left ventricular dysfunction, pleural effusion, fatigue, pyrexia, sinus tachycardia.

The patient was reported to have no significant medical history and described as "previously fit and well". The patient's concomitant medications were not reported.

Dose 1; 20 October 2021. The patient had a history of COVID-19 infection (positive PCR) with mild symptoms 3 weeks prior to hospitalisation (given the reported information the positive PCR would have been approximately 2-3 weeks after dose 1). The patient presented to Accident and Emergency with a sore throat, generalised abdominal pain and in shock. Echocardiogram showed severe LV dysfunction and the patient required inotropes, intubation and ventilation. He was treated with IV antibiotics and high-dose steroids. Suspected paediatric inflammatory multisystem syndrome "associated with COVID-19 or vaccine (or combination of both)".

Significant renal impairment, high CRP (547), high ferritin (6000 µg/L), high D-dimers, creatinine kinase, creatinine and lactate dehydrogenase (values not reported). Echocardiogram (09 Dec 2021) showed severe LV dysfunction (EF 45%). Chest x-ray (06 Dec 2021) showed pleural effusions and possible fluid on the right side. Electrocardiogram (06 Dec 2021): sinus tachycardia, no ST changes. SARS-CoV-2 test (unspecified date) positive; troponin (normal high range 34): 08 Dec 2021 437 ng/L. Patient remained an inpatient at the time of the report and was "for IV Ig".

The patient was less than 12 weeks from COVID-19 infection and there is report of a positive SARS-CoV-2 test at the time of hospitalisation. Given COVID-19 is the current known aetiology of MIS-C this is the most likely cause in this case.

The additional 8 cases do not change the overall assessment of MIS-C/A.

**Rapporteur assessment comment:**
The MAH explained the discrepancy in the number of retrieved MIS-C/-A cases (199 versus 207 cases). The missing 8 cases were previously reported in the 13th and 14th SSRs, and should have been included in Appendix 6A.4 of the PSUR. No new important safety concern could be identified for MIS-C/-A.

**Issue solved**

3. **Regarding myocarditis,**

   a. The MAH is requested to provide detailed information concerning the fatal cases with myocarditis in persons aged 5-11 years, aged 12-15, aged 18-24, aged 25-29, aged 30-39, and aged ≥40 years and perform per case an assessment according to Brighton Collaboration Myocarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

   b. Upon analysis of the requested details of the fatal myocarditis cases the MAH is expected to discuss whether the current risk minimisation measures is still adequate.

**MAH response**

The PBRER described a total of 87 interval myocarditis events reporting a fatal outcome (86 unique cases). After the DLP of the PBRER, 2 of those cases were updated according to follow up data and no longer include a myocarditis event. The remaining 84 cases are analysed in Appendix 3 (see below in comment box) and a case narrative listing is provided in Appendix 5 (not reproduced here). Where cases reporting fatal myocarditis also co-reported a pericarditis event, the pericarditis event is also included in the analysis.

Overall, no new safety information was identified from the analysis of these cases and thus, no changes to the risk management measures are warranted.

**Rapporteur assessment comment:**

Please refer regarding the assessment of the 84 fatal cases to the PRAC Rapporteur's comments in:

[Appendix 3.pdf]

In persons aged 5-11 years, there were 2 fatal cases with myocarditis of which one case is considered BC level 1 and unlikely related to Comirnaty exposure and the other case considered BC level 3 and unassessable.

In persons aged 12-15 years, there were 3 fatal cases with myocarditis of which 1 case is considered BC level 3 and unlikely related to Comirnaty exposure and the other 2 cases considered BC level 4-5.

In persons aged 18-24 years, there were 4 fatal cases with myocarditis of which 2 cases are considered BC level 1 of which one case is considered possible related to Comirnaty exposure and the other case considered unclassified. The remaining 2 cases are considered BC level 4-5.

In persons aged 25-29 years, there were 5 fatal cases with myocarditis of which 3 cases are considered BC level 1 of which one case is considered possible related to Comirnaty exposure, one case unclassified, and one case unassessable. The remaining 2 cases are considered BC level 4.

In persons aged 30-39 years, there were 5 fatal cases with myocarditis of which 3 cases are
considered BC level 1 and all considered unlikely related to Comirnaty exposure. The remaining 2 cases are considered BC level 4-5.

In persons aged ≥40 years, there were 59 fatal cases with myocarditis of which:

- 15 cases are considered BC level 1 of which three cases are considered possible related to Comirnaty exposure, nine cases unlikely, and three cases unassessable;
- 3 cases are considered BC level 2 of which one case is considered unlikely related to Comirnaty, and two cases unassessable;
- 2 cases are considered BC level 3 and considered both unlikely related to Comirnaty;
- the remaining 39 cases are considered BC level 4-5.

The 6 cases with unknown age are considered all BC level 4.

Based on the information provided concerning the fatal cases reporting myocarditis and despite the 5 BC level 1 cases considered possible related, no new important safety information could be identified.

**Issue solved**

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4. **Regarding pericarditis:**

   a. The MAH is requested to provide detailed information concerning the fatal cases with pericarditis in persons aged 18-24 years, 25-29 years, and ≥40 years and perform per case an assessment according to Brighton Collaboration Pericarditis case definition and Level of certainty classification, a WHO causality assessment, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

   b. Upon analysis of the requested details of the fatal pericarditis cases the MAH is expected to discuss whether the current risk minimisation measures is still adequate.

**MAH response**

The PBRER described a total of 19 interval pericarditis events reporting a fatal outcome. Of them, 8 cases co-reported a myocarditis event and were thus analysed in the myocarditis dataset. The remaining 11 cases are analysed in Appendix 4 (see below in comment box) and a case narrative listing is provided in Appendix 6 (not reproduced here).

Overall, no new safety information was identified from the analysis of these cases and thus, no changes to the risk management measures are warranted.

**Rapporteur assessment comment:**

Please refer regarding the assessment of the 11 fatal cases to the PRAC Rapporteur’s comments in:

![Appendix 4.pdf](image)

In persons aged 25-29 years, there were 2 fatal cases with pericarditis of which both cases are considered BC level 1 and unlikely related to Comirnaty exposure.

In persons aged ≥40 years, there were 9 fatal cases with pericarditis of which two cases are considered BC level 1 and both unlikely related to Comirnaty exposure. The remaining 7 cases are considered BC level 4-5.
Based on the information provided concerning the fatal cases reporting pericarditis, no new important safety information could be identified.

**Issue solved**

5. **Regarding glomerulonephritis**, the MAH is requested to provide detailed information concerning the 46 cases with glomerulonephritis in the interval period and perform an WHO-UMC causality assessment per case, and to provide an updated review with DLP 14 Nov 2022 of cases reporting glomerulonephritis from their safety database including a WHO-UMC causality assessment per case regarding Comirnaty exposure.

**MAH response**

**Details of 46 cases**

As of the most current latest information reported in 46 cases as of 18 November 2022:

There were 2 sets of cases which appeared to be duplicates:

- two cases (AER #...; only AER #... will be discussed below in the Table 3).
- two other cases (AER #... and ...; only AER #... will be discussed below in the Table 4).

Nine cases (AERs...) reported a pre-existing medical condition and/or use of co-suspect/concomitant medication representing a reasonable alternative cause of the relevant event Glomerulonephritis: a pre-existing glomerulonephritis, COVID-19, autoimmune conditions of Autoimmune hepatitis, Psoriatic arthropathy, Autoimmune hypothyroidism and/or use of loop diuretics and immunosuppressants. The WHO causality of the relevant event in these 9 cases is considered as Unlikely. For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

Twenty-three cases co-reported other renal serious PTs (eg. nephrotic syndrome, renal failure) and/or other PTs suggesting a possible alternative aetiology for development of the relevant event (eg. Systemic inflammatory response syndrome, Vasculitis). Table 3 ((see below in comment box) presents some details of these 23 cases and WHO causality assessment derived based on details reported (eg age of the patient, latency, medical history, co-reported PTs). For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

**Rapporteur assessment comment:**

Please refer regarding the assessment of the 23 cases reporting glomerulonephritis and/or other PTs suggesting a possible alternative aetiology for development of the relevant event, to the PRAC Rapporteur's comments in:

![Table 3.pdf](image)

Of the 23 cases there were 2 cases considered possible related to Comirnaty exposure, 14 cases unlikely and 7 cases unassessable.
Table 4 (see below in comment box) presents details of the last 12 cases and WHO causality assessment derived based on details reported (e.g., age of the patient, latency, medical history, co-reported PTs). For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

Rapporteur assessment comment:

Please refer regarding the assessment of the 12 remaining cases reporting glomerulonephritis to the PRAC Rapporteur’s comments in:

Table 4.pdf

Of the 12 cases there were 2 cases considered possible related to Comirnaty exposure, 2 cases unlikely and 8 cases unassessable.

Additional two (2) cases from the last PSUR

Two cases (AER 4[REDACTED] and[REDACTED]) were identified as reported in the PSUR #3 and were updated with a newly added PT glomerulonephritis post DLP of 18 June 2022. Based on the information reported in these cases:

1. The WHO causality assessment in case[REDACTED] is Unlikely; the complex medical history and the use of concomitant medication candesartan confound an assessment in a 62 YO male who developed relevant event about 4 months post Dose 2 with many renal co-reported PTs including vasculitis.

2. The WHO causality assessment in case[REDACTED] is Unlikely; the medical history of autoimmune thyroiditis confounds an assessment in a 15 YO female who developed relevant event 2 days post Dose 3 (the case reported exacerbation of haematuria that patient experienced after dose 2 as well).

These 2 cases will not be reported in the upcoming PSUR #4. For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

Rapporteur assessment comment:

MAH’s conclusion is endorsed, these 2 cases are considered unlikely related to Comirnaty exposure.

Cumulative Review Through 14 November 2022

As per the updated search, through 14 Nov 2022, 15 new cases reporting PT glomerulonephritis were identified.

Five cases (AER 4[REDACTED]) reported a pre-existing medical condition representing a reasonable alternative cause of the relevant event: a pre-existing glomerulonephritis, IgA nephropathy, Nephropathy, Pyelonephritis, COVID-19, and/or Systemic lupus erythematosus. The WHO causality of the relevant event in these 5 cases is considered as Unlikely. For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

Table 5 (see below in comment box) presents details of the last 10 cases and WHO causality assessment derived based on details reported (e.g., age of the patient, latency, medical history, co-
reported PTs). For complete narratives of these cases, refer to Appendix 1 Narrative Line Listing (not reproduced here).

**Rapporteur assessment comment:**

Please refer regarding the assessment of the 10 last cases reporting glomerulonephritis to the PRAC Rapporteur’s comments in:

![Table 5.pdf](image)

Of the 10 cases there were 4 cases considered unlikely related to Comirnaty exposure and 6 cases unassessable.

**MAH’s conclusion**

At this time, considering the available information, there are not sufficient data to conclude a causal relationship between COMIRNATY™ and the new onset or exacerbation of Glomerulonephritis.

**Rapporteur assessment comment:**

The MAH provided detailed information concerning the 46 cases with glomerulonephritis in the interval period and performed an WHO-UMC causality assessment per case, and provided an updated review with DLP 14 Nov 2022 of cases reporting glomerulonephritis.

**Interval period of the PSUR**

Of the 46 cases retrieved there were two duplicate cases, resulting in 44 cases. Two additional cases were added to the interval period, resulting finally in 46 cases:

- 4 cases are considered possible related to Comirnaty exposure;
- 27 cases unlikely related;
- 15 cases unassessable.

**Review through 14 Nov 2022**

Retrieved were 15 new cases reporting glomerulonephritis:

- 9 cases are considered unlikely related to Comirnaty exposure;
- 6 cases unassessable.

In conclusion, despite the 4 cases considered possible related, MAH’s conclusion is endorsed that based on the provided data no causal association of Comirnaty with glomerulonephritis can be concluded. No new important information could be identified concerning glomerulonephritis. The MAH should closely monitor any new cases, patterns, or trends of reporting glomerulonephritis through routine pharmacovigilance.

**Issue solved**
6. Regarding **fatal cases reported in paediatric persons**, the MAH is requested to provide detailed information concerning the fatal cases in persons aged 5-11 years and 12-17 years, perform per case a WHO causality assessment, and an assessment according to Brighton Collaboration case definition and Level of certainty classification, if applicable.

**MAH response**

There were 81 fatal cases in persons aged 5-11 years and 12-17 years, one case [redacted] was a 19-year-old subject. A listing of the 82 cases, including narratives, is provided in Appendix 2 (not reproduced here).

Of note, there were 3 reports [redacted] originating from EMA EudraVigilance-WEB that described non-fatal cases of pyrexia and pyrexia with somnolence; these cases were downgraded to non-serious cases.

Of the remaining 79 cases, 56 cases [redacted] were classified as unassessable because the information provided is either insufficient to assess or contradictory and could not be verified.

**Rapporteur assessment comment:**

The MAH considered all 56 cases unassessable. However, the PRAC Rapporteur considered cases [redacted] and [redacted] unclassified.

Of the remaining 23 cases, 6 described myocarditis [redacted] contributing to death and are therefore included in the response to Question 3.

**Rapporteur assessment comment:**

The cases [redacted] and [redacted] were not presented in detail in MAH’s response on question 3 concerning myocarditis cases. These 2 cases are considered the 2 cases that were updated according to follow-up data and no longer include a myocarditis event in MAH’s response on question 3, and considered unclassified.

Of the remaining 4 cases, 3 cases are considered unlikely related to Comirnaty exposure and 1 case unassessable.

The remaining 17 fatal cases are described in Table 6 (see below in comment box).

**Rapporteur assessment comment:**

Please refer regarding the assessment of the 17 remaining fatal cases to the PRAC Rapporteur’s
Of the 17 cases there were 15 cases considered unlikely related to Comirnaty exposure and 2 cases unassessable.

**MAH’s conclusion**

There is no information in the review of these 82 paediatric fatal cases that identifies BNT162b2 as a contributor to the reported deaths. Fatal cases will continue to be monitored via routine pharmacovigilance.

**Rapporteur assessment comment:**

Of the 82 cases retrieved 3 cases were non-fatal cases, resulting in 79 cases:
- 18 cases are considered unlikely related to Comirnaty exposure;
- 4 cases unclassified;
- 57 cases unassessable.

Overall, MAH’s conclusion is endorsed that based on the provided data no causal association of Comirnaty with the reported deaths can be concluded. No new important information could be identified concerning the reported deaths. The MAH should closely monitor any new cases, patterns, or trends of reporting fatal outcome through routine pharmacovigilance.

**Issue solved**

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**7. Comments from member states**

**MS1**

We endorse the PRAC Rapporteur’s assessment of the above-mentioned procedure, and have no further comments.

**Rapporteur assessment comment:**

The endorsement of the PSUR assessment is appreciated.

**MS2**

We agree with the Rapporteur and the MAH that dizziness should be added as an ADR in the Comirnaty SPC. Further, we would like to point out that there are also other events that are currently considered listed by the MAH as anxiety-related reactions, but which we consider not covered by the current SPC.

The table 18 in the PSUR presents the most commonly reported ADRs in the post-marketing data, and in this table dyspnoea and palpitations are marked as *listed or consistent with the listed AEs in the*
current RSI”. We assume that the MAH considers these events to be covered by the text in the section 4.4 describing anxiety-related reactions, as they are not listed in SPC section 4.8. However, based on the medical assessment of the reports received by MS2, in majority of the reports describing dyspnoea or palpitations these symptoms have lasted several days or even weeks, and thus are not considered to be anxiety-related reactions as described by the text in the section 4.4.

Correspondingly, in the section “Vaccination stress/anxiety related ADRs” tachycardia is presented as one of the stress/anxiety-related reactions covered by the current text in the SPC. However, in the reports received by MS2 describing tachycardia/increase in heart rate, these events have lasted several days or even weeks in majority of the reports, and thus are not covered by the text concerning anxiety-related reactions in the section 4.4.

We propose that in the next PSUR the MAH should present cumulative analyses of dyspnoea, palpitations and tachycardia/heart rate increase, with special focus on the duration of the events. The MAH should evaluate whether these events should be added in the section 4.8 of the SPC.

Rapporteur assessment comment:

Here we can agree with the comments. The proposed request for the next PSUR that the MAH should present cumulative analyses of dyspnoea, palpitations and tachycardia/heart rate increase, with special focus on the duration of the events not considered stress/anxiety-related reactions and to evaluate whether these events should be added in the section 4.8 of the SPC, is added.

MS3

The MS3 PRAC Rapporteur notes the request no 1 for supplementary information regarding multiple repeated booster doses.

At the time of approval of the bivalent mRNA vaccines, the MAHs committed to include additional pharmacovigilance activities (the new strains) in relevant PASS protocols at the first regulatory opportunity. These updates are currently ongoing. With this in mind and also given that the vast majority of future post-marketing data are expected to originate from vaccines used as part of multiple repeated booster dose schemes, the PRAC Rapporteur is unsure regarding to which extent a missing information category in the RMP summary of safety concerns will contribute additionally to the PSUR.

If potential safety issues specifically related to the new strains are of interest, a request {confidential information deleted} might be considered:

“Bivalent variant updated Comirnaty vaccines: After the DLP of the PSUR no 3, Comirnaty has been variant updated (bivalent vaccines), approved and is currently being used in the EU. In light of this, in the next PSUR, the MAH is requested to ensure a transparent presentation of cases reported from the newly approved variant vaccines, as well as a continuous comparison of the safety issues identified with the evidence available for the original vaccine. Any differences in occurrence of safety issues, when comparing the variant updated vaccines with the original monovalent Comirnaty or, indeed, when comparing the two different variant updated bivalent Comirnaty vaccines, should be discussed.”

Rapporteur assessment comment:

Please refer to our assessment of MAH’s response on the 1st request for supplementary information in section 6. Proposed request by the MS3 PRAC Rapporteur is added to the requests for next PSUR.

MS4
MS4 endorses the Rapporteur assessment report. However, MS4 has two additional comments regarding cases of “hearing loss” and “acquired haemophilia”.

**Hearing loss**

In MS4, hearing loss following vaccination against COVID-19 with mRNA vaccines are closely monitored and analysed.

A national cross-sectional audiogram-based study was conducted, using the MS4 pharmacovigilance active surveillance system for COVID-19 vaccines. All suspected Sudden Sensorineural Hearing Loss (SSNHL) cases following mRNA COVID-19 vaccination between January 2021 and February 2022 were included. They were retrospectively reviewed based on a comprehensive audiological and medical evaluation by ENT. {confidential information removed}

Over the study period, 97,840,529 doses of Tozinameran (Pfizer-BioNTech BNT162b2) were administered in MS4. The Reporting Rates (RR) of mRNA vaccine-induced SSNHL cases were calculated per 1,000,000 injections. Clinical classification was made according to patient history, unilaterality or bilaterality of the hearing loss, its degree, and recovery after a minimum 3-month follow-up.

For these Tozinameran-induced SSNHL cases, the delay onset was ≤21 days for 108 (76%) cases whose median (range) delay onset was 4 (2.0-9.0) days. Women were concerned in 84 (59%) cases. The median (range) age was 51 (13-83) years, and 98 (69%) patients were in the 30-64 years age class. A total of 50 (35%) patients had a medical history, it was otoneurologic in 17 (12%) cases. The vaccination rank was known for 125 cases, the first injection was involved in 60 (42%) cases. Steroids were administered orally in 67 (47%) cases. SSNHL was unilateral in 142 (79%) cases. Detailed audiometric thresholds were available in 98 (69%) cases, with SSNHL being measured as mild to moderately severe in 61/98 (62%) cases, and as profound in 17 (17%) cases. Tinnitus was associated with SSNHL in 75 (53%) cases and vertigo in 41 (29%) cases. Total recovery was observed in 37 (25%) cases while hearing aid fitting was required in 10 (7%) cases (Table 1). Deafness was more often unilateral than bilateral (p<0.001). Neither sex effect nor vaccination rank effect was found. Case follow-up identified 5 (4%) cases of positive rechallenge (Table 2).

The total RR was estimated at 1.45/1,000,000 doses for Tozinameran.

**Table 1: Characteristics of the included SSNHL cases**

<table>
<thead>
<tr>
<th>Tozinameran</th>
<th>≤21 d</th>
<th>&gt;21 d</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number of patients</strong></td>
<td>108 (76)</td>
<td>34 (24)</td>
<td>142 (100)</td>
</tr>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>43 (40)</td>
<td>15 (44)</td>
<td>58 (41)</td>
</tr>
<tr>
<td>Female</td>
<td>65 (60)</td>
<td>19 (56)</td>
<td>84 (59)</td>
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<tr>
<td><strong>Age, years, median (range)</strong></td>
<td>50 (13-83)</td>
<td>52 (16-72)</td>
<td>51 (13-83)</td>
</tr>
<tr>
<td>0-18</td>
<td>3 (3)</td>
<td>1 (3)</td>
<td>4 (3)</td>
</tr>
<tr>
<td>19-29</td>
<td>7 (6)</td>
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<td>7 (5)</td>
</tr>
<tr>
<td>30-49</td>
<td>43 (40)</td>
<td>12 (35)</td>
<td>55 (39)</td>
</tr>
<tr>
<td>Age Group</td>
<td>Total</td>
<td>Cases</td>
<td>%</td>
</tr>
<tr>
<td>-----------</td>
<td>-------</td>
<td>-------</td>
<td>---</td>
</tr>
<tr>
<td>50-64</td>
<td>26 (24)</td>
<td>17 (50)</td>
<td>43 (30)</td>
</tr>
<tr>
<td>65-74</td>
<td>19 (18)</td>
<td>4 (12)</td>
<td>23 (16)</td>
</tr>
<tr>
<td>≥75</td>
<td>10 (9)</td>
<td>0</td>
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<tr>
<td><strong>Medical History</strong></td>
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</tr>
<tr>
<td>Cardiovascular (CV)</td>
<td>9 (8)</td>
<td>5 (15)</td>
<td>14 (10)</td>
</tr>
<tr>
<td>Otoneurologic (ON)</td>
<td>12 (11)</td>
<td>5 (15)</td>
<td>17 (12)</td>
</tr>
<tr>
<td>Auto Immune disease (AIM)</td>
<td>7 (6)</td>
<td>2 (5)</td>
<td>9 (6)</td>
</tr>
<tr>
<td>CV and ON</td>
<td>4 (4)</td>
<td>0</td>
<td>4 (3)</td>
</tr>
<tr>
<td>ON and AIM</td>
<td>3 (3)</td>
<td>0</td>
<td>3 (2)</td>
</tr>
<tr>
<td>Other etiology</td>
<td>3 (3)</td>
<td>0</td>
<td>3 (2)</td>
</tr>
<tr>
<td><em><em>Delay onset</em>, days, median (range)</em>*</td>
<td>4 (2.0-9.0)</td>
<td>41 (25-67)</td>
<td></td>
</tr>
<tr>
<td><strong>Vaccination Rank</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First Injection</td>
<td>47 (44)</td>
<td>13 (38)</td>
<td>60 (42)</td>
</tr>
<tr>
<td>Second Injection</td>
<td>39 (36)</td>
<td>14 (41)</td>
<td>53 (37)</td>
</tr>
<tr>
<td>Booster</td>
<td>12 (11)</td>
<td>0</td>
<td>12 (9)</td>
</tr>
<tr>
<td>Unknown</td>
<td>10 (9)</td>
<td>7 (21)</td>
<td>17 (12)</td>
</tr>
<tr>
<td><strong>Oral steroid administration</strong></td>
<td>48 (44)</td>
<td>19 (56)</td>
<td>67 (47)</td>
</tr>
<tr>
<td><strong>Laterality</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unilateral</td>
<td>87 (80)</td>
<td>25 (73)</td>
<td>112 (79)</td>
</tr>
<tr>
<td>Bilateral</td>
<td>16 (15)</td>
<td>4 (12)</td>
<td>20 (14)</td>
</tr>
<tr>
<td>Unknown</td>
<td>5 (5)</td>
<td>5 (15)</td>
<td>10 (7)</td>
</tr>
<tr>
<td><strong>Degree of Hearing Loss [Pure Tone Average]</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild [26–40 dB HL]</td>
<td>27 (35)</td>
<td>4 (18)</td>
<td>31 (32)</td>
</tr>
<tr>
<td>Moderate [41–55 dB HL]</td>
<td>14 (18)</td>
<td>5 (22)</td>
<td>19 (20)</td>
</tr>
<tr>
<td>Moderately severe [56–70 dB HL]</td>
<td>8 (11)</td>
<td>3 (14)</td>
<td>11 (11)</td>
</tr>
<tr>
<td>Severe [71–90 dB HL]</td>
<td>11 (15)</td>
<td>4 (18)</td>
<td>15 (15)</td>
</tr>
<tr>
<td>Profound [&gt; 90 dB HL]</td>
<td>12 (16)</td>
<td>5 (23)</td>
<td>17 (17)</td>
</tr>
<tr>
<td>Associated cochleovestibular disorders</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Tinnitus</td>
<td>59 (55)</td>
<td>16 (47)</td>
<td>75 (53)</td>
</tr>
<tr>
<td>Vertigo and balance disorders</td>
<td>33 (30)</td>
<td>8 (24)</td>
<td>41 (29)</td>
</tr>
<tr>
<td><strong>Time to recovery, days, median (range)</strong></td>
<td><strong>15 (5-67.5)</strong></td>
<td><strong>11 (8-22.5)</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Outcome</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total recovery</td>
<td>26 (24)</td>
<td>11 (32)</td>
<td>37 (25)</td>
</tr>
<tr>
<td>No recovery at follow-up</td>
<td>66 (62)</td>
<td>18 (53)</td>
<td>84 (60)</td>
</tr>
<tr>
<td>Hearing aid fitting requirement</td>
<td>8 (7)</td>
<td>2 (6)</td>
<td>10 (7)</td>
</tr>
<tr>
<td>Unknown</td>
<td>8 (7)</td>
<td>3 (9)</td>
<td>11 (8)</td>
</tr>
<tr>
<td><strong>Positive Rechallenge</strong></td>
<td><strong>5 (5)</strong></td>
<td><strong>0</strong></td>
<td><strong>5 (4)</strong></td>
</tr>
</tbody>
</table>

Table 2: Characteristics of patients who experienced a positive rechallenge

<table>
<thead>
<tr>
<th>Cases</th>
<th>Sex</th>
<th>Age</th>
<th>Medical History</th>
<th>Vertigo</th>
<th>Tinnitus</th>
<th>Vaccination rank</th>
<th>Delay onset</th>
<th>Unilateral/bilateral</th>
<th>Deafness degree</th>
<th>Progression</th>
</tr>
</thead>
<tbody>
<tr>
<td>N°1</td>
<td>F</td>
<td>61</td>
<td>0</td>
<td>yes</td>
<td>yes</td>
<td>D1 D2</td>
<td>4 days, 7 days</td>
<td>Unilateral</td>
<td>Moderate</td>
<td>Lasting</td>
</tr>
<tr>
<td>N°2</td>
<td>M</td>
<td>47</td>
<td>0</td>
<td>no</td>
<td>yes</td>
<td>D1 D2</td>
<td>9 days, 9 days</td>
<td>Unilateral</td>
<td>Moderate</td>
<td>Lasting</td>
</tr>
<tr>
<td>N°3</td>
<td>F</td>
<td>74</td>
<td>Fluctuating deafness and vertigo treated by cortisone punctually</td>
<td>no</td>
<td>yes</td>
<td>D1 D2</td>
<td>8 days, 9 days</td>
<td>Bilateral</td>
<td>Moderate</td>
<td>Resolved</td>
</tr>
<tr>
<td>N°4</td>
<td>F</td>
<td>32</td>
<td>Protein S deficiency</td>
<td>no</td>
<td>no</td>
<td>D1 D2</td>
<td>8 days, 2 days</td>
<td>Unilateral</td>
<td>Mild</td>
<td>Resolved</td>
</tr>
<tr>
<td>N°5</td>
<td>F</td>
<td>79</td>
<td>Diabetes stabilized since 1977, Left Meniere's disease since 1977, Hypothyroidism stabilized, High blood pressure stabilized since 1985, Hypercholesterolemia stabilized since 1985</td>
<td>yes</td>
<td>yes</td>
<td>D1 D2 R1</td>
<td>8 days, 8 days, 1 day</td>
<td>Bilateral</td>
<td>Moderate</td>
<td>Resolved</td>
</tr>
</tbody>
</table>
D1: first vaccine injection, D2: second vaccine injection, R1: first booster, R-: no recurrence of deafness

[confidential information removed]

Conclusions

MS4 endorses that a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time.

However, taking into account that there is 5 well documented cases of positive rechallenge with a compatible TTO (≤21 days) of which 2 are bilateral hearing loss and 13 cases of bilateral hearing loss with a compatible TTO (≤21 days), MS4 considers that a causal association between hearing loss and the vaccination cannot be completely ruled out.

Therefore, MS4 considers that the MAH should continue to closely monitor hearing loss and all new cases should be reported and discussed in the future PSURs.

Rapporteur assessment comment:

An interesting submitted manuscript from the MS4 PV experts is shared prior to publication which is appreciated. However, the manuscript is not (yet) peer-reviewed and published. Therefore it is currently unknown whether a peer-review process would introduce changes to the manuscript that would impact on the results of the performed MS4 study. We assume that the described MS4 cases (between Jan 2021 – Feb 2022) had been submitted to EudraVigilance and therefore were also included in the current assessed cumulative review (through 18 Jun 2022). Also, there is endorsement that a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time after a thorough assessment of the current available information through 18 Jun 2022. When the manuscript is published in the near future, the MAH should discuss the study in the PSUR and the new cases reporting hearing loss as appropriate.

Acquired haemophilia

MS4 endorses the Rapporteurs conclusions that a causal association between Comirnaty and acquired haemophilia cannot be concluded at this time.

Moreover, taking into account the recent literature data, for example the review of Franchini et al., and the potential mechanism of action, MS4 endorses the request that the MAH should continue to closely monitor acquired haemophilia and all new cases should be reported and discuss in the future PSURs.


MS5

We overall agree with Rapporteur’s assessment, but would like to highlight some issues:

Autoimmune hepatitis

Overall, we endorse the PRAC Rapporteur assessment report. At this stage, only 99 cases with biopsy results (60 cases) or laboratory data (39 cases) were identified. From the cases with biopsy results, only 7 meet IAIHG-revised scores and 9 meet IAIHG-simplified scores. However, from the cases that meet criteria some of them have a time to onset not compatible with AIH (2 and 4 cases respectively). Regarding the cases with laboratory data available, none had sufficient information to score ≥6 and they cannot be classified as definite or probable cases. As the MAH stated there are several limitations regarding the interpretation of reports submitted voluntarily such as the underreporting and the lack of some important data for a thorough case evaluation, with impact in the O/E analysis.

We want to highlight a pre-proof article (Codoni et al, 2022) performed with cases collected from members of the International AIH Group (IAIHG) and the European Reference Network on Hepatological Diseases (ERN RARE-LIVER). The main advantages of this article are that only cases without known pre-existing liver diseases and transaminase Levels ≥5xULN within 3 months after any anti-SARS-CoV-2 vaccine were considered and that all cases have available liver biopsy. Fifty-nine patients, from 26 centers in 11 countries and exposed to seven different SARS-COV-2 vaccines were recruited. Most of the patients (35) were female patients. Hepatitis was diagnosed after the second vaccine dose in the majority of patients. Although the study included cases with 7 different COVID-19 vaccines, patients with liver injury after mRNA vaccines had higher transaminase levels and higher impairment of coagulation. A comparison between the two mRNA vaccines was limited by the small patient numbers. In three quarters of the cases, liver histology showed a picture of predominant lobular hepatitis while predominant portal hepatitis was present in less than 1/5 of patients. The absence of advanced liver fibrosis in the work-up of an acute liver injury suggests drug-induced liver injury (DILI) or AIH-like DILI as more probable than AIH. In addition, AIH-like is characterized by a low relapse rate after withdrawal of a short-term steroid course. 91% of patients were treated with steroids, ± azathioprine. Serum transaminase levels improved in all cases and normalised 24/58 (41%) after three months, and in 30/46 (65%) after six months. One patient required liver transplantation. Re-exposure to SARS-CoV-2 vaccines of 15 patients resulted in four relapses (three after the same vaccine and one in a heterologous vaccination).

The systematic review mentioned by the MAH (Roy et al) including 23 patients with histopathological data in 13 studies showed similar results to the data observed for Comirnaty and the previous article. The authors of this systematic review stated that biochemically and histologically, most of the cases with ILI resembled AIH.

In summary, AIH should continue to be closely monitored in next PSURs. It would be important to follow-up the cases to differentiate real cases of AIH or AIH-like DILI, and consider the addition of some information in the SmPC.

autoimmune hepatitis should be closely monitored in the PSURs (as is stated in the AR). To differentiate at this stage between cases of AIH or AIH-like DILI without any new important information/signal concerning the occurrence of autoimmune hepatitis after Comirnaty exposure, seems however premature.

Thromboembolic events

We endorse the Rapporteur evaluation of thromboembolic events performed in previous monthly reports and summarized in this PSUR. To note that pulmonary embolism is one the most frequently reported events. Recently, 7 cases included under the HLT pulmonary thrombotic and embolic conditions have been reported in MS5 after the administration of Comirnaty Original/Omicron BA.4-5 within 2 weeks after vaccination. In four of these seven cases, flu vaccine was administered the same day of COVID-19 vaccine and in 1 case two days after. No disproportionality was found in the O/E analyses, but in view of these cases, we consider that pulmonary embolism should be closely monitored in the next PSUR. Particularly, a detailed analysis of the cases occurring with the Comirnaty original/omicron BA 4-5 should be performed.

Rapporteur assessment comment:

The endorsement of the evaluation of thromboembolic events is appreciated. As stated no disproportionality was found in the O/E analyses. Therefore, there is no new important information/signal concerning the occurrence of pulmonary embolism after Comirnaty exposure and no need at the moment to request a review of pulmonary embolism in the next PSUR. AEsIs including thromboembolic events will be closely monitored in PSURs.

Please refer to the MS3 comments regarding a detailed analysis of the cases occurring with the Comirnaty original/omicron BA 4-5, above.

Hearing loss

18 896 spontaneous cases have been identified, 755 medically confirmed. We want to highlight the following considerations regarding these cases:

- Those cases that did not report diagnostic procedures or test have been considered “unassessable” according to the WHO-UMC criteria. In our view, this restrictive approach would not be appropriate, considering that these are cases already medically confirmed and it can be assumed that clinical judgement applies. This would affect around 386 out of 755 medically confirmed cases. It is worrying that serious cases, clinically confirmed may not be taken into account based on these criteria, but the number of serious cases clinically confirmed as well as the seriousness criteria has not been presented by the MAH. Therefore, the MAH should rather conduct the analysis using the Brighton Collaboration Criteria for sensorineural hearing loss.

- It seems that the majority occur after the first and second dose and they do not recover (392 of the 755), they recover or are recovering (292 of the 755). In addition, there are 33 patients that recover but with sequelae. The MAH should provide further information on these patients that recover with sequela to better understand their particular situations and evaluate possible patterns.

Moreover, there are still 9023 cases that although not medically confirmed have not been taken into account despite some of them may have plausible TTO and/or not confounding factors.

Regarding O/E analysis, the assessors noted that the MAH imputed missing values of age, sex, dose and time to onset based on observed cases with known data. Although this may be a correct approach,
this should have been conducted as a sensitivity analysis, to be able to assess differences in O/E rates according to missing values. Nevertheless, it is known that these analysis although support the evaluation, would not confirm a causal association with the vaccine.

Moreover, in our view’s, not all PTs should have the same weight in this analysis, being the sensorineural hearing loss is of utmost importance since the damage can be irreversible, depending on form (unilateral or bilateral) and symptoms onset. A distinction between sudden or progressive sensorineural hearing loss should be considered. In fact, the PRAC already requested to perform the cumulative review of cases of sudden sensorineural hearing loss, but further PTs have been included in this cumulative review. This may have diluted the evaluation of the cases.

In summary, we consider that based on the data presented, a causal association cannot be established nor discarded and changes in the product information are not warranted at this stage, but it should be monitored in the next PSUR. In particular, the MAH should reconsider PTs for the evaluation focussing on sudden sensorineural hearing loss and medically confirmed cases, and provide an updated cumulative review applying the Brighton Collaboration criteria for the evaluation of these cases, indicating serious cases and the seriousness criteria applied, as well as cases with outcome recovered with sequela, and rechallenge cases if new ones arise. Particular attention should be taken with regard to event outcome to determine whether patients recover from the event, and when, or not.

**Rapporteur assessment comment:**

Here we agree that serious clinically confirmed cases may not be taken into account due to that the MAH considered cases that did not report diagnostic procedures or test, as ‘unassessable’. However, the number of serious clinically confirmed cases as well as the seriousness criteria has not been presented by the MAH. Therefore, we endorse that the MAH should conduct the analysis using the Brighton Collaboration Criteria for sensorineural hearing loss in future reviews of cases reporting hearing loss which is added as a request for next PSUR.

Please also refer to the MS4 comments above regarding hearing loss. Requesting the MAH to reconsider PTs for the evaluation focussing on sudden sensorineural hearing loss and medically confirmed cases in the next PSUR is not considered of added value because the O/E analysis showed that all O/E ratios were well below 1 and there is at the moment no new important information/signal concerning the occurrence of hearing loss after Comirnaty exposure.

**IgA nephropathy and other glomerulonephritis**

We consider that this safety topic should be kept as an important potential risk in the PSURs. Although at this stage there is no enough data to establish a causal relationship between Comirnaty and the development of IgA nephropathy, there are several cases with close temporal association. Additionally, a possible mechanism of action is described in the article by Farooq et al. The Rapporteur asked for clarifications regarding the increase in the number of cases of glomerulonephritis cases during this current interval. In MS5, 22 cases with glomerulonephritis were cumulatively reported and only 2 cases were IgA nephropathy. To note that there have been two cases with positive rechallenge. One of them in 53-year-old male patient who suffered macroscopic hematuria and nephritic syndrome after the two doses of Comirnaty and the other one in a 32-year-old male patient that suffered flares of its nephrotic syndrome after the two doses of Comirnaty. Therefore, if no additional conclusions are obtained from the responses required in the comments period, we consider that glomerulonephritis other than IgA nephropathy should be also closely monitored in the next PSUR.

**Rapporteur assessment comment:**
Please refer to the assessment of MAH's response on the 5th request for supplementary information in section 6. Here the PRAC Rapporteur concludes that based on the provided data no new important information could be identified concerning glomerulonephritis. The MAH should closely monitor any new cases, patterns, or trends of reporting glomerulonephritis through routine pharmacovigilance.

8. Late-breaking information PRAC rapporteur

Rapporteur assessment comment:

Regarding post orthostatic tachycardia syndrome (POTS), a recent publication has been noticed:


The authors found, for new diagnoses made after vaccination, that the five conditions with the highest post-vaccination odds of (new) diagnoses were myocarditis, dysautonomia, POTS, mast cell activation syndrome and urinary tract infection (UTI). Overall, the post-vaccination odds of new POTS-associated diagnoses (n = 4,526, odds = 1.33 (1.25–1.41), P < 0.001) was higher than for common primary care (CPC) diagnoses (n = 33,590, odds = 1.21 (1.18–1.23), P < 0.001) but lower than for myocarditis (n = 25, odds = 2.57 (1.02–6.77), P = 0.046). In repeated analyses around receipt of second (rather than the first) vaccination dose, overall similar findings were observed. In summary, POTS-related diagnoses appear to occur with increased frequency in the time period after COVID-19 vaccination as compared to the time period before, particularly when compared to more commonly diagnosed conditions, but at a rate that is approximately five times lower than after SARS-CoV-2 infection.

Therefore, the MAH is requested to discuss the publication of Kwan et al. concerning post orthostatic tachycardia syndrome and Comirnaty exposure and, if applicable, to perform a cumulative review on the association between Comirnaty and post orthostatic tachycardia syndrome. This should include, but not be limited to, information from post-marketing cases, clinical trials, mechanistic studies and literature. The MAH should consider the need for an update of the product information and/or RMP.

Request for next PSUR
PERIODIC SAFETY UPDATE REPORT #3
for
ACTIVE SUBSTANCE: COVID 19 mRNA vaccine (nucleoside modified) (BNT162b2)¹

ATC CODE: J07BX03²

AUTHORISATION PROCEDURE in the EU: Centralised

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EUROPEAN UNION REFERENCE DATE (EURD): 19 DECEMBER 2020

INTERVAL COVERED BY THIS REPORT:
19 DECEMBER 2021 through 18 JUNE 2022

DATE OF THIS REPORT: 18 AUGUST 2022

SIGNATURE: ___________________________ Date: 18 August 2022

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Please note that this report may contain unblinded clinical trial information.

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¹ Change of the name of the active substance from COVID-19 mRNA Vaccine (nucleoside modified) to Tozinameran in EU (EMEA/H/C/005735/X/0044/G).
² Implementation as new ATC code starting from 01 January 2022.
³ Earliest conditional approval date.

CONFIDENTIAL
**EXECUTIVE SUMMARY**

This is the 3rd Periodic Safety Update Report (PSUR) for COVID-19 mRNA vaccine (nucleoside modified) (Coronavirus disease 2019 [COVID-19] mRNA Vaccine, COMIRNATY®, also referred to as BNT162b2), covering the reporting interval 19 December 2021 through 18 June 2022.

A product description is provided in Table 1.

**Table 1.  Product Description**

<table>
<thead>
<tr>
<th>Therapeutic class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>The active substance of the COVID-19 mRNA vaccine is a highly purified single-stranded, 5'-capped mRNA produced using a cell-free in vitro transcription from the corresponding DNA template, encoding the viral spike (S) protein of SARS-CoV-2.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Mechanism of action</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>The nucleoside-modified mRNA is formulated in LNPs, which enable delivery of the RNA into host cells to allow expression of the SARS-CoV-2 S antigen. The vaccine elicits both neutralising antibody and cellular immune responses to the spike (S) antigen, which may contribute to protection against COVID-19.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Indications</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active immunisation to prevent COVID-19 caused by SARS-CoV-2 virus, in individuals 5 years of age and older.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Formulation and route of administration</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>The vaccine is a white to off-white frozen solution, is administered intramuscularly in the deltoid muscle and is available in 3 presentations.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Purple cap (for 12 years of age and older)</th>
<th>Grey cap (for 12 years of age and older)</th>
<th>Orange cap (for age 5 years to &lt;12 years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentrate for dispersion for injection</td>
<td>Dispersion for injection</td>
<td>Concentrate for dispersion for injection</td>
</tr>
<tr>
<td>30 micrograms/dose</td>
<td>30 micrograms/dose</td>
<td>10 micrograms/dose</td>
</tr>
<tr>
<td>Requires dilution</td>
<td>Do not dilute</td>
<td>Requires dilution</td>
</tr>
<tr>
<td>PBS/Sucrose presentation</td>
<td>Tris/Sucrose presentation</td>
<td>Tris/Sucrose presentation</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Posology</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Individuals aged 12 years and older</td>
<td>The 2 formulations (purple cap and grey cap) are administered as 30 μg/dose as a primary series of 2 doses (0.3 mL each) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose (third dose) may be administered approximately 6 months after the second dose in individuals 16 years of age and older.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Individuals aged 5 through 11 years</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Tris/Sucrose formulation (orange cap) is administered after dilution as a primary series of 2 doses (0.2 mL) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose may be administered at least 6 months after the second dose.</td>
<td></td>
</tr>
</tbody>
</table>

---

4 Also referred to as Pfizer-BioNTech COVID-19 vaccine in other Company's documents.
On 17 June 2022, an additional formulation was approved first in the United States (US): the paediatric Tris/Sucrose presentation - maroon cap (3 micrograms/dose) for individuals aged between 6 months and 4 years. This is a concentration for dispersion for injection, to be administered after dilution intramuscularly in the anterolateral aspect of the thigh (or in the deltoid muscle in individuals 1 year of age and older) as a primary series of 3 doses (0.2 mL). The initial 2 doses are administered 3 weeks apart followed by a third dose administered at least 8 weeks after the second dose.

Cumulatively, it is estimated that 66,656\(^5\) participants have received BNT162b2 in sponsor initiated clinical trials worldwide, with 59,260 participants exposed to BNT162b2, 1836 participants exposed to clinical candidates developed as variant vaccines based on BNT162b2 (BNT162b2 [B.1.351], BNT162b2 [B.1.1.7 + B.1.617.2], BNT162b2 [B.1.617.2] and BNT162b2 [B.1.1.7]) and 633 participants exposed to other early development candidates (including BNT162a1 [30], BNT162b1 [411] to, BNT162b3 and BNT162c2 [96 participants each]). There were 7044 participants exposed to blinded therapy and 5871 to placebo.

There were 372 participants who received BNT162b2 as a study vaccine or as a comparator in another Pfizer clinical development program (B747).

From the receipt of the first temporary authorisation for emergency supply on 01 December 2020\(^6\) through 18 June 2022, approximately 3,555,998,805 doses of BNT162b2 were shipped from BioNTech and Pfizer worldwide, corresponding to 2,693,922,584 estimated administered doses.

During the current reporting interval (19 December 2021 through 18 June 2022), approximately 1,115,282,160 doses of BNT162b2 were shipped from BioNTech and Pfizer worldwide, corresponding to 843,724,061 estimated administered doses.\(^7\)

Overall, through 18 June 2022, a total of 143,844,450 adult Tris/Sucrose doses and a total of 229,269,400 paediatric Tris/Sucrose doses were shipped worldwide.

Additionally, as per data provided by license partner (LP) in Hong Kong, Macau, and Taiwan, 27,314,884 doses of BNT162b2 were administered cumulatively through 21 June 2022 and 12,126,713 dose were administered from 19 December 2021 through 21 June 2022.

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\(^5\) Participants to more than one clinical trial (e.g., extension study) are counted once when receiving the same treatment in the parent study.

\(^6\) BNT162b2 received first temporary authorisation for emergency supply under Regulation 174 in the UK on this date.

\(^7\) License Partner data are not included in the reported amount.
Details about BNT162b2 marketing authorisation by type of formulation, and population include:

- The PBS/Sucrose 30 μg formulation for individuals aged 16 years and older has received approvals in 103 countries\(^8\) including full (5), conditional (49), emergency use authorisation (EUA) and other type of approvals (52).

- The PBS/Sucrose 30 μg formulation for individuals aged between 12 and 15 years has received approvals in 81 countries\(^9\) including full (2), conditional (46), EUA and other type of approvals (34).

- The Tris/Sucrose 30 μg formulation for individuals aged 12 years and older has received approvals in 73 countries\(^10\) including full (3), conditional (44), EUA and other type of approvals (28).

- The Tris/Sucrose 10 μg formulation for individuals aged between 5 and 11 years has received approvals in 79 countries\(^9\) including full (2), conditional (43), EUA and other type of approvals (35).

- The Tris/Sucrose 3 μg formulation for individuals aged between 6 months and 4 years has received EUA approval in the US.

- The booster dose has received approvals in 83 countries\(^11\) including full (3), conditional (46), EUA and other type of approvals (36).

The use of BNT162b2 in individuals aged 12 years and older is under EUA in Hong Kong and under a special import permit in Macau and Taiwan. In Hong Kong only the PBS/Sucrose – Purple cap formulation was approved.

The marketing authorisation holders (MAHs) of BNT162b2 are the following: BioNTech (56 countries); Pfizer (40 countries), the local Ministry of Health (MoH) and local Government (3 countries each), the LP Fosun Pharma (2 countries), and the LP Hemas (1 country).

In addition, World Health Organization (WHO) had approved the Emergency Use Listing (EUL) of BNT162b2.

There were no marketing authorisation withdrawals for safety reasons during the reporting interval.

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\(^8\) For this population, both conditional and EUA approvals were granted in the United Kingdom (UK), full and EUA approvals in Singapore and the US.

\(^9\) Both conditional and EUA approvals for this population were granted in the UK.

\(^10\) For this population, both conditional and EUA approvals were granted in the UK, full and EUA approvals in the US.

\(^11\) For this population, both conditional and EUA approvals were granted in the UK, full and EUA approvals in Singapore.
During the reporting period, no actions have been taken with respect to BNT162b2 for safety reasons, either by a Health Authority (HA) or by the MAH. Although not considered by definition a regulatory action taken for safety reasons because it does not significantly impact the benefit risk balance of use of the product in authorised populations, due to the receipt of spontaneous reports of Guillain-Barré syndrome (GBS) after vaccination with mRNA COVID-19 vaccines including BNT162b2, Pharmaceuticals and Medical Devices Agency (PMDA) in Japan has required class changes to include GBS in the important precautions section of the Japan package insert and inclusion of GBS as an important potential risk in the Japan Risk Management Plan (RMP). It should be noted that based on PMDA assessment, the frequency of reported cases of GBS was not significantly higher than the background incidence in any gender or age group and a mechanism is not known.

The reference safety information (RSI) for this PSUR is the COVID 19 mRNA vaccine Core Data Sheet (CDS) version 13.0 dated 10 May 2022, in effect at the end of the reporting period.

Four (4) previous CDS versions (version 9.0 dated 02 December 2021, version 10.0 dated 21 December 2021, version 11.0 dated 14 January 2022 and version 12.0 dated 23 March 2022) were also in effect during the reporting period.

Safety-related changes included updates of the following sections: 4.2 Posology and method of administration (CDS version 13.0), 4.8 Undesirable effects (CDS versions 10.0, 11.0 and 13.0), 5.1 Pharmacodynamic properties (CDS versions 10.0 and 11.0), Appendix A, Appendix B (CDS version 10.0).

During the reporting period, the following signals were addressed:

- Signals determined not to be risks: Appendicitis, Hemolytic anemia, Uveitis, Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders, Capillary leak syndrome (CLS), Corneal graft rejection, Vasculitis, Cerebral venous sinus thrombosis (CVST), Lymphocytic colitis, Chronic urticaria, Polymyalgia rheumatica (PMR), Subacute thyroiditis (SAT), Cerebrovascular accident (CVA)/stroke, Amenorrhea, Heavy menstrual bleeding, Loss of/altered taste and smell.

- Signal determined to be an identified risk (not important): Irritability.

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12 The Japan package insert was updated by the MAH during the current reporting period, on 10 June 2022.

13 Guillain-Barré syndrome was added as important potential risk to the safety concerns in the Japan RMP after Data Lock Point (DLP) of this PSUR, on 22 June 2022.

14 This version of the CDS did not include any safety-related changes.
• Signal determined to be an important identified risk: Myocarditis and pericarditis\(^\text{15}\).

• Ongoing signal: Hearing loss.

Commitments to be addressed in this PSUR were received from European Medicines Agency (EMA), World Health Organization (WHO) and Health Canada. The Pharmacovigilance Risk Assessment Committee (PRAC) requests were included in the Assessment Reports (ARs) of the Summary Safety Reports (SSRs), in the Final AR of PSUR #2 and in signals’ AR. The WHO requests were included in the EUL Procedure. Topics covered in these commitments are summarised in the table below.

<table>
<thead>
<tr>
<th>HA</th>
<th>Commitment(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRAC</td>
<td>Closely monitoring multisystem inflammatory syndrome in children and in adults (MIS-C/A) and reporting of new cases of MIS.</td>
</tr>
<tr>
<td></td>
<td>Observed vs Expected (O/E) analyses using at least no risk window, 14-day risk window and 21-day risk window and sensitivity O/E analyses which include the processed cases plus the backlog cases.</td>
</tr>
<tr>
<td></td>
<td>Assessment of the used study methods in not (yet) peer-reviewed retrieved relevant literature to determine if the study results are valid or not.</td>
</tr>
<tr>
<td></td>
<td>More effort in presenting/evaluating the cases considered to be confounded and present the risk factors for developing the respective conditions.</td>
</tr>
<tr>
<td></td>
<td>Use of follow-up questionnaires anaphylaxis and vaccine associated enhanced disease/vaccine associated enhanced respiratory disease (VAED/VAERD).</td>
</tr>
<tr>
<td></td>
<td>Presentation of all relevant literature concerning the safety of Comirnaty (also besides the database Medline and Embase) during the reporting period.</td>
</tr>
<tr>
<td></td>
<td>Safety evaluation of sudden sensorineural hearing loss, tinnitus, glomerulonephritis and nephrotic syndrome, autoimmune hepatitis, dizziness, acquired haemophilia, IgA nephropathy.</td>
</tr>
<tr>
<td></td>
<td>Continue to report on the number of processed cases downloaded from EudraVigilance.</td>
</tr>
<tr>
<td></td>
<td>Estimate of the exposure of “third doses” in European economic area (EEA) countries, per country and by age group.</td>
</tr>
<tr>
<td></td>
<td>Handling and dosing errors as result of different BNT162b2 formulations on the market.</td>
</tr>
<tr>
<td>WHO</td>
<td>Pregnancy outcome in clinical trials.</td>
</tr>
<tr>
<td></td>
<td>Data on low- and middle-income countries (LMICs) populations with HIV, malnutrition and tuberculosis and other infectious diseases.</td>
</tr>
<tr>
<td>Health</td>
<td>Review on the new variant “Omicron” and other variants.</td>
</tr>
<tr>
<td>Canada</td>
<td>Safety evaluation of tinnitus and hearing loss.</td>
</tr>
</tbody>
</table>

According to the European Risk Management Plan (EU-RMP) version 4.0 adopted on 26 November 2021, in effect at the beginning of the reporting period, safety concerns for BNT162b2 are:

• Important identified risks: Anaphylaxis; Myocarditis and Pericarditis

• Important potential risk: Vaccine-associated enhanced disease (VAED) including Vaccine-associated enhanced respiratory disease (VAERD)

• Missing information: Use in pregnancy and while breast feeding; Use in immunocompromised patients; Use in frail patients with co-morbidities (eg, chronic

\(^{15}\) This refers to the company core list of safety concerns. Myocarditis and pericarditis were already important identified risks in the EU-RMP, US-PVP and many country-level RMP addendums.
obstructive pulmonary disease [COPD], diabetes, chronic neurological disease, cardiovascular disorders); Use in patients with autoimmune or inflammatory disorders; Interaction with other vaccines; Long term safety data.

A summary of the EU-RMP versions and associated procedures, submitted during the reporting period and immediately after the PSUR DLP, are summarised in the table below. Version number of the EU-RMPs was agreed with EMA.

<table>
<thead>
<tr>
<th>Procedure description</th>
<th>Procedure number</th>
<th>Submitted EU-RMP</th>
<th>Approval date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reporting period</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Consolidation of RMPs version 2.6 and 4.0.</td>
<td>EMEA/H/C/005735/II/0087 and EMEA/H/C/005735/X/0077</td>
<td>Version 5.0, submitted on 10 March 2022</td>
<td>10 March 2022</td>
</tr>
<tr>
<td><strong>After the PSUR DLP</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line extension for COMIRNATY® 3 μg Concentrate for dispersion for injection for infants and children between 6 months to 4 years of age.</td>
<td>EMEA/H/C/005735/X/0138</td>
<td>Version 5.1, submitted on 08 July 2022</td>
<td>Ongoing procedure with pending approval.</td>
</tr>
<tr>
<td>Removal of the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP recommendation received on 10 March 2022 with the positive opinion for the Type II Variation 87 (EMEA/H/C/005735/II/0087).</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Variation 0140- To support the extension of the indication to ≥12 years of age to receive an additional booster (fourth) dose of bivalent Omicron-(BA.1) modified vaccine.</td>
<td>EMEA/H/C/005735/II/0140</td>
<td>Version 6.0, submitted on 19 July 2022</td>
<td>Ongoing procedure, with pending assessment.</td>
</tr>
<tr>
<td>To support the extension of the indication to introduce a first or second booster dose of a bivalent Omicron variant-modified vaccine, (BNT162b2 15 μg + BNT162b2 OMI BA.4/5 15 μg, total 30 μg), given ≥3 months after the primary series or ≥4 months after the third dose in individuals ≥12 years of age.</td>
<td>EMEA/H/C/005735/II/0143</td>
<td>Version 7.0, submitted on 15 August 2022</td>
<td>Ongoing procedure, with pending assessment.</td>
</tr>
</tbody>
</table>

In line with the above-mentioned update to the list of safety concerns in the EU-RMP v. 5.1, the MAH proposes to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period, because anaphylaxis is a known risk of vaccines that is understood by healthcare professionals who administer vaccines and patients and does not considerably impact the benefit/risk profile of the vaccine. Product labeling and standards of medical care during the vaccination procedure provide adequate risk mitigation.
After the DLP, an updated CDS (version 14.0) was made effective on 26 July 2022:

- to extend the indication to individuals 6 months of age and older in section 4.1 Therapeutic indications;
- to add the posology and method of administration for the Tris/Sucrose presentation 3 micrograms/dose in section 4.2 Posology and method of administration;
- to add a statement regarding the reporting rates of myocarditis and pericarditis after primary series and booster doses, based on accumulating data, in section 4.4 Special warnings and precautions for use;
- to add clinical data after 3 doses for children 2 through 4 years of age and for children 6 through 23 months of age, irritability and injection site tenderness as adverse drug reactions (ADRs), and myocarditis and pericarditis as ADRs post-authorisation experience in section 4.8 Undesirable effects;
- to add efficacy and immunogenicity data in individuals 6 months through 5 years of age in section 5.1 Pharmacodynamic properties;
- to add myocarditis and pericarditis as ADRs in Appendices A and B.

Risks have been evaluated in the context of the benefits of the vaccine. Based on the available safety and efficacy/effectiveness data from the reporting interval for BNT162b2, the overall benefit-risk profile of BNT162b2 remains favourable. No further changes to the BNT162b2 RSI or additional risk minimisation activities are warranted in addition to those above mentioned.
TABLE OF CONTENTS

EXECUTIVE SUMMARY ........................................................................................................2
LIST OF TABLES ..................................................................................................................13
LIST OF FIGURES ...............................................................................................................17
APPENDICES ......................................................................................................................18
LIST OF ABBREVIATIONS ..................................................................................................20
1. INTRODUCTION ............................................................................................................25
2. WORLDWIDE MARKETING APPROVAL STATUS .........................................................26
3. ACTIONS TAKEN IN THE REPORTING INTERVAL FOR SAFETY REASONS ...........28
4. CHANGES TO REFERENCE SAFETY INFORMATION ..................................................28
5. ESTIMATED EXPOSURE AND USE PATTERNS ..........................................................29
   5.1. Cumulative Subject Exposure in Clinical Trials .......................................................29
   5.2. Cumulative and Interval Patient Exposure from Marketing Experience ............30
6. DATA IN SUMMARY TABULATIONS .............................................................................39
   6.1. Reference Information ............................................................................................39
   6.2. Cumulative Summary Tabulations of Serious Adverse Events from Clinical Trials .........................................................................................................................39
   6.3. Cumulative and Interval Summary Tabulations from Post-Marketing Data Sources ........................................................................................................................................40
       6.3.1. General Overview ............................................................................................40
       6.3.1.1. General Overview of the Safety Database – All Cases ...............................44
       6.3.1.2. General Overview of the Safety Database - Unlocked Cases ..................92
7. SUMMARIES OF SIGNIFICANT SAFETY FINDINGS FROM CLINICAL TRIALS DURING THE REPORTING INTERVAL ..........................................................95
   7.1. Completed Clinical Trials ........................................................................................95
   7.2. Ongoing Clinical Trials ..........................................................................................96
   7.3. Long-term Follow-up ............................................................................................98
   7.4. Other Therapeutic Use of Medicinal Product .......................................................98
   7.5. New Safety Data Related to Fixed Combination Therapies ...............................98
8. FINDINGS FROM NON-INTERVENTIONAL STUDIES .................................................98
   8.1. Completed Non-Interventional Studies ................................................................98
   8.2. Ongoing Non-Interventional Studies ..................................................................100
9. INFORMATION FROM OTHER CLINICAL TRIALS AND SOURCES ....................101
9.1. Other Clinical Trials ................................................................................. 101
9.2. Medication Errors .................................................................................. 102

9.2.1. Errors pertaining to the new formulation of BNT162b2 – Paediatric
Tris/Sucrose Orange Cap presentation (dilute before use) 10 µg/dose for
5 to <12 years of age ................................................................................. 108

9.2.2. Errors pertaining to the new formulation of BNT162b2 – Adult
/Adolescent Tris/Sucrose Grey cap presentation (30 mcg/dose –[Do not
dilute]- in adults and children 12 years and older): .................................. 112

10. NON-CLINICAL DATA ............................................................................. 116
11. LITERATURE .......................................................................................... 116
12. OTHER PERIODIC REPORTS .................................................................. 119
13. LACK OF EFFICACY IN CONTROLLED CLINICAL TRIALS ............. 119
14. LATE-BREAKING INFORMATION ............................................................... 120
15. OVERVIEW OF SIGNALS: NEW, ONGOING, OR CLOSED ................. 121
16. SIGNAL AND RISK EVALUATION ............................................................ 123

16.1. Summary of Safety Concerns .................................................................. 123
16.2. Signal Evaluation .................................................................................. 125

16.2.1. Evaluation of Closed Signals .............................................................. 126
16.2.2. Signal Evaluation Plan for Ongoing Signals ........................................ 134

16.3. Evaluation of Risks and New Information ............................................. 134

16.3.1. Evaluation of Important Identified Risks ............................................ 134

16.3.1.1. Important Identified Risks – Anaphylaxis .................................... 135
16.3.1.2. Important Identified Risks – Myocarditis and Pericarditis .......... 138

16.3.2. Evaluation of Important Potential Risks ............................................. 191
16.3.3. Evaluation of Other Risks (not categorised as important) ................. 197

16.3.3.1. Adverse Events of Special Interest (AESIs) .................................. 197

16.3.3.2. Clinical Reactogenicity Data on Individuals Previously
exposed or not to SARS-CoV-2 ................................................................. 258

16.3.3.3. Local Adverse Reactions ............................................................... 260
16.3.3.4. Systemic Adverse Reactions ......................................................... 264
16.3.3.5. Severe Reactogenicity ................................................................. 273
16.3.3.6. Age-Related Adverse Reactions .................................................. 276
16.3.3.7. Vaccination Stress/Anxiety related ADRs ...................................... 281

16.3.4. Evaluation of Special Situations ......................................................... 285
16.3.4.1. Death ................................................................. 285
16.3.4.2. Overdose .......................................................... 294
16.3.4.3. Abuse, Misuse, and Drug Dependency .................. 296
16.3.4.4. Occupational Exposure ...................................... 299
16.3.4.5. Lack of Therapeutic Efficacy ................................. 300
16.3.4.6. Off-Label Use .................................................. 307
16.3.4.7. Unexpected Therapeutic Effect ............................ 310

16.3.5. Update on Special Patient Populations .......................... 312
16.3.5.1. Use in Elderly Patients ....................................... 312
16.3.5.2. Use in Paediatric Patients .................................... 315
16.3.5.3. Use in Pregnant/Lactating Women ......................... 328
16.3.5.4. Use in Patients with Comorbidities ....................... 340
16.3.5.5. Use in Immunocompromised Patients ...................... 343
16.3.5.6. Use in Patients with Autoimmune or Inflammatory Disorders .............................................................. 347
16.3.5.7. Use in Frail Patients with Comorbidities (e.g. COPD, Diabetes, Chronic Neurological Disease, Cardiovascular Disorders, active Tuberculosis) ................................. 351
16.3.5.8. Interactions with other Vaccines ......................... 355

16.4. Characterisation of Risks ............................................ 357
16.4.1. Characterisation of Important Identified and Potential Risks ................................................................. 357
16.4.1.1. Cumulative Characterisation of Important Identified Risks .................................................................. 358
16.4.1.2. Cumulative Characterisation of Important Potential Risks ................................................................. 361
16.4.2. Description of Missing Information ............................ 362

17. BENEFIT EVALUATION ......................................................... 364
17.1. Important Baseline Efficacy and Effectiveness Information 364
17.2. Newly Identified Information on Efficacy and Effectiveness 373
17.3. Characterisation of Benefits ........................................ 384

18. INTEGRATED BENEFIT-RISK ANALYSIS FOR APPROVED INDICATIONS ........................................ 384
18.1. Benefit-Risk Context – Medical Need and Important Alternatives ......................................................... 384
18.2. Benefit-Risk Analysis Evaluation .................................. 393
18.2.1. Benefits ................................................................. 393
18.2.2. Risks ................................................................. 393
18.2.3. Overall Benefit-Risk ........................................................................................................396

19. CONCLUSION AND ACTIONS .........................................................................................396
LIST OF TABLES

Table 1. Product Descriptiona .........................................................2
Table 2. List of PSURs .................................................................26
Table 3. Marketing Authorisation Holders of BNT162b2 ....................27
Table 4. Safety-Related Changes Made to the RSI After the DLP ........29
Table 5. Cumulative Estimated Shipped and Administered Doses of BNT162b2 by Region Worldwide ....................................................32
Table 6. Cumulative Administered Doses of BNT162b2 – License Partner Data .................................................................33
Table 7. Interval Estimated Shipped and Administered Doses of BNT162b2 by Region Worldwide .........................................................34
Table 8. Interval Administered Doses of BNT162b2 – License Partner Data .................................................................35
Table 9. EU/EEA – Cumulative Number of BNT162b2 Administered Doses by Age Group and Dose Number ........................................36
Table 10. EU/EEA – Cumulative Number of BNT162b2 Administered 3rd and 4th Doses by Age Group and Country ................................37
Table 11. Japan - Cumulative Number of BNT162b2 Administered Doses .................................................................38
Table 12. US - Cumulative Number of BNT162b2 Administered Doses .................................................................38
Table 13. EU/EEA – Interval Number of BNT162b2 Administered Doses by Age Group and Dose Number ........................................39
Table 14. Demographic Information - All Cases Received during the Reporting Interval .................................................................45
Table 15. Clinical Trial Data: Medical History and Co-Suspects ............53
Table 16. Clinical Trial Data: Serious Events Reported in ≥2% Cases ........55
Table 17. Post-Authorisation Data: Medical History and Co-Suspects ....62
Table 18. Post-Authorisation Data: Events Reported in ≥2% Cases ........63
Table 19. Demographic Information – Tris/Sucrose Cases (Orange and Grey Cap) Received during the Reporting Interval ..................74
Table 20. Demographic Information – Comparison of Paediatric (≤ 17 years) Tris/Sucrose (Grey and Orange Cap) versus Paediatric PBS/Sucrose Cases ......76
Table 21. Events Reported in ≥2% Cases - Comparison of Paediatric Tris/Sucrose (Grey and Orange Cap) versus PBS/Sucrose Cases .................77
Table 22. Summary of Approval of Booster Doses in the Reporting Period 79
Table 23. Selected Case Characteristics of CT Data Involving Participants Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022) .........................................81

CONFIDENTIAL
Page 13
Table 24. Selected Case Characteristics of Post-Authorisation Data Involving Individuals Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022) ...................................................... 82

Table 25. Comparison of clinical AEs reported in ≥2% Booster Dose(s) vs Primary Series Cases .......................................................... 90

Table 26. Most Frequently Reported Lot Numbers .................................................................................................................. 91

Table 27. Demographic Information - Unlocked Cases at the End of the Reporting Interval .......................................................... 92

Table 28. Post-Authorisation Data: Events Reported in ≥2% of Unlocked Cases ................................................................. 93

Table 29. Summary of Results from Clinical Trial Completed During the Reporting Period ................................................................. 95

Table 30. Summary of Results from Completed NIS During the Reporting Period .............................................................. 99

Table 31. Clinical Literature Articles that Presented New Safety Information in the Reporting Interval .......................................................... 116

Table 32. Overview of Signals (at DLP 18 June 2022) ...................................................................................................................... 121

Table 33. Ongoing Safety Concerns ........................................................................................................................................ 124

Table 34. Evaluation of Closed Signals During the Reporting Interval ..................................................................................... 126

Table 35. Signal Evaluation Plan for Ongoing Signals .............................................................................................................. 134

Table 36. Myocarditis in Subjects aged 5 – 11 Years (N=48) ........................................................................................................ 143

Table 37. Myocarditis in Subjects aged 12 – 15 Years (N=366) .................................................................................................... 145

Table 38. Myocarditis in Subjects aged 16 – 17 Years (N=345) .................................................................................................... 147

Table 39. Myocarditis in Subjects aged 18 – 24 Years (N=968) .................................................................................................... 150

Table 40. Myocarditis in Subjects aged 25 – 29 Years (N=519) .................................................................................................... 153

Table 41. Myocarditis in Subjects aged 30 – 39 Years (N=983) .................................................................................................... 157

Table 42. Myocarditis in Subjects aged ≥40 Years (N=1752) ....................................................................................................... 161

Table 43. Myocarditis in Subjects of Unknown Age (N=441) ......................................................................................................... 165

Table 44. Myocarditis in Subjects who Received a Booster dose .................................................................................................... 167

Table 45. Pericarditis in Subjects aged 5-11 years (N=30) .................................................................................................................. 170

Table 46. Pericarditis in Subjects aged 12-15 years (N=118) ....................................................................................................... 172

Table 47. Pericarditis in Subjects aged 16-17 years (N=106) ....................................................................................................... 174

Table 48. Pericarditis in Subjects aged 18-24 years (N=479) ....................................................................................................... 176

Table 49. Pericarditis in Subjects aged 25-29 years (N=417) ....................................................................................................... 178

Table 50. Pericarditis in Subjects aged 30-39 years (N=940) ....................................................................................................... 180

Table 51. Pericarditis in Subjects aged ≥ 40 years (N=1756) ....................................................................................................... 186

CONFIDENTIAL

Page 14
Table 52. Pericarditis in Subjects with Unknown Age (N=309) ........................................188
Table 53. Pericarditis in Subjects who Received a Booster Dose ....................................190
Table 54. Analysis of Systemic Adverse Reactions by Age Group, Event Seriousness and Event Outcome .................................................................268
Table 55. Analysis of Systemic Adverse Reactions by Presence of Comorbidities, Event Seriousness and Event Outcome .........................................................270
Table 56. Analysis of Systemic Adverse Reactions by Dose, Event Seriousness and Event Outcome .................................................................272
Table 57. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Adult Age Group Compared with Other Age Groups .................................................................278
Table 58. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Paediatric Group Compared with Other Age Groups .................................................................278
Table 59. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Elderly Age Group Compared with Other Age Groups .................................................................278
Table 60. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Adult Age Group Compared with Other Age Groups .................................................................280
Table 61. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Paediatric Group Compared with Other Age Groups .................................................................280
Table 62. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Elderly Age Group Compared with Other Age Groups .................................................................280
Table 63. Clinical Trial Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group during the Reporting Interval ........................................290
Table 64. Post-Authorisation Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group during the Reporting Interval ........................................290
Table 65. Clinical Trial Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group - Cumulative Reporting Interval ........................................292
Table 66. Post-Authorisation Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group - Cumulative Reporting Interval ........................................293
Table 67. Clinical Trial Data: Pregnancy Outcome - Cumulative Reporting Interval ........332
Table 68. Clinical Trial Data: Pregnancy Outcome during the Reporting Interval ............335
Table 69. Post-Authorisation Data: Pregnancy Outcome during the Reporting Interval* .................................................................339
Table 70. Cumulative Characterisation of Important Identified Risks ................................358
Table 71. Cumulative Characterisation of Important Potential Risks ........................................361
Table 72. Description of Missing Information ........................................................................363
Table 73. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Age Subgroup – Participants without Evidence of Infection and Participants with or without Evidence of Infection prior to 7 Days after Dose 2 – Evaluable Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period ........................................................................365
Table 74. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Risk Status – Participants Without Evidence of Infection* Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population During the Placebo-Controlled Follow-up Period ........................................366
Table 75. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Risk Status – Participants with or without* Evidence of Infection prior to 7 Days after Dose 2 – Evaluable Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period ........................................368
Table 76. Vaccine Efficacy – First Severe COVID-19 Occurrence in Participants with or without* prior SARS-CoV-2 Infection Based on FDA† or Centers for Disease Control and Prevention (CDC)‡ Definition after Dose 1 or from 7 Days after Dose 2 in the Placebo-Controlled Follow-up ........................................370
Table 77. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, Without Evidence of Infection and With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period, Adolescents 12 to 15 Years of Age Evaluable Efficacy (7 Days) Population ........................................................................372
Table 78. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2: Without Evidence of Infection Prior to 7 Days After Dose 2 – Phase 2/3 – Children 5 Through 11 Years of Age Evaluable Efficacy Population ..................................................375
Table 79. Summary of Geometric Mean Ratio for 50% Neutralizing Titer – Comparison of Children 5 Through Less Than 12 Years of Age (Study 3) to Participants 16 Through 25 Years of Age (Study 2) – Participants Without* Evidence of Infection up to 1 Month After Dose 2 – Dose 2 Evaluable Immunogenicity Population ..........................................................377
Table 80. Difference in Percentages of Participants With Seroreponse – Participants Without* Evidence of Infection up to 1 Month After Dose 2 – Immunobridging Subset – Phase 2/3 – Comparison of 5 Through <12 Years of Age to Study 2 Phase 2/3 16 Through 25 Years of Age – Evaluable Immunogenicity Population ..........................................................378
Table 81. Incidence, Prevalence, and Mortality of COVID-19 as of 05 April 2022 ...........386
Table 82. Summary of Important Risks ..............................................................................395
Table 83. Overall Benefit-Risk for BNT162b2 ..........................................................396
LIST OF FIGURES

Figure 1. Clinical Trial Data: Number of SAEs by Age Group and Gender ..............47
Figure 2. Clinical Trial Data: Case Outcome by Presence/Absence of Comorbidities ......49
Figure 3. Clinical Trial Data: Case Outcome by Presence/Absence of Comorbidities and Gender .................................................................50
Figure 4. Clinical Trial Data: Case Outcome by Age Group in Presence of Comorbidities .................................................................51
Figure 5. Clinical Trial Data: Case Outcome by Age Group in Absence of Comorbidities .................................................................52
Figure 6. Post-Authorisation Data: Number of Cases by Age Group and Gender ..........56
Figure 7. Post-Authorisation Data: Case Outcome by Presence/Absence of Comorbidities ...........................................................................58
Figure 8. Post-Authorisation Data: Case Outcome by Presence/Absence of Comorbidities and Gender ...........................................................59
Figure 9. Post-Authorisation Data: Case Outcome by Age Group in Presence of Comorbidities ...........................................................................60
Figure 10. Post-Authorisation Data: Case Outcome by Age Group in Absence of Comorbidities .................................................................61
Figure 11. Post-Authorisation Data: Event Seriousness of the PTs ≥2% of Cases ........66
Figure 12. Post-Authorisation Data: Clinical AEs reported in ≥2% of Cases by Age Group ...............................................................................67
Figure 13. Events Reported in ≥2% of All Clinical Trial Cases by Age Group ..........279
Figure 14. Events Reported in ≥2% of All Post-marketing Cases by Age Group ......281
APPENDICES

APPENDIX 1 REFERENCE INFORMATION (10 MAY 2022) ........................................ 397
APPENDIX 1.1 SAFETY RELATED CHANGES TO RS1 ........................................ 474
APPENDIX 1.2 CORE DATA SHEET (02 DECEMBER 2021) .............................. 494
APPENDIX 1.3 CORE DATA SHEET (21 DECEMBER 2021) ............................. 558
APPENDIX 1.4 CORE DATA SHEET (14 JANUARY 2022) ................................. 626
APPENDIX 1.5 CORE DATA SHEET (23 MARCH 2022) ..................................... 697
APPENDIX 2.1 CUMULATIVE SUMMARY TABULATION OF SERIOUS ADVERSE
EVENTS FROM CLINICAL TRIALS ................................................................. 769
APPENDIX 2.1.1 CUMULATIVE SUMMARY TABULATION OF SERIOUS ADVERSE
EVENTS FROM NON-PFIZER (LICENSING PARTNER)
CLINICAL TRIALS ......................................................................................... 816
APPENDIX 2.2 CUMULATIVE AND INTERVAL SUMMARY TABULATION OF
SERIOUS AND NON-SERIOUS ADVERSE REACTIONS FROM
POST-MARKETING DATA SOURCES ............................................................... 821
APPENDIX 2.3 CUMULATIVE CLINICAL TRIAL EXPOSURE WITH DEMOGRAPHIC
DATA ........................................................................................................... 1214
APPENDIX 2.3B CUMULATIVE CLINICAL TRIAL EXPOSURE WITH DEMOGRAPHIC
DATA FROM BIONTECH DEVELOPMENT PROGRAMS ............................... 1217
APPENDIX 2.3C CUMULATIVE CLINICAL TRIAL EXPOSURE WITH DEMOGRAPHIC
DATA FROM FOSUN DEVELOPMENT PROGRAMS ...................................... 1218
APPENDIX 2.3.1 CUMULATIVE CLINICAL TRIAL EXPOSURE WITH DEMOGRAPHIC
DATA FROM OTHER PFIZER DEVELOPMENT
PROGRAMS .................................................................................................. 1219
APPENDIX 3 TABULAR SUMMARY OF SAFETY SIGNALS................................. 1220
APPENDIX 3.1 SIGNAL EVALUATION ................................................................. NO DATA
APPENDIX 4.1 LISTING OF COMPLETED (CONCLUDED WITH FINAL CLINICAL
STUDY REPORT) INTERVENTIONAL SAFETY STUDIES ............................. NO DATA
APPENDIX 4.2 LISTING OF ONGOING (STARTED OR CONCLUDED WITH NO
FINAL CLINICAL STUDY REPORTS) INTERVENTIONAL
SAFETY STUDIES ....................................................................................... 1235
APPENDIX 4.3 LISTING OF COMPLETED (CONCLUDED WITH FINAL CLINICAL
STUDY REPORT) NON-INTERVENTIONAL SAFETY STUDIES ............. NO DATA
APPENDIX 4.4 LISTING OF ONGOING (STARTED OR CONCLUDED WITH NO
FINAL CLINICAL STUDY REPORTS) NON-INTERVENTIONAL SAFETY
STUDIES .................................................................................................... 1237
APPENDIX 5 LIST OF SOURCES OF INFORMATION USED TO PREPARE THE
PSUR (LITERATURE ABSTRACTS) ................................................................. 1243
APPENDIX 6A LIST OF REGULATORY AUTHORITIES REQUEST(S) ............... 1251
APPENDIX 6A.1 DIZZINESS CUMULATIVE REVIEW ......................................... 1261
APPENDIX 6A.2 ACQUIRED HAEMOPHILIA CUMULATIVE REVIEW .............. 1272

CONFIDENTIAL
Page 18
APPENDIX 6A.3 HEARING LOSS AND TINNITUS CUMULATIVE REVIEW ..........1289
APPENDIX 6A.4 MULTISYSTEM INFLAMMATORY SYNDROME ..................1318
APPENDIX 6A.5 AUTOIMMUNE HEPATITIS CUMULATIVE REVIEW ...........1345
APPENDIX 6B OBSERVED VERSUS EXPECTED ANALYSES FOR ADVERSE EVENTS OF SPECIAL INTEREST ..............................................................................................................1400
APPENDIX 6C.1 INTERVAL SUMMARY TABULATION OF FATAL REPORTS ......1470
APPENDIX 6C.2 CUMULATIVE SUMMARY TABULATION OF FATAL REPORTS .................................................................................................................................1516
APPENDIX 6D TABLES SUPPORTING CLINICAL REACTOGENICITY DATA ON INDIVIDUALS PREVIOUSLY EXPOSED OR NOT TO SARS-COV-2 ..............1599
APPENDIX 6.1 PROPOSED PRODUCT INFORMATION (REGIONAL APPENDIX) .................................................................................................................................NO DATA
APPENDIX 6.2 PROPOSED ADDITIONAL PHARMACOVIGILANCE AND RISK MINIMISATION ACTIVITIES (REGIONAL APPENDIX) ..................NO DATA
APPENDIX 6.3 SUMMARY OF ONGOING SAFETY CONCERNS (REGIONAL APPENDIX) ...................................................................................................................NO DATA
APPENDIX 6.4 REPORTING OF RESULTS FROM POST-AUTHORISATION SAFETY STUDIES (REGIONAL APPENDIX) .................................................................NO DATA
APPENDIX 6.5 EFFECTIVENESS OF RISK MINIMISATION (REGIONAL APPENDIX) ..........................................................................................................................NO DATA
APPENDIX 7 CUMULATIVE MARKETING AUTHORISATION STATUS.......NO DATA
APPENDIX 8 CHARACTERISATION OF IMPORTANT RISKS ...................1658
**LIST OF ABBREVIATIONS**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Term</th>
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<td>HLGOT</td>
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<td>HLT</td>
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<td>IBD</td>
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<td>IC</td>
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<td>International Council for Harmonisation; intracerebral haemorrhage</td>
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<tr>
<td>SE</td>
<td>Sweden</td>
</tr>
<tr>
<td>SFDA</td>
<td>Saudi Food and Drug Authority</td>
</tr>
<tr>
<td>SI</td>
<td>Slovenia</td>
</tr>
<tr>
<td>SIIIV</td>
<td>seasonal inactivated influenza vaccine</td>
</tr>
<tr>
<td>SK</td>
<td>Slovakia</td>
</tr>
<tr>
<td>SmPC</td>
<td>Summary of Product Characteristics</td>
</tr>
<tr>
<td>SMQ</td>
<td>standardised MedDRA Query</td>
</tr>
<tr>
<td>SOC</td>
<td>system organ class</td>
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<tr>
<td>SPEAC</td>
<td>Safety Platform for Emergency vACcines</td>
</tr>
<tr>
<td>SSNHL</td>
<td>sudden sensorineural hearing loss</td>
</tr>
<tr>
<td>SSR</td>
<td>summary safety report</td>
</tr>
<tr>
<td>TGA</td>
<td>Therapeutic Goods Administration</td>
</tr>
<tr>
<td>TME</td>
<td>targeted medical event</td>
</tr>
<tr>
<td>Tris</td>
<td>tromethamine</td>
</tr>
<tr>
<td>U</td>
<td>unknown</td>
</tr>
<tr>
<td>UK</td>
<td>United Kingdom</td>
</tr>
<tr>
<td>UMC</td>
<td>Uppsala Monitoring Centre</td>
</tr>
<tr>
<td>Unk</td>
<td>Unknown</td>
</tr>
<tr>
<td>US</td>
<td>United States</td>
</tr>
<tr>
<td>USG</td>
<td>United States Government</td>
</tr>
<tr>
<td>VACTERL</td>
<td>vertebral defects, anal atresia, cardiac defects, tracheo-esophageal fistula, renal anomalies, and limb abnormalities</td>
</tr>
<tr>
<td>VAED</td>
<td>vaccine associated enhanced disease</td>
</tr>
<tr>
<td>VAERD</td>
<td>vaccine associated enhanced respiratory disease</td>
</tr>
<tr>
<td>VAERS</td>
<td>Vaccine Adverse Event Reporting System</td>
</tr>
<tr>
<td>VE</td>
<td>vaccine efficacy</td>
</tr>
<tr>
<td>VLP</td>
<td>virus-like particle</td>
</tr>
<tr>
<td>VOC</td>
<td>variant of concern</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
</tbody>
</table>
1. INTRODUCTION

This is the 3rd PSUR for the COVID-19 mRNA vaccine (nucleoside modified), COMIRNATY®, also referred to as BNT162b2, covering the reporting interval 19 December 2021 through 18 June 2022.

The format and content of this PSUR is in accordance with the Guideline on GVP Module VII—Periodic safety update report (EMA/816292/2011 [December 2013]), with ICH Guideline E2C (R2) Periodic Benefit-Risk Evaluation Report [Step 5, January 2013]), corePSUR19 guidance (EMA/362988/2021 [08 July 2021]), and Consideration on core requirements for RMPs of COVID-19 vaccines - coreRMP19 guidance v. 3.0 (EMA/PRAC/73244/2022 [08 February 2022]).

BNT162b2 is highly purified single-stranded, 5' capped mRNA produced using a cell-free in vitro transcription from the corresponding DNA templates, encoding the viral S protein of SARS-CoV-2. The nucleoside-modified mRNA is formulated in LNPs, which enable delivery of the RNA into host cells to allow expression of the SARS-CoV-2 S antigen. The vaccine elicits both neutralising antibody and cellular immune responses to the S antigen, which may contribute to protection against COVID-19.

All the BNT162b2 formulations contain: ALC-0315, ALC-0159, DSPC, cholesterol, sucrose and water for injections.

The PBS/Sucrose presentation includes additionally potassium chloride, potassium dihydrogen phosphate, sodium chloride, disodium hydrogen phosphate dihydrate, as excipients. The Tris/Sucrose presentation includes additionally tromethamine, tromethamine hydrochloride as excipients.

BNT162b2 is indicated for active immunisation to prevent COVID-19 caused by SARS-CoV-2 virus, in individuals 5 years of age and older. It is administered intramuscularly in the deltoid muscle.

For individuals aged 12 years and older, the 2 presentations (PBS/Sucrose and Tris/Sucrose) are administered as 30 µg/dose intramuscularly as a primary series of 2 doses (0.3 mL each) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose (third dose) may be administered intramuscularly approximately 6 months after the second dose in individuals 16 years of age and older.

- PBS/Sucrose presentation – Purple cap: dilute before use.
- Tris/Sucrose presentation – Grey cap: do not dilute before use.

For individuals aged 5 through 11 years, the Tris/Sucrose presentation – Orange cap - is administered intramuscularly after dilution as a primary series of 2 doses (0.2 mL) at greater than or equal to 21 days (preferably 3 weeks) apart. A booster dose may be administered intramuscularly at least 6 months after the second dose.

On 17 June 2022, an additional formulation was approved first in the US: the paediatric Tris/Sucrose presentation - maroon cap (3 micrograms/dose) for individuals aged between 6
months and 4 years. This is a concentrate for dispersion for injection, to be administered after dilution intramuscularly in the anterolateral aspect of the thigh (or in the deltoid muscle in individuals 1 year of age and older) as a primary series of 3 doses (0.2 mL). The initial 2 doses are administered 3 weeks apart followed by a third dose administered at least 8 weeks after the second dose.

The list of the PSURs prepared for BNT162b2 is presented in Table 2.

<table>
<thead>
<tr>
<th>PSUR Number</th>
<th>Reporting Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>19 December 2020 through 18 June 2021</td>
</tr>
<tr>
<td>2</td>
<td>19 June 2021 through 18 December 2021</td>
</tr>
</tbody>
</table>

Pfizer is responsible for the preparation of the PSUR on behalf of license partners according to the Pharmacovigilance Agreement(s) in place. Data from respective license partner(s) are included in the report when applicable.

2. WORLDWIDE MARKETING APPROVAL STATUS

BNT162b2 received first temporary authorisation for emergency supply under Regulation 174 in the UK\textsuperscript{16} on 01 December 2020.

BNT162b2 received first regulatory conditional marketing authorisation approval for use in individuals 16 years and older in Switzerland on 19 December 2020. In the European Union, conditional marketing authorisation was granted on 21 December 2021.

BNT162b2 is authorised for the following formulations:

PBS/Sucrose – Purple cap 30 μg formulation:

- in individuals aged 16 years and older in 103 countries\textsuperscript{8} including full (5), conditional\textsuperscript{17} (49), EUA and other type of approvals\textsuperscript{18} (52).

- in individuals aged between 12 and 15 years in 81 countries\textsuperscript{9} including full (2), conditional (46), EUA and other type of approvals (34).

Tris/Sucrose formulation:

\textsuperscript{16} On 01 January 2021, conditional marketing authorisation approval was also granted in the UK and the approval is currently active.

\textsuperscript{17} Including temporary authorisations.

\textsuperscript{18} Including special import licenses.
- Grey cap: at the dosage of 30 μg formulation in individuals aged 12 years and older in 73 countries\textsuperscript{10} including full (3), conditional (44), EUA and other type of approvals (28).

- Orange cap: at the dosage of 10 μg formulation in individuals aged between 5 and 11 years in 79 countries\textsuperscript{9} including full (2), conditional (43), EUA and other type of approvals (35).

- Maroon cap: at the dosage of 3 μg formulation in individuals aged between 6 months and 4 years in the US with EUA\textsuperscript{19}.

- The booster dose has received approvals in 83 countries\textsuperscript{11} including full (3), conditional (46), EUA and other type of approvals (36).

Overall BNT162b2 is authorised in 104 countries/regions; in Table 3 the MAHs and the number of countries where the different MAHs hold the authorisation are presented.

### Table 3. Marketing Authorisation Holders of BNT162b2

<table>
<thead>
<tr>
<th>Marketing Authorisation Holder</th>
<th>Number of Countries/Regions Where the Marketing Authorisation is Held</th>
</tr>
</thead>
<tbody>
<tr>
<td>BioNTech</td>
<td>56</td>
</tr>
<tr>
<td>Pfizer</td>
<td>40</td>
</tr>
<tr>
<td>Fosun Pharma</td>
<td>2</td>
</tr>
<tr>
<td>Local MoH</td>
<td>3</td>
</tr>
<tr>
<td>Local Government</td>
<td>3</td>
</tr>
<tr>
<td>Hemas (LP)</td>
<td>1</td>
</tr>
<tr>
<td>Countries Where BNT162b2 is Authorised</td>
<td>104*</td>
</tr>
</tbody>
</table>

\textsuperscript{a} The sum of the number of the countries where the authorisation is held does not coincide with the total number of countries where BNT162b2 is authorised, because in the UK there are 2 different authorisations: the UK Government is the MAH of the EUA and BioNTech is the MAH of the conditional authorisation.

In addition, WHO had approved the EUL of BNT162b2.

The use of BNT162b2 in individuals aged 12 years and older is under EUA in Hong Kong and under a special import permit in Macau and Taiwan. In Hong Kong only the PBS/Sucrose – Purple cap formulation was approved.

There were no marketing authorisation withdrawals for safety reasons during the reporting interval.

\textsuperscript{19} On 17 June 2022, the paediatric Tris/Sucrose formulation for age 6 months through 4 years was approved first in the US.
3. ACTIONS TAKEN IN THE REPORTING INTERVAL FOR SAFETY REASONS

During the reporting period, no actions have been taken with respect to BNT162b2 for safety reasons, either by a HA or by the MAH.

Although not considered by definition a regulatory action taken for safety reasons because it does not significantly impact the benefit risk balance of use of the product in authorised populations, due to the receipt of spontaneous reports of Guillain-Barré syndrome (GBS) after vaccination with mRNA COVID-19 vaccines including BNT162b2, PMDA in Japan has required class changes to include GBS in the important precautions section of the Japan package insert\textsuperscript{20} and inclusion of GBS as an important potential risk in the Japan RMP\textsuperscript{21}. It should be noted that based on PMDA assessment, the frequency of reported cases of GBS was not significantly higher than the background incidence in any gender or age group and a mechanism is not known.

4. CHANGES TO REFERENCE SAFETY INFORMATION

The RSI for this PSUR is the COVID 19 mRNA vaccine CDS version 13.0 dated 10 May 2022, in effect at the end of the reporting period and included in Appendix 1.

The 4 previous CDS versions (version 9.0 dated 02 December 2021\textsuperscript{22}, version 10.0 dated 21 December 2021, version 11.0 dated 14 January 2022 and version 12.0 dated 23 March 2022\textsuperscript{22}), which were also in effect during the reporting period, are included in Appendix 1.2 through Appendix 1.5.

Safety-related changes included updates of the following sections:

- 4.2 Posology and method of administration (version 13.0),
- 4.8 Undesirable effects (versions 10.0, 11.0 and 13.0),
- 5.1 Pharmacodynamic properties (versions 10.0 and 11.0),
- Appendix A and Appendix B (version 10.0).

Safety-related changes to the RSI are presented in Appendix 1.1.

After the DLP, an updated CDS (version 14.0) was made effective on 26 July 2022; the safety-related changes are summarised in Table 4.

\textsuperscript{20} The Japan package insert was updated by the MAH during the current reporting period, on 10 June 2022.

\textsuperscript{21} Guillain-Barré syndrome was added as important potential risk to the safety concerns in the Japan RMP after DLP of this report on 22 June 2022.

\textsuperscript{22} This version of the CDS did not include any safety-related changes.
Table 4. Safety-Related Changes Made to the RSI After the DLP

<table>
<thead>
<tr>
<th>Section</th>
<th>Revision Type</th>
<th>Revision</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.1 Therapeutic indications</td>
<td>Update</td>
<td>Indication was updated to individuals 6 months of age and older.</td>
</tr>
<tr>
<td>4.2 Posology and method of administration</td>
<td>Addition</td>
<td>Posology and method of administration for the Tris/Sucrose presentation 3 micrograms/dose.</td>
</tr>
<tr>
<td>4.4 Special warnings and precautions for use</td>
<td>Addition</td>
<td>Statement regarding the reporting rates of myocarditis and pericarditis after primary series and booster doses, based on accumulating data.</td>
</tr>
</tbody>
</table>
| 4.8 Undesirable effects                      | Addition      | • Clinical data after 3 doses for children 2 through 4 years of age and for children 6 through 23 months of age  
|                                              |               | • Irritability and injection site tenderness as ADRs  
|                                              |               | • Myocarditis and pericarditis as ADRs post-authorisation experience. |
| 5.1 Pharmacodynamic properties              | Addition      | Efficacy and immunogenicity data in individuals 6 months through 5 years of age. |
| Appendices A and B                           | Addition      | Myocarditis and pericarditis as ADRs.                                     |

5. ESTIMATED EXPOSURE AND USE PATTERNS

5.1. Cumulative Subject Exposure in Clinical Trials

Cumulatively, 66,656 participants have participated in the BNT162b2 clinical development program comprising several clinical candidates, as outlined below:

BNT162b2: 59,260 participants of which
- 33,096 had received BNT162b2;
- 25,205 had received BNT162b2 post-unblinding and had received placebo before;
- 959 had received BNT162b2/placebo.

Variant vaccines based on BNT162b2: 1836 participants of which
- 747 had received BNT162b2 (B.1.351)\(^{23}\);
- 372\(^{24}\) had received BNT162b2 (B.1.617.2);
- 697 had received BNT162b2 (B.1.1.7 + B.1.617.2);
- 20 had received BNT162b2 (B.1.1.7).

\(^{23}\) BNT162b2 (B.1.351), which is also referred as BNT162b2s01 and BNT162b26A.

\(^{24}\) The number of participants exposed to variant vaccine B.1.617.2 is lower compared to the number reported in the 2nd PSUR, since the participants, who were administered an incorrect dose according to trial specific protocol, are not included.
Early development candidates: 633 participants of which
- 30 had received BNT162a1;
- 411 had received BNT162b1;
- 96 had received BNT162b3;
- 96 had received BNT162c2.

Blinded therapy: 7044 participants.

Placebo: 5871 participants.

Participant demographics data (e.g., age, gender, race) for ‘C459’ CTs is presented by treatment group in Appendix 2.3. Cumulative CT exposures with demographic data from BioNTech and Fosun CTs is presented in Appendix 2.3B and Appendix 2.3C.

Of note, BNT162b2 is also being utilised in another Pfizer clinical development program (B747): 372 participants received BNT162b2 as a study vaccine or as a comparator in the clinical study B747102625. Participant demographics data (e.g., age, gender, race) by treatment groups are presented in Appendix 2.3.1.

5.2. Cumulative and Interval Patient Exposure from Marketing Experience

In the current PSUR, the following regulatory requests about the exposure and number of third doses administered are addressed:

EMEA/H/C/005735/MEA/002.8 (9th SMSR), “The MAH should provide an estimate of the exposure of ‘third doses’ in future PSURs separately (reporting period and cumulatively), if applicable.”, and

EMEA/H/C/005735/MEA/002.10 (11th SMSR), “2. The MAH is requested to report the total number of administered Comirnaty dose 3 in the EU/EEA, per country, and by age group.”

Response

It is not possible to determine with certainty the number of subjects who received BNT162b2 during the period of this review, and this applies also to the “third doses”.

The total number of the BNT162b2 third doses administered, downloaded from the HA’s websites (EMA, PMDA and FDA) is provided in Table 9 through Table 13. Details for the cumulative number of third doses administered by age group and during the interval period in the EU/EEA countries are shown in Table 9 and in Table 13.

25 A phase 3, randomized, double blind trial to describe the safety and immunogenicity of 20 valent pneumococcal conjugate vaccine when co-administered with a booster dose of BNT162b2 in adults 65 years of age and older.
Cumulative exposure – MAH and LP Data

The worldwide number of shipped doses may serve as a reasonable indicator of subject exposure, considering that approximately 76% of the shipped doses were administered. This ratio represents the proportion of doses cumulatively administered (as per public available data for the EEA\(^{26}\) countries, the US\(^{27}\), and Japan\(^{28}\)) out of those cumulatively shipped (based on MAH data, according to the shipment tracker [Order Book]\(^{29}\)).

With these caveats in mind, it is estimated that:

Approximately 3,555,998,805 doses of BNT162b2 were shipped worldwide from the receipt of the first temporary authorisation for emergency supply on 01 December 2020 through 18 June 2022, corresponding to 2,693,922,584 estimated administered doses.\(^{30}\)

Overall, through 18 June 2022, a total of 143,844,450 adult Tris/Sucrose doses were shipped worldwide.

Overall, through 18 June 2022, a total of 229,269,400 paediatric Tris/Sucrose doses were shipped worldwide.\(^{31}\)

Cumulative worldwide estimated exposure by dose and region based on or extrapolated from internal data (number of shipped doses) is displayed in Table 5.

\(^{26}\) Approximately 73% of the doses shipped in the EU-EEA countries were administered; this proportion has been calculated considering, out of total number of vaccine doses distributed in the EU-EEA countries, the total number of vaccine doses administered as per report on https://qap.ecdc.europa.eu/public/extensions/COVID-19/vaccine-tracker.html#distribution-tab, as of 16 June 2022.

\(^{27}\) Approximately 77% of the doses shipped in the US were administered; this proportion has been calculated considering, out of total number of vaccine doses distributed in the US, the total number of vaccine doses administered as per report on https://covid.cdc.gov/covid-data-tracker/#vaccinations 17 June 2022.

\(^{28}\) Approximately 81% of the doses shipped in Japan were administered; this proportion this proportion has been calculated considering, out of total number of vaccine doses distributed in the US, the total number of vaccine doses administered as per report on https://www.kantei.go.jp/jp/headline/kansensho/vaccine.html, as of 20 June 2022.

\(^{29}\) The Order Book is the most accurate tracker of shipment used as data source for all the Regions and Countries; US shipment data not available in the Order Book were taken from the Order Management Dashboard and data for Hong Kong, Macau and Taiwan were provided by BioNTech.

\(^{30}\) License Partner data are not included in the reported amount.

\(^{31}\) This total does not include 285,800 doses shipped to US at the DLP of 18 June 2022; on 17 June 2022, FDA approved the Tris/Sucrose presentation - maroon cap (3 micrograms/dose) for individuals aged between 6 months and 4 years.
Table 5. Cumulative Estimated Shipped and Administered Doses of BNT162b2 by Region Worldwide

<table>
<thead>
<tr>
<th>Region/Country</th>
<th>% of Doses</th>
<th>Total Number of Shipped Doses</th>
<th>Total Number of Administered Doses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Europe</td>
<td>33.3</td>
<td>1184790735</td>
<td>873840606</td>
</tr>
<tr>
<td>European Union (27)</td>
<td>24.6</td>
<td>874223640</td>
<td>638183257</td>
</tr>
<tr>
<td>European Economic Area Countries (3)</td>
<td>0.4</td>
<td>12454785</td>
<td>9091993</td>
</tr>
<tr>
<td>Switzerland</td>
<td>0.3</td>
<td>10501650</td>
<td>7981254</td>
</tr>
<tr>
<td>UK</td>
<td>3.4</td>
<td>121752585</td>
<td>92531965</td>
</tr>
<tr>
<td>Other Countries*</td>
<td>3.6</td>
<td>126651915</td>
<td>96235455</td>
</tr>
<tr>
<td>Commonwealth of Independent States</td>
<td>1.1</td>
<td>39206160</td>
<td>29796682</td>
</tr>
<tr>
<td><strong>North America</strong></td>
<td><strong>14.7</strong></td>
<td><strong>523868135</strong></td>
<td><strong>402657330</strong></td>
</tr>
<tr>
<td>US</td>
<td>12.7</td>
<td>451754755</td>
<td>347851161</td>
</tr>
<tr>
<td>Canada</td>
<td>2.0</td>
<td>72111380</td>
<td>54806169</td>
</tr>
<tr>
<td><strong>Central and South America</strong>b</td>
<td><strong>14.4</strong></td>
<td><strong>510365375</strong></td>
<td><strong>387877685</strong></td>
</tr>
<tr>
<td>Asia</td>
<td>29.6</td>
<td>1051292420</td>
<td>812428536</td>
</tr>
<tr>
<td>Japan</td>
<td>7.6</td>
<td>268925940</td>
<td>217830011</td>
</tr>
<tr>
<td>Other Countries*</td>
<td>22.0</td>
<td>782366480</td>
<td>594598525</td>
</tr>
<tr>
<td>Oceania</td>
<td>2.3</td>
<td>81140220</td>
<td>61666567</td>
</tr>
<tr>
<td>Australia/New Zealand</td>
<td>2.3</td>
<td>80243250</td>
<td>60984870</td>
</tr>
<tr>
<td>Other Countries*</td>
<td>0.0</td>
<td>896970</td>
<td>681697</td>
</tr>
<tr>
<td>Africa*</td>
<td>5.8</td>
<td>204541920</td>
<td>155451859</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>100.0</strong></td>
<td><strong>3555998805</strong></td>
<td><strong>269392284</strong></td>
</tr>
</tbody>
</table>

a. Includes the non-EU countries (Albania, Andorra, Bosnia, Kosovo, Montenegro, North Macedonia, Serbia, Turkey and Vatican City) and the Commonwealth of Independent States (Armenia, Azerbaijan, Georgia, Kazakhstan, Kyrgyzstan, Moldova, Tajikistan, Ukraine and Uzbekistan).

b. Includes Antigua & Barbuda, Argentina, Bahamas, Barbados, Belize, Bolivia, Brazil, Chile, Colombia, Costa Rica, Dominica, Dominican Republic, Ecuador, El Salvador, Grenada, Guatemala, Guyana, Haiti, Honduras, Jamaica, Mexico, Nicaragua, Panama, Paraguay, Peru, St Kitts & Nevis, Saint. Lucia, Saint Vincent & the Grenadine, Suriname, Trinidad & Tobago and Uruguay.

c. Includes Bahrain, Bangladesh, Bhutan, Brunei, Cambodia, Indonesia, Iraq, Israel, Jordan, Korea, Kuwait, Laos, Lebanon, Malaysia, Maldives, Mongolia, Nepal, Oman, Pakistan, Palestine, Philippines, Qatar, Saudi Arabia, Singapore, Sri Lanka, Thailand, Timor-Leste, United Arab Emirates and Vietnam.

d. Includes Fiji, Nauru, Samoa, Solomon Islands, Tuvalu.


f. Out of these shipped doses, 35,240,400 doses were shipped for COVAX, 377,151,030 doses were shipped for USG Donation program and 41,807,550 doses were shipped for EC Donation program.

Out of the cumulative estimated shipped and administered doses, 1,948,639,685 and 1,480,966,161 respectively, were shipped to ROW (Non–EEA countries, Canada, Central and South America, Asian countries [excluding Japan], Oceania and Africa).

Cumulative LP (Fosun) data on the number of BNT162b2 doses administered in Hong Kong, Macau and Taiwan is provided in Table 6.
Table 6. Cumulative Administered Doses of BNT162b2 – License Partner Data

<table>
<thead>
<tr>
<th>Region/Country</th>
<th>Total Number of Administered Doses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asia</td>
<td>27314884</td>
</tr>
<tr>
<td>Hong Kong&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>10314451</td>
</tr>
<tr>
<td>Macau&lt;sup&gt;c,d&lt;/sup&gt;</td>
<td>256403</td>
</tr>
<tr>
<td>Taiwan&lt;sup&gt;c,d&lt;/sup&gt;</td>
<td>16744030</td>
</tr>
</tbody>
</table>

a. Cumulative through 18 June 2022
b. Conditional Authorisation under legislation 599K.
c. Special Import Permit.
d. Cumulative through 21 June 2022

Interval exposure – MAH and LP Data

Approximately 1,115,282,160 doses of BNT162b2 were shipped worldwide during the current reporting interval from 19 December 2021 through 18 June 2022, corresponding to 843,724,061 estimated administered doses.<sup>30,32</sup>

During the current reporting interval, a total of 143,844,450 adult Tris/Sucrose doses were shipped worldwide (this number coincides with the total doses shipped worldwide in the cumulative period).<sup>33</sup>

During the current reporting interval, a total of 182,231,200 paediatric Tris/Sucrose doses were shipped worldwide.

Interval worldwide estimated exposure by dose, and region based on or extrapolated from internal data (number of shipped doses) is displayed in Table 7.

---

<sup>32</sup> The same assumptions done for the ratio between the number of the doses shipped and the administered ones for cumulative data are applied also for interval data.

<sup>33</sup> First shipment of Adult Tris/Sucrose was in the US on 27 December 2021.
Table 7. Interval Estimated Shipped and Administered Doses of BNT162b2 by Region Worldwide

<table>
<thead>
<tr>
<th>Region/Country</th>
<th>% of Doses</th>
<th>Total Number of Shipped Doses</th>
<th>Total Number of Administered Doses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Europe</td>
<td>30.0</td>
<td>334826010</td>
<td>246162526</td>
</tr>
<tr>
<td>European Union (27)</td>
<td>24.5</td>
<td>273061770</td>
<td>199335092</td>
</tr>
<tr>
<td>European Economic Area Countries (3)</td>
<td>0.3</td>
<td>3779610</td>
<td>2759115</td>
</tr>
<tr>
<td>Switzerland</td>
<td>0.4</td>
<td>4978080</td>
<td>3783341</td>
</tr>
<tr>
<td>UK</td>
<td>2.2</td>
<td>24785520</td>
<td>18836995</td>
</tr>
<tr>
<td>Other Countries&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.5</td>
<td>16756740</td>
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</tr>
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</tr>
<tr>
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<td>8297986</td>
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<tr>
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<td>8.2</td>
<td>90935210</td>
<td>70020112</td>
</tr>
<tr>
<td>Canada</td>
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<td>17076150</td>
<td>12977874</td>
</tr>
<tr>
<td>Central and South America&lt;sup&gt;b&lt;/sup&gt;</td>
<td>13.5</td>
<td>150188600</td>
<td>114143356</td>
</tr>
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<td>Asia</td>
<td>36.0</td>
<td>401417480</td>
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<tr>
<td>Japan</td>
<td>6.3</td>
<td>70110180</td>
<td>56789246</td>
</tr>
<tr>
<td>Other Countries&lt;sup&gt;c&lt;/sup&gt;</td>
<td>29.7</td>
<td>331307300</td>
<td>251793548</td>
</tr>
<tr>
<td>Oceania</td>
<td>3.0</td>
<td>32918880</td>
<td>25018349</td>
</tr>
<tr>
<td>Australia/New Zealand</td>
<td>2.9</td>
<td>32122530</td>
<td>24413123</td>
</tr>
<tr>
<td>Other Countries&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.1</td>
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<td>605226</td>
</tr>
<tr>
<td>Africa&lt;sup&gt;e&lt;/sup&gt;</td>
<td>7.9</td>
<td>87919830</td>
<td>66819071</td>
</tr>
<tr>
<td>Total</td>
<td>100.0</td>
<td>1115282160</td>
<td>843724061</td>
</tr>
</tbody>
</table>

<sup>a</sup> Includes the non-EU countries (Albania, Bosnia, Kosovo, North Macedonia, Turkey and Vatican City) and the Commonwealth of Independent States (Armenia, Azerbaijan, Georgia, Kazakhstan, Kyrgyzstan, Moldova, Tajikistan, Ukraine and Uzbekistan).

<sup>b</sup> Includes Antigua & Barbuda, Argentina, Bahamas, Barbados, Belize, Bolivia, Brazil, Chile, Colombia, Costa Rica, Dominican Republic, Ecuador, El Salvador, Guatemala, Haiti, Honduras, Jamaica, Mexico, Panama, Paraguay, Peru, Saint Kitts & Nevis, Saint Lucia, Saint Vincent & Grenadine, Suriname, Trinidad & Tobago and Uruguay.

<sup>c</sup> Includes Bahrain, Bangladesh, Bhutan, Brunei, Cambodia, Indonesia, Iraq, Israel, Jordan, Korea, Kuwait, Laos, Lebanon, Malaysia, Maldives, Mongolia, Nepal, Oman, Pakistan, Palestine, Philippines, Qatar, Saudi Arabia, Singapore, Sri Lanka, Thailand, Timor-Leste, United Arab Emirates and Vietnam.

<sup>d</sup> Includes Fiji, Nauru, Samoa, Tonga, Tuvalu.

<sup>e</sup> Includes Angola, Benin, Botswana, Burkina Faso, Cabo Verde, Cameroon, Central Africa Republic, Chad, Comoros, Congo, Djibouti, Democratic Republic of Congo, Egypt, Eswatini, Ethiopia, Gabon, Gambia, Ghana, Ivory Coast, Kenya, Kiribati, Lesotho, Liberia, Libya, Madagascar, Malawi, Mali, Mauritania, Mauritius, Morocco, Namibia, Niger, Nigeria, Rwanda, Senegal, Seychelles, Sierra Leone, Somalia, South Africa, Sudan, Tanzania, Togo, Tunisia, Uganda and Zambia.

<sup>f</sup> Out of these shipped doses, 8,168,940 doses were shipped for COVAX, 186,024,630 doses were shipped for USG Donation program and 34,387,410 doses were shipped for EC Donation program.

During the reporting interval, out of the estimated shipped and administered doses, 677,395,390 and 514,820,496 respectively, were shipped and administered to the ROW.

Interval LP (Fosun) data on the number of BNT162b2 doses administered in Hong Kong, Macau and Taiwan is provided in Table 8 below.
Table 8. Interval Administered Doses of BNT162b2 – License Partner Data

<table>
<thead>
<tr>
<th>Region/Country</th>
<th>Total Number of Administered Doses (30 µg for 12 years and older)</th>
<th>Total Number of Administered Doses (10 µg for 5-11 years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asia</td>
<td>11259174</td>
<td>867539</td>
</tr>
<tr>
<td>Hong Kong&lt;sup&gt;a,b&lt;/sup&gt;</td>
<td>4202582</td>
<td>N/A</td>
</tr>
<tr>
<td>Macau&lt;sup&gt;c,d&lt;/sup&gt;</td>
<td>88514</td>
<td>N/A</td>
</tr>
<tr>
<td>Taiwan&lt;sup&gt;e&lt;/sup&gt;</td>
<td>6968078</td>
<td>867539</td>
</tr>
</tbody>
</table>

a. 19 December 2021 through 18 June 2022.
b. Conditional Authorisation under legislation 599K.
c. Special Import Permit. Only sum of COMIRNATY™ COVID-19 mRNA Vaccine (BNT162b2) 30 mcg and 10 mcg was announced by Macau Government.
d. 22 December 2021 through 21 June 2022
e. 15 December 2021 through 20 June 2022

Cumulative exposure – Health Authority Public Data

Cumulative data about the number of COMIRNATY® doses administered are published for EEA, Japan, and US in the respective Health Authorities’ websites; these data are provided in Table 9 through Table 12.

Table 9 displays the EEA published data with number of doses administered for each age group and by dose number.
Table 9. EU/EEA – Cumulative Number of BNT162b2 Administered Doses by Age Group and Dose Number

<table>
<thead>
<tr>
<th>Age Group</th>
<th>1st Dose</th>
<th>2nd Dose</th>
<th>Dose Unknown</th>
<th>3rd Dose(^a)</th>
<th>4th Dose(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 18 years(^d)</td>
<td>1363639</td>
<td>11718536</td>
<td>982</td>
<td>1885318</td>
<td>1839</td>
</tr>
<tr>
<td>0 – 4 years(^d)</td>
<td>6570</td>
<td>5512</td>
<td>0</td>
<td>123</td>
<td>0</td>
</tr>
<tr>
<td>5 – 9 years(^e)</td>
<td>2400265</td>
<td>1552704</td>
<td>101</td>
<td>1101</td>
<td>0</td>
</tr>
<tr>
<td>10 – 14 years(^e)</td>
<td>4509303</td>
<td>4075582</td>
<td>420</td>
<td>199566</td>
<td>107</td>
</tr>
<tr>
<td>15 – 17 years(^f)</td>
<td>3551465</td>
<td>3307482</td>
<td>704</td>
<td>410503</td>
<td>258</td>
</tr>
<tr>
<td>18 – 24 years(^g)</td>
<td>11371811</td>
<td>10563808</td>
<td>4035</td>
<td>5272098</td>
<td>13263</td>
</tr>
<tr>
<td>25 – 49 years(^g)</td>
<td>51444284</td>
<td>49059427</td>
<td>36983</td>
<td>25414999</td>
<td>112807</td>
</tr>
<tr>
<td>50 – 59 years(^g)</td>
<td>23719359</td>
<td>23084094</td>
<td>25646</td>
<td>14917699</td>
<td>115305</td>
</tr>
<tr>
<td>60 – 69 years(^g)</td>
<td>16347236</td>
<td>16155340</td>
<td>28333</td>
<td>16472372</td>
<td>508401</td>
</tr>
<tr>
<td>70 – 79 years(^g)</td>
<td>15638054</td>
<td>15485654</td>
<td>21790</td>
<td>15020989</td>
<td>843612</td>
</tr>
<tr>
<td>≥ 80 years(^g)</td>
<td>12162934</td>
<td>11939294</td>
<td>9463</td>
<td>10747352</td>
<td>955314</td>
</tr>
<tr>
<td>Age Unknown(^g)</td>
<td>80136</td>
<td>65263</td>
<td>28</td>
<td>18179</td>
<td>59</td>
</tr>
<tr>
<td>EEA – All(^h)</td>
<td>224378211</td>
<td>223231140</td>
<td>126250</td>
<td>151603079</td>
<td>9331517</td>
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</tbody>
</table>

\(^a\) Indicated as Dose Additional 1 in the ECDC webpage.
\(^b\) Indicated as Dose Additional 2 in the ECDC webpage.
\(^c\) Data from 19 countries.
\(^d\) Data from 13 countries.
\(^e\) Data from 17 countries.
\(^f\) Data from 18 countries.
\(^g\) Data from 27 countries.
\(^h\) Data from 30 countries.

Cumulative period up to 2022 week 24 (up to 19 June 2022) – Downloaded on 18 June 2022

Table 10 provides, as per EMEA/H/C/005735/MEA/002.8 (9th SMSR) and EMEA/H/C/005735/MEA/002.10 (11th SMSR) commitments, the cumulative total number of administered Comirnaty dose 3 (Dose additional 1 in the ECDC webpage) in EU/EEA, per country, and by age group. The table contains also data about Dose 4 (reported as Dose Additional 2).
Table 10. EU/EEA – Cumulative Number of BNT162b2 Administered 3rd and 4th Doses by Age Group and Country

<table>
<thead>
<tr>
<th>Age Group by Dose Countries</th>
<th>&lt;18 years</th>
<th>18 - 24 years</th>
<th>25 - 49 years</th>
<th>50 - 59 years</th>
<th>60 - 69 years</th>
<th>70 - 79 years</th>
<th>≥80 years</th>
<th>Age Unknown</th>
<th>ALL</th>
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<td>555191</td>
<td>5714</td>
<td>2488143</td>
<td>43055</td>
<td>1306086</td>
<td>48807</td>
<td>1195270</td>
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<tr>
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<td>280</td>
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<td>261509</td>
<td>560</td>
<td>391338</td>
<td>692</td>
<td>278826</td>
</tr>
</tbody>
</table>

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Page 37
Table 11 below shows the cumulative number of BNT162b2 dose administered in Japan.

Table 11. Japan - Cumulative Number of BNT162b2 Administered Doses

<table>
<thead>
<tr>
<th></th>
<th>1st Dose</th>
<th>2nd Dose</th>
<th>3rd Dose</th>
<th>4th Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>General population*</td>
<td>80859379</td>
<td>80268661</td>
<td>45024154</td>
<td>78384</td>
</tr>
<tr>
<td>Elderly</td>
<td>32248732</td>
<td>32160764</td>
<td>20349603</td>
<td>53808</td>
</tr>
<tr>
<td>Child (5 to &lt; 12 years)</td>
<td>1334886</td>
<td>1185593</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Medical workersb</td>
<td>6378205</td>
<td>5709228</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>All</td>
<td>87237584</td>
<td>85977889</td>
<td>45024154</td>
<td>78384</td>
</tr>
</tbody>
</table>

a. Including elderly and children for all doses. Starting from the 3rd dose, also includes medical workers.
b. Vaccinations for medical workers (1st and 2nd dose) was completed as of 30 July 2021. From the 3rd dose, medical workers are included in the general population.
c. Booster dose in children 5 to < 12 years is not approved in Japan.

Source: Government’s website where this data was downloaded: https://www.kantei.go.jp/jp/headline/kansensho/vaccine.html Download Date: June 20, 2022, 05:00 p.m. [JST]

Table 12 shows the cumulative number of BNT162b2 doses administered in the US.

Table 12. US - Cumulative Number of BNT162b2 Administered Doses

<table>
<thead>
<tr>
<th>Population</th>
<th>No. of Doses</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>349460399</td>
</tr>
<tr>
<td>Fully vaccinated (2 doses)</td>
<td>127603112</td>
</tr>
<tr>
<td>With a 1st booster dose</td>
<td>58521335</td>
</tr>
<tr>
<td>1st Booster dose with BNT162b2 after primary series with BNT162b2 (Homologous Dose Schedule)</td>
<td>49297112</td>
</tr>
<tr>
<td>1st Booster dose with BNT162b2 after primary series with Moderna (Heterologous Dose Schedule)</td>
<td>3156484</td>
</tr>
<tr>
<td>1st Booster dose with BNT162b2 after primary series with J&amp;J (Heterologous Dose Schedule)</td>
<td>2024045</td>
</tr>
<tr>
<td>1st Booster dose with BNT162b2 after primary series with other COVID-19 vaccines (Heterologous Dose Schedule)*</td>
<td>51232</td>
</tr>
<tr>
<td>1st Booster dose with BNT162b2 after primary series with unknown COVID-19 vaccines (Heterologous Dose Schedule)</td>
<td>3992462</td>
</tr>
<tr>
<td>With a 2nd booster dose</td>
<td>8756788</td>
</tr>
</tbody>
</table>

a. Not BNT162b2, Moderna or J&J vaccine.


Interval exposure – Health Authority Public Data

Interval data about the number of COMIRNATY® doses administered are available only for the EEA countries.

Table 13 displays, as per EMEA/H/C/005735/MEA/002.8 (9th SMSR) and EMEA/H/C/005735/MEA/002.10 (11th SMSR) commitments, the interval data with number of doses administered for each age group and by dose number in the EEA countries.
Table 13. EU/EEA – Interval Number of BNT162b2 Administered Doses by Age Group and Dose Number

<table>
<thead>
<tr>
<th>Age Group</th>
<th>1st Dose</th>
<th>2nd Dose</th>
<th>Dose Unknown</th>
<th>3rd Dose</th>
<th>4th Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 18 years</td>
<td>4421971</td>
<td>4233201</td>
<td>611</td>
<td>1833167</td>
<td>1821</td>
</tr>
<tr>
<td>0 – 4 years</td>
<td>5374</td>
<td>5052</td>
<td>0</td>
<td>110</td>
<td>0</td>
</tr>
<tr>
<td>5 – 9 years</td>
<td>2033874</td>
<td>1534791</td>
<td>101</td>
<td>1055</td>
<td>0</td>
</tr>
<tr>
<td>10 – 14 years</td>
<td>1262611</td>
<td>1665408</td>
<td>257</td>
<td>192672</td>
<td>101</td>
</tr>
<tr>
<td>15 – 17 years</td>
<td>160024</td>
<td>272676</td>
<td>290</td>
<td>383121</td>
<td>254</td>
</tr>
<tr>
<td>18 – 24 years</td>
<td>366151</td>
<td>599222</td>
<td>785</td>
<td>4410546</td>
<td>13009</td>
</tr>
<tr>
<td>25 – 49 years</td>
<td>1067139</td>
<td>1916394</td>
<td>3954</td>
<td>17464295</td>
<td>110482</td>
</tr>
<tr>
<td>50 – 59 years</td>
<td>318635</td>
<td>580614</td>
<td>1763</td>
<td>8151219</td>
<td>113819</td>
</tr>
<tr>
<td>60 – 69 years</td>
<td>262799</td>
<td>474033</td>
<td>1789</td>
<td>5777707</td>
<td>506630</td>
</tr>
<tr>
<td>70 – 79 years</td>
<td>163055</td>
<td>285448</td>
<td>993</td>
<td>2887858</td>
<td>842320</td>
</tr>
<tr>
<td>≥ 80 years</td>
<td>146027</td>
<td>229916</td>
<td>661</td>
<td>1652184</td>
<td>934068</td>
</tr>
<tr>
<td>Age Unknown</td>
<td>18890</td>
<td>12175</td>
<td>11</td>
<td>14127</td>
<td>11</td>
</tr>
<tr>
<td>EEA – ALF</td>
<td>5369310</td>
<td>10625902</td>
<td>9945</td>
<td>69607125</td>
<td>9310883</td>
</tr>
</tbody>
</table>

a. Indicated as Dose Additional 1 in the ECDC webpage.
b. Indicated as Dose Additional 2 in the ECDC webpage.
c. Data from 19 countries.
d. Data from 13 countries.
e. Data from 17 countries.
f. Data from 18 countries.
g. Data from 27 countries.
h. Data from 16 countries.
i. Data from 30 countries.

Interval reporting period including 2021, week 51 through 2022 week 24 (up to 19 June 2022) – Downloaded on 18 June 2022.

Currently there are no available public data that allow to estimate the COMIRNATY®
exposure by gender.

6. DATA IN SUMMARY TABULATIONS

6.1. Reference Information

The MedDRA version 25.0 has been used to code adverse events/reactions in summary tabulations.

6.2. Cumulative Summary Tabulations of Serious Adverse Events from Clinical Trials

Appendix 2.1 provides a cumulative summary tabulation, from the MAH’s safety database, of SAEs reported in Pfizer clinical trial cases received by the MAH. This appendix is organised according to MedDRA SOC. This appendix includes SAEs originated from the following studies: C4591001, C4591005, C4591007, C4591015, C4591017, C4591020, C4591024, C4591030 and C4591031.

Appendix 2.1.1 provides a cumulative summary tabulation, from the MAH’s safety database, of SAEs reported in BioNTech and Fosun clinical trial cases. This appendix includes SAEs
originated from the following studies: BNT162-01, BNT162-03, BNT162-04, BNT162-06, BNT162-14 and BNT162-17.

6.3. Cumulative and Interval Summary Tabulations from Post-Marketing Data Sources

Appendix 2.2 provides a cumulative and interval summary tabulation of adverse drug reactions by PT from post-marketing sources. This tabulation includes serious and non-serious reactions from spontaneous sources, as well as serious adverse reactions from non-interventional studies and other non-interventional solicited sources. The cumulative data include all data up to 18 June 2022 and the interval data are from 19 December 2021 to 18 June 2022. This appendix is organised according to SOC and presents data for spontaneous cases (including regulatory authority and literature cases) separately from non-interventional sources.

Please note that adverse event totals presented for safety topic evaluations in Section 16 Signal and Risk Evaluation, may differ from Appendix 2.2 totals, due to the fact that Appendix 2.2 only displays the number of serious reactions from non-interventional studies and solicited sources as described above, whereas the safety topic evaluation includes all reported events. Cases from non-interventional studies and other non-interventional solicited sources must contain at least 1 serious related event to meet PSUR inclusion criteria and may also contain additional events that are considered unrelated, all of which would be evaluated.

6.3.1. General Overview

The list of regulatory commitments received from EMA (included in the PRAC ARs on PSURs, SMMRs/SBSRs or signal assessments), WHO and Health Canada to be addressed in the PSUR is detailed below.

Responses are provided in Appendix 6A and in the relevant sections/appendices cross-referenced below, apart from the response to request 5 of the AR of the 2nd PSUR (EMEA/H/C/PSUSA/00010898/202112) that is included in the eCTD sequence with the submission of the current PSUR.

As part of the PSURs AR, the PRAC requested the MAH to address the following requests:

- EMEA/H/C/PSUSA/00010898/202112 (2nd PSUR - reporting period 19 June 2021 through 18 December 2021)

1. The MAH is requested to continue to report on the number of processed cases downloaded from EudraVigilance in the PSUR.

2. The MAH is requested to re-assess the need for continuing the follow-up questionnaires anaphylaxis and VAED/VAERD and provide process data (e.g., response rate, extent of additional information collected) separately for cases reporting anaphylaxis and cases reporting VAED/VAERD, if applicable.

3. The MAH is requested to provide a cumulative review of cases reporting dizziness after Comirnaty exposure outside the context of anxiety/stress-related reactions (as already labelled in the Comirnaty SmPC section 4.4) and a discussion on the need to
add dizziness (including a proposal for the frequency of occurrence) to the ADR table of the Comirnaty SmPC section 4.8, as applicable. (Appendix 6A.1)

4. The MAH is requested to provide a cumulative review of cases reporting acquired haemophilia, including details of the underlying condition(s), time to onset, duration, outcome and an assessment of the causal relationship with the vaccination. The MAH should also provide a comprehensive discussion of the literature, mechanism of action and a discussion on the need to update the product information of Comirnaty, if applicable. (Appendix 6A.2)

5. The MAH is requested to provide a cumulative review of cases reporting IgA nephropathy, including details of the underlying condition(s), time to onset, duration, outcome and an assessment of the causal relationship with the vaccination. The MAH should also provide a comprehensive discussion of the literature, mechanism of action and a discussion on the need to update the product information of Comirnaty, if applicable.

- EMEA/H/C/PSUSA/00010898/202106 (1\textsuperscript{st} PSUR - reporting period 19 December 2020 through 18 June 2021)

Of concern are the backlog cases and the impact thereof on the O/E analyses. Besides the O/E analyses that include the processed cases, no sensitivity O/E analysis is presented which include the processed cases plus the backlog cases. In future PSURs and similar to the O/E analyses reported in the MSSRs, the MAH is requested to perform overall O/E analyses using at least no risk window, 14-day risk window and 21-day risk window and additionally perform sensitivity O/E analyses which include the processed cases plus the backlog cases. (Appendix 6B)

As part of the SMSR/SBSR assessment reports, the PRAC requested:

- EMEA/H/C/005735/MEM/002.8 (9\textsuperscript{th} SMSR)

The MAH should provide an estimate of the exposure of "third doses" in future PSURs separately (reporting period and cumulatively), if applicable. (Section 5.2 Cumulative and Interval Patient Exposure from Marketing Experience)

- EMEA/H/C/005735/MEM/002.10 (11\textsuperscript{th} SMSR)

1. The MAH should report on handling and dosing errors as a result of the different Comirnaty formulations on the market. (Section 9.2 Medication Errors)

2. The MAH is requested to report the total number of administered Comirnaty dose 3 in the EU/EEA, per country, and by age group. (Section 5.2 Cumulative and Interval Patient Exposure from Marketing Experience)
• EMA/PRAC/202255/2022 (13th SSR-2nd SBSR)

1. The MAH is requested to discuss the following publications regarding sudden sensorineural hearing loss (SSNHL) in association with COVID-19 vaccination:


Furthermore, the MAH is requested to conduct age-stratified O/E analyses for the AESI of sudden hearing loss using the age-specific background incidence rates of SSNHL reported in the following publication: Alexander T and Harris J. Incidence of Sudden Sensorineural Hearing Loss. Otol Neurotol. 2013 Dec; 34(9): 1586-9. doi:10.1097/MAO.0000000000000222. (Appendix 6A.3)

3. The MAH is requested in future SSRs and PSURs to present all relevant literature concerning the safety of Comirnaty (also besides the database Medline and Embase) that is published during the reporting period.

4. The MAH should continue to closely monitor MIS-C/A as outlined in PRAC’s signal recommendation (EPIT 19732). All new cases of MIS should be reported in the SSRs and PSURs. (Appendix 6A.4)

• EMA/PRAC/577594/2022 (14th SSR – 3rd SBSR)

1. In not (yet) peer-reviewed retrieved relevant literature, the MAH is requested to assess the used study methods to determine if the study results are valid or not, for further characterisation of a particular safety issue (e.g., myocarditis and pericarditis). (Appendix 6A.3)

2. The MAH is requested to perform a cumulative review on the association between sudden sensorineural hearing loss and Comirnaty exposure, including a review of the relevant new literature published after the reporting period of the 14th SSR, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable. (Appendix 6A.3)

3. In general the MAH should perform more effort in presenting/evaluating the cases that they have considered to be confounded and should present the risk factors for developing the respective conditions. If patient groups/patients with underlying disease have a higher risk of developing the condition this information is important to communicate. (NO) (Section 16.3.1 Evaluation of Important Identified Risks,
Section 16.3.2 Evaluation of Important Potential Risks, Section 16.3.4.1 Death and Section 16.3.5.2 Use in Paediatric Patients

As per signal assessment reports, the PRAC requested:

- Signal assessment report on Glomerulonephritis and nephrotic syndrome with tozinameran EMA/PRAC/416198/2021 – EPITT 19722

Having considered the available evidence from the cumulative review submitted by the Marketing Authorisation Holder (MAH), the PRAC has agreed that the MAH of the COVID-19 mRNA vaccine (nucleoside-modified) COMIRNATY® (BioNTech Manufacturing GmbH) should closely monitor the issue of 'glomerulonephritis/nephrotic syndrome', including exacerbations, and present a cumulative review of cases from all sources and relevant literature in the upcoming PSUR submissions. However, if new relevant information becomes available earlier that would support an association with the vaccine, the MAH should propose updates of the product information accordingly and without delay.


The MAH should continue to closely monitor this safety issue and new cases of MIS-C/A should be reported in the MSSRs and PSURs. A dedicated questionnaire should be implemented to retrieve an appropriate level of information to facilitate the assessment of the cases. The MAH should focus on the well described index case(s) and less on quantity and numbers. A few well described cases may be sufficient in our opinion to indicate a causal association for a very rare serious event. (Appendix 6A.4)

- Signal assessment on Autoimmune hepatitis with tozinameran EMA/PRAC/632042/2021 - EPITT 19749

The MAH should provide in the next PSUR (submission date 27 August 2022) a cumulative review of all cases of autoimmune hepatitis, including any relevant new data, from all available sources. The cumulative review should include, but not be limited to, data from clinical trials, post-marketing cases and any relevant articles from literature, using a data lock-point as recent as possible. (Appendix 6A.5)

WHO approval letter for the emergency use of Tozinameran - COVID-19 mRNA vaccine (nucleoside modified) - COMIRNATY®

The MAH was requested to provide additional data on vaccine immunogenicity, effectiveness and safety on population groups represented in Low- and Middle-Income Countries (LMICs) including individuals with conditions such as malnutrition and populations with existing co-morbidities such as tuberculosis, human immunodeficiency virus (HIV) infection and other high prevalent infectious diseases. (Section 16.3.3.1.21 AESIs in subjects with Malnutrition; HIV infection)
The MAH was requested to present the outcome of the cases of pregnancy observed in the clinical studies. (Section 16.3.5.3 Use in Pregnant/Lactating Women)

Health Canada (29 November 2021)

Future updates of the Cumulative Review on the new variant “Omicron” and other variants should be included in the Monthly Summary Safety Report (MSSRs) and the Periodic Benefit-Risk Evaluation Reports (PBRER) for Comirnaty Pfizer-BioNTech COVID-19 Vaccine (tozinameran). (Section 16.3.4.5 Lack of Therapeutic Efficacy and Section 17.2 Newly Identified Information on Efficacy and Effectiveness)

Health Canada (31 May 2022 – 2nd SBSR AR)

Tinnitus and hearing loss: The WHO recently published an update regarding COVID-19 vaccines and hearing loss. Signal detection activities at the UMC up to 22 Feb 2022 retrieved 164 cases with HLT Hearing losses (142 cases for Comirnaty) and 367 cases with the PT Tinnitus (293 for Comirnaty) with COVID-19 vaccines. Based on well documented cases, alternative causes were not identified for most of the patients and a plausible mechanism of action has been suggested. As such, provide a cumulative review of all cases of tinnitus and hearing loss. This cumulative review should include analyses of all cases, stratified by age, gender, doses administered, time to onset, and any other relevant information. An observed-to-expected analysis should be provided including the appropriate risk window. An appropriate case definition including a causality assessment should also be provided. (Appendix 6A.3)

6.3.1.1. General Overview of the Safety Database – All Cases

As per PRAC assessment report of the 2nd PSUR (procedure EMEA/H/C/PSUSA/00010898/202112):

1. The MAH is requested to continue to report on the number of processed cases downloaded from EudraVigilance in the PSUR.”

Response

Please refer to Appendix 6A.

General Overview – All Cases

A total of 508,351 case reports (668 from CT34 and 507,683 from PM) containing 1,597,673 events fulfilled criteria for inclusion in this PSUR reporting period, compared to 658,249 case

34 Clinical Trials cases include:

- 630 cases originated from 6 interventional trials (C4591001, C4591001-OPENLABEL, C4591007, C4591007-OPENLABEL, C4591015, C4591024, C4591030, C4591031, C4591031-OPENLABEL) for which BioNTech is the Sponsor and Pfizer acts as lead development party.
- 17 cases from 2 BioNTech interventional trials (BNT162-14 and BNT162-17), and
reports retrieved in the PSUR #2. Refer to Appendix 2.1 and Appendix 2.1.1 for the cumulative summary tabulation of all CT cases and to Appendix 2.2 for the summary tabulation of all PM cases received during the current reporting period and cumulatively.

Demographic information of all cases included in the safety database and received during the reporting interval are shown in Table 14.

**Table 14. Demographic Information - All Cases Received during the Reporting Interval**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>All No. of Cases (%)</th>
<th>CT No. of Cases (%)</th>
<th>PM No. of Cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EudraVigilance Cases</td>
<td>N=508,351*</td>
<td>N=668</td>
<td>N=507,683*</td>
</tr>
<tr>
<td>Yes</td>
<td>309,455</td>
<td>0</td>
<td>309,455</td>
</tr>
<tr>
<td>No</td>
<td>303,769 (59.8)</td>
<td>668 (100)</td>
<td>203,914 (40.2)</td>
</tr>
<tr>
<td>Country of occurrence</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Germany*</td>
<td>114,573 (22.5)</td>
<td>27 (40)</td>
<td>114,546 (22.6)</td>
</tr>
<tr>
<td>Austria*</td>
<td>55,474 (10.9)</td>
<td>0 (0)</td>
<td>55,474 (10.9)</td>
</tr>
<tr>
<td>Netherlands*</td>
<td>38,790 (7.6)</td>
<td>0 (0)</td>
<td>38,790 (7.6)</td>
</tr>
<tr>
<td>France*</td>
<td>38,418 (7.6)</td>
<td>0 (0)</td>
<td>38,418 (7.6)</td>
</tr>
<tr>
<td>UK*</td>
<td>36,833 (7.2)</td>
<td>1 (0.1)</td>
<td>36,832 (7.3)</td>
</tr>
<tr>
<td>Australia*</td>
<td>23,189 (4.6)</td>
<td>0 (0)</td>
<td>23,189 (4.6)</td>
</tr>
<tr>
<td>US*</td>
<td>22,605 (4.4)</td>
<td>425 (63.6)</td>
<td>22,180 (4.4)</td>
</tr>
<tr>
<td>Philippines*</td>
<td>17,981 (3.5)</td>
<td>0 (0)</td>
<td>17,981 (3.5)</td>
</tr>
<tr>
<td>Sweden*</td>
<td>16,862 (3.3)</td>
<td>0 (0)</td>
<td>16,862 (3.3)</td>
</tr>
<tr>
<td>Italy*</td>
<td>12,201 (2.4)</td>
<td>0 (0)</td>
<td>12,201 (2.4)</td>
</tr>
<tr>
<td>Japan*</td>
<td>12,080 (2.4)</td>
<td>0 (0)</td>
<td>12,080 (2.4)</td>
</tr>
<tr>
<td>Norway*</td>
<td>10,868 (2.1)</td>
<td>0 (0)</td>
<td>10,868 (2.1)</td>
</tr>
<tr>
<td>Malaysia*</td>
<td>10,009 (2.0)</td>
<td>0 (0)</td>
<td>10,009 (2.0)</td>
</tr>
<tr>
<td>Other countries</td>
<td>98,468 (19.4)</td>
<td>215 (32.2)</td>
<td>98,253 (19.4)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>324,059 (63.7)</td>
<td>304 (45.5)</td>
<td>323,755 (63.8)</td>
</tr>
<tr>
<td>Male</td>
<td>149,371 (29.4)</td>
<td>360 (53.9)</td>
<td>149,011 (29.4)</td>
</tr>
<tr>
<td>Unknown/No Data</td>
<td>34,921 (6.9)</td>
<td>4 (0.6)</td>
<td>34,917 (6.9)</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>447,034</td>
<td>649</td>
<td>446,385</td>
</tr>
<tr>
<td>Min-Max</td>
<td>0.01-120</td>
<td>0.58-87</td>
<td>0.01-120</td>
</tr>
<tr>
<td>Mean</td>
<td>42.5</td>
<td>49.8</td>
<td>42.5</td>
</tr>
<tr>
<td>Median</td>
<td>41</td>
<td>57</td>
<td>41</td>
</tr>
<tr>
<td>Age Range</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 17 years</td>
<td>33,236 a (6.5)</td>
<td>117 (17.5)</td>
<td>33,119 a (6.5)</td>
</tr>
<tr>
<td>[31,927] a</td>
<td>[102] a</td>
<td>[31,825] a</td>
<td></td>
</tr>
<tr>
<td>0 to 27 days</td>
<td>145 (0.03)</td>
<td>9 (1.3)</td>
<td>136 (0.03)</td>
</tr>
</tbody>
</table>

- 21 cases from 1 Fosun (BioNTech License Partner) interventional trial (BNT162-06) with BioNTech third party acting as lead development party.
- After DLP, an additional case originated from the BioNTech interventional trial BNT162-14 was identified, but it is not included among the CT cases, because the Case Report Type was erroneously reported as non-MAH sponsored interventional study. This case is from a 66-year-old male participant, who developed atrial fibrillation 175 days after vaccination with BNT162b2101. The SAE resolved and was assessed as unrelated to the vaccine by the investigator and the Sponsor.
Table 14. Demographic Information - All Cases Received during the Reporting Interval

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>All No. of Cases (%) N=508,351</th>
<th>CT No. of Cases (%) N=668</th>
<th>PM No. of Cases (%) N=507,683</th>
</tr>
</thead>
<tbody>
<tr>
<td>28 days to 23 months</td>
<td>1071 (0.2) 102 [98641] e</td>
<td>28 (4.2) 22 [65] e</td>
<td>1043 (0.2) [80] e</td>
</tr>
<tr>
<td>2-11 years</td>
<td>10,044 (2.0) 1044 [98641] e</td>
<td>65 (9.7) 65 [65] e</td>
<td>9979 (2.0) [9799] e</td>
</tr>
<tr>
<td>12-17 years</td>
<td>21,976 (4.3) 21,976 [21,958] e</td>
<td>15 (2.2) 15 [15] e</td>
<td>21,961 (4.3) [21,943] e</td>
</tr>
<tr>
<td>18-30 years</td>
<td>91,380 (18.0) 91,380 [21,958] e</td>
<td>37 (5.5) 37 [37] e</td>
<td>91,343 (18.0) [21,943] e</td>
</tr>
<tr>
<td>31-50 years</td>
<td>183,209 (36.0) 183,209 [183,209]</td>
<td>128 (19.2) 128 [128] e</td>
<td>183,081 (36.1) [183,079] e</td>
</tr>
<tr>
<td>51-64 years</td>
<td>86,885 (17.1) 86,885 [86,885]</td>
<td>171 (25.6) 171 [171] e</td>
<td>86,714 (17.1) [86,705] e</td>
</tr>
<tr>
<td>≥ 75 years</td>
<td>21,598 (4.2) 21,598 [21,598]</td>
<td>79 (11.8) 79 [79] e</td>
<td>21,519 (4.2) [21,509] e</td>
</tr>
<tr>
<td>Unknown</td>
<td>56,648 (11.1) 56,648 [56,648]</td>
<td>1 (0.1) 1 [1] e</td>
<td>56,647 (11.2) [56,637] e</td>
</tr>
<tr>
<td>N/A</td>
<td>194 (&lt;0.1) 194 [194]</td>
<td>3 (0.4) 3 [3] e</td>
<td>191 (&lt;0.1) 191 [191]</td>
</tr>
<tr>
<td>Case Seriousness</td>
<td>Serious 152,093 (29.9) 152,093 [152,093] 668 (100) 668 [668] 151,425 (29.8) 151,425 [151,425]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Non-serious 356,258 (70.1) 356,258 [356,258] 0 (0) 0 [0] 356,258 (70.2) 356,258 [356,258]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Case Outcome</td>
<td>Fatal 3,280 (0.6) 3,280 [3,280] 35 (5.2) 35 [35] 3,245 (0.6) 3,245 [3,245]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Recovered/ Recovering 178,812 (35.2) 178,812 [178,812] 499 (74.7) 499 [499] 178,313 (35.1) 178,313 [178,313]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Recovered with sequelae 9,451 (1.9) 9,451 [9,451] 39 (5.8) 39 [39] 9,412 (1.9) 9,412 [9,412]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Unknown 155,903 (30.7) 155,903 [155,903] 2 (0.3) 2 [2] 155,901 (30.7) 155,901 [155,901]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Presence of comorbidities b</td>
<td>Yes 38,787 (7.6) 38,787 [38,787] 259 (38.8) 259 [259] 38,528 (7.6) 38,528 [38,528]</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>No 469,564 (92.4) 469,564 [469,564] 409 (61.2) 409 [409] 469,155 (92.4) 469,155 [469,155]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a. The sum of percentages may not exactly match 100% due to rounding in calculations.
b. Cases retrieved applying the criteria in place to identify the reports involving the special populations of Immunocompromised patients, Patients with autoimmune or inflammatory disorders, Frail patients with comorbidities (e.g. COPD, diabetes, chronic neurological disease, cardiovascular disorders, active tuberculosis) described as special populations in Section 16.3.5.5, Section 16.3.5.6, and Section 16.3.5.7, respectively.
c. It includes 2 pregnant female subjects aged 16 and 17 years.
d. Foetus cases-Age range only applies to post-birth subjects.
e. Numbers of squared brackets include number of participants/subjects who received BNT162b2.
Numbers of squared brackets do not include non-participants (foetus/neonates) exposed before or during the mother’s pregnancy or babies exposed through breastfeeding. Number of cases reported in this table may not match with number of cases evaluated in individual Sections due to case by case review that is not possible to implement in the overall dataset.
f. It includes 284 cases where the age is reported as “Adolescent”.
g. There were 4 cases involving pregnant subjects, whose ages were erroneously reported under the elderly age group. These 4 cases are excluded in the Section 16.3.5.1 Use in Elderly Patients.
MC = Medically confirmed; N: Number of cases; N/A=Not applicable; CT=Clinical trial; PM=Post-marketing

6.3.1.1.1. General Overview of the Safety Database - Clinical Trials Data

During the reporting period, in the CT dataset, the number of male participants was slightly higher than female (53.9% vs 45.5%); the number of SAEs experienced by male participants...
is slightly higher than female (482 vs 391); in the 18 - 30 years and the 31 - 50 years age groups, the number of SAEs reported in females was higher than in males, while in the paediatric population, in 51-64 years and in the elderly (≥ 65 years) age groups, the SAEs reported in male participants was higher than in females (Figure 1).

**Figure 1. Clinical Trial Data: Number of SAEs by Age Group and Gender**

![Bar chart showing the number of SAEs by age group and gender.]

Case outcomes by presence/absence of comorbidities, by gender and age group in clinical trial cases are presented in Figure 2 through Figure 5. Overall, the proportion of cases with comorbidities is lower than cases without comorbidities and this is reflected also among the cases with a fatal outcome (Figure 2). A slightly higher number of male participants (7 cases, 1.0% of the total CT dataset) than female participants (6 cases, 0.9% of the total CT dataset) experienced a fatal outcome in the presence of comorbidities (Figure 3), and more male (18 cases, 2.7% of the total CT dataset) than female participants (4 cases, 0.6% of the total CT dataset) experienced a fatal outcome in the absence of comorbidities (Figure 3). When comorbidities are reported, the age group 51-64 years is the most represented across the case outcomes of recovered/recovering (55), recovered with sequelae (5), and not recovered (16). Among the cases with a fatal outcome, most cases were presented in the age group ≥ 75 years (5); furthermore, the same number of occurrences was reported in the age groups 31-50 years and 51-64 years, as shown in Figure 4. In cases without comorbidities (Figure 5), most of the

---

35 Cases retrieved applying the criteria in place to identify the reports involving the special populations of Immunocompromised patients, Patients with autoimmune or inflammatory disorders, Frail patients with comorbidities (e.g. COPD, diabetes, chronic neurological disease, cardiovascular disorders, active tuberculosis) described as special populations in Section 16.3.5.5, Section 16.3.5.6 and Section 16.3.5.7, respectively.
cases had a favourable outcome across all age groups at the time of reporting with the paediatric participants as more represented group, whose recovered case outcome involved 68 cases; the highest number of fatal outcomes occurred in the 31-50 years and 65-74 years followed by the 51-64 years.
Figure 2. Clinical Trial Data: Case Outcome by Presence/Absence of Comorbidities

- **FATAL**
  - With Comorbidities: 13
  - Without Comorbidities: 22

- **NOT RECOVERED/NOT RESOLVED**
  - With Comorbidities: 42
  - Without Comorbidities: 51

- **RECOVERED/RESOLVED**
  - With Comorbidities: 146
  - Without Comorbidities: 234

- **RECOVERED/RESOLVED WITH SEQUELAE**
  - With Comorbidities: 14
  - Without Comorbidities: 25

- **RECOVERING/RESOLVING**
  - With Comorbidities: 44
  - Without Comorbidities: 75

- **UNKNOWN**
  - With Comorbidities: 2
  - Without Comorbidities: 0
Figure 3. Clinical Trial Data: Case Outcome by Presence/Absence of Comorbidities and Gender

<table>
<thead>
<tr>
<th>Case Outcome</th>
<th>With Comorbidities</th>
<th>Without Comorbidities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>Female</td>
<td>14</td>
<td>20</td>
</tr>
<tr>
<td>Male</td>
<td>18</td>
<td>7</td>
</tr>
<tr>
<td>Not Recovered/Not Resolved</td>
<td>28</td>
<td>30</td>
</tr>
<tr>
<td>Female</td>
<td>64</td>
<td>107</td>
</tr>
<tr>
<td>Male</td>
<td>124</td>
<td>82</td>
</tr>
<tr>
<td>Recovered/Resolved</td>
<td>3</td>
<td>10</td>
</tr>
<tr>
<td>Female</td>
<td>3</td>
<td>11</td>
</tr>
<tr>
<td>Male</td>
<td>25</td>
<td>19</td>
</tr>
<tr>
<td>Recovering/Resolving</td>
<td>11</td>
<td>15</td>
</tr>
<tr>
<td>Female</td>
<td>25</td>
<td>43</td>
</tr>
<tr>
<td>Male</td>
<td>32</td>
<td>19</td>
</tr>
<tr>
<td>Unknown</td>
<td>3</td>
<td>2</td>
</tr>
</tbody>
</table>

CONFIDENTIAL
Page 50
Figure 4. Clinical Trial Data: Case Outcome by Age Group in Presence of Comorbidities

- 18 - 30 years
- 31 - 50 years
- 51 - 64 years
- 65 - 74 years
- Greater than or equal to 75 years
- Less than or equal to 17 years

<table>
<thead>
<tr>
<th>Case Outcome</th>
<th>18 - 30 years</th>
<th>31 - 50 years</th>
<th>51 - 64 years</th>
<th>65 - 74 years</th>
<th>Greater than or equal to 75 years</th>
<th>Less than or equal to 17 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>2</td>
<td>1</td>
<td>4</td>
<td>5</td>
<td>9</td>
<td>10</td>
</tr>
<tr>
<td>Not Recovered/Not Resolved</td>
<td>16</td>
<td>15</td>
<td>9</td>
<td>4</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Recovered/Resolved</td>
<td>18</td>
<td>21</td>
<td>30</td>
<td>28</td>
<td>45</td>
<td>21</td>
</tr>
<tr>
<td>Recovered/Resolved with Sequelae</td>
<td>1</td>
<td>5</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Recovering/Resolving</td>
<td>9</td>
<td>10</td>
<td>14</td>
<td>6</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

No. of Cases
Figure 5. Clinical Trial Data: Case Outcome by Age Group in Absence of Comorbidities
The summary of medical history and co-suspects reported in the CT cases is provided in Table 15.

Table 15. Clinical Trial Data: Medical History and Co-Suspects

<table>
<thead>
<tr>
<th>Medical History and Co-Suspects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most frequently reported (≥2% medical history (HLGT): Vascular hypertensive disorders (192), Lipid metabolism disorders (131), Appetite and general nutritional disorders, Glucose metabolism disorders (incl diabetes mellitus) (90 each), Allergic conditions (87), Joint disorders (86), Lifestyle issues (83), Gastrointestinal motility and defaecation conditions (76), Depressed mood disorders and disturbances (70), Anxiety disorders and symptoms (68), Bronchial disorders (excl neoplasms) (62), Gastrointestinal therapeutic procedures (57), Sleep disorders and disturbances (48), Thyroid gland disorders (46), Infections - pathogen unspecified (44), Cardiac arrhythmias, Coronary artery disorders (41 each), Obstetric and gynaecological therapeutic procedures (39), Bone and joint therapeutic procedures (35), Respiratory disorders NEC (34), Viral infectious disorders, Therapeutic procedures and supportive care NEC (31 each), Musculoskeletal and connective tissue disorders NEC, Headaches, Peripheral neuropathies, Vascular therapeutic procedures (27 each), Pregnancy, labour, delivery and postpartum conditions, Musculoskeletal and connective tissue deformities (incl intervertebral disc disorders) (25 each), Epidermal and dermal conditions, Male genital tract therapeutic procedures (24 each), Age related factors (23), Vision disorders, Injuries NEC, Bone disorders (excl congenital and fractures), Prostatic disorders (excl infections and inflammations) (22 each), Upper respiratory tract disorders (excl infections), Bone and joint injuries (21 each), General system disorders NEC, Hepatobiliary therapeutic procedures (20 each), Gallbladder disorders, Renal and urinary tract therapeutic procedures (19 each), Abdominal hernias and other abdominal wall conditions, Muscle disorders (18 each), Lipid analyses, Central nervous system vascular disorders, Nervous system, skull and spine therapeutic procedures (16 each), Anterior eye structural change, deposit and degeneration, Renal disorders (excl nephropathies) (15 each), Anaemias nonhaemolytic and normoerythrocytic, Pyridoxine and pyridoxal phosphate metabolism disorders, Urinary tract signs and symptoms, and Cardiac therapeutic procedures (14 each), Breast therapeutic procedures, and Gastrointestinal signs and symptoms (13 each).</td>
</tr>
</tbody>
</table>

The medical conditions (PT's History) reported in more than 2% of the cases included Hypertension (186), Obesity (77), Type 2 diabetes mellitus (71), Depression (63), Osteoarthritis, Seasonal allergy (62 each), Gastroesophageal reflux disease (60), Anxiety (56), Hypercholesterolaemia (51), Hyperlipidaemia (48), Insomnia (45), Hypothyroidism (43), Asthma (34), Non-tobacco user, Dyslipidaemia (29 each), Coronary artery disease (28), Pregnancy, Atrial fibrillation (25 each), Ex-smoker user (24), Chronic obstructive pulmonary disease (23), Back pain (22), Benign prostatic hyperplasia (20), Cholecystectomy (19), Appendicectomy, Migraine, Postmenopause (18 each), Tobacco user (16), Rhinitis allergic, Choledolithiasis, Neuropathy peripheral, Osteoporosis (15 each), Blood cholesterol increased, Cataract, Diabetes mellitus, Drug hypersensitivity, Hysterectomy, and Vasectomy (14 each), Sleep apnoea syndrome, Ex-alcohol user, Live birth, and Caesarean section (13 each). |


Most frequently reported (≥ 2 cases) co-suspect medications: amlodipine besilate and metformin (2 each).
Adverse Event Data

A total of 879 SAEs were reported in 668 cases.

The MedDRA SOCs containing the greatest number of reported events (≥2%) from clinical trial data were Infections and infestations (158), Injury, poisoning and procedural complications (100), Neoplasms benign, malignant and unspecified (incl cysts and polyps) (99), Cardiac disorders (66), Nervous system disorders (64), Gastrointestinal disorders (61), General disorders and administration site conditions (54), Respiratory, thoracic and mediastinal disorders (42), Musculoskeletal and connective tissue disorders (40), Hepatobiliary disorders (30), Psychiatric disorders (28), Vascular disorders (25), Pregnancy, puerperium and perinatal conditions (24), Metabolism and nutrition disorders, Renal and urinary disorders (22 each).

The overall safety evaluation includes a review of the most frequently reported serious events by SOC and PT for events reported in ≥2% of all clinical trial cases during the reporting interval as compared to the cumulative period through 18 June 2022, as summarised in Table 16.

---

36 Of note, multiple adverse events may be reported in a single case.
Table 16. Clinical Trial Data: Serious Events Reported in ≥2% Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>Reporting Period</th>
<th>Cumulatively through</th>
<th>MedDRA PT</th>
<th>All Cases&lt;sup&gt;a&lt;/sup&gt;</th>
<th>BNT162b2 / b2b01 / BT Cases</th>
<th>All Cases&lt;sup&gt;b&lt;/sup&gt;</th>
<th>BNT162b1 / b2 / b2b01 / b3 / c2&lt;sup&gt;c&lt;/sup&gt;</th>
<th>BT Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maternal exposure during pregnancy&lt;sup&gt;d&lt;/sup&gt;</td>
<td>25 (3.7)</td>
<td>25 (3.8)</td>
<td>121 (5.0)</td>
<td>113 (5.0)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Condition aggravated</td>
<td>24 (3.6)</td>
<td>23 (3.5)</td>
<td>79 (3.3)</td>
<td>72 (3.2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infections and infestations</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pneumonia</td>
<td>17 (2.5)</td>
<td>17 (2.6)</td>
<td>56 (2.3)</td>
<td>54 (2.4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gastroenteritis</td>
<td>15 (2.3)</td>
<td>15 (2.3)</td>
<td>22 (0.9)</td>
<td>21 (0.9)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Appendicitis</td>
<td>14 (2.1)</td>
<td>13 (2.0)</td>
<td>58 (2.4)</td>
<td>53 (2.3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>16 (2.4)</td>
<td>16 (2.4)</td>
<td>47 (1.9)</td>
<td>46 (2.0)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cerebrovascular accident</td>
<td>13 (2.0)</td>
<td>13 (2.0)</td>
<td>40 (1.6)</td>
<td>39 (1.7)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neoplasms benign, malignant and unspecified (incl cysts and polyps)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prostate cancer</td>
<td>13 (2.0)</td>
<td>13 (2.0)</td>
<td>32 (1.3)</td>
<td>32 (1.4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Includes BNT162b2 (b2), BNT162b2b201 (b2b01), BT, and Placebo.
<sup>b</sup> Includes BNT162b1, b2, b2b01, b3, BNT162c2 (c2), BT and Placebo.
<sup>c</sup> The variant vaccines b1 and c2 are study drugs in study BNT162-01, b2b01 in Study BNT162-14 and b3 in Study BNT162-04, respectively. Please refer to Section 7 for details on these studies.
<sup>d</sup> Reporting proportion calculated as n/N (% of cases) in the current reporting period or cumulatively.
<sup>e</sup> Reported as serious occurrence associated to SAEs. This PT is coded in maternal cases, and in foetal cases when a foetal AE is reported. For associated SAEs, refer to Section 16.3.5.3, Use in Pregnant/Lactating Women.

AE = Adverse Event; AERP = Adverse Event Reporting Proportion; BT = Blinded Therapy; MedDRA = Medical Dictionary for Regulatory Activities; N = Number of cases; n = Number of events; PT = Preferred Term; SOC = Summary Organ Class

During the reporting period, the most frequently reported serious adverse events in the clinical trials are not expected or consistent with expected events as per the current Investigator's Brochure except Appendicitis. Among these most frequently reported serious adverse events, the reporting proportion of the PT Gastroenteritis during the reporting interval was higher (2.3%) compared to its proportion in the cumulative dataset (0.9%). Upon review, all SAEs of Gastroenteritis during the reporting interval are assessed as unrelated by the investigator and the Sponsor. Event outcomes were resolved (14) and resolved with sequelae (1).
There were 2 SAEs assessed as related to BNT162b2 during the reporting interval:

- Dehydration was assessed as related by both the Investigator and the Sponsor.
- Abortion spontaneous was assessed as related by the Investigator and unrelated by the Sponsor.

**Conclusion**

Based on the review of the CT cases, no new safety issues were identified.

**6.3.1.1.2. General Overview of the Safety Database - Post-Authorisation Data**

During the reporting period, in the PM dataset the number of female subjects was 2.2 times the number of male subjects (63.8% vs 29.4%); across the different age groups the ratio of female/male cases ranged between 1.1 in the less than or equal to 17 years to 2.7 in the 31-50 years group (Figure 6).

**Figure 6. Post-Authorisation Data: Number of Cases by Age Group and Gender**

![Bar Chart](image-url)
Case outcomes by presence/absence of comorbidities\textsuperscript{35}, by gender and age group in PM cases are presented in Figure 7 through Figure 10. Overall, the proportion of cases with comorbidities was 7.6% of the total PM dataset; Figure 7 shows the case outcome by presence/absence of comorbidities.
Figure 7. Post-Authorisation Data: Case Outcome by Presence/Absence of Comorbidities

- With Comorbidities
- Without Comorbidities

FATAL

NOT RECOVERED/NOT RESOLVED

RECOVERED/RESOLVED

RECOVERED/RESOLVED WITH SEQUELAE

RECOVERING/RESOLVING

UNKNOWN

A slightly higher number of male subjects experienced a fatal outcome independently from the presence of comorbidities (Figure 8).
The age group 31-50 years represents 36.1% of the PM cases; this age group is the most represented one across all but fatal case outcomes, both in presence and in absence of comorbidities (Figure 9 and Figure 10). Among the cases with a fatal outcome, the age group more represented is the one including elderly aged at least 75 years, both in presence or in absence of comorbidities, as shown in Figure 9 and in Figure 10.
Figure 9. Post-Authorisation Data: Case Outcome by Age Group in Presence of Comorbidities
Figure 10. Post-Authorisation Data: Case Outcome by Age Group in Absence of Comorbidities

The summary of medical history and co-suspects reported in the PM cases is provided in Table 17
Table 17. Post-Authorisation Data: Medical History and Co-Suspects

Most frequently reported (≥2%) medical history (HLGT): Viral infectious disorders (22,612), Allergic conditions (19,947), Vascular hypertensive disorders (13,073), and Lifestyle issues (11,228).

The medical conditions (PTs History) reported in more than 2% of the cases included PT Hypertension (12,888).

COVID-19 medical history: COVID-19 (14,526), Suspected COVID-19 (5971), Post-acute COVID-19 syndrome (254), COVID-19 pneumonia (111), SARS-CoV-2 test positive (90), Coronavirus infection (77), Exposure to SARS-CoV-2 (71), Asymptomatic COVID-19 (52), SARS-CoV-2 antibody test positive (6), Coronavirus test positive, Occupational exposure to SARS-CoV-2 (3 each), Breakthrough COVID-19 (2), and COVID-19 treatment (1).

Most frequently reported (≥40) co-suspect vaccines/medications (other than COVID-19 vaccines): adalimumab (626), influenza vaccine (465), influenza vaccine inact SAG 4V (149), influenza vaccine inact SPLIT 4V (143), ocrelizumab (92), upadacitinib (80), influenza vaccine inact SPLIT 3V (44), Risankizumab (42), pneumococcal vaccine polysacch 23V (41), apixaban, and ethinylestradiol/levonorgestrel (40 each).

Most frequently reported (≥42) co-suspect COVID-19 vaccines: COVID-19 vaccine (1929), COVID-19 vaccine mRNA (mRNA 1273) (1608), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (1040), JNJ 78436735 (144), and COVID-19 vaccine inact (vero) CZ02 (42).

Adverse Event Data

A total of 1,596,793 AEs (of which 439,443 were serious and 1,158,240 non-serious\textsuperscript{37}) were reported in 507,683 PM cases.

The MedDRA SOCs containing the greatest number of events (≥2%) were General disorders and administration site conditions (459,731), Nervous system disorders (204,185), Musculoskeletal and connective tissue disorders (148,849), Injury, poisoning and procedural complications (130,333), Infections and infestations (82,131), Gastrointestinal disorders (81,816), Reproductive system and breast disorders (77,917), Skin and subcutaneous tissue disorders (62,405), Respiratory, thoracic and mediastinal disorders (56,663), Cardiac disorders (54,208), Surgical and medical procedures (52,531), and Blood and lymphatic system disorders (38,366).

The overall safety evaluation includes a review of the most frequently reported events by SOC and by PT for events reported in ≥2% of all post-marketing cases during the interval period as compared to the cumulative period through 18 June 2022.

\textsuperscript{37} Multiple episodes of the same event were reported with a different seriousness in some cases hence the sum of the events seriousness exceeds the total number of events.
## Table 18. Post-Authorisation Data: Events Reported in ≥2% Cases

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>All Cases</strong></td>
<td><strong>Serious Cases</strong></td>
<td><strong>All Cases</strong></td>
</tr>
<tr>
<td>(N=507,683)</td>
<td>(N=151,420)</td>
<td>(N=1,484,945)</td>
</tr>
<tr>
<td>AEs</td>
<td>Serious AEs</td>
<td>AEs</td>
</tr>
<tr>
<td>(n=1,596,793)</td>
<td>(n=439,443)</td>
<td>(n=4,974,391)</td>
</tr>
<tr>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
</tr>
<tr>
<td><strong>Nervous system disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache b</td>
<td>77,974 (15.4)</td>
<td>297,293 (20.0)</td>
</tr>
<tr>
<td></td>
<td>9451 (6.2)</td>
<td>41,338 (9.7)</td>
</tr>
<tr>
<td>Dizziness b</td>
<td>30,880 (6.1)</td>
<td>93,304 (6.3)</td>
</tr>
<tr>
<td></td>
<td>5418 (3.6)</td>
<td>20,903 (4.9)</td>
</tr>
<tr>
<td>Paraesthesia a</td>
<td>14,993 (3.0)</td>
<td>44,666 (3.0)</td>
</tr>
<tr>
<td></td>
<td>3018 (2.0)</td>
<td>10,640 (2.5)</td>
</tr>
<tr>
<td><strong>General disorders and administration site conditions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatigue b</td>
<td>67,879 (13.4)</td>
<td>235,562 (15.9)</td>
</tr>
<tr>
<td></td>
<td>8675 (5.7)</td>
<td>34,742 (8.2)</td>
</tr>
<tr>
<td>Pyrexia b</td>
<td>57,746 (11.4)</td>
<td>228,574 (15.4)</td>
</tr>
<tr>
<td></td>
<td>6642 (4.4)</td>
<td>29,973 (7.0)</td>
</tr>
<tr>
<td>Vaccination site pain b</td>
<td>49,263 (9.7)</td>
<td>190,875 (12.9)</td>
</tr>
<tr>
<td></td>
<td>2199 (1.5)</td>
<td>9,703 (2.3)</td>
</tr>
<tr>
<td>Chills b</td>
<td>33,542 (6.6)</td>
<td>128,602 (8.7)</td>
</tr>
<tr>
<td></td>
<td>2895 (1.9)</td>
<td>14,687 (3.5)</td>
</tr>
<tr>
<td>Malaise b</td>
<td>32,701 (6.4)</td>
<td>142,545 (9.6)</td>
</tr>
<tr>
<td></td>
<td>3327 (2.2)</td>
<td>15,085 (3.5)</td>
</tr>
<tr>
<td>Drug ineffective b</td>
<td>26,688 (5.3)</td>
<td>41,566 (2.8)</td>
</tr>
<tr>
<td></td>
<td>26,664 (17.6)</td>
<td>41,515 (9.8)</td>
</tr>
<tr>
<td>Vaccination failure d</td>
<td>24,419 (4.8)</td>
<td>37,933 (2.6)</td>
</tr>
<tr>
<td></td>
<td>24,415 (16.1)</td>
<td>37,926 (8.9)</td>
</tr>
<tr>
<td>Chest pain i</td>
<td>17,945 (3.5)</td>
<td>40,839 (2.8)</td>
</tr>
<tr>
<td></td>
<td>5694 (3.8)</td>
<td>15,623 (3.7)</td>
</tr>
<tr>
<td>Pain b</td>
<td>16,529 (3.3)</td>
<td>80,302 (5.4)</td>
</tr>
<tr>
<td></td>
<td>3618 (2.4)</td>
<td>14,660 (3.4)</td>
</tr>
<tr>
<td>Anemia b</td>
<td>13,703 (2.7)</td>
<td>59,692 (4.0)</td>
</tr>
<tr>
<td></td>
<td>2793 (1.8)</td>
<td>11,424 (2.7)</td>
</tr>
<tr>
<td>Vaccination site swelling b</td>
<td>10,670 (2.1)</td>
<td>40,218 (2.7)</td>
</tr>
<tr>
<td></td>
<td>446 (0.3)</td>
<td>1,954 (0.5)</td>
</tr>
<tr>
<td><strong>Infections and infestations</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>COVID-19 e</td>
<td>47,988 (9.5)</td>
<td>76,044 (5.1)</td>
</tr>
<tr>
<td></td>
<td>47,449 (31.3)</td>
<td>72,718 (17.1)</td>
</tr>
<tr>
<td><strong>Musculoskeletal and connective tissue disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myalgia b</td>
<td>43,916 (8.7)</td>
<td>178,198 (12.0)</td>
</tr>
<tr>
<td></td>
<td>4451 (2.9)</td>
<td>18,937 (4.5)</td>
</tr>
<tr>
<td>Arthralgia b</td>
<td>29,430 (5.8)</td>
<td>121,898 (8.2)</td>
</tr>
<tr>
<td></td>
<td>4702 (3.1)</td>
<td>18,152 (4.3)</td>
</tr>
<tr>
<td>Pain in extremity b</td>
<td>25,090 (4.9)</td>
<td>93,467 (6.3)</td>
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<tr>
<td></td>
<td>4584 (3.0)</td>
<td>18,828 (4.4)</td>
</tr>
<tr>
<td>Limb discomfort b</td>
<td>11,578 (2.3)</td>
<td>23,939 (1.6)</td>
</tr>
<tr>
<td></td>
<td>670 (0.4)</td>
<td>2,558 (0.6)</td>
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<tr>
<td><strong>Injury, poisoning and procedural complications</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inappropriate schedule of product administration d</td>
<td>35,318 (7.0)</td>
<td>57,719 (3.9)</td>
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<tr>
<td></td>
<td>466 (0.3)</td>
<td>1,020 (0.2)</td>
</tr>
<tr>
<td>Off label use d</td>
<td>29,927 (5.9)</td>
<td>54,754 (3.7)</td>
</tr>
<tr>
<td></td>
<td>10,293 (6.8)</td>
<td>16,400 (3.9)</td>
</tr>
<tr>
<td>Poor quality product administered j</td>
<td>17,859 (3.5)</td>
<td>30,830 (2.1)</td>
</tr>
<tr>
<td></td>
<td>4 (0.003)</td>
<td>14 (0.004)</td>
</tr>
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</table>
Table 18. Post-Authorisation Data: Events Reported in ≥2% Cases

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All Cases (N=507,683) AEs (n=1,596,793)</td>
<td>Serious Cases (N=151,420) Serious AEs (n=439,443)</td>
</tr>
<tr>
<td></td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
</tr>
<tr>
<td>Blood and lymphatic system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lymphadenopathy b</td>
<td>31,132 (6.1)</td>
<td>2794 (1.9)</td>
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<tr>
<td>Gastrointestinal disorders</td>
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<td></td>
</tr>
<tr>
<td>Nausea b</td>
<td>30,670 (6.0)</td>
<td>4338 (2.9)</td>
</tr>
<tr>
<td>Vomiting b</td>
<td>11,424 (2.3)</td>
<td>2454 (1.6)</td>
</tr>
<tr>
<td>Diarrhoea b</td>
<td>10,211 (2.0)</td>
<td>1644 (1.1)</td>
</tr>
<tr>
<td>Surgical and medical procedures</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immunisation c</td>
<td>25776 (5.1)</td>
<td>11,063 (7.3)</td>
</tr>
<tr>
<td>Interchange of vaccine products c</td>
<td>25,233 (5.0)</td>
<td>9397 (6.2)</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dyspnoea b</td>
<td>21,736 (4.3)</td>
<td>6947 (4.6)</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash b</td>
<td>13,640 (2.7)</td>
<td>1802 (1.2)</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palpitations b</td>
<td>13,071 (2.6)</td>
<td>4231 (2.8)</td>
</tr>
<tr>
<td>Tachycardia d</td>
<td>10,914 (2.2)</td>
<td>3028 (2.0)</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heavy menstrual bleeding i</td>
<td>12,905 (2.5)</td>
<td>1711 (1.1)</td>
</tr>
<tr>
<td>Menstrual disorder i</td>
<td>12,579 (2.5)</td>
<td>871 (0.6)</td>
</tr>
</tbody>
</table>
Table 18. Post-Authorisation Data: Events Reported in ≥2% Cases

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All Cases (N=507,683)</td>
<td>All Cases (N=1,484,945)</td>
</tr>
<tr>
<td></td>
<td>Serious Cases (N=151,420)</td>
<td>Serious Cases (N=425,314)</td>
</tr>
<tr>
<td></td>
<td>Serious AEs *(n=1,596,793)</td>
<td>Serious AEs *(n=4,974,391)</td>
</tr>
<tr>
<td></td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
</tr>
<tr>
<td></td>
<td>Serious AEs *(n=439,443)</td>
<td>Serious AEs *(n=1,326,116)</td>
</tr>
<tr>
<td></td>
<td>n (AERP, %)</td>
<td>n (AERP, %)</td>
</tr>
</tbody>
</table>

a. Non-serious events are not included.
b. Listed or consistent with listed AEs in current RSI.
c. Listed per case processing conventions, except for fatal cases.
d. Listed per case processing conventions.
e. PTs selected per case processing conventions to indicate cases reporting third/booster doses.
f. Reporting proportion calculated as n/N (% of all incremental cases, incremental serious cases and all cumulative cases).
g. Paresthesia / Hypoesthesia were included as ADRs in the EU-SmPC Section 4.8 as per PRAC recommendation (Procedure number EMEA/H/C/005735/II/0080).
h. Drug ineffective represents efficacy-related conditions.
i. Unlisted in the current RSI.
j. Follow the listedness of the associated AE.

N=Number of cases; n=Number of events; MedDRA=Medical Dictionary for Regulatory Activities; SOC=System Organ Class; PT=Preferred Term; AE=Adverse Event; AERP=Adverse Event Reporting Proportion; RSI=Reference Safety Information.
Most of the frequently reported events are listed or consistent with listed events as per the current RSI.

Out of the 1,596,793 AEs in the PM dataset, 72.5% of them were non-serious. Figure 11 shows the seriousness of the PTs reported in more than 2% of the cases where most of the occurrences were non-serious with the exception of COVID-19, Drug ineffective, and Vaccination failure.

Figure 12 provides information about the age breakdown in the clinical AEs reported in more than 2% of the cases by SOC; the age group 31-50 years is the one reporting higher proportion of events than other age groups and this is consistent being the largest group in terms of number of cases.

**Figure 11. Post-Authorisation Data: Event Seriousness of the PTs ≥2% of Cases**
Figure 12. Post-Authorisation Data: Clinical AEs reported in ≥2% of Cases by Age Group

Nervous System Disorders
- Less than or equal to 17 years
- 18 - 39 years
- 40 - 59 years
- 60 - 64 years
- 65 - 74 years
- Greater than or equal to 75 years
- Unknown

General disorders and administration site conditions
- Less than or equal to 17 years
- 18 - 39 years
- 40 - 59 years
- 60 - 64 years
- 65 - 74 years
- Greater than or equal to 75 years
- Unknown
Conclusion

Overall, during the reporting period, the serious cases represented 29.8% of the total PM; fatal outcomes occurred in less than 1% of the cases. About two-thirds of the cases occurred in female subjects and the age group 31-50 years was the group most frequently reporting AEs. The most frequently reported (≥2%) AEs (listed in the current RSI) are in majority non-serious.

Based on the review of the PM cases, no new safety issues were identified.

6.3.1.1.2.1. Analysis by Dose

Potential for local and systemic adverse reactions are analysed by dose of the vaccine in Section 16.3.3.3 Local Adverse Reactions and Section 16.3.3.4 Systemic Adverse Reactions.

6.3.1.1.2.2. Tris/Sucrose Presentation

The currently authorised presentations of BNT162b2 that use tromethamine (Tris) buffer are the following:

- Grey cap: multidose vial, formulated to provide, without need for dilution, 6 doses (each 0.3 mL dose containing 30 μg modRNA) for individuals 12 years of age and older. This presentation was approved first in the US on 29 October 2021 and then in other countries worldwide.

- Orange cap: multidose vial, formulated to provide, after dilution, 10 doses (each 0.2 mL dose containing 10 μg modRNA) for individuals 5 through 11 years of age. This paediatric presentation was approved first in the US on 29 October 2021 and then in other countries worldwide.

- Maroon cap: multidose vial, formulated to provide, after dilution, 10 doses (each 0.2 mL dose containing 3 μg modRNA) for individuals 6 months through 4 years of age. This paediatric presentation was approved first in the US on 17 June 2022.

A total of 19,789 case reports with Tris/Sucrose formulation\(^{38}\) containing 38,950 events (3.9% of the total PM dataset) fulfilled criteria for inclusion in this PSUR reporting period. Data presented in Table 19 through Table 23 refer to the paediatric 5-11 years old orange cap and ≥ 12 years grey cap presentations.\(^{39}\) Demographic information of Tris/Sucrose cases received during the reporting interval are shown in Table 19. Most cases (9055 cases, 45.8%) were reported in paediatric subjects (aged ≤ 17 years); a demographic and the comparison with AEs reporting rate between Tris/Sucrose cases and paediatric PBS/Sucrose cases is provided in Table 22 and Table 23. There were no significant differences in the demographic data between paediatric subjects receiving Tris/Sucrose formulation and those receiving PBS.

\(^{38}\) Search criteria: “EUA Tris” and “BLA Tris” in the concentration field.

\(^{39}\) No data on the paediatric 6 months through 4 years presentation are available since it was first approved on 17 June 2022.
A higher percentage of medication error cases was reported in the Tris/Sucrose paediatric group and this may reflect initial difficulties in managing the new formulation (cross-referenced with Section 9.2, Medication Errors for errors related to Tris/Sucrose formulation). Routine pharmacovigilance activities to mitigate these medication errors, including label information (vial differentiation, instructions for reconstitution and administration, vaccination scheme, storage conditions for each formulation and available dosage), educational materials for healthcare providers, medical information call centers and traceability are listed in the approved version 5.0 of the EU-RMP adopted on 10 March 2022. The approved BLA US-PVP version 1.4.1 dated 29 April 2022 includes as routine pharmacovigilance activities label information on vial differentiation.

With regard to the reported medical events, the majority were reported in lower proportion in the Tris/Sucrose group compared to the PBS/Sucrose group although there were 7 events (Vaccination site pain, Vomiting, Abdominal pain, Diarrhoea, Rash, Urticaria, Pruritus) with a higher AERP (11.4%, 7.5%, 3.8%, 2.6%, 5.5%, 2.8%, and 2.6%, respectively) in the Tris/Sucrose paediatric group. On review, few occurrences were serious (as important medical events – 97 for PT Vomiting, 49 for Abdominal pain, 42 for Rash, 41 for Urticaria, 24 for Diarrhoea, 20 for Pruritus, and 15 for Vaccination site pain). The clinical outcome of the serious occurrences was resolved/resolving (188), resolved with sequelae (5), not resolved (39), unknown (51), and fatal (5) at the time of reporting. In the 4 cases recording Abdominal pain, Vomiting (2 each), and Diarrhoea (1) as the fatal events, limited information was provided in 4 paediatric subjects. In these 4 cases, it is not clear whether the subjects had any underlying diseases or conditions, and date of death was unknown. In the paediatric PBS/Sucrose cases, these events were assessed as serious as follows: PTs Vomiting (325), Abdominal pain (103), Rash (133), Urticaria (76), Diarrhoea (93), PT Pruritus (76), and PT Vaccination site pain (101). These serious events had the report proportion ≤ 0.5% of the total number of events among all paediatric PBS/Sucrose cases.

Table 19. Demographic Information – Tris/Sucrose Cases (Orange and Grey Cap) Received during the Reporting Interval

<table>
<thead>
<tr>
<th>Country of occurrence</th>
<th>Tris/Sucrose No. of Cases (%)* [Direct exposure]°</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Orange Cap N=17,450 Grey Cap N=2,340</td>
</tr>
<tr>
<td><strong>MC cases</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Yes</strong></td>
<td>7902 (45.3)</td>
</tr>
<tr>
<td></td>
<td>1597 (68.2)</td>
</tr>
<tr>
<td><strong>No</strong></td>
<td>9548 (54.7)</td>
</tr>
<tr>
<td></td>
<td>743 (31.8)</td>
</tr>
<tr>
<td><strong>Country of occurrence</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Philippines</strong></td>
<td>7220 (41.4)</td>
</tr>
<tr>
<td></td>
<td>1 (0.04)</td>
</tr>
<tr>
<td><strong>US</strong></td>
<td>4463 (25.6)</td>
</tr>
<tr>
<td></td>
<td>2183 (93.3)</td>
</tr>
<tr>
<td><strong>Australia</strong></td>
<td>1522 (8.7)</td>
</tr>
<tr>
<td></td>
<td>-</td>
</tr>
<tr>
<td><strong>Germany</strong></td>
<td>917 (5.3)</td>
</tr>
<tr>
<td></td>
<td>50 (2.1)</td>
</tr>
<tr>
<td><strong>Japan</strong></td>
<td>896 (5.1)</td>
</tr>
<tr>
<td></td>
<td>1 (0.04)</td>
</tr>
<tr>
<td><strong>Italy</strong></td>
<td>424 (2.4)</td>
</tr>
<tr>
<td></td>
<td>45 (1.9)</td>
</tr>
<tr>
<td><strong>Spain</strong></td>
<td>378 (2.2)</td>
</tr>
<tr>
<td></td>
<td>3 (0.1)</td>
</tr>
<tr>
<td><strong>Canada</strong></td>
<td>171 (1.0)</td>
</tr>
<tr>
<td></td>
<td>2 (0.09)</td>
</tr>
<tr>
<td><strong>Denmark</strong></td>
<td>147 (0.8)</td>
</tr>
<tr>
<td></td>
<td>-</td>
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<tr>
<td><strong>France</strong></td>
<td>141 (0.8)</td>
</tr>
<tr>
<td></td>
<td>34 (1.5)</td>
</tr>
<tr>
<td><strong>Total Others</strong></td>
<td>1171 (6.7)</td>
</tr>
<tr>
<td></td>
<td>21 (0.9)</td>
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# Table 19. Demographic Information – Tris/Sucrose Cases (Orange and Grey Cap) Received during the Reporting Interval

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<tr>
<th></th>
<th>Tris/Sucrose</th>
<th></th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>No. of Cases* (%)†</td>
<td>Direct exposure‡</td>
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</tr>
<tr>
<td></td>
<td>Orange Cap</td>
<td>Grey Cap</td>
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</tr>
<tr>
<td>Gender</td>
<td></td>
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<tr>
<td>Female</td>
<td>6550 (37.5)</td>
<td>361 (15.4)</td>
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<tr>
<td>Male</td>
<td>6007 (34.4)</td>
<td>240 (10.3)</td>
<td></td>
</tr>
<tr>
<td>Unknown/No Data</td>
<td>4893 (28.0)</td>
<td>1739 (74.3)</td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>9472</td>
<td>567</td>
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</tr>
<tr>
<td>Min-Max</td>
<td>0.06-100</td>
<td>0.5-99</td>
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<tr>
<td>Mean / Median</td>
<td>15.5/9</td>
<td>45.5/44</td>
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</tr>
<tr>
<td>Age Range</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 17 years</td>
<td>9028 (51.7) [9015]</td>
<td>42 (1.8) [40]</td>
<td></td>
</tr>
<tr>
<td>0 to 27 days</td>
<td>2 (0.01) [0]</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>28 days to 23 months</td>
<td>20 (0.1) [11]</td>
<td>2 (0.09) [0]</td>
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<tr>
<td>2-11 years</td>
<td>8380 (48.0) [8378]</td>
<td>12 (0.5) [12]</td>
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<tr>
<td>12-17 years</td>
<td>626 (3.6) [626]</td>
<td>28 (1.2) [28]</td>
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<tr>
<td>18-30 years</td>
<td>123 (0.7)</td>
<td>127 (5.4)</td>
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<tr>
<td>31-50 years</td>
<td>239 (1.4)</td>
<td>164 (7.0)</td>
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<tr>
<td>51-64 years</td>
<td>382 (2.2)</td>
<td>118 (5.0)</td>
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</tr>
<tr>
<td>65-74 years</td>
<td>341 (2.0)</td>
<td>67 (2.9)</td>
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</tr>
<tr>
<td>≥ 75 years</td>
<td>209 (1.2)</td>
<td>53 (2.3)</td>
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<tr>
<td>Unknown</td>
<td>7127 (40.8)</td>
<td>1769 (75.6)</td>
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<tr>
<td>Case Seriousness</td>
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<tr>
<td>Serious</td>
<td>2070 (11.9)</td>
<td>80 (3.4)</td>
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<tr>
<td>Non-serious</td>
<td>15,380 (88.1)</td>
<td>2260 (96.6)</td>
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<td>Case Outcome</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>30 (0.2)</td>
<td>3 (0.1)</td>
<td></td>
</tr>
<tr>
<td>Not resolved</td>
<td>1618 (9.3)</td>
<td>101 (4.3)</td>
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</tr>
<tr>
<td>Resolved/Resolving</td>
<td>4511 (25.9)</td>
<td>78 (3.3)</td>
<td></td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>40 (0.2)</td>
<td>5 (0.2)</td>
<td></td>
</tr>
<tr>
<td>Unknown/No data</td>
<td>11,251 (64.5)</td>
<td>2153 (92.0)</td>
<td></td>
</tr>
<tr>
<td>Presence of comorbidities ‡</td>
<td>Yes</td>
<td>617 (3.5)</td>
<td>55 (2.4)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>16,833 (96.5)</td>
<td>2285 (97.6)</td>
</tr>
</tbody>
</table>

a. Includes all subjects to whom BNT162b2 (Tris/Sucrose formulation) was administered.
b. Due to rounding, sum of percentages may not match 100%.
c. Includes only subjects to whom BNT162b2 (Tris/Sucrose formulation) was administered directly; does not include reports of foetus/neonates exposed during the mother's pregnancy or babies exposed through breastfeeding.
d. Cases retrieved applying the criteria in place to identify the reports involving the special populations of Immunocompromised patients, Patients with autoimmune or inflammatory disorders, Frail patients with comorbidities (e.g., COPD, diabetes, chronic neurological disease, cardiovascular disorders, active tuberculosis) described as special populations in Section 16.3.5.5, Section 16.3.5.6, and Section 16.3.5.7, respectively.
e. In 1 case, the subject received orange and grey cap separately in different periods.
N=Number of cases; MC=Minedically confirmed; Min=Minimum; Max=Maximum
Table 20. Demographic Information – Comparison of Paediatric (≤ 17 years) Tris/Sucrose (Grey and Orange Cap) versus Paediatric PBS/Sucrose Cases

<table>
<thead>
<tr>
<th></th>
<th>Tris/Sucrose (Grey and Orange Cap) No. of Cases (%)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>PBS/Sucrose No. of Cases (%)&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N=9055</td>
<td>N=22,772</td>
</tr>
<tr>
<td>MC cases</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>6471 (71.5)</td>
<td>13,659 (60.0)</td>
</tr>
<tr>
<td>No</td>
<td>2584 (28.5)</td>
<td>9113 (40.0)</td>
</tr>
<tr>
<td>Country of occurrence</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Germany</td>
<td>773 (8.5)</td>
<td>3916 (17.2)</td>
</tr>
<tr>
<td>Philippines</td>
<td>1229 (13.6)</td>
<td>3064 (13.5)</td>
</tr>
<tr>
<td>Australia</td>
<td>1508 (16.7)</td>
<td>2155 (9.5)</td>
</tr>
<tr>
<td>Malaysia</td>
<td>7 (0.08)</td>
<td>1190 (5.2)</td>
</tr>
<tr>
<td>Taiwan, Province of China</td>
<td>10 (0.1)</td>
<td>1188 (5.2)</td>
</tr>
<tr>
<td>France</td>
<td>102 (1.1)</td>
<td>1080 (4.7)</td>
</tr>
<tr>
<td>US</td>
<td>2537 (28.0)</td>
<td>1009 (4.4)</td>
</tr>
<tr>
<td>Japan</td>
<td>835 (9.2)</td>
<td>510 (2.2)</td>
</tr>
<tr>
<td>Italy</td>
<td>406 (4.5)</td>
<td>634 (2.8)</td>
</tr>
<tr>
<td>Spain</td>
<td>377 (4.2)</td>
<td>370 (1.6)</td>
</tr>
<tr>
<td>Denmark</td>
<td>146 (1.6)</td>
<td>160 (0.7)</td>
</tr>
<tr>
<td>Canada</td>
<td>134 (1.5)</td>
<td>175 (0.8)</td>
</tr>
<tr>
<td>Ireland</td>
<td>121 (1.3)</td>
<td>171 (0.8)</td>
</tr>
<tr>
<td>Total Others</td>
<td>870 (9.6)</td>
<td>7150 (31.4)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>3804 (42.0)</td>
<td>11,896 (52.2)</td>
</tr>
<tr>
<td>Male</td>
<td>4078 (45.0)</td>
<td>10,012 (44.0)</td>
</tr>
<tr>
<td>Unknown/No Data</td>
<td>1173 (13.0)</td>
<td>864 (3.8)</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>8213</td>
<td>22,089</td>
</tr>
<tr>
<td>Min-Max</td>
<td>0.58-17</td>
<td>0.01-17</td>
</tr>
<tr>
<td>Mean / Median</td>
<td>8.7/9</td>
<td>14.3/15</td>
</tr>
<tr>
<td>Age Range</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0-27 days</td>
<td>0</td>
<td>3 (0.01)</td>
</tr>
<tr>
<td>28 days to 23 months</td>
<td>11 (0.1)</td>
<td>69 (0.3)</td>
</tr>
<tr>
<td>2-11 years</td>
<td>8390 (92.7)</td>
<td>1410 (6.2)</td>
</tr>
<tr>
<td>12-17 years</td>
<td>654 (7.2)</td>
<td>21,290 (93.5)</td>
</tr>
<tr>
<td>Case Seriousness</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serious</td>
<td>1589 (17.5)</td>
<td>6956 (30.5)</td>
</tr>
<tr>
<td>Non-serious</td>
<td>7466 (82.5)</td>
<td>15,816 (69.5)</td>
</tr>
<tr>
<td>Case Outcome</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>17 (0.2)</td>
<td>65 (0.3)</td>
</tr>
<tr>
<td>Not resolved</td>
<td>1288 (14.2)</td>
<td>5013 (22.0)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>4205 (46.4)</td>
<td>11,567 (50.8)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>24 (0.3)</td>
<td>142 (0.6)</td>
</tr>
<tr>
<td>Unknown/No data</td>
<td>3521 (38.9)</td>
<td>5985 (26.3)</td>
</tr>
<tr>
<td>Presence of comorbidities&lt;sup&gt;e&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>289 (3.2)</td>
<td>642 (2.8)</td>
</tr>
<tr>
<td>No</td>
<td>8766 (96.8)</td>
<td>22,130 (97.2)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Only paediatric subject received BNT162b2.
<sup>b</sup> Due to rounding, sum of percentages may not match 100%.
<sup>c</sup> Cases retrieved applying the criteria in place to identify the reports involving the special populations of Immunocompromised patients, Patients with autoimmune or inflammatory disorders, Frail patients with comorbidities (e.g., COPD, diabetes, chronic neurological disease, cardiovascular disorders, active tuberculosis) described as special populations in Section 16.3.5.5, Section 16.3.5.6, and Section 16.3.5.7, respectively.

N=Number of cases; MC=Medically confirmed; Min=Minimum; Max=Maximum
### Table 21. Events Reported in ≥2%* Cases - Comparison of Paediatric Tris/Sucrose (Grey and Orange Cap) versus PBS/Sucrose Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>Tris/Sucrose (Grey and Orange Cap)</th>
<th>PBS/Sucrose</th>
</tr>
</thead>
<tbody>
<tr>
<td>MedDRA PT</td>
<td>N=9055 n (AERP, %)</td>
<td>N=22,772 n (AERP, %)</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications&lt;sup&gt;40&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poor quality product administered</td>
<td>1222 (13.5)</td>
<td>289 (1.3)</td>
</tr>
<tr>
<td>Product administration error</td>
<td>888 (9.8)</td>
<td>162 (0.7)</td>
</tr>
<tr>
<td>Product administered to patient of inappropriate age</td>
<td>503 (5.6)</td>
<td>1364 (6.0)</td>
</tr>
<tr>
<td>Overdose</td>
<td>478 (5.3)</td>
<td>125 (0.6)</td>
</tr>
<tr>
<td>Product preparation error</td>
<td>433 (4.8)</td>
<td>28 (0.1)</td>
</tr>
<tr>
<td>Underdose</td>
<td>289 (3.2)</td>
<td>241 (1.1)</td>
</tr>
<tr>
<td>Inappropriate schedule of product administration</td>
<td>243 (2.7)</td>
<td>952 (4.2)</td>
</tr>
<tr>
<td>Product preparation issue</td>
<td>234 (2.6)</td>
<td>99 (0.4)</td>
</tr>
<tr>
<td>Expired product administered&lt;sup&gt;b&lt;/sup&gt;</td>
<td>215 (2.4)</td>
<td>72 (0.3)</td>
</tr>
<tr>
<td>Vaccination error</td>
<td>208 (2.3)</td>
<td>258 (1.1)</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pyrexia</td>
<td>1222 (13.5)</td>
<td>3507 (15.4)</td>
</tr>
<tr>
<td>Vaccination site pain</td>
<td>1032 (11.4)</td>
<td>2073 (9.1)</td>
</tr>
<tr>
<td>Fatigue</td>
<td>371 (4.1)</td>
<td>2010 (8.8)</td>
</tr>
<tr>
<td>Chest pain</td>
<td>334 (3.7)</td>
<td>1959 (8.6)</td>
</tr>
<tr>
<td>Malaise</td>
<td>287 (3.2)</td>
<td>1000 (4.4)</td>
</tr>
<tr>
<td>Drug ineffective</td>
<td>218 (2.4)</td>
<td>601 (2.6)</td>
</tr>
<tr>
<td>Vaccination failure</td>
<td>46 (0.5)</td>
<td>1061 (4.7)</td>
</tr>
<tr>
<td>Chills</td>
<td>143 (1.6)</td>
<td>978 (4.3)</td>
</tr>
<tr>
<td>Chest discomfort</td>
<td>63 (0.7)</td>
<td>822 (3.6)</td>
</tr>
<tr>
<td>Asthenia</td>
<td>117 (1.3)</td>
<td>582 (2.6)</td>
</tr>
<tr>
<td>Pain</td>
<td>161 (1.8)</td>
<td>528 (2.3)</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>909 (10.0)</td>
<td>3576 (15.7)</td>
</tr>
<tr>
<td>Dizziness</td>
<td>355 (3.9)</td>
<td>2390 (10.5)</td>
</tr>
<tr>
<td>Syncope</td>
<td>211 (2.3)</td>
<td>685 (3.0)</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vomiting</td>
<td>683 (7.5)</td>
<td>1275 (5.6)</td>
</tr>
<tr>
<td>Nausea</td>
<td>386 (4.3)</td>
<td>1699 (7.5)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>348 (3.8)</td>
<td>506 (2.2)</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>236 (2.6)</td>
<td>448 (2.0)</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rash</td>
<td>501 (5.5)</td>
<td>1037 (4.6)</td>
</tr>
<tr>
<td>Urticaria</td>
<td>254 (2.8)</td>
<td>427 (1.9)</td>
</tr>
<tr>
<td>Pruritus</td>
<td>234 (2.6)</td>
<td>484 (2.1)</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain in extremity</td>
<td>278 (3.1)</td>
<td>768 (3.4)</td>
</tr>
<tr>
<td>Myalgia</td>
<td>172 (1.9)</td>
<td>992 (4.4)</td>
</tr>
</tbody>
</table>

<sup>40</sup> The number of PTs indicative of medication errors may not match with the numbers reported in Section 9.2 Medication Errors, since the adopted search criteria are different and the cases retrieved in this table involved only paediatric subjects.
Table 21. Events Reported in ≥2%* Cases - Comparison of Paediatric Tris/Sucrose (Grey and Orange Cap) versus PBS/Sucrose Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>Tris/Sucrose (Grey and Orange Cap)</th>
<th>PBS/Sucrose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arthralgia</td>
<td>N=9055 n (AERP, %)</td>
<td>N=22,772 n (AERP, %)</td>
</tr>
<tr>
<td>Product issues</td>
<td>123 (1.4)</td>
<td>491 (2.2)</td>
</tr>
<tr>
<td>Product temperature excursion issue</td>
<td>253 (2.8)</td>
<td>114 (0.5)</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td>227 (2.5)</td>
<td>1269 (5.6)</td>
</tr>
<tr>
<td>Dyspnoea</td>
<td>172 (1.9)</td>
<td>445 (2.0)</td>
</tr>
<tr>
<td>Cough</td>
<td>203 (2.2)</td>
<td>903 (4.0)</td>
</tr>
<tr>
<td>Lymphadenopathy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palpitations</td>
<td>85 (0.9)</td>
<td>691 (3.0)</td>
</tr>
<tr>
<td>Myocarditis</td>
<td>38 (0.4)</td>
<td>632 (2.8)</td>
</tr>
<tr>
<td>Tachycardia</td>
<td>82 (0.9)</td>
<td>467 (2.1)</td>
</tr>
<tr>
<td>Total number of events</td>
<td>14,457</td>
<td>38,010</td>
</tr>
</tbody>
</table>

a. Reporting proportion (% of total PM cases) in one or both paediatric populations.
b. Majority of the cases reported uncertain expiry dates.

Conclusion

Overall, more than 45% of the Tris/Sucrose cases was reported in paediatric subjects; the most frequently reported AEs in this population do not differ from the paediatric PBS/Sucrose formulation. A higher percentage of medication error cases was reported in the Tris/Sucrose paediatric group and this may reflect initial difficulties in managing the new formulation (Section 9.2, Medication Errors). Routine pharmacovigilance activities to mitigate these medication errors are listed in the approved version 5.0 of the EU-RMP adopted on 10 March 2022.

Based on the review of the cases reported with Tris/Sucrose formulation, no new safety issues were identified.

6.3.1.2.3. Booster Doses (Third and Fourth Doses)

A summary of the approvals of booster doses for the different age groups and associated regulatory procedures is provided in Table 22 for the reporting period.

First booster is indeed the third dose after completing a 2-dose primary series of BNT162b2 (as a homologous booster dose), or the first booster following completion of primary vaccination with another authorised COVID-19 vaccine (as a heterologous booster dose).

Second booster is indeed the fourth dose after completing a 2-dose primary series and the first booster dose with BNT162b2 (as a homologous booster dose) or the second booster dose following completion of primary vaccination and of a first booster dose with any authorised COVID-19 vaccine (as a heterologous booster dose).
<table>
<thead>
<tr>
<th>Age Group</th>
<th>Procedure and Description</th>
<th>Approval Date</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First booster</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16+ years</td>
<td>EMEA/H/C/005735/II/0104*</td>
<td>CHMP Opinion: 22 April 2022&lt;br&gt;EC decision: 04 May 2022</td>
</tr>
<tr>
<td></td>
<td>• Heterologous vaccination (both in the primary series and for booster vaccinations) PI update based on cumulative review of available immunogenicity, safety and efficacy data.</td>
<td>Procedure ongoing pending approval.</td>
</tr>
<tr>
<td></td>
<td>• Further PI update to implement booster (dose) interval reduction from 6 months to 3 months per Agency request based on the totality of available evidence (not MAH’s own clinical data).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>EMEA/H/C/005735/II/0139&lt;br&gt;PI update regarding individuals 16+ years based on six-month post (booster) dose 3 data from Studies C4591001 and C4591031 data.</td>
<td></td>
</tr>
<tr>
<td>12-15 years</td>
<td>EMEA/H/C/005735/II/0104*</td>
<td>CHMP Opinion: 22 April 2022&lt;br&gt;EC decision: 04 May 2022</td>
</tr>
<tr>
<td></td>
<td>• Heterologous vaccination (both in the primary series and for booster vaccinations) PI update based on cumulative review of available immunogenicity, safety and efficacy data.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Further PI update to implement booster (dose) interval reduction from 6 months to 3 months per Agency request based on the totality of available evidence (not MAH’s own clinical data).</td>
<td></td>
</tr>
<tr>
<td></td>
<td>EMEA/H/C/005735/II/0111&lt;br&gt;PI update to lower the age of the booster dose to patients 12 years of age and older based on RWE data from MoH Israel.</td>
<td></td>
</tr>
<tr>
<td>5-11 years</td>
<td>EMEA/H/C/005735/II/0129&lt;br&gt;5-11 year PI update – one month post dose (booster) dose 3 (1MPD3) based on clinical study C4591007 data.</td>
<td>CHMP Opinion: 24 February 2022&lt;br&gt;EC decision: 28 February 2022</td>
</tr>
<tr>
<td></td>
<td>Procedure ongoing pending approval.</td>
<td></td>
</tr>
<tr>
<td><strong>Second booster</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16+ years and 12-15 years</td>
<td>EMEA/H/C/005735/II/0140&lt;br&gt;Bivalent Original/Omicron BA.1 as from 12+ years- Rolling submission.</td>
<td>Procedure ongoing pending approval.</td>
</tr>
<tr>
<td>5-11 years</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 22. Summary of Approval of Booster Doses in the Reporting Period

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Procedure and Description</th>
<th>Approval Date</th>
</tr>
</thead>
</table>
| First booster | **16+ years**  
US FDA lowered the authorised dosing interval of the homologous booster dose to at least 5 months after completion of the primary series.                                                                                                   | 03 January 2022     |
|            | **12-15 years**  
US FDA authorised the use of the vaccine as a single booster dose at least 5 months after completion of a primary series with BNT162b2.                                                                                               | 03 January 2022     |
|            | **5-11 years**  
US FDA authorised a third primary series dose of the vaccine administered at least 28 days following the two-dose regimen of BNT162b2 in individuals 5 through 11 years of age who have undergone solid organ transplantation or are diagnosed with conditions that are considered to have an equivalent level of immunocompromise.  
EUA was granted for a single booster dose at least 5 months after completing a primary series with BNT162b2.                              | 17 May 2022         |
| Second booster | **50+ years**  
EUA was granted for the use of BNT162b2 as second booster for individuals 50+ years at least 4 months after receipt of a first booster dose of any FDA-authorised or approved COVID-19 vaccine.                                                                          | 29 March 2022       |
|            | **12+ years**  
The US FDA issues an EUA to approve the use of BNT162b2 as second booster at least 4 months after a first booster dose with any FDA authorised or approved COVID-19 vaccine for 12+ years individuals, who have undergone solid organ transplantation or have been diagnosed with conditions that are considered to have an equivalent level of immunocompromise. | 29 March 2022       |
|            | **5-11 years**  | -                                                                                                                                  | -                   |

*: The same procedure was issued for two different age groups due to a combined claim in the PI for both age groups (Type II variation 104 for heterologous boosting).

Analysis of Booster Doses

Search criteria: Dose number equal to 3 or Dose number equal to 4 OR Dose Description containing the word "BOOSTER" OR LLT equal to BOOSTER.

The search yielded 119,601 cases (491 CT cases and 119,110 PM cases).

Upon review,

- 455 cases (1 CT and 454 PM) involving babies were excluded due to indirect exposure (transplacental/transmammary) to BNT162b2.
- 906 PM cases were determined to be non-contributory and were not included in the discussion since in these cases the booster dose administered was not BNT162b2 (904 cases) or the case did not contain any information that the individuals received a booster dose (2 cases).
**Overall**

**CT data**

Table 23. Selected Case Characteristics of CT Data Involving Participants Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022)

<table>
<thead>
<tr>
<th>Case characteristics</th>
<th>Overall participants who received at least one booster dose of BNT162b2</th>
<th>Participants who received single booster dose of BNT162b2</th>
<th>Participants who received &gt;1 booster doses of BNT162b2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cases</td>
<td>490 (BNT162b2 [441], blinded therapy [46] and placebo [3])</td>
<td>423 (BNT162b2 [375], blinded therapy [45] and placebo [3])</td>
<td>67 (BNT162b2 [66] and blinded therapy [1])</td>
</tr>
<tr>
<td>Cases by protocol ID41</td>
<td>C4591001 (347), C4591031 (101), C4591024 (16), C4591007 (14), BNT162-17 (11) and C4591030 (1)</td>
<td>C4591001 (301), C4591031 (86), C4591024 (16), BNT162-17 (10) and C4591007 (10)</td>
<td>C4591001 (46), C4591031 (15), C4591007 (4), BNT162-17 and C4591030 (1 each)</td>
</tr>
<tr>
<td>Country of incidence (≥2 cases)</td>
<td>US (358), Argentina (73), Brazil (31), Germany (15), South Africa (4), and Israel (2)</td>
<td>US (310), Argentina (67), Brazil (25), Germany (14), and South Africa (3)</td>
<td>US (48), Argentina, Brazil (6 each), and Israel (2)</td>
</tr>
<tr>
<td>Participants’ gender</td>
<td>female (228) and male (262)</td>
<td>female (198) and male (225)</td>
<td>female (30) and male (37)</td>
</tr>
<tr>
<td>Participants’ age in years</td>
<td>n = 490; range: 1.4-87.0; mean = 54.8; median = 61.0</td>
<td>n = 423; range: 1.4-87.0; mean = 55.2; median = 61.0</td>
<td>n = 67; range: 2.0-83.0; mean = 51.8; median = 58.0</td>
</tr>
<tr>
<td>Case outcome</td>
<td>fatal (29), resolved/resolving (354), resolved with sequelae (33), not resolved (73), and unknown (1)</td>
<td>fatal (22), resolved/resolving (304), resolved with sequelae (30), not resolved (66), and unknown (1)</td>
<td>fatal (7), resolved/resolving (50), resolved with sequelae (3), and not resolved (7)</td>
</tr>
<tr>
<td>Most frequently reported PTs41 (≥2%)</td>
<td>Condition aggravated (20), Atrial fibrillation (14), Pneumonia (12), Appendicitis, Cerebrovascular accident (11 each), and Prostate cancer (10)</td>
<td>Condition aggravated (19), Atrial fibrillation (12), Cerebrovascular accident (11), Appendicitis, Prostate cancer (10 each), and Pneumonia (9)</td>
<td>Pneumonia (3), Atrial fibrillation, Febrile convulsion, Injury, Osteoarthritis, and Syncope (2 each)</td>
</tr>
</tbody>
</table>

a. Includes all SAEs irrespective of relatedness to BNT162b2/blinded therapy/placebo

---

41 Please refer to Section 7.2 Ongoing Clinical Trials for details about these clinical trials and to Section 17.2 Newly Identified Information on Efficacy and Effectiveness for the newly identified information on efficacy and effectiveness from the interim analysis of studies C4591031 and C4591007.
# Post-Authorisation Data

## Table 24. Selected Case Characteristics of Post-Authorisation Data Involving Individuals Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022)

<table>
<thead>
<tr>
<th>Case characteristics</th>
<th>Overall individuals who received at least one booster dose of BNT162b2</th>
<th>Individuals who received single booster dose of BNT162b2&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Individuals who received &gt;1 booster doses of COVID-19 vaccine (the latest booster dose was BNT162b2)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Individuals who received unknown booster dose(s) of BNT162b2&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number of cases</strong></td>
<td>117,750</td>
<td>106,889</td>
<td>3427</td>
<td>7434</td>
</tr>
<tr>
<td><strong>Total number of events</strong></td>
<td>484,959</td>
<td>451,970</td>
<td>13,627</td>
<td>19,362</td>
</tr>
<tr>
<td><strong>MC/NMC cases</strong></td>
<td>MC: 28,695; NMC: 89,055</td>
<td>MC: 25,113; NMC: 81,776</td>
<td>MC: 1104; NMC: 2323</td>
<td>MC: 2478; NMC: 4956</td>
</tr>
<tr>
<td><strong>Country of incidence (≥2%)</strong></td>
<td>Germany (25,532), Netherlands (21,182), UK (18,523), France (10,285), US (7953), Austria (6391), and Japan (4930)</td>
<td>Germany (24,922), Netherlands (20,795), UK (17,614), Austria (6382), US (5773), France (5028), and Japan (4895)</td>
<td>US (1403), UK (639), Germany (372), France (192), Netherlands (182), Canada (103), and Sweden (76)</td>
<td>France (5065), US (777), Ireland (528), UK (270), Germany (238), and Netherlands (205)</td>
</tr>
<tr>
<td><strong>Subjects’ gender</strong></td>
<td>female (80,563), male (33,585), and unknown (3602)</td>
<td>female (73,716), male (30,196), and unknown (2977)</td>
<td>female (1936), male (1215), and unknown (276)</td>
<td>female (4911), male (2174), and unknown (349)</td>
</tr>
<tr>
<td><strong>Subjects’ age in years</strong></td>
<td>n = 108,038; range: 0.3-120.0; mean = 45.5; median = 43.0</td>
<td>n = 98,367; range: 0.3-120.0; mean = 44.8; median = 42.0</td>
<td>n = 2894; range: 2.0-102.0; mean = 65.4; median = 68.0</td>
<td>n = 6777; range: 0.5-103.0; mean = 47.0; median = 46.0</td>
</tr>
<tr>
<td><strong>Case seriousness</strong></td>
<td>serious = 42,918; non-serious = 74,832</td>
<td>serious = 39,781; non-serious = 67,108</td>
<td>serious = 1722; non-serious = 1705</td>
<td>serious = 1415; non-serious = 6019</td>
</tr>
<tr>
<td><strong>Case outcome</strong></td>
<td>fatal (1225), resolved/resolving (41,604), resolved with sequelae (1805), not resolved (51,914), and unknown (21,202)</td>
<td>fatal (1093), resolved/resolving (37,881), resolved with sequelae (1722), not resolved (47,915), and unknown (18,278)</td>
<td>fatal (72), resolved/resolving (894), resolved with sequelae (32), not resolved (958), and unknown (1471)</td>
<td>fatal (60), resolved/resolving (2829), resolved with sequelae (51), not resolved (3041), and unknown (1453)</td>
</tr>
<tr>
<td><strong>Most frequently reported PTs (≥2%)</strong></td>
<td>Immunisation&lt;sup&gt;d&lt;/sup&gt; (25,650), Headache (24,152), Off label use (22,894), Fatigue (21,550), Interchange of vaccine product&lt;sup&gt;c&lt;/sup&gt; (20,384), Pyrexia (16,639),</td>
<td>Immunisation&lt;sup&gt;d&lt;/sup&gt; (23,292), Headache (23,112), Off label use (20,889), Fatigue (20,740), Interchange of vaccine product&lt;sup&gt;c&lt;/sup&gt; (19,823), Pyrexia (15,894),</td>
<td>Off label use (1627), Immunisation&lt;sup&gt;d&lt;/sup&gt; (1403), COVID-19 (704), Drug ineffective (568), Incorrect dose administered (512), Fatigue (377), Headache (348),</td>
<td>Immunisation&lt;sup&gt;d&lt;/sup&gt; (955), Immunisation reaction (873), Lymphadenopathy (858), Headache (692), Fatigue (433), Pyrexia (401), Influenza like illness (396),</td>
</tr>
</tbody>
</table>
Table 24. Selected Case Characteristics of Post-Authorisation Data Involving Individuals Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022)

<table>
<thead>
<tr>
<th>Case characteristics</th>
<th>Overall individuals who received at least one booster dose of BNT162b2</th>
<th>Individuals who received single booster dose of BNT162b2*</th>
<th>Individuals who received &gt;1 booster doses of COVID-19 vaccine (the latest booster dose was BNT162b2)*</th>
<th>Individuals who received unknown booster dose(s) of BNT162b2*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymphadenopathy (16,442), Vaccination site pain (15,397), Myalgia (15,323), Malaise (14,845), Chills (13,836), Arthralgia (10,331), Nausea (10,055), COVID-19 (9701), Pain in extremity (6792), Dizziness (6314), Drug ineffective (5830), Vaccination site swelling (5289), Dyspnoea (5079), Vaccination failure (4942), Pain (4748), Chest pain (3988), Limb discomfort (3472), Vaccination site lymphadenopathy (3433), Vaccination site erythema (3386), Vaccination site inflammation (3329), Axillary pain (3141), Rash (3129), Asthenia (3007), Palpitations (2947), Vaccination site warmth (2288), Parasthesia (2813), Vomiting (2740), Tachycardia (2641), Diarrhoea (2517), and Heavy menstrual bleeding (2368).</td>
<td>Lymphadenopathy (15,510), Vaccination site pain (14,853), Myalgia (14,789), Malaise (14,333), Chills (13,377), Arthralgia (9806), Nausea (9564), COVID-19 (8789), Pain in extremity (6213), Dizziness (5956), Vaccination site swelling (5190), Drug ineffective (5054), Dyspnoea (4775), Vaccination failure (4706), Pain (4546), Chest pain (3681), Limb discomfort (3403), Vaccination site lymphadenopathy (3398), Vaccination site erythema (3281), Vaccination site inflammation (3252), Axillary pain (3022), Rash (2899), Palpitations (2778), Vaccination site warmth (2777), Parasthesia (2568), Asthenia (2542), Tachycardia (2490), Vomiting (2479), Diarrhoea (2319), Heavy menstrual bleeding (2253), and Influenza (2226).</td>
<td>Pyrexia (344), Interchange of vaccine products* (267), Chills (239), Vaccination failure (231), Pain in extremity (203), Pain (202), Malaise (199), Myalgia, Vaccination site pain (188 each), Poor quality product administered (185), Arthralgia (172), Nausea (169), Dizziness (159), Product administration error (116), Dyspnoea (104), Asthenia (99), Vomiting (96), Suspected COVID-19 (85), Expired product administered (81), Diarrhoea (80), Chest pain (76), Lymphadenopathy (74), and Feeling abnormal (69).</td>
<td>Off-label use (378), Pain in extremity (376), Asthenia (366), Vaccination site pain (356), Arthralgia (353), Myalgia (346), Nausea (322), Malaise (313), Interchange of vaccine products* (294), Chest pain (231), Drug ineffective (228), Poor quality product administered (226), Chills, Dyspnoea, Pain (220 each), Dizziness (219), Parasthesia (209), COVID-19 (208), Herpes zoster (204), Menstrual disorder (199), Rash (168), Tinnitus (167), and Vomiting (165).</td>
<td></td>
</tr>
</tbody>
</table>

a. Indicates individuals who received one additional (booster) dose of BNT162b2 after completing the primary series of any authorised COVID-19 vaccine

b. Indicates individuals who received 2 or more additional (booster) doses of COVID-19 vaccine (the latest booster dose was BNT162b2) after completing the primary series of any authorised COVID-19 vaccine

CONFIDENTIAL
Page 83
Table 24. Selected Case Characteristics of Post-Authorisation Data Involving Individuals Who Received Booster Dose(s) of BNT162b2 During The Reporting Period (19 December 2021 through 18 June 2022)

<table>
<thead>
<tr>
<th>Case characteristics</th>
<th>Overall individuals who received at least one booster dose of BNT162b2</th>
<th>Individuals who received single booster dose of BNT162b2*</th>
<th>Individuals who received &gt;1 booster doses of COVID-19 vaccine (the latest booster dose was BNT162b2)*</th>
<th>Individuals who received unknown booster dose(s) of BNT162b2*</th>
</tr>
</thead>
</table>

c. Indicates individuals who received unknown additional (booster) dose(s) of BNT162b2 after completing the primary series of any authorised COVID-19 vaccine.
d. PT Immunisation, initially selected per case processing conventions to retrieve cases reporting third/booster doses was no longer applied after 31 January 2022. Following that, “booster” was added in the dose description field.
e. PT Interchange of vaccine products, initially selected per case processing conventions to retrieve cases reporting heterologous administration of third/booster doses was no longer applied after 31 January 2022. After that, the LT Interchange of vaccine products was added in the relevant medical history section.
Of the relevant 490 CT cases, all participants received homologous doses schedule (primary series and booster with BNT162b2). While among the relevant 117,750 PM cases, 47,759 cases received homologous doses schedule, 23,252 cases received heterologous doses schedule (primary series with a specified non-BNT162b2 COVID-19 vaccine and booster with BNT162b2), and 46,739 cases received booster doses of BNT162b2 administered after an unspecified primary COVID-19 vaccination series. The details of these cases are as follows:

**Homologous doses schedule (primary series and booster with BNT162b2)**

**Clinical Trial Data**

- Number of cases: 490 (BNT162b2 [441], blinded therapy [46] and placebo [3]) (73.4% of 668 cases, the total CT dataset). Please refer Table 23 for further details.

**Post-Authorisation Data**

- Number of cases: 47,759 (9.4% of 507,683 cases, the total PM dataset; 40.6% of the PM booster dataset).
- MC cases (13,848), NMC cases (33,911).
- Country of incidence (≥2%): Netherlands (15,076), UK (6998), US (4904), Austria (3222), Germany (2855), France (2377), Japan (1930), Spain (1131), Italy (1030), and Belgium (1008).
- Subjects' gender: female (33,157), male (13,463) and unknown (1139).
- Subjects' age in years (n = 43,778), range: 0.5–120.0, mean: 45.2, median: 41.0.
- Case outcome: fatal (550), resolved/resolving (15,853), resolved with sequelae (480), not resolved (21,330), and unknown (9546).
- In 550 cases (reporting 1604 events with a fatal outcome), the reported causes of death (≥20 cases) were coded to the PTs COVID-19 (86), Vaccination failure (62), Cardiac arrest (52), COVID-19 pneumonia (46), Sudden death (31), Cardio-respiratory arrest (27), Cardiac failure, Myocardial infarction (21 each), Cerebral haemorrhage and Pulmonary embolism (20 each). Of note, in 99 cases limited information regarding the cause of death was provided (PT Death).
- Medical history (n = 16,928): the most frequently (≥2% of homologous doses schedule PM cases) reported medical conditions included Disease risk factor (2101), Hypertension (2081), Asthma (1015), Drug hypersensitivity (827), Hypothyroidism (526), Seasonal allergy (509), Food allergy (483), Diabetes mellitus (480), Hypersensitivity (478), Depression (418), and Immunodeficiency (336).
- COVID-19 Medical history (n = 3615): COVID-19 (2165), Suspected COVID-19 (1437), Post-acute COVID-19 syndrome (33), Exposure to SARS-CoV-2 (20), SARS-CoV-2 test positive (11), COVID-19 pneumonia (7), Asymptomatic COVID-19, Coronavirus infection (4 each), and Occupational exposure to SARS-CoV-2 (3).
- Number of events: 190,262.
• Event seriousness42: serious (63,265), non-serious (127,091).

• The most reported (≥2% of homologous doses schedule PM cases) PTs were Headache (10,390), Immunosuppression43 (9993), Fatigue (9945), Malaise (8187), Myalgia (7932), COVID-19 (7123), Pyrexia (6602), Vaccination site pain (6486), Chills (6360), Lymphadenopathy (6287), Arthralgia (5138), Vaccination failure (4891), Nausea (4672), Drug ineffective (3064), Vaccination site swelling (2813), Pain in extremity (2459), Vaccination site inflammation (2312), Vaccination site lymphadenopathy (2099), Vaccination site warmth (1900), Pain (1887), Dyspnoea (1815), Dizziness (1794), Vaccination site erythema (1774), Chest pain (1616), Axillary pain (1385), Off label use (1171), Palpitations (1112), and Heavy menstrual bleeding (1034).

**Heterologous doses schedule (primary series with a specified non-BNT162b2 COVID-19 vaccine and booster with BNT162b2)**

**Post-Authorisation Data**

• Number of cases: 23,252 (4.6% of 507,683 cases, the total PM dataset; 19.7% of the PM booster dataset).

• MC cases (3665), NMC cases (19,587).

• Country of incidence (≥2%): UK (9601), Netherlands (5987), Germany (1801), France (1192), Belgium (500), and US (496).

• Subjects' gender: female (16,361), male (6296) and unknown (595).

• Subjects' age in years (n = 20,855), range: 0.3 – 102.0, mean: 46.5, median: 45.0.

• Case outcome: fatal (162), resolved/resolving (7179), resolved with sequelae (322), not resolved (12,429), and unknown (3160).

• In 162 cases (reporting 781 events with a fatal outcome), the reported causes of death (≥5 cases) were coded to the PTs Interchange of vaccine products (21), Off label use (20), Cardiac arrest (14), Sudden death (12), Cerebrovascular accident, Dyspnoea, Immunisation (11 each), Pulmonary embolism (9), Malaise, Myocardial infarction (7 each), Cerebral haemorrhage, COVID-19, Drug ineffective, Myocardial ischaemia, Pneumonia, Thrombosis (6 each), Myocarditis, Oxygen saturation decreased, and Septic shock (5 each). Of note, in 32 cases limited information regarding the cause of death was provided (PT Death).

• Medical history (n = 10,734): the most frequently (≥2%) reported medical conditions included Disease risk factor (1596), Hypertension (888), Interchange of vaccine products (805), Asthma (684), Immunodeficiency (502), Hypersensitivity (289), Diabetes

42 Multiple episodes of the same event were reported with a different seriousness in some cases hence the sum of the events seriousness exceeds the total number of events.

43 PT selected per case processing conventions to collect cases reporting third/booster doses; as of 31 January 2022 this convention was no more applicable.
mellitus (278), Hypothyroidism (271), Steroid therapy (253), Depression (241), Drug hypersensitivity (229) and Clinical trial participant (224).

- COVID-19 Medical history \( (n = 2649) \): Suspected COVID-19 (1346), COVID-19 (1340), Post-acute COVID-19 syndrome (22), SARS-CoV-2 test positive (18), COVID-19 pneumonia (6), Coronavirus infection (4), Asymptomatic COVID-19 (2) and Exposure to SARS-CoV-2 (1).

- Among the 23,252 cases reporting administration of heterologous booster dose(s) of BNT162b2 following a specified non-BNT162b2 COVID-19 vaccine, the previous vaccine series consisted of:
  - 9651 subjects immunised with AstraZeneca vaccine;
  - 5334 subjects immunised with Moderna vaccine;
  - 5214 subjects immunised with unknown non-Pfizer-BioNTech COVID-19 vaccine;
  - 2427 subjects immunised with Johnson and Johnson vaccine;
  - 417 subjects immunised with Coronavac (Sinovac) vaccine;
  - 88 subjects immunised with Sinopharm vaccine;
  - 76 subjects immunised with Sputnik vaccine;
  - 29 subjects immunised with Novavax vaccine;
  - 9 subjects immunised with Fiocruz vaccine;
  - 2 subjects each immunised with Medicago-Clinical study and Medigen vaccine;
  - 1 subject each immunised with Cansino vaccine, Covaxin vaccine, and Valneva vaccine.

- Number of events: 140,835.

- Event seriousness\(^{42}\): serious (61,291), non-serious (79,594).

- The most reported \( (> 2\% \) of heterologous dose schedule PM cases) PTs were Off label use (20,437), Interchange of vaccine products\(^{43}\) (20,376), Immunisation (9982), Headache (5229), Fatigue (4854), Myalgia (3412), Malaise (3362), Pyrexia (3144), Lymphadenopathy (3139), Vaccination site pain (2926), Chills (2918), Arthralgia (2578), Nausea (2382), Pain in extremity (1848), Pain (1362), Drug ineffective (1223), COVID-19 (1192), Dizziness (1161), Vaccination site swelling (1140), Dyspnoea (1091), Chest pain (991), Axillary pain (948), Vaccination site inflammation (942), Palpitations (850), Vaccination site warmth (830), Vaccination site lymphadenopathy (815), Vaccination site erythema (751), Pruritus (630), Rash (617), Swelling (610), Heavy menstrual
bleeding (608), Asthenia, Diarrhoea (565 each), Peripheral swelling (557), Paraesthesia (548), Vomiting (516), and Tachycardia (462).

**Booster doses of BNT162b2 administered after an unspecified primary COVID-19 vaccination series**

**Post-Authorisation Data**

- Number of cases: 46,739 (9.2% of 507,683 cases, the total PM dataset; 39.7% of the PM booster dataset).
- MC cases (11,182), NMC cases (35,557).
- Country of incidence (≥2%): Germany (20,876), France (6716), Japan (2922), Austria (2833), US (2553), and UK (1924).
- Subjects' gender: female (31,045), male (13,826) and unknown (1868).
- Subjects' age in years (n = 43,405), range: 1.0 – 104.0, mean: 45.3, median: 43.0.
- Case outcome: fatal (513), resolved/resolving (18,572), resolved with sequelae (1003), not resolved (18,155), and unknown (8496).
- In 513 cases (reporting 1318 events with a fatal outcome), the reported causes of death (≥15 cases) were coded to the PTs Cardiac arrest (38), Cardio-respiratory arrest, Myocardial infarction (35 each), Sudden death (25), Pulmonary embolism (24), Cardiac failure (22), Dyspnoea (19), Cerebral haemorrhage (17), and Acute myocardial infarction (15). Of note, in 150 cases limited information regarding the cause of death was provided (PT Death).
- Medical history (n = 11,782): the most frequently (≥2%) reported medical conditions included Hypertension (1856), Asthma (836), Drug hypersensitivity (642), Seasonal allergy (625), Hypersensitivity (456), Diabetes mellitus (426), Hypothyroidism (384), Obesity (336), Food allergy (314), Type 2 diabetes mellitus (295), Atrial fibrillation and Depression (241 each).
- Number of events: 153,862.
- Event seriousness: serious (35,762), non-serious (118,147).
- The most reported (≥2%) PTs were Headache (8533), Lymphadenopathy (7016), Pyrexia (6893), Fatigue (6751), Vaccination site pain (5985), Immunisation (5675), Chills (4558), Myalgia (3979), Dizziness (3359), Malaise (3296), Nausea (3001), Limb discomfort (2798), Arthralgia (2615), Pain in extremity (2485), Dyspnoea (2173), Influenza (1877), Rash (1738), Drug ineffective (1563), Tachycardia (1577), Asthenia (1512), Pain (1499), Paraesthesia (1416), COVID-19 (1386), Chest pain (1381), Vaccination site swelling (1336), Vomiting (1325), Off label use (1286), Herpes...
zoster (1203), Menstrual disorder (1101), Diarrhoea (1062), Poor quality product administered (1061), Feeling hot (1017), Immunisation reaction (1016), Influenza like illness (1006), and Palpitations (985).

Analysis booster doses versus primary vaccination series

There were 117,750 PM cases of subjects who received at least one booster dose of BNT162b2.

Among the 117,750 PM cases,

- 106,889 PM cases involved subjects who received single booster dose of BNT162b2
- 3427 PM cases involved subjects who received >1 booster doses of COVID-19 vaccine (the latest booster dose was BNT162b2) and
- 7434 cases involved subjects who received unknown booster dose(s) of BNT162b2.
- The most frequently (≥2%) reported clinical AEs 44 in PM cases of subjects who received the booster dose(s) of BNT162b2 are largely reflective of reactogenicity and events associated with the immunisation process.
- The most frequently (≥2%) reported clinical AEs in PM cases of subjects who received booster dose(s) of BNT162b2 were consistent with those reported in subjects receiving primary vaccination series, as shown in Table 25.
- A higher AERP rate 45 was observed for 9 PTs (Lymphadenopathy [14.0% vs 3.8%], Malaise [12.6% vs 4.6%], Chills [11.8% vs 5.1%], Vaccination site swelling [4.5% vs 1.4%], Vaccination site erythema [2.9% vs 1.0%], Vaccination site lymphadenopathy [2.9% vs 0.2%], Vaccination site inflammation [2.8% vs 0.3%], Axillary pain [2.7% vs 0.5%], and Vaccination site warmth [2.4% vs 0.3%]) was observed in subjects who received the booster dose(s) of BNT162b2 compared to subjects receiving the primary vaccination series. This is consistent with the known BNT162b2 safety profile (as per the RSI), where higher rates of lymphadenopathy and reactogenicity reactions in booster doses versus primary doses were observed in interventional clinical studies.
- No clinically significant differences were noted in the other events.

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44 The PT Immunisation, Interchange of vaccine products, and Off label use are not included in the Table 25.

45 A PT was considered to have a higher AERP rate if the ratio (AERP of the PT in the booster dataset/AERP of the PT in the primary series dataset) is ≥2.
Table 25. Comparison of clinical AEs reported in ≥2% Booster Dose(s) vs Primary Series Cases

<table>
<thead>
<tr>
<th>PT Decode (Event)</th>
<th>Booster dose(s) Cases N = 117,750</th>
<th>Primary Series Cases N = 385,599</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>AERF$^a$ (%)</td>
</tr>
<tr>
<td>Headache</td>
<td>24,152</td>
<td>20.5</td>
</tr>
<tr>
<td>Fatigue</td>
<td>21,550</td>
<td>18.3</td>
</tr>
<tr>
<td>Pyrexia</td>
<td>16,639</td>
<td>14.1</td>
</tr>
<tr>
<td>Lymphadenopathy</td>
<td>16,442</td>
<td>14.0</td>
</tr>
<tr>
<td>Vaccination site pain</td>
<td>15,397</td>
<td>13.1</td>
</tr>
<tr>
<td>Myalgia</td>
<td>15,323</td>
<td>13.0</td>
</tr>
<tr>
<td>Malaise</td>
<td>14,845</td>
<td>12.6</td>
</tr>
<tr>
<td>Chills</td>
<td>13,836</td>
<td>11.8</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>10,331</td>
<td>8.8</td>
</tr>
<tr>
<td>Nausea</td>
<td>10,055</td>
<td>8.5</td>
</tr>
<tr>
<td>COVID-19</td>
<td>9,701</td>
<td>8.2</td>
</tr>
<tr>
<td>Pain in extremity</td>
<td>6,792</td>
<td>5.8</td>
</tr>
<tr>
<td>Dizziness</td>
<td>6,314</td>
<td>5.4</td>
</tr>
<tr>
<td>Drug ineffective</td>
<td>5,850</td>
<td>5.0</td>
</tr>
<tr>
<td>Vaccination site swelling</td>
<td>5,289</td>
<td>4.5</td>
</tr>
<tr>
<td>Dyspnoea</td>
<td>5,079</td>
<td>4.3</td>
</tr>
<tr>
<td>Vaccination failure</td>
<td>4,945</td>
<td>4.2</td>
</tr>
<tr>
<td>Pain</td>
<td>4,748</td>
<td>4.0</td>
</tr>
<tr>
<td>Chest pain</td>
<td>3,988</td>
<td>3.4</td>
</tr>
<tr>
<td>Limb discomfort</td>
<td>3,472</td>
<td>2.9</td>
</tr>
<tr>
<td>Vaccination site lymphadenopathy</td>
<td>3,433</td>
<td>2.9</td>
</tr>
<tr>
<td>Vaccination site erythema</td>
<td>3,386</td>
<td>2.9</td>
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<tr>
<td>Vaccination site inflammation</td>
<td>3,329</td>
<td>2.8</td>
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<tr>
<td>Axillary pain</td>
<td>3,141</td>
<td>2.7</td>
</tr>
<tr>
<td>Rash</td>
<td>3,129</td>
<td>2.7</td>
</tr>
<tr>
<td>Asthenia</td>
<td>3,007</td>
<td>2.6</td>
</tr>
<tr>
<td>Palpitations</td>
<td>2,947</td>
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</tr>
<tr>
<td>Vaccination site warmth</td>
<td>2,828</td>
<td>2.4</td>
</tr>
<tr>
<td>Paraesthesia</td>
<td>2,813</td>
<td>2.4</td>
</tr>
<tr>
<td>Vomiting</td>
<td>2,740</td>
<td>2.3</td>
</tr>
<tr>
<td>Tachycardia</td>
<td>2,641</td>
<td>2.2</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>2,517</td>
<td>2.1</td>
</tr>
<tr>
<td>Heavy menstrual bleeding</td>
<td>2,368</td>
<td>2.0</td>
</tr>
</tbody>
</table>

a. Calculated as n/N.

Conclusion

Based on the review of the cases reported with the booster dose(s), no new safety issues were identified.

6.3.1.1.2.4. Batch-Related issues

The most frequently reported lot numbers in PM case reports (≥3000 cases) are listed in Table 26 below.
Table 26. Most Frequently Reported Lot Numbers

<table>
<thead>
<tr>
<th>Lot Number</th>
<th>Number of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>FD6840</td>
<td>9639</td>
</tr>
<tr>
<td>FE6208</td>
<td>8777</td>
</tr>
<tr>
<td>FD4555</td>
<td>7568</td>
</tr>
<tr>
<td>FD0168</td>
<td>7566</td>
</tr>
<tr>
<td>FD1921</td>
<td>7499</td>
</tr>
<tr>
<td>FR8392</td>
<td>6749</td>
</tr>
<tr>
<td>FF2382</td>
<td>5860</td>
</tr>
<tr>
<td>FF0680</td>
<td>4582</td>
</tr>
<tr>
<td>FC0681</td>
<td>4416</td>
</tr>
<tr>
<td>FC2493</td>
<td>4358</td>
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<tr>
<td>EY7015</td>
<td>4195</td>
</tr>
<tr>
<td>FA4598</td>
<td>4168</td>
</tr>
<tr>
<td>FG7911</td>
<td>4035</td>
</tr>
<tr>
<td>EY3014</td>
<td>3886</td>
</tr>
<tr>
<td>FT3318</td>
<td>3731</td>
</tr>
<tr>
<td>FH9951</td>
<td>3377</td>
</tr>
</tbody>
</table>

*The lots/batches reported in the table were all manufactured at Pfizer Puurs (Belgium).*

The AEs most frequently reported (≥ 4%) with these lot numbers do not differ from those reported in the overall incremental dataset except for the events coded to the PT Product administered to patient of inappropriate age.

During the current reporting period, on a total of 331,982 PM cases reporting a lot/batch number, there were 8068 PM cases including events coded to the PT Product administered to patient of inappropriate age, representing 1.6% of the overall incremental dataset and 2.4% of the total number of cases with lot numbers. These 8068 cases are also included in Section 9.2 Medication Errors or Section 16.3.4.6 Off-Label Use. The majority of these cases (5988 cases with unknown age) was non-serious and non-medically confirmed and originated from lot FR8392 which contains Tris/Sucrose presentation, 10 micrograms/dose, formulation indicated for age 5 years to < 12 years, per the current BNT162b2 RSI. This lot was administered in the Philippines to populations with an unknown age. Upon review, in 2640 cases, it was reported that the orange cap formulation was administered to adult individuals; while in 3348 cases, it was reported that the orange cap formulation was administered to individuals above 12 years old. No clinical AEs were co-reported.

Overall, there were no safety issues related to quality identified during product complaint investigations.

Surveillance for any potential product quality issues includes review of quarterly AE/PC reports and monthly SAE/PC reports, and review of weekly AE-batch/lot trending reports. In support to this process as needed, a review of AE data related to respective PCs may be requested to support trend analysis and notifications. Alerts in the AE/PC reports are reviewed and closed or escalated based on clinical judgement and product knowledge. Any potential signals indicating a potential relationship between a safety issue and a particular batch lot, and that was not already evaluated as part of other signal activities, would undergo evaluation and escalation as per standard procedures.
Conclusion

Based on the review of the cases with the most frequently reported lot numbers, no new safety issues were identified.

6.3.1.2. General Overview of the Safety Database - Unlocked Cases

A total of 2441 (0.5%) unlocked\textsuperscript{46} case reports (4 from CT and 2437 from PM) containing 8399 events fulfilled criteria for inclusion in this PSUR, compared to 139,698 (21.2%) case reports retrieved in the PSUR #2. Table 27 displays demographic information of the unlocked cases at the end of the reporting interval.

Table 27. Demographic Information - Unlocked Cases at the End of the Reporting Interval

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>All No. of Cases (%)</th>
<th>CT No. of Cases (%)</th>
<th>PM No. of Cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N=2441</td>
<td>N=4</td>
<td>N=2437</td>
</tr>
<tr>
<td>No. of Cases</td>
<td>2441</td>
<td>4</td>
<td>2437</td>
</tr>
<tr>
<td>MC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>1203 (49.3)</td>
<td>4 (100)</td>
<td>1199 (49.2)</td>
</tr>
<tr>
<td>No</td>
<td>1238 (50.7)</td>
<td>0</td>
<td>1238 (50.8)</td>
</tr>
<tr>
<td>Country of occurrence (≥2% of all cases)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Austria</td>
<td>388 (15.9)</td>
<td>0</td>
<td>388 (15.9)</td>
</tr>
<tr>
<td>Germany</td>
<td>334 (13.7)</td>
<td>0</td>
<td>334 (13.7)</td>
</tr>
<tr>
<td>France</td>
<td>284 (11.6)</td>
<td>0</td>
<td>284 (11.7)</td>
</tr>
<tr>
<td>US</td>
<td>277 (11.3)</td>
<td>4 (100)</td>
<td>273 (11.2)</td>
</tr>
<tr>
<td>Italy</td>
<td>175 (7.2)</td>
<td>0</td>
<td>175 (7.2)</td>
</tr>
<tr>
<td>Sweden</td>
<td>127 (5.2)</td>
<td>0</td>
<td>127 (5.2)</td>
</tr>
<tr>
<td>Romania</td>
<td>95 (3.9)</td>
<td>0</td>
<td>95 (3.9)</td>
</tr>
<tr>
<td>UK</td>
<td>87 (3.6)</td>
<td>0</td>
<td>87 (3.6)</td>
</tr>
<tr>
<td>Philippines</td>
<td>83 (3.4)</td>
<td>0</td>
<td>83 (3.4)</td>
</tr>
<tr>
<td>Japan</td>
<td>78 (3.2)</td>
<td>0</td>
<td>78 (3.2)</td>
</tr>
<tr>
<td>Greece</td>
<td>65 (2.7)</td>
<td>0</td>
<td>65 (2.7)</td>
</tr>
<tr>
<td>Netherlands</td>
<td>62 (2.5)</td>
<td>0</td>
<td>62 (2.5)</td>
</tr>
<tr>
<td>Norway</td>
<td>51 (2.1)</td>
<td>0</td>
<td>51 (2.1)</td>
</tr>
<tr>
<td>Other countries</td>
<td>335 (13.7)</td>
<td>0</td>
<td>335 (13.7)</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1430 (58.6)</td>
<td>1 (25)</td>
<td>1429 (58.6)</td>
</tr>
<tr>
<td>Male</td>
<td>796 (32.6)</td>
<td>3 (75)</td>
<td>793 (32.5)</td>
</tr>
<tr>
<td>Unknown/No Data</td>
<td>215 (8.8)</td>
<td>0</td>
<td>215 (8.9)</td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>2160</td>
<td>4</td>
<td>2156</td>
</tr>
<tr>
<td>Min-Max</td>
<td>0.04-99</td>
<td>6-63</td>
<td>0.04-99</td>
</tr>
<tr>
<td>Mean\textsuperscript{b}</td>
<td>48.9</td>
<td>45.3</td>
<td>48.9</td>
</tr>
<tr>
<td>Median\textsuperscript{b}</td>
<td>48.5</td>
<td>56</td>
<td>48</td>
</tr>
<tr>
<td>Age Range</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 17</td>
<td>143 (5.9)</td>
<td>1 (25)</td>
<td>142 (5.8)</td>
</tr>
<tr>
<td>18-30</td>
<td>262 (10.7)</td>
<td>0</td>
<td>262 (10.8)</td>
</tr>
<tr>
<td>31-50</td>
<td>749 (30.7)</td>
<td>1 (25)</td>
<td>748 (30.7)</td>
</tr>
<tr>
<td>51-64</td>
<td>516 (21.1)</td>
<td>2 (50)</td>
<td>514 (21.1)</td>
</tr>
<tr>
<td>65-74</td>
<td>254 (10.4)</td>
<td>0</td>
<td>254 (10.4)</td>
</tr>
</tbody>
</table>

\textsuperscript{46} Unlocked cases are those cases either in the Drug Safety Unit, Primary Review or the Medical Review workflows that are not yet in the Distribution workflow, which locks the cases, and the system automatically runs reporting rules, schedules and subsequently generates expedited reports as appropriate.
Table 27. Demographic Information - Unlocked Cases at the End of the Reporting Interval

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>All No. of Cases (%) N=2441</th>
<th>CT No. of Cases (%) N=4</th>
<th>PM No. of Cases (%) N=2437</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt; 75</td>
<td>247 (10.1)</td>
<td>0</td>
<td>247 (10.1)</td>
</tr>
<tr>
<td>Unknown</td>
<td>269 (11.0)</td>
<td>0</td>
<td>269 (11.0)</td>
</tr>
<tr>
<td>Not applicable</td>
<td>1 (0.0)</td>
<td>0</td>
<td>1 (0.0)</td>
</tr>
<tr>
<td>Case Seriousness</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serious</td>
<td>1541 (63.1)</td>
<td>4 (100)</td>
<td>1537 (63.1)</td>
</tr>
<tr>
<td>Non-serious</td>
<td>900 (36.9)</td>
<td>0</td>
<td>900 (36.9)</td>
</tr>
<tr>
<td>Case Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>32 (1.3)</td>
<td>1 (25)</td>
<td>31 (1.3)</td>
</tr>
<tr>
<td>Not resolved</td>
<td>747 (30.6)</td>
<td>1 (25)</td>
<td>746 (30.6)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>642 (26.3)</td>
<td>2 (50)</td>
<td>640 (26.3)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>47 (1.9)</td>
<td>0</td>
<td>47 (1.9)</td>
</tr>
<tr>
<td>Unknown</td>
<td>973 (39.9)</td>
<td>0</td>
<td>973 (39.9)</td>
</tr>
<tr>
<td>Presence of comorbidities</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>366 (15.0)</td>
<td>1 (25)</td>
<td>365 (15.0)</td>
</tr>
<tr>
<td>No</td>
<td>2075 (85.0)</td>
<td>3 (75)</td>
<td>2072 (85.0)</td>
</tr>
</tbody>
</table>

a. The sum of percentages may not exactly match 100% due to rounding in calculations.
b. Values referred to all cases with an age in years not null.

6.3.1.2.1. General Overview of the Safety Database Unlocked Cases - Clinical Trials Data

The events reported in clinical trial cases that were unlocked at the end of the reporting interval were coded to the PTs Appendicitis, Foreign body in gastrointestinal tract, Overdose, and Rheumatoid arthritis (1 each).

6.3.1.2.2. General Overview of the Safety Database Unlocked Cases - Post-Authorisation Data

The overall safety evaluation includes a review of the reported events by SOC and by PT for events reported in $\geq 2\%$ of unlocked cases at the end of the reporting interval.

Table 28. Post-Authorisation Data: Events Reported in $\geq 2\%$ of Unlocked Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>MedDRA PTs</th>
<th>N=2437 n (AERP, $^a$ %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood and lymphatic system disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lymphadenopathy$^b$</td>
<td></td>
<td>61 (2.5)</td>
</tr>
<tr>
<td>Cardiac</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palpitations$^c$</td>
<td></td>
<td>55 (2.3)</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nausea$^b$</td>
<td></td>
<td>95 (3.9)</td>
</tr>
<tr>
<td>Diarrhoea$^b$</td>
<td></td>
<td>54 (2.2)</td>
</tr>
<tr>
<td>Vomiting$^b$</td>
<td></td>
<td>50 (2.1)</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drug ineffective$^d$</td>
<td></td>
<td>343 (14.1)</td>
</tr>
<tr>
<td>Vaccination failure$^d$</td>
<td></td>
<td>265 (10.9)</td>
</tr>
<tr>
<td>Fatigue$^b$</td>
<td></td>
<td>228 (9.4)</td>
</tr>
<tr>
<td>Pyrexia$^b$</td>
<td></td>
<td>213 (8.7)</td>
</tr>
</tbody>
</table>
Table 28. Post-Authorisation Data: Events Reported in ≥2% of Unlocked Cases

<table>
<thead>
<tr>
<th>MedDRA SOC</th>
<th>N=2437 n (AERP, %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthenia b</td>
<td>110 (4.5)</td>
</tr>
<tr>
<td>Vaccination site pain b</td>
<td>106 (4.4)</td>
</tr>
<tr>
<td>Pain c</td>
<td>97 (4.0)</td>
</tr>
<tr>
<td>Chills b</td>
<td>96 (3.9)</td>
</tr>
<tr>
<td>Malaise b</td>
<td>95 (3.9)</td>
</tr>
<tr>
<td>Chest pain c</td>
<td>79 (3.2)</td>
</tr>
</tbody>
</table>

Infections and infestations
COVID-19d

Injury, poisoning and procedural complications
Inappropriate schedule of product administration a
Off label use e
Poor quality product administered e
Expired product administered e
Product administration error e

Musculoskeletal and connective tissue disorders
Pain in extremity b
Myalgia b
Arthralgia b

Nervous system disorders
Headache b
Dizziness e
Paraesthesia e
Hypoesthesia e
Syncope e

Product issues
Product temperature excursion issue e

Reproductive system and breast disorders
Heavy menstrual bleeding c

Respiratory, thoracic and mediastinal disorders
Dyspnoea c

Surgical and medical procedures
Interchange of vaccine products f
Immunisation g

Total number of events
4304

a. Reporting proportion (% of total cases) calculated as n/N at the end of the current reporting period.
N: Number of Cases; n: Number of events.
b. Listed or consistent with listed AEs in current RSI.
c. Unlisted in the current RSI.
d. Listed per case processing convention.
e. Per case processing convention, the PT term follows the most conservative listedness assessment of the associated AEs. If the term is reported alone or without associated AEs, the term is assessed as listed.
f. Of note, a majority of the cases reporting interchange of vaccine products (75/78; 96.2%) co-reported Off label use. These cases described use of vaccines from different manufacturers. Per case processing convention, the PT term follows the most conservative listedness assessment of the associated AEs. If the term is reported alone or without associated AEs, the term is assessed as listed.
g. Of note, a majority of the cases reporting Immunisation (40/49; 81.6%) co-reported Off label use. These cases report a booster being administered and it is captured as an event when it is off-label per the local label. Per case processing convention, the PT terms follows the most conservative listedness assessment of the associated AEs. If the term is reported alone or without associated AEs, the term is assessed as listed.
Conclusion

The data contained in the unlocked cases are consistent with the overall dataset.

7. SUMMARIES OF SIGNIFICANT SAFETY FINDINGS FROM CLINICAL TRIALS DURING THE REPORTING INTERVAL

Appendix 4.2 provides a list of ongoing interventional safety studies. No interventional safety studies were completed during the reporting interval.

7.1. Completed Clinical Trials

Safety Trials

During the reporting period, no interventional safety studies were completed with a final CSR.

Other Trials that reported new significant efficacy information

During the reporting interval, there was a completed clinical trial (C4591017) with a final CSR (available upon request). No clinically important new information has emerged from this clinical trial; overall conclusions for the study are provided below.

Table 29. Summary of Results from Clinical Trial Completed During the Reporting Period

<table>
<thead>
<tr>
<th>Protocol ID</th>
<th>Protocol Title</th>
<th>Conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4591017</td>
<td>A phase 3, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of multiple production lots and dose levels of the vaccine candidate BNT162b2 against COVID-19 in healthy participants 12 through 50 years of age and the safety, tolerability, and immunogenicity of BNT162b2 RNA-based COVID-19 vaccine candidates as a booster dose in healthy participants 18 through 50 years of age.</td>
<td>The 3 US lots in the primary study met the 1.5-fold equivalence criteria for all 3 between-lot comparisons based on full-length S-binding IgG levels and were considered similar. Safety profiles across all vaccine groups were similar with no safety issues identified in both the primary and booster study. Vaccines in all arms of the study were well-tolerated in both the primary and booster study. The safety profile of BNT162b2 was consistent with previous studies.</td>
</tr>
</tbody>
</table>
7.2. Ongoing Clinical Trials

During the reporting period, there were 14 ongoing\textsuperscript{47} sponsor-initiated clinical trials.

**Safety Trials** (see Appendix 4.2 for a list of ongoing interventional safety studies):

**PASS:**

- \textit{C4591015 [A phase 2/3, placebo-controlled, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate (BNT162b2) against COVID-19 in healthy pregnant women 18 years of age and older]} is an ongoing PASS. No clinically important new information has emerged from this ongoing PASS.

- \textit{C4591024 [A phase 2b, open-label study to evaluate the safety, tolerability, and immunogenicity of vaccine candidate BNT162b2 in immunocompromised participants \textgreater{}2 years of age]} is an ongoing PASS. No clinically important new information has emerged from this ongoing PASS.

Other trials where the primary aim of the trial was to identify, characterise or quantify a safety hazard or confirm the safety profile of the medicinal product: None.

**Other Trials** that reported new significant efficacy information

There were 8 ongoing clinical trials:

- \textit{C4591001\textsuperscript{48}, A phase 1/2/3, placebo-controlled, randomized, observer-blind, dose-finding study to evaluate the safety, tolerability, immunogenicity, and efficacy of SARS-CoV-2 RNA vaccine candidates against COVID-19 in healthy individuals.}

- \textit{C4591007, A phase 1, open-label dose-finding study to evaluate safety, tolerability, and immunogenicity and phase 2/3 placebo-controlled, observer-blinded safety, tolerability, and immunogenicity study of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy children \textless{}12 years of age.}

- \textit{C4591031,\textsuperscript{49} A phase 3 master protocol to evaluate additional dose(s) of BNT162b2 in healthy individuals previously vaccinated with BNT162b2.}

\textsuperscript{47} Includes ongoing studies as well as studies in which participant enrollment and follow-up have been completed, but the analysis and CSR are in-progress.

\textsuperscript{48} Two interim CSRs were issued for Study C4591001 during the reporting interval [BNT162b2 (30 \mu g) Booster (Dose 3) – Phase 1 (4-Month Update) and Phase 3 (6-Month Update) v. 1.0 dated 19 May 2022 and BNT162b2SA VOC Booster Subset v. 1.0 dated 20 May 2022].

\textsuperscript{49} Two interim CSRs were issued for Substudy A of Study C4591031 (v. 1.0 on 25 April 2022 and v. 2.0 on 07 June 2022) in the reporting period.

- BNT162-04, A multi-site, phase II, 2-part, dose-escalation trial investigating the safety and immunogenicity of four prophylactic SARS-CoV-2 RNA vaccines against COVID-19 using different dosing regimens in healthy and immunocompromised adults.


- BNT162-14, A Phase II, open-label rollover trial to evaluate the safety and immunogenicity of one or two boosting doses of Comirnaty or one dose of BNT162b2 at BNT162-01 trial subjects, or two boosting doses of Comirnaty in BNT162-04 trial subjects.

No clinically important new safety information has emerged from ongoing clinical trials.

Remaining Trials

There were 4 ongoing clinical trials:

- C4591005, A phase 1/2, placebo-controlled, randomized, and observer-blind study to evaluate the safety, tolerability, and immunogenicity of a SARS-CoV-2 RNA vaccine candidate against COVID-19 in healthy Japanese adults.

- C4591020, A phase 3, randomized, observer-blind study to evaluate the safety, tolerability, and immunogenicity of multiple formulations of the vaccine candidate BNT162b2 against COVID-19 in healthy adults 18 through 55 years of age.

- C4591030, A phase 3, randomized, observer-blind trial to evaluate the safety and immunogenicity of BNT162b2 when co-administered with seasonal inactivated influenza vaccine (SIIV) in adults 18 through 64 years of age.

- BNT162-17, A Phase II trial to evaluate the safety and immunogenicity of SARS-CoV-2, monovalent and multivalent RNA-based vaccines in healthy subjects.

---

50 Last subject last visit occurred during the reporting interval for the following studies: BNT162-01 (13 Apr 2022); BNT162-04 (07 Feb 2022); BNT162-06 (09 Jan 2022).

51 This study is conducted by Shanghai Fosun Pharmaceutical Development, Inc. and sponsored by BioNTech SE.
No clinically important new safety information has emerged from these ongoing clinical trials.

7.3. Long-term Follow-up
There is no new significant safety information with regards to long-term follow-up of clinical trial participants for this reporting period.

7.4. Other Therapeutic Use of Medicinal Product
BNT162b2 was also utilised in another Pfizer-sponsored clinical development program (B747). The study B7471026 “A phase 3, randomized, double blind trial to describe the safety and immunogenicity of 20 valent pneumococcal conjugate vaccine when coadministered with a booster dose of BNT162b2 in adults 65 years of age and older” was completed during the reporting period.

There was no new clinically important safety information identified for this reporting period. The CSR for this trial is available upon request.

7.5. New Safety Data Related to Fixed Combination Therapies
BNT162b2 is not used in fixed or multi-drug combination with other compounds.

8. FINDINGS FROM NON-INTERVENTIONAL STUDIES
During the reporting period, there were 11 ongoing sponsor-initiated non-interventional studies and one non-interventional study (C4591035) was completed.

8.1. Completed Non-Interventional Studies
Safety studies

Neither PASS nor other studies where the primary aim of the trial was to identify, characterise or quantify a safety hazard or confirm the safety profile of the medicinal product were completed during the reporting period.

Other studies

During the reporting period, the study C4591035 was completed. No new safety information emerged from this non-interventional study; the summary of results from this study is provided in Table 30.
Table 30. Summary of Results from Completed NIS During the Reporting Period

<table>
<thead>
<tr>
<th>Protocol ID</th>
<th>Protocol Title</th>
<th>Conclusions*</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4591035</td>
<td>Coronavirus Disease 2019 (COVID-19) Vaccination and Breakthrough Infections Among Persons with Immunocompromising Conditions in the US.</td>
<td>This study contained the largest population of fully vaccinated IC individuals to date in whom COVID-19 vaccine breakthrough infections have been evaluated. The study findings show that breakthrough infections are generally rare but are more common and severe in people with certain IC conditions. The conclusions support the FDA authorisation and CDC recommendations to offer a 3rd vaccine dose to increase protection among IC individuals and the need for vigilant efforts to maximize vaccine uptake among the IC, especially in the context of waning duration of protection and emerging SARS-CoV-2 variants. Moreover, the findings from this study suggest that breakthrough infections can occur regardless of active treatment status in the IC and that there may be additional vulnerable IC groups that could benefit from increased protection. This study advances the understanding of post-vaccination outcomes across multiple IC condition groups in a real-world setting and may help healthcare providers in the decision-making process when vaccinating and treating patients at high-risk for COVID-19.</td>
</tr>
</tbody>
</table>

---


On 04 May 2022 the CSR was finalised with confirmation that this non-interventional study manuscript, which is final in content and has been printed from its definitive source, is a complete and accurate representation of the data and statistical analyses from study C4591035.
8.2. Ongoing Non-Interventional Studies

Safety Studies (see Appendix 4.4 for a list of ongoing non-interventional safety studies and their protocol titles):

PASS\textsuperscript{52}: Non-interventional studies C4591006\textsuperscript{53}, C4591009\textsuperscript{54}, C4591010\textsuperscript{55}, C4591012\textsuperscript{54}, C4591021\textsuperscript{54} and C4591022\textsuperscript{54} are PASS. No clinically important information has emerged from PASS.

Other studies where the primary aim of the trial was to identify, characterise or quantify a safety hazard or confirm the safety profile of the medicinal product: None.

Other Studies

There were 5 ongoing non-interventional studies:

- \textit{C4591006\textsuperscript{56}} General Investigation of COMIRNATY intramuscular injection (follow-up study for subjects [healthcare professionals] who are vaccinated at an early post-approval stage).
- \textit{C4591014,\textsuperscript{57} Pfizer-BioNTech COVID-19 BNT162b2 Vaccine Effectiveness Study - Kaiser Permanente Southern California}.
- \textit{C4591019, Special investigation of COMIRNATY Intramuscular Injection (Investigation of Patients with Underlying Disease Considered to be at High Risk of Aggravation of COVID-19)}
- \textit{C4591025,\textsuperscript{58} A prospective, single-arm, open-label, non-interventional, multicenter to assess the safety of BNT162b2 in domestic post-marketing surveillance}.
- \textit{C4591034, Patient-Reported Health-Related Quality of Life Associated With COVID-19: A Prospective Survey Study on Symptomatic Adults Confirmed With RT-PCR From Outpatient Settings in the US}.

During the reporting period, no new significant - safety information non-interventional studies was reported.

\begin{footnotesize}
\begin{itemize}
\item \textsuperscript{52} During the reporting period, interim CSRcs were issued for the studies C4591010 (17 February 2022), C4591021 (23 March 2022) and C4591022 (25 January 2022).
\item \textsuperscript{53} Study C4591008 is a voluntary study; it is included in the US-PVP as post-authorisation safety study addressing the important potential risk of VAED/VAERD.
\item \textsuperscript{54} Studies C4591009, C4591012, C4591021 and C4591022 are commitments to the US FDA and are Category 3 commitments in the EU-RMP v.5.0.
\item \textsuperscript{55} Study C4591010 is Category 3 commitment in the EU-RMP v. 5.0.
\item \textsuperscript{56} Studies C4591006 and C4591019 are commitments to the Japanese regulatory.
\item \textsuperscript{57} PAM-MEA-013.
\item \textsuperscript{58} Study C4591025 is a committed study, which was requested by the Ministry of Food and Drug Safety in Korea.
\end{itemize}
\end{footnotesize}
9. INFORMATION FROM OTHER CLINICAL TRIALS AND SOURCES

9.1. Other Clinical Trials

During the reporting interval, there were 6 relevant cases that originated from non-Pfizer and non-BNT clinical trials. In 3 of these cases, BNT162b2 was a study drug, while in the other 3 cases BNT162b2 was co-administered with the study drug.

Four (4) cases originated from the following non-Pfizer and non-BNT trials:


- DE-EORTC-1745-186-1 - A phase II study of adjuvant palbociclib as an alternative to chemotherapy in elderly patients with high-risk ER+/HER2- early breast cancer (appalaches) (SAE: Thrombophlebitis).

- RBR-9NN3SCW - Phase 4, randomized, controlled, single-blind study to assess the immunogenicity and safety of a third dose of heterologous booster with recombinant COVID-19 vaccine (Astra Zeneca/Fiocruz), COVID-19 mRNA vaccine (Pfizer/Wyeth) or vaccine recombinant COVID-19 (Janssen) in subjects previously vaccinated against COVID-19 with two doses of Sinovac/Butantan compared to a third homologous booster dose of adsorbed inactivated COVID-19 vaccine (Sinovac/Butantan) in adults (SAEs: Deep vein thrombosis, Pulmonary embolism).

- H3B-6545-G000-102 - An open-label multicenter phase 1b study of H3B-6545 in combination with palbociclib in women with advanced or metastatic estrogen receptor-positive HER2-negative breast cancer (SAE: Myocarditis). This case referred to a 72-year-old female participant with breast cancer and with a history of left ventricular failure; she received the 4th dose of BNT162b2 7 days before the beginning of treatment with palbociclib and 28 days before the beginning of the treatment with H3B-6545. Nine (9) days after the vaccination, she had palpitations and 5 days later she had abnormal electrocardiogram (inverted T waves in V5/V6, which later flattened, and raised troponin); she was hospitalised with focal myocarditis. Palbociclib was interrupted on the same day. Five (5) days after the hospital admission, she recovered from myocarditis.

The SAEs reported in these 4 cases were assessed as related to the BNT162b2 by the investigators and the MAH agreed except for the case reporting Deep vein thrombosis and Pulmonary embolism, where it was considered that there was not a reasonable possibility that the events were related to vaccine administration, based on the absence of a plausible pathophysiological mechanism.

Two (2) cases originated from the following non-Pfizer, non-BNT trials:

- 3101-304-002 - Phase 3, multicentre, randomized, double-blind, placebo-controlled, parallel-group Study to evaluate the efficacy, safety, and tolerability of oral atogepant
for the prophylaxis of migraine in participants with episodic migraine who have previously failed 2 to 4 classes of oral prophylactic treatments (ELEVATE) [SAEs: Immunisation, Overdose (0.5 ml of Pfizer-BioNTech COVID-19 vaccine), Ventricular tachycardia].

- 21-0012 - A phase 1/2 study of delayed heterologous SARs-CoV-2 vaccine dosing (boost) after receipt of EUA vaccines (SAEs: Condition aggravated, Endometrial thickening).

The investigator’s assessment for Ventricular tachycardia was not provided; Endometrial thickening was considered unrelated to BNT162b2 by the investigator; in both cases the MAH considered the SAEs as unrelated to the vaccine.

During this reporting period, there was no new significant safety information reported from other non-Pfizer, non-BNT sponsored clinical trials/studies.

9.2. Medication Errors

As part of the AR of the 11th SMSR (Procedure EMEA/H/C/005735/MEA/002.10), the following commitment is included: “The MAH should report on handling and dosing errors as a result of the different Comirnaty formulations on the market.”

Response

Please refer to Appendix 6A for details.

Analysis of the Medication Errors

Cases potentially indicative of medication errors\(^\text{59}\) that occurred in the reporting period are summarised below.

\(^{59}\) Search criteria: MedDRA (version 25.0): HLTs (All paths): Accidental exposures to product; Product administration errors and issues; Product confusion errors and issues; Product dispensing errors and issues; Product label issues; Product monitoring errors and issues; Product preparation errors and issues; Product prescribing errors and issues; Product selection errors and issues; Product storage errors and issues in the product use system; Product transcribing errors and communication issues, OR PTs: Accidental poisoning; Circumstance or information capable of leading to device use error; Circumstance or information capable of leading to medication error; Contraindicated device used; Contraindication to vaccination; Deprescribing error; Device use error; Dose calculation error; Drug titration error; Expired device used; Exposure via direct contact; Exposure via eye contact; Exposure via mucosa; Exposure via skin contact; Failure of child resistant product closure; Inadequate aseptic technique in use of product; Incorrect disposal of product; Interpreted medication error; Interpreted product prescribing error; Medication error; Multiple use of single-use product; Product advertising issue; Product distribution issue; Product prescribing error; Product prescribing issue; Product substitution error; Product temperature excursion issue; Product use in unapproved therapeutic environment; Radiation underdose; Underdose; Unintentional medical device removal; Unintentional use for unapproved indication; Vaccination error; Wrong device used; Wrong dosage form; Wrong dosage formulation; Wrong dose; Wrong drug; Wrong patient; Wrong product procured; Wrong product stored; Wrong rate; Wrong route; Wrong schedule; Wrong strength; Wrong technique in device usage process; Wrong technique in product usage
Clinical Trial Data

During the reporting period, there were 2 serious cases (0.3% of 668 cases, the total CT dataset) indicative of medication errors (PTs: Accidental overdose and Inappropriate schedule of product administration). In the first case, the accidental overdose referred to paracetamol and not to BNT162b2 and in the remaining case reporting inappropriate schedule of product administration, the investigator assessed the event as not related to BNT162b2. There was 1 serious case retrieved during the reporting period of the PSUR #2.

Post-Authorisation Data

From the global safety database, 68,025 cases (13.4% of 507,683 cases, the total PM dataset) potentially indicative of medication errors were retrieved during the reporting period.

Of the 68,025 cases, 1261 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

- Off-label use or intentional use rather than medication error was reported in 515 cases;
- Reporting information was no longer consistent to meet medication error criteria in 11 cases;
- Cases consisted of questions or requests for information about the scheduling of the 2 doses of BNT162b2 or the second dose (not administered yet at the time of reporting) or scheduling outside the prescribed dosing window were reported in 700 cases;
- Cases consisted of booster dose scheduling outside the prescribing window were reported in 6 cases;
- Medical inquiries only were reported in 4 cases;
- The subject intentionally refused to be vaccinated or was not able to receive the scheduled BNT162b2 in 7 cases;
- In 7 cases, the reported errors were not due to BNT162b2;
- In 6 cases, subjects were exposed to the vaccine during breastfeeding;
- An unspecified number of subjects were described in 5 cases.

The potentially relevant medication error cases during the reporting period were 66,764 (13.1%) reporting 87,307 events, compared to 33,834 relevant cases (5.1%) analysed in the PSUR #2.

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60 Among the most commonly reported countries (≥2% of cases), significant increase in proportion of cases observed from Austria (21.5 vs 6.9%), Philippines (9.3% vs 0.07%), Sweden (11.5 vs 4.5%) and Australia (2.3% vs 0.4%) when compared to cases reported in PSUR #2. Most of the cases from these 4 countries reported scheduling, formulation or storage errors. In comparison with PSUR #2, there was a significant increase in number of cases identified with new Tris/Sucrose formulation in the current reporting period: Orange cap (755 cases vs 9426 cases)/Grey cap (0 vs 2750 cases), respectively.
The 66,764 relevant medication error cases originated mostly (≥2% of cases) from the following countries: Austria (14,339 cases), US (9592 cases), Germany (8074 cases), Sweden (7666 cases), Philippines (6182 cases), UK (4531 cases), France (2923 cases), Netherlands (2586 cases), Japan (1630 cases), Australia (1506 cases), and Canada (1425 cases).

The most frequently reported (≥2% of cases) medication error PTs were: Inappropriate schedule of product administration (34,486 events), Poor quality product administered (17,837 events), Product administered to patient of inappropriate age (7952 events), Product administration error (6946 events), Product storage error (5882 events), Product temperature excursion issue (4626 events), Expired product administered (2060 events), and Incorrect route of product administration (1889 events).

In some instances, clusters of medication errors were reported. During the reporting interval, 5 different types of medication error cases (>1000 occurrences) were identified coded to the PTs Product storage error, Poor quality product administered, and Product administered to patient of inappropriate age.

All cases demonstrated no-harm and had no co-reported events:

- In 3348 cases, Tris/Sucrose Orange cap presentation of BNT162b2 was given to adolescent patients.
- In 3011 cases, BNT162b2 was thawed and kept in the refrigerator longer than the recommended period.
- In 2640 cases, Tris/Sucrose Orange cap presentation of BNT162b2 was given to adult patients.
- In 1169 cases, BNT162b2 was stored in the cold storage (15 to 25 degrees Celsius) 20 days longer than the recommended 2 weeks period prior to use and the vaccine was administered.
- In 1169 cases, BNT162b2 was stored in the cold storage (15 to 25 degrees Celsius) 13 days longer than the recommended 2 weeks period prior to use and the vaccine was administered.

Medication Errors Analysis

Among the relevant medication error cases (66,764 cases61), the following scenarios, were described:

- Medication errors associated with harm [i.e., resulting in adverse reaction(s)] according to EMA guidance “Good practice guide on recording, coding, reporting and assessment of medication errors” (EMA/762563/2014) were reported in 1326 cases (2% of relevant

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61 Relevant medication error cases 66,764 (1326 harm + 65,350 no-harm + 87 potential + 1 intercepted errors).
medication error cases) compared to 879 cases (2.6% of relevant medication error cases) analysed in the PSUR #2.

Of note, some cases involved more than one medication error.

**Medication errors associated with harm (1398 medication error events in 1326 cases)**

Of the 1326 cases, 472 were medically confirmed, and 854 were non-medically confirmed. Cases were mostly (>50 occurrences) reported from Germany (390 cases), Poland (206 cases), Iraq (191 cases), France (91 cases), UK (73 cases), Italy (69 cases), US (63 cases).

There were 899 females and 392 male subjects, whereas the gender was not specified for 35 subjects. When provided (n = 1250), the age ranged from 1.0 month to 98.0 years with a mean age of 40.5 years and a median of 40.0 years.

Of the 1398 medication error events, 78 were serious and 1320 were non-serious. The relevant event outcome was reported as fatal (2), resolved/resolving (29), not resolved (20), and unknown (1347).

There were 2 fatal medication error events reported in 2 cases and they were coded to the PT Incorrect dose administered Section 16.3.4.1 *Death*.

- **First case**: An 84-year-old male subject received BNT162b2 as dose 4 (booster) for COVID-19 immunisation. The subject’s relevant medical history included: artificial cardiac pacemaker wearer (ongoing) and prostate problems. Vaccination history included: 3 doses of BNT162b2 and the subject experienced several adverse events after each vaccination including intestinal haemorrhage, headache, vomiting/urge to vomit, malaise, increased bleeding under the skin, and petechiae. It was reported that the subject received an incorrect fourth dose of BNT162b2 (dose unspecified) and experienced a fatal event of intestinal haemorrhage. The other events reported were haemorrhage subcutaneous, petechiae, headache, vomiting, malaise, and contusion. The cause of death was unknown, and it was not reported if an autopsy was performed.

- **Second case**: An 89-year-old female subject received BNT162b2 second booster dose by mistake (PT Incorrect dose administered) and died due to cardiac failure and acute pulmonary oedema 2 days after the time of vaccination. It was reported that the subject had multiple pre-existing conditions including cardiac diseases, follicular lymphoma, pulmonary tuberculosis, lumbar spinal stenosis, and breast cancer. It was unknown if an autopsy was performed. No further information was available.

**Serious medication errors**

In 70 cases (involving 78 medication error events; 5.3% of 1326 cases; this includes 2 fatal cases [described above]), serious medication errors potentially contributed to the occurrence of SAEs when compared to 59 serious cases (involving 69 medication error events; 6.7% of 879 cases) analysed in the PSUR #2.
Cases (≥2 occurrences) originated from UK (43 cases), Germany (7 cases), France (4 cases), Poland and Belgium (2 cases each).

The serious events (≥3 occurrences) indicative of medication errors were Medication error (19 events), Product administered at inappropriate site (15 events), Incorrect route of product administration (12 events), Incorrect dose administered (10 events), Product administration error (9 events), Wrong technique in product usage process (5 events), and Expired product administered (3 events).

The most frequently (≥6 occurrences) co-reported clinical events were Headache (19 events), Pain in extremity (16 events), Pain (11 events), Arthralgia, Nausea (10 events each), Fatigue (8 events), Myalgia, Paraesthesia, Pyrexia (7 events each), Chills, Dizziness, Malaise, Periarthritis, Shoulder injury related to vaccine administration, and Vaccination site pain (6 events each).

Upon review of medication error serious events:

*Vaccine administration errors (56 events, 50 cases)*

Events described in these cases were: errors of vaccination at the wrong anatomical site, errors in administration technique, errors in route of administration, errors in vaccine dosage, and other administration errors.

- **Errors of vaccination at the wrong anatomical site (18 events)**
  - The vaccine was administered in the shoulder, arm site other than deltid (8 events each), leg, and bursa (1 event each).

- **Errors in vaccine dosage administered (11 events)**
  - Errors included administration of fourth/second booster dose (3 events), additional dose (1 event), and incorrect unspecified dose administered (7 events).

- **Errors in the route of administration (10 events)**
  - The route of administration for the vaccine was subcutaneous (5 events), intravenous (3 events), intrameningeal, and intradermal (1 event each).

- **Administration technique errors (8 events)**
  - Errors included injury to nerve/blood vessel (3 events), moving the needle around in the arm, injected with force, poor injection technique (1 event each), unspecified technique errors (2 events).

- **Other vaccine administration errors (9 events)**
  - These events reported administration of expired vaccines (3 events), administration of wrong vaccine (1 event), unspecified administration errors (5 events).
Vaccine preparation errors (2 events, 2 cases)

- Errors included dilution before use was not performed or improper dilution (1 event each).

Other errors (20 events, 18 cases)

- Errors included unspecified medication error/unspecified vaccination error (20 events).

Non-serious medication errors

In 1256 cases (involving 1320 medication error events), non-serious medication errors potentially contributed to the occurrence of non-serious AEs, when compared to 820 cases (involving 886 non-serious medication error events) analysed in the PSUR #2. Most of the cases (≥60 occurrences) originated from Germany (383 cases), Poland (204 cases), Iraq (190 cases), France (87 cases), Italy (69 cases), and US (62 cases).

The most frequently (≥105 occurrences) co-reported clinical events were Pyrexia (323 events), Headache (265 events), Vaccination site pain (245 events), Fatigue (178 events), Pain in extremity (150 events), Myalgia (126 events), Asthenia (114 events), and Chills (105 events).

- Vaccine administration errors\(^{62}\) (1224 events): Events mainly described errors in route of administration, errors in the volume or dosage of the vaccine administered, errors in administration of vaccine dose not adequately prepared, errors of vaccination at the wrong anatomical site, errors in the administration of incorrect product, and other administration errors.

- Vaccine preparation errors\(^{63}\) (19 events): Events mainly described errors during dilution and other preparation errors.

- Vaccine scheduling errors\(^{64}\) (1 event): Event described error in scheduling of second dose administration

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\(^{62}\) PTs Incorrect route of product administration, Incorrect dose administered, Product administered at inappropriate site, Product administered to patient of inappropriate age, Underdose, Expired product administered, Poor quality product administered, Product administration error, Wrong product administered, Accidental overdose, Accidental exposure to product, Extra dose administered, Product administration interrupted.

\(^{63}\) PTs Product preparation error, Product preparation issue, Underdose, Accidental overdose (if associated to other PT indicative of erroneous preparation).

\(^{64}\) PTs Inappropriate schedule of product administration.
Other medication errors (76 events): Events mainly described temperature excursion, vaccine administration technique, lot number, vaccination error, and other errors.

The summary of analysis of medication errors pertaining to the new BNT162b2 formulations (Tris/Sucrose presentation) is presented below.

9.2.1. Errors pertaining to the new formulation of BNT162b2 – Paediatric Tris/Sucrose Orange Cap presentation (dilute before use) 10 μg/dose for 5 to <12 years of age

Search criteria used for selecting the below cases for discussion: Paediatric subjects 5 to <12 years of age received Tris/Sucrose Orange cap (10 mcg/dose) presentation; Tris/Sucrose Orange cap formulation was used or administered to age groups other than 5 to <12 years instead of PBS/Sucrose presentation.

There were 9426 cases reporting 12,051 events indicative of medication errors per the medication error MedDRA search strategy related to Tris/Sucrose Orange cap presentation (paediatric formulation). The majority of these cases (>30 cases) describing medication errors were from Philippines (5994 cases), US (2640 cases), Australia (276 cases), Germany (109 cases), Canada (94 cases), Japan (68 cases), Spain (61 cases), and Korea, Republic of (South Korea) (32 cases).

The events (>200 occurrences) indicative of medication error were coded to the PTs Product administered to patient of inappropriate age (6485 events), Poor quality product administered (1710 events), Product administration error (1143 events), Product preparation error (550 events), Product temperature excursion issue (466 events), Underdose (387 events), Product preparation issue (302 events), Expired product administered (260 events), Vaccination error (214 events), Inappropriate schedule of product administration (204 events).

Medication Errors Harm Analysis

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65 PTs Wrong technique in product usage process, Medication error, Vaccination error, Product temperature excursion issue

66 Among the reviewed cases, there were additional 1119 cases reported errors in paediatric subjects of 5 to <12 years, who are the authorised individuals to receive Tris/Sucrose Orange cap presentation, but these subjects received PBS/Sucrose Purple cap presentation. As these subjects did not receive Orange cap presentation, these cases are not included in Section 9.2.1. Of note, 24 of 1119 cases involved intentional administration and they were excluded from the analysis of the overall medication error dataset.

67 Among the reviewed cases, there were additional 199 cases involving children aged 5 - <12 years who were vaccinated with the adult formulation (30 micrograms/dose, PBS presentation) before the approval of the appropriate Tris/Sucrose pediatric presentation in their respective countries; these cases were hence not consistent with medication errors pertaining to the new tris-sucrose pediatric presentation. Of note, 6 of 199 cases involved intentional administration and they were excluded from the analysis of the overall medication error dataset.
Among the medication error cases, the following scenarios, categorised according to the EMA guidance “Good practice guide on recording, coding, reporting and assessment of medication errors” (EMA/762563/2014) were described:

- Medication errors associated with harm [i.e., resulting in adverse reaction(s)] were reported in 48 cases (0.5% of medication error cases relevant to Tris/Sucrose Orange cap presentation).

- Medication errors without harm [i.e., not resulting in adverse reaction(s)] were reported in 9367 cases (99.4% of medication error cases relevant to Tris/Sucrose Orange cap presentation).

- Potential errors were reported in 11 cases (0.1% of medication error cases relevant to Tris/Sucrose Orange cap presentation).

- There were no cases reporting intercepted medication errors during the reporting interval.

Of note, some cases involved more than one medication error.

**Medication errors with harm (55 medication errors in 48 cases)**

In 48 cases involving 55 medication error events, 1 event was assessed as serious and the remaining 54 medication errors were assessed as non-serious. These 48 cases originated from Germany (15 cases), US (9 cases), Italy (7 cases), Poland (6 cases), Australia (5 cases), Philippines, Japan (2 cases each), Spain and Lithuania (1 case each).

**Serious medication error**

In 1 case, a serious medication error was reported that potentially contributed to the occurrence of SAEs.

A consumer reported that a female child subject of unspecified age received Tris/Sucrose Orange cap presentation for COVID-19 immunization. The treating nurse administered the wrong dose of vaccine to the child. She was supposed to dilute it, instead she gave a full vial of the COVID Vaccine (PTs: Product preparation error; Overdose). Following administration, the subject experienced dizziness, nausea, and increased heart rate. These events were considered serious as the subject requiring hospitalisation and it was unknown if the subject recovered from these events. No further information was available.

**Non-serious medication errors**

In 47 cases (involving 54 medication error events), non-serious medication errors potentially contributed to the occurrence of non-serious AEs.

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68 AEs may be co-reported in a case, but they are not considered to be a result of the medication error.
The most frequently reported clinical events (≥4 events) in these cases were coded to the PTs Pyrexia (15 events), Vaccination site pain (11 events), Pain in extremity (10 events), Headache (9 events), Vomiting (5 events), Asthenia and Fatigue (4 events).

- **Errors in the administration of incorrect formulation and/or errors in the administration vaccine dosage due to incorrect formulation (4 events)**
  - Errors in the choice of formulation included adolescent (2)/adult (1) subjects who received paediatric formulation (Tris/Sucrose Orange cap presentation) (3 events).
  - Error included adolescent subject receiving paediatric formulation (Tris/Sucrose Orange cap presentation) that led to error in vaccine dosage (1 event).

- **Errors in the preparation / administration of incorrectly prepared vaccine (6 events)**
  - Errors in preparation: Errors included dilution before use not performed (4 events) or incorrect dilution (1 event)
  - Error in administration of inadequately prepared vaccine: Error involved in administering undiluted vaccine (1 event)

- **Other errors (44 events)**
  - Errors of vaccine dosage included smaller doses (6 events), receiving lower volume of correct dosage, additional dose (1 event each), or unspecified inappropriate dose (3 events)
  - Other errors included unintentional administrations of vaccine to paediatric individuals below 5 years of age (15 events), errors in the route of administration (10 events), administered expired vaccine (3 events), administration technique errors (1 event) or unspecified administration/vaccination errors (4 events).

**Medication errors without harm (11,983 medication errors in 9367 cases)**

There were 9367 cases involving 11,983 medication error events without harm. The cases (≥32 cases) originated from Philippines (5992 cases), US (2623 cases), Australia (271 cases), Germany (94 cases), Canada (93 cases), Japan (66 cases), Spain (60 cases), and Korea, Republic of (South Korea) (32 cases).

- **Errors in the administration of incorrect formulation and/or errors in administration of vaccine dosage due to incorrect formulation (6682 events)**
  - Errors in the choice of formulation included adolescent (3615)/adult subjects (2839) receiving paediatric formulation (Tris/Sucrose Orange cap presentation) (6454 events), or elderly subjects receiving paediatric formulation (Tris/Sucrose Orange cap presentation) (31 events)
Errors included administration of smaller doses or errors in vaccine dosage as the adult/adolescent subjects receiving paediatric formulation (Tris/Sucrose Orange cap presentation) (195 events), administration of smaller doses as a consequence of giving paediatric formulation (Tris/Sucrose Orange cap presentation) to elderly subjects (2 events).

- **Errors in preparation and/or administration of incorrectly prepared vaccine (1095 events)**
  - **Errors in preparation**: Errors included dilution before use not performed (412 events), dilution before use with larger/smaller/incorrect volume of diluent (144 events), incorrect/improper dilution (125 events), smaller doses due to use of large volume of diluent (119 events), diluent from the same vial was used for multiple vials (108 events), improper dilution that led to error in vaccine dosage (30 events), smaller doses due to vial mixing error/improper dilution (23 events), dilution before use with a different solvent (21 events), diluted orange cap was given to adult subject (12 events), less/more doses in the vial after preparation (6 events), Orange cap formulation was mixed with Purple cap formulation (5 events), larger doses due to dilution error/improper dilution (3 events), Orange cap presentation was mixed with the amount of diluent recommended for Purple cap presentation, larger doses as the dilution before use was not performed (2 events each), or unspecified preparation/dilution errors (10 events).
  - **Errors in administration of incorrectly prepared vaccine**: Errors involved administration of undiluted vaccine (33 events), administered vaccine after diluting with incorrect diluent (30 events), administered after improper dilution (8 events), administered orange cap formulation after mixing with purple cap formulation (2 events).

- **Other errors (4206 events)**
  - Errors of vaccine dosage included smaller doses due to product leakage / syringe issue / subject non-compliance / needle issue or due to other issues (72 events), unspecified inappropriate dose (35 events), larger doses (19 events), administration of incorrect dose/booster dose/invalid dose/additional doses (10 events), not enough doses or incorrect dosage in the vial, administration of fourth and fifth doses together/half of the booster dose (2 events each) or given smaller volume of correct dosage (1 event),
  - Other errors reported administration of expired vaccine (1320 events), temperature excursion or administration of vaccine after temperature excursion (946 events), administration of poor quality vaccine/defective material (813 events), inadequate storage or administration of vaccine after inadequate storage (471 events), unspecified vaccination/administration errors (211 events), Inappropriate schedule/administration of second dose of the vaccine earlier or later than the 3-week schedule (204 events),
These events included unintentional administrations of vaccine to paediatric individual below 5 years of age (33 events), administration technique errors (17 events), wrong product/vaccine administered (15 events), errors of vaccination at the wrong anatomical site (13 events), accidental exposure (9 events), incorrect route of administration (6 events), dispensed vaccine after expiry (2 events), product not completely administered due to interruption, multiple use of single-use product (details unspecified), unspecified prescribing error, product leakage during administration, or the syringe was shaken before administration (1 event each).

Potential medication errors (13 medication errors in 11 cases)

There were 11 cases from US (8 cases), UK, Canada, and Italy (1 case each)

- The potential errors were described as the confusion with the expiration date/formulation to be used (4 events), user requesting clarification if the paediatric vial stored in the standard freezer can be used/about storage instructions in the label and/or package (3 events), beyond use date was not written on the box, user was unsure if the vaccine was given via appropriate route, printing error in the label, reporter was unsure if the subject received vaccine after dilution, adult vial with paediatric label sticker, expiration date in the carton and on the vial was different (1 event each).

9.2.2. Errors pertaining to the new formulation of BNT162b2 – Adult/Adolescent
Tris/Sucrose Grey cap presentation (30 mcg/dose –[Do not dilute]- in adults and children 12 years and older):

Search criteria used for selecting the below cases for discussion: Adult/adolescent/elderly subjects received Tris/Sucrose Grey cap (30 mcg/dose) presentation; Tris/Sucrose Grey cap formulation was used or administered to paediatric age group of 5 to <12 years instead of Tris/Sucrose Orange cap presentation.

There were 2750 cases (4.1% of the relevant medication error cases) reporting 5496 events indicative of medication errors related to Tris/Sucrose Grey cap presentation (adult/adolescent formulation). Of the 2750 cases, in 53 cases paediatric subjects of 5 to <12 years of age range received Tris/Sucrose Grey cap presentation instead of Tris/Sucrose Orange cap presentation by mistake. These 2750 cases originated from the US (2684 cases), France (36 cases), Germany (17 cases), Spain, Canada, UK, Australia, Italy (2 cases each), Puerto Rico, Finland and Israel (1 case each).

The medication errors (≥42 occurrences) reported in 2750 cases were coded to the PTs Poor quality product administered (2429 events), Product administration error (1665 events), Product temperature excursion issue (733 events), Expired product administered (193 events), Underdose (96 events), Product preparation issue (94 events), Product storage error (86 events), Inappropriate schedule of product administration (45 events), Product administered to patient of inappropriate age (43 events), and Product preparation error (42 events).
Medication Errors Harm Analysis

Among the medication error cases, the following scenarios, categorized according to the EMA guidance “Good practice guide on recording, coding, reporting and assessment of medication errors” (EMA/762563/2014) were described:

- Medication errors associated with harm [i.e., resulting in adverse reaction(s)] were reported in 12 cases (0.4% of medication error cases relevant to Tris/Sucrose Grey cap presentation)

- Medication errors without harm [i.e., not resulting in adverse reaction(s)]\textsuperscript{69} were reported in 2727 cases (99.2% of medication error cases relevant to Tris/Sucrose Grey cap presentation)

- Potential error was reported in 11 cases (0.4% of medication error cases relevant to Tris/Sucrose Grey cap presentation)

- There were no cases reporting intercepted medication errors during the reporting interval.

Of note, some cases involved more than one medication error.

\textit{Medication errors with harm (17 medication errors in 12 cases)}

In 12 cases involving 17 medication error events, all the events were assessed as non-serious. These 12 cases originated from US (10 cases), Australia, and France (1 case each).

\textbf{Serious medication error}

There were no cases that reported serious medication error.

\textbf{Non-serious medication errors}

In 12 cases (involving 17 medication error events), non-serious medication errors potentially contributed to the occurrence of non-serious AEs.

The clinical events (\(\geq 2\) events) reported in these cases were coded to the PTs Pyrexia (3 events) and Dizziness (2 events).

- \textit{Errors in the administration of incorrect formulation (1 event)}
  
  - Errors in the choice of formulation included paediatric subjects receiving adult/adolescent formulation (Tris/Sucrose Grey cap presentation [1 event])

\textsuperscript{69} AEs may be co-reported in a case, but they are not considered to be a result of the medication error.
• **Other errors (16 events)**

  - Errors in vaccine dosage included the 3rd booster dose/additional dose, smaller doses, or unspecified inappropriate doses (2 events each)

  - Other errors included incorrect route of administration (intradermal/IV), administered after expiry/administered expired vaccine, administered poor quality vaccine (2 events each), administration of vaccine 12 hours after first puncture, administration of vaccine after inadequate storage, administration of wrong product, and unspecified vaccination error (1 event each).

**Medication errors without harm (5467 medication errors in 2727 cases)**

There were 2727 cases involving 5467 medication error events without harm and they are categorised into:

• **Errors in the administration of incorrect formulation and/or errors in the administration vaccine dosage due to incorrect formulation (51 events)**

  - Errors in the choice of formulation included paediatric subjects receiving adult/adolescent formulation (Tris/Sucrose Grey cap presentation) (48 events)

  - Errors included paediatric subject receiving adult/adolescent formulation (Tris/Sucrose Grey cap presentation) that led to error in vaccine dosage (3 events).

• **Errors in the preparation / administration of incorrectly prepared vaccine (206 events)**

  - **Errors in preparation:** Errors included dilution of vaccine before use (83 events), subjects received smaller doses as the vaccine was diluted before use (59 events), smaller doses due to preparation error (20 events), incorrect/improper dilution (17 events), Grey cap presentation was diluted and given to paediatric subject (9 events), undiluted Grey cap presentation was given to paediatric subject (4 events), Grey cap presentation was diluted with Orange cap presentation (1 event).

  - **Errors in administration of incorrectly prepared vaccine:** Errors involved administration of diluted vaccine/administered after improper dilution (13 events)

• **Other Errors (5210 events)**

  - Errors of vaccine dosage included administration of doses from 2 different vials (20 events), smaller doses/smaller doses due to improper injection/smaller doses due to syringe leakage/smaller doses as the part of the liquid was not administered (16 events), third/fourth/fifth dose (7 events), additional or booster doses (4 events), larger doses (3 events), doses left in the vial after completion of 6 doses (2 events) or unspecified inappropriate dose (5 events)
Other errors included temperature excursion or administration of vaccine after temperature excursion (1508 events), administration of poor quality vaccine/defective material (1330 events), administered after expiry/administration of expired vaccine (1237 events), inadequate storage or administration of vaccine after inadequate storage (877 events), vaccine given after 12 hours of initial vial puncture (131 events), inappropriate schedule/second dose was given earlier or later than the 3-week schedule (45 events), technique errors or other administration errors (12 events), incorrect route of administration or errors of vaccination at the wrong anatomical site (8 events), administered vaccine from the old vial, administration of wrong vaccine (1 event each) or other unspecified error (3 events).

**Potential medication error (12 medication errors in 11 cases)**

- There were 11 cases from US (9 cases), France and Germany (1 case each)

- The potential errors were described as no expiration date on the vial (7 events), confusion in the formulation or expiration date (2 events), reporter was unsure if the subject received vaccine after dilution, user requesting clarification about the expiration date on the label, user suggested to include information note about dilution to avoid confusion as the dilution was performed before use by mistake (1 event each).

**Conclusion**

Overall, among the 66,764 relevant medication error PM cases, 1326 cases (0.3% of the total interval cases, 2.0% of total relevant medication error cases) were considered harmful, 70 of which (0.1% of total relevant cases) were serious and most of them originated from vaccine administration issues (50 cases of 70 serious cases with harm).

The potential for medication errors with the new presentations are mitigated through the information in the vaccine labelling and available educational materials for the HCPs administering the vaccine.

Some medication errors are expected to occur despite written instructions for handling the vaccine (thawing, dilution, preparation) and educational activities for the HCPs or due to national or regional recommendations regarding dosing schedules that may differ from recommendations in the product labelling. The number and seriousness of the reported medication errors events do not indicate any trend and potential needs for any additional mitigation activity.
10. NON-CLINICAL DATA

During the reporting period, no new nonclinical safety findings were identified.

11. LITERATURE

Nonclinical (Published)

A search of the Medline and Embase databases identified no nonclinical studies that presented important new safety findings for BNT162b2.

Clinical (Published)

A search of the Medline and Embase databases identified 8 clinical trials that presented important new safety findings for BNT162b2. These are presented in Table 31 below grouped as follows: a) At risk patients; b) Special patient population/Pregnancy; c) Efficacy and Effectiveness and d) Other Safety Information (citations with a brief comment).

See Appendix 5 for the abstracts. Full publications are available upon request.

Table 31. Clinical Literature Articles that Presented New Safety Information in the Reporting Interval

<table>
<thead>
<tr>
<th>Citation/Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) At Risk patients</td>
</tr>
<tr>
<td>This article described a reduced immunoresponse to BNT162b2 in patients treated with immunosuppressants. Section 4.4. Special warnings and precautions for use (Immunocompromised individuals) of the EU SmPC includes a warning regarding vaccination in immunocompromised patients, as follows, “The efficacy and safety of the vaccine has not been assessed in immunocompromised individuals, including those receiving immunosuppressant therapy. The efficacy of Comirnaty may be lower in immunocompromised individuals.”</td>
</tr>
<tr>
<td>Use in immunocompromised patients is considered missing information for BNT162b2; please refer to Section 16.4.2 Description of Missing Information for the description of this topic and to Section 16.3.5.5 Use in Immunocompromised Patients for the summary of the cases received during the reporting period.</td>
</tr>
<tr>
<td>b) Special Patients Population (Pregnancy)</td>
</tr>
<tr>
<td>This article contributes to the growing evidence that risk of spontaneous abortion after COVID-19 vaccine immunisation during the first trimester of pregnancy is commensurate with the predicted risk in non-vaccinated pregnant women.</td>
</tr>
<tr>
<td>Use in pregnancy and while breastfeeding patients is considered missing information for BNT162b2; please refer to Section 16.4.2 Description of Missing Information for the description of this topic and to Section 16.3.5.3 Use in Pregnant/Lactating Women for the summary of the cases received during the reporting period.</td>
</tr>
</tbody>
</table>
Table 31. Clinical Literature Articles that Present New Safety Information in the Reporting Interval

<table>
<thead>
<tr>
<th>Citation/Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>c) Efficacy and Effectiveness</td>
</tr>
<tr>
<td>5. Kliker L, Zuckerman N, Atari N et al. COVID-19 vaccination and BA.1 breakthrough infection induce neutralising antibodies which are less efficient against BA.4 and BA.5 Omicron variants, Israel, March to June 2022. <a href="http://www.eurosurveillance.org">www.eurosurveillance.org</a> submitted on 12 Jul 2022 / accepted on 28 Jul 2022 / published on 28 Jul 2022.</td>
</tr>
</tbody>
</table>

Please refer to Section 17.2 Newly Identified Information on Efficacy and Effectiveness for the comments on these articles and to Section 16.3.4.5. Lack of Therapeutic Efficacy for the review of the cases indicative of LOE reported in the current interval period.

d) Other Safety Information

<table>
<thead>
<tr>
<th>Citation/Comment</th>
</tr>
</thead>
</table>

This study suggests that the COVID-19 vaccine might be associated with increased risk of Sudden Sensorineural Hearing Loss; however, the effect size is very small. The study had various limitations and no causality assessment has been conducted. The MAH will continue to monitor using routine pharmacovigilance.

Please refer to Appendix 6A.3 for further discussion of this article and for cumulative review of cases indicative of hearing loss.

<table>
<thead>
<tr>
<th>Citation/Comment</th>
</tr>
</thead>
</table>

---

70 Articles 5 and 6 were published/posted after the DLP, but they include information relevant to the reporting period of this PSUR.
Table 31. Clinical Literature Articles that Presented New Safety Information in the Reporting Interval

<table>
<thead>
<tr>
<th>Citation/Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>In this study, BNT162b2 was associated with an immediate negative effect on anticoagulation control in patients treated with vitamin K antagonists. The author though cannot exclude the possibility that the effect on anticoagulation control was due to dose adjustments to avoid complications and patients themselves could have decided to decrease the dosage in the days following COVID-19 vaccination as they might be afraid for bleeding complications after intramuscular injection. This could result in a higher percentage of subtherapeutic INRs after vaccination. In addition, the authors use a surrogate variable for bleeding complications (INR &gt;5). The possible effects of vaccines on anticoagulation control remain debated even though several prospective studies have been performed (mostly on the effect of the influenza vaccine on anticoagulation control), but overall results were conflicting. As of now, there is no biological or pharmacological plausibility for a vaccine – drug interaction. The MAH will continue to monitor using routine pharmacovigilance.</td>
</tr>
<tr>
<td>Please refer to Section 16.3.3.1.19 Thromboembolic AESIs for the summary of cases indicative of coagulopathy received in the reporting period.</td>
</tr>
</tbody>
</table>

All Other Published Sources

In the final AR for PAM-MEA-002.11 - 12. SMSR (1st SBSR) received on 09 February 2022 (EMEA/H/C/005735/MEA/002.11), the MAH was requested to include in the 2nd SBSR the following article, published in the reporting interval of the PSUR # 3:

  https://www.medrxiv.org/content/10.1101/2022.01.17.22269263v1.article-metrics.

The above article was included and discussed in the SBSR no. 2 dated 04 March 2022.

In the final AR for PAM-MEA-002.12 13th SMSR (2nd SBSR) dated 05 April 2022 (EMA/PRAC/202255/2022), the MAH was requested to discuss the following publication regarding SSNHL in association with COVID-19 vaccination:


The abstract of the above article and the discussion are available in Appendix 6A.3.

Unpublished manuscripts

In the final assessment report for PAM-MEA-002.12 13th SSR (2nd SBSR) dated 05 April 2022 (EMA/PRAC/202255/2022), the MAH was requested to include in the 3rd SBSR the following ACIP presentation, presented in the reporting interval of the PSUR # 3:

The above presentation on myocarditis outcomes following mRNA COVID-19 vaccination was included and discussed in the SBSR no. 3 dated 06 May 2022.

12. OTHER PERIODIC REPORTS

During the reporting period, the MAH did not submit another PSUR for BNT162b2. However, the MAH was requested to prepare Summary Monthly Safety Update Reports (SMSRs), in accordance with the EMA coreRMP19 Guidance (EMA/544966/2020) and, as applicable, with ICH Guideline E2C (R2) Periodic Benefit-Risk Evaluation Report [Step 5, January 2013] considering the information for the evidence from post-EUA/conditional marketing authorisation approval data sources.

Following the proposal of discontinuation of SSR submission by PRAC included in the final PRAC AR of the 3rd SBSR (Report EMA/PRAC/577594/2022 dated 08 June 2022), the preparation of the SBSR was discontinued.

The list of periodic reports prepared and submitted by the MAH during the reporting period is provided below.

<table>
<thead>
<tr>
<th>Periodic Report Type</th>
<th>No.</th>
<th>Reporting Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summary Bimonthly Safety Report (SBSR)</td>
<td>2</td>
<td>16 December 2021 through 15 February 2022</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>16 February 2022 through 15 April 2022</td>
</tr>
<tr>
<td>Abbreviated SMRSa</td>
<td>2</td>
<td>16 December 2021 through 15 January 2022</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>16 February 2022 through 15 March 2022</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>16 April 2022 through 15 May 2022</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>16 May 2022 through 15 June 2022</td>
</tr>
</tbody>
</table>

a. Submitted to non-EEA countries.

During the reporting period, no new significant safety findings were identified for BNT162b2 in other periodic reports prepared by the MAH.

13. LACK OF EFFICACY IN CONTROLLED CLINICAL TRIALS

Study C4591007 is the ongoing, randomised, placebo-controlled, Phase 1/2/3 paediatric study in healthy children from 6 months to <12 years of age. The Phase 2/3 primary immunogenicity objective in children from 6 months to <5 years of age was immunobridging the immune responses against SARS-CoV-2 wild-type strain from children 2 to <5 years and 6 months to <2 years of age in Study C4591007 compared to young adults 16 to 25 years of age in the Phase 3 efficacy Study C4591001. Immunobridging data after Dose 2 met success criteria for the 6 months to <2 years group and did not meet GMR success criteria (but met seroresponse criteria) for the 2 to <5 years of group, compared to young adults 16 to 25 years of age. Given emerging real-world data in the Omicron wave that two-dose protection against symptomatic infection was only modest, a third dose was evaluated for children <5 years of age. Immunobridging data after Dose 3 met success criteria for the 6 months to <5 years age group, compared to young adults 16 to 25 years of age.
The observed VE from at least 7 days after Dose 2 to before Dose 3 for BNT162b2 3-μg administered to children 6 months to <5 years of age without prior evidence of SARS-CoV-2 infection before or during the vaccination regimen was 28.3% (2-sided 95% CI: 8.0%, 43.9%) based on 163 cases in the BNT162b2 group and 113 cases in the placebo group, adjusted for surveillance time (noting 2:1 randomisation of vaccine:placebo). In this population, observed VE against Delta and Omicron was 70.2% (2-sided 95% CI: 27.2%, 88.5%) and 21.8% (2-sided 95% CI: -1.7%, 39.7%), respectively. Note that most of the cases across this age population that were confirmed post-Dose 2 to before Dose 3 were reported in January 2022.

The observed VE from at least 7 days after Dose 3 to the cutoff date (29 April 2022) across the total population of children 6 months to <5 years of age was 80.3% (2-sided 95% CI: 13.9%, 96.7%) based on 3 cases in the BNT162b2 group and 7 cases in the placebo group, adjusted for surveillance time (noting 2:1 randomization of vaccine:placebo). Note that all post-Dose 3 cases were reported in February through April 2022.

The observed VE after 3 doses in children 6 months to <5 years of age in Study C4591007 is consistent with real-world effectiveness data for older age groups, which indicate that in adolescents (12 to 17 years of age) and adults (18 years of age and older), three doses of BNT162b2 are needed to provide a high level of protection against symptomatic disease due to Omicron.  

14. LATE-BREAKING INFORMATION

As reported in Section 16.1 Summary of Safety Concerns, on 08 July 2022 (after the DLP of this PSUR), the MAH submitted the EU RMP version 5.1 to remove the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP.


recommendation received in March 2022 with the positive opinion for the Type II Variation 87 (EMEA/H/C/005735/II/0087).

After the DLP, an updated CDS (version 14.0) was made effective on 26 July 2022 to extend the indication of BNT162b2 to individuals 6 months of age and older, to include posology and method of administration for the Tris/Sucrose presentation 3 micrograms/dose, and to add irritability, injection site tenderness, myocarditis and pericarditis as ADRs in section 4.8 Undesirable effects. Please refer to Section 4 Changes to the Reference Safety Information for additional details on the changes.

15. OVERVIEW OF SIGNALS: NEW, ONGOING, OR CLOSED

Signal Overview

Signals detected for BNT162b2 during the reporting interval are presented below in Table 32 along with the ongoing signals and signals closed during the reporting interval.

Appendix 3 provides a summary of the safety signals that were new, ongoing, or closed during the reporting interval. See Section 16.2.1 for evaluation of signals that were closed during the reporting interval and Section 16.3 for evaluation of new information for previously known risks not considered to constitute a newly identified signal.

**Table 32. Overview of Signals (at DLP 18 June 2022)**

<table>
<thead>
<tr>
<th>Signal</th>
<th>Signal Status*</th>
<th>Source</th>
<th>Category*</th>
<th>EMA Regulatory Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myocarditis and Pericarditis</td>
<td>Closed</td>
<td>Other: Routine safety surveillance</td>
<td>Important identified risk</td>
<td>-</td>
</tr>
<tr>
<td>Irritability</td>
<td>New and closed</td>
<td>Clinical Trial C4591007 unblinded review of data in 6 months to &lt;5-year-old (Pfizer)</td>
<td>Identified risk (not &quot;important&quot;)</td>
<td>-</td>
</tr>
<tr>
<td>Appendicitis</td>
<td>Re-opened and closed</td>
<td>Inquiry from a competent authority (Singapore BoH)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Hemolytic anemia</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (Saudi Arabia SFDA)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Uveitis</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (Health Canada)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Capillary Leak Syndrome (CLS)</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>PAM-SDA-051 EPITT 19743</td>
</tr>
</tbody>
</table>
Table 32. Overview of Signals (at DLP 18 June 2022)

<table>
<thead>
<tr>
<th>Signal</th>
<th>Signal Status*</th>
<th>Source</th>
<th>Category*</th>
<th>EMA Regulatory Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corneal Graft Rejection</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>PAM-SDA-055</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EPTIT 19789</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>Ongoing and</td>
<td>Notification from a competent authority (Netherlands Lareb)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>closed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cerebral venous sinus thrombosis (CVST)</td>
<td>Ongoing and</td>
<td>Inquiry from a competent authority (Switzerland Swissmedic)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>closed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lymphocytic colitis</td>
<td>New and closed</td>
<td>Scientific literature(^2)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Chronic Urticaria</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Polymyalgia Rheumatica (PMR)</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Subacute Thyroiditis (SAT)</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Cerebrovascular Accident (CVA)/Stroke</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (Australia TGA)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td>Amenorrhea</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>PAM-SDA-052</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EPTIT 19784</td>
</tr>
<tr>
<td>Heavy Menstrual Bleeding</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (EMA PRAC)</td>
<td>No risk</td>
<td>PAM-SDA-053</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>EPTIT 19783</td>
</tr>
<tr>
<td>Loss of/Altered Taste and Smell</td>
<td>New and closed</td>
<td>Inquiry from a competent authority (Australia TGA)</td>
<td>No risk</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hearing Loss</td>
<td>Re-opened and</td>
<td>Inquiry from a competent authority</td>
<td>Not yet</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Ongoing</td>
<td></td>
<td>determined</td>
<td></td>
</tr>
</tbody>
</table>

\(^*\) Reflects the MAH position and entry into MAH signal log following evaluation. This may not be the same as the position of the competent authority.

Other Safety Topics Not Considered Signals

Continued monitoring or a cumulative review was requested in an assessment report or recommended in a previous SMSR of the following unlisted events/topics that a competent

authority did not consider a signal and were also determined to not be a safety signal by the MAH.

Factors that were considered in coming to this conclusion included one or more of the following:

- Whether the AE is new for the product;
- Seriousness, severity, increased frequency or medical significance of the data;
- High or rapidly increasing statistical disproportionality score;
- Potential public health impact;
- Factors suggestive of a relationship to the drug when considering disease knowledge, biological plausibility, mechanism of action of the drug or the drug class, alternative etiologies based on clinical and scientific experience, and temporal clustering of events.

The safety topics reviewed are the following:

Dizziness (Appendix 6A.1)
Acquired haemophilia\(^\text{73}\) (Appendix 6A.2)
MIS-C/A (Appendix 6A.4)
Autoimmune hepatitis (Appendix 6A.5)

16. SIGNAL AND RISK EVALUATION

16.1. Summary of Safety Concerns

Table 33 summarises the important risks and missing information for BNT162b2 at the beginning of the reporting interval, as per EU-RMP version 4.0 adopted on 26 November 2021 (Procedure number: EMEA/H/C/005735/X/0077).

There were no changes to the safety concerns during the reporting period.

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\(^{73}\) This was considered a signal after the DLP. Please refer to the evaluation in Appendix 6A.2.
Table 33. Ongoing Safety Concerns

<table>
<thead>
<tr>
<th>Important identified risks</th>
<th>Anaphylaxis</th>
</tr>
</thead>
</table>
|                           | Myocarditis and Pericarditis

<table>
<thead>
<tr>
<th>Important potential risks</th>
<th>Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Missing information</td>
<td>Use in pregnancy and while breast feeding</td>
</tr>
<tr>
<td></td>
<td>Use in immunocompromised patients</td>
</tr>
<tr>
<td></td>
<td>Use in frail patients with co-morbidities (e.g. chronic obstructive pulmonary disease [COPD], diabetes, chronic neurological disease, cardiovascular disorders)</td>
</tr>
<tr>
<td></td>
<td>Use in patients with autoimmune or inflammatory disorders</td>
</tr>
<tr>
<td></td>
<td>Interaction with other vaccines</td>
</tr>
<tr>
<td></td>
<td>Long-term safety data</td>
</tr>
</tbody>
</table>

During the reporting period, the MAH submitted the EU-RMP version 5.0:

- Consolidation of the RMPs versions 2.6 (procedure EMEA/H/C/005735/II/0087) and 4.0 (Procedure number: EMEA/H/C/005735/X/0077), adopted on 10 March 2022.

After the DLP of this PSUR, the MAH submitted 3 additional EU-RMPs:

1. version 5.1 on 08 July 2022 (Procedure Number: EMEA/H/C/005735/X/0138):
   - to include the 6 months to <2 years and 2 years to <5 years phase 1 and phase 2/3 data from interventional clinical study C4591007 for the line extension of COMIRNATY® 3 µg Concentrate for dispersion for injection for infants and children between 6 months to 4 years of age;
   - to remove the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP recommendation received on March 2022 with the positive opinion for the Type II Variation 87 (EMEA/H/C/005735/II/0087).

2. Version 6.0 on 19 July 2022 (Procedure Number: EMEA/H/C/005735/II/0140):
   - To support the extension of the indication to ≥12 years of age to receive an additional booster (fourth) dose of bivalent Omicron-(BA.1) modified vaccine.

   - To support the extension of the indication to introduce a first or second booster dose of a bivalent Omicron variant-modified vaccine, (BNT162b2 15 µg +

---

34 "Myocarditis and Pericarditis" has been added as important identified risk in the EU-RMP version 2.3 submitted on 05 August 2021; this version received a positive CHMP opinion on 30 September 2021.
BNT162b2 OMI BA.4/5 15 μg, total 30 μg), given ≥3 months after the primary series or ≥4 months after the third dose in individuals ≥12 years of age.

There are no further changes to propose with regard to the safety concerns in the EU-RMP.

16.2. Signal Evaluation

Please refer to Table 32 for signals that were ongoing and closed during the reporting interval.

Post-approval regulatory requests (worldwide)

According to the corePSUR19 guidance75, the conclusion of the evaluation resulting from the safety review of hearing loss requested by the PRAC and Health Canada in the context of final ARs of the 13th and 14th SMSRs is summarised below; the complete review is reported in Appendix 6A.3.

Procedure no EMA/PRAC/202255/2022 (13th SMSR-2nd SBSR):
The MAH is requested to discuss the following publications regarding sudden sensorineural hearing loss (SSNHL) in association with COVID-19 vaccination:


Furthermore, the MAH is requested to conduct age-stratified O/E analyses for the AESI of sudden hearing loss using the age-specific background incidence rates of SSNHL reported in the following publication: Alexander T and Harris J. Incidence of Sudden Sensorineural Hearing Loss. Otol Neurotol. 2013 Dec;34(9):1586-9. doi:10.1097/MAO.0000000000000222.

Procedure no. EMA/PRAC/577594/2022 (14th SMSR-3rd SBSR): The MAH is requested to perform a cumulative review on the association between sudden sensorineural hearing loss and Comirnaty exposure, including a review of the relevant new literature published after the reporting period of the 14th SSR, and a discussion on the need to update of the Comirnaty product information and/or RMP, if applicable.

Health Canada 31 May 2022 (13th SMSR-2nd SBSR): The WHO recently published an update regarding COVID-19 vaccines and hearing loss. Signal detection activities at the

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UMC up to 22 Feb 2022 retrieved 164 cases with HLT Hearing losses (142 cases for Comirnaty) and 367 cases with the PT Tinnitus (293 for Comirnaty) with COVID-19 vaccines. Based on well documented cases, alternative causes were not identified for most of the patients and a plausible mechanism of action has been suggested. As such, provide a cumulative review of all cases of tinnitus and hearing loss. This cumulative review should include analyses of all cases, stratified by age, gender, doses administered, time to onset, and any other relevant information. An observed-to-expected analysis should be provided including the appropriate risk window. An appropriate case definition including a causality assessment should also be provided.

Conclusion

Taking into account the totality of the data available, a causal association between Comirnaty and hearing loss or tinnitus cannot be concluded at this time. Routine pharmacovigilance will continue.

16.2.1. Evaluation of Closed Signals

Table 34 provides the evaluation of the signals closed during the reporting period.

<table>
<thead>
<tr>
<th>Signal</th>
<th>Evaluation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signals Determined to Not be Risks</td>
<td>Appendixitis was identified as a signal during the reporting period based on a competent authority (Singapore BoH) inquiry following 18 local reports. The Pfizer safety database search through 01 April 2022 revealed 690 cases and those with sufficient information provided did not show any trends considered inconsistent with the underlying epidemiology and/or natural course of the condition. Placebo-controlled clinical trial data from the pivotal Pfizer-run studies did not reveal any clinically meaningful difference between the BNT162b2 and placebo groups. Of 3 large population-based studies from the US, Israel and Sweden, 2 showed no increase in appendicitis after vaccination while 1 showed a slightly increased risk ratio in the vaccinated group compared to the unvaccinated group. Observed to expected analyses conducted were well below 1. A plausible mechanism by which BNT162b2 could cause appendicitis is unknown. The totality of the information was not supportive of a causal association between BNT162b2 and appendicitis signal was closed by the MAH.</td>
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</table>
| Hemolytic anemia     | Haemolytic anaemia was identified as a signal during the reporting period based on a competent authority (Saudi FDA) request for an evaluation. The Pfizer safety database search through 13 January 2022 yielded 176 cases, most of which were confounded or contained insufficient information. Among the cases with no obvious confounder or trigger, a definitive causal association could not be concluded. There were no events of haemolytic anaemia reported in the pivotal clinical trial C4591001. The medical literature yielded one case report and one prospective study of 108 patients with autoimmune cytopenias (56 with autoimmune haemolytic anaemia [AIHA]) who were vaccinated with Pfizer/BNT, Moderna or Astra-Zeneca COVID-19 vaccines. Four elderly patients with AIHA had a clinically significant haemoglobin reduction requiring treatment adjustment (2 had received Pfizer/BNT vaccine). Notably, autoimmune cytopenia recurrences were not predictable, since they occurred in both patients on active treatment and off therapy, independently from AIHA type, after either the first or
### Table 34. Evaluation of Closed Signals During the Reporting Interval

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<th>Signal</th>
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<td>the second dose, and regardless of vaccine type. Observed to expected analyses conducted were below 1. Based on the totality of available information, a causal association between BNT162b2 and haemolytic anaemia could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Uveitis</td>
<td>Uveitis was identified as a signal during the reporting period based on a competent authority (Health Canada) request for a cumulative review. The Pfizer safety database search through 04 April 2022 yielded 538 cases, 121 of which were medically confirmed and did not report confounding factors or an implausible time to onset. Of these, 9 were determined to have a possible causality based on individual assessment. During the placebo-controlled period in the pivotal clinical trial C4591001, one case was reported in the placebo group and no cases were reported in the BNT162b2 group from Dose 1 to 1 month after Dose 2. The medical literature consisted of case reports and case series descriptions with one population-based study estimating the prevalence rates of uveitis coincident with COVID-19 vaccination as 0.9 cases per million doses or less. Observed to expected analyses conducted using both a low and high range of background rates were well below 1 overall, by dose and within age and sex strata. Based on the totality of available information, a causal association between BNT162b2 could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders</td>
<td>A cumulative review of autoimmune and inflammatory disorder exacerbations was requested in an updated PSUR Assessment Report received from EMA/PRAC during the reporting period (30 December 2021). The search of the safety database used SMQ Immune-mediated/autoimmune disorders (narrow terms), HLT Autoimmune disorders, HLT Immune disorders NEC and HLT Neuromuscular junction dysfunction. There were 2223 cases describing a medical history and adverse event of an autoimmune disease (indicating a potential exacerbation). Overall, cases lacked information to ascertain baseline disease status, treatment and other factors which may affect underlying disease activity, despite most reporting exacerbations or potential relapses within 2 days of vaccination. In the placebo-controlled portion of pivotal clinical trial C4591001, 2955/21926 BNT162b2 participants and 2977/21921 placebo participants had underlying autoimmune conditions. Of these, 7 (0.2%) and 4 (0.1%) of BNT and placebo participants, respectively, reported potential aggravations of their autoimmune disorder from Dose 1 to 1 month post Dose 2. From Dose 1 to unblinding, 8 participants in each group (IR/100 person years = 0.7) reported potential aggravations. The medical literature search yielded many studies of COVID-19 vaccination in patients with underlying autoimmune disorders. The findings consistently showed that reported post-vaccination adverse events were similar to those of healthy vaccinees. Most studies did not have control groups of participants with autoimmune disorders who did not receive COVID-19 vaccination, although those that did, did not report that vaccinees had more exacerbations than non-vaccinees. Based on the totality of the available information, a causal association between BNT162b2 and autoimmune disorder exacerbations could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Capillary leak syndrome (CLS)</td>
<td>Capillary leak syndrome, or Systemic capillary leak syndrome (SCLS), was identified as a signal by PRAC on 13 January 2022. The safety database search yielded 44 cases, 2 of which were literature case reports, which occurred in individuals from 20 to 101 years of age. Four cases described a medical history of SCLS. The majority of cases lacked clinical details or provided evidence of an alternative etiology other than vaccination. There were no reported events of SCLS in the placebo (21921) or BNT162b2 group (21926) in the placebo-controlled portion of C4591001 in participants 16 years and older from dose 1 to 1 month</td>
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<td>after dose 2. The medical literature has described cases of CLS occurring after COVID-19 infection and there were only individual case reports of CLS occurring after COVID-19 vaccination. Based on the totality of the available information, a causal association between BNT162b2 and CLS/SCLS could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Corneal graft rejection</td>
<td>Corneal graft rejection was identified as a signal by PRAC on 07 April 2022. The safety database search through 14 April 2022 yielded 42 potential cases describing 40 unique individuals, all adults or elderly. There was no distinguishing trend in the cases with regard to sex, age, dose number, age of graft or time to onset. Of 12 cases with a plausible temporal relationship to vaccination, only 2 did not have reported risk factors for rejection (e.g., increased age of transplant, possible infection, graft surgery complications). Data from large clinical studies C4591001 (snapshot date of 11 April 2022), C4591031 (cut-off date of 08 February 2022) and C4591007 (cut-off dates of 08 October 2021 and 22 March 2022) were searched for PTs, corneal graft rejection and corneal graft failure. Neither of these PTs were reported in the unblinded data from the placebo-controlled portions of the studies. There were 32 clinical trial participants, all ≥16 years of age, who reported a history of corneal transplant or keratoplasty in either Study C4591001 and/or Study C4591031. There were no participants in C4591007 who reported a history of corneal transplant or keratoplasty. The medical literature consisted of case reports which were included in the safety database. There were no mechanistic studies, rather various hypotheses were theorized such as increased vascular permeability, immune responses and immune system deregulation. Based on the totality of the available information, a causal association between BNT162b2 and corneal graft rejection could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Vasculitis</td>
<td>During the reporting period, vasculitis was reviewed initially following a signal noted by the Lareb (Netherlands) and through 15 April at the request of PRAC in the Assessment Report for SBSR 2. Through 15 April 2022, a search of the safety database yielded 868 reports with individual ages ranging from 2 to 98 years. Reported vasculitides included vasculitis (not otherwise described), giant cell arteritis and Henoch-Schonlein purpura. The cases were generally confounded or lacked necessary details to confirm the diagnoses and/or a causal relationship. Phase 2/3 Study C4591001 placebo-controlled unblinded adverse events in participants 16 years and older from Dose 1 to 1 month after Dose 2 (data cutoff date 13 March 2021) was also reviewed for the PT Vasculitis. In the Phase 2/3 safety population, vasculitis was not reported in any of 21926 participants in the BNT162b2 (Pfizer-BioNTech COVID-19 Vaccine) group or in any of 21921 participants in the placebo group. Observed to expected analyses for the 3 most common subtypes of vasculitis (Henoch-Schonlein purpura, Giant cell arteritis, Skin manifestations of vasculitis) have repeatedly been less than one. Based on the totality of the available information, a causal association between BNT162b2 and vasculitis could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Cerebral venous sinus thrombosis (CVST)</td>
<td>Previous to the current PSUR reporting period, a cumulative review of CVST through 24 November 2021 was conducted by the MAH (SBSR 2, Appendix 3.4) in response to a request from a competent authority (Swissmedic). In the SBSR 2 PRAC Assessment Report, the MAH was requested to provide more detail on some of the cases and a further cumulative review of the topic. A search of the safety database yielded 527 cases that were reported through 24 November 2021 and 297 from 25 November 2021 to 15 April 2022. Of 37 cases in patients younger than 75 years of age with no medical history or information pertaining an increased risk for the development reported through 24 November 2021, only 3 were assessed as possible (the remaining were unassessable or unlikely per the WHO-UMC case</td>
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### Table 34. Evaluation of Closed Signals During the Reporting Interval

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<tr>
<td>Lymphocytic colitis</td>
<td>During the reporting period, this signal was identified from a published (literature) case report of a 69-year-old woman who presented for evaluation of severe abdominal pain, nausea, and diarrhea after her second vaccination with Pfizer/BNT COVID-19 vaccine. Within 24 hours of vaccination, she reported onset of diarrhea (2-3 loose to watery stools per day). Symptoms intensified over the next several days to 3-5 watery stools per day with incontinence, abdominal cramping, and nausea. GI PCR and COVID testing were negative and ondansetron and loperamide were started with minimal benefit. Two-months later, a GI consultation was obtained due to persistent symptoms. Work up demonstrated no anemia with normal CRP, celiac serologies, and GI PCR. Colonoscopy on day 98 post-onset revealed patchy erythema in the descending colon and rectosigmoid. Histologic evaluation of mucosal biopsies revealed lymphocytic colitis characterized by numerous lymphocytes infiltrating the epithelium and abundant plasma cells in the lamina propria. A previous colonoscopy performed in 2012 was unremarkable. At her most recent follow-up on day 113 post-onset, the patient reported gradual improvement of abdominal symptoms and diarrhea. This case report was recorded in the Pfizer safety database. There was no other relevant literature information on lymphocytic colitis and COVID-19 vaccination. A search of the safety database through 20 Jan 2022 yielded 40 cases for review (incl index case); in all the cases, there was either no Pfizer/BNT COVID-19 vaccine used, an unconfirmed diagnosis, lack of clinical detail, or the presence of alternative explanations or risk factors for lymphocytic colitis. Based on the totality of the available information, a causal association between BNT162b2 and lymphocytic colitis could not be concluded, and the signal was closed by the MAH.</td>
</tr>
<tr>
<td>Chronic urticaria</td>
<td>This signal was identified following a request for a cumulative review on the subject from EMA FRAC on 09 May 2022. A search of the Pfizer safety database through 09 May 2022 yielded 244 cases; 31 of which described medical histories of chronic urticaria. Of the cases of new onset chronic urticaria, time to onset ranged from 0 to 90 days post vaccination, cases were reported after dose 1, dose 2</td>
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<td>and booster doses, 26 cases described the background of underlying autoimmune disorders, 26 reported hypersensitivity conditions and 14 reported histories of COVID-19. In all, 35 cases specifically reported that the urticaria lasted more than 6 weeks (meeting criteria for chronic urticaria). Sixteen of these reported a time to onset that was reasonably temporally associated with vaccination, and among them only 7 (43.7%) of the cases provided a medical history (85.7% of which implied a predisposition to urticaria or an alternate trigger for it). During the placebo-controlled unblinded period of pivotal study C4591001 (data cut-off 15 April 2022), of participants 12 years and older, chronic urticaria was not reported in any of the 23,068 participants in the BNT162b2 group or the 23,063 participants in the placebo group from Dose 1 to data cutoff date. Observed versus expected analyses were &lt; 1 overall, by dose and within strata of age groups. The MAH considers urticaria as an adverse reaction of BNT162b2, however, a causal association between the vaccine and chronic urticaria was not supported based on the available information.</td>
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<tr>
<td>Polymyalgia rheumatica (PMR)</td>
<td>This signal was identified following a request for a cumulative review on the subject by EMA PRAC in the PSUR 1 Assessment Report. A search of the Pfizer safety database through 18 December 2021 yielded 628 cases, the majority of which were excluded from further consideration due to implausible time to onset, medical history or conditions confounding assessment or lack of clinical detail supporting the diagnosis of PMR. Of the reports providing laboratory data (CRP and/or ESR) supportive of the diagnosis of PMR, most were unassessable or unlikely per WHO-UMC causality criteria. The 54 cases reporting an exacerbation of PMR following vaccination were similarly hindered by lack of information. There were no reports of PMR in the pivotal Phase 2/3 Study C4591001 of individuals 16 years of age (21926 vaccine/21921 placebo) and older from Dose 1 to 1 month after Dose 2 that was placebo-controlled (data cutoff date 13 March 2021). Ten subjects reported polymyalgia rheumatica in the medical history and none of these subjects reported a flare up of the underlying disease. The medical literature search yielded several studies that did not support an association between vaccination and autoimmune disorders or flares. Based on the totality of the available information, a causal association between BNT162b2 and PMR could not be concluded, and the signal was closed by the MAH.</td>
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| Subacute thyroiditis | This signal was identified following a request for a cumulative review on the topic by EMA PRAC on 18 January 2022 following the assessment of PSUR 2. The Pfizer safety database search yielded 498 cases through 18 December 2021. There was a similar number of cases reporting each of the Pts: Thyroiditis subacute, Autoimmune thyroiditis and Thyroiditis. The majority of reports described underlying thyroid disorder or concomitant disorders that represented confounding factors and/or did not provide a sufficient amount of information (medical history, laboratory and other diagnostic data) to allow a proper evaluation. The details of 38 cases that included laboratory work ups confirming hyperthyroid activity, did not provide enough relevant information to confirm the causal association with the vaccine. In the placebo-controlled portion of clinical trial C4591001, in the safety population of participants 16 years and older, there was 1 case of autoimmune thyroiditis reported among 21926 participants in the BNT162b2 group compared with 1 case of autoimmune thyroiditis among 21921 participants in the placebo group from dose 1 to 1 month after dose 2 (data cutoff date 13 March 2021). The medical literature consisted of case reports and case series of patients who developed thyroiditis following vaccination with various COVID-19 vaccines, including BNT162b2. Observed to expected analyses were conducted and ratios were below 1 for all age groups, doses and gender strata. Based on the totality of...
Table 34. Evaluation of Closed Signals During the Reporting Interval

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<td>Cerebrovascular accident (CVA)/Stroke</td>
<td>This signal was identified following a request from a competent authority (Australia, TGA) for an analysis on 27 January 2022. The Pfizer safety database search using a search strategy covering ischemic and haemorrhagic strokes, yielded 8934 cases through 18 December 2021. There were 4719 reports of females and 4024 of males, when sex was provided and the mean and median ages were 66.9 and 70 years of age, respectively. Cases were reviewed by age group. As would be expected based on the known epidemiology of stroke, the number of reported cases was highest in the oldest individuals and proportionally decreased with age; the number of ischemic strokes was greater than haemorrhagic strokes. Most of the reports described known risk factors for stroke and many of the cases in younger individuals were inconsistent with actual strokes upon individual case review. Of the 10 relevant studies in the medical literature, two studies (Shimazawa R et al [case reports] and Hippisley-Cox J et al) reported a correlative association between BNT162b2 vaccine and ischemic or haemorrhagic stroke. The study by Shimazawa R et al was based on 10 post-vaccination fatalities in Japan. Of these 10 cases, 4 females died of ICH. Insufficient details precluded further assessment. The study by Hippisley-Cox et al reported an increased risk of ischaemic stroke after a first dose of BNT162b2 but contextualized that this risk is far greater with COVID-19, emphasizing the importance of vaccination. Seven studies (Jabagi MJ et al, Simpson CR et al, Carl L et al, Barba N et al, Koh JS et al, Klein NP et al, and Sessa M et al) did not support an association between BNT162b2 vaccine and haemorrhagic or ischemic stroke. In the remaining publication by Patone M et al, an increased risk of haemorrhagic stroke after BNT162b2 vaccination was reported in a study in England but was not replicated in a Scottish study that was somewhat smaller. Overall, the literature data does not support a clear causal association between BNT162b2 and stroke. The medical literature also did not provide a plausible mechanism for how BNT162b2 could increase the risk of haemorrhagic or ischemic strokes. All observed to expected ratios across all age, sex and dose stratifications were below 1 for haemorrhagic and ischemic strokes. Based on the totality of the available information, a causal association between BNT162b2 and hemorrhagic and ischemic stroke could not be concluded, and the signal was closed by the MAH.</td>
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<tr>
<td>Amenorrhea</td>
<td>Amenorrhea was identified as a signal by PRAC on 14 February 2022. A search of the Pfizer safety database through 15 February 2022 yielded 9634 reports which were mostly non-serious and non-medically confirmed. Ages ranged from 11 to 66 years (mean 33.4). Of the cases without confounders and occurring in women younger than 45 years of age, causality assessments using the WHO-UMC criteria were all unlikely or unassessable. In the placebo-controlled portion of the C4591001 study, prior to treatment assignment unblinding, there were 8 events of amenorrhea, with an equal split of 4 events after receipt of placebo vaccination and 4 events after receipt of active vaccination. Participants were followed up for a mean period of 135.8 days following the second dose of blinded study vaccine until unblinding (median 145 days; range 85-171 days). The medical literature consisted of studies obtaining mostly self-reported data. One study of almost 4000 women in the US found COVID-19 vaccination (mostly mRNA vaccines) was associated with a &lt; 1 day change in cycle length for dose 1 and dose 2 compared to pre-vaccine cycles. The Lareb (Netherlands) reported amenorrhea as the most reported category of menstrual abnormalities although changes were small and quickly reversed. A retrospective study in the UK of &gt;1200 women 18 and older did not find an association between COVID-19 vaccination and menstrual changes.</td>
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<td>Based on the totality of the available information, a causal association between BNT162b2 and amenorrhea could not be concluded, and the signal was closed by the MAH. On 13 June 2022, the PRAC requested that an updated analysis of the topic be submitted in the PSUR 4.</td>
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<tr>
<td>Heavy menstrual bleeding</td>
<td>Heavy menstrual bleeding (HMB) was identified as a signal by PRAC on 14 February 2022. A search of the Pfizer safety database through 15 February 2022 yielded 23,659 cases of heavy menstrual bleeding. The majority are non-serious and non-medically confirmed. Of the much smaller subset of serious, medically confirmed reports that provided information about menstrual patterns, 4 were assessed as possibly related to vaccine using the WHO-UMC causality criteria as requested; one of the 4 was 1 of 2 cases that described a rechallenge. In addition, the O/E ratios do not indicate that reported events are higher than expected based on background incidence rates. In the placebo-controlled portion of the C4591001 study, prior to treatment assignment unblinding, there were 6 events of heavy menstrual bleeding; 4 of these events were after receipt of active vaccination and 2 events after receipt of placebo. The event in the C4591031 study occurred during the placebo-controlled portion of the study and was in a participant who received placebo. In the C4591001 study, participants were followed up for a mean period of 137.5 days following the second dose during the placebo-controlled follow-up period until unblinding (median 132 days; range 89 – 176 days). The participant in the C4591031 study was followed up for 96 days from blinded study vaccine (placebo) until unblinding. The medical literature on the topic reveals that menstrual abnormalities in general are very common and there have been correlations between SARS-CoV-2 pandemic stress, anxiety, and depression with menstrual cycle abnormalities. A clear pathophysiological mechanism for heavy menstrual bleeding itself is not understood. A well-designed US study of self-reported menstrual cycle data by Alison Edelman et al. did not support a significant effect of vaccination on the number of days of menstrual bleeding. Studies are limited by their retrospective nature and self-reporting. While most menstruating women do not report menstrual changes associated with COVID-19 vaccination, it seems that variables such as age, BMI, changes in cycle over the previous year and the presence of fibroids and smoking may be playing a role. Based on the totality of the available information, a causal association between BNT162b2 and HMB could not be concluded, and the signal was closed by the MAH. On 13 June 2022, the PRAC responded with a list of questions that the MAH is in the process of preparing for submission by 24 August 2022.</td>
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<tr>
<td>Loss of/altered taste and smell</td>
<td>This was identified as a signal during the reporting period following a request for a competent authority (Australia, TGA) for an analysis of the topic. A search of the safety database through 01 March 2022, yielded 12,140 potentially relevant cases (1% of all AE reports for BNT162b2). There were 17 fatal cases all unrelated to vaccination but related to intercurrent diseases. To enable a focused review of the most informative cases, the MAH applied an exclusion algorithm focusing on serious and medically confirmed cases, excluding most confounding conditions and concomitant medications which could have contributed to the events. The identified 154 were further reviewed. 76 cases had insufficient information to make a thorough medical evaluation. 67 cases had alternative explanations or confounding factors which could have contributed to the events and there were 11 remaining cases, only five out of the 11 cases where judged “possible related”</td>
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### Table 34. Evaluation of Closed Signals During the Reporting Interval

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<td>according to WHO causality assessment. There were no cases with a probable or definite relationship. Review of post-marketing data did not support a causal relationship between vaccination with BNT162b2 and the development of taste and smell disorders. Based on the mid-range background rates from ACCESS, O/E ratios were greater than 1 overall for all ages globally using the 7-day risk window. Using both mid- and high-range ACCESS background rates, the O/E ratios were &gt;1 for certain age groups using both the 7- and 21-day risk windows, suggesting that the number of reported cases may be higher than expected compared to unvaccinated persons. The O/E ratios for these events may be overestimated for a few reasons. First, these pre-COVID (2017-2019) background rates from ACCESS reflect the incidence of anosmia and/or ageusia that was treated in either an inpatient or outpatient setting. Given that these symptoms may be mild in some cases, these rates might underestimate the overall incidence (and thus expected cases) of these conditions in a general, unvaccinated populations if not all cases are reported to healthcare providers. Second, they may also underestimate the incidence rate during the COVID-era since anosmia and/or ageusia are symptoms of SARS-COV-2 infection. For example, the reported incidence rates for anosmia/ageusia were 1.4-2.3 times higher in 2020 than during 2017-2019 in both of the ACCESS sources used for background estimates. In 2020, the ES_SIDIAP_PC database rate was 35.23/100,000 persons per year, and in the ES-FISABIO database rate was 67.5/100,000 persons per year. Third, observed cases may include cases of anosmia or ageusia due to recent or current SARS-COV-2 infection that was not documented or diagnosed during the risk window period. Fourth, due to the unprecedented attention to COVID-19 vaccination and outreach to encourage AE reporting, the long-held assumption that reported cases are an underestimation of actual cases may be incorrect. Finally, the ACCESS rates for anosmia and ageusia were defined with ICD codes that captured anosmia, parosmia, and parageusia, while the observed case definition included PTs for additional related conditions. In the pivotal clinical trial C4591001, during the placebo-controlled follow up period from Dose 1 to 1 month after Dose 2 of BNT162b2, 17 events of interest were reported in the vaccine group (N=21926) and 10 in the placebo group (N=21921) in participants ≥16 years of age. None of the 5 relevant PTs were reported by 12-15 year old from Dose 1 to 1 month post-Dose 2 in C4591001. None of the 5 PTs were reported by 5 to &lt;12 year old from Dose 1 to 1 month Dose 2 in C4591007. There was a limited amount of medical literature on this topic and it was not supportive of a known relationship between vaccination and loss of taste or smell. Based on the totality of the available information, a causal association between BNT162b2 and anosmia and ageusia could not be concluded, and the signal was closed by the MAH.</td>
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**Important Risks**

| Myocarditis and Pericarditis | During the reporting period, myocarditis and pericarditis, which have been considered important identified risks in the US-PVP and EU-PV, were moved from important potential risks to important identified risks in the company core list of safety concerns. After the DLP of this PSUR, they were also added as adverse reactions to the company CDS v. 14.0 dated 26 July 2022 (Section 4.8, Appendix A and Appendix B). The changes to the core list of safety concerns and CDS were made based on the summation of data that has accumulated in the surveillance of this issue, including the published incidence and reporting rates from multiple sources with consistent findings. |
Table 34. Evaluation of Closed Signals During the Reporting Interval

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<td>Irritability</td>
<td>During the reporting period, placebo-controlled safety data from Clinical Trial C4591007 was unblinded for submission to regulatory authorities to support authorisation of vaccination in individuals 6 months to &lt; 5 years of age. Irritability was the most frequently reported systemic event reported within 7 days after each of the 3 doses of BNT162b2 (3 µg) for the 6 months to &lt; 2 years age group. Irritability was reported by 51.2%, 47.4% and 43.6% of participants in the BNT162b2 group and 47.2%, 40.7% and 37.6% in the placebo group, after dose 1, 2 and 3, respectively. Based on these data, irritability was determined to be an adverse reaction for the age group 6 months to &lt; 2 years.</td>
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16.2.2. Signal Evaluation Plan for Ongoing Signals

The table below provides the evaluation plan for signals in which the evaluation was still ongoing (i.e., not closed) at the cut-off date of this PSUR.

Table 35. Signal Evaluation Plan for Ongoing Signals

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<th>Signal</th>
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<tr>
<td>Hearing loss</td>
<td>Following enquiry from a competent authority (EMA PRAC and Health Canada) this signal was reopened during the reporting period and is under evaluation at the cut-off date of this PSUR (18 June 2022). The requested cumulative review is in Appendix 6A.3.</td>
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16.3. Evaluation of Risks and New Information

Evaluation of new information for previously recognised important identified and important potential risks, other risks (not categorised as important), special situations, and special patient populations for BNT162b2 is provided below in Section 16.3.1, Section 16.3.2, Section 16.3.3, Section 16.3.4 and Section 16.3.5, respectively.

As part of the AR of the 2nd PSUR (Procedure number EMEA/H/C/PSUSA/00010898/202112), the following request was received:

2. The MAH is requested to re-assess the need for continuing the follow-up questionnaires anaphylaxis and VAED/VAERD and provide process data (e.g., response rate, extent of additional information collected) separately for cases reporting anaphylaxis and cases reporting VAED/VAERD, if applicable.

Response

Please refer to Appendix 6A.

16.3.1. Evaluation of Important Identified Risks

Evaluation of incremental data for the important identified risks Anaphylaxis, Myocarditis and Pericarditis is provided below.
16.3.1.1. Important Identified Risks – Anaphylaxis

Search criteria - PTs: Anaphylactic reaction; Anaphylactic shock; Anaphylactoid reaction; Anaphylactoid shock. 77

Clinical Trial Data

- Number of cases: 3 (0.45% of 668 cases of the total CT dataset), compared to 2 cases (0.28%) retrieved in the PSUR #2.

The investigator and the Sponsor reported that there was not a reasonable possibility that the events anaphylactic reactions in all cases were related to the blinded study vaccine/BNT162b2, or clinical trial procedure. In 2 cases the anaphylaxis reactions were associated with food allergies and in the remaining case anaphylaxis reaction was attributed to another product (etoricoxib).

Post-Authorisation Data

- Number of cases: 1037 (0.20% of 507,683 cases, the total PM dataset), compared to 3507 cases (0.53%) retrieved in the PSUR #2.
- MC cases (690), NMC cases (347).
- Country of incidence (top 10): Japan (184), Germany (158), Australia (113), UK (105), US (59), Poland (48), France (47), New Zealand (41), Philippines (26), and Sweden (24); the remaining 232 cases were distributed among 34 countries.
- Subjects’ gender: female (768), male (219) and unknown (50).
- Subjects’ age in years: n = 949, range: 5 - 99, mean: 40.2, median: 40.0.
- Medical history (n = 422): the most frequently (≥ 10 occurrences) reported medical conditions Asthma (90), Food allergy (83), Drug hypersensitivity (66), Hypersensitivity (55), Hypertension (42), Seasonal allergy (38), Anaphylactic reaction (30), COVID-19 (21), Mite allergy (20), Allergy to arthropod sting (19), Allergy to animal (14), Dermatitis contact (14), Contrast media allergy (12), Mast cell activation syndrome (12), Multiple allergies (12), Diabetes mellitus (11), Urticaria (11), Allergy to chemicals (10), Anaphylactic shock (10), Migraine (10), Obesity (10), and Rubber sensitivity (10).
- Co-suspects (n = 21 cases): Relevant co-suspect vaccines/medications reported more than once were: adalimumab, herbal pollen NOS, JNJ 78436735 (2 each).
- Number of relevant events: 1073.
- Relevant event seriousness: serious (1073).

77 According to the search criteria specified for Anaphylaxis in the EU-RMP v 4.0.
- Reported relevant PTs: Anaphylactic reaction (802), Anaphylactic shock (238), Anaphylactoid reaction (33).

- Time to event onset (n = 781), range: <24 hours to 365 days, median: 0 days.
  - <24 hours: 659 events (6 of which had a fatal outcome);
  - 1 day: 46 events;
  - 2-7 days: 36 events;
  - 8-14 days: 14 events;
  - 15-30 days: 8 events;
  - 31-180 days: 17 events;
  - > 181 days: 1 event.

- Duration of relevant events (n = 249 out of 1037 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 246 days, median 0 days.
  - <24 hours: 152 events;
  - 1 day: 47 events;
  - 2-7 days: 38 events;
  - 8-14 days: 3 events;
  - 15-31 days: 3 events;
  - 32-181 days: 4 events;
  - > 181 days 2 events.

- Relevant event outcome\(^{78}\): fatal (8\(^{79}\)), resolved/resolving (647), resolved with sequelae (23), not resolved (134), unknown (263).

Of the 7 cases reporting relevant events with fatal outcome, 4 cases reported limited information regarding one or more of the following: medical history, concomitant medication, and clinical course of events, precluding a meaningful medical assessment. The remaining 3 cases are described below.

In a non-medically confirmed case, a 96-year-old male subject became “ill” 35 minutes following BNT162b2 vaccination (unknown dose number). The subject was found collapsed in parked car and resuscitation was attempted. According to the coroner, the subject died of anaphylaxis reaction. Ischaemic heart disease and old age were secondary causes.

An 80-year-old female subject fainted while shopping about 39 minutes after dose 1 and, despite resuscitation, died at the hospital. Her primary physician reported that she was treated at the cardiology clinic but had not had an ordered ECG and had stopped taking her

\(^{78}\) Multiple episodes of the same event were reported with a different outcome in some cases hence the sum of the events outcome exceeds the total number of events.

\(^{79}\) Of note, after the DLP, 2 fatal cases were found to be duplicates and one of them was made invalid. Excluding the invalid case, there were 7 cases reporting events with a fatal outcome.
anthypertensive medication. Autopsy results noted that the primary cause of death was atherosclerosis and secondary cause was aortic aneurysm dissection to the pleural cavities.

A 67-year-old male subject with a history of diabetes and idiopathic thrombocytopenic purpura on multiple medications, received dose 3 and less than 30 minutes later presented to the hospital with chest and GI discomfort. His systolic blood pressure was 72 and a diagnosis of anaphylaxis was made. He was given adrenaline with no response and shortly thereafter an ECG showed signs of a myocardial infarction. He was transferred to another hospital for treatment but had a cardiac arrest during angiography. He remained hospitalised until he died due to mesenteric ischemia 12 days post vaccination.

Of the 433 cases reporting medical history/co-suspects, 324 cases reported relevant medical history/risk factors (e.g., asthma, drug hypersensitivity, food allergies, autoimmune disorders, hypersensitivity, prior anaphylactic reactions) and/or co-suspect (e.g., adalimumab, infliximab, influenza vaccine inact SAG 4V, herbal pollen NOS, JNJ 78436735, immunoglobulin human normal), which may have contributed to the anaphylaxis related events.

Analysis by age group

PM: Paediatric (120), Adults (751), Elderly (80) and Unknown (86).

- No significant difference was observed in the reporting proportion of anaphylaxis relevant PTs between paediatric, adult and elderly populations (0.38% in paediatric vs 0.21% in adults vs 0.14% in elderly).

Analysis by presence of comorbidities

Number of subjects with comorbidities: 181 (17.5% of the cases reporting anaphylaxis).

- The reporting proportion of anaphylaxis related events with fatal outcome with comorbid conditions is 1.7% compared to the reporting proportion of 3.3% observed in the individuals without comorbidities. A meaningful comparison is not possible due to the low number of fatal anaphylactic related cases.

Literature Data

During the reporting interval, there were no new significant data received from literature sources.

O/E Analysis

O/E analysis was performed for Anaphylaxis (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

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80 CT and PM pooled data.
Risk Assessment of New Information

Based on the interval data, no new significant safety information was identified pertaining to the risk of anaphylaxis with BNT162b2.

This risk is communicated in the BNT162b2 CDS, Section 4.4, Special warnings and precautions for use, which includes information on appropriate action to be taken, as follows: "As with all injectable vaccines, appropriate medical treatment and supervision must always be readily available in case of a rare anaphylactic event following the administration of the vaccine." This risk is also listed in the CDS Section 4.8, Undesirable effects, Appendix A, Appendix B.

In line with the removal of anaphylaxis from the list of safety concerns in the EU-RMP v. 5.1 submitted on 08 July 2022, the MAH proposes to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period, because anaphylaxis is a known risk of vaccines that is understood by HCPs who administer vaccines and patients and does not considerably impact the benefit/risk profile of the vaccine. Product labeling and standards of medical care during the vaccination procedure provide adequate risk mitigation.

This risk will continue to be monitored through routine pharmacovigilance.

16.3.1.2. Important Identified Risks – Myocarditis and Pericarditis

There were 8533 potentially relevant cases of Myocarditis and Pericarditis: 5423 cases reported myocarditis and 4156 cases reported pericarditis (in 1046 of these 8533 cases, the subjects developed both myocarditis and pericarditis).

For the incremental evaluation of Myocarditis and Pericarditis cases, please refer to Section 16.3.1.2.1 and Section 16.3.1.2.2, respectively.

Literature Data

During the reporting period an unpublished presentation including significant information on myocarditis was reviewed. Please refer to Section 11 Literature for details.

Risk Assessment of New Information

Based on the interval data, no new safety information was identified pertaining to the risk of myocarditis and pericarditis with BNT162b2.

This risk is communicated in the BNT162b2 CDS, Section 4.4, General recommendations, which includes information on appropriate action to be taken, as follows: "Very rare cases of myocarditis and pericarditis have been reported following vaccination with TRADENAME. Typically, the cases have occurred more often in younger men and after the second dose of the vaccine and within 14 days after vaccination. These are generally mild cases and individuals tend to recover within a short time following standard treatment and rest."
Healthcare professionals should be alert to the signs and symptoms of myocarditis and pericarditis in vaccine recipients. This risk will continue to be monitored through routine pharmacovigilance.

16.3.1.2.1. Important Identified Risks – Myocarditis

Search criteria - PTs: Autoimmune myocarditis; Carditis; Chronic myocarditis; Eosinophilic myocarditis; Giant cell myocarditis; Hypersensitivity myocarditis; Immune-mediated myocarditis; Myopericarditis; Myocarditis.

Overall - All Ages

Clinical Trial Data

- Number of cases: 1 case of BNT162b2 (0.15% of 668 cases of the total CT dataset), compared to 2 cases (0.3% of 721 cases of the total CT dataset) retrieved in the PSUR #2.
- Country of incidence: US.
- Subject’s gender: Male (1).
- Subject’s age in years: 43 years.
- Medical history: PT Abstains from alcohol, Abstains from recreational drugs, Anxiety, Attention deficit hyperactivity disorder, Clinical trial participant, Dyspepsia, Gastroesophageal reflux disease, Neuralgia, Non-tobacco user, Postural orthostatic tachycardia syndrome, Prophylaxis, Seasonal allergy, Stress, Tachycardia, Thyroiditis, Vasectomy (1 each).
- Co-suspects: None.
- Number of relevant serious events: 1.
- Reported relevant PTs: Myocarditis (not related to BNT162b2).
- Relevant event outcome: Resolved (1).
- Time to onset of relevant events: 98 days after dose 3.
- Duration of myocarditis was reported as 2 days.

81 Myocarditis and pericarditis are listed in Section 4.8 of the EU-SmPC and have been added as ADRs in Section 4.8, Appendices A and B of the CDS v. 14.0 made effective on 26 July 2022, after the DLP of this PSUR (please refer to Section 4 Changes to Reference Safety Information and Section 14 Late-breaking Information).

82 The SMQ (narrow) Noninfectious myocarditis/pericarditis that became available upon the upversioning to MedDRA v. 25.0 is used as search criteria. Three new PTs (Carditis, Chronic myocarditis and Myopericarditis) are included in the search criteria for myocarditis, compared to the criteria specified for myocarditis in the EU-RMP v 4.0, used in the previous PSUR.
Post-Authorisation Data

- Number of cases: 5422 (1.1% of 507,683 cases of the total PM dataset), compared to 6347 cases (1.0%) retrieved in the PSUR #2.83

- Country/region of incidence (≥10): Germany (1342), UK (1230), Australia (509), France (344), Taiwan, Province Of China (280), Canada (216), Austria (193), Japan (163), Italy (151), Sweden (119), US (118), New Zealand (107), Greece (77), Israel (64), Finland (51), Spain (45), Netherlands (44), Hong Kong (41), Poland (35), Belgium, Denmark (29 each), Switzerland (25), Norway, Portugal (24 each), Malaysia (22), Ireland (21), Czech Republic (16), Brazil (15), Romania (10). The remaining 78 cases were distributed among 25 countries.

- MC (2710), NMC (2712).

- Subjects’ gender: female (1997), male (3307) and unknown (118).

- Subjects’ age in years: n = 4981, range: 6 - 98, mean: 35.3, median: 32.

- Medical history (n = 1699): the most frequently (≥50 occurrences) reported medical conditions included Hypertension (214), Asthma (140), Seasonal allergy (130), Tobacco user (96), Drug hypersensitivity (67), Immunodeficiency (64), Obesity (62), Hypothyroidism, Non-tobacco user (59 each), and Food allergy (58).


- Number of relevant events: 5458.

- Relevant event seriousness: serious (5458).

- Reported relevant PTs: Myocarditis (4639), Myopericarditis (697), Carditis (113), Eosinophilic myocarditis (4), Giant cell myocarditis, Hypersensitivity myocarditis (2 each), and Immune-mediated myocarditis (1).

- Relevant event outcome78: fatal (87), resolved/resolving (1925), resolved with sequelae (160), not resolved (1608), unknown (1682).

- Of the 5422 cases, in 1108 cases (20.4% of the cases reporting myocarditis related events) the events were confounded by subject’ s relevant medical history (1016 cases; e.g., COVID-19, seasonal allergy, tobacco user, drug hypersensitivity, food allergy, myocarditis, mite allergy, autoimmune thyroiditis, cardiac failure, allergy to animal,

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83 During the reporting period of PSUR #2 there were 6347 events of myocarditis: Eosinophilic myocarditis (4), Autoimmune myocarditis, Hypersensitivity myocarditis (3 each), Immune-mediated myocarditis (1).
overweight, alcohol use, cardiac disorder, pericarditis, allergy to metals, breast cancer, chemotherapy, allergy to plants, dust allergy, rheumatoid arthritis, influenza, myopericarditis, systemic lupus erythematosus, mycotic allergy, radiotherapy, allergy to arthropod sting, allergy to chemicals, Epstein Barr virus infection, autoimmune disorder, Lyme disease, rheumatic disorder) and/or relevant co-suspect/concomitant medications (92 cases; e.g., influenza vaccine, isotretinoin, mesalazine, olanzapine, quetiapine, rituximab, cyclophosphamide, epirubicin, hepatitis B vaccine RHBSAG (yeast), minocycline, norepinephrine, sulfasalazine, zuclopenthixol, COVID-19 vaccine mRNA (mRNA 1273), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19), COVID-19 vaccine, pembrolizumab, clozapine, hepatitis A vaccine, influenza vaccine INACT SAG 3V, ipilimumab, JNJ 78436735, nivolumab). Of the 5422 cases, 236 cases involved elderly (age >70 years) subjects and 61% cases involved male subjects.

*Age-stratified data* 84

**Subjects aged less than 5 years**

**Clinical Trial Data**
- Number of cases: none. No cases were retrieved in the PSUR #2.

**Post-Authorisation Data**
- Number of cases: none. No cases were retrieved in the PSUR #2.

**Subjects aged 5 - 11 years**

**Clinical Trial Data**
- Number of cases: none. No cases were retrieved in the PSUR #2.

**Post-Authorisation Data**
- Number of cases: 48 cases (0.01 % of 507,683 cases of the total PM dataset; 0.6 % of the 8375 subjects aged 5-11 years); 10 cases (0.002%) were retrieved in the PSUR #2.
- Country/region of incidence: Australia (15), Canada, Japan (6 each), Italy, Portugal, Spain (3 each), Greece, Taiwan, Province of China (2 each), Austria, Denmark, Finland, France, New Zealand, Philippines, UK, US (1 each).
- Medical history (n = 8): Asthma, Atrioventricular block, Attention deficit hyperactivity disorder, Autoimmune thyroiditis, Cardiac failure, Cerebral palsy, Condition aggravated, Dependence on respirator, Ejection fraction decreased, Hypoxic-ischaemic encephalopathy, Intellectual disability, Motor dysfunction, Myocarditis, Neonatal

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84 Cases where the age was reported as Child (3 cases), Adolescent (23 cases), Adult (103 cases) and Elderly (11 cases) are included in the subgroup of Age Unknown age and in the overall.
asphyxia, Non-tobacco user, Obesity, Respiratory tract infection, Rhinitis allergic, Type 1 diabetes mellitus (1 each).

- Co-suspect vaccine/medications: None.
- Most frequently co-reported PTs (>5 occurrences): Chest pain (28), Dyspnoea, Pyrexia (10 each), Troponin increased (7), Chest discomfort, Electrocardiogram abnormal, Tachycardia (6 each).

**Fatal myocarditis cases in subjects aged 5-11 years (2 cases, medically confirmed)**

A 6-year-old male subject from Portugal:

- Medical history: Autoimmune thyroiditis, Rhinitis allergic, Type 1 diabetes mellitus.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis, Cardio-respiratory arrest, COVID-19.
- Time to onset (myocarditis): 7 days after dose 1.
- Causes of death: Cardio-respiratory arrest; Myocarditis.
- Autopsy: Results awaited at the time of reporting.

An 11-year-old female subject from Japan:

- Co-suspect medications: None.
- PTs with fatal outcome: Blood pressure decreased, Blood pressure immeasurable, Bradycardia, Cardio-respiratory arrest, Cyanosis, Heart rate decreased, Myocarditis, Respiratory failure.
- Time to onset (myocarditis): 1 day after dose 2.
- Causes of death: Blood pressure decreased; Blood pressure immeasurable; Bradycardia; Cardiac failure acute; Cardio-respiratory arrest; Cyanosis; Heart rate decreased; Myocarditis; Respiratory failure.
- Autopsy: Pleural X-ray was performed as autopsy imaging and did not show abnormal findings.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 36 below.
Table 36. Myocarditis in Subjects aged 5 – 11 Years (N=48)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>16</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td>Myocarditis</td>
<td>13</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>Myopericarditis</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Carditis</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>5</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>14</td>
<td>18</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>4</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Time to Onset n=39</td>
<td>&lt; 24 hours</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>1-5 days</td>
<td>7</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>14-20 days</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Event Outcome</td>
<td>Fatal</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>4</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Duration of eventb n=4, median = 1 day</td>
<td>Up to 3 days</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>4-6 days</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>7-25 days</td>
<td>0</td>
<td>2</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Subjects aged 12 - 15 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 366 (0.07% of 507,683 cases of the total PM dataset; 2.7% of the 13,366 subjects aged 12-15 years), compared to 488 cases (0.07% of all cases in the total PM dataset) retrieved in the PSUR #2.
- Country/region of incidence (≥10): Taiwan, Province of China (87), Germany (64), UK (30), Australia, Japan (23 each), Canada (18), France (15), Italy (14), Israel, Malaysia (11 each), Hong Kong (10). The remaining 60 cases were distributed among 20 countries.
- Medical history (n = 72): the most frequently (≥2 occurrence) reported medical conditions included Asthma (7), Attention deficit hyperactivity disorder, Food allergy, Obesity, Seasonal allergy (4 each), Glucose-6-phosphate dehydrogenase deficiency, Hypersensitivity, Migraine, Non-tobacco user, Pericarditis, Rhinitis allergic (3 each), Anxiety, Autism spectrum disorder, Childhood asthma, Cough, Dermatitis atopic, Mite allergy, Pneumonia, Prophylaxis, Tonsillectomy (2 each).


- Co-suspect vaccine/medications: None.

- Most frequently co-reported PTs (>5 occurrences): Chest pain (174), Pyrexia (80), Chest discomfort (64), Dyspnoea (60), Headache (39), Palpitations (36), Pericarditis (34), Fatigue (31), Tachycardia (25), Inappropriate schedule of product administration, Troponin increased (23 each), Vomiting (20), Electrocardiogram ST segment elevation (18), Dizziness, Nausea (17 each), Malaise (16), C-reactive protein increased (15), Myalgia, Troponin I increased (14 each), Cough (13), Asthenia, Off label use, Pain in extremity (11 each), Blood creatine phosphokinase MB increased, Vaccination site pain (10 each), Chills (9), Blood creatine phosphokinase increased, Diarrhoea, Pain (8 each), Decreased appetite, Immunisation, Pericardial effusion, Syncope (7 each), Arthralgia, Electrocardiogram abnormal, Heart rate increased, Multisystem inflammatory syndrome in children, Nasopharyngitis (6 each).

*Fatal myocarditis cases in subjects aged 12-15 years (2 cases, medically confirmed; 1 case non-medically confirmed)*

A 13-year-old male subject from Taiwan, Province of China:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardiac failure, Myocarditis.
- Time to onset (myocarditis): 69 days after dose 2.
- Causes of death: Cardiac failure; Myocarditis.
- Autopsy: Not reported if autopsy was performed.

A 13-year-old male subject from UK:
- Medical history: Abdominal pain, Chest pain.
- Co-suspect medications: None.
- PTs with fatal outcome: Anuria, Asthenia, Cardiac arrest, Compartment syndrome, Enterovirus infection, Malaise, Multi-organ disorder, Multiple organ dysfunction syndrome, Myocarditis, Pulseless electrical activity, Renal failure, Rhinovirus infection, Ventricular tachycardia.
- Time to onset (myocarditis): 5 days after dose 2.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

- Causes of death: Asthenia; Cardiac arrest; Compartment syndrome; Enterovirus infection; Malaise; Multi-organ disorder; Myocarditis; Pulseless electrical activity; Renal failure; Rhinovirus infection; Ventricular tachycardia.
- Autopsy: Not reported if autopsy was performed.

A 13-year-old female subject from Philippines:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): 6 days after dose 1.
- Causes of death: Myocarditis.
- Autopsy: Adverse event following immunisation.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 37 below.

Table 37. Myocarditis in Subjects aged 12 – 15 Years (N=366)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>42</td>
<td>232</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>19</td>
<td>71</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Myocarditis</td>
<td>56</td>
<td>269</td>
</tr>
<tr>
<td></td>
<td>Myopericarditis</td>
<td>5</td>
<td>31</td>
</tr>
<tr>
<td></td>
<td>Hypersensitivity myocarditis</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Carditis</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>32</td>
<td>226</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>29</td>
<td>77</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>16</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>33</td>
<td>186</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>4</td>
<td>43</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>8</td>
<td>25</td>
</tr>
<tr>
<td>Time to Onset n=274</td>
<td>≤ 24 hours</td>
<td>2</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>1-5 days</td>
<td>29</td>
<td>158</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>7</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>14-21 days</td>
<td>2</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>22-31 days</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>&gt;31 days</td>
<td>6</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>14</td>
<td>78</td>
</tr>
<tr>
<td>Event Outcome</td>
<td>Fatal</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>11</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>21</td>
<td>88</td>
</tr>
<tr>
<td></td>
<td>Resolved with sequelae</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>21</td>
<td>104</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>7</td>
<td>63</td>
</tr>
</tbody>
</table>
Table 37. Myocarditis in Subjects aged 12 – 15 Years (N=366)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration of eventb n=39, median=7 days</td>
<td>Up to 3 days 3 8 0</td>
<td>4-6 days 0 6 0</td>
<td>7-25 days 4 11 0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Subjects aged 16 - 17 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 345 (0.07 % of 507,683 cases of the total PM dataset; 4.2 % of the 8313 subjects aged 16-17 years), compared to 470 cases (0.07%) retrieved in the PSUR #2.
- Country of incidence (≥10): Germany (88), Taiwan, Province of China (55), UK (29), Australia (18), Austria (17), France, Poland (14 each), Italy, Japan (12 each), Greece (11). The remaining 75 cases were distributed among 26 countries.
- Subjects' age in years: n = 345, range: 16 -17, mean: 16.6, median: 17.0.
- Medical history (n = 71): the most frequently (>2 occurrence) reported medical conditions included Seasonal allergy (9), Asthma, Obesity (4 each), Food allergy, Mite allergy, Nasopharyngitis, Overweight (3 each).
- Co-suspect vaccine/medications: Clonazepam, infliximab, and levomethadone (1 each).
- Most frequently co-reported PTs (>5 occurrences): Chest pain (147), Pyrexia (72), Dyspnoea (53), Chest discomfort (43), Palpitations (34), Tachycardia (33), Fatigue (29), Headache (27), Inappropriate schedule of product administration (26), Pericarditis (24), Troponin increased (23), Dizziness, Vomiting (18 each), Malaise, Nausea (17 each), Asthenia (12), Chills, Immunisation, Off label use (10 each), Arrhythmia, Blood creatine phosphokinase increased, C-reactive protein increased, Electrocardiogram ST segment elevation, Pain in extremity, Pericardial effusion, Syncope, Troponin I increased (9 each), Cough (8), Myocardial necrosis marker increased, Pain, Troponin T increased (7 each), Angina pectoris, Back pain, Blood creatine phosphokinase MB increased, Diarrhoea, Electrocardiogram abnormal, Lethargy (6 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 38 below.
### Table 38. Myocarditis in Subjects aged 16 – 17 Years (N=345)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>28</td>
<td>202</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>23</td>
<td>90</td>
<td>0</td>
</tr>
<tr>
<td>Relevant PT(^a)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>48</td>
<td>243</td>
<td>2</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>4</td>
<td>47</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>0</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>27</td>
<td>223</td>
<td>0</td>
</tr>
<tr>
<td>No</td>
<td>24</td>
<td>69</td>
<td>2</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>13</td>
<td>39</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>23</td>
<td>154</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>9</td>
<td>65</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>6</td>
<td>34</td>
<td>0</td>
</tr>
</tbody>
</table>

| Time to Onset n=249                    |                     |                   |                      |
| <24 hours                              | 7                   | 18                | 1                    |
| 1-5 days                               | 13                  | 149               | 1                    |
| 6-13 days                              | 6                   | 19                | 0                    |
| 14-21 days                             | 1                   | 7                 | 0                    |
| 22-31 days                             | 2                   | 3                 | 0                    |
| 32-90 days                             | 4                   | 14                | 0                    |
| 91-150 days                            | 1                   | 5                 | 0                    |
| Unknown                                | 18                  | 81                | 0                    |

| Event Outcome                          |                     |                   |                      |
| Fatal                                  | 0                   | 0                 | 0                    |
| Not resolved                           | 11                  | 60                | 0                    |
| Resolved                               | 9                   | 78                | 1                    |
| Resolved with sequelae                 | 1                   | 2                 | 0                    |
| Resolving                              | 11                  | 80                | 0                    |
| Unknown                                | 20                  | 75                | 1                    |

| Duration of event\(^b\) n=30, median= 8 days |                     |                   |                      |
| Up to 3 days                            | 1                   | 7                 | 0                    |
| 4-6 days                                | 0                   | 3                 | 0                    |
| 7-25 days                               | 2                   | 13                | 0                    |
| 26-68 days                              | 0                   | 4                 | 0                    |

---

a. All serious occurrences.

b. For those cases where the event resolved.

### Subjects aged 18 - 24 years

#### Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

#### Post-Authorisation Data

- Number of cases: 968 (0.2 % of 507,683 cases of the total PM dataset, 0.18 % of the 38293 subjects aged 18-24 years), compared to 1187 cases (2.3%) retrieved in the PSUR #2.
• Country of incidence (≥10): Germany (289), UK (114), France (105), Australia (99), Taiwan, Province of China (46), Italy (43), Austria (38), Sweden (32), Japan (25), US (18), Greece, New Zealand (16 each), Israel (14), Canada, Spain (13 each), Finland (11), Denmark (10). The remaining 66 cases were distributed among 18 countries.

• Subjects’ age in years: n = 968, range: 18-24, mean: 21, median: 21.

• Medical history (n = 237): the most frequently (>2 occurrence) reported medical conditions included PT Tobacco user (28), Asthma (23), Seasonal allergy (16), Myocarditis (12), Hypertension, Non-tobacco user (11 each), Immunodeficiency, Obesity (9 each), Attention deficit hyperactivity disorder (8), Hypersensitivity, Nicotine dependence (7 each), Alcohol use, Contraception, Mite allergy (6 each), Acne, Crohn’s disease, Drug hypersensitivity, Migraine, Overweight, Substance use (5 each), Anaemia, Autism spectrum disorder, Epstein-Barr virus infection (4 each), Appendectomy, Chest pain, Food allergy, Hypothyroidism, Oral contraception, Pericarditis, Pharyngitis, Psoriasis, Syncope, Wisdom teeth removal (3 each).

• COVID-19 Medical history (n = 52): COVID-19 (26), Suspected COVID-19 (22), SARS-CoV-2 test positive (3), and Asymptomatic COVID-19 (1).

• Co-suspect vaccine/medications: Drug COVID-19 vaccine, COVID-19 vaccine MRNA (MRNA 1273), COVID-19 vaccine NRVV AD (CHADOXI NCOV-19), insulin, levothyroxine, and zuclopenthixol (1 each).

• Most frequently co-reported PTs (>5 occurrences): Chest pain (354), Dyspnoea (168), Pyrexia (134), Pericarditis (116), Fatigue (115), Palpitations (112), Chest discomfort (104), Troponin increased (75), Tachycardia (74), Headache (61), Inappropriate schedule of product administration (59), Off label use (48), Dizziness, Immunisation (46 each), Interchange of vaccine products (38), Asthenia (33), Chills (32), Malaise, Myalgia (29 each), Angina pectoris (28), Pain, Syncope (25 each), Nausea (23), Arrhythmia, Dyspnoea exertional (21 each), Pericardial effusion (20), Influenza like illness (19), Cough, Vomiting (18 each), Pain in extremity (17), C-reactive protein increased, Heart rate increased (15 each), Electrocardiogram ST segment elevation, Hyperhidrosis, Lethargy (14 each), Electrocardiogram abnormal (13), Diarrhoea (12), Oropharyngeal pain (11), Arthralgia, Blood creatine phosphokinase increased, COVID-19, Myocardial infarction, Myocardial necrosis marker increased, Troponin I increased, Troponin T increased (10 each), Acute myocardial infarction, Paraesthesia (9 each), Abdominal pain, Abdominal pain upper, Back pain, Cardiac failure, Drug ineffective, Feeling abnormal, Inflammation, Sinus tachycardia (8 each), Hypertension, Hypoesthesia, Limb discomfort, Lymphadenopathy, Night sweats, Pulmonary embolism, Vaccination site pain, Ventricular hypokinesia (7 each), Costochondritis, Feeling hot, Incorrect route of product administration, Insomnia, Left ventricular dysfunction, Loss of consciousness, Somnolence (6 each).

**Fatal myocarditis cases in subjects aged 18-24 years (4 cases, medically confirmed)**

A 23-year-old male subject from Estonia:

- Medical history: Non-tobacco user.
- Co-suspect medications: None.
PTs with fatal outcome: Circulatory collapse, Endocarditis, Myocarditis, Sudden cardiac death.
- Time to onset (myocarditis): unknown days after dose 2.
- Causes of death: Circulatory collapse; Endocarditis; Myocarditis; Sudden cardiac death.
- Autopsy: Autopsy was performed, results were not provided at the time of reporting.

A 20-year-old male subject from Taiwan, Province of China:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): 20 days after dose 1.
- Cause of death: Myocarditis.
- Autopsy: Autopsy results showed cause of death as myocarditis.

A 19-year-old male subject from Japan:
- Co-suspect medications: None.
- PTs with fatal outcome: Arrhythmia, Hernia, Hypoxia, Loss of consciousness, Myocardial necrosis, Myocardial necrosis marker increased, Myocarditis, Sudden death, Ventricular hypokinesia.
- Time to onset (myocarditis): 3 days after dose 3.
- Causes of death: Arrhythmia; Hernia; Hypoxia; Loss of consciousness; Myocardial necrosis; Myocardial necrosis marker increased; Myocarditis; Sudden death; Ventricular hypokinesia.
- Autopsy: The autopsy revealed extensive necrosis of the left ventricular myocardium (myocardial necrosis); myocarditis/fulminant myocarditis.

A 23-year-old male subject from Germany:
- Medical history: Hypertension, Obesity.
- Co-suspect medications: None.
- PTs with fatal outcome: Death, Myocarditis.
- Time to onset (myocarditis): 16 days after dose 3.
- Cause of death: Myocarditis.
- Autopsy: Information not available.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 39 below.
Table 39. Myocarditis in Subjects aged 18 – 24 Years (N=968)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>110</td>
<td>471</td>
<td>3</td>
</tr>
<tr>
<td>No</td>
<td>97</td>
<td>283</td>
<td>4</td>
</tr>
<tr>
<td>Relevant PT&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>179</td>
<td>600</td>
<td>7</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>23</td>
<td>147</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>5</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/extended</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>108</td>
<td>484</td>
<td>5</td>
</tr>
<tr>
<td>No</td>
<td>99</td>
<td>271</td>
<td>2</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>48</td>
<td>145</td>
<td>3</td>
</tr>
<tr>
<td>Dose 2</td>
<td>84</td>
<td>316</td>
<td>1</td>
</tr>
<tr>
<td>Dose 3</td>
<td>55</td>
<td>235</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>22</td>
<td>60</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset n= 724</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;24 hours</td>
<td>18</td>
<td>39</td>
<td>1</td>
</tr>
<tr>
<td>1-5 days</td>
<td>73</td>
<td>362</td>
<td>2</td>
</tr>
<tr>
<td>6-13 days</td>
<td>15</td>
<td>57</td>
<td>1</td>
</tr>
<tr>
<td>14-21 days</td>
<td>9</td>
<td>43</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>4</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>6</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>61-220 days</td>
<td>13</td>
<td>34</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>69</td>
<td>179</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>0</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>66</td>
<td>234</td>
<td>1</td>
</tr>
<tr>
<td>Resolved</td>
<td>25</td>
<td>139</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>10</td>
<td>27</td>
<td>1</td>
</tr>
<tr>
<td>Resolving</td>
<td>52</td>
<td>195</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>54</td>
<td>162</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event&lt;sup&gt;b&lt;/sup&gt; n= 71, median=7 days</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>4</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>4</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>1</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>26-195 days</td>
<td>2</td>
<td>18</td>
<td>1</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.
Subjects aged 25 - 29 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 519 (0.1% of 507,683 cases of the total PM dataset, 1.2% of the 43,518 subjects aged 25-29 years), compared to 589 cases (0.09%) retrieved in the PSUR #2.
- Country of incidence (≥10): Germany (150), UK (113), Australia (54), France (39), Austria (27), Sweden, Taiwan, Province of China (17 each), Japan (14), Italy (13), New Zealand (11). The remaining 64 cases were distributed among 25 countries.
- Subjects' age in years: n = 519, range: 25-29, mean: 27.1, median: 27.
- Medical history (n = 141): the most frequently (>2 occurrence) reported medical conditions included Seasonal allergy (20), Asthma (14), Tobacco user (13), Food allergy (10), Hypertension, Mite allergy (8 each), Allergy to animal, Chest pain (7 each), Nontobacco user (6), Hypersensitivity, Hypothyroidism (5 each), Anxiety, Depression, Drug hypersensitivity, Migraine, Myocarditis, Steroid therapy (4 each), Autoimmune thyroiditis, Contraception, Gastroesophageal reflux disease, Polycystic ovaries (3 each).
- COVID-19 Medical history (n = 43): COVID-19 (22), Suspected COVID-19 (21), Post-acute COVID-19 syndrome (2), and SARS-CoV-2 test positive (1).
- Co-suspect vaccine/medications (n = 8): COVID-19 vaccine mRNA (MRNA 1273) (3), adalimumab, fluticasone, influenza vaccine, levotheroxine, and methylphenidate (1 each).
- Most frequently co-reported PTs (>5 occurrences): Chest pain (217), Dyspnoea (153), Palpitations (122), Fatigue (110), Tachycardia (98), Pericarditis (88), Chest discomfort, Pyrexia (71 each), Headache (47), Immunisation (42), Dizziness (39), Arrhythmia (31), Myalgia (25), Off label use, Pain in extremity (24 each), Interchange of vaccine products, Troponin increased (23 each), Heart rate increased, Inappropriate schedule of product administration, Malaise, Pain (22 each), Angina pectoris (21), Lymphadenopathy (18), Asthenia, Nausea (16 each), Chills, Pericardial effusion (15 each), Influenza like illness, Paraoesthesia, Syncope, Vaccination site pain (14 each), Arthralgia (13), Lethargy (10), COVID-19, Dyspnoea exertional, Influenza, Migraine, Vomiting (9 each), Back pain, Cardiac flutter, Heart rate irregular, Tremor (8 each), Abdominal pain upper, Blood pressure increased, Cardiac disorder, Diarrhoea, Extrasystoles, Hyperhidrosis, Peripheral swelling (7 each), Abdominal pain, Cough, Electrocardiogram ST segment elevation, Feeling abnormal, Hypertension, Inflammation, Myocardial infarction, Rash, Sleep disorder (6 each).

Fatal myocarditis cases in subjects aged 25-29 years (2 cases, medically confirmed; 3 case non-medically confirmed)

A 29-year-old male subject from Japan:

- Medical history: Hepatic steatosis.
- Co-suspect medications: COVID-19 vaccine mRNA (MRNA 1273).
- PTs with fatal outcome: Arrhythmia, Myocarditis.
- Time to onset (myocarditis): unknown days after dose 2.
- Causes of death: Arrhythmia; Myocarditis.
- Autopsy: Autopsy revealed arrhythmia

A 27-year-old male subject from Brazil:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use, Chest pain.
- Time to onset (myocarditis): 10 days after dose 3.
- Causes of death: Myocarditis.
- Autopsy: Not reported if autopsy was performed

A 26-year-old male subject from US:

- Medical history: Aneurysm, Surgery, Vein of Galen aneurysmal malformation.
- Co-suspect medications: Influenza vaccine.
- PTs with fatal outcome: Myocarditis, Arrhythmia, Inflammation, Left ventricular dysfunction.
- Time to onset (myocarditis): 4 days after dose 3.
- Causes of death: Arrhythmia; Inflammation; Left ventricular dysfunction; Myocarditis.
- Autopsy: Autopsy results showed myocarditis.

A 26-year-old female subject from Germany:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use.
- Time to onset (myocarditis): unknown days after dose 2.
- Causes of death: Myocarditis.
- Autopsy: Autopsy results showed myocarditis.

A 27-year-old female subject from Germany:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis, Interchange of vaccine products, Off label use.
- Time to onset (myocarditis): unknown days after dose 2.
- Causes of death: Myocarditis.
- Autopsy: Autopsy results showed myocarditis.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 40 below.
Table 40. Myocarditis in Subjects aged 25 – 29 Years (N=519)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>69</td>
<td>169</td>
<td>0</td>
</tr>
<tr>
<td>No</td>
<td>110</td>
<td>168</td>
<td>3</td>
</tr>
<tr>
<td>Relevant PTs^</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>150</td>
<td>285</td>
<td>3</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>27</td>
<td>48</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>4</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/longed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>58</td>
<td>171</td>
<td>1</td>
</tr>
<tr>
<td>No</td>
<td>121</td>
<td>166</td>
<td>2</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>62</td>
<td>93</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>43</td>
<td>108</td>
<td>1</td>
</tr>
<tr>
<td>Dose 3</td>
<td>52</td>
<td>103</td>
<td>1</td>
</tr>
<tr>
<td>Dose 4</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>22</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>Time to Onset ( n = 335 )</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(&lt; 24 ) hours</td>
<td>16</td>
<td>23</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>49</td>
<td>131</td>
<td>0</td>
</tr>
<tr>
<td>6-13 days</td>
<td>12</td>
<td>27</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>15</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>3</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>5</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>61-366 days</td>
<td>3</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>78</td>
<td>106</td>
<td>3</td>
</tr>
<tr>
<td>Event Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>70</td>
<td>108</td>
<td>0</td>
</tr>
<tr>
<td>Resolved</td>
<td>21</td>
<td>41</td>
<td>0</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>9</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>27</td>
<td>82</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>52</td>
<td>92</td>
<td>2</td>
</tr>
<tr>
<td>Duration of event^b (n = 34, median=27)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Up to 3 days</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>1</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>4</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>26-259 days</td>
<td>10</td>
<td>9</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.  
b. For those cases where the event resolved.

Subjects aged 30 - 39 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 983 (0.2 % of 507,683 cases of the total PM dataset, 1.0 % of the 97870 subjects aged 30-39), compared to 995 cases (0.15%) retrieved in the PSUR #2.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

- Country of incidence (≥10): UK (310), Germany (247), Australia (114), France (51),
  Austria (35), Taiwan, Province of China (33), New Zealand, Sweden (18 each), Italy, US
  (17 each), Finland (13), Canada, Greece (12 each), Japan (11), and Belgium (10). The
  remaining 65 cases were distributed among 23 countries.

- Subjects' age in years: n = 983, range: 30-39, mean: 34.3, median: 34.

- Medical history (n = 290): the most frequently (>2 occurrence) reported medical
  conditions included Asthma (27), Seasonal allergy (26), Hypothyroidism (18), Tobacco
  user (17), Immunodeficiency (14), Drug hypersensitivity (13), Hypertension, Migraine,
  Myocarditis, Non-tobacco user (12 each), Food allergy (11), Breast feeding, Clinical trial
  participant (10 each), Diabetes mellitus, Dyspnoea, Obesity, Pregnancy (9 each),
  Autoimmune thyroiditis, Steroid therapy (6 each), Alcohol use, Colitis ulcerative, Dust
  allergy, Fibromyalgia, Histamine intolerance, Hyperhidrosis, Malaise, Pain, Pericarditis
  (5 each), Chest pain, Coeliac disease, Depression, Headache, Lymphadenopathy, Mast
  cell activation syndrome, Mite allergy, Pneumonia, Polycystic ovaries, Post viral fatigue
  syndrome (4 each), Allergy to animal, Allergy to metals, Cardiac disorder, Crohn's
  disease, Drug intolerance, Fatigue, Gastrooesophageal reflux disease, Hypersensitivity,
  Hypophosphataemia, Lactose intolerance, Multiple sclerosis, Muscular weakness,
  Mycotic allergy, Myocardial infarction, Nicotine dependence, Osteoporosis, Pancreatic
  failure, Postural orthostatic tachycardia syndrome, Pulmonary embolism, Small fibre
  neuropathy (3 each).

- COVID-19 Medical history (n = 82): COVID-19 (41), Suspected COVID-19 (39), Post-
  acute COVID-19 syndrome, SARS-CoV-2 test positive (1 each).

- Co-suspect vaccine/medications: Drug COVID-19 vaccine mRNA (MRNA 1273) (3),
  Amoxicillin, clozapine, colchicine, COVID-19 VACCINE NRV V AD (CHADOX1
  NCOV-19), ipilimumab, losartan, nivolumab, propranolol (1 each).

- Most frequently co-reported PTs (>5 occurrences): Chest pain (383), Dyspnoea (294),
  Palpitations (284), Fatigue (276), Pericarditis (240), Tachycardia (215), Pyrexia (128),
  Chest discomfort (110), Headache (104), Immunisation (100), Dizziness (92), Off label
  use (78), Inappropriate schedule of product administration (60), Interchange of vaccine
  products (58), Arrhythmia (54), Pain in extremity (51), Heart rate increased (48), Malaise,
  Myalgia (46 each), Pain (44), Asthenia (40), Syncope (39), Paraesthesia (38), COVID-19
  (36), Drug ineffective (35), Angina pectoris, Troponin increased (34 each), Arthralgia,
  Hypoesthesia (33 each), Chills, Nausea (32 each), Hyperhidrosis, Lymphadenopathy,
  Vomiting (25 each), Dyspnoea exertional (24), Cardiac flutter (23), Vaccination site pain
  (22), Cough, Feeling abnormal, Pericardial effusion (21 each), Exercise tolerance
decreased (20), Discomfort, Influenza like illness (19 each), Anxiety, Back pain,
  Hypertension (18 each), Diarrhoea (16), Heavy menstrual bleeding, Insomnia, Neck pain
  (15 each), Burning sensation (14), Hypotension, Loss of personal independence in daily
  activities, Oropharyngeal pain (13 each), Heart rate irregular, Menstruation irregular,
  Presyncope, Product use issue (12 each), Cardiac discomfort, Condition aggravated,
  Extrasystoles, Inflammation, Lethargy, Muscle twitching, Myocardial infarction,
  Pulmonary oedema, Rash (11 each), Cardiac disorder, Cardiomegaly, Disturbance in
  attention, Tinnitus (10 each), Atrial fibrillation, Cardiac failure, Electrocardiogram
  abnormal, Maternal exposure during pregnancy, Migraine, Muscular weakness, Panic
attack, Pruritus, Somnolence, Supraventricular tachycardia, Thrombosis, Tremor (9 each), Abdominal pain upper, Fibrin D dimer increased, Influenza, Limb discomfort, Musculoskeletal stiffness, Night sweats, Pleural effusion, Vision blurred (8 each), Abdominal pain, Amenorrhoea, Body temperature increased, Ejection fraction decreased, Heart rate decreased, Loss of consciousness, Muscle spasms, Sleep disorder, Suspected COVID-19, Ventricular extrasystoles (7 each), Asthma, Blood pressure increased, Cardiac arrest, Cardiovascular disorder, Congestive cardiomyopathy, Eczema, Feeling cold, Gait disturbance, Haemorrhage, Heart rate, Illness, Menstrual disorder, Nasopharyngitis, Pulmonary embolism (6 each).

Fatal myocarditis cases in subjects aged 30–39 years (4 cases, medically confirmed; 1 case non-medically confirmed)

A 36-year-old male subject from Japan:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardio-respiratory arrest, Myocarditis.
- Time to onset (myocarditis): 68 days after unknown dose.
- Causes of death: Cardio-respiratory arrest; Myocarditis.
- Autopsy: Autopsy revealed myocarditis and cardio-respiratory arrest.

A 33-year-old female subject from Germany:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Abdominal pain, Arrhythmia, Cardiac arrest, Chest pain, Circulatory collapse, Myocarditis, Resuscitation.
- Time to onset (myocarditis): 20 days after dose 1.
- Causes of death: Abdominal pain; Arrhythmia; Cardiac arrest; Chest pain; Circulatory collapse; Myocarditis.
- Autopsy: Autopsy information was not reported.

A 34-year-old male subject from UK:

- Medical history: Dyspnoea, Malaise.
- Co-suspect medications: None.
- PTs with fatal outcome: Arrhythmia, Cardiac arrest, Cardiogenic shock, Circulatory collapse, Dyspnoea, Hypertension, Hypoxia, Left ventricular dysfunction, Myocarditis, Pulmonary oedema, Syncope.
- Time to onset (myocarditis): Unknown days after unknown dose.
- Causes of death: Arrhythmia; Cardiac arrest; Cardiogenic shock; Circulatory collapse; Dyspnoea; Hypertension; Hypoxia; Left ventricular dysfunction; Pulmonary oedema; Syncope.
- Autopsy: Autopsy revealed cause of death as myocarditis.
A 36-year-old female subject from UK:

- Medical history: Depressed mood, Familial risk factor, Perinatal depression, Pregnancy, Tobacco user.
- Co-suspect medications: None.
- PTs with fatal outcome: Hypoaesthesia, Menstruation irregular, Myocardial injury, Myocarditis, Myopericarditis, Neck pain, Pain in extremity, Pain in jaw, Paraesthesia, Pleural effusion, Thrombosis, Vaccination site pain.
- Time to onset (myocarditis and myopericarditis): Unknown duration after first dose.
- Autopsy: Autopsy revealed extensive and severe bilateral lung congestion but no evidence of ischemic, hypertensive or valvular heart disease. No evidence of subarachnoid haemorrhage was present. COVID-19 swabs were negative. Histology showed a single focus of myocarditis, with extensive lung congestion suggestive of sudden cardiac death and smoking related changes.

A 38-year-old female subject from Finland:

- Medical history: Cerebral palsy.
- Co-suspect medications: None.
- PTs with fatal outcome: Back pain, Diarrhoea, Dyspepsia, Myocarditis, Pain, Pain in extremity.
- Time to onset (myocarditis): 41 days after dose 3.
- Causes of death: Myocarditis.
- Autopsy: Autopsy revealed cause of death as myocarditis.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 41 below.
### Table 41. Myocarditis in Subjects aged 30 – 39 Years (N=983)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>125</td>
<td>263</td>
<td>6</td>
</tr>
<tr>
<td>No</td>
<td>276</td>
<td>305</td>
<td>8</td>
</tr>
<tr>
<td>Relevant PT&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>363</td>
<td>505</td>
<td>12</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>27</td>
<td>51</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
<td>13</td>
<td>13</td>
<td>2</td>
</tr>
<tr>
<td>Immune-mediated myocarditis</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/extended</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>104</td>
<td>233</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>297</td>
<td>335</td>
<td>12</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>132</td>
<td>178</td>
<td>7</td>
</tr>
<tr>
<td>Dose 2</td>
<td>116</td>
<td>204</td>
<td>2</td>
</tr>
<tr>
<td>Dose 3</td>
<td>111</td>
<td>143</td>
<td>3</td>
</tr>
<tr>
<td>Dose 4</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>42</td>
<td>42</td>
<td>2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤24 hours</td>
<td>23</td>
<td>31</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>78</td>
<td>156</td>
<td>2</td>
</tr>
<tr>
<td>6-13 days</td>
<td>28</td>
<td>70</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>16</td>
<td>31</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>17</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>12</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>61-449 days</td>
<td>12</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>218</td>
<td>210</td>
<td>12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Event Outcome</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatal</td>
<td>4</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>123</td>
<td>209</td>
<td>4</td>
</tr>
<tr>
<td>Resolved</td>
<td>43</td>
<td>75</td>
<td>3</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>11</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>70</td>
<td>98</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>153</td>
<td>171</td>
<td>7</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration of event&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>2</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>2</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>7-25 days</td>
<td>6</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>26-128 days</td>
<td>10</td>
<td>11</td>
<td>0</td>
</tr>
</tbody>
</table>

---

<sup>a</sup> All serious occurrences.

<sup>b</sup> For those cases where the event resolved.

### Subjects aged ≥40 years

**Clinical Trial Data**

- Number of cases: 1 case of BNT162b2 (0.15 % of 668 cases of the total CT dataset); 1 case (0.14%) was retrieved in the PSUR #2.
- Country of incidence: .
- Subject's gender: Male (1).
Subject’s age in years: 43 years.
• Medical history: PT Abstains from alcohol, Abstains from recreational drugs, Anxiety, Attention deficit hyperactivity disorder, Clinical trial participant, Dyspepsia, Gastroesophageal reflux disease, Neuralgia, Non-tobacco user, Postural orthostatic tachycardia syndrome, Prophylaxis, Seasonal allergy, Stress, Tachycardia, Thyroiditis, Vasectomy (1 each).
• COVID-19 Medical history: COVID-19 (1).
• Co-suspects: None.
• Number of relevant serious events: 1.
• Reported relevant PTs: Myocarditis (not related to BNT162b2).
• Relevant event outcome: Resolved (1).
• Time to onset of relevant events: 98 days.
• Duration of myocarditis was reported as 2 days.

Post-Authorisation Data

• Number of cases: 1752 (0.3 % of 507,683 cases of the total PM dataset, 0.7 % of the 236404 subjects ≥ 40 years), compared to 1876 cases (0.3%) retrieved in the PSUR #2.
• Country of incidence (≥10): Germany (472), UK (464), Australia (168), France (116), Austria (66), Japan (58), New Zealand (53), Italy (41), Taiwan, Province of China (39), Canada (38), Sweden (37), Greece, US (25 each), Norway (16), Netherlands (15), Finland (14), Israel (12), Spain (11), Belgium, Denmark (10 each). The remaining 62 cases were distributed among 21 countries.
• Subjects’ age in years: n = 1752, range: 40-98, mean: 55, median: 53.
• Medical history (n = 754): the most frequently (>5 occurrences) reported medical conditions included Hypertension (164), Seasonal allergy (53), Asthma (50), Drug hypersensitivity (35), Immunodeficiency (34), Obesity (32), Hypothyroidism, Tobacco user (30 each), Atrial fibrillation (29), Diabetes mellitus (28), Cardiac failure (26), Dyslipidaemia (25), Food allergy, Non-tobacco user, Type 2 diabetes mellitus (23 each), Hypersensitivity (21), Gastroesophageal reflux disease (18), Anxiety, Depression (17 each), Clinical trial participant (16), Autoimmune thyroiditis, Breast cancer, Chronic obstructive pulmonary disease, Hyperlipidaemia (15 each), Migraine, Myocarditis (14 each), Coronary artery disease (13), Cardiac disorder, Chemotherapy, Ex-tobacco user, Hypercholesterolaemia, Overweight (12 each), Myocardial infarction, Sleep apnoea syndrome, Thyroidectomy, Tobacco abuse (11 each), Allergy to animal, Allergy to metals, Fibromyalgia, Rubber sensitivity (10 each), Appendectomy, Arteriosclerosis, Fatigue, Interchange of vaccine products, Rheumatoid arthritis (9 each), Alcohol use, Menopause, Mite allergy, Osteoporosis, Radiotherapy, Systemic lupus erythematosus (8 each), Blood cholesterol increased, Cardiac ablation, Dyspnoea, Gout, Hysterectomy, Mitral valve incompetence, Nasopharyngitis, Pulmonary embolism, Steroid therapy, Surgery (7 each), Abstains from alcohol, Allergy to arthropod sting, Allergy to plants,
Arrhythmia, Arthritis, Blood cholesterol abnormal, Cerebrovascular accident, Cholecystectomy, Chronic kidney disease, Colitis ulcerative, Hormone replacement therapy, Inflammatory bowel disease, Influenza, Insomnia, Neoplasm, Nicotine dependence, Osteoarthritis, Supraventricular tachycardia, Tachycardia (6 each).


- Co-suspect vaccine/medications: Influenza vaccine (3), pembrolizumab (2), adalimumab, cisplatin, COVID-19 vaccine, COVID-19 vaccine mRNA (MRNA 1273), COVID-19 vaccine NRV\(_V\) AD (CHADOX1 NCOV-19), gabapentin, glyceryl trinitrate, hepatitis A vaccine, ibuprofen, influenza vaccine INACT SAG 3V, paracetamol, risankizumab, rivaroxaban, vinorelbine, vitamins NOS (1 each).

- Most frequently co-reported PTs (>5 occurrences): Chest pain (572), Dyspnoea (509), Fatigue (493), Palpitations (463), Pericarditis (407), Tachycardia (339), Off label use (285), Interchange of vaccine products (266), Immunisation (264), Pyrexia (235), Headache (189), Chest discomfort (180), Dizziness (174), Arrhythmia (139), Asthenia (101), Malaise, Syncope (89 each), Inappropriate schedule of product administration, Pain in extremity (88 each), Nausea (86), Pain (83), Angina pectoris (73), Cardiac failure (71), Chills (67), Myalgia (66), Arthralgia (64), Pericardial effusion (61), Heart rate increased (58), Dyspnoea exertional, Troponin increased (57 each), Atrial fibrillation (53), Hyperhidrosis (52), Myocardial infarction (50), Cough (49), Back pain (45), Hypertension (43), Paraesthesia (42), Vomiting (40), Diarrhoea, Lethargy (39 each), COVID-19, Lymphadenopathy (35 each), Cardiac flutter, Vaccination site pain (33 each), Extrasystoles, Influenza like illness (32 each), Cardiac disorder, Hypoesthesia (30 each), Decreased appetite (28), Drug ineffective, Insomnia (27 each), Thrombosis (26), Abdominal pain upper, Blood pressure increased (25 each), Cardiomyopathy, Condition aggravated, C-reactive protein increased, Exercise tolerance decreased (24 each), Anxiety, Feeling abnormal, Neck pain (23 each), Electrocardiogram abnormal, Somnolence, Vertigo (21 each), Abdominal pain, Acute myocardial infarction, Cardiac discomfort, Inflammation, Pulmonary embolism, Tremor (20 each), Cardiac arrest, Gait disturbance, Muscular weakness, Ventricular extrasystoles (19 each), Hypertension (18), Cerebrovascular accident, Limb discomfort, Rash (17 each), Cardiomegaly, N-terminal prohormone brain natriuretic peptide increased, Peripheral swelling, Swelling, Ventricular tachycardia, Vision blurred (16 each), Breast pain, Bundle branch block left, Dyspepsia, Heart rate irregular, Impaired work ability, Musculoskeletal chest pain, Pneumonia, Presyncope (15 each), Axillary pain, Congestive cardiomyopathy, Coronary artery disease, Discomfort, Feeling hot, Influenza, Oedema peripheral, Pulmonary oedema, Tinnitus, Troponin T increased (14 each), Disturbance in attention, Ejec- tion fraction decreased, Loss of personal independence in daily activities, Oedema, Pain in jaw, Sinus tachycardia (13 each), Acute coronary syndrome, Blood creatine phosphokinase increased, Feeling cold, Illness, Left ventricular dysfunction, Oropharyngeal pain, Performance status decreased, Suspected COVID-19, Weight

\(^{25}\) More than 1 COVID-19 medical history is reported in some cases.
COVID-19 mRNA vaccine (nucleoside modified)  Reporting Period
Periodic Safety Update Report (PSUR) 3  19 December 2021 through 18 June 2022

decreased (12 each), Atrial flutter, Atrioventricular block, Migraine, Musculoskeletal pain, Pleural effusion (11 each), Cardiogenic shock, Fibrin D dimer increased, Heart rate abnormal, Joint swelling, Memory impairment, Supraventricular tachycardia, Ventricular hypokinesia (10 each), Bradycardia, Confusional state, Electrocardiogram ST segment elevation, Feeling of body temperature change, Head discomfort, Lymph node pain, Mitral valve incompetence, Muscle spasms, Muscle twitching, Myositis, Orthopnoea, Sleep disorder, Stress, Throat tightness (9 each), Acute kidney injury, Blood pressure decreased, Burning sensation, Cold sweat, Cyanosis, Death, Depression, General physical health deterioration, Heavy menstrual bleeding, Hypokinesia, Left ventricular failure, Nasopharyngitis, Respiratory failure, Transient ischaemic attack, Urticaria, Wheezing (8 each), Abdominal discomfort, Amnesia, Arthritis, Brain natriuretic peptide increased, Bronchospasm, Cardio-respiratory arrest, Disease recurrence, Ear pain, Echocardiogram abnormal, Electrocardiogram ST segment depression, Herpes zoster, Hypersensitivity, Loss of consciousness, Menstrual disorder, Myocardial necrosis marker increased, Pallor, Thrombocytopenia, Visual impairment (7 each), Blood pressure abnormal, Body temperature increased, Bronchitis, Cardiac dysfunction, Cardiac failure acute, Depressed level of consciousness, Dysgeusia, Fall, Heart rate decreased, Mobility decreased, Night sweats, Pleuritic pain, Pruritus, Sepsis, Vaccination failure, Vaccination site swelling, Ventricular arrhythmia (6 each).

**Fatal myocarditis cases in subjects aged >40 Years**

There were 59 cases that reported 59 relevant events with fatal outcome in this age group. Of the 59 cases, 40 cases were medically confirmed and 19 were non-medically confirmed cases. There were 23 female and 36 male subjects. Subjects' ages ranged from 40 years to 96 years. The cases were reported from Japan (17), Germany (12), UK (7), Australia (5), Austria, France, Sweden (3 each), New Zealand, Taiwan, Province of China (2 each), Hong Kong, Italy, Netherlands, Norway, and Switzerland (1 each).

The fatal events in these cases were coded to the PTs Abdominal pain upper, Acute coronary syndrome, Acute myocardial infarction, Amnesia, Aortic dissection, Aortic rupture, Aortitis, Arrhythmia, Arteriosclerosis coronary artery, Arteritis coronary, Arthralgia, Asthenia, Atrial fibrillation, Atrioventricular block complete, Back pain, Bacteraemia, Basal ganglia haemorrhage, Blood creatine phosphokinase increased, Blood creatinine increased, Blood lactic acid, Bradycardia, Brain injury, Cardiac arrest, Cardiac disorder, Cardiac dysfunction, Cardiac failure, Cardiac failure high output, Cardiogenic shock, Cardiomegaly, Cardiomyopathy, Cardio-respiratory arrest, Cerebral haemorrhage, Chest pain, Chronic kidney disease, Circulatory collapse, Colitis, Coma, Coronary artery stenosis, C-reactive protein increased, Cytology abnormal, Death, Dizziness, Dyspnoea, Dyspnoea exertional, Electrocardiogram ST segment depression, Embolism, Encephalitis, Encephalomalacia, Endocarditis, Eosinophilic myocarditis, Fatigue, Haemorrhage, Haemosiderosis, Hepatotoxicity, Hyperhidrosis, Hypersensitivity myocarditis, Immunisation, Infection, Inflammation, Influenza like illness, Interchange of vaccine products, Internal haemorrhage, Intracranial pressure increased, Ischaemic cardiomyopathy, Malaise, Memory impairment, Multiple organ dysfunction syndrome, Myalgia, Myocardial fibrosis, Myocardial infarction, Myocardial necrosis, Myocarditis, Myopericarditis, Myositis, Obstruction, Off label use, Pain in extremity, Palpitations, Pericarditis, Peripheral coldness, pH body fluid, Pneumonia, Pneumonia aspiration, Pulmonary artery thrombosis, Pulmonary embolism, Pulmonary

CONFIDENTIAL
Page 160
hypertension, Pulmonary oedema, Pulseless electrical activity, Pyrexia, Respiration abnormal, Respiratory failure, Right ventricular failure, Sepsis, Spinal cord haemorrhage, Sudden death, Syncope, Tachycardia, Tachypnoea, Thrombocytopenia, Thrombosis, Troponin I, Troponin increased, Vasculitis, Vasculitis necrotising, Ventricular fibrillation, Ventricular hypokinesia, Viral myocarditis, Vomiting (1 each). Only 1 case reported a co-suspect medication (pembrolizumab). The most frequently reported (>1 occurrence) medical histories were coded to the PTs Hypertension (8), Cardiac failure, Diabetes mellitus, Obesity (4 each), Cardiac disorder (3), Cardiac failure chronic, Dyslipidaemia, and Type 2 diabetes mellitus (2 each). The most frequently reported (>2 occurrence) cause of death in these cases were coded to the PTs Myocarditis (47), Cardiac arrest (9), Death (7), Cardiac failure, Pericarditis (5 each), Cardio-respiratory arrest, Chest pain, Dyspnoea, Sudden death (4 each), Pneumonia, Syncope (3 each).

Myocarditis relevant data in this subgroup of subjects are summarised in Table 42 below.

Table 42. Myocarditis in Subjects aged ≥40 Years (N=1752)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>357</td>
<td>374</td>
<td>3</td>
</tr>
<tr>
<td>No</td>
<td>565</td>
<td>437</td>
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<tr>
<td>Relevant PTs*</td>
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</tr>
<tr>
<td>Myocarditis</td>
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<td>19</td>
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<tr>
<td>Myopericarditis</td>
<td>110</td>
<td>81</td>
<td>0</td>
</tr>
<tr>
<td>Carditis</td>
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<td>Eosinophilic myocarditis</td>
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</tr>
<tr>
<td>Hypersensitivity myocarditis</td>
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<td>Hospitalisation required/prolonged</td>
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<td></td>
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<td>Yes</td>
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<td>Relevant suspect dose</td>
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</tr>
<tr>
<td>Dose 1</td>
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</tr>
<tr>
<td>Dose 2</td>
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<td>9</td>
</tr>
<tr>
<td>Dose 3</td>
<td>346</td>
<td>308</td>
<td>6</td>
</tr>
<tr>
<td>Dose 4</td>
<td>8</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>83</td>
<td>77</td>
<td>1</td>
</tr>
</tbody>
</table>

| Time to Onset n=958                  |                     |                   |                      |
| ≤24 hours                            | 60                  | 36                | 0                    |
| 1-5 days                             | 167                 | 166               | 1                    |
| 6-13 days                            | 88                  | 90                | 4                    |
| 14-21 days                           | 58                  | 56                | 1                    |
| 22-31 days                           | 34                  | 26                | 0                    |
| 32-60 days                           | 40                  | 42                | 0                    |
| 61-367 days                          | 45                  | 43                | 1                    |
| Unknown                              | 435                 | 359               | 12                   |
Table 42. Myocarditis in Subjects aged ≥40 Years (N=1752)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Event Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>23</td>
<td>36</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>294</td>
<td>250</td>
<td>4</td>
</tr>
<tr>
<td>Resolved</td>
<td>97</td>
<td>107</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>33</td>
<td>32</td>
<td>2</td>
</tr>
<tr>
<td>Resolving</td>
<td>143</td>
<td>147</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>337</td>
<td>245</td>
<td>12</td>
</tr>
<tr>
<td>Duration of eventb</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n=89, median=34 days</td>
<td>Up to 3 days</td>
<td>2</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>4-6 days</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>7-25 days</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>26-170 days</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>171-822 days</td>
<td>10</td>
<td>5</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved.

Subjects with Unknown Age

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 441 (0.08% of 507,683 cases of the total PM dataset, 0.7% of the 60379 subjects with unknown age), compared to 732 cases (0.11%) retrieved in the PSUR #2.
- Country of incidence (≥10): UK (169), Canada (119), US (36), Germany (32), Australia (18), Israel, Japan (14 each). The remaining 39 cases were distributed among 17 countries.
- Subjects’ age in years: Unknown
- Medical history (n = 125): the most frequently (>2 occurrence) reported medical conditions included Hypertension (16), Asthma (14), Anxiety, Attention deficit hyperactivity disorder, Diabetes mellitus, Drug hypersensitivity, Gastroesophageal reflux disease, Tobacco user (7 each), Immunodeficiency, Steroid therapy (5 each), Depression, Fibromyalgia, Food allergy, Palpitations (4 each), Angina pectoris, Chest pain, Endometriosis, Hypothyroidism, Insomnia, Migraine, Nephrolithiasis, Obstructive sleep apnoea syndrome, Pregnancy, Rheumatoid arthritis (3 each).
- COVID-19 Medical history (n = 37): Suspected COVID-19 (23), COVID-19 (14)
- Co-suspect vaccine/medications: COVID-19 vaccine mRNA (MRNA 1273) (2), COVID-19 vaccine NVRV AD (CHADOX1 NCOV-19), influenza vaccine, and JNJ 78436735 (1 each).
Most frequently co-reported PTs (>5 occurrences): Chest pain (171), Pericarditis (132), Dyspnoea (124), Fatigue (122), Palpitations (113), Tachycardia (74), Immunisation (69), Off label use (55), Pyrexia (54), Interchange of vaccine products (51), Headache (37), Chest discomfort (36), Dizziness (26), Nausea (25), Pain in extremity (19), Arthralgia, Syncope (18 each), Pain (17), Asthenia, Malaise (16 each), Cough (14), Hypoaesthesia (13), Chills, Vomiting (12 each), Heart rate increased, Lymphadenopathy (11 each), Arrhythmia, COVID-19, Myalgia, Paraesthesia (10 each), Cardiac flutter, Diarrhoea, Drug ineffective, Feeling abnormal (9 each), Angina pectoris, Hyperhidrosis, Insomnia, Migraine, Peripheral swelling (8 each), Abdominal pain, Axillary pain, Inappropriate schedule of product administration, Troponin increased (7 each), Back pain, Decreased appetite, Heavy menstrual bleeding, Hypertension, Influenza like illness, Muscular weakness, Pericardial effusion, Pleuritic pain, Tremor (6 each).

Fatal myocarditis cases in subjects of unknown age (6 cases, medically confirmed; 2 cases non-medically confirmed)

A male subject from US:

- Medical history: Attention deficit hyperactivity disorder.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardiac arrest, Myocardial injury, Myocarditis, Toxic cardiomyopathy.
- Time to onset (myocarditis): Unknown days after dose 2.
- Causes of death: Cardiac arrest; Myocardial injury; Myocarditis; Toxic cardiomyopathy.
- Autopsy: Autopsy revealed stress cardiomyopathy.

A male subject from US:

- Medical history: Obesity.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocardial injury, Myocarditis, Stress cardiomyopathy, Toxic cardiomyopathy.
- Time to onset (myocarditis): Unknown days after dose 2.
- Causes of death: Myocardial injury; Myocarditis; Stress cardiomyopathy.
- Autopsy: The autopsy revealed biventricular dilatation (dilatation ventricular); pulmonary oedema (pulmonary oedema); global myocardial injury (myocardial injury); toxic cardiomyopathy (toxic cardiomyopathy).

A subject of unknown gender from US:

- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): Unknown days after unknown dose.
- Causes of death: Myocarditis.
Autopsy: It was not reported if an autopsy was performed.

A male subject from US:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): Unknown days after unknown dose.
- Causes of death: Myocarditis.
- Autopsy: It was not reported if an autopsy was performed.

A male subject from US:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Myocarditis.
- Time to onset (myocarditis): Unknown days after unknown dose.
- Causes of death: Myocarditis.
- Autopsy: It was not reported if an autopsy was performed.

A subject of unknown gender from Japan:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardio-respiratory arrest, Myocarditis.
- Time to onset (myocarditis): Unknown days after dose 3.
- Causes of death: Cardio-respiratory arrest; Myocarditis.
- Autopsy: It was not reported if an autopsy was performed.

A female subject from US:
- Medical history: None.
- Co-suspect medications: None.
- PTs with fatal outcome: Pneumonitis, Myocarditis.
- Time to onset (myocarditis): Unknown days after dose 2.
- Causes of death: Myocarditis; Pneumonitis.
- Autopsy: It was not reported if an autopsy was performed.

A subject of unknown gender from UK:
- Medical history: None
- Co-suspect medications: None
- PTs with fatal outcome: Immunisation, Myocarditis, Colitis, Liver injury.
- Time to onset (myocarditis): Unknown days after unknown dose.
- Causes of death: Colitis; Liver injury; Myocarditis.
Autopsy: It was not reported if an autopsy was performed.

Myocarditis relevant data in this subgroup of subjects are summarised in Table 43 below.

**Table 43. Myocarditis in Subjects of Unknown Age (N=441)**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>60</td>
<td>121</td>
<td>30</td>
</tr>
<tr>
<td>No</td>
<td>97</td>
<td>92</td>
<td>41</td>
</tr>
<tr>
<td>Relevant PT&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myocarditis</td>
<td>132</td>
<td>160</td>
<td>62</td>
</tr>
<tr>
<td>Myopericarditis</td>
<td>28</td>
<td>51</td>
<td>7</td>
</tr>
<tr>
<td>Carditis</td>
<td>0</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>37</td>
<td>84</td>
<td>8</td>
</tr>
<tr>
<td>No</td>
<td>120</td>
<td>129</td>
<td>63</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>37</td>
<td>49</td>
<td>14</td>
</tr>
<tr>
<td>Dose 2</td>
<td>53</td>
<td>92</td>
<td>11</td>
</tr>
<tr>
<td>Dose 3</td>
<td>54</td>
<td>44</td>
<td>22</td>
</tr>
<tr>
<td>Dose 4</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>13</td>
<td>27</td>
<td>24</td>
</tr>
</tbody>
</table>

| Time to Onset n=133                      |                     |                   |                      |
| ≤24 hours                                | 9                   | 10                | 0                    |
| 1-5 days                                 | 18                  | 44                | 3                    |
| 6-13 days                                | 3                   | 16                | 2                    |
| 14-21 days                               | 5                   | 7                 | 1                    |
| 22-31 days                               | 1                   | 3                 | 1                    |
| 32-149 days                              | 4                   | 5                 | 1                    |
| Unknown                                  | 120                 | 134               | 64                   |

| Event Outcome                            |                     |                   |                      |
| Fatal                                    | 1                   | 4                 | 3                    |
| Not resolved                             | 47                  | 55                | 8                    |
| Resolved                                 | 26                  | 55                | 8                    |
| Resolved with sequelae                   | 2                   | 1                 | 0                    |
| Resolving                                | 5                   | 16                | 3                    |
| Unknown                                  | 79                  | 87                | 50                   |

| Duration of event b n=7, median=3 days   |                     |                   |                      |
| Up to 3 days                             | 0                   | 3                 | 0                    |
| 4-6 days                                 | 1                   | 0                 | 0                    |
| 7-146 days                               | 1                   | 1                 | 1                    |

a. All serious occurrences.

b. For those cases where the event resolved.

**Subjects with booster dose**

**Clinical Trial Data**

- Number of cases: 1 case (0.15% of 668 cases of the total CT dataset, 0.2% of the 490 subjects who received a booster dose), compared to 2 cases (0.3%) in the PSUR #2.
The case involved a 43-year-old male participant, who received homologous booster dose. Please see above the "Overall-All Ages" subsection for complete details.

Post-Authorisation Data

- Number of cases: 1682 (0.3% of 507,683 cases of the total PM dataset, 1.4% of the 117750 subjects who received a booster dose), compared to 381 cases (0.06%) in the PSUR #2.

- Country/region of incidence (≥10): UK (617), Germany (422), France (113), Austria (72), Italy (53), Japan (48), Israel (44), New Zealand (41), US (32), Greece, Sweden (21 each), Finland (19), Netherlands, Taiwan, Province of China (17 Each), Denmark (16), Australia, Hong Kong (15 each), Switzerland (13), Spain (12), Ireland (10); the remaining 64 cases were distributed among 19 countries.

- MC (702), NMC (980).

- Subjects' gender: female (656), male (988), and unknown (38).


- Medical history (n = 633): the medical conditions reported (≥4 occurrence) included Hypertension (87), Asthma (46), Immunodeficiency (34), Tobacco user (31), Seasonal allergy (30), Hypothyroidism (28), Clinical trial participant (22), Myocarditis (21), Diabetes mellitus (20), Atrial fibrillation, Non-tobacco user (19), Obesity (18), Depression, Migraine (17 each), Steroid therapy (14), Food allergy (13), Anxiety, Dyslipidaemia (12 each), Drug hypersensitivity, Gastroesophageal reflux disease, Interchange of vaccine products (11 each), Chest pain, Mite allergy, Overweight (10 each), Rheumatoid arthritis, Type 2 diabetes mellitus (9 each), Alcohol use, Chronic obstructive pulmonary disease, Dyspnoea, Fibromyalgia, Hyperlipidaemia, Myocardial infarction (8 each), Autoimmune thyroiditis, Cardiac disorder, Coronary artery disease, Ex-tobacco user, Nasopharyngitis (7 each), Cerebrovascular accident, Colitis ulcerative, Crohn's disease, Hypersensitivity, Inflammatory bowel disease, Insomnia, Mitral valve incompetence, Neoplasm, Nephrolithiasis, Nicotine dependence, Osteoarthritis, Pain, Pericarditis, Pneumonia, Pulmonary embolism, Sleep apnoea syndrome (6 each), Abstains from alcohol, Allergy to animal, Appendicectomy, Arteriosclerosis, Attention deficit hyperactivity disorder, Coeliac disease, Congestive cardiomyopathy, Contraception, Endometriosis, Epstein-Barr virus infection, Fatigue, Gastritis, Gout, Hodgkin's disease, Hormone replacement therapy, Menopause, Myopericarditis, Osteoporosis, Palpitations, Pregnancy, Radiotherapy, Supraventricular tachycardia, Surgery, Urinary tract infection (5 each).


- Co-suspect vaccines (n= 20) reported more than once: Influenza vaccine (5), pembrolizumab (2), amoxicillin, cispaltin, COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19), fluticasone, gabapentin, hepatitis A vaccine, infliximab, influenza vaccine...
INACT SAG 3V, JNJ 78436735, paracetamol, propranolol, vinorelbine, zuclopenthixol (1 each).

- Number of relevant events: 1696.
- Relevant event seriousness: all serious.
- Reported relevant PTs: Myocarditis (1458), Myopericarditis (211), Carditis (25), and Eosinophilic myocarditis (2).
- Relevant event outcome: fatal (39), resolved/resolving (528); resolved with sequelae (29), not resolved (453), unknown (649).
- Most frequently co-reported PTs (>20 occurrence): Chest pain (691), Immunisation (537), Fatigue (494), Pericarditis (467), Dyspnoea (466), Palpitations (443), Off label use (442), Interchange of vaccine products (379), Tachycardia (358), Pyrexia (311), Headache (174), Chest discomfort (163), Dizziness (111), Malaise, Pain (86 each), Pain in extremity (84), Nausea (77), Syncope (73), Arrhythmia (71), Chills, Heart rate increased (70 each), Angina pectoris (66), Myalgia (65), Asthenia (59), Arthralgia (58), Troponin increased (52), Lymphadenopathy (51), Vomiting (45), Dyspnoea exertional (43), Back pain, Pericardial effusion (41 each), Hypertension (38), Diarrhoea, Influenza like illness (37), Atrial fibrillation (36), Cough (35), Cardiac flutter, Hyperhidrosis (34 each), Cardiac failure, Vaccination site pain (30), COVID-19 (27), Oropharyngeal pain (26), C-reactive protein increased, Neck pain (24 each), Insomnia (23), Axillary pain, Hypoesthesia, Paresthesia (22 each), Cardiac disorder, Myocardial infarction (21 each), Blood pressure increased, Extrasystoles (20 each).

The number of myocarditis cases occurred after a booster dose in each age group is reported in Table 44 below by gender.

### Table 44. Myocarditis in Subjects who Received a Booster dose

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Heterologous Booster dose No. of Cases</th>
<th>Homologous Booster dose No. of Cases</th>
<th>Unknown dose No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F  M  U</td>
<td>F  M  U</td>
<td>F  M  U</td>
</tr>
<tr>
<td>Age group</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 to 17 years</td>
<td>1  3  0</td>
<td>6  42  0</td>
<td>7  69  1</td>
</tr>
<tr>
<td>18 to 24 years</td>
<td>4  30  0</td>
<td>31  109  0</td>
<td>22  108  0</td>
</tr>
<tr>
<td>25 to 29 years</td>
<td>7  10  0</td>
<td>29  55  0</td>
<td>17  42  1</td>
</tr>
<tr>
<td>30 to 39 years</td>
<td>25  24  0</td>
<td>60  65  3</td>
<td>28  61  0</td>
</tr>
<tr>
<td>40 years and older</td>
<td>142 109 5</td>
<td>134 113 2</td>
<td>86 101 0</td>
</tr>
<tr>
<td>Unknown</td>
<td>30  12  7</td>
<td>21  25  8</td>
<td>6  10  11</td>
</tr>
<tr>
<td>TOTAL</td>
<td>209 188 12</td>
<td>281 409 13</td>
<td>166 391 13</td>
</tr>
</tbody>
</table>

F=female; M=male; U=unknown

During the reporting period there were 1639 cases of medically confirmed myocarditis with a latency 21 days or less in subjects receiving booster dose. All cases were assessed as serious due to hospitalisation and/or medically significant (1002) or due to medically significant (637). In 1314 cases myocarditis occurred within 1 week post vaccine administration. In
most of these cases, the insufficient description of cardiovascular and/or non-cardiovascular medical history and the lack of diagnostic tests confirming the aetiologies of myocarditis preclude a clear causality assessment on an individual case basis.

16.3.1.2.2. Important Identified Risks – Pericarditis

Search criteria - PTs: Autoimmune pericarditis; Pericarditis; Pericarditis adhesive; Pericarditis constrictive; Pleuropericarditis.

Overall - All Ages

Clinical Trial Data

- Number of cases: No cases were retrieved during the current reporting period, compared to 1 case (0.14%) retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 4156 (0.8% of 507,683 cases of the total PM dataset), compared to 5311 cases paid (0.8%) retrieved in the PSUR #2.

- Country of incidence: Australia (1085), UK (903), France (580), Italy (281), Germany (271), Canada (174), New Zealand (111), Netherlands (97), Sweden (71), Japan (68). The remaining 515 cases were distributed among 44 countries.

- MC (2370), NMC (1786).

- Subjects' gender: female (2049), male (2017) and unknown (90).

- Subjects' age in years: n = 3847, range: 4-98 years, mean: 39.8, median: 37.0.

- Medical history: (n = 1292) the most frequently (≥1%) reported relevant medical history included: Hypertension (154), Asthma (109), Pericarditis (95), Seasonal allergy (60), Drug hypersensitivity, Tobacco user (58 each), Immunodeficiency (54), Hypothyroidism (51), Obesity (46), Hypersensitivity, Non-tobacco user (40 each).


- Co-suspects (n=48 cases): frequently (?3 occurrences) reported relevant co-suspect vaccines/medications were COVID-19 vaccine mRNA (mRNA 1273), Influenza vaccine (8 each), COVID-19 vaccine, Influenza vaccine INACT SAG 3V, Influenza vaccine INACT SPLIT 4V (3 each).

- Number of relevant events: 4164.

During the reporting period of PSUR #2 there were 5311 events of pericarditis [Pericarditis (5274), Pleuropericarditis (34), Pericarditis constrictive (11), Pericarditis adhesive (1)].
COVID-19 mRNA vaccine (nucleoside modified)
Periodic Safety Update Report (PSUR) 3
19 December 2021 through 18 June 2022

- Relevant event seriousness: serious (4164).
- Reported relevant PTs: Pericarditis (4133), Pleuropericarditis (26), Pericarditis constrictive (5).
- Relevant event outcome\(^78\): fatal (19), resolved/resolving (1311), resolved with sequelae (82), not resolved (1428), unknown (1325).

Age-stratified data\(^87\)

**Subjects aged less than 5 years**

**Clinical Trial Data**

- Number of cases: none. No cases were retrieved in the PSUR #2.

**Post-Authorisation Data**

- Number of cases: 1; 1 case was retrieved in the PSUR #2.
- Country of incidence: [blank]
- Subject’s age in year: 4.
- Gender: female.
- Medical history: unknown.
- Co-suspects: none.
- Relevant PT: Pericarditis
- Medically Confirmed: yes.
- Hospitalisation required: no
- Time to onset (pericarditis): \(\leq 24\) hours after the 1st dose.
- Co-reported PTs: Chest discomfort, Chest pain, Dyspnoea, Fatigue, Headache, Myalgia, Pyrexia, and Product administered to patient of inappropriate age.

**Subjects aged 5 - 11 years**

**Clinical Trial Data**

- Number of cases: none. No cases were retrieved in the PSUR #2.

---

\(^{78}\) Cases where the age was reported as Child (1 case), Adolescent (9 cases), Adult (62 cases) and Elderly (8 cases) are included in the subgroup of unknown age and in the overall.
Post-Authorisation Data

- Number of cases: 30 (0.006 % of 507,683 cases of the total PM dataset, 0.4 % of the 8375 subjects aged 5-11 years); 4 cases (0.0006%) were retrieved in the PSUR #2.
- Country of incidence: Australia (19), Canada (3), Italy, Japan (2 each), Germany, Israel, New Zealand, UK (1 each).
- Subjects’ age in year: n = 30, range: 5 -11, mean: 9.4, median: 10.0.
- Medical history: Coeliac disease, Kawasaki’s disease, Urinary tract infection viral (1 each).
- COVID-19 Medical history: COVID-19 (1)
- Co-suspects: none.
- Most frequently co-reported PTs (>2 occurrences): Chest pain (24), Dyspnoea (12), Electrocardiogram abnormal (7), Chest discomfort (6), Palpitations (5), Myocarditis (4), Pyrexia (3).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 45.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>7</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Pericarditis</td>
<td>9</td>
<td>21</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>8</td>
<td>19</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>7</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Time to Onset</td>
<td>≤ 24 hours</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>1-5 days</td>
<td>4</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>14-21 days</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>22-31 days</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Event Outcome</td>
<td>Fatal</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Duration of event*</td>
<td>4-6 days</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>11-26 days</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.
Subjects aged 12 - 15 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 118 (0.02 % of 507,683 cases of the total PM dataset, 0.9 % of the 13,366 subjects aged 12-15 years), compared to 215 cases (0.03%) retrieved in the PSUR #2.
- Country of incidence: Australia (31), UK (13), Taiwan, Province of China (11), France, Japan (8 each), Canada, Italy (7 each), Malaysia (6). The remaining 27 cases were distributed among 12 countries.
- Medical history (n = 20): the medical conditions reported more than once included Adenotonsillectomy, Asthma, Glucose-6-phosphate dehydrogenase deficiency, and Hypersensitivity (2 each).
- Co-suspects: none.
- Most frequently co-reported PTs (≥2%): Chest pain (60), Myocarditis (34), Dyspnoea (25), Palpitations (23), Pyrexia (22), Chest discomfort, Fatigue (15 each), Headache, Tachycardia (9 each), Dizziness, Malaise (7 each), Asthenia, Inappropriate schedule of product administration (6 each), Cough, Heart rate increased, Nausea, Pain, Pericardial effusion (5 each), Dyspnoea exertional, Syncope, Vomiting (4 each), Arthralgia, Chills, COVID-19, Electrocardiogram abnormal, Electrocardiogram ST segment elevation, Troponin increased (3 each), Angina pectoris, Back pain, Drug ineffective, Electrocardiogram ambulatory abnormal, Exercise tolerance decreased, Immune system disorder, Lethargy, Musculoskeletal chest pain, Myalgia, Nasopharyngitis, Oropharyngeal pain, Pleural effusion, Pleuritic pain, and Sinus tachycardia (2 each).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 46.
### Table 46. Pericarditis in Subjects aged 12-15 years (N=118)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>15</td>
<td>72</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>9</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>Relevant PT&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>24</td>
<td>91</td>
<td>2</td>
</tr>
<tr>
<td>Pleuropericarditis</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>4</td>
<td>42</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>20</td>
<td>50</td>
<td>0</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>11</td>
<td>38</td>
<td>0</td>
</tr>
<tr>
<td>Dose 2</td>
<td>8</td>
<td>42</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>2</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>3</td>
<td>8</td>
<td>2</td>
</tr>
</tbody>
</table>

| Time to Onset n=118                     |                     |                   |                     |
| < 24 hours                              | 3                   | 5                 | 0                   |
| 1-5 days                                | 10                  | 35                | 0                   |
| 6-13 days                               | 4                   | 9                 | 0                   |
| 14-21 days                              | 1                   | 2                 | 0                   |
| 22-31 days                              | 0                   | 3                 | 0                   |
| 32-60 days                              | 0                   | 2                 | 0                   |
| 61-180 days                             | 0                   | 3                 | 0                   |
| Unknown                                 | 6                   | 33                | 2                   |

| Event Outcome                           |                     |                   |                     |
| Fatal                                   | 0                   | 0                 | 0                   |
| Not resolved                            | 11                  | 25                | 0                   |
| Resolved                                | 4                   | 12                | 1                   |
| Resolved with sequelae                  | 0                   | 1                 | 0                   |
| Resolving                               | 4                   | 29                | 0                   |
| Unknown                                 | 5                   | 25                | 1                   |

| Duration of event<sup>b</sup> n=6, median: 9 |                     |                   |                     |
| 4-6 days                                 | 0                   | 2                 | 0                   |
| 7-10 days                                | 0                   | 2                 | 0                   |
| 11-26 days                               | 1                   | 0                 | 0                   |
| 27-57 days                               | 1                   | 0                 | 0                   |

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

### Subjects aged 16 - 17 years

#### Clinical Trial Data

- **Number of cases:** none. No cases were retrieved in the PSUR #2.

#### Post-Authorisation Data

- **Number of cases:** 106 (0.02% of 507,683 cases of the total PM dataset, 1.3% of the 8313 subjects aged 16-17 years), compared to 174 cases (0.03%) retrieved in the PSUR #2.
- Country of incidence: Australia (25), UK (20), France (15), Italy (11), Germany (6), Taiwan, province of China (5). The remaining 24 cases were distributed among 14 countries.
- Subjects' age in years: $n = 106$, range: 16-17, mean: 16.5, median: 16.0.
- Medical history ($n = 17$): the medical conditions reported more than once included the PTs Asthma, Food allergy, Pericarditis, Seasonal allergy (2 each).
- Co-suspects ($n=2$ cases): COVID-19 vaccine mRNA (MRNA 1273), HPV vaccine VLP RL1 9V (yeast), Influenza vaccine INACT SPLIT 4V, Pneumococcal vaccine polysacch 23V (1 each).
- Most frequently co-reported PTs ($\geq 2\%$): Chest pain (56), Dyspnoea (25), Myocarditis, Pyrexia (22 each), Fatigue, Palpitations (19 each), Tachycardia (15), Chest discomfort (14), Inappropriate schedule of product administration, Nausea, Pain (7 each), Headache, Pericardial effusion, Vomiting (6 each), Electrocardiogram abnormal, Malaise, Myopericarditis (5 each), Chills, Cough, Dizziness, Troponin increased, Abdominal pain upper, Influenza like illness, Lethargy, Pain in extremity (3 each), Asthenia, Back pain, Cellulitis, Cold sweat, C-reactive protein increased, Decreased appetite, Feeling hot, Heart rate irregular, Hyperhidrosis, Interchange of vaccine products, Myalgia, Off label use, Product use issue, Syncope, Vaccination site pain (2 each).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 47.
Table 47. Pericarditis in Subjects aged 16-17 years (N=106)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>20</td>
<td>50</td>
<td>1</td>
</tr>
<tr>
<td>No</td>
<td>15</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>35</td>
<td>70</td>
<td>1</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>7</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>No</td>
<td>28</td>
<td>49</td>
<td>1</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>12</td>
<td>23</td>
<td>1</td>
</tr>
<tr>
<td>Dose 2</td>
<td>20</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>2</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>1</td>
<td>8</td>
<td>0</td>
</tr>
</tbody>
</table>

| Time to Onset n=106                  |                     |                   |                      |
| < 24 hours                           | 1                   | 6                 | 0                    |
| 1-5 days                             | 9                   | 20                | 1                    |
| 6-13 days                            | 3                   | 5                 | 0                    |
| 14-21 days                           | 1                   | 3                 | 0                    |
| 22-31 days                           | 2                   | 3                 | 0                    |
| 32-60 days                           | 5                   | 2                 | 0                    |
| 61-180 days                          | 2                   | 3                 | 0                    |
| 181-375 days                         | 0                   | 1                 | 0                    |
| Unknown                              | 12                  | 27                | 0                    |
| Event Outcome                        |                     |                   |                      |
| Fatal                                | 0                   | 0                 | 0                    |
| Not resolved                         | 11                  | 17                | 1                    |
| Resolved                             | 4                   | 19                | 0                    |
| Resolved with sequelae               | 0                   | 1                 | 0                    |
| Resolving                            | 11                  | 13                | 0                    |
| Unknown                              | 9                   | 20                | 0                    |
| Duration of eventb n=7, median: 10   |                     |                   |                      |
| Up to 3 days                         | 1                   | 2                 | 0                    |
| 7-10 days                            | 0                   | 1                 | 0                    |
| 11-26 days                           | 0                   | 1                 | 0                    |
| 27-57 days                           | 0                   | 1                 | 0                    |
| 58-180 days                          | 0                   | 1                 | 0                    |

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

Subjects aged 18 - 24 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 479 (0.09 % of 507,683 cases of the total PM dataset, 1.3% of the 38,293 subjects aged 18-24 years), compared to 659 cases (0.10%) retrieved in the PSUR #2.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

- Country of incidence: Australia (135), France (79), UK (73), Germany (44), Italy (33), New Zealand (19), Japan (14), Netherlands, Sweden (12 each), Norway (11), US (6). The remaining 41 cases were distributed among 16 countries.
- Medical history (n = 120): the medical conditions reported more than twice included Asthma (21), Immunodeficiency, Pericarditis (6 each), Attention deficit hyperactivity disorder, Mite allergy, Non-tobacco user, Obesity, Overweight, Tobacco user (5 each), Food allergy, Irritable bowel syndrome (4 each), Disease risk factor, Drug hypersensitivity, Endometriosis, Hospitalisation, Hypersensitivity, Hypothyroidism, Migraine, Seasonal allergy, and Substance use (3 each).
- Co-suspects (n= 8 cases): COVID-19 vaccine MRNA (MRNA 1273) (2), COVID-19 vaccine, dupilumab, Influenza vaccine INACT SPLIT 4V, insulin, levothyroxine, salbutamol, zuclopenthixol (1 each).
- Most frequently co-reported PTs (≥2%): Dyspnoea (141), Myocarditis (112), Palpitations (95), Fatigue (83), Chest discomfort (75), Pyrexia (66), Tachycardia (65), Headache, Pericardial effusion (35 each), Dizziness (33), Inappropriate schedule of product administration (26), Electrocardiogram abnormal (24), Pain (23), Immunisation, Myalgia (20 each), Malaise, Off label use (19 each), Interchange of vaccine products, Syncope (18 each), Asthenia (17), Pain in extremity (16), Nausea (15), Angina pectoris, Chills, Vomiting (14 each), Cough (13), C-reactive protein increased, Dyspnoea exertional, Hyperhidrosis, Lethargy (12 each), Anxiety, Sinus tachycardia (10 each), Arthralgia, Heart rate increased (9 each), Back pain, Electrocardiogram ST segment elevation, Parasthesia, Troponin increased (8 each).
- Pericarditis events with fatal outcome (1).

**Fatal pericarditis cases in adult (18-24 years of age) (1 case, medically confirmed)**

A 22-year-old male subject from Israel:

- Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardiac tamponade, Multiple organ dysfunction syndrome, Pericardial effusion, Pericardial mass, Pericardial mesothelioma malignant, Pericarditis, Right ventricular dysfunction, Right ventricular failure.
- Time to onset (pericarditis): 31 days after dose 2.
- Causes of death: all the above events.

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 48.

CONFIDENTIAL
Page 175
### Table 48. Pericarditis in Subjects aged 18-24 years (N=479)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>120</td>
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<tr>
<td>No</td>
<td>72</td>
<td>110</td>
<td>2</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>192</td>
<td>279</td>
<td>8</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>46</td>
<td>96</td>
<td>3</td>
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<tr>
<td>No</td>
<td>146</td>
<td>183</td>
<td>5</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>79</td>
<td>102</td>
<td>4</td>
</tr>
<tr>
<td>Dose 2</td>
<td>52</td>
<td>96</td>
<td>0</td>
</tr>
<tr>
<td>Dose 3</td>
<td>49</td>
<td>60</td>
<td>0</td>
</tr>
<tr>
<td>Dose 4</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>11</td>
<td>19</td>
<td>4</td>
</tr>
<tr>
<td>Time to Onset</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n=479</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 24 hours</td>
<td>27</td>
<td>18</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>66</td>
<td>93</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>22</td>
<td>34</td>
<td>2</td>
</tr>
<tr>
<td>14-21 days</td>
<td>5</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>5</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>1</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>61-180 days</td>
<td>7</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>181-375 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>59</td>
<td>86</td>
<td>4</td>
</tr>
<tr>
<td>Event Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>74</td>
<td>97</td>
<td>2</td>
</tr>
<tr>
<td>Resolved</td>
<td>20</td>
<td>33</td>
<td>2</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>7</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>Resolving</td>
<td>47</td>
<td>65</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>44</td>
<td>76</td>
<td>3</td>
</tr>
<tr>
<td>Duration of eventb n=18, median: 21</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Up to 3 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>11-26 days</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>27-57 days</td>
<td>3</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>58-180 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

### Subjects aged 25 - 29 years

#### Clinical Trial Data

- Number of cases: none; no cases were retrieved in the PSUR #2.
Post-Authorisation Data

- Number of cases: 417 (0.08 % of 507,683 cases of the total PM dataset, 1.0 % of the 43,518 subjects aged 25-29 years), compared to 614 cases (0.09%) retrieved in the PSUR #2.
- Country of incidence: Australia (136), UK (75), France (71), Germany (21), Italy, Netherlands (18 each), New Zealand (16), Sweden (8), Japan, Spain (7 each), Denmark (6), Canada (5). The remaining 29 cases were distributed among 17 countries.
- Subjects’ age in years: n = 417, range: 25 -29, mean: 27.0, median: 27.0.
- Medical history (n = 87): the medical conditions reported more than twice included Asthma (10), Tobacco user (7), Obesity (6), Disease risk factor, Drug hypersensitivity, Non-tobacco user, Pericarditis (4 each), Abstains from alcohol, Contraception, Gastritis, Steroid therapy (3 each).
- Co-suspects (n= 3 cases): COVID-19 vaccine (2), and Methylphenidate (1).
- Most frequently co-reported PTs (>2%): Dyspnoea (137), Palpitations (101), Fatigue (94), Myocarditis (87), Tachycardia (67), Chest discomfort (60), Pyrexia (49), Headache (40), Dizziness, Immunisation (33 each), Nausea (22), Pain (21), Off label use (20), Interchange of vaccine products (19), Malaise, Pericardial effusion (17 each), Syncope (16), Electrocardiogram abnormal, Myalgia (14 each), Angina pectoris (13), Arthralgia, Asthenia, Heart rate increased (12 each), Dyspnoea exertional, Pain in extremity, Paraesthesia, Vaccination site pain (11 each), Lethargy, Lymphadenopathy (10 each), Inappropriate schedule of product administration (9), Troponin increased (8), Cardiac flutter, Diarrhoea, Vomiting (7 each).
- Pericarditis events with fatal outcome (1).

Fatal pericarditis cases in adult (25-29 years of age) (1 case, medically confirmed)

A 29-year-old male subject from Finland:

- Medical history: Hypoventilation, Obesity, Pulmonary fibrosis, Sleep apnoea syndrome, Still's disease
- Co-suspect medications: None
- PTs with fatal outcome: Acute kidney injury, Aortic dissection, Chest pain, Hypoventilation, Inflammatory marker increased, Multiple organ dysfunction syndrome, Pericardial disease, Pericarditis, Respiratory failure, Sepsis.
- Time to onset (pericarditis): 6 days after dose 3.
- Causes of death: Multiple organ dysfunction syndrome; Sepsis; Still's disease.

88 In 4 cases, more than one COVID-19 history are reported.
Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 49.

Table 49. Pericarditis in Subjects aged 25-29 years (N=417)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>102</td>
<td>146</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>71</td>
<td>91</td>
<td>5</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>171</td>
<td>237</td>
<td>7</td>
</tr>
<tr>
<td>Pleuropericarditis</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>41</td>
<td>47</td>
<td>2</td>
</tr>
<tr>
<td>No</td>
<td>132</td>
<td>190</td>
<td>5</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>68</td>
<td>100</td>
<td>4</td>
</tr>
<tr>
<td>Dose 2</td>
<td>40</td>
<td>61</td>
<td>1</td>
</tr>
<tr>
<td>Dose 3</td>
<td>50</td>
<td>58</td>
<td>2</td>
</tr>
<tr>
<td>Dose 4</td>
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<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>15</td>
<td>17</td>
<td>0</td>
</tr>
<tr>
<td>Time to Onset n=418</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 24 hours</td>
<td>15</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>42</td>
<td>76</td>
<td>1</td>
</tr>
<tr>
<td>6-13 days</td>
<td>23</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>14-21 days</td>
<td>15</td>
<td>20</td>
<td>1</td>
</tr>
<tr>
<td>22-31 days</td>
<td>6</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>8</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>61-180 days</td>
<td>11</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>181-375 days</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>52</td>
<td>73</td>
<td>4</td>
</tr>
<tr>
<td>Event Outcome</td>
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<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>78</td>
<td>84</td>
<td>1</td>
</tr>
<tr>
<td>Resolved</td>
<td>12</td>
<td>37</td>
<td>2</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>34</td>
<td>47</td>
<td>1</td>
</tr>
<tr>
<td>Unknown</td>
<td>47</td>
<td>65</td>
<td>3</td>
</tr>
<tr>
<td>Duration of eventb n=13, median: 13</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Up to 3 days</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>11-26 days</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>27-57 days</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>58-180 days</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

Subjects aged 30 - 39 years

Clinical Trial Data

- Number of cases: none. No cases were retrieved in the PSUR #2.
Post-Authorisation Data

- Number of cases: 940 (0.2 % of 507,683 cases of the total PM dataset; 1.0 % of the 97,870 subjects aged 30-39), compared to 1222 cases (0.2%) retrieved in the PSUR #2.

- Country/region of incidence: Australia (356), UK (217), France (114), Germany (46), Italy (39), New Zealand (26), Netherlands (21), Canada (18), Norway (16), Sweden (13), Belgium (9), Greece, US (8 each), Denmark, Japan (6 each), Austria, and Hong Kong (5 each). The remaining 27 cases were distributed among 17 different countries.

- Subjects’ age in years: n = 940, range: 30 -39, mean: 34.3, median: 34.0.

- Medical history (n = 217): the medical conditions reported more than 5 times included the PTs Pericarditis (27), Asthma (20), Drug hypersensitivity (18), Seasonal allergy (13), Mite allergy, Non-tobacco user, Pregnancy, Tobacco user (11 each), Migraine (10), Chest pain, Hypothyroidism (8 each), Anxiety, Clinical trial participant, Immunodeficiency (7 each), Alcohol use, Eczema, Obesity (6 each).

- COVID-19 Medical history (n = 70): COVID-19 (43), Suspected COVID-19 (24), SARS-CoV-2 test positive (3).

- Co-suspect vaccines/medications (n=6): colchicine (2), amoxicillin, interferon Beta-1A, iron isomaltoside 1000, propranolol (1 each).

- Most frequently co-reported PTs (≥2%): Chest pain (547), Dyspnoea (345), Palpitations (260), Fatigue (239), Myocarditis (236), Tachycardia (179), Chest discomfort (125), Pyrexia (113), Headache (93), Dizziness (75), Immunisation (74), Malaise, Pain in extremity (54 each), Nausea, Paraesthesia (48 each), Arthralgia, Pain (46 each), Off label use (45), Myalgia (44), Inappropriate schedule of product administration (40), Heart rate increased, Interchange of vaccine products (39 each), Hypoesthesia, Pericardial effusion (38 each), Asthenia (32), Hyperhidrosis, Syncope (29 each), Electrocardiogram abnormal, Influenza like illness (25 each), Cardiac flutter (24), Arrhythmia (23), Chills, Lethargy (21 each), Feeling abnormal, Vaccination site pain (20 each), Diarrhoea, Exercise tolerance decreased, Vomiting (18 each), Cough (17), Anxiety, Back pain, Dyspnoea exertional, Lymphadenopathy, Neck pain (16 each).

Pericarditis relevant data in this subgroup of subjects are summarised in below Table 50.
## Table 50. Pericarditis in Subjects aged 30-39 years (N=940)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>277</td>
<td>284</td>
<td>5</td>
</tr>
<tr>
<td>No</td>
<td>193</td>
<td>179</td>
<td>2</td>
</tr>
<tr>
<td>Relevant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>470</td>
<td>462</td>
<td>7</td>
</tr>
<tr>
<td>Pleuropericarditis</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>68</td>
<td>96</td>
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<tr>
<td>No</td>
<td>402</td>
<td>367</td>
<td>7</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>223</td>
<td>243</td>
<td>3</td>
</tr>
<tr>
<td>Dose 2</td>
<td>119</td>
<td>111</td>
<td>2</td>
</tr>
<tr>
<td>Dose 3</td>
<td>108</td>
<td>85</td>
<td>2</td>
</tr>
<tr>
<td>Unknown</td>
<td>20</td>
<td>24</td>
<td>0</td>
</tr>
<tr>
<td>Time to Onset n=941</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤ 24 hours</td>
<td>37</td>
<td>35</td>
<td>0</td>
</tr>
<tr>
<td>1-5 days</td>
<td>105</td>
<td>120</td>
<td>0</td>
</tr>
<tr>
<td>6-13 days</td>
<td>47</td>
<td>52</td>
<td>1</td>
</tr>
<tr>
<td>14-21 days</td>
<td>27</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>17</td>
<td>17</td>
<td>0</td>
</tr>
<tr>
<td>32-60 days</td>
<td>17</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td>61-180 days</td>
<td>21</td>
<td>12</td>
<td>1</td>
</tr>
<tr>
<td>181-375 days</td>
<td>5</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>194</td>
<td>187</td>
<td>5</td>
</tr>
<tr>
<td>Event Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>181</td>
<td>189</td>
<td>3</td>
</tr>
<tr>
<td>Resolved</td>
<td>37</td>
<td>63</td>
<td>1</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>6</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>80</td>
<td>65</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>166</td>
<td>144</td>
<td>3</td>
</tr>
</tbody>
</table>

**Duration of eventb n=27, median: 15**

<table>
<thead>
<tr>
<th></th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 3 days</td>
<td>0</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>4-6 days</td>
<td>1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>7-10 days</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>11-26 days</td>
<td>5</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>27-57 days</td>
<td>4</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>58-180 days</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

---

a. All serious occurrences.
b. For those cases where the event resolved or resolved with sequelae.

### Subjects aged >40 years

#### Clinical Trial Data

- Number of cases: none. One (1) case (0.14%) retrieved in the PSUR #2. Please see above the "Overall – All Ages" subsection.
Post-Authorisation Data

- Number of cases: 1756 (0.3 % of 507,683 cases of the total PM dataset, 0.7% of the 236,404 subjects ≥ 40 years), compared to 2059 cases (0.3%) retrieved in the PSUR #2.

- Country of incidence: UK (375), Australia (333), France (288), Italy (169), Germany (137), Canada (61), New Zealand (44), Netherlands (41), Greece (40), Sweden (35), Austria, Norway (28 each), Japan (25), Denmark (20). The remaining 132 cases were distributed among 25 different countries.

- Subjects' age in years: n = 1756, range: 40-98, mean: 54.6, median: 52.0.

- Medical history (n = 738): the medical conditions reported more than 10 times included PTs Hypertension (133), Pericarditis (47), Asthma (43), Immunodeficiency, Seasonal allergy (36 each), Hypothyroidism (34), Hypersensitivity (29), Obesity, Tobacco user, Type 2 diabetes mellitus (28 each), Diabetes mellitus (27), Drug hypersensitivity, Gastroesophageal reflux disease (25 each), Depression (20), Atrial fibrillation (19), Dyslipidaemia, Rheumatoid arthritis (18 each), Anxiety (17), Breast cancer, Dyspnoea, Hypercholesterolaemia, Myocardial ischaemia, Non-tobacco user (16 each), Chronic kidney disease (15), Myocardial infarction (14), Chronic obstructive pulmonary disease, Food allergy, Gastritis (13 each), Autoimmune thyroiditis, Chest pain, Systemic lupus erythematosus (12 each), Overweight, Palpitations, Psoriasis, Steroid therapy (11 each).


- Co-suspect vaccines/medications (n= 24): Influenza vaccine (7), COVID-19 vaccine MRNA (MRNA 1273), Influenza vaccine INACT SAG 3V (3 each), Adalimumab, Apixaban, COVID-19 vaccine NRVV AD (CHADOXI NCOV-19), Etanercept, Gliceryl trinitrate, Influenza vaccine INACT SPLIT 4V, Levetiracetam, Peginterferon alfa-2A, Pemrolizumab, Rivaroxaban, Sotrovimab (1 each).

- Most frequently co-reported PTs (≥2%): Chest pain (787), Dyspnoea (567), Fatigue (455), Myocarditis (396), Palpitations (372), Tachycardia (286), Off label use (272), Interchange of vaccine products (246), Immunisation (243), Pyrexia (223), Chest discomfort (199), Pericardial effusion (167), Headache (157), Dizziness (131), Malaise (87), Pain in extremity (83), Asthenia (82), Arthralgia (74), Nausea (73), Pain (71), Inappropriate schedule of product administration (70), Syncope (69), Myalgia (67), Angina pectoris, Paraesthesia (60 each), Arrhythmia, Cough (57 each), Lymphadenopathy (50), Heart rate increased (49), Chills (46), Hyperhidrosis (44), Electrocardiogram abnormal (43), Back pain (42), Hypertension (41), Lethargy (40), Pleural effusion, Vaccination site pain (39 each), Atrial fibrillation, Diarrhoea (38 each), Dyspnoea exertional (37), Influenza like illness (36), Myocardial infarction (33), Neck pain (32), Cardiac flutter, Condition aggravated (31 each), C-reactive protein increased, Vomiting (29 each).
- Pericarditis events with fatal outcome (17) occurred in subjects aged ≥40 years (n=17, ranged between 41 to 92 years of age).

**Fatal Pericarditis cases in adult (40-50 years of age) (4 cases; 2 cases medically confirmed and 2 non-medically confirmed)**

- **2 cases medically confirmed:**

  A 43-year-old male subject from Japan.
  - Medical history: Diabetes mellitus, Obesity.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Myocarditis, Pericarditis, Sudden death.
  - Time to onset (pericarditis and myocarditis): On the same day of receiving dose 3, the patient died.
  - Cause of death: Myocarditis, Pericarditis, Sudden death.

  A 48-year-old male subject from Hong Kong.
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Brain stem haemorrhage, Pericarditis.
  - Time to onset (pericarditis): 17 days after dose 2.
  - Cause of death: Both the above events.

- **2 cases non-medically confirmed:**

  A 41-year-old male subject from UK.
  - Medical history: Congestive cardiomyopathy, Huntington's disease, Positive airway pressure therapy, Sleep apnoea syndrome, Type 2 diabetes mellitus.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Interchange of vaccine products, Myocarditis, Off label use, Pericarditis, Sudden death.
  - Time to onset (pericarditis and myocarditis): 11.5 hours after dose 3, the patient died.
  - Cause of death: Myocarditis; Pericarditis; Sudden death.

  A 49-year-old male subject from UK.
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Aortic rupture, Back pain, Cardiomegaly, Internal haemorrhage, Myocarditis, Pericarditis, Pyrexia, Syncope, Vomiting.
  - Time to onset (pericarditis): ~50 days after dose 1, the patient died due to the above events.
- Cause of death: Cardiomegaly.

**Fatal Pericarditis cases in adult (51-64 years of age) (7 cases; 5 cases medically confirmed and 2 non-medically confirmed)**

- **5 cases medically confirmed:**

  A 56-year-old male subject from Australia.
  
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Malaise, Pericarditis.
  - Time to onset (pericarditis): On the same day of receiving dose 1.
  - Cause of death: Both the above events.

  A 57-year-old female subject from Austria.
  
  - Medical history: Thyroid cancer.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Pericarditis.
  - Time to onset (pericarditis): Unspecified days after dose 3.
  - Cause of death: Pericarditis.

  A 59-year-old female subject from Australia.
  
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Atrial fibrillation, Atrioventricular block complete, Cardiac arrest, Chest pain, Electrocardiogram ST segment depression, Myocarditis, Pericarditis, Troponin increased.
  - Time to onset (pericarditis): 67 days after dose 3.
  - Cause of death: All the above events.

  A 61-year-old female subject from Japan.
  
  - Medical history: Cerebrovascular accident, Syncope, Thymic carcinoma, Thymoma.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Cardio-respiratory arrest, Coronary artery stenosis, Endocarditis, Myocarditis, Pericarditis, Right ventricular failure, Sudden death.
  - Time to onset (pericarditis): 11 days after dose 3.
  - Cause of death: All the above events.

  A 62-year-old male subject from Austria.
  
  - Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Cardiac failure, Pericarditis.
- Time to onset (pericarditis): 14 days after dose 2.
- Cause of death: Both the above events.

- 2 cases non-medically confirmed:

A 53-year-old male subject from Italy.

- Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Pericarditis.
- Time to onset (pericarditis): Within 7 days after dose 1.
- Cause of death: Pericarditis.

A 62-year-old male subject from UK.

- Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Abdominal pain upper, Cardiac arrest, Chest pain, Dizziness, Dyspnoea, Fatigue, Immunisation, Myocarditis, Pain in extremity, Palpitations, Pericarditis, Thrombosis.
- Time to onset (pericarditis): 6 days after dose 3.
- Cause of death: All the above clinical events.

**Fatal Pericarditis cases in elderly (65-74 years of age) (2 cases, both non-medically confirmed)**

A 69-year-old female subject from UK.

- Medical history: Arthralgia, Brain neoplasm, Hypertension.
- Co-suspect medications: None.
- PTs with fatal outcome: Amnesia, Death, Interchange of vaccine products, Memory impairment, Myocarditis, Off label use, Pericarditis.
- Time to onset (pericarditis and myocarditis): Unspecified days after the dose 3.
- Causes of death: Brain neoplasm.

A 71-year-old male subject from UK.

- Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Chest pain, Death, Dyspnoea, Fatigue, Myocarditis, Palpitations, Pericarditis, Pulmonary embolism, Thrombosis.
- Time to onset (pericarditis and myocarditis): Unspecified days after the dose 2.
Causes of death: Death; Thrombosis.

**Fatal Pericarditis cases in elderly (> 75 years of age – 4 cases; 2 cases medically confirmed and 2 cases non-medically confirmed)**

- **2 cases medically confirmed:**

  An 89-year-old female subject from Spain.
  
  - Medical history: Anaemia megaloblastic, Aphasia, Arthropathy, Atrial fibrillation, Benign tumour excision, Cardiac assistance device user, Cerebrovascular accident, Chronic gastritis, Cognitive disorder, Diverticulum, Dyslipidaemia, Hypertension, Neoplasm, Oropharyngeal surgery, Parotitis, Salivary gland neoplasm, Sinus node dysfunction, Type 2 diabetes mellitus.
  - Co-suspect medications: Influenza vaccine INACT SAG 3V.
  - PTs with fatal outcome: Pericarditis.
  - Time to onset (pericarditis): 2 days after dose 3.
  - Causes of death: Pericarditis

  A 92-year-old male subject from Japan.
  
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Aortic dissection, Cardiac failure, Cardiac tamponade, Pericarditis.
  - Time to onset (pericarditis): Unspecified days after dose 3.
  - Causes of death: All the above events.

- **2 cases not medically confirmed:**

  An 81-year-old male subject, from Italy.
  
  - Medical history: Unknown.
  - Co-suspect medications: None.
  - PTs with fatal outcome: Pericarditis.
  - Time to onset (pericarditis): 60 days after dose 3.

A 78-year-old male subject from Slovakia.

- Medical history: Unknown.
- Co-suspect medications: None.
- PTs with fatal outcome: Multiple organ dysfunction syndrome, Myocardial infarction, Pericarditis.
- Time to onset (pericarditis): 8 days after dose 3.
- Causes of death: Due to all the above events.

Pericarditis relevant data in this subgroup of subjects are summarised in Table 51 below.

**Table 51. Pericarditis in Subjects aged ≥ 40 years (N=1756)**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td>Yes</td>
<td>538</td>
<td>388</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>474</td>
<td>338</td>
</tr>
<tr>
<td>Relevant PT*</td>
<td>Pericarditis</td>
<td>1001</td>
<td>720</td>
</tr>
<tr>
<td></td>
<td>Pericarditis constrictive</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Pleuropericarditis</td>
<td>15</td>
<td>4</td>
</tr>
<tr>
<td>Hospitalisation required/prolonged</td>
<td>Yes</td>
<td>225</td>
<td>264</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>789</td>
<td>462</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td>Dose 1</td>
<td>296</td>
<td>224</td>
</tr>
<tr>
<td></td>
<td>Dose 2</td>
<td>295</td>
<td>210</td>
</tr>
<tr>
<td></td>
<td>Dose 3</td>
<td>348</td>
<td>241</td>
</tr>
<tr>
<td></td>
<td>Dose 4</td>
<td>15</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>58</td>
<td>43</td>
</tr>
<tr>
<td>Time to Onset</td>
<td>&lt; 24 hours</td>
<td>53</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td>1-5 days</td>
<td>190</td>
<td>122</td>
</tr>
<tr>
<td></td>
<td>6-13 days</td>
<td>119</td>
<td>99</td>
</tr>
<tr>
<td></td>
<td>14-21 days</td>
<td>77</td>
<td>74</td>
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<td></td>
<td>22-31 days</td>
<td>57</td>
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</tr>
<tr>
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<td>32-60 days</td>
<td>64</td>
<td>44</td>
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<tr>
<td></td>
<td>61-180 days</td>
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<td>181-375 days</td>
<td>17</td>
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</tr>
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<td></td>
<td>Unknown</td>
<td>374</td>
<td>257</td>
</tr>
<tr>
<td>Event Outcome</td>
<td>Fatal</td>
<td>5</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Not resolved</td>
<td>355</td>
<td>207</td>
</tr>
<tr>
<td></td>
<td>Resolved</td>
<td>114</td>
<td>121</td>
</tr>
<tr>
<td></td>
<td>Resolved with sequelae</td>
<td>26</td>
<td>21</td>
</tr>
<tr>
<td></td>
<td>Resolving</td>
<td>215</td>
<td>148</td>
</tr>
<tr>
<td></td>
<td>Unknown</td>
<td>304</td>
<td>218</td>
</tr>
<tr>
<td>Duration of event</td>
<td>Up to 3 days</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>4-6 days</td>
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<td>7-10 days</td>
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<td>11-26 days</td>
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<td></td>
<td>27-57 days</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>58-180 days</td>
<td>16</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>181-265 days</td>
<td>3</td>
<td>0</td>
</tr>
</tbody>
</table>

a. All serious occurrences.
b. Multiple episodes of the same PT event were reported with a different clinical outcome in one case hence the sum of the events outcome exceeds the total number of PT events.
c. For those cases where the event resolved or resolved with sequelae.
Subjects with Unknown Age

Clinical Trial Data

- Number of cases: none; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 309 (0.06% of 507,683 cases of the total PM dataset, 0.5% of the 60,379 subjects with unknown age), compared to 363 (0.06%) cases retrieved in the PSUR #2.

- Country of incidence: UK (129), Canada (73), Australia (49), US (14), Germany (12), France, Greece (5 each). The remaining 22 cases were distributed among 13 countries.

- Subjects’ age in years: Unknown.

- Medical history (n = 90): the medical conditions reported more than twice included the PTs Hypertension (12), Asthma (11), Pericarditis (8), Drug hypersensitivity, Tobacco user (7), Palpitations (5), Anxiety, Fibromyalgia, Hypothyroidism, Immunodeficiency, Seasonal allergy, Steroid therapy (4 each), Blood cholesterol increased, Depression, Gastrooesophageal reflux disease, Insomnia, Non-tobacco user, Pain, and Rheumatoid arthritis (3 each).


- Co-suspects (n= 5 cases): COVID-19 vaccine MRNA (MRNA 1273) (2), Clozapine, COBID-19 vaccine NRV V AD (CHADOXI NCOV-19), Influenza vaccine (1 each).

- Most frequently co-reported PTs (≥2%): Chest pain (167), Palpitations (120), Myocarditis (116), Dyspnoea (109), Fatigue (108), Tachycardia (72), Immunisation (46), Interchange of vaccine products, Off label use (44 each), Pyrexia (41), Chest discomfort (33), Headache (26), Dizziness (25), Myopericarditis, Syncope (17 each), Nausea, Pain in extremity (16 each), Malaise (15), Pain (14), Arthralgia, Asthenia, Pericardial effusion (12 each), Hypoaesthesia (11), Chills, COVID-19, Inappropriate schedule of product administration, Myalgia, Paraesthesia (10 each), Heart rate increased (9), Angina pectoris, Drug ineffective, Hyperhidrosis, Migraine (8 each), Back pain, Cardiac flutter, Cough, Influenza like illness, Loss of personal independence in daily activities (7 each), Arrhythmia, Axillary pain, Heart rate abnormal, Insomnia, Loss of consciousness, Lymphadenopathy, Muscular weakness, Musculoskeletal chest pain, Pleuritic pain (6 each), Condition aggravated, Discomfort, Gait disturbance, Oxygen saturation decreased, Tremor, Vaccination site pain, Vision blurred, and Vomiting (5 each).

Pericarditis relevant data in this subgroup of subjects are summarised in the below Table 52.
Table 52. Pericarditis in Subjects with Unknown Age (N=309)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Female No. of Cases</th>
<th>Male No. of Cases</th>
<th>Unknown No. of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medically Confirmed</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>46</td>
<td>79</td>
<td>14</td>
</tr>
<tr>
<td>No</td>
<td>87</td>
<td>50</td>
<td>33</td>
</tr>
<tr>
<td>Re却ant PTa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pericarditis</td>
<td>132</td>
<td>129</td>
<td>47</td>
</tr>
<tr>
<td>Pleuropericarditis</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Hospitalisation required/ prolonged</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>27</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>No</td>
<td>106</td>
<td>99</td>
<td>44</td>
</tr>
<tr>
<td>Relevant suspect dose</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dose 1</td>
<td>38</td>
<td>50</td>
<td>13</td>
</tr>
<tr>
<td>Dose 2</td>
<td>43</td>
<td>44</td>
<td>8</td>
</tr>
<tr>
<td>Dose 3</td>
<td>47</td>
<td>25</td>
<td>11</td>
</tr>
<tr>
<td>Unknown</td>
<td>5</td>
<td>10</td>
<td>15</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Time to Onset (n=309)</th>
<th>Female No. of Events</th>
<th>Male No. of Events</th>
<th>Unknown No. of Events</th>
</tr>
</thead>
<tbody>
<tr>
<td>(&lt;24 \text{ hours})</td>
<td>4</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>1-5 days</td>
<td>22</td>
<td>28</td>
<td>3</td>
</tr>
<tr>
<td>6-13 days</td>
<td>2</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>14-21 days</td>
<td>5</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>22-31 days</td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>32-60 days</td>
<td>5</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>61-180 days</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>181-375 days</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>89</td>
<td>72</td>
<td>39</td>
</tr>
<tr>
<td>Event Outcome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fatal</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Not resolved</td>
<td>35</td>
<td>44</td>
<td>4</td>
</tr>
<tr>
<td>Resolved</td>
<td>6</td>
<td>19</td>
<td>4</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Resolving</td>
<td>11</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Unknown</td>
<td>79</td>
<td>57</td>
<td>37</td>
</tr>
</tbody>
</table>

\(a\). All serious occurrences.

\(b\). For those cases where the event resolved or resolved with sequelae.

Subjects with booster dose

Clinical Trial Data

- Number of cases: none; one (1) case was retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 1216 (0.2% of 507,683 cases of the total PM dataset, 1.0% of the 117,750 subjects who received a booster dose), compared to 283 cases (0.04%) in the PSUR #2.

CONFIDENTIAL
Page 188
COVID-19 mRNA vaccine (nucleoside modified)  
Periodic Safety Update Report (PSUR) 3  
19 December 2021 through 18 June 2022

- Country of incidence: UK (474), France (202), Germany (94), Italy (93), Netherlands (46), New Zealand (38), Norway (32), Japan (30), Israel, Sweden (21 each); the remaining 165 cases were distributed among 24 countries.

- MC (500), NMC (716).

- Subjects’ gender: female (661), male (531), and unknown (24).

- Subjects’ age in year: n = 1130, range: 13 -93, mean: 45.1, median: 44.0

- Medical history (n = 566): the medical conditions reported more or equal to 10 times included the PTs Hypertension (79), Pericarditis (40), Asthma (38), Immunodeficiency (32), Hypothyroidism (29), Obesity (22), Diabetes mellitus, Drug hypersensitivity (19 each), Seasonal allergy (18), Depression, Steroid therapy, Tobacco user (15 each), Atrial fibrillation, Non-tobacco user, Type 2 diabetes mellitus (14 each), Anxiety, Dyslipidaemia, Gastroesophageal reflux disease (13 each), Clinical trial participant, Disease risk factor, Migraine (12 each), Myocardial infarction, Rheumatoid arthritis (11 each), Chronic kidney disease, Food allergy (10 each).


- Co-suspects (n=20 cases): Influenza vaccine (6), Influenza vaccine INACT SAG 3V (3), Adalimumab, amoxicillin, Apixaban, Colchicine, COVID-19 vaccine, COVID-19 vaccine MRNA (MRNA 1273), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19), Etanercept, Propranolol, Salbutamol, Zuclopenthixol (1 each).

- Number of relevant events: 1220.

- Relevant event seriousness: all serious.

- Reported relevant PTs: Pericarditis (1212), Pleuropericarditis (8)

- Relevant event outcome78: fatal (12), resolved/resolving (414), resolved with sequelae (21), not resolved (296), unknown (478).

- Most frequently co-reported PTs (>3%): Chest pain (620), Myocarditis (461), Dyspnoea (448), Fatigue (427), Immunisation (418), Off label use (365), Palpitations (363), Interchange of vaccine products (332), Tachycardia (291), Pyrexia (218), Chest discomfort (151), Headache (124), Pericardial effusion (88), Malaise (87), Pain (82), Dizziness (74), Pain in extremity (66), Syncope (61), Arthralgia (58), Heart rate increased (57), Angina pectoris, Nausea (50 each), Asthenia (48), Arrhythmia (41), Chills, Lymphadenopathy, Myalgia (40 each), Back pain, Vaccination site pain (34 each), Cough (33).

The number of pericarditis cases occurred after a booster dose in each age group is reported in the below Table 53 by gender.
Table 53. Pericarditis in Subjects who Received a Booster Dose

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Heterologous Booster dose No. of Cases</th>
<th>Homologous Booster dose No. of Cases</th>
<th>Unknown dose No. of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F</td>
<td>M</td>
<td>U</td>
</tr>
<tr>
<td>Age group</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 to 17 years</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>18 to 24 years</td>
<td>4</td>
<td>13</td>
<td>0</td>
</tr>
<tr>
<td>25 to 29 years</td>
<td>8</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>30 to 39 years</td>
<td>23</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>40 years and older</td>
<td>140</td>
<td>94</td>
<td>4</td>
</tr>
<tr>
<td>Unknown</td>
<td>26</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>202</td>
<td>140</td>
<td>9</td>
</tr>
</tbody>
</table>

F=female; M=male; U=unknown

During the reporting period, of the 4156 cases reported, there were 1319 cases of medically confirmed pericarditis with a latency 21 days or less, of which in 975 cases pericarditis occurred within 1 week post vaccine administration. The majority (1255) of the cases were assessed as serious due to hospitalisation and/or medically significant. In 58 other cases, the seriousness criterion was reported as disability or life threatening, and in 6 cases, a fatal outcome was reported, which are reviewed above in the age-stratified sections.

Cumulatively, there were 9896 cases of pericarditis which constitute 0.7% of the overall PM dataset. During the current reporting period, there were 4156 cases reported which constitute 0.8% of 507,683 cases of the total PM dataset, and majority (~99%) of these cases were spontaneously reported, in which 43% of the cases were non-medically confirmed cases. Upon review of these 4156 cases, the majority of the cases (56.7%) were reported from adult population with the age group ranging from 30 to 64 years of age, where the female subjects (55.1%) were reported higher than the male subjects (44.1%). In the majority (66.2%) of the cases, the event of pericarditis was reported after the 1st dose (37.6%) or the 2nd dose (28.5%) and relatively less after the 3rd/booster dose (26.8%). Approximately 35% of the cases reported medical history such as hypertension, asthma and pericarditis, which might be attributed to the event of pericarditis. However, in the remaining cases, insufficient description of the cardiovascular and/or non-cardiovascular medical history and diagnostic to rule out other aetiologies in majority of the cases continues to preclude proper medical adjudication of causality assessment between administration of the vaccine and occurrence of pericarditis.

O/E Analysis

O/E analysis was performed for Myocarditis/Pericarditis (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Evaluation of Myocarditis and Pericarditis did not reveal any significant new safety information for this interval. Considering the accumulating data from post-authorisation use of the vaccine, including the consistent findings from passive and active surveillance
databases of increased occurrences of myocarditis and pericarditis following vaccination with BNT162b2, myocarditis and pericarditis have been added as ADRs in section 4.8, in Appendix A and Appendix B of the CDS v. 14.0, made effective on 26 July 2022, after the DLP.

16.3.2. Evaluation of Important Potential Risks

Evaluation of incremental data for the important potential risk VAED/VAERD is provided below.

Search criteria:

1. PTs Vaccine associated enhanced respiratory disease OR Vaccine associated enhanced disease OR

2. Standard Decreased Therapeutic Response Search (Drug ineffective OR Vaccination failure) AND 1 among the following PTs: Dyspnœa; Tachypnoea; Hypoxia; COVID-19 pneumonia; Respiratory failure; Acute respiratory distress syndrome; Cardiac failure; Cardiogenic shock; Acute myocardial infarction; Arrhythmia; Myocarditis; Vomiting; Diarrhoea; Abdominal pain; Jaundice; Acute hepatic failure; Deep vein thrombosis; Pulmonary embolism; Peripheral ischaemia; Vasculitis; Shock; Acute kidney injury; Renal failure; Altered state of consciousness; Seizure; Encephalopathy; Meningitis; Cerebrovascular accident; Thrombocytopenia; Disseminated intravascular coagulation; Chillblains; Erythema multiforme; Multiple organ dysfunction syndrome; Multisystem inflammatory syndrome in children.

VAED is a modified and/or severe presentation of an infectious disease affecting individuals exposed to the wild-type pathogen after having received vaccine designed to prevent infection.89

As noted by the Brighton Collaboration, there is currently no uniformly accepted definition of VAED (or VAERD) and the BC working group considers that a definitive case of VAED (Level 1 diagnostic certainty) cannot be ascertained with current knowledge of the mechanisms of pathogenesis of the condition; they have provided guidance on levels of diagnostic certainty of VAED cases based on various laboratory and clinical findings.

No post-authorisation AE reports have been identified as cases of VAED/VAERD, therefore, there is no observed data at this time. An expected rate of VAED is difficult to establish so a meaningful O/E analysis cannot be conducted at this point based on available data. The feasibility of conducting such an analysis will be re-evaluated on an ongoing basis as data on the virus grows and the vaccine safety data continue to accrue.


CONFIDENTIAL
Page 191
Of note, there were 9 cases reporting the PTs Vaccine associated enhanced disease, and Vaccine associated enhanced respiratory disease. None of them met the criteria to be considered as a true VAED case.

Clinical Trial Data

There were no cases reporting COVID-19 infection associated to one of the PTs utilised to identify potential severe or atypical cases of COVID-19.

Post-Authorisation Data

Of the 1278 cases retrieved based on search strategy, 10 cases were determined to be non-contributory and are not included in the discussion for the following reasons:

- In 4 cases the PT indicative of lack of efficacy did not refer to BNT162b2 vaccine.
- In 3 cases, the subjects developed SARS-CoV-2 infection during the early days from the 1st dose (days 1 – 13); therefore, the vaccine has not had sufficient time to stimulate the immune system and, consequently, the development of a vaccine preventable, even if severe, cannot be considered a potential case of enhanced disease.
- In 3 cases the PT Drug ineffective was erroneously coded; upon review, none of them developed COVID-19 infection.

Overview

- Number of cases: 1268 (0.2% of 507,683 cases, the total PM dataset), compared to 1490 (0.2%) retrieved in the PSUR # 2. All cases are serious.
- MC cases (878), NMC cases (390).
- Country of incidence: France (346), Spain (142), UK (139), US (117), Italy (105), Estonia (94), Germany (66), Philippines (45), Australia, Canada (19 each), Switzerland (18), Portugal, (17), Netherlands (14), Austria (10); the remaining 117 cases originated from 117 different countries.
- Gender: female (636), male (604), and unknown (28).
- Age in years (n = 1215), range: 5 – 102, mean: 61.4, median: 65.0.
- Relevant event seriousness: 1295 serious, 406 non-serious.
- Reported relevant PTs by organ system:
  - Respiratory system PTs (163): COVID-19 pneumonia (524), Dyspnoea (398), Respiratory failure (48), Acute respiratory distress syndrome (42), Pulmonary embolism (40), Hypoxia (28), and Tachypnoea (27).
  - Gastrointestinal/Hepatic system PTs (288): Diarrhoea (139), Vomiting (88), Abdominal pain (54), and Jaundice (7).
  - Cardiovascular system PTs (143): Myocarditis (85), Arrhythmia (32), Cardiac failure (18), Acute myocardial infarction (6), and Cardiogenic shock (2).
Renal and urinary system PTs (39): Acute kidney injury (27), and Renal failure (12).

Nervous system PTs (47): Seizure (21), Cerebrovascular accident (18), Encephalopathy (6), and Altered state of consciousness (2).

Vascular system PTs (23): Deep vein thrombosis (12), Shock (6), Vasculitis (3), and Peripheral ischaemia (2).

Blood and lymphatic system PTs (14): Thrombocytopenia (12), and Disseminated intravascular coagulation (2).

Immune system PTs (30): Vaccine associated enhanced disease (12), and Multisystem inflammatory syndrome in children (9), and Vaccine associated enhanced respiratory disease (9 each).

Other PTs (24): Multiple organ dysfunction syndrome (13), Chillblains (5), Meningitis (4), and Erythema multiforme (2).

Case outcome: fatal (184), not resolved (329), resolved/resolving (582), resolved with sequelae (31), and unknown (142).

COVID-19 positivity and severity of events

Suspected COVID-19 infection: 188 [no information on confirmatory tests performed or test negative; LOE coded to Drug ineffective (180 cases) or to Vaccination failure (8 cases, 2 of these cases also co-reported Vaccine associated enhance disease, Vaccine associated enhanced respiratory disease)]

Confirmed COVID-19 infection: 1080 [test positive or implied COVID-19 infection; LOE coded to Drug ineffective (524 cases) or Vaccination failure (556 cases, 7 of these cases also co-reported Vaccine associated enhanced disease, or Vaccine associate enhanced respiratory disease)].

Seriousness criteria for the total 1080 cases:
- Medically significant: 325;
- Hospitalisation required (non-fatal/non-life threatening): 521;
- Life threatening: 60;
- Death: 174.

Seriousness criteria: medically significant (450)

In 325 of 450 cases where the seriousness criterion was "medically significant", the subjects had a confirmed COVID-19 infection after vaccination, while 125 subjects had suspected COVID-19 infection. These 125 subjects did not require hospitalisation.
• In the 325 confirmed COVID-19 cases, subjects’ age ranged from 7 to 100 years (n = 304, mean: 47.0 years, median: 44.0 years) (14 paediatrics, 225 adults, 66 elderly, 20 unknown); gender was reported as female (212), male (101), and unknown (12).

• Time to event onset of the COVID-19 infection was reported for 250 of these 325 cases:
  - Day 13 to 344 after dose 1 (59 cases);
  - Day 0 to 492 after dose 2 (79 cases);
  - Day 0 to 329 after dose 3 (87 cases);
  - Day 6 to 95 after dose 4 (5 cases);
  - Day 12 to 431 after vaccination [dose number not reported] (20 cases).

• These 325 cases reported 419 relevant events. The most commonly (≥17 occurrences) reported relevant PTs Dyspnoea (137), Diarrhoea (76), Myocarditis (45), Vomiting (38), Abdominal pain (33), and COVID-19 pneumonia (17).

• The outcome of the COVID-19 infection related events reported in these 325 cases was: resolved/resolving (129), resolved with sequelae (2), not resolved (92), and unknown (197).

**Seriousness criteria: hospitalisation (non-fatal, non-life threatening) (568)**

• Hospitalisation occurred in 521 subjects, for 47 of them the COVID-19 infection was not confirmed.

• In the 521 COVID-19 confirmed cases, subjects’ age (n = 515) ranged from 10 to 102 years, (mean: 69.2 years, median: 73.0 years) (6 paediatrics, 166 adults, 344 elderly, 5 unknown); gender was reported as female (235), male (279), and unknown (7).

• Time to event onset of the COVID-19 infection was reported for 464 of these 521 cases.
  - Day 13 to 264 after dose 1 (20 cases);
  - Day 3 to 424 days after dose 2 (223 cases);
  - Day 0 to 232 days after dose 3 (199 cases);
  - Day 2 to 71 days after dose 4 (15 cases);
  - Day 9 to 249 after vaccination [dose number not reported] (7 cases).

• These 521 cases reported 695 relevant events. The most commonly (≥27 occurrences) reported relevant PTs COVID-19 pneumonia (344), Dyspnoea (146), Respiratory failure (29), and Diarrhoea (27).

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90 PTs included in the search strategy excluding Drug ineffective and Vaccination failure.
• The outcome of the COVID-19 infection related events reported in these 521 cases was: resolved/resolving (402), not resolved (86), resolved with sequelae (7), and unknown (200).

**Seriousness criteria: life-threatening (non-fatal) (66)**

• In 60 of the 66 cases characterised as life-threatening, the subjects had a confirmed COVID-19 infection after vaccination, while 6 subjects had suspected COVID-19 infection.

• In these 60 confirmed COVID-19 cases, subject’s age ranged from 5 to 94 years (n = 59), (mean: 57.2 years, median: 60.0 years), (4 paediatrics, 30 adults, 25 elderly, 1 unknown); gender was reported as female (17), and male (43).

• Time to event onset of the COVID-19 infection was reported for 47 of these 60 cases.
  - Day 17 to 244 after dose 1 (8 cases);
  - Day 26 to 377 after dose 2 (25 cases);
  - Day 1 to 184 after dose 3 (13 cases);
  - Day 18 after vaccination [dose number not reported] (1 case).

• These 60 cases reported 85 relevant events. The most commonly (≥6 occurrences) reported relevant PTs COVID-19 pneumonia (24), Dyspnoea (9), Pulmonary embolism (8), Acute respiratory distress syndrome, and Myocarditis (6 each).

• The outcome of the COVID-19 infection related events reported in these 60 cases was: resolved/resolving (30), not resolved (11), resolved with sequelae (2), and unknown (42).

**Seriousness criteria: Death (184 cases)**

One-hundred and eighty-four (184) subjects died, of which COVID-19 was not confirmed in 10 cases; the remaining 174 confirmed cases are described below.

• Age: 11 to 99 years (n = 171), Mean = 77.6 years, Median = 80.0 years.

• Country of incidence: France (78), Spain (26), Estonia (21), Italy (7), Germany, Malaysia, Philippines, Switzerland (4 each), Hungary, Portugal (3 each), Bulgaria, Norway, Slovenia, South Africa, UK, US (2 each), Austria, Czech Republic, Greece, Iceland, Lithuania, Luxembourg, Netherlands, and Sweden (1 each).

• Gender: female (65), male (107), and unknown (2).
Medical history (n = 153) included PTs in the following SOCs; Most frequently (≥4 occurrences) reported PTs by SOC are presented below:

- Vascular disorders - 93 cases (60.8%): Hypertension (81), Deep vein thrombosis (6), Aortic aneurysm, Aortic stenosis, and Peripheral venous disease (4 each);
- Metabolism and nutrition disorders - 74 cases (48.3%): Dyslipidaemia (28), Type 2 diabetes mellitus (23), Diabetes mellitus (13), Gout, Obesity (10 each), and Hypercholesterolaemia (5);
- Cardiac disorders - 71 cases (46.4%): Atrial fibrillation (32), Myocardial ischaemia (17), Cardiac failure (15), Myocardial infarction (11), and Hypertensive heart disease (8);
- Nervous system disorders - 52 cases (34.0%): Cognitive disorder (11), Cerebrovascular accident, Dementia Alzheimer’s type (7 each), Ischaemic stroke (6), Parkinson’s disease, and Transient ischaemic attack (4 each);
- Surgical and medical procedures - 43 cases (28.1%): Cardiac pacemaker insertion, Hysterectomy, and Stent placement (4 each);
- Neoplasms benign, malignant and unspecified (incl cysts and polyps) - 42 cases (27.4%): Chronic lymphocytic leukaemia (6), Plasma cell myeloma, and Prostate cancer (5 each);
- Respiratory, thoracic and mediastinal disorders - 31 cases (20.3%): Chronic obstructive pulmonary disease (8), Emphysema, Pulmonary embolism, and Sleep apnoea syndrome (4 each);
- Other medical histories were reported under the following SOCs: Musculoskeletal and connective tissue disorders (28), Gastrointestinal disorders, Social circumstances (26 each), Infections and infestations (21), Psychiatric disorders (20), Endocrine disorders (16), Injury, poisoning and procedural complications (15), Eye disorders (11), Hepatobiliary disorders, Immune system disorders, Reproductive system and breast disorders (9 each), Ear and labyrinth disorders (8), Skin and subcutaneous tissue disorders (7), Blood and lymphatic system disorders, General disorders and administration site conditions (6 each), Congenital, familial and genetic disorders (5), Investigations (2).

Latency of the COVID-19 occurrence was reported in 147 of the 174 cases:

- Day 13 to 29 after dose 1 (7 cases);
- Day 0 to 346 after dose 2 (64 cases);
- Day 2 to 214 after dose 3 (63 cases);
- Day 10 to 84 after dose 4 (3 cases);
- Day 5 to 192 after vaccination [dose number not reported] (10 cases).

The most frequently (>10 occurrences) reported causes of death in these 174 cases coded to the PTs COVID-19 pneumonia (112), COVID-19 (58), Vaccination failure (45), Drug ineffective (41), Acute respiratory distress syndrome, Dyspnoea (14), Multiple organ
dysfunction syndrome, and Respiratory failure (11 each). Of note, in 11 cases limited information regarding the cause of death was reported (PT Death).

- In 93 of the 174 fatal cases, vaccination failure was reported (cross referenced with Section 16.3.4.5 Lack of Therapeutic Efficacy).
- One hundred and fifty-two (152) of these 174 cases involved elderly subjects (aged 65 to 74 years [43] or ≥75 years [109]), including 138 subjects with underlying medical history of clinical significance.
- Among the remaining 22 cases; 15 of them had concurrent medical histories (less than or equal to 17 years [1], 18 to 30 years [1], 31 to 50 years [3], 51 to 64 years [9], and unknown [1]) that could impact the severity and evolution of the COVID-19 infection, including but not limited to cardiac history (carotid arteriosclerosis, hypertension, peripheral arterial occlusive disease, ventricular arrhythmia), renal disorders (chronic kidney disease) respiratory disorders (COVID-19, ex-tobacco user, tobacco user, acute respiratory distress syndrome, asphyxiating thoracic dystrophy, chronic respiratory failure, pleural thickening), and immunodeficient conditions (kidney transplant rejection, pulmonary mass, renal transplant, solid organ transplant).
- Of the remaining 7 cases where medical history was not reported, none of these 7 cases reported concomitant medications. The causes of death were reported as Dyspnoea (4), COVID-19 (2), Asthenia, Cerebral haemorrhage, Circulatory collapse, Cough, Hypoxia, Pain, Pyrexia, Vaccination failure, and Vomiting (1 each). Of note, in 1 case provided limited information regarding cause of death (PT Death). In 2 cases, the latency to onset of COVID-19 infection was not reported in the remaining 5 cases the latency was reported from dose 1 was 17 days, from dose 2 was: 59, 92, and 139 days, and from an unknown dose was reported as 5 days in 1 case.

Conclusion

The purpose of this review of subjects with COVID-19 following vaccination is to identify cases of potential vaccine-associated enhanced disease. The nature of spontaneously reported data provides a challenge because of the lack of a comparison group. Further, the background rate of VAED is not known. Considering the limitations of the review, VAED/VAERD remains a theoretical risk for the vaccine. Surveillance will continue.

16.3.3. Evaluation of Other Risks (not categorised as important)

There were no other risks that were classified as listed adverse events in which a SMSR or SMR assessment report recommended/requested continued monitoring in future PSURs and/or risks not categorised as important in which new information has become available during the reporting interval that allows further characterisation of a previously recognised risk.

16.3.3.1. Adverse Events of Special Interest (AESIs)

The company's AESI list takes into consideration the lists of AESIs from several expert groups and regulatory authorities including but not limited to the following: Brighton
Collaboration (SPEAC), ACCESS protocol, US CDC (preliminary list of AESI for VAERS surveillance), MHRA (unpublished guideline).

The AESI terms are incorporated into a TME list and include events of interest due to their association with severe COVID-19 and events of interest for vaccines in general. The AESI list includes MedDRA PTs, HLTs, HLGTs or MedDRA SMQs and will be changed as appropriate based on the evolving safety profile of the vaccine.

Overlapping terms among multiple categories were assigned to one category only based on their most clinical relevance.

Please refer to Appendix 6B for the observed versus expected analysis for the AESIs.

16.3.3.1.1. Anaphylactic AESIs

Please refer to the Risk ‘Anaphylaxis’ in Section 16.3.1 Evaluation of Important Identified Risks.

16.3.3.1.2. Cardiovascular AESIs

Search criteria – PTs: Acute myocardial infarction; Arrhythmia; Cardiac failure; Cardiac failure acute; Cardiogenic shock; Chest pain; Coronary artery disease; Myocardial infarction; Postural orthostatic tachycardia syndrome; Stress cardiomyopathy; Tachycardia.

Clinical Trial Data

- Number of cases: 27 (blinded therapy [6], and BNT162b2 [21]) (4.0 % of 668 cases, the total CT dataset) compared to 35 cases (4.9%) retrieved in the PSUR #2.
- Country of incidence: US (22), Argentina (3), Germany, South Africa (1 each).
- Subjects' gender: female (7), male (20).
- Subjects' age in years: (n = 27), range: 3-80, mean: 61.7, median: 64.
- Medical history (n = 23): the reported relevant medical conditions (>2 occurrences) included Hypertension (13), Type 2 diabetes mellitus (7), Hypercholesterolaemia, Hyperlipidaemia, Obesity (5 each), Coronary artery disease, Dyslipidaemia, Gastrooesophageal reflux disease (4 each), Anxiety, Depression, Osteoarthritis (3 each).
- COVID-19 medical history: None.
- Co-suspect medications: None.
- Reported relevant PTs: Myocardial infarction (9), Chest pain (8), Coronary artery disease (5), Acute myocardial infarction (4), Cardiac failure, Tachycardia (1 each).
- Relevant event outcome: fatal (2), resolved/resolving (21), resolved with sequelae (4), not resolved (1).
- None of the events were related BNT162b2 or blinded therapy.
Post-Authorisation Data

- Number of cases: 32,712 (6.4% of 507,683 cases, the total PM dataset), compared to 29,486 (4.5%) cases retrieved in the PSUR #2.
- MC cases (11,952), NMC cases (20760).
- Country of incidence (>16 occurrences): Germany (11,180), Australia (4456), UK (3049), France (2612), Taiwan Province of China (1393), Italy (1334), Netherlands (810), Austria (677), Malaysia (591), Philippines (475), US (422), Japan (397), New Zealand (396), Norway (369), Finland (338), Sweden (333), Belgium (328), Canada (317), Poland (307), Iraq (298), Greece (275), Spain (271), Ireland (237), Romania (217), Czech Republic (198), Denmark (125), Switzerland (119), Brazil (118), Lithuania (110), Croatia (92), Estonia (88), Portugal (82), Egypt (77), Israel (74), Slovenia (70), Mexico (62), Iceland (52), Slovakia (51), Hungary (50), Singapore (35), South Africa (30), Georgia (24), Latvia (24), Luxembourg (22), Turkey (20), Bulgaria (18), Cyprus (17); the remaining 72 cases were distributed among 31 countries.
- Subjects' gender: female (19,730), male (12,424) and unknown (558).
- Subjects’ age in years (n = 31,124), range: 2 months-99, mean: 40.3, median: 39.
- Medical history (n = 9348): the most frequently (>2%) reported relevant medical conditions included Hypertension (1349), Seasonal allergy (1121), Asthma (839), Drug hypersensitivity (758), Hypersensitivity (511), Food allergy (502), Mite allergy (387), Hypothyroidism (365), Tobacco user (290), Allergy to animal (285), Autoimmune thyroiditis (280), Diabetes mellitus (274), Obesity (253), Atrial fibrillation (232), Nontobacco user (232), Arrhythmia (220), Allergy to metals (216), Migraine (214).
- COVID-19 Medical history (n = 1546): the medical conditions reported included COVID-19 (1021), Suspected COVID-19 (492), Post-acute COVID-19 syndrome (40), COVID-19 pneumonia (16), SARS-CoV-2 test positive (14), Coronavirus infection (7), Asymptomatic COVID-19 (6), Exposure to SARS-CoV-2 (4), and Coronavirus pneumonia (1).
- Co-suspects (n = 295 cases): the frequently (>12 occurrences) reported relevant co-suspect medications were COVID-19 vaccine mRNA (MRNA 1273) (66), COVID-19 vaccine (34), COVID-19 vaccine NRV V AD (CHADOXI NCOV-19) (20), INFLUENZA VACCINE (19), adalimumab (13).
- Number of relevant events: 36,790.
- Relevant event seriousness37: serious (16,539), non-serious (20,268).
- Relevant PTs: Chest pain (17,945), Tachycardia (10,914), Arrhythmia (5627), Myocardial infarction (921), Cardiac failure (583), Acute myocardial infarction (364), Postural orthostatic tachycardia syndrome (149), Coronary artery disease (114), Cardiogenic shock (72), Cardiac failure acute (57), Stress cardiomyopathy (44).
- Time to event onset (n = 26,744 occurrences), range: <24 hours to 382 days, median: 1 day.
  - <24 hours: 7337 events (19 fatal events);
  - 1 day: 5830 events (56 fatal events);
  - 2-7 days: 7426 events (83 fatal events);
  - 8-14 days: 2202 events (45 fatal events);
  - 15-30 days: 1887 events (42 fatal events);
  - 31-181 days: 1892 events (68 fatal events);
  - 182-382 days: 170 events (6 fatal events).

- Duration of relevant events (n = 8262 out of 8906 occurrences with outcome of resolved and resolved with sequelae), range: <24 hours to 430 days, median: 4 days.
  - <24 hours: 816 events;
  - 1 day: 681 events;
  - 2-7 days: 1379 events;
  - 8-14 days: 476 events;
  - 15-30 days: 487 events;
  - 31-181 days: 904 events;
  - 182-430 days: 158 events.

- Relevant event outcome: fatal (496), resolved/resolving (13,937), resolved with sequelae (1321), not resolved (12,839), unknown (8437).

In 449 cases (reporting 496 relevant events with fatal outcome), the reported causes of death (>10 occurrences) were coded to the PTs Myocardial infarction (147), Cardiac failure (94), Chest pain (55), Acute myocardial infarction (54), Dyspnoea (43), Cardiac arrest (42), Arrhythmia (34), Cardiac failure acute (26), Myocarditis (22), Cardiogenic shock, Cardio-respiratory arrest (20 each), Thrombosis (15), Malaise (13), Loss of consciousness, Pulmonary embolism, Pulmonary oedema, Tachycardia (12 each), Pneumonia, Respiratory failure (11 each). Of note, in 16 cases limited information regarding the cause of death was provided (PT Death [11]; PT Sudden death [1]; Unknown [4]). Most (250 of 449 cases) of the fatal cases involved elderly subjects. When the medical history was provided (253 cases), the most frequently (≥ 9 occurrences) relevant medical conditions included events coded to the PTs Hypertension (91), Diabetes mellitus (32), Atrial fibrillation (28), Obesity (24), Cardiac failure (23), Type 2 diabetes mellitus (16), Coronary artery disease (15), Dyslipidaemia, Myocardial infarction (13 each), Chronic kidney disease, Chronic obstructive pulmonary disease (12 each), Cardiac disorder, Tobacco user (11 each), Cardiac failure chronic, Hyperlipidaemia (10 each), Arteriosclerosis, Asthma, Cerebral infarction, Myocardial ischaemia (9 each).

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91 This number does not include 10,261 events for which partial administration or event onset dates were reported or events did not have a meaningful time to onset value provided in the reported information.
Analysis by age group

CT: Paediatric (1), Adults (14), and Elderly (12).

- A meaningful comparison between the different age groups is not possible due to the low number of cases.

PM: Paediatric (2808), Adults (25850), Elderly (2996) and Unknown (1058).

- Higher reporting proportion of events coded to the PTs Arrhythmia [4.6% in paediatrics vs 18.2% in adults vs 24.3% in elderly], Cardiac failure [0.5% in paediatrics vs 0.8% in adults vs 11.5% in elderly], Myocardial infarction [0.3% in paediatrics vs 2.2% in adults vs 9.0% in elderly], Cardiogenic shock [0.1% in paediatrics vs 0.2% in adults vs 0.8% in elderly], Acute myocardial infarction [0.1% in paediatrics vs 0.8% in adults vs 4.8% in elderly], Cardiac failure acute [0.1% in paediatrics vs 0.1% in adults vs 0.8% in elderly], Stress cardiomyopathy [0.04% in paediatrics vs 0.1% in adults vs 0.7% in elderly], and Coronary artery disease [0% in paediatrics vs 0.3% in adults vs 1.5% in elderly] and was reported in elderly population when compared to adult and paediatric population. Higher reporting proportion of PT Chest pain [81.6% in paediatrics vs 53.5% in adults vs 36.3% in elderly] was reported in paediatrics compared to adults and elderly subjects. Higher reporting proportion of PT Tachycardia [19.6% in paediatrics vs 36.5% in adults vs 22.7% in elderly] was reported in adults compared to paediatrics and elderly subjects. The PT Postural orthostatic tachycardia syndrome was reported among the paediatric and adult subjects only (0.4% each).

Analysis by presence of comorbidities

Number of subjects with comorbidities: 3726 (0.7% of 507,683 cases, the total dataset).

No significant difference was observed in the reporting proportion of cardiovascular AESIs with fatal outcome in individuals with comorbid conditions (0.4% of events with fatal outcome) when compared to the reporting proportion observed in the individuals without comorbidities (0.9% of events with fatal outcome).

O/E Analysis

O/E analysis was performed for Acute myocardial infarction/Myocardial infarction; Arrhythmia; Coronary artery disease; Heart failure; Postural orthostatic tachycardia syndrome; Stress cardiomyopathy (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.
16.3.3.1.3. Haematological AESIs

Search criteria\(^92\) - HLTs (All Path) Leukopenias NEC; Neutropenias OR SMQ Haemorrhage terms (excl laboratory terms) OR PT Acquired haemophilia.

**Clinical Trial Data**

- Number of cases: 15 (BNT162b2 [12] and blinded therapy [3]) (2.2 % of 668 cases, the total CT dataset) compared to 19 cases (2.4%) retrieved in the PSUR #2\(^93\).
- Country of incidence: US (11), Argentina, Brazil, China, and Germany (1 each).
- Subjects’ gender: female (9), male (6).
- Subjects’ age in years (n = 15), range: 14-79, mean: 51.5, median: 49.0.
- Medical history (n = 15): the relevant medical conditions reported more than twice were coded to the PTs Hypertension (6), Depression, Type 2 diabetes mellitus, and Hypercholesterolaemia (3 each).
- COVID-19 Medical history: none.
- Co-suspects: none.
- Number of relevant events: 16.
- Reported relevant PTs (≥2 occurrences): Subdural haematoma (3), Haematoma, and Lower gastrointestinal haemorrhage (2 each). None of these SAEs were assessed as related to BNT162b2/blinded therapy.
- Relevant event outcome: fatal (1), resolved/resolving (13), resolved with sequelae (1), not resolved (1).

**Post-Authorisation Data**

- Number of relevant cases: 30,302 (5.9% of 507,683 cases, the total PM dataset), compared to 37,327 cases (5.7%) retrieved in the PSUR #2\(^93\).
- MC cases (4952), NMC cases (25,350).

\(^{92}\) The PT Acquired haemophilia has been added and PT Thrombocytopenia has been reassigned to Immune-mediated/autoimmune AESIs category.

\(^{93}\) The change to the search criteria should be considered when comparing the cases retrieved in the current PSUR and in PSUR #2.
COVID-19 mRNA vaccine (nucleoside modified)  
Periodic Safety Update Report (PSUR) 3  
19 December 2021 through 18 June 2022

- Country of incidence: Germany (7802), Netherlands (6166), UK (3266), Norway (2930), France (2905), Australia (1010), Spain (626), Italy (579), Sweden (557), Belgium (438); the remaining 4023 cases were distributed among 63 countries.

- Subjects' age in years (n = 28,488), range: 5 months-100 years, mean: 38.9, median: 37.0.

- Medical history (n = 10,294): the most frequently (≥200 occurrences) reported relevant medical conditions were coded to the PTs Disease risk factor (957), Hypertension (648), Menopause (620), Asthma (508), Seasonal allergy (427), Drug hypersensitivity (419), Amenorrhoea (404), Hypersensitivity (379), Hypothyroidism (295), Endometriosis (225), Food allergy (223), Migraine (213), and Contraception (200).

- COVID-19 Medical history (n = 2397): Medical conditions reported more than once were coded to the PTs COVID-19 (1636), Suspected COVID-19 (730), Post-acute COVID-19 syndrome (12), SARS-CoV-2 test positive (8), COVID-19 pneumonia (5), Coronavirus infection, Exposure to SARS-CoV-2 (2 each), Asymptomatic COVID-19, and Coronavirus test positive (1 each).

- Co-suspects: the most frequently (≥10 occurrences) reported relevant co-suspect vaccines/medications were COVID-19 vaccine MRNA (97), adalimumab (43), Influenza vaccine (36), COVID-19 vaccine NRV (31), levonorgestrel (24), and COVID-19 vaccine (19).

- Number of relevant events: 33,677.

- Relevant event seriousness: serious (8090) and non-serious (25,587).

- Most frequently reported relevant PTs (≥2%): Heavy menstrual bleeding (12,905), Intermenstrual bleeding (6088), Vaginal haemorrhage (1759), Epistaxis (1645), Contusion (1450), Vaccination site haematoma (1137), Postmenopausal haemorrhage (1137), Haematoma (944), and Haemorrhage (677).

- Time to event onset (n = 24,005 events), range: <24 hours to 7337 days, median: 3 days.
  - <24 hours: 4020 events (7 of which had a fatal outcome);
  - 1 day: 3680 events (10 of which had a fatal outcome);
  - 2-7 days: 6477 events (23 of which had a fatal outcome);
  - 8-14 days: 3113 events (15 of which had a fatal outcome);
  - 15-30 days: 3566 events (15 of which had a fatal outcome);
  - 31-181 days: 3003 events (24 of which had a fatal outcome);
  - ≥182 days: 146 events (5 of which had a fatal outcome).

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94 This number does not include 45 events for which partial administration or event onset dates were reported or events did not have a meaningful time to onset value provided in the reports.
• Duration of relevant events (n = 240 out of 572 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 21,170 days.
  - 1 day: 512 events;
  - 2-7 days: 1767 events;
  - 8-14 days: 522 events;
  - 15-30 days: 352 events;
  - 31-181 days: 1311 events;
  - 182-235 days: 64 events;
  - >235 days: 30 events.

• Relevant event outcome: fatal (146), resolved/resolving (11,605), resolved with sequelae (571), not resolved (13,999), and unknown (7480).
  - In the 174 fatal cases (including 146 relevant events with fatal outcome, reported in 114 cases), the reported causes of death (>8 occurrences) were coded to the PTs Haemorrhage (12), Gastrointestinal haemorrhage, Haematemesis, and Pericardial haemorrhage (9 each). Of note, in 19 cases limited information regarding the cause of death was provided (PT Death). Most (122 of 174 cases) of the fatal cases involved elderly subjects. When the medical history was provided (114 cases), the most frequently (>10 occurrences) relevant medical conditions included the PTs Hypertension (44), Cardiac arrest (16), Myocardial infarction (14), Cardiac failure, Cardio-respiratory arrest, Haemorrhage, and Myocardial ischaemia (10 each).

Analysis by age group

• CT: Adults (9) and Elderly (5).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

• PM: Paediatric (1044), Adults (26,592), Elderly (1731) and Unknown (935).
  - A significantly higher reporting proportion of events coded to the PTs Heavy menstrual bleeding and Intermenstrual bleeding was observed in paediatric and adult population when compared to elderly population (Heavy menstrual bleeding [33.9 % in paediatrics vs 45.7% in adults vs 0.2 % in elderly] and Intermenstrual bleeding [8.3% in paediatrics vs 22.1 % in adults vs 1.2 % in elderly]). The reporting proportion of the PT Epistaxis was significantly higher in paediatric and elderly population when compared to adult population (21.3 % in paediatrics vs 14.3 % in elderly vs 4.2 % in adults). The reporting proportion of PT Haematoma was higher in elderly population (12.1 %) when compared to paediatrics (1.2 %) and adult (2.0 %) population. The comparative differences in reporting proportions are not unexpected given the generally expected medical issues affecting each age group (paediatrics, adults, elderly).
Analysis by presence of comorbidities

- Number of subjects with comorbidities: 2542 (8.4% of the CT and PM cases reporting haematological AESIs).

- The reporting proportion of haematological AESIs with fatal outcome (1.6%) is higher in individuals with comorbid conditions when compared to the reporting proportion observed in the individuals without comorbidities (0.3%).

O/E Analysis

O/E analysis was performed for Acquired haemophilia and Haemorrhage (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Acquired haemophilia was evaluated during this reporting period (please refer to Section 15 Overview of Signals: New, Ongoing, or Closed and to Appendix 6A.2 for cumulative review of cases indicative of acquired haemophilia). No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

16.3.3.1.4. COVID-19 AESIs

Search criteria – SMQ COVID-19 (Narrow and Broad) OR PTs: Ageusia; Anosmia; Vaccine associated enhanced disease; Vaccine associated enhanced respiratory disease.95. Cases reporting long COVID (PT: Post-acute COVID-19 syndrome) are reviewed in this section. Please refer also to Section 18.1 Benefit-Risk Context – Medical Need and Important Alternatives (Complications of COVID-19 and Post-acute COVID).

Clinical Trial Data

- Number of cases: 7 (blinded therapy [3] and BNT162b2 [4]) (1.0 % of 668 cases, the total CT dataset) compared to 3 cases (0.4%) retrieved in the PSUR #293.

- Country of incidence: US (3), Argentina, Dominican Republic, South Africa and Spain (1 each).

- Subjects' gender: female (5), male (2).

- Subjects' age in years: (n = 6), range: 2-77, mean: 38.0, median: 33.0

95 The PTs Vaccine associated enhanced disease and Vaccine associated enhanced respiratory disease are evaluated in Section 16.3.2. Evaluation of Important Potential Risks, as overlapping terms with the important potential risk Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD).
• Medical history (n = 7): the reported relevant medical conditions included the PTs Asthma, Hypothyroidism (2 each), Abscess, Anxiety disorder, Basal cell carcinoma, Bipolar disorder, Catheterisation cardiac, Concussion, Coronary arterial stent insertion, Depression, Epilepsy, Ex-alcohol user, Glucose tolerance impaired, Headache, Heart transplant, Hypercholesterolaemia, Hyperlipidaemia, Hypertension, Hypocalcaemia, Medical procedure, Migraine, Non-tobacco user, Pain, Palpitations, Pulmonary valve stenosis, Seasonal allergy, Sinus bradycardia, Stress, Surgery, Varicella (1 each).

• COVID-19 Medical history: none.

• Co-suspects: none.

• Reported relevant PTs: COVID-19 (6), COVID-19 pneumonia (1). None of the events were related to BNT162b2 or blinded therapy.

• Relevant event outcome: fatal (1), resolved/resolving (6).

Post-Authorisation Data

• Number of relevant cases: 54,335 (10.7% of 507,683 cases, the total PM dataset), compared to 25,453 cases (3.9%) retrieved in the PSUR #2\(^3\). The increase in the number of cases reported during the current PSUR is attributed to the increase in cases reported from Austria (9068 cases in the PSUR #2 vs 31,769 cases in the current PSUR #3) due to active solicitation of LOE cases from the Austrian BoH.

• MC cases (40,416); NMC cases (13,919).

• Country of incidence (≥ 2%): Austria (31,769), US (4874), UK (2725), Germany (2386), France (1934), Netherlands (1495); the remaining 9152 cases were distributed among 77 countries.

• Subjects' gender: female (29,370), male (22,867) and unknown (2098).

• Subjects' age in years: (n = 51,267), range: 6 months – 107 years, mean: 47.1, median: 46.0.

• Medical history (n = 8328): the most frequently (≥2%) reported relevant medical conditions included Hypertension (1429), Asthma (766), Drug hypersensitivity (617).

• COVID-19 Medical history: COVID-19 (1018), Suspected COVID-19 (361), Exposure to SARS-CoV-2 (49), Post-acute COVID-19 syndrome (48), COVID-19 pneumonia (10), SARS-CoV-2 test positive (9), Asymptomatic COVID-19, Coronavirus infection (3 each), Coronavirus test positive, Occupational exposure to SARS-CoV-2 (2 each).

• Co-suspects (n = 3995 cases): the most frequently (≥10) reported relevant co-suspect vaccines/medications were COVID-19 vaccine (1861), COVID-19 vaccine NRVV AD (CHADOXI NCOV-19) (798), COVID-19 vaccine mRNA (MRNA 1273) (768), Adalimumab (256), JNJ 78436735 (100), Ocrelizumab (35), Influenza vaccine (34),
Upadacitinib (31), COVID-19 vaccine INACT (VERO) CZO2 (25), Risankizumab (21), Prednisone (18), Casirivimab/Imdevimab, Rituximab (13 each), Mycophenolate (10).

- Number of relevant events: 55,437.
- Relevant event seriousness: serious (52,185), non-serious (3254).
- Most frequently reported relevant PTs (≥2%): COVID-19 (47,981) Suspected COVID-19 (3002), and Ageusia (1094).
- Time to event onset (n = 46,269\textsuperscript{96}), range: <24 hours to 564 days, median: 117 days.
  - <24 hours: 828 events (7 fatal events);
  - 1 day: 598 events (2 fatal events);
  - 2-7 days: 2061 events (24 fatal events);
  - 8-14 days: 1832 events (36 fatal events);
  - 15-30 days: 1812 events (35 fatal events);
  - 31-181 days: 36339 events (256 fatal events);
  - ≥ 182 days: 2799 events (53 fatal events).
- Duration of relevant events (n = 1968 out of 4800 occurrences with outcome of resolved/resolved with sequelae), range: 24 hours to 373 days, median: 9 days:
  - <24 hours: 71 events
  - 1 day: 46 events
  - 2-7 days: 631 events
  - 8-14 days: 848 events
  - 15-30 days: 275 events
  - 31-181 days: 76 events
  - ≥ 182 days: 9 events
- Relevant event outcome\textsuperscript{78}: fatal (506), resolved/resolving (7289), resolved with sequelae (296), not resolved (3281), unknown (44071).
  - In 493 cases (reporting 543 relevant events of which 506 relevant events reported a fatal outcome), the reported causes of death (≥20 occurrences) were coded to the PTs COVID-19 (297), Vaccination failure (143), Drug ineffective (131), COVID-19 pneumonia (127), Death (46), Dyspnoea (21). Of note, in 39 cases limited information regarding the cause of death was provided (PT Death [38] and Sudden death [1]). Most (406 of 493 cases) of the fatal cases involved elderly subjects. When the medical history was provided (272 cases), the most frequently (≥20 occurrences) relevant medical conditions included the PTs Hypertension (117), Atrial

\textsuperscript{96} This number does not include 28 events for which a meaningful time to onset value was not provided in the reported information.
fibrillation (53), Chronic kidney disease, Dyslipidaemia (33 each), Type 2 diabetes mellitus (29), Myocardial ischaemia (25), Cardiac failure (24), COVID-19 (22), and Diabetes mellitus (21).

Analysis by age group

- CT: Paediatric (1), Adults (5), Elderly (1).
  - Due to low volume of paediatric cases, a meaningful comparison of the same with the other age groups is not possible.

- PM: Paediatric (2158), Adults (39,726), Elderly (9566).
  - No significant difference was observed in the reporting proportion of the most frequently reported COVID-19 AEs (≥2%) between adult, elderly and paediatric population.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 3846 (0.8% of 507,683 cases, the total dataset).

  The reporting proportion of COVID-19 AEsIs with fatal outcome (5.6% [230 of 4093 events]) is higher in subjects with comorbid conditions, compared to the reporting proportion observed in the individuals without comorbidities (0.5% [276 of 51,344 cases] of fatal events).

LONG COVID

Clinical Trial Data

- Number of cases: none.

Post-Authorisation Data

- Number of relevant cases: 200 (0.04% of 507,683 cases, the total PM dataset), compared to 72 cases (0.3% of 25,453 cases) retrieved in the PSUR #2.

- MC cases (62); NMC cases (138).

- Country of incidence: Germany (106), France (15), UK (14), Austria (13), Sweden (9), Australia, Finland (8 each), Netherlands (6), Italy (4), Ireland, US (3 each), Belgium, Hungary, New Zealand, Spain (2 each), Brazil, Greece and Luxembourg (1 each).

- Subjects' gender: female (151), male (46) and unknown (3).

- Subjects' age in years: (n = 174), range: 9 – 85 years, mean: 43.6, median: 45.0. Of these 174 subjects, there were 10 paediatric, 156 adults, and 8 elderly subjects.
• Medical history (n = 104): the most frequently (≥2%) reported medical conditions included Asthma, Drug hypersensitivity (8 each), and Seasonal allergy (7).


**Analysis by presence of co-morbidities:**

• Number of subjects with comorbidities: 33 (16.5% of the 200 cases).

• Co-suspects (n = 9 cases): the reported co-suspect vaccines/medications were COVID-19 vaccine (5), COVID-19 vaccine mRNA (MRNA 1273) (3), Rupatadine (1).

• Co-reported events (≥20 occurrences): Fatigue (82), Headache (46), Condition aggravated (35), Dyspnoea (33), Disturbance in attention, Pyrexia (31 each), Drug ineffective (29), Asthenia (28), Myalgia (26), Palpitations (24), and Chest pain (20).

• Relevant event seriousness: serious (93), non-serious (107).

• Relevant event outcome: Resolved/resolving (22), Resolved with sequelae (9), Not resolved (98), Unknown (70) and Fatal (1).

The case reporting the fatal outcome was spontaneously reported, that involved an 83-year-old female patient with co-morbid conditions (such as angina pectoris, macular degeneration, mitral valve incompetence, myalgia, pneumonia bacterial, respiratory failure) who received 2 doses of Comirnaty (on 27 January 2021 and on 05 March 2021) for immunisation and reported multiple serious events coded to the PTs Post-acute COVID-19 syndrome, Cough, Pneumonia, Hypoxia, Myocardial infarction, Acute myocardial infarction, Angina pectoris, and Cardiac failure. In this case all the above fatal events were likely attributable to the patient’s medical history and an individual contributory role of Comirnaty cannot be established.

• Time to event onset (n = 95), range: <24 hours to 251 days, median: 2 days.
  - <24 hours: 25 events;
  - 1 day: 18 events;
  - 2-7 days: 20 events;
  - 8-14 days: 7 events;
  - 15-30 days: 15 events;
  - 31-181 days: 9 events;
  - ≥182 days: 1 event.

• Duration of relevant events (n = 7 out of 20 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 209 days, median: 4 days:
  - 1 day: 1 event
  - 8-14 days: 1 event
- 31-181 days: 4 events
- ≥ 182 days: 1 event

O/E Analysis

O/E analysis was performed for Ageusia/anosmia (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Loss of/Altered Taste and Smell was evaluated as signal during the reporting period and determined not to be a risk (please refer to Section 16.2.1 Evaluation of Closed Signals).

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

16.3.3.1.5. Dermatological AEsIs

Search criteria - PTs: Chillblains; Erythema multiforme.

Clinical Trial Data

- During the reporting period no serious cases from the CT dataset were reported; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 284 (0.06% of 507,683 cases, the total PM dataset), compared to 339 (0.05%) cases retrieved in the PSUR #2.
- MC cases (158), NMC cases (126).
- Country of incidence: France (72), Germany (38), UK (25), Italy (24), Singapore (18), Japan, Poland (11 each), the Netherlands, US (9 each), Australia (8), Belgium (7), Canada, New Zealand, Spain (6 each); the remaining 34 cases were distributed among 18 countries.
- Subjects' gender: female (182), male (93) and unknown (9).
- Subjects' age in years: (n = 269), range: 7-89, mean: 46.4, median: 46.
- Medical history (n = 102): the most frequently (≥ 4 occurrences) reported relevant medical conditions included Hypertension (16), COVID-19 (12), Asthma (8), Drug hypersensitivity, Suspected COVID-19 (6 each), Diabetes mellitus (5), Cerebrovascular accident, Food allergy, Herpes simplex, Hypothyroidism, Type 2 diabetes mellitus (4 each).
- COVID-19 Medical history (n = 17): COVID-19 (12), Suspected COVID-19 (6), and Post-acute COVID-19 syndrome (1).
• Co-suspects (n = 4 cases): Acetylcysteine/benzalkoniumchloride/tiaminoheptane sulfate, Albendazole, Dextromethorphan, Ibuprofen, Ketoprofen, Ocrelizumab, Prednisolone metasulfobenzoate sodium, Sulfasalazine (1 each).

• Number of events: 284.

• Relevant event seriousness: serious (206), non-serious (78).

• Reported relevant PTs: Erythema multiforme (181), Chillblains (103).

• Time to event onset (n = 72), range: <24 hours to 262 days, median: 4 days.
  - <24 hours: 26 events;
  - 1 day: 35 events;
  - 2-7 days: 69 events;
  - 8-14 days: 31 events;
  - 15-30 days: 26 events;
  - 31-180 days: 24 events.

• Duration of relevant events (n = 14 out of 53 occurrences with outcome of resolved/resolved with sequelae), range: 0 days to 67 days, median: 20.5 days:
  - 1-7 days: 2 events;
  - 8-14 days: 2 events;
  - 15-30 days: 5 events;
  - 31-180 days: 3 events.

• Relevant event outcome: resolved/resolving (108), resolved with sequelae (8), not resolved (118), unknown (50). No fatal events were reported.

Analysis by age group

• PM: Paediatric (31), Adults (183), Elderly (60) and Unknown (10).
  - Due to low volume of paediatric cases a meaningful comparison of the same with the other age groups is not possible. No significant difference observed in the reporting proportion of events chillblains and erythema multiforme between adult and elderly population.

Analysis by presence of comorbidities

• Number of subjects reporting comorbidities: 53 (18.7 % of the cases reporting dermatological AESIs). A higher reporting proportion of dermatological AESIs was reported in subjects without significant comorbidities (81.3 %) when compared to subjects with significant comorbidities.
O/E Analysis

O/E analysis was performed for Chillblains and Erythema multiforme (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No other safety signals have emerged based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

16.3.3.1.6. Facial Paralysis

Search criteria – PTs: Bell’s palsy, Facial paralysis, Facial paresis, Oculofacial paralysis.

Clinical Trial Data

- Number of cases: 1 (BNT162b2 [1]) (0.1% of 668 cases, the total CT dataset) compared to no cases retrieved in the PSUR #2.

- Country of incidence: US (1).

- Subject’s gender: female (1).

- Subject’s age: 75 years (1).

- Medical history (n = 1): Blood cholesterol increased, Cataract, Dyspepsia, Foot fracture, Gastroesophageal reflux disease, Headache, Hypermetropia, Hypertension, Hypertonic bladder, Myopia, Overweight, and Squamous cell carcinoma of skin (1 each).

- COVID-19 Medical history: none.

- Co-suspects: none.

- Reported relevant PT: Bell’s palsy (1), not related to BNT162b2.

- Relevant event outcome: resolved (1).

Post-Authorisation Data

- Number of cases: 2589 (0.5% of 507,683 cases, the total PM dataset), compared to 4515 cases (0.7%) retrieved in the PSUR #2.

- MC cases (1105), NMC cases (1484).

- Country/region of incidence: Germany (714), France (387), UK (229), Australia (184), Italy (112), Austria (97), Sweden (87), Hong Kong (70), Taiwan, Province of China (69), US (54); the remaining 586 cases were distributed among 40 countries.

- Subjects’ gender: female (1487), male (1060), and unknown (42).
Subjects’ age in years: \( n = 2473 \), range: 1.42 – 99, mean 47.3, median 47.0.

Medical history \( n = 934 \): the most frequently (>2%) reported relevant medical conditions were coded to the PTs Hypertension (185), Asthma (89), Seasonal allergy (85), Drug hypersensitivity (63), Hypersensitivity (57), Diabetes mellitus (55), Type 2 diabetes mellitus (40), Obesity (35), Food allergy (34), Hypothyroidism (32), Facial paralysis, Mite allergy (28 each), Allergy to animal (26), Bell’s palsy (25), Hypercholesterolaemia (24), Chronic obstructive pulmonary disease, Tobacco user (23 each), Migraine (20), and Coronary artery disease (19).

COVID-19 Medical history \( n = 133 \): reported medical conditions were coded to the PTs COVID-19 (100), Suspected COVID-19 (31), Post-acute COVID-19 syndrome (4), SARS-CoV-2 test positive (2), Asymptomatic COVID-19, Coronavirus infection, and COVID-19 pneumonia (1 each).

Co-suspects \( n = 33 \): the relevant co-suspect vaccines/medications were diphtheria vaccine toxoid/ polio vaccine inact 3V (vero)/ tetanus vaccine toxoid and meningococcal group C tetanus toxoid conjugate vaccine (1 each).

Number of relevant events: 2706.

Reported relevant seriousness: \( n = 42 \) serious (2431) and non-serious (543).

Reported relevant PTs: Facial paralysis (1428), Bell’s palsy (733), Facial paresis (543), and Oculofacial paralysis (2).

Time to event onset \( n = 2152 \) events, range: <24 hours to 389 days, median 7 days.

- <24 hours: 351 events (none of which had a fatal outcome);
- 1 day: 233 events (1 of which had a fatal outcome);
- 2-7 days: 529 events (1 of which had a fatal outcome);
- 8-14 days: 265 events (none of which had a fatal outcome);
- 15-30 days: 332 events (2 of which had a fatal outcome);
- 31-181 days: 414 events (1 of which had a fatal outcome);
- 182-389 days: 27 events (none of which had a fatal outcome).

Duration of relevant events \( n = 286 \) out of 613 occurrences with outcome of resolved/resolved with sequelae, range: <24 hours to 246 days, median: 6 days.

- <24 hours: 48 events;
- 1 day: 32 events:
- 2-7 days: 69 events;
- 8-14 days: 18 events;

\(^{97}\) This number does not include 2 events for which partial administration and/or event onset date was reported.
- 15-30 days: 41 events;
- 31-181 days: 71 events.
- 182-246 days: 7 events.

- Relevant event outcome: fatal (6), resolved/resolving (999), resolved with sequelae (111), not resolved at the time of reporting (1063), and unknown (534).

In 6 cases (reporting 6 relevant events with a fatal outcome), the causes of death (≥2 occurrences) were coded to the PTs: Facial paralysis (3), Cerebrovascular accident and Death (2 each). Of note, in 2 cases limited information regarding the cause of death was provided (PT Death). All of the patients were >60 years of age (range 61 to 99 years). When the medical history was provided (4 cases), significant medical conditions reported: Arthralgia, Cerebral infarction, Diverticulitis, Lung adenocarcinoma, Neoplasm malignant, Pemphigoid, and Pulmonary embolism (1 each).

Analysis by age group

- PM: Paediatric (146), Adults (1914), Elderly (420), and Unknown (109).
  - There was no significant difference observed in the reporting proportion of facial paralysis events between age groups.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 407 (15.7% of the CT and PM cases reporting facial paralysis).
  - The reporting proportion of cases reporting a facial paralysis events with a fatal outcome is higher in subjects with comorbid conditions (0.74%) when compared to the reporting proportion observed in the subjects without comorbidities (0.14%).

O/E Analysis

O/E analysis was performed for Bell’s palsy (PTs: Bell’s palsy, Facial paralysis, Facial paresis, Oculo-facial paralysis) (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.
16.3.3.1.7. Hepatic AESIs

Search criteria - SMQ Liver related investigations, signs and symptoms (Narrow and Broad) OR PTs Autoimmune hepatitis\(^98\), Liver injury.

Upon review, 2 cases were determined to be non-contributory and were not included in the discussion since these cases involved exposure to the vaccine during the mother’s pregnancy or through breastfeeding\(^99\).

**Clinical Trial Data**

- During the reporting period no serious cases from the CT dataset were reported, compared to 2 cases (0.28%) retrieved in the PSUR #2\(^93\).

**Post-Authorisation Data**

- Number of relevant cases: 1091 (0.2 % of 507,683 cases, the total PM dataset), compared to 1393 cases (0.2%) retrieved in the PSUR #2\(^93\).

- MC cases (560), NMC cases (531).

- Country of incidence: Germany (276), Japan (157), France (152), UK (71), Australia (66), US (55), Italy (44), Austria (35), Spain (29), Taiwan, province of China (26), Netherlands (17), Sweden (15), Belgium, Finland (14 each), New Zealand (13), Greece (11), Canada (10), Denmark (9), Czech Republic (8), Norway, Poland (6 each), Croatia, Ireland, Romania (5 each), Switzerland (4), Brazil, Latvia, Philippines, Portugal, Slovakia, Slovenia (3 each), Estonia, Hungary, Lithuania, Malaysia, Mexico (2 each); the remaining 10 cases were distributed among 10 countries.

- Subjects' gender: female (661), male (406) and unknown (24).

- Subjects' age in years (n = 1017), range: 5 - 94, mean: 49.3, median: 51.0.

- Medical history (n = 518): the most frequently reported relevant medical conditions (≥ 5 occurrences) included Hypertension (80), Drug hypersensitivity (38), Hypothyroidism, Seasonal allergy (33), Asthma (32), Food allergy (23), Autoimmune thyroiditis, Type 2 diabetes mellitus (21 each), Allergy to animal (20), Allergy to metals, Dyslipidaemia (19 each), Mite allergy (17), Atrial fibrillation (16), Diabetes mellitus, Hepatic steatosis, Tobacco user (15 each), Obesity (14), Hypercholesterolaemia, Hypersensitivity (13 each), Gastroesophageal reflux disease, Rheumatoid arthritis (12 each), Breast cancer, Mycotic allergy, Non-tobacco user, Tonsillectomy (11 each), Autoimmune hepatitis, Depression, Ovarian cystectomy (10 each), Interchange of vaccine products, Osteoporosis, Salivary gland operation (9 each), Cardiac failure, Cholecystectomy, Liver disorder, Migraine (8

\(^98\) The PT Autoimmune hepatitis has been added, compared to search criteria used in PSUR #2 (cross-referenced with Section 15 Overview Of Signals: New, Ongoing, Or Closed).

\(^99\) These cases are included in Section 16.3.5.3 Use in Pregnant/Lactating Women.
each), Abstains from alcohol, Alcohol use, Allergy to plants, Anxiety, Arrhythmia, Epstein-Barr virus infection, Headache, Hysterectomy, Pyrexia, Type 1 diabetes mellitus (7 each), Coeliac disease, Disease risk factor, Hepatic cirrhosis, Hyperuricaemia, Thyroid disorder, Weight decreased (6 each), Colon cancer, Dermatitis contact, Diverticulum intestinal, Epilepsy, Hepatic function abnormal, Hepatitis, Hyperlipidaemia, Immune-deficiency, Insomnia, Nephrolithiasis, Neuropathy peripheral, Pericarditis, Primary biliary cholangitis, Sinus operation, Sjogren’s syndrome, Sleep apnoea syndrome, and Thyroid cancer (5 each).

- COVID-19 Medical history (n = 46): the medical conditions reported included COVID-19 (34), Post-acute COVID-19 syndrome, Suspected COVID-19, (5 each), Asymptomatic COVID-19, and COVID-19 pneumonia (1 each).

- Co-suspects (n = 58): the relevant co-suspect medications reported were adalimumab (10), upadacitinib (3), atorvastatin, hepatic A vaccine, methotrexate, paracetamol (2 each), amlodipine, amoxicillin, cabozantinib, cefuroxime, certolizumab, clopidogrel, clozapine, colchicine, drosopirone, ethinylestradiol, ebastine, ethinylestradiol gestodene, exemestane, fingolimod, ibuprofen, ipilimumab, lanreotide, nitrofurantoin, nivolumab, paclitaxel, ribociclib, rosuvastatin, sorafenib, spironolactone, teriflunomide, torasemide, and valsartan (1 each).

- Number of relevant events: 1422.

- Relevant event seriousness: serious (676) and non-serious (746).

- Most frequently reported relevant PTs (≥50 occurrences): Hepatic enzyme increased (131), Alanine aminotransferase increased (126), Hepatic function abnormal (124), Liver function test abnormal (119), Aspartate aminotransferase increased (110), Autoimmune hepatitis (99), Hepatic pain (98), Gamma-glutamyltransferase increased (86), Liver function test increased (79), Transaminases increased (72), Ascites (60).

- Time to event onset (n = 876 events)\(^{100}\), range: <24 hours to 177 days, median: 7 days.
  - <24 hours: 83 events (of which 1 had a fatal outcome);
  - 1 day: 77 events;
  - 2-7 days: 290 events (of which 3 had a fatal outcome);
  - 8-14 days: 133 events (of which 4 had a fatal outcome);
  - 15-30 days: 132 events (of which 2 had a fatal outcome);
  - 31-180 days: 161 events (of which 2 had a fatal outcome).

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\(^{100}\) This number is not including 546 events for which partial administration and/or event onset dates were reported or events did not have a meaningful time to onset value provided in the reported information.
• Duration of relevant events (n = 120 out of 1425 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 210 days, median: 23 days.
  – <24 hours: 4 events
  – 1 day: 2 events;
  – 2-7 days: 24 events;
  – 8-14 days: 18 events;
  – 15-30 days: 22 events;
  – 31-180 days: 49 events;
  – >180 days: 1 event.

• Relevant event outcome78: fatal (23), resolved/resolving (426), resolved with sequelae (46), not resolved at the time of reporting (343), and unknown (586).

In 22 cases with fatal outcome (reporting 23 relevant events with fatal outcome), the reported causes of death were coded to Ascites (5), Congestive hepatopathy, Hepatic function abnormal, Hepatic pain, Hypertransaminasaemia (2 each), Alanine aminotransferase increased, Autoimmune hepatitis, Blood bilirubin increased, Hepatic enzyme increased, Hepatic mass, Hepatomegaly, Hepatosplenomegaly, Hypoalbuminaemia, Liver function test abnormal, and Liver injury (1 each). Of note, in 6 cases limited information regarding the cause of death was provided (Alanine aminotransferase increased, Ascites, Hepatic mass, Hepatic pain, Hypertransaminasaemia, Liver injury (1 each). Most (13 of 22 cases) of the fatal cases involved subjects who were ≥60 years of age.

When the medical history was provided (13 cases), the relevant medical conditions included Hepatic steatosis, Type 2 diabetes mellitus (3 each), Diabetes mellitus (2), Autoimmune hepatitis, and Hepatic function abnormal (1 each).

Analysis by age group

• PM: Paediatric (66), Adults (712), Elderly (243) and No data (70).
  – Among the frequently (≥2%) reported relevant hepatic events, Hepatic pain was reported significantly higher in the adult population when compared to elderly population (25.5% in adult vs 6.3% in elderly). Upon further review, the majority of the events of hepatic pain were assessed as non-serious in the adult population (63 of 84 events).

Analysis by presence of comorbidities

• Number of subjects with comorbidities: 265 (24.3% of the CT and PM cases reporting hepatic AESIs).

• The reporting proportion of hepatic AESIs with fatal outcome (2.1%) is slightly higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subjects without comorbidities (1.4%)
O/E Analysis

O/E analysis was performed for Acute liver injury/Liver injury and Autoimmune hepatitis (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

A cumulative review of Autoimmune hepatitis has been performed (please refer to Section 15 Overview of Signals: New, Ongoing, or Closed and to Appendix 6A.5 for a cumulative review of cases indicative of autoimmune hepatitis). No other safety signals have emerged based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

16.3.3.1.8. Immune-mediated/autoimmune AESIs

Search criteria\textsuperscript{101} - SMQ Immune-mediated/autoimmune disorders (Narrow and Broad) OR HLGT (All Path) Autoimmune disorders OR PTs Cytokine storm; Hypersensitivity; Thyroiditis subacute; Thrombocytopenia.

Clinical Trial Data

- Number of cases: 19 (BNT162b2 [17] and blinded therapy [2] (2.8% of 668 cases, the total CT dataset) compared to 20 cases (2.8%) retrieved in the PSUR #2\textsuperscript{93}.
- Country of incidence: US (14), Brazil (3), Argentina and China (1 each).
- Subjects' gender: female (7) and male (12).
- Subjects' age in years (n = 19), range: 6 – 79, mean 39.8, median 45.0.
- Medical history (n = 18): the relevant medical conditions reported more than once were coded to the PTs Dermatomyositis, Diabetes mellitus, Hypothyroidism, Seasonal allergy, and Type 1 diabetes mellitus (2 each).
- COVID-19 Medical history: None.
- Co-suspects: None.
- Number of relevant events: 19.

\textsuperscript{101} The PTs indicative of myocarditis and pericarditis have been moved from Immune-mediated/autoimmune AESIs to the newly added Section 16.3.3.1.11 Myocarditis and Pericarditis AESIs that is cross-referenced to Section 16.3.1.2 Important Identified Risks - Myocarditis and Pericarditis. The PT Thrombocytopenia has been reassigned from Haematological AESIs to Immune-mediated/autoimmune AESIs category.
• Reported relevant PTs (≥2 occurrences): Colitis (5), Diabetic ketoacidosis (3), Dermatomyositis, and Pancreatitis (2 each). All SAEs were assessed as not related to BNT162b2 or blinded therapy.

• Relevant event outcome: resolved/resolving (15), resolved with sequelae (1), not resolved (3).

Post-Authorisation Data

Number of cases: 11,729. Upon review, 3 cases were determined to be non-contributory and were not included in the discussion since these 3 cases involved exposures to the vaccine during the mothers' pregnancy or through breastfeeding.\textsuperscript{99}

• Number of cases: 11,726 (2.3% of 507,683 cases of the total PM dataset), compared to 21,994 cases (3.3%) retrieved in the PSUR #2\textsuperscript{99}.

• MC cases (4822), NMC cases (6904).

• Country of incidence: Germany (3094), France (1474), UK (1038), US (718), Italy (582), Japan (490), Australia (461), Netherlands (370), Austria (362), Sweden (250), Belgium (237), Norway (230), Greece (229), Finland (216), Poland, Spain (184 each), Taiwan, Province of China (156), Canada (145), New Zealand (125); the remaining 1181 cases were distributed among 64 countries.

• Subjects' gender: female (7678), male (3661), and unknown (387).

• Subjects' age in years (n = 10,827), range: 5 – 98, mean: 47.5, median: 47.0.

• Medical history (n = 4887): the most frequently (≥150 occurrences) reported relevant medical conditions were coded to the PTs Seasonal allergy (400), Asthma (378), Drug hypersensitivity (317), Hypersensitivity (306), Psoriasis (269), Hypothyroidism (252), Autoimmune thyroiditis (237), Food allergy (235), Diabetes mellitus (186), and Colitis ulcerative (158).

• COVID-19 Medical history (n = 507): the reported medical conditions were coded to the PTs COVID-19 (382), Suspected COVID-19 (115), COVID-19 pneumonia (11), Post-acute COVID-19 syndrome (6), Coronavirus infection (4), SARS-CoV-2 test positive (3), and Asymptomatic COVID-19 (2).

• Co-suspects (n = 460): the most frequently (≥10 occurrences) reported relevant co-suspects were adalimumab (168), COVID-19 vaccine MRNA (MRNA 1273) (36), influenza vaccine (26), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (19), influenza vaccine inact split 4V, and Risankizumab (13 each).

• Number of relevant events: 12,795.

• Relevant event seriousness:\textsuperscript{42} serious (8445) and non-serious (4356).
Most frequently reported relevant PTs (≥2%): Hypersensitivity (2393), Psoriasis (660), Thrombocytopenia (487), Polymyalgia rheumatica (431), Dermatitis (305), Rheumatic disorder (286), and Alopecia areata (281).

Time to event onset (n = 7591), \(^{102}\) range: <24 hours to 499 days, median: 6 days.

- <24 hours: 1451 events (5 of which had a fatal outcome);
- 1 day: 847 events (1 of which had a fatal outcome);
- 2-7 days: 1845 events (20 of which had a fatal outcome);
- 8-14 days: 924 events (9 of which had a fatal outcome);
- 15-30 days: 1032 events (18 of which had a fatal outcome);
- 31-181 days: 1348 events (23 of which had a fatal outcome);
- 182-499 days: 144 events (3 of which had a fatal outcome).

Duration of relevant events (n = 969 out of 2334 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 419 days, median 14 days.

- <24 hours: 145 events;
- 1 day: 68 events;
- 2-7 days: 186 events;
- 8-14 days: 90 events;
- 15-30 days: 106 events;
- 31-181 days: 311 events;
- 182-419 days: 63 events.

Relevant event outcome: fatal (133), resolved/resolving (3786), resolved with sequelae (664), not resolved at the time of reporting (4934), and unknown (3304).

- In 112 cases (reporting 133 relevant events with a fatal outcome), the reported causes of death (≥5 occurrences) were coded to the PTs Thrombocytopenia (19), Death, Interstitial lung disease (13 each), Haemophagocytic lymphohistiocytosis, Immune thrombocytopenia (8 each), Cerebral haemorrhage, Encephalitis, Multiple organ dysfunction syndrome, Renal failure (7 each), Pneumonia, Respiratory failure (6 each), and Pulmonary embolism (5). Of note, in 13 cases limited information regarding the cause of death was provided (PT Death). Most (78 of 104 cases that provided age) of the fatal cases involved elderly subjects. When the medical history was provided (67 cases), significant medical conditions reported in more than 3 cases included Hypertension (23), Atrial fibrillation (9), Osteoporosis (7), Dyslipidaemia (6), Diabetes mellitus, Hyperlipidemia, Type 2 diabetes mellitus (5 each), Hypothyroidism, Myocardial infarction, Radiotherapy, and Thrombocytopenia (4 each).

\(^{102}\) This number does not include 23 events for which partial administration and/or event onset dates were reported.
Analysis by age group

- CT: Paediatric (5), Adults (11), and Elderly (3).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM: Paediatric (591), Adults (8319), Elderly (2125) and Unknown (691).
  - Among the frequently (>2%) reported immune mediated/autoimmune AESIs, a higher reporting proportion of events coded to the PT Polymyalgia rheumatica were observed in the elderly population when compared to paediatric and adult populations (none in paediatrics vs 1.7% in adults vs 12.9% in elderly). A higher reporting proportion of events coded to the PTs Hypersensitivity and Alopecia areata were observed in the paediatric and adult populations when compared to the elderly population (Hypersensitivity [24.2% in paediatrics vs 20.8% in adults vs 11.8% in elderly], Alopecia areata [3.6% in paediatrics vs 2.7% in adults vs 0.9% in elderly]). A higher reporting proportion of events coded to the PTs Psoriasis and Rheumatic disorder were observed in the adult and elderly populations when compared to the paediatric population (Psoriasis [2.2% in paediatrics vs 5.8% in adults vs 5.9% in elderly], Rheumatic disorder [0.5% in paediatrics vs 2.3% in adults vs 3.4% in elderly]). A higher reporting proportion of events coded to the PT Thrombocytopenia were observed in the paediatric and elderly populations when compared to the adult population (8.8% in paediatrics vs 3.3% in adults vs 6.3% in elderly).

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 3199 (27.2% of the CT and PM cases reporting immune mediated/autoimmune AESIs).

The reporting proportion of immune mediated/autoimmune AESIs with a fatal outcome (2.6%) is higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subjects without comorbidities (0.6% of events with fatal outcome).

O/E Analysis

O/E analysis was performed for Acute disseminated encephalomyelitis (ADEM), ADEM and encephalitis, Autoimmune thyroiditis, Myasthenia gravis, Polymyalgia rheumatica, and Type 1 diabetes mellitus (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Polymyalgia rheumatica, Uveitis and Subacute Thyroiditis (SAT) were evaluated as signals in the reporting period and determined not to be risks (please refer to Section 16.2.1 Evaluation of Closed Signals).
No new safety signals have emerged based on a review of the remaining events and on O/E analysis. Safety surveillance will continue.

16.3.3.1.9. Multisystem Inflammatory Syndrome in Children / Adults

Search Criteria\(^{103}\): PTs Cytokine release syndrome; Distributive shock; Multiple organ dysfunction syndrome; Multisystem inflammatory syndrome; Multisystem inflammatory syndrome in adults; Multisystem inflammatory syndrome in children; Systemic inflammatory response syndrome.

Clinical Trial Data

- During the reporting period, no serious cases from the CT dataset were reported. For comparison, 2 cases (0.3%) were retrieved in the PSUR #2\(^{93}\).

Post-Authorisation Data

- Number of relevant cases: 207 (0.04% of 507,683 cases in the total PM dataset), compared to 438 (0.07%) retrieved in PSUR #2\(^{93}\).
- MC cases (170), NMC cases (37).
- Country of incidence (≥5 occurrences): France (55), Germany (27), UK (18), Australia (15), US (14), Japan (12), Norway (6), Spain (5); the remaining 55 cases were distributed among 30 countries.
- Subjects’ gender: female (92), male (109), unknown (6).
- Subjects’ age in years (n = 196), range: 3 – 95, mean: 46.6, median: 50.
- Medical history (n = 132): the most frequently (≥5 occurrences) reported medical conditions included the PTs Hypertension (40), Obesity (11), Diabetes mellitus (9), Ex-tobacco user, Hypothyroidism (8 each), Atrial fibrillation, Tobacco user, Type 2 diabetes mellitus (7 each), Alcohol use, Osteoporosis, Pyrexia, Sleep apnoea syndrome (6 each), Asthma, Non-tobacco user, Prostate cancer, and Rheumatoid arthritis (5 each).

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\(^{103}\) The MAH proposed to consider the PTs Cytokine release syndrome, Distributive shock, Multiple organ dysfunction syndrome, Multisystem inflammatory syndrome, Multisystem inflammatory syndrome in adults, Systemic inflammatory response syndrome as search strategy for the identification of potential cases of MIS-C/A going forward was endorsed by the PRAC as per EMA PRAC Assessment for the SBSR 2 (Product No. EMEA/H/C/005735/MEA/002.12). The other PTs included in the TME List and previously reviewed under the MIS-C/A AESIs category (Autoinflammatory disease, Cytokine storm, Haemophagocytic lymphohistiocytosis, Hypotensive crisis, Kawasaki’s disease, Vaccine associated enhanced disease, Vaccine associated enhanced respiratory disease) were re-assigned and are reviewed under a different AESIs category based on their clinical relevance; the remaining PTs previously reviewed under the MIS-C/A AESIs category but not included in the TME List (Macrophage activation, Macrophages increased, Septic shock, Toxic shock syndrome) will no longer be reviewed in the AESIs review.

Co-suspects (n = 16 cases): COVID-19 vaccine mRNA (mRNA 1273) (2), carboplatin, cefotaxime, ciclosporin, colchicine, COVID-19 vaccine, eltrombopag, enoxaparin, everolimus, mesalazine, methotrexate, pembrolizumab, pemtrexed, rituximab, treprostinil (1 each).

Number of relevant events: 210.

Relevant event seriousness: serious (210).

Relevant PTs: Multiple organ dysfunction syndrome (82), Multisystem inflammatory syndrome (43), Multisystem inflammatory syndrome in children (38), Systemic inflammatory response syndrome (32), Multisystem inflammatory syndrome in adults (10), Cytokine release syndrome (5).

Time to event onset (n = 115),\textsuperscript{104} range: \(< 24\) hours to 234 days, median: 15 days.

- \(< 24\) hours: 8 events (2 of which had a fatal outcome);
- 1 day: 5 events (1 of which had a fatal outcome);
- 2-7 days: 27 events (7 of which had a fatal outcome);
- 8-14 days: 13 events (2 of which had a fatal outcome);
- 15-30 days: 27 events (8 of which had a fatal outcome);
- 31-180 days: 33 events (11 of which had a fatal outcome);
- \(> 180\) days: 2 events.

Duration of relevant events (n = 12 out of 39 occurrences with outcome of resolved or resolved with sequelae), range: 3 days to 57 days, median: 16 days.

- 2-7 days: 5 events;
- 8-14 days: 1 event;
- 15-30 days: 2 events;
- 31-180 days: 4 events.

Relevant event outcome:\textsuperscript{105} fatal (57), resolved/resolving (61), resolved with sequelae (3), not resolved (20), unknown (72).

\textsuperscript{104} This number does not include 98 events for which administration and/or event onset dates were not provided or were incomplete. Please note, multiple episodes of the same PT event were reported with different latencies within some cases hence the sum of latencies exceeds the total number of PT events.

\textsuperscript{105} Multiple episodes of the same PT event were reported with a different clinical outcome within some cases hence the sum of the events outcome exceeds the total number of PT events.
COVID-19 mRNA vaccine (nucleoside modified)           Reporting Period
Periodic Safety Update Report (PSUR) 3          19 December 2021 through 18 June 2022

- In 56 cases (reporting 57 relevant events with fatal outcome), the reported causes of
death (≥5 occurrences) were coded to Multiple organ dysfunction syndrome (55), Septic
shock (10), Renal failure (9), Immunisation, Sepsis (8 each), Pneumonia (7), Acute
respiratory distress syndrome (6), Acute kidney injury, Cardiac arrest, COVID-19,
COVID-19 pneumonia, Drug ineffective, Hepatic failure, Respiratory failure, and
Vaccination failure (5 each).106 Most (35 of 56 cases) of the fatal cases involved elderly
subjects. When the medical history was provided (43 cases), the most
frequently (≥3 occurrences) medical conditions included Hypertension (19), Diabetes
mellitus, Obesity (6 each), COVID-19 (5), Atrial fibrillation, Ex-tobacco user,
Hypothyroidism, Osteoarthritis, Renal transplant, and Tobacco user (3 each).

Analysis by age group

- PM: Paediatric (46 [16 Child, 30 Adolescent]), Adult (84), Elderly (69), Unknown (8).

  - Among the relevant multisystem inflammatory syndrome events, it was observed that:

    - PT Multiple organ dysfunction syndrome was reported at a higher frequency in
      the elderly population compared to the adult and paediatric populations (62.3% of
      the elderly population vs 39.3% of the adult population and 4.3% of the paediatric
      population).
    - PT Multisystem inflammatory syndrome was reported at a higher frequency in
      the adult population compared to the elderly and paediatric populations (29.8% of
      the adult population vs 14.5% of the elderly population and 15.2% of the paediatric
      population).
    - PT Multisystem inflammatory syndrome in children was reported, as expected,
      primarily in the paediatric population (80.4% were in the paediatric population).
    - PT Systemic inflammatory response syndrome was reported at similar frequency
      in the adults and the elderly population (21.4% of the adult population vs 18.8%
      of the elderly population; no cases in paediatric population).
    - PT Multisystem inflammatory syndrome in adults was reported, as expected,
      primarily in the adult population and elderly population (9.5% of the adult
      population and 2.9% of the elderly population; no cases in paediatric population).
    - PT Cytokine release syndrome was observed only in the adult and elderly
      populations (2.4% in the adult population and 1.4% in the elderly population; no
      cases in paediatric population).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 73 (35.3% of the 207 cases reporting
Multisystem Inflammatory Syndrome AESIs).

106 A case may report multiple causes of death (i.e., other causes of death in addition to the fatal relevant
PT[s]).
• Of the PM cases that reported medical histories, the percentage of cases with a fatal Multisystem Inflammatory Syndrome AESI is higher in subjects with comorbid conditions (60.5%) when compared to the percentage of cases with a fatal Multisystem Inflammatory Syndrome AESI in subjects without comorbidities (39.5%).

• Upon review of the relevant events in PM cases that recorded medical histories, no Multisystem Inflammatory Syndrome AESIs had a significant proportional reporting ratio of >3:1 in subjects with comorbidities compared to subjects without comorbidities.

O/E Analysis

O/E analysis was performed for Multisystem inflammatory syndrome (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest). As in the most recent SBSR #3, the 21-24 years age group using the 21-day risk window meets the signal criteria with an O/E ratio >1, however, the result if not statistically significant as the 95% CI includes 1.

Conclusion

Cases of potential MIS in adults (MIS-A) and children (MIS-C) reported during this interval period are assessed in Appendix 6A.4). During the reporting period, an article including important safety information on MIS-C was reviewed. Please refer to Section 11 Literature for details.

No new safety signals have emerged based on a review of these cases, literature or of the O/E analysis. The MAH will continue to monitor MIS.

16.3.3.1.10. Musculoskeletal AESIs

Search Criteria: PTs Arthralgia; Arthritis; Chronic fatigue syndrome; Juvenile idiopathic arthritis\textsuperscript{107}; Polyarthritis; Post viral fatigue syndrome; Rhabdomyolysis; Rheumatoid arthritis.

Clinical Trial Data

• Number of cases: 6 (BNT162b2) (0.9% of 668 cases, the total CT dataset) compared to 4 cases (0.6%) retrieved in the PSUR #2\textsuperscript{93}.

• Country of incidence: US (6).

• Subjects' gender: female (3), male (3).

• Subjects' age in years (n = 6), range: 50-79, mean: 70.8, median: 74.0.

\textsuperscript{107} The PT Juvenile idiopathic arthritis has been moved from Immune-mediated/autoimmune AESIs to Musculoskeletal AESIs category.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

- Medical history (n=6): the most frequently (>1 occurrence) reported medical conditions included Hypertension (4), Insomnia (3), Anaemia (2), Arthritis (2), Gastroesophageal reflux disease (2), and Hyperlipidaemia (2).

- COVID-19 Medical history: None.

- Co-suspects: None.

- Reported relevant PTs (6): Arthritis (3), Arthritis (1), Rhabdomyolysis (1) and Rheumatoid arthritis (1), not related to BNT162b2.

- Relevant event outcome: resolved/resolving (5), not resolved (1).

Post-Authorisation Data

- Number of relevant cases: 31,012 (6.1% of 507,683 cases, the total PM dataset), compared to 58,250 cases (8.9 %) retrieved in the PSUR #2.

- MC cases (8400), NMC cases (22,612).

- Country of incidence (>105 occurrences): Netherlands (6107), Germany (4444), UK (2391), Belgium (1821), France (1733), Australia (1498), Iraq (1334), Austria (1248), Sweden (1114), Italy (986), Japan (893), Poland (722), US (653), Romania (620), Slovenia (613), Norway (606), Czech Republic (474), Finland (425), Spain (364), Denmark (358), Malaysia (296), Ireland (269), Portugal (215), Philippines (211), Canada (168), New Zealand (153), Lithuania (119), Taiwan, Province Of China (105); the remaining 1072 cases were distributed among 51 countries.

- Subjects' gender: female (22,130), male (8359) and unknown (523).

- Subjects' age in years (n = 29,340), range: 0.08-97 years, mean: 44.9, median: 44.

- Medical history (n = 7884 cases): the most frequently (> 100 occurrences) reported medical conditions included Disease risk factor (1020), Hypertension (874), Asthma (620), Seasonal allergy (479), Drug hypersensitivity (461), Hypersensitivity (389), Rheumatoid arthritis (323), Hypothyroidism (308), Food allergy (255), Diabetes mellitus (249), Fibromyalgia (213), Arthritis (190), Depression (187), Migraine (183), Osteoarthritis (177), Pain (173), Autoimmune thyroiditis (164), Immunodeficiency (157), Arthritis (154), Mite allergy (138), Type 2 diabetes mellitus (132), Non-tobacco user (126), Tobacco user (113), Anxiety (110), Psoriasis (109), Gastroesophageal reflux disease (104), Allergy to animal (103), Interchange of vaccine products (103), Hypercholesterolaemia (101).

Co-suspects (n = 478 cases): the frequently (>5 occurrences) reported co-suspect vaccines/medications included adalimumab (119), influenza vaccine (48), COVID-19 vaccine mRNA (mRNA 1273) (44), upadacitinib (38), COVID-19 vaccine NRVV AD (Chadorl NCOV-19) (25), COVID-19 vaccine (15), influenza vaccine inact split 3v (14), pneumococcal vaccine polysacchar 23v (11), etanercept (8), tocilizumab (8), influenza vaccine inact split 4v (7), ocrelizumab (7), influenza vaccine inact sag 4v (6).

- Number of relevant events: 31,633.
- Relevant event seriousness: 42 serious (6164), non-serious (25,510).
- Relevant PTs: Arthralgia (29,429), Arthritis (996), Rheumatoid arthritis (660), Chronic fatigue syndrome (219), Polyarthritis (145), Post viral fatigue syndrome (92), Rhabdomyolysis (78), Juvenile idiopathic arthritis (14).

- Time to event onset (n = 24,700\textsuperscript{108}), range: <24 hours to 3654 days, median: 0 days.
  - <24 hours: 8655 events (1 of which had a fatal outcome;
  - 1 day: 8837 events;
  - 2-7 days: 4004 events;
  - 8-14 days: 1105 events;
  - 15-30 days: 959 events;
  - 31-180 days: 1044 events;
  - 181-3654 days: 96 events.

- Duration of relevant events (n = 5620 out of 31,633 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 353 days, median 1 day.
  - <24 hours: 394 events;
  - 1 day: 1656 events;
  - 2 - 7 days: 2840 events;
  - 8-14 days: 211 events;
  - 15-31 days: 157 events;
  - 32-181 days: 286 events;
  - 182-353 days: 89 events.

- Relevant event outcome (31,815): fatal (22), resolved/resolving (14,443), resolved with sequelae (512), not resolved (12,142), unknown (4696).

In 22 cases (reporting 22 relevant events with fatal outcome), the reported causes of death were coded to the PTs Arthralgia (14), Rhabdomyolysis (5), Pyrexia (2), Arthritis (1), Inflammation (1), Polyarthritus (1), and Rheumatoid arthritis (1). Most (14 of 22 cases) of

\textsuperscript{108} This number does not include 40 events for which partial administration and/or event onset dates were reported or events with a not meaningful time to onset value as per reported information.
the fatal cases involved elderly subjects. When the medical history was provided (18 cases), the most frequently (≥ 2 occurrences) relevant medical conditions included Hypertension (9), Parkinson’s disease (3), Cerebrovascular accident (2), Coronary arterial stent insertion (2), Diabetes mellitus (2), Myocardial infarction (2), Osteoarthritis (2).

Analysis by age group

- CT: Adult (1) and Elderly (5).
- PM: Paediatric (664), Adult (25,307), Elderly (3469), Unknown (1572).
  - Higher reporting proportion of events coded to the PT Rheumatoid arthritis was reported in the elderly population when compared to adult and paediatric population ([1.8 % in adults vs 0.8% in paediatrics vs 4.6 % in elderly]).

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 3193 (10.3% of the cases reporting musculoskeletal AESIs).
- A higher reporting proportion of musculoskeletal AESIs was reported in subjects without significant comorbidities 27,825 (89.7%) when compared to subjects with significant comorbidities.
- The reporting proportion of musculoskeletal AESIs with outcome resolved (91.4%) is higher in subjects without comorbid conditions when compared to the reporting proportion observed in the subjects with comorbidities (8.6% of events with resolved).

O/E Analysis

O/E analysis was performed for Chronic fatigue syndrome/MED/PVFS, Rhabdomyolysis, Rheumatoid arthritis, polyarthritis, juvenile idiopathic arthritis (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. The majority of the events reported in this category are arthralgia which is considered to be an adverse reaction for the vaccine and is labelled as such. Arthralgia will be removed from the search strategy in the next PSUR. Safety surveillance will continue.
16.3.3.1.11. Myocarditis and Pericarditis AEsIs\textsuperscript{109}

Please refer to the Risk ‘Myocarditis and Pericarditis’ in Section 16.3.1.2 Important Identified Risks – Myocarditis.

16.3.3.1.12. Neurological AEsIs (including demyelination)

Search Criteria\textsuperscript{110}: SMQ Generalised convulsive seizures following immunisation (Narrow) OR SMQ Demyelination (Narrow and Broad) OR PTs Ataxia; Cataplexy; Fibromyalgia; Intracranial pressure increased; Meningitis; Meningitis aseptic; Narcolepsy; Neuropathy peripheral; Polyneuropathy.

Clinical Trial Data

- Number of cases: 15 cases (BNT162b2 [11], blinded therapy [4]; 2.2% of 668 cases in the total CT dataset) compared to 7 cases (0.97%) retrieved in the PSUR #2\textsuperscript{93}.
- Country of incidence: US (10), Brazil, Poland (2 each), Argentina (1).
- Subjects’ gender: female (7), male (8).
- Subjects’ age in years (n = 15), range: 0.58 – 79, mean: 36.1, median: 43.
- Medical history (n = 9): medical conditions reported more than once were coded to the PTs Insomnia (3), Blood cholesterol increased, Hypertension, and Seizure (2 each).
- COVID-19 medical history: None.
- Co-suspects: None.
- Reported relevant PTs: Seizure (6), Febrile convulsion (3), Meningitis, Myelitis transverse, Optic neuritis, Polyneuropathy, Toxic leukoencephalopathy (1 each). However, none of these SAEs were assessed as related to BNT162b2/blinded therapy.
- Relevant event outcome: resolved/resolving (12), resolved with sequelae (2), not resolved (2).

Post-Authorisation Data

- Number of relevant cases: 5111 (1.0% of 507,683 cases in the total PM dataset), compared to 7197 cases (1.1%) retrieved in the PSUR #2\textsuperscript{93}.

\textsuperscript{109} The PTs indicative of myocarditis and pericarditis have been moved from Immune-mediated/autoimmune AEsIs to this newly added AEsIs.

\textsuperscript{110} The SMQ Convulsions (Narrow and Broad) has been replaced with the SMQ Generalized convulsive seizures following immunisation (Narrow) to be more vaccine-focused.
• MC cases (2245), NMC cases (2866).

• Country of incidence (> 56 occurrences): Germany (1258), France (552), UK (455), Italy (288), US (248), Australia (240), Japan (227), Austria (178), Poland (177), Netherlands (158), Finland (96), Norway (92), Sweden (81), Canada (79), Philippines (78), New Zealand (76), Spain (74), Greece (65), Taiwan (64), Belgium, Ireland (60 each), Czech Republic (57); the remaining 448 cases were distributed among 51 countries.

• Subjects' gender: female (3163), male (1810), unknown (138).

• Subjects' age in years (n = 4811), range: 2.33 – 100, mean: 44.0, median: 44.

• Medical history (n = 2493): the most frequently (>51 occurrences) reported medical conditions included the PTs Hypertension (333), Epilepsy (253), Multiple sclerosis (241), Seasonal allergy (172), Drug hypersensitivity (160), Fibromyalgia (156), Asthma (150), Depression (98), Hypothyroidism (92), Food allergy, Hypersensitivity (91 each), Seizure (82), Diabetes mellitus (81), Migraine (62), Mite allergy (61), Obesity (60), Type 2 diabetes mellitus (54), and Pain (52).


• Co-suspects (n = 150 cases): the reported co-suspect medications (≥3 occurrence) were ocrelizumab (15), COVID-19 vaccine mRNA (mRNA 1273) (12), adalimumab (11), COVID-19 vaccine NRVV AD, influenza vaccine INACT SPLIT 4V (10 each), influenza vaccine (9), COVID-19 vaccine (6), apixaban, levetiracetam, teriflunomide (4 each), cannabidiol/dronabinol, lamotrigine, and natalizumab (3 each).

• Number of relevant events: 5501.

• Relevant event seriousness:69 serious (4973), non-serious (530).

• Most frequently (>58 occurrences) reported relevant PTs: Seizure (1282), Epilepsy (540), Neuropathy peripheral (528), Guillain-Barre syndrome (524), Trigeminal neuralgia (341), Fibromyalgia (314), Multiple sclerosis (265), Polyneuropathy (250), Multiple sclerosis relapse (199), Optic neuritis (182), Generalised tonic-clonic seizure (175), Ataxia (127), Myelitis transverse (90), Meningitis (79), Febrile convulsion (76), Demyelination, Intracranial pressure increased (59 each).
Time to event onset (n = 3717),\(^{111}\) range: <24 hours to 391 days, median: 3 days.

- <24 hours: 883 events (7 of which had a fatal outcome);
- 1 day: 579 events (5 of which had a fatal outcome);
- 2-7 days: 862 events (12 of which had a fatal outcome);
- 8-14 days: 383 events (7 of which had a fatal outcome);
- 15-30 days: 414 events (6 of which had a fatal outcome);
- 31-180 days: 557 events (12 of which had a fatal outcome);
- >180 days: 39 events.

Duration of relevant events (n = 619 out of 1404 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 330 days, median 1 day.

- <24 hours: 269 events;
- 1 day: 60 events;
- 2-7 days: 102 events;
- 8-14 days: 42 events;
- 15-30 days: 43 events;
- 31-180 days: 88 events;
- >180 days: 15 events.

Relevant event outcome: fatal (67), resolved/resolving (1890), resolved with sequelae (306), not resolved (1839), unknown (1418).

In 61 cases (reporting 67 relevant events with fatal outcome), the reported causes of death (>3 occurrences) were coded to the PTs Seizure (34), Guillain-Barre syndrome (11), Headache (8), Cardiac arrest, Off label use (7 each), Epilepsy (6), Dyspnoea, Interchange of vaccine products, Pneumonia (5 each), Intracranial pressure increased, Loss of consciousness, Pyrexia, and Sudden death (4 each).\(^{106}\) Over half (31 of 61 cases) of the fatal cases involved elderly subjects. When the medical history was provided (33 cases), the most frequent (>3 occurrences) medical conditions included the PTs Hypertension (14), Cardiac failure, COVID-19, Diabetes mellitus (4 each), Atrial fibrillation, Chronic obstructive pulmonary disease, and Seizure (3 each).

Analysis by age group

- CT: Paediatric (Infant [2], Child [4]), Adult (6), Elderly (3).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

\(^{111}\) This number does not include 1808 events for which administration and/or event onset dates were not provided or were incomplete; or events without a meaningful time to onset value as per reported information. Please note, multiple episodes of the same PT event were reported with different latencies within some cases hence the sum of latencies exceeds the total number of PT events.
PM: Paediatric (523 [162 Child, 361 Adolescent]), Adult (3574), Elderly (787), Unknown (227).

- Among the most frequently (>50 occurrences) reported relevant neurological events, it was observed that:
  - The PTs Seizure, Generalised tonic-clonic seizure, and Febrile convulsion were reported at higher frequencies in the paediatric population compared to the adult population and the elderly population (57.9%, 7.5%, and 6.1% of the paediatric population vs 22.0%, 3.2%, and 0.8% of the adult population, and 15.5%, 2.2%, and 1.7% of the elderly population, respectively). This pattern is consistent with the known epidemiology of seizures and epilepsy.
  - The PTs Guillain-Barre syndrome, Polynuropathy, and Ataxia were reported at higher frequencies in the elderly population compared to the paediatric population and the adult population (17.2%, 8.4%, and 4.7% of the elderly population vs 7.1%, 1.3%, and 1.1% of the paediatric population, and 9.0%, 4.7%, and 2.3% of the adult population, respectively).
  - The PTs Neuropathy peripheral, Trigeminal neuralgia, and Fibromyalgia were reported at higher frequencies in the adult population and the elderly population compared to the paediatric population (11.3%, 7.9%, and 7.2% of the adult population and 11.3%, 5.8%, and 5.3% of the elderly population vs 1.3%, 0.0%, and 0.4% of the paediatric population, respectively).
  - The PTs Multiple sclerosis, Multiple sclerosis relapse, and Optic neuritis were reported at higher frequencies in the adult population compared to the paediatric population and the elderly population (6.4%, 4.8%, and 4.4% of the adult population vs 1.0%, 0.6%, and 2.3% of the paediatric population, and 1.8%, 2.0%, and 1.4% of the elderly population, respectively).

**Analysis by presence of comorbidities**

- Number of PM subjects reporting comorbidities: 1201 (23.5% of the 5111 cases reporting Neurological AESIs).

- Of the PM cases that reported medical histories, the percentage of cases with a fatal Neurological AESI is higher in subjects with comorbid conditions (71.9%) when compared to the percentage of cases with a fatal Neurological AESI in subjects without comorbidities (28.1%).

Upon review of the most frequent (>50 occurrences) relevant events in PM cases that recorded medical histories, the PTs Multiple sclerosis and Multiple sclerosis relapse were the only PTs that had a significant proportional reporting ratio of >3:1 in subjects with comorbidities compared to subjects without comorbidities.
O/E Analysis

O/E analysis was performed for Generalized convulsive, Fibromyalgia, Guillain-Barré syndrome, Meningitis, Narcolepsy, Multiple sclerosis (MS) and Polyneuropathy (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

16.3.3.1.13. Other AESIs

Search Criteria\textsuperscript{112}: HLT (All Path) Herpes viral infections OR PTs Adverse event following immunisation; Appendicectomy; Appendicitis; Appendiceal abscess; Appendicitis perforated; Complicated appendicitis; Deafness; Deafness bilateral; Deafness neurosensory; Deafness permanent; Deafness transitory; Deafness unilateral; Hypoacusis; Inflammation; Manufacturing laboratory analytical testing issue; Manufacturing materials issue; Manufacturing production issue; MERS-CoV test; MERS-CoV test negative; MERS-CoV test positive; Occupational exposure to communicable disease; Patient isolation; Product availability issue; Product distribution issue; Product supply issue; Pyrexia; Quarantine; SARS-CoV-1 test; SARS-CoV-1 test negative; SARS-CoV-1 test positive; Sudden hearing loss.

Upon review, 245 PM cases were determined to be non-contributory and were not included in the discussion since these 245 cases involved exposures to the vaccine during the mother’s pregnancy or through breastfeeding.\textsuperscript{99}

Clinical Trial Data

\begin{itemize}
\item Number of cases: 26 (BNT162b2 [22], blinded therapy [3] and placebo [1]) (3.9\% of 668 cases, the total CT dataset) compared to 2 cases (0.28\%) retrieved in the PSUR #2\textsuperscript{93}.
\item Country of incidence: US (14), Argentina (5), Germany (3), Brazil, China, Finland, and Poland (1 each).
\item Subjects’ gender: female (14), male (12).
\item Subjects’ age in years (n = 26), range: 23 months – 79 years, mean: 25.5, median: 22.0
\item Relevant Medical history: Appendicectomy (2) and Wiskott-Aldrich syndrome (1)
\end{itemize}

\textsuperscript{112} The PTs Appendicectomy; Appendicitis; Appendiceal abscess; Appendicitis perforated; Complicated appendicitis; Deafness; Deafness bilateral; Deafness neurosensory; Deafness permanent; Deafness transitory; Deafness unilateral; Hypoacusis; Sudden hearing loss have been added. Hearing loss and Appendicitis were reviewed as signals during the reporting period (see Appendix 6A.3 and Section 15).
COVID-19 Medical history: None.

Co-suspects: None.

Reported relevant PTs: Appendicitis (14), Pyrexia (6), Deafness neurosensory (2), Complicated appendicitis, Exanthema subitem, Herpes zoster, and Sudden hearing loss (1 each). None of the SAEs were assessed as related to BNT162b2 or blinded therapy or placebo.

Relevant event outcome: resolved/resolving (23), not resolved (3).

Post-Authorisation Data

- Number of cases: 68,548 (13.5% of 507,683 cases, the total PM dataset), compared to 118,843 cases (18.1%) retrieved in the PSUR #293.

- MC cases (25,353), NMC cases (43,195).

- Country of incidence (≥100 occurrences): Germany (14,740), Netherlands (5318), Iraq (4917), Japan (3834), UK (3932), Spain (2839), France (2686), Australia (2634), Poland (2374), Italy (2362), Sweden (2297), Philippines (2234), Belgium (2144), Romania (1847), Austria (1829), Malaysia (1620), Egypt (1468), US (1282), Norway (1013), Finland (833), Denmark (747), Czech Republic (640), Taiwan, Province of China (534), Ireland (538), Lithuania (445), Greece (377), Portugal (375), Croatia (347), Canada (313), Switzerland (298), Georgia (285), New Zealand (279), Brazil (276), Estonia (193), Mexico (138), and Hungary (107); the remaining 893 cases were distributed among 50 countries.

- Subjects' gender: female (45,066), male (21,639) and unknown (1843).

- Subjects' age in years (n = 63640), range: 2 days -104 years, mean: 41.1, median: 39.

- Medical history (n = 15037): the most frequently (≥50 occurrences) relevant medical conditions included Immunodeficiency (242), Herpes zoster (176), Breast cancer (155), Neoplasm malignant (70).

- COVID-19 Medical history (n = 3,535): the most frequently (≥10 occurrences) reported medical conditions included COVID-19 (2452), Suspected COVID-19 (938), Post-acute COVID-19 syndrome (59), COVID-19 pneumonia (33), Coronavirus infection (24), Asymptomatic COVID-19, SARS-CoV-2 test positive (10 each), Exposure to SARS-CoV-2 (6), Coronavirus test positive, Occupational exposure to SARS-CoV-2, and SARS-CoV-2 antibody test positive (1 each).

- Co-suspects (n = 677): the reported relevant co-suspect medications were Adalimumab (64), Methotrexate (5) Apixaban (4), Etanercept (3), Rituximab (2), Infliximab (1).

- Number of relevant events: 69,859.
• Relevant event seriousness: serious (10,676), non-serious (59,208).

• Most frequently reported relevant PTs (≥ 50 occurrences): Pyrexia (57,474), Herpes zoster (6216), Inflammation (1585), Oral herpes (794), Hypoacusis (744), Deafness (488), Sudden hearing loss (370), Herpes virus infection (297), Appendicitis (254), Deafness unilateral, Ophthalmic herpes zoster (222 each), Herpes simplex (206), Adverse event following immunisation (188), Genital herpes (163), Herpes ophthalmic (70), Deafness neurosensory (63), Herpes zoster oticus (61), Herpes zoster reactivation, and Varicella (59 each).

• Time to event onset (n = 55,778),\(^{113}\) range: <24 hours to 180 days, median: 1 day.
  - <24 hours: 21,103 events (17 of which had a fatal outcome);
  - 1 day: 20491 events (25 of which had a fatal outcome);
  - 2-7 days: 6833 events (27 of which had a fatal outcome);
  - 8-14 days: 2154 events (14 of which had a fatal outcome);
  - 15-30 days: 2229 events (8 of which had a fatal outcome);
  - 31-180 days: 2968 events (20 of which had a fatal outcome).

• Duration of relevant events (n = 19,717 out of 70,158 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 179 days, median 1 day.
  - <24 hours: 2487 events;
  - 1 day: 7397 events;
  - 2 - 7 days: 8026 events;
  - 8-14 days: 724 events;
  - 15-31 days: 547 events;
  - 32-181 days: 536 events.

• Relevant event outcome:\(^{78}\) fatal (166), resolved/resolving (43,472), resolved with sequelae (855), not resolved (13,677), unknown (11,880).

• In 164 cases (reporting 166 relevant events with fatal outcome), the reported causes of death (>10 occurrences) were coded to the PTs Pyrexia (118), Adverse event following immunisation (27), and Inflammation (11). Most (82 of 164 cases) of the fatal cases involved elderly subjects. When the medical history was provided (151 cases), the most frequently (≥ 20 occurrences) relevant medical conditions included Hypertension (52) and Diabetes mellitus (27).

\(^{113}\) This number does not include 13,972 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.
Analysis by age group

CT: Adult (14), Paediatric (10), Elderly (2).

- A meaningful comparison between the different age groups is not possible due to the low number of cases.

PM: Paediatric (5092), Adult (53,918), Elderly (6771), and Unknown (2767).

- Among the frequently (≥2%) reported relevant Other AESI events, PTs Herpes zoster and Inflammation were reported significantly higher in elderly population when compared to adult population (Herpes zoster [37% in adults vs 58.7% in elderly], Inflammation [0.9% in adult vs 2.7% in elderly]). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 5215 (7.6% of the cases reporting other AESI). A higher reporting proportion of other AESIs was reported in population without significant comorbidities (92.4%) when compared to population with significant comorbidities.

- The reporting proportion of other AESIs with fatal outcome (0.9%) is higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subjects without comorbidities (0.2% of events with fatal outcome).

O/E Analysis

O/E analysis was performed on Appendicitis, Herpes zoster and Sudden hearing loss (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Hearing loss was evaluated as a signal during the reporting period (please refer to Appendix 6A.3 for cumulative review of cases indicative of hearing loss).

Appendicitis was evaluated as signal during the reporting period (please refer to Section 16.2.1 Evaluation of Closed Signals).

No other safety signals than those mentioned have emerged based on the review of these cases, or from the O/E analysis. No risks have been identified following the evaluations of appendicitis and hearing loss. Safety surveillance will continue.

16.3.3.1.14. Pregnancy related AESIs

Search criteria - PTs Amniotic cavity infection; Caesarean section; Congenital anomaly; Death neonatal; Eclampsia; Foetal distress syndrome; Low birth weight baby; Maternal
exposure during pregnancy; Placenta praevia; Pre-eclampsia; Premature labour; Renal failure neonatal; Renal impairment neonatal; Stillbirth; Uterine rupture; Vasa praevia.

For relevant cases, please refer to the Section 16.3.5.3 Use in Pregnant/Lactating Women.

16.3.3.1.15. Glomerulonephritis and Nephrotic Syndrome AESIs

Search criteria - HLT Glomerulonephritis and nephrotic syndrome.

Upon review, 2 cases were determined to be noncontributory and were not included in the discussion since these cases involved exposures to the vaccine during the mother’s pregnancy or through breastfeeding.

Clinical Trial Data

- During the reporting period no serious cases from the CT dataset were reported. No comparison with PSUR #2 is possible due to the change in the search criteria.

Post-Authorisation Data

- Number of cases: 276 (0.05% of 507,683 cases, the total PM dataset). No comparison with PSUR #2 is possible.
- MC cases (172), NMC cases (104).
- Country of incidence: Germany (74), Japan (50), France (29), Australia (13), Italy, UK (11 each); the remaining 88 cases were distributed among 28 countries.
- Subjects' gender: female (150), male (124) and unknown (2).
- Subjects' age in years (n = 270), range: 5 – 88, mean: 44.2, median: 43.0.
- Medical history (n = 148): the most frequently (≥5 occurrences) reported relevant medical conditions included Hypertension (25), Nephrotic syndrome (12), Hypercholesterolaemia (8), Dyslipidaemia, Glomerulonephritis, Haematuria, IgA nephropathy (7 each), Proteinuria (6).
- Co-suspects (n= 3 cases): the reported relevant co-suspect medications included Hepatitis A vaccine, Influenza vaccine, and Tocilizumab (1 each).
- Number of relevant events: 323
- Relevant event seriousness: serious (318), non-serious (5).

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114 The PTs Acute kidney injury and Renal failure have been removed from the search criteria and replaced with a more focused search of glomerulonephritis and nephrotic syndrome based on the evolving pharmacovigilance and medical literature. An evaluation of IgA nephropathy, as requested by EMA in the PSUR 2 assessment report is ongoing and will be provided to EMA under separate cover from the PSUR.
Most frequently reported relevant PTs: Nephrotic syndrome (99), IgA nephropathy (47), Glomerulonephritis (46), Glomerulonephritis minimal lesion (25), Granulomatosis with polyangiitis (22), Microscopic polyangiitis (14), Glomerulonephritis membranous (12), Focal segmental glomerulosclerosis, and Glomerulonephritis rapidly progressive (10 each).

- Time to event onset (n = 172),\textsuperscript{115} range: <24 hours to 172 days, median: 12 days.
  - <24 hours: 7 events;
  - 1 day: 22 events;
  - 2-7 days: 43 events (1 of which had a fatal outcome);
  - 8-14 days: 23 events;
  - 15-30 days: 32 events;
  - 31-180 days: 45 events (1 of which had a fatal outcome).

- Duration of relevant events (n = 12 out of 323 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 137 days, median 48 days.
  - <24 hours: 1 event;
  - 1 day: 0 events;
  - 2-7 days: 1 event;
  - 8-14 days: 0 events;
  - 15-31 days: 1 event;
  - 32-181 days: 9 events.

- Relevant event outcome: fatal (2), resolved/resolving (95), resolved with sequelae (23), not resolved (111), unknown (92).

- In 2 cases (reporting 2 relevant events with fatal outcome), the reported causes of death were coded to Glomerulonephritis and Granulomatosis with polyangiitis (1 each). Both fatal cases involved elderly subjects. Medical history was provided in both cases and included Autoimmune hypothyroidism, Hypertension and Obesity (1 each).

### Analysis by age group

PM: Paediatric (33), Adult (177), Elderly (62) and Unknown (4).

- Among the frequently (>2%) reported relevant events Glomerulonephritis and Nephrotic Syndrome AESIs, the PT Renal failure was reported significantly higher in elderly population when compared to adult population (2.8% in adults vs 8.4% in elderly). A higher reporting proportion of events coded to the PTs Haematuria and IgA nephropathy were observed in the adult population when compared to the elderly population (Haematuria [10.4% in adults vs 2.1% in elderly] and IgA nephropathy

\textsuperscript{115} This number does not include 151 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.
[11.1% in adults vs 1.1% in elderly]). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 14 (5.1% of the cases reporting Glomerulonephritis and nephrotic syndrome AESIs).

- The reporting proportion of Glomerulonephritis and nephrotic syndrome AESIs with fatal outcome is 0.4% in subjects without comorbid conditions. There were no fatal outcomes in the subjects with comorbidities.

O/E Analysis

O/E analysis was performed for Glomerulonephritis/nephrotic syndrome (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

Please refer to Section 15 Overview of Signals: New, Ongoing, or Closed and to Appendix 6A for the response to the PRAC request (EMA/PRAC/416198/2021 – EPITT 19722).

No safety signals have emerged based on the review of these cases, or from the O/E analysis. The ongoing evaluation of IgA nephropathy will be submitted separately from the PSUR. Safety surveillance will continue.

16.3.3.1.16. Respiratory AESIs

Search criteria - HLTs (All Path) Lower respiratory tract infections NEC; Respiratory failures (excl neonatal); Viral lower respiratory tract infections OR PTs Acute respiratory distress syndrome; Endotracheal intubation; Hypoxia; Respiratory disorder.

Upon review, 4 cases were determined to be non-contributory and were not included in the discussion since these cases involved exposures to the vaccine during the mother’s pregnancy or through breastfeeding.

Clinical Trial Data

- Number of cases: 33 (Blinded therapy [10], BNT162b2 [23]) (4.9% of 668 cases, the total CT dataset) compared to 38 cases (5.3%) retrieved in the PSUR #2.

- Country of incidence: US (24), Argentina (4), Poland, Spain (2 each), and Brazil (1).

- Subjects' gender: female (15), male (18).

- Subjects’ age in years (n = 33), range: 17 months – 81 years, mean: 49.3, median: 63.0.
• Medical history (n = 25): the relevant medical conditions included Chronic obstructive pulmonary disease (4), Seasonal allergy (3), Asthma, Bronchitis chronic, Lung neoplasm malignant, and Upper respiratory tract infection (1 each).

• COVID-19 medical history (n = 1): COVID-19 (1).

• Co-suspects: None.

• Reported relevant PTs (35): Pneumonia (15), Acute respiratory failure (6), Bronchitis (4), Hypoxia (3), Lower respiratory tract infection, Metapneumovirus infection (2 each), Cardio-respiratory arrest, Respiratory failure, and Respiratory syncytial virus bronchitis (1 each).

• Relevant event outcome: fatal (4), resolved/resolving (30), not resolved (1).

• Of the above SAEs, all were assessed as not related to BNT162B2 or Blinded therapy.

Post-Authorisation Data

• Number of cases: 2188 (0.4% of 507,683 cases, the total PM dataset), compared to 3356 cases (0.51%) retrieved in the PSUR #2.

• MC cases (186), NMC cases (1002).

• Country of incidence: Germany (374), France (311), UK (193), Japan (156), Australia (145), Belgium (108), US (106), Italy (102), Austria (82), Spain (66), Philippines (58); the remaining 487 cases were distributed among 43 countries.

• Subjects' gender: female (1189), male (948) and unknown (51).

• Subjects' age in years (n = 2064), range: 5 - 106, mean: 56.4, median: 58.0

• Medical history (n = 1168): the most frequently (≥5 occurrences) reported medical conditions included Asthma (140), Chronic obstructive pulmonary disease (70), Seasonal allergy (42), Pneumonia (29), Pulmonary embolism (20), Sleep apnoea syndrome (14), Bronchitis, Emphysema (11 each), Chronic respiratory failure, Lung disorder (8 each), Bronchiectasis, Lower respiratory tract infection, Obstructive sleep apnoea syndrome, Pulmonary fibrosis (7 each), Bronchitis chronic, and Respiratory disorder (6 each).

• COVID-19 Medical history (n = 123): the most frequently reported medical conditions included COVID-19 (88), Suspected COVID-19 (24), COVID-19 pneumonia (6), Post-acute COVID-19 syndrome (2), Coronavirus infection, Exposure to SARS-CoV-2, SARS-CoV-2 test positive (1 each).

• Co-suspects (n = 138 cases): the reported relevant co-suspect medications included Adalimumab (25), Rituximab (3), Casirivimab, Imdevimab (2), Atenolol, Bromazepam, Durvalumab, Methotrexate, Salbutamol, and Terbutaline (1 each).

• Number of relevant events: 2383.

• Relevant event seriousness: serious (1873), non-serious (510).
• Most frequently reported relevant PTs (≥ 100 occurrences): Pneumonia (809), Respiratory disorder (325), Bronchitis (303), Respiratory failure (213), Lower respiratory tract infection (175), Cardio-respiratory arrest (140), and Hypoxia (133).

• Time to event onset (n = 1422)\textsuperscript{116} range: <24 hours to 437 days, median: 5 days.
  - <24 hours: 216 events (21 of which had a fatal outcome);
  - 1 day: 230 events (52 of which had a fatal outcome);
  - 2-7 days: 330 events (50 of which had a fatal outcome);
  - 8-14 days: 170 events (29 of which had a fatal outcome);
  - 15-30 days: 146 events (27 of which had a fatal outcome);
  - 31-180 days: 272 events (44 of which had a fatal outcome)
  - 181-437 days: 58 events (10 of which had a fatal outcome).

• Duration of relevant events (n = 176 out of 2395 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 288 days, median 7 days.
  - <24 hours: 26 events;
  - 1 day: 24 events;
  - 2-7 days: 43 events;
  - 8-14 days: 29 events;
  - 15-30 days: 29 events
  - 31-180 days: 21 events;
  - 181-288 days: 4 events.

• Relevant event outcome:\textsuperscript{78} fatal (363), resolved/resolving (694), resolved with sequelae (53), not resolved (544), unknown (738).
  - In 318 cases (reporting 363 relevant events with fatal outcome), the reported causes of death (≥20 occurrences) were coded to the PTs Cardio-respiratory arrest (99), Pneumonia (74), Respiratory failure (63), Acute respiratory failure (31), Hypoxia (26), and Acute respiratory distress syndrome (22). Most (235 of 318 cases) of the fatal cases involved elderly subjects. When the medical history was provided (234 cases), the most frequently (≥ 20 occurrences) relevant medical conditions included Hypertension (104), Atrial fibrillation (39), Diabetes mellitus (27), Type 2 diabetes mellitus (25), Chronic obstructive pulmonary disease, and Dyslipidaemia (23 each).

\textsuperscript{116} This number does not include 973 events for which partial administration and/or event onset dates were reported or events with a not meaningful time to onset value as per reported information.
Analysis by age group

- CT: Paediatric (8), Adult (12) and Elderly (13).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM: Paediatric (83), Adult (1168), Elderly (836) and Unknown (101).
  - A higher reporting proportion of events coded to the PTs Acute respiratory distress syndrome and Respiratory failure was observed in elderly population when compared to the adult population (Acute respiratory distress syndrome [5.7% vs 1.3%], Respiratory failure [12.7% vs 4.6%]. Additionally, a higher reporting proportion of events coded to the PTs Cardio-respiratory arrest and Respiratory disorder was observed in the adult population when compared to elderly (Cardio-respiratory arrest 10% vs 1.4% and Respiratory disorder 15.2% vs 5.6%). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

Analysis by presence of comorbidities

- Number of subjects reporting comorbidities: 113 (5.16 % of the cases reporting respiratory AESIs).

- The reporting proportion of respiratory events with a fatal outcome (17.4 %) is higher in individuals with comorbid conditions when compared to the reporting proportion observed in the individuals without comorbidities (15.1 % of events with resolved).

O/E Analysis

O/E analysis was performed for Acute respiratory distress syndrome (ARDS) (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue. Respiratory events were originally included in the AESI list in order to capture potential cases of respiratory failure that may occur in cases of severe COVID-19. The search strategy will be amended to focus on acute respiratory distress syndrome and respiratory failure for the next PSUR.

16.3.3.1.17. Stroke

Search criteria - HLT Central nervous system haemorrhages and cerebrovascular accidents (All path); Cerebrovascular venous and sinus thrombosis (Primary Path).
Clinical Trial Data

- Number of cases: 19 cases (BNT162b2 [18], blinded therapy [1]; 2.8% of 668 cases in the total CT dataset) compared to 19 cases (2.6%) retrieved in the PSUR #2.

- Country of incidence: US (15), Argentina (2), Brazil, China (1 each).

- Subjects' gender: female (5), male (14).

- Subjects' age in years (n = 19), range: 36 – 85, mean: 67.4, median: 71.

- Medical history (n = 18): medical conditions reported more than twice were coded to the PTs Hypertension (12), Type 2 diabetes mellitus (5), Hypercholesterolaemia (4), Obesity, and Osteoarthritis (3 each).

- COVID-19 Medical history: None.

- Co-suspects (n= 1 case): amlodipine, metoprolol (1 each).

- Reported relevant PTs: Cerebrovascular accident (13), Ischaemic stroke (3), Cerebral infarction, Embolic stroke, Haemorrhagic stroke (1 each). However, none of these SAEs were assessed as related to BNT162b2/blinded therapy.

- Relevant event outcome: fatal (1), resolved/resolving (12), resolved with sequelae (6).

Post-Authorisation Data

- Number of cases: 3091 (0.6% of 507,683 cases in the total PM dataset), compared to 4834 cases (0.7%) retrieved in the PSUR #2.

- MC cases (1418), NMC cases (1673).

- Country of incidence (>50 occurrences): Germany (1011), France (353), UK (219), Japan (143), Austria (138), Australia (127), US (116), Italy (108), Sweden (89), Taiwan (83), Netherlands (81), Poland (57), Denmark (51); the remaining 515 cases were distributed among 40 countries.

- Subjects' gender: female (1555), male (1467), unknown (69).

- Subjects' age in years (n = 2915), range: 5 – 101, mean: 60.8, median: 62.

- Medical history (n = 1593): the most frequently (>55 occurrences) reported medical conditions were coded to the PTs Hypertension (590), Diabetes mellitus (119), Atrial fibrillation (111), Tobacco user (98), Type 2 diabetes mellitus (96), Obesity (86), Cerebrovascular accident (83), Dyslipidaemia, Seasonal allergy (74 each), Non-tobacco user (71), Hypercholesterolaemia (63), Asthma (58), Drug hypersensitivity, and Hypothyroidism (56 each).


- Co-suspects (n = 96 cases): Most frequently (>3 occurrences) reported co-suspect medications were COVID-19 vaccine mRNA (mRNA 1273) (10), influenza vaccine, influenza vaccine INACT SPLIT 4V (7 each), adalimumab, apixaban (6 each), COVID-
19 vaccine, ethinylestradiol/levonorgestrel (4 each), acetylsalicylic acid, clopidogrel, COVID-19 vaccine NRVV AD, and rivaroxaban (3 each).

- Number of relevant events: 3532.
- Relevant event seriousness: serious (3532).
- Most frequently (>25 occurrences) reported relevant PTs: Cerebrovascular accident (1363), Cerebral infarction (416), Ischaemic stroke (367), Cerebral haemorrhage (306), Cerebral venous sinus thrombosis (166), Cerebral thrombosis (93), Cerebral ischaemia (76), Subarachnoid haemorrhage (72), Cerebral venous thrombosis (68), Cerebellar infarction (42), Brain stem infarction, Haemorrhage intracranial (35 each), Ischaemic cerebral infarction (33), Embolic stroke (31), Haemorrhagic stroke (29), Thalamic infarction (26).
- Time to event onset (n = 2626), 117 range: <24 hours to 402 days, median: 12 days.
  - <24 hours: 183 events (18 of which had a fatal outcome);
  - 1 day: 212 events (23 of which had a fatal outcome);
  - 2-7 days: 649 events (67 of which had a fatal outcome);
  - 8-14 days: 388 events (34 of which had a fatal outcome);
  - 15-30 days: 508 events (37 of which had a fatal outcome);
  - 31-180 days: 625 events (53 of which had a fatal outcome);
  - >180 days: 61 events (6 of which had a fatal outcome).
- Duration of relevant events (n = 201 out of 833 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 350 days, median 5 days.
  - <24 hours: 50 events;
  - 1 day: 16 events;
  - 2-7 days: 50 events;
  - 8-14 days: 16 events;
  - 15-30 days: 18 events;
  - 31-181 days: 41 events;
  - >180 days: 10 events.
- Relevant event outcome:78 fatal (314), resolved/resolving (1013), resolved with sequelae (504), not resolved (804), unknown (910).
- In 267 cases (reporting 314 relevant events with fatal outcome), the reported causes of death (≥10 occurrences) were coded to the PTs Cerebrovascular accident (94), Cerebral haemorrhage (77), Cerebral infarction (36), Immunisation (27), Off label use (26), Interchange of vaccine products (23), Ischaemic stroke (18), Headache (17), Death,

117 This number does not include 923 events for which administration and/or event onset dates were not provided or were incomplete; or events without a meaningful time to onset value as per reported information. Please note, multiple episodes of the same PT event were reported with different latencies within some cases hence the sum of latencies exceeds the total number of PT events.
Subarachnoid haemorrhage (16 each), Thrombosis (14), Myocardial infarction (13), Brain oedema, Cerebral thrombosis, Loss of consciousness (12 each), and Cardiac arrest (10). Most (183 of 267 cases) of the fatal cases involved elderly subjects. When the medical history was provided (162 cases), the most frequently (>5 occurrences) medical conditions included the PTs Hypertension (59), Diabetes mellitus (19), Atrial fibrillation (18), Cerebrovascular accident (12), COVID-19 (9), Cardiac failure, Cerebral infarction, Dyslipidaemia, Type 2 diabetes mellitus (8 each), Tobacco user (7), Chronic obstructive pulmonary disease, Cognitive disorder, Depression, Obesity, and Pulmonary embolism (6 each).

Analysis by age group

- CT: Adult (6), Elderly (13).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM: Paediatric (33 [9 Child, 24 Adolescent]), Adult (1575), Elderly (1352), Unknown (131).
  - Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible. Among the most frequently (>25 occurrences) reported relevant stroke-related events, the PTs Cerebral venous sinus thrombosis and Cerebral venous thrombosis had a greater than 3-fold reporting proportion in the adult population (8.2% and 3.7%, respectively) when compared to the elderly population (1.6% and 0.6%, respectively). Conversely, among the most frequently reported relevant stroke-related events, the PT Haemorrhagic stroke had a greater than 3-fold reporting proportion in the elderly population (1.7%) when compared to the adult population (0.4%).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 719 (23.3% of the 3091 cases reporting stroke-related events).

- Of the PM cases that reported medical histories, the percentage of cases with a fatal stroke-related event is higher in subjects with comorbid conditions (55.6%) when compared to the percentage of cases with a fatal stroke-related event in subjects without comorbidities (44.4%).

- Upon review of the most frequent (>25 occurrences) relevant events in PM cases that recorded medical histories, no relevant stroke-related events had a significant proportional reporting ratio of >3:1 in subjects with comorbidities compared to subjects without comorbidities.
O/E Analysis

O/E analysis was performed for Cerebral venous sinus thrombosis, Ischemic stroke and Hemorrhagic stroke (see Appendix 6B *Observed versus Expected Analyses for Adverse Events of Special Interest*). For CVST, some age groups had O/E greater than 1 when the low background rate was used in the analysis. However, the 95% CIs did not all include 1 (indicating non statistical significance). The O/E were similar to the most recent SBSR #3. Using the mid-range background rate, all stratifications have an O/E ratio less than 1.

Conclusion

Cerebral venous sinus thrombosis (CVST) and Cerebrovascular Accident (CVA)/Stroke were evaluated as signals during the reporting period and were not determined to be risks causally associated with the vaccine (please refer to Section 16.2.1 *Evaluation of Closed Signals*).

No additional safety signals other than those mentioned above have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

16.3.3.1.18. Sudden Death

Search criteria - PT Sudden Death.

Please refer to Section 16.3.4.1 *Death*.

16.3.3.1.19. Thromboembolic AESIs

Search criteria - HLGT (All path) Embolism and thrombosis (excluding PTs reviewed as Stroke AESIs) OR PT Coagulopathy.

Clinical Trial Data

- Number of cases: 17 (BNT162b2 [16], blinded therapy [1]; 2.5% of 668 cases in the total CT dataset) compared to 15 cases (2.1%) retrieved in the PSUR #2.

- Country of incidence: US (13), Argentina (3), Brazil (1).

- Subjects' gender: female (10), male (7).

- Subjects' age in years (n = 17), range: 18 – 81, mean: 56.4, median: 55.

- Medical history (n = 14): medical conditions reported more than once were coded to the PTs: Obesity (6), Hypertension (4), Depression (3), Anxiety, Cholecystectomy, Deep vein thrombosis, Gastroesophageal reflux disease, Hyperlipidaemia, Hypothyroidism, Osteoarthritis, Type 2 diabetes mellitus, and Vasectomy (2 each).

- COVID-19 medical history: None.

- Co-suspects: None.

- Reported relevant PTs: Pulmonary embolism (8), Deep vein thrombosis (6), Thrombosis (2), Coagulopathy, Embolism, Peripheral artery thrombosis, Portal vein
thrombosis (1 each). However, none of these SAEs were assessed as related to BNT162b2/blinded therapy.

- Relevant event outcome: fatal (1), resolved/resolving (13), resolved with sequelae (1), not resolved (5).

Post-Authorisation Data

- Number of cases: 6102 (1.2% of 507,683 cases in the total PM dataset), compared to 6507 cases (1.0%) retrieved in the PSUR #2.
- MC cases (2944), NMC cases (3158).
- Country of incidence (>50 occurrences): Germany (1882), France (910), UK (499), Australia (314), Italy (294), Sweden (278), Austria (264), US (200), Netherlands (129), Japan (109), New Zealand, Poland (93 each), Greece, Spain (90 each), Finland (86), Czech Republic (85), Belgium (82), Norway (78), Denmark (72), Taiwan (52); the remaining 402 cases were distributed among 41 countries.
- Subjects' gender: female (3322), male (2682), unknown (98).
- Subjects’ age in years (n = 5794), range: 5 – 102, mean: 55.0, median: 55.
- Medical history (n = 3054): the most frequently (>100 occurrences) reported medical conditions were coded to the PTs Hypertension (738), Non-tobacco user (203), Obesity (199), Asthma (184), Seasonal allergy (173), Deep vein thrombosis, Drug hypersensitivity (151 each), Tobacco user (137), Hypothyroidism (135), Pulmonary embolism (119), Type 2 diabetes mellitus (115), and Diabetes mellitus (113).
- Co-suspects (n = 162 cases): the most frequently (>3 occurrences) reported co-suspect medications were ethinylestradiol/levonorgestrel (14), COVID-19 vaccine mRNA (mRNA 1273) (13), adalimumab (12), COVID-19 vaccine, influenza vaccine INACT SPLIT 4V (11 each), apixaban, influenza vaccine (9 each), COVID-19 vaccine NRVV AD, JNJ 78436735 (6 each), rivaroxaban (5), influenza vaccine INACT SAG 4V (4), enoxaparin, ethinylestradiol/etinogestrel, and ethinylestradiol/gestodene (3 each).
- Number of relevant events: 7194.
- Relevant event seriousness: serious (6724), non-serious (470).
- Most frequently (>50 occurrences) reported relevant PTs: Pulmonary embolism (2068), Thrombosis (1461), Deep vein thrombosis (1321), Thrombophlebitis (285), Venous thrombosis limb (276), Superficial vein thrombosis (258), Venous thrombosis (173), Coagulopathy (164), Retinal vein occlusion (127), Embolism (103), Pulmonary thrombosis (77), Ophthalmic vein thrombosis (74), Retinal vein thrombosis (54), Retinal artery occlusion (52), Portal vein thrombosis (50).
• Time to event onset (n = 5217),\textsuperscript{118} range: < 24 hours to 375 days, median: 12 days.
  
  - <24 hours: 321 events (14 of which had a fatal outcome);
  - 1 day: 368 events (12 of which had a fatal outcome);
  - 2-7 days: 1358 events (37 of which had a fatal outcome);
  - 8-14 days: 810 events (36 of which had a fatal outcome);
  - 15-30 days: 1007 events (19 of which had a fatal outcome);
  - 31-180 days: 1243 events (37 of which had a fatal outcome);
  - >180 days: 110 events (6 of which had a fatal outcome).

• Duration of relevant events (n = 442 out of 1325 occurrences with outcome of resolved/resolved with sequelae), range: < 24 hours to 329 days, median 24.5 days.
  
  - <24 hours: 25 events;
  - 1 day: 11 events;
  - 2-7 days: 92 events;
  - 8-14 days: 51 events;
  - 15-30 days: 61 events;
  - 31-180 days: 165 events;
  - >180 days: 37 events.

• Relevant event outcome:\textsuperscript{78} fatal (265), resolved/resolving (2521), resolved with sequelae (506), not resolved (2148), unknown (1774).

• In 236 cases (reporting 265 relevant events with fatal outcome), the reported causes of death (>10 occurrences) were coded to the PTs Pulmonary embolism (116), Thrombosis (62), Cardiac arrest (27), Immunisation (26), Dyspnoea (24), Off label use (19), Myocardial infarction (18), Interchange of vaccine products (17), Deep vein thrombosis (15), Cardio-respiratory arrest, Cerebrovascular accident (14 each), Embolism (13), and Loss of consciousness (11).\textsuperscript{106} Most (153 of 236 cases) of the fatal cases involved elderly subjects. When the medical history was provided (153 cases), the most frequently (>5 occurrences) medical conditions included the PTs Hypertension (51), Diabetes mellitus (13), Atrial fibrillation, Obesity (12 each), Chronic obstructive pulmonary disease, Osteoporosis, Type 2 diabetes mellitus (9 each), Arteriosclerosis, Tobacco user (8 each), Asthma, Cholecystectomy, COVID-19, Deep vein thrombosis, Hypothyroidism, Myocardial infarction, Pulmonary embolism (7 each), Cerebral infarction, Dementia, Depression, Ex-tobacco user, Hospitalisation, Hypercholesterolaemia, Osteoarthritis, Overweight, and Surgery (6 each).

\textsuperscript{118} This number does not include 2001 events for which administration and/or event onset dates were not provided or were incomplete; or events without a meaningful time to onset value as per reported information. Please note, multiple episodes of the same PT event were reported with different latencies within some cases hence the sum of latencies exceeds the total number of PT events.
Analysis by age group

- CT: Adults (12), Elderly (5).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM: Paediatric (79 [7 Child, 72 Adolescent]), Adults (3833), Elderly (1966), Unknown (224).
  - Except for the PT Coagulopathy, no significant difference was observed in the reporting proportion of the most frequently (≥50 occurrences) reported thromboembolic AESIs, between the paediatric, adult and elderly populations. The reporting proportion of the PT Coagulopathy was significantly higher in the paediatric population (11.4%) when compared to the adult and elderly populations (2.7% and 2.1%, respectively).

Analysis by presence of comorbidities

- Number of PM subjects reporting comorbidities: 1356 (22.2% of the 6102 cases reporting thromboembolic AESIs).

- Of the PM cases that reported medical histories, the percentage of cases with a fatal thromboembolic AESI is higher in subjects with comorbid conditions (65.4%) when compared to the percentage of cases with a fatal thromboembolic AESI in subjects without comorbidities (34.6%).

- Upon review of the most frequent (≥50 occurrences) relevant events in cases that recorded medical histories, no thromboembolic AESIs had a significant proportional reporting ratio of >3:1 in subjects with comorbidities compared to subjects without comorbidities.

O/E Analysis

O/E analysis was performed for Arterial thromboembolism, Deep vein thrombosis, Disseminated intravascular coagulation, Thrombotic thrombocytopenia syndrome and Venous thromboembolism (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.

16.3.3.1.20. Vascular events

Search criteria - HLT (All Path) Vasculitides OR PTs Microangiopathy; Peripheral ischaemia.
Clinical Trial Data

- During the reporting period no serious cases from the CT dataset were reported; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 612 (0.12% of 507,683 cases, the total PM dataset), compared to 854 cases (0.13%) retrieved in the PSUR #2.
- MC cases (375), NMC cases (237).
- Country of incidence: Germany (154), France (112), Japan (57), UK (34), Italy (28), Australia (26), US (20), Austria (19), Netherlands (16), Taiwan, Province of China (15), Denmark, Greece, Norway (11 each); the remaining 98 cases were distributed among 31 countries.
- Subjects' gender: female (363), male (234) and unknown (15).
- Subjects' age in years (n = 576), range: 2 – 97, mean: 53.3, median: 58.0.
- Medical history (n = 332): the most frequently (≥ 5 occurrences) reported relevant medical conditions included Hypertension (78), Tobacco user (22), Type 2 diabetes mellitus (19), Diabetes mellitus (16), Obesity (13), Hypercholesterolaemia (12), Giant cell arteritis, Vasculitis (11 each), Dyslipidaemia, Henoch-Schonlein purpura (10), Autoimmune thyroiditis, Hyperlipidaemia, Hypersensitivity, Rheumatoid arthritis (9 each), Drug hypersensitivity (8), Raynaud's phenomenon (7), Autoimmune disorder (6), Polymyalgia rheumatica, Pulmonary embolism, Tobacco abuse, and Uveitis (5 each).
- Co-suspects (n = 20 cases): relevant co-suspect included Adalimumab (3).
- Number of events: 648.
- Relevant event seriousness: serious (455) and nonserious (193).
- Most frequently reported relevant PTs (≥20 occurrences): Vasculitis (267), Giant cell arteritis (102), Henoch-Schonlein purpura (66), Peripheral ischaemia (60).
- Time to event onset (n = 390)\(^{119}\), range: range: <24 hours to 178 days, median: 10 days.
  - <24 hours: 33 events (1 of which had a fatal outcome);
  - 1 day: 43 events;
  - 2-7 days: 105 events (2 of which had a fatal outcome);
  - 8-14 days: 51 events (1 of which had a fatal outcome);
  - 15-30 days: 63 events (1 of which had a fatal outcome);

\(^{119}\) This number does not include 259 events for which partial administration or event onset date was reported.
- 31-180 days: 95 events (6 of which had a fatal outcome).

- Duration of relevant events (n = 49 out of 649 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 172 days, median 25 days.
  - <24 hours: 2 events;
  - 1 day: 1 event;
  - 2 - 7 days: 11 events;
  - 8-14 days: 5 events;
  - 15-30 days: 9 events;
  - 31-180 days: 21 events.

- Relevant event outcome: fatal (17), resolved/resolving (218), resolved with sequelae (53), not resolved (195), unknown (165).

- In 17 cases (reporting 17 relevant events with fatal outcome), the reported causes of death (>2 occurrences) were coded to Vasculitis (7), Eosinophilic granulomatosis with polyangiitis, Peripheral ischaemia (2 each). Most (13 of 17 cases) of the fatal cases involved elderly subjects. When the medical history was provided (14 cases), the most frequently (>2 occurrences) relevant medical conditions included Hypertension (3), Diabetes mellitus and Obesity (2 each).

Analysis by age group

- PM: Paediatric (62), Adults (317), Elderly (208) and Unknown (25).
  - Among the frequently (>2%) reported relevant PT, the reporting proportion of PT Anti-neutrophil cytoplasmic antibody positive vasculitis was higher in elderly population when compared to adult population (47.1% in elderly vs 38.5% in adult). No paediatric cases reported PT Anti-neutrophil cytoplasmic antibody positive vasculitis which is consistent with the known epidemiology.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 224 (36.6 % of the PM cases reporting vasculitic events).

- The reporting proportion of vasculitic AESIs with a fatal outcome (3.4 %) is higher in subjects with comorbid conditions when compared to the reporting proportion observed in the subject without comorbidities (2.2 % for fatal outcome).

O/E Analysis

O/E analysis was performed for Behcet’s syndrome, Giant cell arteritis, Henoch-Schonlein purpura, Limb ischaemia, and Vasculitis (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).
Conclusion

Vasculitis was evaluated as signal during the reporting period and was determined to not be a risk (please refer to Section 16.2.1 Evaluation of Closed Signals).

No new significant safety information was identified based on a review of these cases or of the O/E analysis. Safety surveillance will continue.

16.3.3.1.21. AESIs in subjects with Malnutrition; HIV infection

As part of the approval letter for the emergency use of Tozinameran - COVID-19 mRNA vaccine (nucleoside modified) - COMIRNATY®, the WHO requested the MAH to provide additional data on vaccine immunogenicity, effectiveness and safety on population groups represented in Low- and Middle-Income Countries (LMICs) including individuals with conditions such as malnutrition and populations with existing co-morbidities such as tuberculosis, human immunodeficiency virus (HIV) infection and other high prevalent infectious diseases.

Search criteria - PT Decode (History): Acute HIV infection; Asymptomatic HIV infection; Congenital HIV infection; HIV infection; HIV infection CDC Group I; HIV infection CDC Group II; HIV infection CDC Group III; HIV infection CDC Group IV subgroup A; HIV infection CDC Group IV subgroup B; HIV infection CDC Group IV subgroup C1; HIV infection CDC Group IV subgroup C2; HIV infection CDC Group IV subgroup D; HIV infection CDC Group IV subgroup E; HIV infection CDC category A; HIV infection CDC category B; HIV infection CDC category C; HIV infection CDC group IV; HIV infection WHO clinical stage I; HIV infection WHO clinical stage II; HIV infection WHO clinical stage III; HIV infection WHO clinical stage IV; Malnutrition; Perinatal HIV infection; Prophylaxis against HIV infection; Tuberculosis.

Clinical Trial Data

- Number of cases: 11 (blinded therapy [2], BNT162b2 [9]) (1.6% of 668 cases, the total CT dataset, compared to 7 cases (1.0%) retrieved in the PSUR #2.
- Country of incidence: US (4), Brazil, Germany, South Africa (2 each), and Argentina (1).
- Subjects' gender: female (2), male (9).
- Subjects' age in years (n = 11), range: 6 – 71, mean: 40, median: 39.
- Medical history (n = 11): HIV infection (7), Malnutrition, Tuberculosis (2 each).
- COVID-19 Medical history: None.
- Co-suspects: None.
- Reported PTs (16): Condition aggravated, Maternal exposure during pregnancy120, Mental disorder (2 each), Atrial fibrillation, Cephalo-pelvic disproportion, Constipation,

120 Maternal cases with no exposure in-utero reported.
Cranio-cerebral injury, Failed trial of labour, Headache, Intestinal obstruction, Lumbar spinal stenosis, Prostate cancer, and Spinal claudication (1 each). None of the events were related to BNT162b2 or blinded therapy.

- Relevant event outcome: resolved/resolving (14), resolved with sequelae (2).

Post-Authorisation Data

- Number of cases: 197 (0.04% of 507,683 cases, the total PM dataset), compared to 393 cases (0.06%) retrieved in the PSUR #2.

Patients with pre-existing HIV Infection: 107 (0.02% of 507,683 cases, the total PM dataset).

- MC cases (50), NMC cases (57).
- Country of incidence\textsuperscript{121}: France (29), Italy (21), Germany (17), US (12), UK (7), Brazil, Netherlands (3 each), Mexico, Romania, and Sweden (2 each); the remaining 9 cases were distributed among 9 countries.
- Subjects' gender: female (19), male (79) and unknown (9).
- Subjects' age in years (n = 98), range: 16 – 81, mean: 50.2, median: 51.
- Co-suspect vaccines/medications (3): COVID-19 vaccine (unspecified), dolutegravir sodium/rilpivirine hydrochloride, influenza vaccine inactive Split 3V (1 each).
- Of the 107 cases reporting a pre-existing HIV condition, 6 subjects reported cardiac disorders. The events (10) in these cases were coded to the PTs Myocarditis, Pericarditis, Tachycardia (2 each), Arrhythmia, Cardiovascular disorder, Endocarditis fibroplastica, and Palpitations (1 each). Of the 10 events, 8 were assessed as serious and 2 events were non-serious. Outcome of the events\textsuperscript{122} was reported as resolved/resolving (3), not resolved (2), and unknown (6).
- Of the 107 cases, 35 subjects reported nervous system disorders. The events (49) reported more than once in these cases were coded to the PTs Headache (15), Dizziness (7), Facial paralysis (3), Bell's palsy, Cerebrovascular accident, Disturbance in attention, Hypoaesthesia, Paraesthesia, and Speech disorder (2 each); Of the 49 events, 23 were assessed as serious and 26 events as non-serious. Outcome was reported as resolved/resolving (21), not resolved (17), and unknown (11).
- Of the 107 cases, 30 subjects reported infectious events. The events (30) in these cases were coded to PTs COVID-19 (16), Herpes zoster (2), Arthritis bacterial, Asymptomatic COVID-19, Bacterial sepsis, Encephalitis, Fungal infection, Herpes simplex encephalitis, ...

\textsuperscript{121} There were 7 cases reported from low- and middle-income countries (Brazil [3], Mexico [2], Serbia, South Africa [1 each]).

\textsuperscript{122} One event reported more than 1 outcome.
Herpes zoster reactivation, HIV peripheral neuropathy, Myelitis, Oral fungal infection, Post viral fatigue syndrome, Virologic failure (1 each). Of the 30 events, 25 were assessed as serious and 5 events were non-serious. Outcome of the events was reported as resolved with sequelae (1), resolved/resolving (8), not resolved (6), and unknown (15).

- Time to event onset (n = 175\textsuperscript{123}), range: <24 hours to 255 days, median: 2 days (no events with a fatal outcome).
  - <24 hours: 47 events;
  - 1 day: 26 events;
  - 2-7 days: 45 events;
  - 8-14 days: 14 events;
  - 15-30 days: 8 events;
  - 31-90 days: 6 events;
  - 91-255 days: 19 events.

- Duration of relevant events (n = 22\textsuperscript{124}, of which 21 events reported an outcome of resolved/resolving/resolved with sequelae), range: 2 hours to 288 days, median: 2 days.
  - <24 hours: 1 event;
  - 1 day: 3 events;
  - 2-7 days: 10 events;
  - 8-14 days: 3 event;
  - 15-31 days: 4 events;
  - 288 days: 1 events.

- There was no trend in the reporting of infections, cardiac disorders and nervous disorders in subjects with pre-existing HIV infection when compared to the subjects without the disease.

- Of the 107 cases, 88 cases involved adults, 9 cases involved elderly and in 9 cases age group was not reported. Due to the low volume of cases reported in elderly, it was not possible to make a meaningful comparison between the adults and elderly patient population.

Patients with pre-existing tuberculosis: 67 (0.01% of 507,683 cases, the total PM dataset).

- MC cases (37), NMC cases (30).

\textsuperscript{123} This number does not include events which occurred prior to vaccine administration.

\textsuperscript{124} This number does not include events for which event onset dates or event cessation dates were not reported or events with a not meaningful time to event cessation value as per reported information.
COVID-19 mRNA vaccine (nucleoside modified)  
Periodic Safety Update Report (PSUR) 3  
19 December 2021 through 18 June 2022

- Country of incidence\textsuperscript{125}: France (35), UK (7), Germany, Brazil, US (4 each), Canada, Netherlands, South Africa, Taiwan (Province of China) (2 each), Bulgaria, Japan, New Zealand, Philippines, Sweden (1 each).

- Subjects' gender: female (46), male (21).

- Subjects' age in years (n = 66), range: 8 - 94, mean: 59.3, median: 62.


- Co-suspect vaccines/medications (7): COVID-19 vaccine (unspecified), COVID-19 MRNA 1273, COVID-19 vaccine NRVAD (CHADOX1 NCOV-19), Influenza vaccine, Influenza vaccine INACT SPLIT 4V, Ranibizumab, varicella zoster vaccine live (OKA/Merck) (1 each).

- Of the 67 cases reporting pre-existing tuberculosis, 13 subjects reported cardiac disorders. The events (21) in these cases were coded to the PTs Pericarditis (4), Myopericarditis, Palpitations (2 each), Arrhythmia, Atrioventricular block, Bradycardia, Cardiac arrest, Cardiac failure, Cardiac failure acute, Cardiovascular disorder, Early repolarisation syndrome, Myocarditis, Pericardial effusion, Pericardial fibrosis, Tachycardia, and Ventricular hypokinesia (1 each). Of the 21 events, 19 were assessed as serious and 2 events were non-serious. Outcome of the events\textsuperscript{78} was reported as fatal (1), resolved with sequelae (1), resolved/resolving (8), not resolved (5), and unknown (7).

- Of the 67 cases, 19 subjects reported nervous system disorders. The events (32) in these cases were coded to PTs Headache (7), Dizziness, Somnolence (3 each), Hypoaesthesia, Speech disorder (2 each), Aphasia, Burning sensation, Cerebral infarction, Dysstasia, Head discomfort, Hypokinesia, Irregular sleep wake rhythm disorder, Ischaemic stroke, Loss of consciousness, Migraine, Neuralgia, Paraesthesia, Sensory disturbance, Sensory loss, and Syncope (1 each). Of the 32 events, 14 were assessed as serious and 18 events were non-serious. Outcome of the events was reported as fatal (1), resolved/resolving (11), not resolved (6), resolved with sequelae (1), and unknown (13).

- Of the 67 cases, 13 subjects reported infectious events. The events (15) in these cases were coded to the PTs COVID-19 (4), Influenza, Nasopharyngitis (2 each), Bronchitis, Chorioretinitis, COVID-19 pneumonia, Herpes virus infection, Herpes zoster, Pancreatic abscess, Tuberculosis (1 each). Of the 15 events, 9 were assessed as serious and 6 events were non-serious. Outcome of the events was reported as resolved with sequelae (1), resolved/resolving (4), not resolved (1), and unknown (9).

- Time to event onset (n = 235), range: <24 hours to 377 days, median: 2 days.
  - <24 hours: 70 events (none of which had a fatal outcome);
  - 1 day: 22 events (1 of which had a fatal outcome);
  - 2-7 days: 55 events (none of which had a fatal outcome);
  - 8-14 days: 12 events (none of which had a fatal outcome);

\textsuperscript{125} There were 8 cases reported from low- and middle-income countries (Brazil [4], South Africa [2], Bulgaria, and Philippines [1 each]).
15-30 days: 27 events (1 of which had a fatal outcome);
31-180 days: 44 events (4 of which had a fatal outcome);
181-377 days: 5 events (none of which had a fatal outcome).

- Duration of relevant events (n = 13126, out of which 12 occurrences were reported with outcome of resolved/resolved with sequelae/resolving), range: 2 to 300 days, median 12 days.
  - 2 - 7 days: 5 events;
  - 8-14 days: 3 events;
  - 15-31 days: 2 events;
  - 32-300 days: 3 events.

- There was no trend in the reporting of infections, cardiac disorders and nervous disorders in subjects with pre-existing tuberculosis when compared to the subjects without the disease.

- Of the 67 cases, 36 cases involved adults, and 29 cases involved elderly, and the age group was not reported in 1 case. The reporting proportion of cases involving infectious events was higher in adult population (14.9%) when compared to the elderly (4.5%); and more adult subjects reported cases involving nervous system disorders as compared to the elderly (17.9% in adults vs 10.4% in elderly). No significant difference was observed in the reporting proportion of cases involving cardiac events (10.4% in adults vs 9.0% in elderly) between the elderly and adult population.

Patients with pre-existing malnutrition: 23 (<0.01% of 507,683 cases, the total PM dataset).

- MC cases (13), NMC cases (10).
- Country of incidence127: France (8), Germany (5), Sweden, Switzerland (3 each), Finland, Japan, Latvia, and US (1 each).
- Subjects’ gender: female (13), male (10).
- COVID-19 Medical history (n = 2): COVID-19, COVID-19 pneumonia (1 each).
- Co-suspect medications (2): bevacizumab, and COVID-19 vaccine MRNA (MRNA 1273) (1 each).
- In these 23 cases, the most frequently reported events (119, ≥3 occurrences) were coded to the PTs General physical health deterioration (5), Headache, Inappropriate schedule of

126 This number does not include events for which event onset dates or event cessation dates were not reported or events without a meaningful time to event cessation value as per reported information.
127 There was 1 case reported from a low- and middle-income country (Latvia).
product administration, Interchange of vaccine products, Off label use, Pyrexia (4 each), Condition aggravated, Fatigue, Vaccination site pain (3 each).

- Of the 23 cases reporting pre-existing malnutrition, 9 subjects reported PTs General physical health deterioration (5), Condition aggravated, Fatigue (3 each), Anaemia, Asthenia, Dehydration, and Marasmus (1 each). Of the total 15 events, 7 events were assessed as serious, and 8 events were non-serious. Outcome of the events was reported as fatal (1), resolved/resolving (4), not resolved (4), and unknown (6).

- Time to event onset (n = 81), range: <24 hours to 107 days, median: 1 day.
  - <24 hours: 38 events (3 of which had a fatal outcome);
  - 1 day: 7 events (none of which had a fatal outcome);
  - 2-7 days: 10 events (1 of which had a fatal outcome);
  - 8-14 days: 9 events (2 of which had a fatal outcome);
  - 15-30 days: 10 events (3 of which had a fatal outcome);
  - 31-107 days: 7 events (2 of which had a fatal outcome).

- Duration of relevant events (n = 9128 all occurrences with outcome of resolved), range: <24 hours to 3 days, median 1 day.
  - <24 hours: 1 event;
  - 1 day: 7 events;
  - 3 days: 1 event;

Of the 23 cases, 10 were reported in elderly and 11 cases involved adults, the age group was not reported in 1 case. Due to the low volume of cases (1 case) reporting cardiac disorders, it was not possible to make a meaningful comparison between the adults and elderly patient population. The reporting proportion of cases involving infectious events was higher in the elderly population (13.0%) when compared to adults (4.3%). No significant difference was observed in the reporting proportion of cases involving nervous system disorders between the elderly (21.7%) and adult population (17.4%). Generally, there was a low volume of cases reporting malnutrition in the current dataset.

**Conclusion**

No safety signals have emerged based on the review of these cases. Safety surveillance will continue.

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128 This number does not include events for which event onset dates or event cessation dates were not reported or events without a meaningful time to event cessation value as per reported information.
16.3.3.2. Clinical Reactogenicity Data on Individuals Previously exposed or not to SARS-COV-2

Data are available from 3 analyses: children 2 to <5 years and children 6 months to <2 years receiving up to 3 primary doses of BNT162b2 3 μg, and adults 18-55 years receiving a fourth dose booster of either the current vaccine or a monovalent Omicron-modified vaccine, both at 30 μg.

Children 6 months to <2 years (from C4591007)

Subgroups of Phase 2/3 pediatric participants 6 months to <2 years of age had similar reactogenicity, with regard to local reactions, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution. There were no meaningful differences in the overall patterns of local reactions across these subgroups.

There were 88 BNT162b2 participants with baseline positive SARS-CoV-2 status and 1078 BNT162b2 participants with baseline negative SARS-CoV-2 status who reported e-diary data in the 6 months to <2 years of age group. The frequencies of local reactions reported after any dose of BNT162b2 in the SARS-CoV-2 baseline status subgroups were:

- Tenderness: baseline positive: 28.4%, baseline negative: 26.3%
- Redness: baseline positive: 13.6%, baseline negative: 18.0%
- Swelling: baseline positive: 6.8%, baseline negative: 7.4%.

After any of three doses of BNT162b2 3-μg in the 6 months to <2 years of age group, the frequency and pattern of local reactions in baseline positive children was similar to those who were baseline negative and did not suggest any clinically meaningful difference based on prior infection status.

Subgroups of Phase 2/3 pediatric participants 6 months to <2 years of age had similar reactogenicity, with regard to systemic events, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS-CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution. There were no meaningful differences in the overall patterns of systemic events across these subgroups.

The frequencies of systemic events reported after any dose of BNT162b2 in the SARS-CoV-2 baseline status (positive or negative) subgroups were:

- Irritability: baseline positive: 63.6%, baseline negative: 68.6%
- Drowsiness: baseline positive: 47.7%, baseline negative: 40.9%
- Appetite: baseline positive: 46.6%, baseline negative: 37.9%
- Fever: baseline positive: 14.8%, baseline negative: 14.2%.

After any of three doses of BNT162b2 3-μg in the 6 months to <2 years of age group, the frequency and pattern of systemic events in baseline positive children was similar to those
who were baseline negative and did not suggest any clinically meaningful difference based on prior infection status.

The reactogenicity data for participant subgroups based on their baseline SARS-CoV-2 positive or negative status are shown as tabular summary data for local reactions (Appendix 6D -Table 5 and Table 6) and systemic events (Appendix 6D -Table 7 and Table 8).

**Children** 2 to <5 years (from C4591007)

Subgroups of Phase 2/3 pediatric participants 2 to <5 years of age had similar reactogenicity, with regard to local reactions, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution. There were no meaningful differences in the overall patterns of local reactions across these subgroups.

There were 231 BNT162b2 participants with baseline positive SARS-CoV-2 status and 1597 BNT162b2 participants with baseline negative SARS-CoV-2 status who reported e-diary data in the 2 to <5 years of age group. The frequencies of local reactions reported after any dose of BNT162b2 in the SARS-CoV-2 baseline status subgroups were:

- Pain: baseline positive: 45.9%, baseline negative: 47.2%
- Redness: baseline positive: 15.2%, baseline negative: 19.4%
- Swelling: baseline positive: 9.1%, baseline negative: 8.3%.

After any of three doses of BNT162b2 3-μg, the frequency and pattern of local reactions in baseline positive children was similar to those who were baseline negative and did not suggest any clinically meaningful difference based on prior infection status.

Subgroups of Phase 2/3 pediatric participants 2 to <5 years of age had similar reactogenicity, with regard to systemic events, across the BNT162b2 and placebo groups when evaluated by baseline SARS-CoV-2 status. The subgroup of baseline SARS-CoV-2 status (positive) included a limited number of participants, and results should be interpreted with caution. There were no meaningful differences in the overall patterns of systemic events across these subgroups.

The frequencies of common systemic events reported after any dose of BNT162b2 in the SARS-CoV-2 baseline status (positive or negative) subgroups were:

- Fatigue: baseline positive: 35.5%, baseline negative: 46.0%
- Fever: baseline positive: 10.0%, baseline negative: 10.6%
- Headache: baseline positive: 10.4%, baseline negative: 8.4%.

After any of three doses of BNT162b2 3-μg in the 2 to <5 years of age group, the frequency and pattern of systemic events in baseline positive children was similar to those who were
baseline negative and did not suggest any clinically meaningful difference based on prior infection status.

The reactogenicity data for participant subgroups based on their baseline SARS-CoV-2 positive or negative status are shown as tabular summary data for local reactions (Appendix 6D - Table 1 and Table 2) and systemic events (Appendix 6D - Table 3 and Table 4).

**Adults 18 through 55 years (from C4591031 Substudy D)**

Any local reactions reported within 7 days after first study (Dose 4) vaccination for Cohort 2 participants were similar in the BNT162b2 OMI (78.6%) and BNT162b2 (79.4%) groups, and most events were mild or moderate in severity. No Grade 4 local reactions were reported.

Across the BNT162b2 OMI and BNT162b2 vaccine groups, the frequencies of the most commonly reported local reaction of pain at the injection site were ≤78.0% for baseline positive and ≤80.8% baseline negative participants, respectively. The baseline positive subgroup included a limited number of participants, and their results should be interpreted with caution. Overall, numerical differences in any of the local reactions by SARS-CoV-2 baseline status were not considered clinically meaningful.

Any systemic events reported within 7 days after first study (Dose 4) vaccination for Cohort 2 participants were similar in the BNT162b2 OMI (77.6%) and BNT162b2 (72.9%) groups, and most events were mild or moderate in severity. No Grade 4 systemic events were reported.

Across the BNT162b2 OMI and BNT162b2 vaccine groups, the frequencies of headache were ≤46.7% for baseline positive and ≤48.4% for baseline negative participants, respectively. For fatigue, the other most commonly reported systemic event, the frequencies in baseline positive participants were 70.0% (2-sided 95% CI: 55.4, 82.1) for BNT162b2 OMI compared to 44.4% (2-sided 95% CI: 29.6, 60.0) for BNT162b2. Frequencies of fatigue in baseline negative participants were 63.1% (2-sided 95% CI: 56.7, 69.2) and 63.2% (2-sided 95% CI: 57.1, 69.1) for the BNT162b2 OMI and BNT162b2 groups, respectively. The baseline positive subgroup included a limited number of participants, which contributed to wide confidence intervals around the point estimate, and their results should be interpreted with caution. Overall, numerical differences in any of the systemic events by SARS-CoV-2 baseline status were not considered clinically meaningful.

The reactogenicity data for participant subgroups based on their baseline SARS-CoV-2 positive or negative status are shown as tabular summary data for local reactions (Appendix 6D - Table 9 and Table 10) and systemic events (Appendix 6D - Table 11 and Table 12).

**16.3.3.3. Local Adverse Reactions**

Search criteria - PTs Erythema; Injection site erythema; Injection site pain; Injection site swelling; Swelling.
Of the 8654 cases, 57 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

- In 4 case the event of interest was due to underlying conditions (rheumatoid arthritis flare-up (2 cases), superficial vein thrombosis, and total knee replacement)
- In 15 cases the event of interest was attributed to another co-suspect drug and not COVID-19 mRNA vaccine
- Foetuses, neonates, or infants exposed to the vaccine during the mother’ pregnancy or exposed through breastfeeding were reported in 38 cases (cases reporting exposure in utero or exposure during lactation are reviewed in Section 16.3.5.3 Use in Pregnant/Lactating Women).

Therefore, 8597 cases are included in the analysis below.

Clinical Trial Data

- There were no serious clinical trial cases of local reactions reported during the reporting interval; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 8597 (1.7% of 507,683 cases, the total PM dataset), compared to 21,240 cases (3.2%) retrieved in the PSUR #2.
- MC cases (3250), NMC cases (5347).
- Country of incidence (>2%): UK (1629), Germany (1121), Malaysia (781), Japan (368), US (361), France (360), Italy (305), Poland (294), Australia (275), Netherlands (253), Sweden (247), Ireland (219), Philippines (213), Belgium (189); the remaining 1982 cases were distributed among 54 countries.
- Subjects' gender: female (6266), male (2050) and unknown (281).
- Subjects' age in years (n = 7836), range: 2 - 98, mean: 43.7, median: 43.0.
- Medical history (n = 2461): the most frequently (>50) reported medical conditions included Hypertension (266), Asthma (215), Drug hypersensitivity (189), Hypersensitivity (158), Seasonal allergy (154), Food allergy (131), Hypothyroidism (104), Immunodeficiency (77), Depression (72), Fibromyalgia (64), Diabetes mellitus (60), Migraine (54), Anxiety (52), and Gastroesophageal reflux disease (51).
- Co-suspect vaccines/medications (n = 121): those reported in ≥ 2 cases included adalimumab (15), influenza vaccine (9), COVID-19 vaccine mRNA [MRNA 1273] (8),

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129 Some cases reported more than 1 medical history event.

- Number of relevant events: 9243.
- Relevant event seriousness: 42 serious (1868), non-serious (7380).
- Most frequently reported relevant PTs (≥2%): Erythema (4137), Swelling (4036), Injection site pain (690), and Injection site swelling (212).
- Most frequently co-reported PTs (>5%): Headache (1404), Pruritus (1402), Pyrexia (1302), Immunisation (1187), Pain (1160), Fatigue (1108), Pain in extremity (1017), Off label use (1007), Lymphadenopathy (970), Interchange of vaccine products (916), Rash (848), Myalgia (782), Axillary pain (778), Vaccination site pain (778), Arthralgia (723), Injection site pain (690), Peripheral swelling (676), Chills (644), Nausea (559), Dizziness (549), Malaise (530), and Dyspnoea (514).
- Time to event onset (n = 5683)\textsuperscript{130} range: range: <24 hours to 366 days, median: 1 day.
  - <24 hours: 2059 events (1 of which had a fatal outcome)\textsuperscript{131};
  - 1 day: 1673 events;
  - 2-7 days: 1334 events;
  - 8-14 days: 264 events;
  - 15-30 days: 172 events;
  - 31-181 days: 181 events.
- Duration of relevant events (n = 1328 out of 9440 occurrences with outcome of resolved/resolved with sequelae), range = <24 hours to 233 days, median 3 days.
  - <24 hours: 310 events;
  - 1 day: 150 events;
  - 2-7 days: 499 events;
  - 8-14 days: 130 events;
  - 15-30 days: 67 events;
  - 31-180 days: 167 events;
  - >180 days: 5 events.

\textsuperscript{130} This number does not include 3598 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\textsuperscript{131} In this case, the subject died from a fatal anaphylactic reaction.
• Relevant event outcome: 78 fatal (6), resolved/resolving (4065), resolved with sequelae (99), not resolved (2582), unknown (2524).
  - There were 6 cases reporting fatal events of interest (Erythema [4 cases] and Swelling [2 case]) in elderly (4 cases) and adult (2 cases) patients. Time to onset of fatal events were < 24 hours (1 event), 1 day (1 event), 3 days (1 event), 4 days (1 event), and unknown days (2 events). Review of these cases identified additional fatal adverse events reported in these cases and the local adverse reactions were not the primary cause of death in these cases.

Analysis by age group

PM: Paediatric (512), Adults (6972), Elderly (373) and Unknown (740).

<table>
<thead>
<tr>
<th>Event of Interest</th>
<th>Paediatric (n%)</th>
<th>Adult (n%)</th>
<th>Elderly (n%)</th>
<th>Unknown (n%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythema</td>
<td>323 (59.7)</td>
<td>2885 (41.2)</td>
<td>678 (63.8)</td>
<td>251 (39.7)</td>
</tr>
<tr>
<td>Injection site erythema</td>
<td>9 (1.7)</td>
<td>137 (2.0)</td>
<td>14 (1.3)</td>
<td>8 (1.3)</td>
</tr>
<tr>
<td>Injection site pain</td>
<td>16 (3.0)</td>
<td>617 (8.8)</td>
<td>47 (4.4)</td>
<td>10 (1.6)</td>
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<tr>
<td>Injection site swelling</td>
<td>10 (1.8)</td>
<td>183 (2.6)</td>
<td>15 (1.4)</td>
<td>4 (0.6)</td>
</tr>
<tr>
<td>Swelling</td>
<td>183 (33.8)</td>
<td>3185 (45.5)</td>
<td>309 (29.1)</td>
<td>359 (56.8)</td>
</tr>
<tr>
<td>Total</td>
<td>541</td>
<td>7007</td>
<td>1063</td>
<td>632</td>
</tr>
</tbody>
</table>

  a. Some cases reported more than 1 event.

• In general, the events of interest were similar by percentage across age group, with Erythema, Injection site pain, and Swelling more frequently reported.

Analysis by presence of comorbidities

• PM
  - Number of subjects with comorbidities: 38,787 (7.6% of 507,683 cases, the total PM dataset). Subjects with comorbidities were reported in (902/10.5 %) of the Local Adverse Reactions dataset. Given the nature of the adverse events of interest reported (Erythema, Injection site erythema, Injection site pain, Injection site swelling, Swelling) and the percentage of patients with comorbidities in the dataset, there were no differences between the group with comorbidities and the one without comorbidities.
Analysis by dose

- PM

  - Number of post-authorisation vaccine doses\textsuperscript{132} administered at the time of the event onset: Dose 1 in 2140 cases, Dose 2 in 1874 cases, Dose 3 in 2627 cases, Dose 4 in 77 case, Dose 5 in 1 case, and the dose number was not specified in 2039 cases.

<table>
<thead>
<tr>
<th>PT</th>
<th>Dose 1(^a) (n/%)</th>
<th>Dose 2(^a) (n/%)</th>
<th>Dose 3(^a) (n/%)</th>
<th>Dose 4(^a) (n/%)</th>
<th>Dose 5(^a) (n/-)</th>
<th>Dose Unspecified(^a) (n/%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythema</td>
<td>1200 (53.2)</td>
<td>998 (50.0)</td>
<td>1068 (38.5)</td>
<td>47 (58.8)</td>
<td>-</td>
<td>917 (39.7)</td>
</tr>
<tr>
<td>Injection site erythema</td>
<td>23 (1.0)</td>
<td>45 (2.3)</td>
<td>28 (1.0)</td>
<td>2 (2.5)</td>
<td>-</td>
<td>74 (3.2)</td>
</tr>
<tr>
<td>Injection site pain</td>
<td>91 (4.0)</td>
<td>61 (3.1)</td>
<td>47 (1.7)</td>
<td>3 (3.8)</td>
<td>-</td>
<td>492 (21.3)</td>
</tr>
<tr>
<td>Injection site swelling</td>
<td>12 (0.5)</td>
<td>14 (0.7)</td>
<td>11 (0.4)</td>
<td>1 (1.3)</td>
<td>-</td>
<td>117 (5.1)</td>
</tr>
<tr>
<td>Swelling</td>
<td>930 (41.2)</td>
<td>877 (44.0)</td>
<td>1619 (58.4)</td>
<td>27 (33.8)</td>
<td>1 (100)</td>
<td>648 (28.1)</td>
</tr>
<tr>
<td>Total</td>
<td>2256</td>
<td>1995</td>
<td>2773</td>
<td>80</td>
<td>1</td>
<td>2308</td>
</tr>
</tbody>
</table>

\(^a\) Vaccine dose count by PT differs than vaccine dose count by case given that some cases reported more than 1 PT.

The majority of post-authorisation events reported across doses were similar with the exception of injection site pain being reported more frequently in the unspecified dose group.

Conclusion

Local adverse reactions were reported in 8597 relevant cases representing 1.7% of the cases in the reporting period. The majority of events (79.8%) were non-serious events with 44.9% of the events resolved, resolved with sequelae or resolving at the time of reporting. There were 9 fatal cases describing fatal local adverse reactions in 6 cases; two were in adult and 4 were elderly subjects. Three of the 9 fatal cases did not report fatal local adverse reaction events. Review of these cases indicated that there were additional fatal adverse events reported and the event of interests (Erythema, Swelling) were not the primary cause of death in these subjects. When reported, the majority onset of events occurred within to <24 hours, with durations lasting <24 hours to 7 days.

The PM data appears to differ from the clinical trial data where injection site pain is generally the most frequently reported local reactogenicity event in adults and children. However, this is considered to be an effect of coding conventions given that commonly co-reported PTs in the cases are: Pain, Pain in extremity and Vaccination site pain. Evaluation of local adverse reaction cases did not reveal any significant new safety information. Local adverse reactions are appropriately described in the RSI. Surveillance of local adverse reactions will continue.

16.3.3.4. Systemic Adverse Reactions

Search criteria - PTs Arthralgia; Chills; Fatigue; Headache; Myalgia; Pyrexia.

\(^{132}\) Number of vaccine doses is reported by case number.
Of the 167,869 cases, 98 cases were determined to be non-contributory and were not included in the discussion due to involving neonate, or infants exposed to the vaccine through breastfeeding.

Clinical Trial Data

- Number of cases: 11 (BNT162b2 [10], and blinded therapy [1]) (1.6% of 668 cases, the total CT dataset) compared to 4 cases (0.6%) retrieved in the PSUR #2.
- Country of incidence: US (7), Germany (2), Finland, Spain (1 each).
- Subjects' gender: male (11).
- Subjects' age (n = 11), range: 23 months to 79 years; median 46 years.
- Medical history (n = 10, >1 occurrence): Hypertension (3), Dermatitis atopic, Gastroesophageal reflux disease, Insomnia (2 each).
- Co-suspects: None.
- Number of relevant events: 11.
- Relevant PTs: Pyrexia (6), Arthralgia (3), Headache, and Myalgia (1 each), none of which were assessed as related to BNT162b2 by the investigator and Sponsor.
- Time to event onset (n = 11): range: 21 to 282 days, median: 114 days (none of the events had a fatal outcome).
  - 21-30 days: 2 events;
  - 31-90 days: 3 events;
  - 91-180 days: 4 events;
  - 282 days: 1 event.
- Duration of relevant events (n = 9, all of which were reported with an outcome of resolved), range: 1 day 6 hours to 17 days, median 4 days.
  - 1-7 days: 7 events;
  - 8-17 days: 2 events.
- Relevant event outcome: resolved/resolving (11).

Post-Authorisation Data

- Number of cases: 167,760 (33% of 507,683 cases in the total PM dataset), compared to 279,184 (42.5% retrieved in the PSUR #2).
- MC cases (47,132), NMC cases (120,628).
COVID-19 mRNA vaccine (nucleoside modified)
Periodic Safety Update Report (PSUR) 3
19 December 2021 through 18 June 2022

- Country of incidence (top 10 countries): Germany (45,946), Netherlands (20,067), UK (10,862), Australia (7333), Iraq (6827), France (6453), Belgium (5932), Sweden (5441), Austria (4818), Japan (4718); the remaining cases were distributed among 89 countries.

- Subjects' gender: female (116,859), male (47,526) and unknown (3375).

- Subjects' age in years (n = 156,917), range: 3 days – 104 years, mean: 41.2; median: 40.0.

- Medical history (n = 39604): the most frequently (>1000 cases) reported medical conditions included Hypertension (4039), Disease risk factor (3139), Asthma (3026), Seasonal allergy (2591), Drug hypersensitivity (2144), Hypersensitivity (1801), Hypothyroidism (1404), Food allergy (1363), Diabetes mellitus (1005).

- COVID-19 Medical history (n = 9126): COVID-19 (6235), Suspected COVID-19 (2812), Post-acute COVID-19 syndrome (157), COVID-19 pneumonia (46), Coronavirus infection (44), SARS-CoV-2 test positive (33), Asymptomatic COVID-19, Exposure to SARS-CoV-2 (18 each), SARS-CoV-2 antibody test positive (2), Breakthrough COVID-19, Coronavirus pneumonia, Coronavirus test positive, and COVID-19 treatment (1 each).

- Co-suspects (n = 1652): the most frequently (≥10 occurrences) reported co-suspect medications included COVID-10 vaccine MRNA (MRNA 1273) (268), influenza vaccine (181), adalimumab (167), COVID-10 vaccine NRV V AD (CHADOX1 NCOV19) (108), COVID-10 vaccine (97), influenza vaccine INACT SAG 4V (53), influenza vaccine INACT SAG 4V (46), ocrelizumab (40), upadacitinib (26), pneumococcal vaccine polysacch 23V (23), influenza vaccine INACT SPLIT 3V (22), levithroxine, paracetamol (17 each), INJ 78436735 (16), ethinylestradiol, levonorgestrel (11), acetylsalicylic acid, desogestrel, hepatitis A vaccine (10 each).

- Number of relevant events: 310,383.

- Relevant event seriousness42: serious (36801), non-serious (273,863).

- Relevant PTs: Headache (77,970), Fatigue (67,855), Pyrexia (57,671), Myalgia (43,916), Chills (33,541), and Arthralgia (29,430).

- Time to event onset (n = 253,501133) range: <24 hours to 3654 days, median: 1 day.
  - <24 hours: 106,574 events (42 of which had a fatal outcome);
  - 1 day: 97,138 events (41 of which had a fatal outcome);
  - 2-7 days: 30,202 events (42 of which had a fatal outcome);
  - 8-14 days: 6818 events (21 of which had a fatal outcome);
  - 15-30 days: 5752 events (18 of which had a fatal outcome);
  - 31-180 days: 6175 events (40 of which had a fatal outcome);
  - 181-240 days: 472 events (none of which had a fatal outcome);
  - 241-365 days: 303 events (none of which had a fatal outcome);
  - 366-500 days: 56 events (none of which had a fatal outcome);
  - 501-3654 days: 11 events (none of which had a fatal outcome).

133 This number does not include events which occurred prior to vaccine administration.
• Duration of relevant events (n = 76,627\textsuperscript{134}, out of which 76,067 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 1 year, 2 months 8 days, median 2 days.
  - <24 hours: 6027 events;
  - 1-29 days: 67,465 events;
  - 30-365 days: 3126 events;
  - >365 days = 9 events.
• Relevant event outcome\textsuperscript{78}: fatal (300), resolved/resolving (175,756), resolved with sequelae (3756), not resolved (86,147), unknown (45,676).
  - In 233 cases, the following relevant events (300) were reported as fatal: PTs Pyrexia (119), Fatigue (63), Headache (55), Chills (29), Myalgia (20), and Arthralgia (14). More than half (124 of 233 cases, 53.2\%) of the cases with a fatal outcome involved elderly subjects.

Analysis by age group

CT: Paediatric (5, PTs Pyrexia [4], Myalgia [1]), Adults (1, PT Headache), Elderly (5, PTs Arthralgia [3], Pyrexia [2]).

  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

PM

  - An analysis of relevant PM events by age group, event seriousness and event outcome are provided in Table 54. Per the RSI, (CDS version 13.0, dated 10 May 2022), the most frequent systemic adverse reactions in subjects 16 years of age and older (in order from highest to lowest frequencies) after 2 doses were injection site pain (>80\%), fatigue (>60\%), headache (>50\%), myalgia (>40\%), chills (>30\%), arthralgia (>20\%), pyrexia and injection site swelling (>10\% each); and after booster dose, were injection site pain (>80\%), fatigue (>60\%), headache (>40\%), myalgia (>30\%), chills and arthralgia (>20\%). In adolescent subjects 12 through 15 years of age after 2 doses were injection site pain (>90\%), fatigue and headache (>70\%), myalgia and chills (>40\%), arthralgia and pyrexia (>20\%). In children 5 through <12 years of age after 2 doses were injection site pain (>80\%), fatigue (>50\%), headache (>30\%), injection site redness and swelling (>20\%), myalgia and chills (>10\%); and after booster dose

\textsuperscript{134} This number does not include events for which event onset dates or event cessation dates were not reported or events with a not meaningful time to event cessation value as per reported information.
were injection site pain (>70%), fatigue (>40%), headache (>30%), myalgia, chills, injection site redness, and swelling (>10%).

- In the current reporting interval, the most frequent systemic adverse reactions in subjects 16 years of age and older (in order from highest to lowest frequencies) were Pts Headache (69,392), Fatigue (61,567), Pyrexia (48,928), Myalgia (40,707), Chills (30,837) and Arthralgia (27,333); the most frequent systemic adverse reactions in adolescents 12 through 15 years of age were Pts Headache (4485), Pyrexia (4727), Fatigue (2381), Myalgia (1164), Chills (1121), Arthralgia (614). Across the age groups in the table below, the greatest number of events were reported in the adult population, followed by the elderly. In general, relevant events were more likely to be assessed as non-serious and/or associated with a resolving outcome with increasing age. Generally, there were less relevant events associated with a worse outcome (not resolved/fatal).

<table>
<thead>
<tr>
<th>Table 54. Analysis of Systemic Adverse Reactions by Age Group, Event seriousness and Event Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td><strong>Arthralgia</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 614 (4.2%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 118 (0.8 %)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 0 (0.0%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 200 (1.4%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 1 (&lt;0.1 %)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 294 (2.0%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 120 (0.8%)</td>
</tr>
<tr>
<td><strong>Arthralgia</strong> <strong>Total Events</strong> 24196 (9.4%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 3605 (1.4%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 5 (&lt;0.01 %)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 9051 (3.5%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 335 (0.1%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 11886 (4.6%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 3052 (1.2%)</td>
</tr>
<tr>
<td><strong>Arthralgia</strong> <strong>Total Events</strong> 3137 (14.0%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 665 (3.0%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 9 (&lt;0.1 %)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 1333 (5.9%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 72 (0.3%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 1197 (5.3%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 547 (2.4%)</td>
</tr>
<tr>
<td><strong>Arthralgia</strong> <strong>Total Events</strong> 1482 (8.7%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 314 (1.8%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 0 (0.0%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 502 (2.9%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 9 (0.1%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 501 (2.9%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 478 (2.8%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 1121 (7.7%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 144 (1.0%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 13 (&lt;0.1 %)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 232 (1.6%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 230 (1.9%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 707 (4.9%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 180 (1.2%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 28825 (11.2%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 2261 (0.9%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 15 (&lt;0.1 %)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 5390 (2.1%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 23 (0.1%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 20191 (7.9%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 3081 (1.2%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 2012 (9.0%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 278 (1.2%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 15 (0.1%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 405 (1.8%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 23 (0.1%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 1237 (5.5%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 334 (1.5%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 1582 (9.2%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 211 (1.2%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 0 (0.0%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 220 (1.3%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 5 (&lt;0.1 %)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 923 (5.4%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 436 (2.5%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 2381 (16.4%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 427 (2.9%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 17 (0.1%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 696 (4.8%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 170 (0.3%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 28495 (11.1%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 496 (3.4%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 4865 (21.7%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 949 (4.2%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 43 (0.2%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 1648 (7.4%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 93 (0.4%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 2065 (9.2%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 1812 (10.6%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 3904 (22.8%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 648 (3.8%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 3 (0.0%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 853 (5.0%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 17 (0.1%)</td>
</tr>
<tr>
<td><strong>Resolved/Resolving</strong> 1812 (10.6%)</td>
</tr>
<tr>
<td><strong>Unknown</strong> 1240 (7.2%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
</tr>
<tr>
<td><strong>Total Events</strong> 4090 (23.9%)</td>
</tr>
<tr>
<td><strong>Serious Events</strong> 4713 (21.0%)</td>
</tr>
<tr>
<td><strong>Event Outcome</strong>: Fatal 1055 (4.7%)</td>
</tr>
<tr>
<td><strong>Not Resolved</strong> 1240 (7.2%)</td>
</tr>
<tr>
<td><strong>Resolved with sequelae</strong> 4090 (23.9%)</td>
</tr>
</tbody>
</table>

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Page 268
Table 54. Analysis of Systemic Adverse Reactions by Age Group, Event seriousness and Event Outcome

<table>
<thead>
<tr>
<th></th>
<th>Paediatric N = 14,492</th>
<th>Adults N = 256,344</th>
<th>Elderly N = 22,420</th>
<th>Unknown N = 17,114</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serious Events</td>
<td>735 (5.1%)</td>
<td>7237 (2.8%)</td>
<td>824 (3.7%)</td>
<td>654 (3.8%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>6 (&lt;0.1%)</td>
<td>29 (&lt;0.1%)</td>
<td>18 (0.1%)</td>
<td>2 (&lt;0.1%)</td>
</tr>
<tr>
<td>NotResolved</td>
<td>1110 (7.7%)</td>
<td>19818 (7.7%)</td>
<td>1361 (6.1%)</td>
<td>973 (5.7%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>24 (0.2%)</td>
<td>931 (0.4%)</td>
<td>98 (0.4%)</td>
<td>26 (0.2%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>2523 (17.4%)</td>
<td>35703 (13.9%)</td>
<td>2412 (10.8%)</td>
<td>1916 (11.2%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>829 (5.7%)</td>
<td>8421 (3.3%)</td>
<td>834 (3.7%)</td>
<td>1186 (6.9%)</td>
</tr>
</tbody>
</table>

**Myalgia**

<table>
<thead>
<tr>
<th></th>
<th>Total Events</th>
<th>Serious Events</th>
<th>Event Outcome: Fatal</th>
<th>NotResolved</th>
<th>Resolved with sequelae</th>
<th>Resolved/Resolving</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Events</td>
<td>1164 (8.0%)</td>
<td>185 (1.3%)</td>
<td>12 (&lt;0.1%)</td>
<td>322 (2.2%)</td>
<td>5 (0.03%)</td>
<td>612 (4.2%)</td>
<td>226 (1.6%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>37276 (14.5%)</td>
<td>3465 (1.4%)</td>
<td>12 (&lt;0.1%)</td>
<td>11931 (4.7%)</td>
<td>587 (0.2%)</td>
<td>20522 (8.0%)</td>
<td>4306 (1.7%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>3431 (15.3%)</td>
<td>535 (2.4%)</td>
<td>7 (&lt;0.1%)</td>
<td>1371 (6.1%)</td>
<td>84 (0.4%)</td>
<td>1466 (6.6%)</td>
<td>492 (2.2%)</td>
</tr>
<tr>
<td>NotResolved</td>
<td>2043 (11.9%)</td>
<td>266 (1.6%)</td>
<td>1 (&lt;0.1%)</td>
<td>513 (3.0%)</td>
<td>16 (0.1%)</td>
<td>989 (5.8%)</td>
<td>528 (3.1%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>2685 (10.9%)</td>
<td>295 (1.1%)</td>
<td>2 (&lt;0.1%)</td>
<td>434 (1.9%)</td>
<td>10 (0.1%)</td>
<td>524 (3.1%)</td>
<td>369 (2.2%)</td>
</tr>
</tbody>
</table>

**Pyrexia**

<table>
<thead>
<tr>
<th></th>
<th>Total Events</th>
<th>Serious Events</th>
<th>Event Outcome: Fatal</th>
<th>NotResolved</th>
<th>Resolved with sequelae</th>
<th>Resolved/Resolving</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Events</td>
<td>4727 (32.6%)</td>
<td>838 (5.8%)</td>
<td>10 (&lt;0.1%)</td>
<td>704 (4.9%)</td>
<td>13 (0.1%)</td>
<td>3202 (22.1%)</td>
<td>1201 (7.7%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>44666 (17.4%)</td>
<td>4608 (1.8%)</td>
<td>45 (&lt;0.1%)</td>
<td>7166 (2.8%)</td>
<td>348 (0.1%)</td>
<td>30692 (12.0%)</td>
<td>6532 (2.5%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>4262 (19.0%)</td>
<td>756 (3.4%)</td>
<td>61 (0.3%)</td>
<td>561 (2.5%)</td>
<td>50 (0.2%)</td>
<td>2623 (11.7%)</td>
<td>974 (4.3%)</td>
</tr>
<tr>
<td>NotResolved</td>
<td>4013 (23.4%)</td>
<td>429 (2.5%)</td>
<td>3 (&lt;0.1%)</td>
<td>361 (2.1%)</td>
<td>9 (0.1%)</td>
<td>2439 (14.3%)</td>
<td>1215 (7.1%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>989 (5.8%)</td>
<td>951 (3.5%)</td>
<td>3 (&lt;0.1%)</td>
<td>486 (2.7%)</td>
<td>5 (0.3%)</td>
<td>834 (4.6%)</td>
<td>157 (0.9%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>528 (3.1%)</td>
<td>369 (2.2%)</td>
<td>1 (&lt;0.1%)</td>
<td>342 (1.9%)</td>
<td>5 (0.3%)</td>
<td>528 (3.1%)</td>
<td>157 (0.9%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>1201 (7.7%)</td>
<td>1201 (7.7%)</td>
<td>2 (&lt;0.1%)</td>
<td>157 (0.9%)</td>
<td>5 (0.3%)</td>
<td>1201 (7.7%)</td>
<td>157 (0.9%)</td>
</tr>
</tbody>
</table>

*a.* Multiple episodes of the same event were reported with different clinical outcomes in some cases hence the sum of the events for outcome may differ.

**N:** Total number of events in the population subset; **n:** number of events; percentage (%) calculated as **n/N.**

**Analysis by presence of comorbidities**

Number of subjects with comorbidities: 13,030 (2.6% of 508,351 cases in the total dataset and 7.8% of 167,771 [11 CT and 167,760 PM] cases reporting systemic adverse reactions).

**CT:**

- None of the CT cases reported selected comorbidities.

**PM:**

- An analysis of relevant PM events by presence of selected comorbidities, event seriousness and event outcome is provided in Table 55. The total proportion of relevant events were generally evenly distributed among subjects that reported selected comorbidities and subjects that did not report selected comorbidities. In subjects with selected comorbidities, the relevant event was more likely to be assessed as non-serious and/or with a resolved or resolving event outcome. Of note, subjects
that reported comorbidities were more likely to be of advanced age, polypharmacy users, report more AEs on average (e.g., concurrent conditions) and/or prone to hospitalisation; therefore, assessment of the contributory role of BNT162b2 on the seriousness and outcome of these relevant events is confounded.

Table 55. Analysis of Systemic Adverse Reactions by Presence of Comorbidities, Event Seriousness and Event Outcome

<table>
<thead>
<tr>
<th></th>
<th>Without Comorbidities</th>
<th>With Comorbidities</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N = 288685</td>
<td>N = 22298</td>
</tr>
<tr>
<td></td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td><strong>Arthralgia</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>26701 (9.3%)</td>
<td>2729 (12.2%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>23750 (8.2%)</td>
<td>1019 (4.6%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>13 (&lt;0.1%)</td>
<td>1 (&lt;0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>9994 (3.5%)</td>
<td>1092 (4.9%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>368 (0.1%)</td>
<td>49 (0.2%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>12832 (4.5%)</td>
<td>1059 (4.7%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3638 (1.3%)</td>
<td>559 (2.5%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>31535 (10.9%)</td>
<td>2006 (9.0%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>2384 (0.8%)</td>
<td>510 (2.3%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>19 (&lt;0.1%)</td>
<td>10 (&lt;0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>5844 (2.0%)</td>
<td>404 (1.8%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>190 (0.1%)</td>
<td>30 (0.1%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>21867 (7.6%)</td>
<td>1209 (5.4%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3671 (1.3%)</td>
<td>360 (1.6%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>62443 (21.7%)</td>
<td>5412 (24.3%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>6977 (2.4%)</td>
<td>1696 (7.6%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>37 (&lt;0.1%)</td>
<td>26 (0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>20820 (7.2%)</td>
<td>1800 (8.1%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>815 (0.3%)</td>
<td>113 (0.5%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>31487 (10.9%)</td>
<td>2109 (9.5%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>9670 (3.4%)</td>
<td>1430 (6.4%)</td>
</tr>
<tr>
<td><strong>Headache</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>72452 (25.1%)</td>
<td>5518 (24.7%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>7797 (2.7%)</td>
<td>1653 (7.4%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>45 (&lt;0.1%)</td>
<td>10 (&lt;0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>21581 (7.5%)</td>
<td>1682 (7.5%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>984 (0.3%)</td>
<td>95 (0.4%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>39917 (13.9%)</td>
<td>2673 (12.0%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>10186 (3.5%)</td>
<td>1086 (4.9%)</td>
</tr>
<tr>
<td><strong>Myalgia</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>41272 (14.3%)</td>
<td>2644 (11.9%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>3630 (1.3%)</td>
<td>821 (3.7%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>16 (&lt;0.1%)</td>
<td>4 (&lt;0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>13190 (4.6%)</td>
<td>947 (4.2%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>624 (0.2%)</td>
<td>68 (0.3%)</td>
</tr>
</tbody>
</table>
Table 55. Analysis of Systemic Adverse Reactions by Presence of Comorbidities, Event Seriousness and Event Outcome

<table>
<thead>
<tr>
<th></th>
<th>Without Comorbidities</th>
<th>With Comorbidities</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N = 288085</td>
<td>N = 22298</td>
</tr>
<tr>
<td></td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>22437 (7.8%)</td>
<td>1189 (5.3%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>5109 (1.8%)</td>
<td>444 (2.0%)</td>
</tr>
</tbody>
</table>

Pyrexia

<table>
<thead>
<tr>
<th></th>
<th>Without Comorbidities</th>
<th>With Comorbidities</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N = 38562</td>
<td>N = 3989</td>
</tr>
<tr>
<td></td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td>Total Events</td>
<td>53682 (18.6%)</td>
<td>3989 (17.9%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>5362 (1.9%)</td>
<td>1269 (5.7%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>77 (&lt;0.1%)</td>
<td>42 (0.2%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>8104 (2.8%)</td>
<td>689 (3.1%)</td>
</tr>
<tr>
<td>Resolved with sequela</td>
<td>367 (0.1%)</td>
<td>53 (0.2%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>36678 (12.7%)</td>
<td>2299 (10.3%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>8594 (3.0%)</td>
<td>929 (4.2%)</td>
</tr>
</tbody>
</table>

a. Multiple episodes of the same event were reported with different clinical outcomes in some cases hence the sum of the events for outcome may differ.
N: Total number of events in the population subset; n: number of events; percentage (%) calculated as n/N.

Analysis by dose

Number of vaccine doses administered: 1 dose in 47,268 cases, 2 doses in 49,553 cases; 3 doses in 44,738 cases, 4 doses in 893 cases, and in 25,515 cases the dose was either not specified or reported as others.

CT:
- Vaccination dose number: 2 doses (3), 3 doses (7) and 4 doses (1).
- A meaningful comparison by dose is not possible due to the low number of CT cases.

PM:
- An analysis of relevant PM events by dose, event seriousness and event outcome are provided in Table 56. In general, the total proportion of relevant events, event seriousness, and event outcome were highest in those subjects who had received three doses of the vaccine; following this, most events were reported in those who had received two doses of the vaccine.
Table 56. Analysis of Systemic Adverse Reactions by Dose, Event Seriousness and Event Outcome

<table>
<thead>
<tr>
<th></th>
<th>1 Dose N = 78166 n (%)</th>
<th>2 Doses N = 92344 n (%)</th>
<th>3 Doses N = 94444 n (%)</th>
<th>4 Doses N = 1607 n (%)</th>
<th>Dose Not Specified/Other N = 44080 n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Arthralgia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>7235 (9.3%)</td>
<td>8451 (9.2%)</td>
<td>9471 (10.0%)</td>
<td>164 (10.2%)</td>
<td>4147 (9.4%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>925 (1.2%)</td>
<td>1497 (1.6%)</td>
<td>1920 (2.0%)</td>
<td>61 (3.8%)</td>
<td>312 (0.7%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>2 (0.03%)</td>
<td>2 (0.002%)</td>
<td>2 (0.002%)</td>
<td>0 (0.0%)</td>
<td>8 (0.02%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>2866 (3.7%)</td>
<td>3143 (3.4%)</td>
<td>3590 (3.8%)</td>
<td>66 (4.1%)</td>
<td>1442 (3.3%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>138 (0.2%)</td>
<td>167 (0.2%)</td>
<td>75 (0.1%)</td>
<td>1 (0.1%)</td>
<td>40 (0.1%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>3128 (4.0%)</td>
<td>3795 (4.1%)</td>
<td>4810 (5.1%)</td>
<td>63 (3.9%)</td>
<td>2099 (4.8%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>1143 (1.5%)</td>
<td>1405 (1.5%)</td>
<td>1054 (1.1%)</td>
<td>38 (2.4%)</td>
<td>571 (1.3%)</td>
</tr>
<tr>
<td><strong>Chills</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>5860 (7.5%)</td>
<td>10310 (11.2%)</td>
<td>12888 (13.6%)</td>
<td>228 (14.2%)</td>
<td>4270 (9.7%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>403 (0.5%)</td>
<td>731 (0.8%)</td>
<td>1506 (1.6%)</td>
<td>66 (4.1%)</td>
<td>191 (0.4%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>3 (0.004%)</td>
<td>6 (0.01%)</td>
<td>7 (0.01%)</td>
<td>1 (0.1%)</td>
<td>12 (0.03%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>1312 (1.7%)</td>
<td>1617 (1.8%)</td>
<td>2591 (2.7%)</td>
<td>43 (2.7%)</td>
<td>689 (1.6%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>68 (0.1%)</td>
<td>81 (0.1%)</td>
<td>45 (0.05%)</td>
<td>1 (0.1%)</td>
<td>26 (0.1%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>3650 (4.7%)</td>
<td>7163 (7.8%)</td>
<td>9223 (9.8%)</td>
<td>124 (7.7%)</td>
<td>2923 (6.6%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>833 (1.1%)</td>
<td>1472 (1.6%)</td>
<td>1048 (1.1%)</td>
<td>59 (3.7%)</td>
<td>624 (1.4%)</td>
</tr>
<tr>
<td><strong>Fatigue</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>19275 (24.7%)</td>
<td>20318 (22.0%)</td>
<td>20040 (21.2%)</td>
<td>365 (22.7%)</td>
<td>7923 (18.0%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>1821 (2.3%)</td>
<td>2636 (2.9%)</td>
<td>3578 (3.8%)</td>
<td>100 (6.2%)</td>
<td>557 (1.3%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>14 (0.02%)</td>
<td>15 (0.02%)</td>
<td>20 (0.02%)</td>
<td>4 (0.2%)</td>
<td>10 (0.02%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>6141 (7.9%)</td>
<td>6625 (7.2%)</td>
<td>7214 (7.6%)</td>
<td>95 (5.9%)</td>
<td>2567 (5.8%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>333 (0.4%)</td>
<td>354 (0.4%)</td>
<td>140 (0.1%)</td>
<td>6 (0.4%)</td>
<td>100 (0.2%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>9654 (12.4%)</td>
<td>9752 (10.6%)</td>
<td>10117 (10.7%)</td>
<td>141 (8.8%)</td>
<td>3956 (9.0%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3224 (4.1%)</td>
<td>3744 (4.1%)</td>
<td>2694 (2.9%)</td>
<td>124 (7.7%)</td>
<td>1325 (3.0%)</td>
</tr>
<tr>
<td><strong>Headache</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>21087 (27.0%)</td>
<td>22687 (24.6%)</td>
<td>22345 (23.7%)</td>
<td>336 (20.9%)</td>
<td>11546 (26.2%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>2017 (2.6%)</td>
<td>2789 (3.0%)</td>
<td>3801 (4.0%)</td>
<td>100 (6.2%)</td>
<td>754 (1.7%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>5 (0.01%)</td>
<td>16 (0.02%)</td>
<td>14 (0.01%)</td>
<td>0 (0.0%)</td>
<td>20 (0.05%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>7047 (9.0%)</td>
<td>6399 (6.9%)</td>
<td>6940 (7.3%)</td>
<td>86 (5.4%)</td>
<td>2801 (6.4%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>402 (0.5%)</td>
<td>389 (0.4%)</td>
<td>173 (0.2%)</td>
<td>2 (0.1%)</td>
<td>119 (0.3%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>10387 (13.3%)</td>
<td>12253 (13.3%)</td>
<td>12977 (13.7%)</td>
<td>167 (10.4%)</td>
<td>6815 (15.5%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>3312 (4.2%)</td>
<td>3756 (4.1%)</td>
<td>2322 (2.3%)</td>
<td>82 (5.1%)</td>
<td>1808 (4.1%)</td>
</tr>
</tbody>
</table>
Table 56. Analysis of Systemic Adverse Reactions by Dose, Event Seriousness and Event Outcome

<table>
<thead>
<tr>
<th></th>
<th>1 Dose N = 78166 n (%)</th>
<th>2 Doses N = 92344 n (%)</th>
<th>3 Doses N = 94444 n (%)</th>
<th>4 Doses N = 1607 n (%)</th>
<th>Dose Not Specified/Other N = 44080 n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Myalgia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>10242 (13.1%)</td>
<td>13060 (14.1%)</td>
<td>14189 (15.0%)</td>
<td>179 (11.1%)</td>
<td>6262 (14.2%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>875 (1.1%)</td>
<td>1442 (1.6%)</td>
<td>1820 (1.9%)</td>
<td>48 (3.0%)</td>
<td>274 (0.6%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>3 (0.004%)</td>
<td>4 (0.004%)</td>
<td>8 (0.01%)</td>
<td>1 (0.1%)</td>
<td>4 (0.01%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>3622 (4.6%)</td>
<td>3972 (4.3%)</td>
<td>4818 (5.1%)</td>
<td>56 (3.5%)</td>
<td>1677 (3.8%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>228 (0.3%)</td>
<td>296 (0.3%)</td>
<td>97 (0.1%)</td>
<td>1 (0.1%)</td>
<td>73 (0.2%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>4740 (6.1%)</td>
<td>6786 (7.3%)</td>
<td>8202 (8.7%)</td>
<td>86 (5.4%)</td>
<td>3811 (8.6%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>1676 (2.1%)</td>
<td>2047 (2.2%)</td>
<td>1095 (1.2%)</td>
<td>35 (2.2%)</td>
<td>706 (1.6%)</td>
</tr>
<tr>
<td><strong>Pyrexia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total Events</td>
<td>14467 (18.5%)</td>
<td>17518 (19.0%)</td>
<td>15511 (16.4%)</td>
<td>335 (20.8%)</td>
<td>9932 (22.5%)</td>
</tr>
<tr>
<td>Serious Events</td>
<td>1129 (1.4%)</td>
<td>2027 (2.2%)</td>
<td>2800 (3.0%)</td>
<td>114 (7.1%)</td>
<td>575 (1.3%)</td>
</tr>
<tr>
<td>Event Outcome: Fatal</td>
<td>13 (0.02%)</td>
<td>29 (0.03%)</td>
<td>30 (0.03%)</td>
<td>6 (0.4%)</td>
<td>42 (0.1%)</td>
</tr>
<tr>
<td>Not Resolved</td>
<td>2382 (3.0%)</td>
<td>2363 (2.6%)</td>
<td>2861 (3.0%)</td>
<td>46 (2.9%)</td>
<td>1159 (2.6%)</td>
</tr>
<tr>
<td>Resolved with sequelae</td>
<td>104 (0.1%)</td>
<td>149 (0.2%)</td>
<td>99 (0.1%)</td>
<td>6 (0.4%)</td>
<td>62 (0.1%)</td>
</tr>
<tr>
<td>Resolved/Resolving</td>
<td>9807 (12.5%)</td>
<td>11559 (12.5%)</td>
<td>10224 (10.8%)</td>
<td>185 (11.5%)</td>
<td>7250 (16.4%)</td>
</tr>
<tr>
<td>Unknown</td>
<td>2200 (2.8%)</td>
<td>3464 (3.8%)</td>
<td>2364 (2.5%)</td>
<td>94 (5.8%)</td>
<td>1427 (3.2%)</td>
</tr>
</tbody>
</table>

a. Multiple episodes of the same event were reported with different clinical outcomes in some cases hence the sum of the events for outcome may differ.

N: Total number of events in the population subset; n: number of events; percentage (%) calculated as n/N.

Conclusion

Systemic adverse reactions were reported in 167,771 (11 CT and 167,760 PM) cases representing 33.0% of the cases in the total dataset for the reporting period. The majority of events (88.2%) were non-serious events with 57.8% of the events resolved, resolved with sequelae or resolving at the time of reporting. Evaluation of systemic adverse reaction cases did not reveal any significant new safety information based on analysis by age group, by presence of comorbidities or by dose. Systemic adverse reactions are appropriately described in the RSI. Surveillance of systemic adverse reactions will continue.

16.3.3.5. Severe Reactogenicity

Search criteria - PT Extensive swelling of vaccinated limb.

Upon review, 4 cases were determined to be non-contributory and were not included in the discussion since they involved exposure to the vaccine during the mother’s pregnancy or through breastfeeding.
Clinical Trial Data

During the current reporting interval, there were no serious CT cases indicative of extensive swelling of vaccinated limb; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 1613 (0.32% of 507,683 cases, the total PM dataset), compared to 1558 cases (0.24%) retrieved in the PSUR #2.
- MC cases (196), NMC cases (1417).
- Country of incidence: Netherlands (921), Belgium (590), Iraq (26), Australia (24), UK (12), France, Germany (8 each), Philippines (5); the remaining 19 cases were distributed among 10 countries.
- Subjects' gender: female (1310), male (300) and unknown (3).
- Subjects' age in years (n = 1536), range: 7 – 94, mean: 38.3, median: 36.0.
- Medical history (n = 497): the relevant reported medical conditions included Drug hypersensitivity (24), Hypersensitivity (8), Allergic reaction to excipient, Allergy to vaccine, Reaction to preservatives (1 each).
- COVID-19 Medical history (n = 219): medical conditions reported included COVID-19 (162), Suspected COVID-19 (54), Post-acute COVID-19 syndrome (2), and SARS-CoV-2 test positive (1).
- Co-suspects (n= 17 cases): Influenza vaccine (6), Pneumococcal vaccine polysacch 23V (2).
- Number of relevant events: 1613
- Relevant event seriousness: serious (202), non-serious (1,411).
- Time to event onset (n = 1450)\(^\text{135}\), range: range: <24 hours to 175 days, median: 1 day.
  - <24 hours: 589 events;
  - 1 day: 649 events;
  - 2-7 days: 185 events;
  - 8-14 days: 9 events;
  - 15-30 days: 7 events;
  - 31-180 days: 11 events.
- Duration of relevant events (n = 375 out of 1,615 occurrences with outcome of resolved/resolved with sequelae), range: <24 hours to 157 days, median 4 days.
  - <24 hours: 6 events;

\(^{135}\) This number does not include 165 events for which partial administration and/or event onset dates were reported or events with a not meaningful time to onset value as per reported information.
COVID-19 mRNA vaccine (nucleoside modified)
Periodic Safety Update Report (PSUR) 3
19 December 2021 through 18 June 2022

- 1 day: 43 events;
- 2 - 7 days: 278 events;
- 8-14 days: 30 events;
- 15-30 days: 7 events;
- 31-180 days: 11 events.

- Relevant event outcome: 78 fatal (1), resolved/resolving (910), resolved with sequelae (8), not resolved (583), unknown (112).

The reported relevant PT included Extensive swelling of vaccinated limb (1613). During the reporting period, 1 case was received from a Health Authority, reporting the relevant PT Extensive swelling of vaccinated limb with a fatal outcome. This case described a 14-year-old male patient who received BNT162b2 intramuscularly for COVID-19 immunisation and experienced swelling of limb. The patient also experienced difficulty breathing (PT Dyspnoea), cyanosis (PT Cyanosis) and oedematous lower extremities (PT Oedema), all of which were reported as non-serious events. The reported cause of death was peripheral swelling. Limited information was provided in this case precluding a meaningful medical assessment, including a lack of event onset dates, event details, test results, medical history, and concomitant medications.

A majority of the cases did not describe the type or extent of swelling and reported (verbatim) terms such as, “extensive swelling of the arm, reaction at or around the injection site, swelling limb, or extended swelling of the arm: extensive swelling of vaccinated limb”. Many cases also reported additional events related to pain, warmth, or erythema at the injection site, with no additional relevant details. Most cases described localized redness or swelling limited to the injection site and/or reports of lymph node swelling with no evidence in the case detail regarding any additional extensive swelling. For those cases reporting details of swelling, most appeared limited to the area surrounding the injection site with little evidence of additional extensive swelling of the rest of the limb. In a majority of the cases reporting swelling associated with the injection site, it was not reported if treatment was required, and no case reported long lasting or permanent sequelae following the event.

Analysis by age group

PM: Paediatric (25), Adult (1506), Elderly (65), Unknown (17).

A higher reporting proportion of events coded to the PT Extensive swelling of vaccinated limb was observed in elderly versus adult population (26.5% in elderly vs 20.3% in adults). Due to the relative low volume of cases in the paediatric population, a meaningful comparison of the same with the other age groups was not possible.

Analysis by presence of comorbidities

Number of subjects reporting comorbidities: 51 (3.2% of the cases reporting the event severe reactogenicity). A higher reporting proportion of severe reactogenicity was reported in patients without significant comorbidities (96.9%) when compared to patients with significant comorbidities.

CONFIDENTIAL
Page 275
The reporting proportion of the event severe reactogenicity with the outcome of resolved/resolving (58.8%) is slightly higher in individuals with comorbid conditions when compared to the reporting proportion observed in the individuals without comorbidities (56.3 % of events with resolved/resolving).

Conclusion

There was a total of 1613 cases in the safety database reporting the PT Extensive swelling of vaccinated limb with the use of BNT162b2 which were mostly reported from the Netherlands (921) and Belgium (590). A majority of the cases involved females (1310, 81.2%) and were reported in subjects aged 31-50 years (793, 49.2%). Two-hundred and two (202; 12.5%) of the events were assessed as serious due to meeting medically significant criteria (there were 6 hospitalisations due to reported events). There was 1 case reporting a fatal outcome. One thousand two hundred and thirty-seven (1237) cases reported time to onset of the event as the same day or the day following vaccination. The majority of cases reporting swelling associated with the injection site, did not report that treatment was required, and no case reported long lasting or permanent sequelae following the event.

Injection site swelling and lymphadenopathy are listed adverse drug reactions in the RSI for BNT162b2, and based on the data reviewed, there is insufficient evidence from reported cases to date that would warrant a change to the existing product information.

16.3.3.6. Age-Related Adverse Reactions

All adverse events reported during the reporting period were reviewed in the context of age categories. For the overall demographic information for all CT and PM cases refer to Section 6.3.1.1 General Overview of the Safety Database – All Cases.

Clinical Trial Data

- Number of cases: 668 (cross-referenced to Section 6.3.1.1.1 General Overview of the Safety Database - Clinical Trials Data)

- Time to event onset (n = 793), range: <24 hours to 558 days, median: 116 days.
  - <24 hours: 5 events (none of which had a fatal outcome);
  - 1 day: 8 events;
  - 2-7 days: 28 events;
  - 8-14 days: 11 events;
  - 15-30 days: 47 events;
  - 31-180 days: 548 events;
  - >181 days: 146 events.

- Relevant event outcome: fatal (50), resolved/resolving (663), resolved with sequelae (49), not resolved (115), unknown (3).
Post-Authorisation Data

- Number of cases: 507,683 (cross-referenced to Section 6.3.1.1.2 General Overview of the Safety Database – Post-Authorisation Data)
- Time to event onset (n = 1,196,069), range: <24 hours to 7337 days, median: 1 day.
  - <24 hours: 477,739 events (1067 of which had a fatal outcome);
  - 1 day: 284,078 events;
  - 2-7 days: 182,163 events;
  - 8-14 days: 57,900 events;
  - 15-30 days: 54,875 events;
  - 31-180 days: 127,948 events;
  - >181 days: 11,366 events.

- Relevant event outcome: 78% fatal (8526), resolved/resolving (595,395), resolved with sequelae (26,518), not resolved (434,513), unknown (536,733).

Analysis by age group

- CT: Paediatric (103), Adults (336), Elderly (211) and Unknown (1).

The 5 MedDRA SOCs with the most frequently reported events for the current reporting period for each age group is presented are Table 57, Table 58 and Table 59. Of note, 139 cases reported 151 events pertaining the Infections and infestations SOC, which was included among the SOCs of the most frequently reported AEs in all 3 age groups.

There were 59 cases reporting 65 events in the Cardiac disorders SOC for the adult and elderly age group. Fortyt-five (45) cases reported relevant medical history (e.g., coronary artery disease, atrial fibrillation, congestive cardiac failure, cardiovascular disorder), which may have contributed to the relevant events. The most frequently reported events (≥3 occurrences) in the Cardiac disorders SOC for the adult and elderly age group were Atrial fibrillation (16), Myocardial infarction (9), Cardiac failure congestive, Coronary artery disease (5 each), Acute coronary syndrome, Acute myocardial infarction (4 each), Angina pectoris and Angina unstable (3).

There were 96 cases reporting 98 events in the Neoplasms benign, malignant and unspecified (incl cysts and polyps) SOC for the adult and elderly age group. Twenty-eight (28) cases reported pre-existing medical history of cancer (e.g., basal cell carcinoma, neoplasm malignant, pituitary tumour benign, prostate cancer). The most frequently reported events (≥3 occurrences) in the Neoplasms benign, malignant and unspecified (incl cysts and polyps) SOC for the adult and elderly age group were Prostate cancer (13), Adenocarcinoma of colon, Breast cancer, Pancreatic carcinoma (5 each), Brain neoplasm (4), Invasive ductal breast carcinoma, and Oesophageal carcinoma (3 each). When reported, latency ranged from 1 day to 437 days with a median of 104 days. Of the 78 events reporting latency, the majority of the relevant event latency (65 events) was reported between 1 day to 6 months.
There were 7 cases reporting 9 events in the Psychiatric disorders SOC for the paediatric age group. The 9 events reported were Depression, Suicidal ideation, Suicide attempt (2 each), Depression suicidal, Major depression and Mental status changes (1 each). The events were assessed as unrelated to BNT162b2/Blinded therapy by the investigator and the Sponsor.

Table 57. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Adult Age Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Adult</th>
<th>Paediatric</th>
<th>Elderly</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>76</td>
<td>50</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>58</td>
<td>9</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>Neoplasms benign, malignant and unspecified (incl cysts and polyps)</td>
<td>44</td>
<td>1</td>
<td>54</td>
<td>0</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>33</td>
<td>1</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>31</td>
<td>11</td>
<td>19</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 58. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Paediatric Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Paediatric</th>
<th>Adult</th>
<th>Elderly</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>50</td>
<td>76</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>11</td>
<td>31</td>
<td>19</td>
<td>0</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>9</td>
<td>58</td>
<td>16</td>
<td>0</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>9</td>
<td>28</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>Psychiatric disorders</td>
<td>9</td>
<td>18</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 59. Clinical Trial Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Elderly Age Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Elderly</th>
<th>Adult</th>
<th>Paediatric</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neoplasms benign, malignant and unspecified (incl cysts and polyps)</td>
<td>54</td>
<td>44</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>32</td>
<td>33</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>26</td>
<td>28</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>25</td>
<td>76</td>
<td>50</td>
<td>0</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>22</td>
<td>16</td>
<td>2</td>
<td>0</td>
</tr>
</tbody>
</table>

The distribution of the most frequently reported serious PTs (≥ 2%) by age group in the 651 CT cases where the participants were directly exposed to BNT162b2, is shown in Figure 13 below.
Figure 13. Events Reported in ≥2% of All Clinical Trial Cases by Age Group

- PM: Paediatric (31,832), Adults (361,138), Elderly (56,588) and Unknown (56,647).

The top 5 MedDRA SOCs with the most frequently reported events for the current reporting period for each age group are presented in Table 60, Table 61, and Table 62. The top 5 SOCs were generally comparable for all age groups except Reproductive system and breast disorders in the adult age group, Skin and subcutaneous tissue disorders in the paediatric age group and Infections and infestations in the elderly age group.

In the Reproductive system and breast disorders SOC for adult age group, event seriousness was assessed as serious (8609) and non-serious (61,891). Event outcome was reported as resolved/resolving (19,328), not resolved (33,732), resolved with sequelae (1,390), unknown (16,268), and fatal (6). The most commonly reported PTs (>1000 occurrences) in Reproductive system and breast disorders for the adult age group were Heavy menstrual bleeding (11,691), Menstrual disorder (11,655), Menstruation irregular (6481), Dysmenorrhea (5824), Intermenstrual bleeding (5650), Amenorrhea (5267), Polymenorrhoea (4522), Menstruation delayed (4500), Oligomenorrhea (1818), Breast pain (1816), Vaginal haemorrhage (1588), and Postmenopausal haemorrhage (1028). It is not unexpected for these events of reproductive system and breast disorders to be reported more frequently in adult subjects compared to elderly and paediatric subjects (males or females of non-puberty age).

In the Skin and subcutaneous tissue disorders SOC for paediatric age group, event seriousness was assessed as serious (966) and non-serious (4194). Event outcome was reported as resolved/resolving (2903), not resolved (1161), resolved with sequelae (20), unknown (1085), and fatal (3). The fatal cases are reviewed in Section 16.3.4.1 Death. The
most commonly reported PTs (≥110 occurrences) in Skin and subcutaneous tissue disorders for the paediatric age group were Rash (1538), Pruritus (718), Urticaria (681), Erythema (326), Hyperhidrosis (270), Rash pruritic (198), Cold sweat (131), and Sensitive skin (110). Most of these events are listed or consistent with listed events as per the current RSI.

In the Infections and infestations SOC for elderly age group, event seriousness was assessed as serious (11,447) and non-serious (2756). Event outcome was reported as resolved/resolving (3096), not resolved (2014), resolved with sequelae (157), unknown (8305), and fatal (649). The fatal cases are reviewed in Section 16.3.4.1 Death. The most commonly reported PTs (>250 occurrences) in Infections and infestations for the elderly age group were coded to the PTs COVID-19 (8394), Herpes zoster (1771), Suspected COVID-19 (462), COVID-19 pneumonia (408), Influenza (407), Pneumonia (346), and Nasopharyngitis (251). It is not unexpected for these events to be reported more frequently in elderly subjects compared to adult and paediatric age groups.

Table 60. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Adult Age Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Adult</th>
<th>Elderly</th>
<th>Paediatric</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>367,077</td>
<td>42,066</td>
<td>23,304</td>
<td>27,093</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>160,280</td>
<td>20,808</td>
<td>12,369</td>
<td>10,612</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>120,125</td>
<td>15,881</td>
<td>4,256</td>
<td>8,578</td>
</tr>
<tr>
<td>Reproductive system and breast disorders</td>
<td>70,479</td>
<td>508</td>
<td>2,048</td>
<td>4,688</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>62,657</td>
<td>8,009</td>
<td>6,830</td>
<td>4,159</td>
</tr>
</tbody>
</table>

Table 61. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Paediatric Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Paediatric</th>
<th>Adult</th>
<th>Elderly</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>23,304</td>
<td>367,077</td>
<td>42,066</td>
<td>27,093</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>12,369</td>
<td>160,280</td>
<td>20,808</td>
<td>10,612</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>9,921</td>
<td>61,730</td>
<td>12,498</td>
<td>44,688</td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>6,830</td>
<td>62,657</td>
<td>8,009</td>
<td>4,159</td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td>5,158</td>
<td>44,803</td>
<td>8,423</td>
<td>3,972</td>
</tr>
</tbody>
</table>

Table 62. Post-Authorisation Data: Number of AEs in the Top 5 Frequently Reported System Organ Classes for Elderly Age Group Compared with Other Age Groups

<table>
<thead>
<tr>
<th>SOC</th>
<th>Elderly</th>
<th>Adult</th>
<th>Paediatric</th>
<th>Unknown</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>42,066</td>
<td>367,077</td>
<td>23,304</td>
<td>27,093</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>20,808</td>
<td>160,280</td>
<td>12,369</td>
<td>10,612</td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>15,881</td>
<td>120,125</td>
<td>4,256</td>
<td>8,578</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>14,201</td>
<td>60,151</td>
<td>3,655</td>
<td>4,094</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>12,498</td>
<td>61,730</td>
<td>9,921</td>
<td>44,688</td>
</tr>
</tbody>
</table>
The distribution of the most frequently reported overall PTs (≥ 2%) by age group is shown in Figure 14. Most of these events are listed or consistent with listed events as per the current RSI.

**Figure 14. Events Reported in ≥ 2% of All Post-marketing Cases by Age Group**

![Graph showing distribution of events by age group](image)

**Conclusion**

The most frequently reported SOCs and overall PTs with the COVID-19 vaccine across the age groups were listed or consistent with listed events as per the current RSI. The analysis of age-related AEs did not identify any new significant safety information.

**16.3.3.7. Vaccination Stress/Anxiety related ADRs**

Search criteria - PTs: Anxiety; Blood pressure decreased; Blood pressure increased; Dizziness; Dyspnoea; Hyperhidrosis; Loss of consciousness; Palpitations; Paraesthesia; Paraesthesia oral; Syncope; Tachycardia (reported in very close temporal proximity to
vaccination, e.g., when time to event onset for the relevant PTs is same day or 1 day after vaccination\(^{136}\).

Of the 82,924 cases including PTs indicative of vaccination stress/anxiety related ADRs, 43,122 cases were determined to be non-contributory and were not included in the discussion:

- since exposure to the vaccine occurred during the mother’s pregnancy or through breastfeeding (4 cases)
- since the relevant PTs were reported with time to event onset ≥ 2 days, unknown or with meaningless values (43,118 cases).

**Clinical Trial Data**

- Number of cases: 2, both involving BNT162b2 (0.3 % of 668 cases in the total CT dataset) compared to no cases\(^{137}\) retrieved in the PSUR #2.
- Country of incidence: Israel, Poland (1 each).
- Subjects’ gender: male (2).
- Subjects’ age in years (n = 2), 10 years and 73 years.
- Medical history: Hypertension, Hypercholesterolaemia, Myocardial ischaemia and Glucose-6-phosphate dehydrogenase deficiency (1 each).
- COVID-19 Medical history: none.
- Co-suspects: none.
- Reported relevant PTs: Dyspnoea and Syncope (1 each).
- Time to event onset: 1 days for both the relevant events.
- Duration of relevant events: 1 day for the event Syncope, 5 days for the event Dyspnoea.
- Relevant event outcome: resolved (2).

**Post-Authorisation Data**

- Number of relevant cases: 39,800 (7.8% of 507,683 cases, the total PM dataset), compared to 56,230 cases\(^{138}\) (8.6%) retrieved in the PSUR #2.

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\(^{136}\) To have consistency with the concept of vaccination stress/anxiety related ADRs, the search criteria has been restricted to relevant PTs with time to event onset equal to same day or 1 day after vaccination.

\(^{137}\) In PSUR #2, 15 cases originating from clinical trials reported the relevant PTs, but none of these cases included events indicative of vaccination stress/anxiety related ADRs with time to event onset ≤ 1 day.

\(^{138}\) In PSUR #2, 104,405 cases originating from PM sources reported the relevant PTs, and 56,230 cases included 69,338 events indicative of vaccination stress/anxiety related ADRs with time to event onset ≤ 1 day.
MC cases (13,225), NMC cases (26,575).

- Country of incidence (≥2%): Germany (13,472), Philippines (2881), Australia (2621), UK (2310), France (2286), Italy (1507), Netherlands (1483), Denmark (1229), Sweden (1047), Austria (1038), Romania (940), Poland (873); the remaining 8113 cases were distributed among 58 countries.

- Subjects' gender: female (28,653), male (10,626) and unknown (521).

- Subjects' age (n = 38,473), range: 10 weeks\textsuperscript{139} – 100 years, mean: 39.8 years, median: 39.0 years.

- Medical history (n = 21,834): the most frequently (≥2%) reported relevant medical conditions included Asthma (1033), Hypertension (999), Seasonal allergy (917), Drug hypersensitivity (793), Hypersensitivity (633), Food allergy (548).


- Co-suspects (n = 151 cases): the most frequently (≥ 8 occurrences) reported co-suspect vaccines/medications included influenza vaccine (41) and mestranol/norethisterone (8).

- Number of relevant events: 50,360.

- Relevant event seriousness:\textsuperscript{42} serious (12,116), non-serious (38,264).

- Most frequently reported relevant PTs (≥2%): Dizziness (16,611), Dyspnoea (7875), Paraesthesia (6846), Tachycardia (4757), Palpitations (4754), Blood pressure increased (2539), Hyperhidrosis (2339), Syncope (2294) and Loss of consciousness (1037).

- Time to event onset:
  - ≤24 hours: 31,865 events (48 of which had a fatal outcome);
  - 1 day: 18,534 events (33 of which had a fatal outcome).

- Duration of event (n = 12,385 of 18,547 relevant events with outcome of resolved/resolved with sequelae), range: < 24 hours to 447 days, median: 1 day.
  - ≤24 hours: 4908 events;
  - 1 day: 2265 events;
  - 2-7 days: 3328 events;
  - 8-14 days: 548 events;
  - 15-30 days: 443 events;
  - 31-181 days: 722 events;
  - 182-240 days: 103 events;

\textsuperscript{139} This infant subject received the vaccination for adult (cross-referenced to Section 9.2 Medication errors).
- 241-447 days: 68 events.

- Relevant event outcome: 78 fatal (81), resolved/resolving (26,704), resolved with sequelae (1170), not resolved (16,695), unknown (5789).

- In 73 cases (reporting 81 relevant events with fatal outcome), the reported causes of death (≥18 occurrences) were coded to the PTs Dyspnoea (39) and Loss of consciousness (18). Most (49 of 81 cases) of the fatal cases involved elderly subjects. When the medical history was provided (41 cases), the most frequently (≥5 occurrences) relevant medical conditions included hypertension (19), diabetes mellitus (11), cardiac failure and chronic obstructive pulmonary disease (5 each).

**Analysis by age group**

- **CT Data:** Paediatric (1) and Adults (1).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- **PM Data:** Paediatric (3681), Adults (31,950), Elderly (2921) and Unknown (1248).
  - No significant difference was observed in the reporting proportion of frequently (≥2%) reported relevant events between the adult and elderly populations. A higher reporting proportion of relevant PT Syncope was observed in the paediatric population when compared to the adult or elderly population (15.2% in paediatric vs 4.6% in adult vs 6.1% in elderly subjects). This is consistent with expectations based on age-related event reports from other vaccines.  

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 3277 (0.8% of the cases reporting stress/anxiety ADRs).

  - Upon review, no significant difference in the occurrence of the most frequently reported AEs related to vaccination stress/anxiety and in relevant AEs with fatal outcome in the subjects with comorbidities compared to the population without underlying diseases was identified, apart from the event syncope that was reported with higher proportion (6.3%) in subjects with comorbidities with respect to subjects without comorbidities (0.6%). The subjects’ underlying conditions are likely to be contributory to the occurrence of syncope in these cases.

**Conclusion**

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No new significant safety information was identified based on a review of these cases.

16.3.4. Evaluation of Special Situations

New data identified during the reporting interval for use of BNT162b2 by special subject situations is described below.

16.3.4.1. Death

Search criteria - Death cases are identified based on the following criteria:

- If the case or event outcome is “Fatal”.
- If the date of death field has a value.
- If any of the history type values is “Death” or “Autopsy”.
- If the death field is set to “Yes”.
- If the case contains one of the following terms: High Level Group Term: Fatal outcomes; Preferred Terms: Assisted suicide, Completed suicide.

Clinical Trial Data

- Number of cases: 34141 (blinded therapy [4] and BNT162b2 [30]) (5.1 % of 668 cases, the total CT dataset) compared to 44 cases (6.1%) retrieved in the PSUR #2.
- Country of incidence: the US (29), Brazil (2), Argentina, South Africa and Turkey (1 each).
- Subjects' gender: female (10) and male (24).
- Subjects' age in years (n = 34), range: 19.0 – 87.0 years, mean: 59.8 years, median: 63.5 years.
- Medical history (n = 28): the most frequently (>3 occurrences) reported medical conditions included Hypertension (16), Depression (12), Anxiety (7), Type 2 diabetes mellitus (6), Seasonal allergy (5), and Osteoarthritis (4).
- COVID-19 Medical history: None.
- Causes of death most frequently reported (>2 occurrences): Death (6), Disease progression (5), and Completed suicide (4).
- Autopsy results: None
- Events with a fatal outcome (n = 48): The most frequently reported PTs (>2 occurrences): Death (6) and Completed suicide (4). None of the fatal events are considered related to blinded therapy/BNT162b2.

141 During the current reporting interval, there were 3 additional cases reporting subjects’ death that were excluded from further analysis in this subsection as: death was mentioned as an incidental information only with none of the reported events presenting a fatal outcome (2) and a case which involved transplacental exposure is reviewed in Section 16.3.5.3 Use in Pregnant/Lactating Women.
• Co-suspects (n= 1 case): Alprazolam, bupropion, cyclobenzaprine, trazadone, venlafaxine (1 each).

• Time to fatal event onset (n = 40),\(^{142}\) range: 6 – 348 days, median: 119 days.
  - 2-7 days: 1 event;
  - 31-181 days: 30 events;
  - 182-240 days: 5 events;
  - 241-365 days: 4 events

Post-Authorisation Data

• Number of cases: 3163\(^{143}\) (0.6% of 507,683 cases, the total PM dataset) compared to 5215 (0.8%) analysed in the PSUR #2.

• MC cases (2061), NMC cases (1102).

• Country of incidence (≥107 occurrences): Germany (655), France (304), Japan (252), Philippines (205), Austria (194), the UK (164), Malaysia (151), the US (138), Australia (122), and Italy (107).

• Subjects’ gender: female (1304), male (1722), unknown (137).

• Subjects’ age in years (n = 2901), range: 5.0 – 107.0 years, mean: 68.0 years, median: 73.0 years.

• Medical history (n = 1631)\(^{144}\): The most frequently reported (>70 occurrences) medical conditions included cardiac and vascular disorders [e.g., Hypertension (588), Atrial fibrillation (171), Cardiac failure (113), Dyslipidaemia (80), and Myocardial ischaemia (72)]. Other most frequently reported (>70 occurrences) medical conditions included Diabetes mellitus (169), Type 2 diabetes mellitus (117), Obesity (102), Chronic obstructive pulmonary disease (95), Dementia (83), and Chronic kidney disease (72).

• COVID-19 Medical history (n = 98): COVID-19 (86), Suspected COVID-19 (9), COVID-19 pneumonia (8), Coronavirus infection, Post-acute COVID-19 syndrome, and SARS-CoV-2 antibody test positive (1 each).

• Causes of death most frequently reported (>100 occurrences): Death (739), COVID-19 (301), Cardiac arrest (215), Dyspnoea (185), Myocardial infarction (154), Vaccination

\(^{142}\) This number does not include 6 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\(^{143}\) During the current reporting interval, there were 159 additional cases reporting subjects' death that were excluded from further analysis in this subsection as death was mentioned as incidental information only with none of the reported events having a fatal outcome (77) and cases which reported foetal death/spontaneous abortion-involved transplacental exposure are reviewed in Section 16.3.5.3 Use in Pregnant/Lactating Women (82).

\(^{144}\) This list excluded the medical history terms indicative of COVID-19. Of note, more than 1 medical history was reported in some cases.
failure (144), Drug ineffective (131), COVID-19 pneumonia (129), Sudden death (110), Pulmonary embolism (105), Cardio-respiratory arrest (102), and Cardiac failure (101).

- Autopsy results were provided in 165 cases and the most commonly reported (≥7 occurrences) were: Pulmonary embolism (22), Myocarditis (18), Pulmonary oedema (12), Arteriosclerosis coronary artery, Myocardial infarction, Myocardial ischaemia (10 each), Acute myocardial infarction, Arteriosclerosis (9 each), Arrhythmia, Death (8 each), Cardiac failure (7).

- Co-suspect vaccines/medications (n = 144): the most frequently reported (>3 occurrences) were COVID-19 vaccine (25), influenza vaccine (16), COVID-19 vaccine MRNA (MRNA 1273) (15), COVID-19 vaccine NRV VAD (CHADOX1 NCOV-19) (14), influenza vaccine INACT SPLIT 4V (8), influenza vaccine INACT SAG 4V (6), casirivimab/imdevimab (5), apixaban, furosemide, and lenalidomide (4 each).

- Cases with confounders and risk factors: 1726 fatal cases included one or more contributing factors, which precluded a meaningful causality assessment: co-suspect (144 cases), concomitant drugs (638 cases) and/or underlying medical history/risk factors (1652 cases).

- Events with a fatal outcome (n = 8335): The most frequently reported (>100 occurrences) fatal events were coded to the PTs: Death (652), COVID-19 (340), Immunisation (240), Cardiac arrest (222), Vaccination failure (218), Dyspnoea (217), Drug ineffective (209), Off label use (193), Myocardial infarction (155), Interchange of vaccine products (144), Sudden death (140), COVID-19 pneumonia (137), Pyrexia (119), Pulmonary embolism (116), and Cardiac failure (102).

- Time to fatal event onset (n = 5580),\textsuperscript{145} range: <24 hours to 365 days, median: 8 days.
  - Same day: 1030 events;
  - 1 day: 592 events;
  - 2-7 days: 1058 events;
  - 8-14 days: 608 events;
  - 15-30 days: 621 events;
  - 31-181 days: 1463 events;
  - 182-240 days: 117 events
  - 241-365 days: 91 events

Analysis by age group

- CT: Adults (18-64) (17) and Elderly (65 years and older) (17).
  - A meaningful comparison between age groups is not possible due to the low number of cases with a fatal outcome.

\textsuperscript{145} This number does not include 4 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
PM: Paediatric (17 years and under) (82), Adults (18-64 years) (932), Elderly (65 years and older) (1946), and Unknown (203).

- There is a significant difference observed in the reporting proportion for the majority of the frequently reported fatal events (>100 occurrences) in the elderly population when compared to the adult population due to a higher proportion of fatal cases reported in subjects over 64 years of age (61.5% vs 29.5%, respectively). There is no meaningful comparison between elderly vs paediatric population possible due to the low number of paediatric fatal cases reported (2.6% vs 61.5%, respectively).

Most of the cases reporting a fatal outcome (42.1%) were in subjects over 75 years of age. The elderly population is generally considered a priority group targeted for vaccination by many regions and countries, including Europe and the US (with various lower age cut-offs across countries), because of their higher risk of severe disease and mortality if infected with SARS-CoV-2.\textsuperscript{146,147,148}

**Analysis by presence of comorbidities**

- Number of subjects with comorbidities: 1094 (0.2% of 508,351, the total dataset) when compared to 2090 (0.3% of 658,249 cases) in the PSUR #2.

- Upon review, there were no significant differences observed in the patterns of the most frequently reported fatal events (>100 occurrences) between the group with comorbidities and the one without comorbidities.

**Analysis by dose**

- Number of vaccine doses administered at the time of the subjects’ death:
  - First dose (378 cases)
  - Second dose (934 cases). Of the 934 cases, 163 cases (17.5%) reported a latency of same day to 3 days after vaccination. There were 2477 fatal events. The most frequently reported (>100 occurrences) fatal events were coded to the PTs COVID-19 (178), Death (154), Drug ineffective (113), Vaccination failure (111).
  - Third dose (1084 cases). Majority of these cases (>50 occurrences) originated from Germany (240), Japan (151), France (139), the UK (78), and Austria (56). There were 3267 fatal events. The most frequently reported (>100 occurrences) fatal events were

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\textsuperscript{146} Overview of the implementation of COVID-19 vaccination strategies and vaccine deployment plans in the EU/EEA. ECDC, February 2021.

\textsuperscript{147} https://www.cdc.gov/vaccines/hcp/accip-recs/vacc-specific/COVID-19/evidence-table-phase-1b-1c.html.

coded to the PTs Death (206) Immunisation\textsuperscript{43} (188), Off label use (117), COVID-19 (112), Interchange of vaccine products (107), and Vaccination failure (101).

- Fourth dose (71 cases). Majority of these cases (>10 occurrences) originated from Germany (23), France, and the UK (11 each). There were 254 fatal events. The most frequently reported (>20 occurrences) fatal events were coded to the PTs Off label use (42) Immunisation\textsuperscript{43} (39), and Death (22).

- Fifth dose (1 case). This is a spontaneous case reported by a consumer. In this case, a 66-year-old male subject received BNT162b2, as dose 5 (booster), for COVID-19 immunisation (Off label use). Relevant medical history included interchange of vaccine products (first 2 doses with Coronavac; third and fourth doses with BNT162b2) and hospitalisation for the drop in oxygen saturation. The subject’s condition worsened after receiving the fifth dose and he experienced immunisation reaction such as low oxygen saturation, lung oedema, abnormal lung function and shortness of breath, and he died 3 days later. Oxygen deficiency and failure of the lungs to function were cited as the cause of death. It was unknown if an autopsy was performed.

- In the remaining cases (695), dose number was not specified at the time of the subject’s death.

**Literature**

Review of the literature did not identify any significant new information regarding the use of BNT162b2 and death.

**Conclusion**

No new risks were identified following review of fatal cases.

16.3.4.1.1. Death Review by Age Group

This is a high-level overview of the 3197 cases in the interval reporting period (see Section 16.3.4.1 for further details). According to the corePSUR\textsuperscript{125} summary tabulation of fatal reports by Age groups and SOCs is provided in Appendix 6.C.\textsuperscript{149}.

**Interval Reporting Period**

- CT (34 cases): Adults (18-64) (17) and Elderly (65 years and older) (17)

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\textsuperscript{149} Please note that the numbers of AEs reported in the appendix may not match with the numbers of AEs reported in Table 63 and Table 64, since in the appendix all the events included in the fatal cases are reported, while in the above mentioned tables, only AEs with fatal outcome are reported.
The top 6 MedDRA SOCsWith the most frequently reported (>3 occurrences) events with the total number of fatal events in the interval period by age group is presented in the table below.

**Table 63. Clinical Trial Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group during the Reporting Interval**

<table>
<thead>
<tr>
<th>SOC</th>
<th>Total number of events</th>
<th>18-24 years</th>
<th>25-49 years</th>
<th>50-59 years</th>
<th>60-69 years</th>
<th>70+ years</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>10</td>
<td>0</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>8</td>
<td>0</td>
<td>5</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>5</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Neoplasms benign, malignant and unspecified (incl cysts and polyps)</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>4</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Psychiatric disorders</td>
<td>4</td>
<td>1</td>
<td>3</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Of note, multiple AEs may be reported in a single case.

- PM (3163 cases): Paediatric (17 years and under) (82), Adults (18-64 years) (932), Elderly (65 years and older) (1946), and Unknown (203).

The top 5 MedDRA SOCs with the most frequently reported (>500 occurrences) events with a fatal outcome cumulative by age group in the post-authorisation data are presented in the table below.

**Table 64. Post-Authorisation Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group during the Reporting Interval**

<table>
<thead>
<tr>
<th>SOC</th>
<th>Total number of events</th>
<th>≤ 17 years</th>
<th>18-24 years</th>
<th>25-49 years</th>
<th>50-59 years</th>
<th>60-69 years</th>
<th>70+ years</th>
<th>Unk</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>2059</td>
<td>52</td>
<td>33</td>
<td>242</td>
<td>172</td>
<td>278</td>
<td>1161</td>
<td>121</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>1197</td>
<td>42</td>
<td>30</td>
<td>168</td>
<td>133</td>
<td>192</td>
<td>588</td>
<td>44</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>950</td>
<td>30</td>
<td>36</td>
<td>120</td>
<td>96</td>
<td>151</td>
<td>483</td>
<td>34</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>824</td>
<td>16</td>
<td>11</td>
<td>42</td>
<td>42</td>
<td>106</td>
<td>580</td>
<td>27</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td>779</td>
<td>29</td>
<td>16</td>
<td>97</td>
<td>75</td>
<td>129</td>
<td>413</td>
<td>20</td>
</tr>
</tbody>
</table>

Of note, multiple AEs may be reported in a single case.
Cumulative Reporting Period

This is a high-level overview of the 13,659 relevant cumulative cases with a fatal outcome. According to the corePSUR19 guidance, a summary tabulation of fatal reports by age groups and SOC's is provided in Appendix 6C.2.

Clinical Trial Data

- Number of cases: 150 (6.2% of 2426 cases, the total CT dataset; 143 cases involved blinded therapy [67]/BNT162b2 [76]). In the remaining 7 cases subjects received placebo.
- Causes of death most frequently reported (>7 occurrences): Disease progression (29), Cardiac arrest (15), Death (14), Completed suicide (10), Cardio-respiratory arrest, Myocardial infarction (8 each).
- Autopsy results were provided in 10 cases and the most commonly (>2 occurrences) reported were: Arteriosclerosis, Hypertensive heart disease, Pulmonary embolism (2 each).
- Events with a fatal outcome (n = 198): The most frequently reported PTs (>5 occurrences) were: Death (14), Completed suicide (10), Cardio-respiratory arrest (9), Cardiac arrest, Myocardial infarction (8 each), Pulmonary embolism (6), Acute respiratory failure, COVID-19, COVID-19 pneumonia, and Septic shock (5 each). None of these events are considered related to blinded therapy/BNT162b2.

Post-Authorisation Data

- Number of cases: 13,509 (0.9 % of 1,484,945 cases, the total cumulative PM dataset).
- MC cases (9582), NMC cases (3927).
- Causes of death most frequently reported (>500 occurrences): Death (3145), COVID-19 (1296), Cardiac arrest (892), Dyspnoea (725), Sudden death (618), Myocardial infarction (610), Vaccination failure (574), Cardio-respiratory arrest (557), Pyrexia (541), Drug ineffective (517), and Pulmonary embolism (514).

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150 Please note that the numbers of AEs reported in the appendix may not match with the numbers of AEs reported in Table 65 and Table 66, since in the appendix all the events included in the fatal cases are reported, while in the above mentioned tables, only AEs with fatal outcome are reported.

151 There were 16 additional cases reporting subject deaths that were excluded from further analysis in this subsection because: death was mentioned as incidental information only with none of the reported events presenting a fatal outcome (12) and cases which involved transplacental exposure/baby cases (4) are reviewed in Section 16.3.5.3 Use in Pregnant/Lactating Women.

152 During the current reporting interval, there were 492 additional cases reporting subject deaths that were excluded from further analysis in this subsection because: death was mentioned as incidental information only with none of the reported events presenting a fatal outcome (227) and cases which reported foetal death/still birth/spontaneous abortion/involved transplacental or trans-mammary exposure are reviewed in Section 16.3.5.3 Use in Pregnant/Lactating Women (265).
Autopsy results were provided in 725 cases and the most commonly reported (>30 occurrences) were: Pulmonary embolism (82), Pulmonary oedema (61), Arteriosclerosis (54), Myocardial infarction (50), Arteriosclerosis coronary artery (48), Acute myocardial infarction (46), Myocarditis (39), Cardiac hypertrophy (33), and Cardiomegaly (31).

Events with a fatal outcome (n = 32,992): The most frequently reported (>500 occurrences) events were coded to the PTs: Death (3016), COVID-19 (1389), Cardiac arrest (911), Dyspnoea (813), Vaccination failure (737), Drug ineffective (716), Sudden death (704), Pyrexia (622), Myocardial infarction (619), Cardio-respiratory arrest (575), and Pulmonary embolism (536).

Analysis by age group:

CT: Adults (79), and Elderly (71).

The top 6 MedDRA SOC's with the most frequently reported (≥15 occurrences) events with a fatal outcome cumulative by age group is presented in the table below.

### Table 65. Clinical Trial Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group - Cumulative Reporting Interval

<table>
<thead>
<tr>
<th>SOC</th>
<th>Total number of events</th>
<th>18-24 years</th>
<th>25-49 years</th>
<th>50-59 years</th>
<th>60-69 years</th>
<th>70+ years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>35</td>
<td>0</td>
<td>5</td>
<td>9</td>
<td>11</td>
<td>10</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>34</td>
<td>0</td>
<td>3</td>
<td>8</td>
<td>10</td>
<td>13</td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td>25</td>
<td>0</td>
<td>5</td>
<td>5</td>
<td>8</td>
<td>7</td>
</tr>
<tr>
<td>Neoplasms benign, malignant and unspecified (incl cysts and polyps)</td>
<td>24</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>7</td>
<td>15</td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>18</td>
<td>1</td>
<td>9</td>
<td>2</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td>17</td>
<td>0</td>
<td>2</td>
<td>3</td>
<td>9</td>
<td>3</td>
</tr>
</tbody>
</table>

Of note, multiple AEs may be reported in a single case.

- A meaningful comparison between age groups is not possible due to the low number of cases with a fatal outcome.

PM: Paediatric (17 years and under) (161), Adults (18-64 years) (2708), Elderly (65 years and older) (9568) and Unknown (1072).
The top 5 MedDRA SOCs with the most frequently reported (>3000 occurrences) events with a fatal outcome cumulative by age group in the PM data are presented in the table below.

Table 66. Post-Authorisation Data: Total Number of AEs with a Fatal Outcome by SOCs and by Age Group - Cumulative Reporting Interval

<table>
<thead>
<tr>
<th>SOC</th>
<th>Total number of events</th>
<th>≤17 years</th>
<th>18-24 years</th>
<th>25-49 years</th>
<th>50-59 years</th>
<th>60-69 years</th>
<th>70+ years</th>
<th>Unk</th>
</tr>
</thead>
<tbody>
<tr>
<td>General disorders and administration site conditions</td>
<td>8735</td>
<td>115</td>
<td>65</td>
<td>612</td>
<td>534</td>
<td>956</td>
<td>5720</td>
<td>733</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td>5014</td>
<td>78</td>
<td>59</td>
<td>539</td>
<td>433</td>
<td>700</td>
<td>3075</td>
<td>130</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>3818</td>
<td>56</td>
<td>63</td>
<td>336</td>
<td>311</td>
<td>516</td>
<td>2433</td>
<td>103</td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td>3535</td>
<td>62</td>
<td>32</td>
<td>282</td>
<td>251</td>
<td>444</td>
<td>2395</td>
<td>69</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>3473</td>
<td>28</td>
<td>21</td>
<td>108</td>
<td>118</td>
<td>355</td>
<td>2596</td>
<td>247</td>
</tr>
</tbody>
</table>

Of note, multiple AEs may be reported in a single case.

- There is a significant difference observed in the reporting proportion of most frequently reported fatal events (listed above) in the elderly population when compared to the adult population (70.8% vs 20.0%, respectively). A meaningful comparison between the elderly vs paediatric population is not possible due to the low number of paediatric fatal cases reported (1.2% vs 70.8%, respectively).

- Most of the cases reporting a fatal outcome (53.7%) were in subjects over 75 years of age. The elderly population were generally considered a priority group targeted for vaccination by many regions and countries, including Europe and the US (with various lower age cut-offs across countries), because of their higher risk of severe disease and mortality if infected with SARS-CoV-2.\textsuperscript{146,147,148}

O/E Analysis

O/E analysis was performed for events with a fatal outcome (see Appendix 6B Observed versus Expected Analyses for Adverse Events of Special Interest).

Conclusion

No safety signals have emerged based on the review of these cases, or from the O/E analysis. Safety surveillance will continue.
16.3.4.2. Overdose

Search criteria - HLT Overdoses NEC OR PT Accidental overdose.

Of the 1605 cases, 9 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

- Overdose was not implied (i.e., inquiry, past expiry, underdose\textsuperscript{153}) in 7 cases
- the reported PTs of overdose referred to vitamin K/digoxin or occurred 17 days following vaccination in 2 cases.

Clinical Trial Data

There were no\textsuperscript{154} serious clinical trial cases of overdose of the vaccine reported during the current interval period, similar to no cases in the PSUR #2.

Post-Authorisation Data

- Number of cases: 1595\textsuperscript{155} (0.3% of 507,683 cases, the total PM dataset), compared to 1985 cases (0.3%) retrieved in the PSUR #2.
- MC cases (1237), NMC cases (358).
- Country of incidence (≥2%): US (769), Germany (168), Taiwan (100), Canada (92), France, Italy (82 each), Poland (48), UK (44), Portugal (32); the remaining 178 cases were distributed among 25 countries.
- Subjects’ gender: female (572), male (444), and unknown (579).
- Subjects’ age in years (n = 1029), range: 1 – 101 years, mean: 27.6 years, median: 17 years.
- Medical history (n = 185): the most frequently (≥4 occurrences) reported medical conditions included: Hypertension (26), Asthma, COVID-19 (20 each), Diabetes mellitus, Hypersensitivity (10 each), Obesity (9), Food allergy (8), Anxiety, Attention deficit hyperactivity disorder, Drug hypersensitivity (7 each), Depression, Interchange of vaccine products (6 each), Autoimmune thyroiditis, Type 2 diabetes mellitus (4 each).
- Co-suspect vaccines/medications: COVID-19 vaccine MRNA (MRNA 1273) (4), Influenza vaccine (2), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19), Diphtheria vaccine toxoid/pertussis vaccine acellular/tetanus vaccine toxoid,

\textsuperscript{153} A 13 year old subject received the “orange cap” booster (for ages 5 to <12) rather than the ≥ 12 years dose.

\textsuperscript{154} Two cases involved paracetamol and cocaine.

\textsuperscript{155} Among these cases, 239 involved the Tris/Sucrose formulation (cross-referenced to Section 6.3.1.1.2.2 Tris/Sucrose Presentation).
enzalutamide, HPV vaccine VLP RL1 4V (yeast), Hydrocortisone, JNJ 78436735, Mirogabalin besilate, Pneumococcal vaccine conj 20V (CRM197) (1 each).

- Number of relevant events: 1595.
- Relevant event seriousness: serious (83), non-serious (1512).
- Relevant PTs: Overdose (1510), Accidental overdose (81), and Intentional overdose (4).
- Relevant event outcome: resolved/resolving (68), not resolved (12), fatal (3), resolved with sequelae (2), unknown (1510).
- Most frequently co-reported PTs (≥2%): Product preparation error (663), Product preparation issue (237), Product administered to patient of inappropriate age (121), Poor quality product administered (107), Expired product administered (96), Pyrexia (88), Headache (79), Pain in extremity (62), Vaccination site pain (59), Product temperature excursion issue (55), Chest pain (45), Fatigue (37), Product administration error (36), Asthenia, Off label use (35 each), Dizziness (34), Chest discomfort, Immunisation, Incorrect dose administered (33 each).

Analysis by age group

- Paediatric (630), Adults (420), Elderly (89) and Unknown (456).
  - Upon review, no significant differences in the reporting proportion of the most frequently co-reported AEs were noted between the different age groups.

Analysis by presence of comorbidities

- Number of subjects with comorbidities: 63 (4.0% of the total cases reporting overdose).

- Upon review, no significant differences in the occurrence of the most frequently co-reported AEs in the subjects with comorbidities compared to the population without underlying diseases was identified.

Literature

Review of the literature did not identify any significant new information regarding overdoses of BNT162b2.

Conclusion

The most frequently reported reasons (≥2%) for overdose were:

- administration of incorrect dose of diluted vaccine, different from the recommended 0.3 ml for the subjects aged ≥ 12 years and 0.2 ml for the paediatric subjects aged 5 through 11 years (411; 25.8% of the total cases reporting overdose);

- administration of undiluted vaccine (582; 36.5% of the total cases reporting overdose);
- dilution with a volume of sodium chloride different from the recommended 1.8 ml for the subjects aged ≥ 12 years and 1.3 ml for the paediatric subjects aged 5 through 11 years (170; 10.7% of the total cases reporting overdose);

- administration of more than 1 dose of vaccine (39; 2.4% of the total cases reporting overdose);

- incorrect vaccine formulation administered to paediatric subjects aged 5 through 11 years instead of the recommended 10 mcg dosage (71; 4.4% of the total cases reporting overdose).

In most cases, the incorrect preparation and/or administration of vaccine occurred by mistake. In 218 cases, the reason for overdose was not reported or unclear, 2 of which reported the PT Intentional overdose. In the remaining 2 cases reporting intentional overdose, an administration of 30 mcg in children (aged 9 and 10 years old) was reported. No new significant safety information was identified based on the review of these cases. The majority of the most frequently co-reported AEs other than overdose and medication error PTs were events consistent with the known reactogenicity of the vaccine and listed in Section 4.8 Undesirable effects of the CDS.

16.3.4.3. Abuse, Misuse, and Drug Dependency

Abuse Search Criteria: PTs Alcohol use disorder; Dependence; Disturbance in social behaviour; Dopamine dysregulation syndrome; Drug abuse; Drug abuser; Drug dependence; Drug dependence, antepartum; Drug dependence, postpartum; Drug detoxification; Drug diversion; Drug level above therapeutic; Drug level increased; Drug rehabilitation; Drug screen; Drug screen positive; Drug tolerance; Drug tolerance decreased; Drug tolerance increased; Drug use disorder; Drug use disorder, antepartum; Drug use disorder, postpartum; Drug withdrawal convulsions; Drug withdrawal headache; Drug withdrawal maintenance therapy; Drug withdrawal syndrome; Drug withdrawal syndrome neonatal; Maternal use of illicit drugs; Needle track marks; Neonatal complications of substance abuse; Pharmaceutical nomadism; Substance abuse; Substance abuser; Substance dependence; Substance use; Substance use disorder; Toxicity to various agents; Withdrawal syndrome.

Misuse Search Criteria: Intentional product misuses; Intentional device use issue; Intentional dose omission; Intentional medical device removal by patient; Intentional product use issue; Intentional removal of drug delivery system by patient; Intentional overdose; Performance enhancing product use; Prescription drug used without a prescription; Treatment noncompliance.

Of the 55 cases, 44 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

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156 This PT has been added to the Abuse Derived MedDRA Condition upon MedDRA upversioning to v.25.0.
Eleven (11) cases reported the PT Toxicity to various agents. These cases described adverse events experienced by the subjects but did not involve abuse, intentional, excessive or non-therapeutic use of BNT162b2;

Six (6) cases involved the abuse of illicit substances, including cannabis (2), delorazepam, “pain killers”, and “used to be a drug addict before he had the vaccine”, and multiple drugs [morphine, cannabis, methadone, amphetamines] (1 each);

Four (4) cases reported the PT Intentional dose omission; one (1) case reported Intentional dose omission and Product dose omission issue; and one (1) case reported Product dose omission issue and Intentional product use issue. Each case involved drugs other than BNT162b2 (methotrexate (1), adalimumab (1), lenalidomide (1), and in 3 cases it was unclear as to which drug dose was omitted);

Four (4) cases reported the PT Needle track marks following administration of BNT162b2; however, there was no information regarding intravenous drug abuse/substance use in these cases;

Four (4) cases reported the PT Withdrawal syndrome; and two (2) cases reported the PT Disturbance in social behaviour. Each case described adverse events after receiving BNT162b2 by the subjects but did not involve abuse, intentional, excessive or non-therapeutic use of BNT162b2;

Three (3) cases reported the PTs (Drug level increased [2 cases] and Drug level above therapeutic [1 case]) regarding co-suspect drugs clozapine (2) and citalopram hydrochloride (1);

Two (2) cases involved Intentional product use issue; and one (1) case reported Intentional product misuse. Each case involved drugs other than BNT162b2 (apixaban [2], and adalimumab [1]);

Two (2) cases reported Treatment noncompliance involving drugs other than BNT162b2 (brodalumab [1], and an unspecified drug used to treat an allergic reaction);

One (1) case reported Drug dependence after receiving a single dose but did not report intentional, excessive or non-therapeutic use of BNT162b2;

One (1) case reported the PT Drug withdrawal, which was associated with opiate withdrawal;

One (1) case reported the PT Drug tolerance described as “built up tolerance to Humira medication”.
Clinical Trial Data

There were no serious clinical trial cases of abuse or misuse of the vaccine reported during the reporting period; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 11 (0.002% of 507,683 cases, the total PM dataset), compared to 45 cases (0.01%) retrieved in the PSUR #2.
- MC cases (8), NMC cases (3).
- Country of incidence: Canada (3), Australia (2), Finland, Germany, Ireland, Japan, New Zealand, US (1 each).
- Subjects' gender: female (8), male (2), and unknown (1).
- Subjects' age in years (n = 9), range: 4 - 73, mean: 40.4, median: 49.0.
- Medical history (n = 7): Cardiac disorder, Drug hypersensitivity, Hypertension, Intracranial aneurysm, Malnutrition, Migraine with aura, Rheumatoid arthritis (1 each).
- COVID-19 Medical history: None.
- Co-suspect vaccines/medications (n = 8): abatacept, COVID-19 vaccine MRNA (MRNA 1273), dabrafenib, gabapentin, natalizumab, nivolumab, perindopril, trametinib (1 each).
- Number of events: 92 (of which 11 were events of interest).
- Relevant event seriousness: serious (5), non-serious (6).
- Relevant PTs: Intentional product misuse, Intentional product use issue (4 each), Intentional underdose (3).
- Co-reported AEs (≥2): Fatigue, Nausea, Off label use (3 each), Chest pain, Headache, Malaise, Poor quality product administered, Product storage error (2).
- Time to event onset (n = 3), range: < 24 hours, median: 0 days.
  - <24 hours: 3 cases.
- Relevant event outcome: fatal (1), not resolved (3), unknown (7).

In the case involving the fatal outcome, a female subject (age unknown) received three vaccines COVID-19 (BNT162b2), pneumonia vaccine (unspecified) and the flu vaccine at one time. The patient experienced a myocardial infarction and died. Onset date of myocardial infarction was not reported.

Analysis by age group

- PM: Paediatric (2), Adults (5), Elderly (2), and Unknown (2).
  - There was no meaningful difference between different age groups.
Analysis by dose

- PM: Number of vaccine doses administered at the time of the event onset: dose 1 in 1 case, dose 2 in 1 case, dose 3 in 1 case, and number of doses was not specified in 8 cases.
  - There are no differences between the AEs that occurred after the first, the second and the booster dose.

Literature

Review of the literature did not identify any significant new information regarding the use of BNT162b2 and abuse, dependence or misuse.

Conclusion

Overall, there were 11 cases representing 0.002% of the overall post-marketing dataset, that reported events indicative of misuse. These cases involved either improper storage, improper dilution of vaccine, administration of vaccine to unapproved age groups or administration of vaccine at a dose lower than the recommended dose. In general, the most frequently co-reported events observed in these cases were consistent with those observed in the overall population. No safety signals have emerged that would be considered specific to this population.

16.3.4.4. Occupational Exposure

Search criteria - PTs Exposure to contaminated device; Occupational exposure to product; Occupational exposure to radiation; Occupational exposure to toxic agent.

Clinical Trial Data

- There were no serious clinical trial cases indicative of occupational exposure during the reporting period; no cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 20 (0.004% of 507,683 cases, the total PM dataset), compared to 41 cases (0.01%) retrieved in the PSUR #2.
- MC cases (18), NMC cases (2).
- Country of incidence: US (7), Germany (6), Australia (5), Brazil, Italy (1 each).
- Subjects’ gender: female (14), male (4) and unknown (2).
- Subjects’ age in years (n = 7), range: 2 – 58 years, mean: 37.1 years, median: 43.0 years.
- Medical history (n = 3): reported medical conditions included Abnormal behaviour, Anxiety, Asthma, Autism spectrum disorder, Cerebral palsy, Chronic active Epstein-Barr virus infection, COVID-19, Cyclic vomiting syndrome, Depression, Drug hypersensitivity, Gastroesophageal reflux disease, Gene mutation, Gluten sensitivity,
Hypothyroidism, Iodine deficiency, Irritable bowel syndrome, Neurodermatitis, Obesity, Pain, Rubber sensitivity, Vitamin D deficiency (1 each).

- Co-suspect vaccines/medications (n = 0).
- Number of events: 33 (of which 20 were events of interest).
- Relevant event seriousness: serious (1), non-serious (19).
- Relevant PTs: Occupational exposure to product (20).
- Co-reported AEs: Abdominal pain, Amylase decreased, Decreased appetite, Exposure via skin contact, Fatigue, Headache, Nausea, Ocular hyperaemia, Off label use, Pain, Product use issue, Underdose, Weight decreased (1 each).
- Time to event onset (n = 5), range: 0 and 224 days.
  - <24 hours: 4 events (none of which had a fatal outcome);
  - 1 day: 0 events;
  - 2-7 days: 0 events;
  - 8-14 days: 0 events;
  - 15-30 days: 0 events;
  - 31-180 days: 0 events.
- Relevant event outcome: resolved/resolving (2), resolved with sequelae (1), unknown (17).

Analysis by age group

- PM: Paediatric (2), Adults (5), Elderly (0) and Unknown (13).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

Literature

Review of the literature did not identify any significant new information regarding the use of BNT162b2 and occupational exposure.

Conclusion

Overall, there were 20 cases representing 0.004% % of the overall post-marketing dataset, that reported events indicative of occupational exposure. Review of the cases did not identify any significant new information regarding the use of BNT162b2 and occupational exposure. No safety signals have emerged that would be considered specific to this population.

16.3.4.5. Lack of Therapeutic Efficacy

Company conventions for MedDRA coding of cases indicative of lack of efficacy:
The coding conventions for COVID-19 vaccine cases indicative of lack of efficacy was revised on 27 Sep 2021, as shown below:

- PT “Vaccination failure” is coded when ALL of the following criteria are met:
  - The subject received the appropriate series of two doses based on the CDS.
  - At least 7 days have elapsed since administration of the second dose.
  - The subject experiences COVID-19 infection (confirmed by laboratory tests or reported by HCP).

- PT “Drug ineffective” is coded when any of the following applies:
  - The COVID-19 infection is not reported by HCP or not confirmed through laboratory tests (irrespective of the vaccination schedule). This includes scenarios where LOE is stated or implied by consumers, e.g., “the vaccine did not work”, “I got COVID-19”.
  - It is unknown:
    - Whether the subject has received the two doses within the correct intervals based on the labeling instructions;
    - How many days have passed since the first dose (including unspecified number of days like “a few days”, “some days”, etc.);
    - If 7 days have passed since the second dose of vaccine.
  - The subject experiences COVID-19 infection 14 days after receiving the first dose up to and through 6 days after receipt of the second dose.

- Note: A case is considered a potential LOE case after the immune system has had sufficient time (14 days) to respond to the vaccine, even if the vaccination course is not complete.

This is the summary of the coding conventions based on the timing of vaccination:

<table>
<thead>
<tr>
<th>From 1st dose to day 13 post 1st dose</th>
<th>From day 14 post 1st dose to day 6 post 2nd dose</th>
<th>From day 7 post 2nd dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Code only the events describing the COVID-19 infection</td>
<td>Code “Drug ineffective”</td>
<td>Code “Vaccination failure”</td>
</tr>
<tr>
<td>Scenario not considered LOE</td>
<td>Scenario considered LOE as “Drug ineffective”</td>
<td>Scenario considered LOE as “Vaccination failure”</td>
</tr>
</tbody>
</table>
Lack of efficacy cases\textsuperscript{157}

Search criteria - PTs Drug ineffective; Vaccination failure.

- Of the 51,107 cases, 79 cases were determined to be non-contributory and were not included in the discussion for the following reasons:
  - 18 cases are not considered true LOE cases because the subjects developed SARS-CoV-2 infection days 1-13 from the first dose.
  - 6 cases were invalidated in the safety database after the PSUR DLP.
  - 18 cases were not LOE reports (subjects did not develop SARS-CoV-2 infection).
  - In 37 cases, the LOE PT did not refer to BNT162b2 vaccine.

Clinical Trial Data

There were no lack of efficacy cases in the clinical trial dataset for this reporting period or for the reporting period of PSUR #2.

Post-Authorisation Data

- Number of cases: 51,028 (10.1% of 507,683 cases, the total PM dataset), compared to 21,457 cases (3.3%) in PSUR #2. The increase in the reporting proportion of LOE cases was multifactorial. A high number of cases were reported from Austria (31,629 cases in the current PSUR), as compared to the previous PSURs (9009 cases in PSUR #2 and 204 cases in PSUR #1) due to the active solicitation of LOE cases, including retrospective cases, by the Austrian Board of Health starting from August 2021. In addition, the epidemiology of the virus has changed since December 2021 in that the Omicron variant has become predominant in most regions. BNT162b2 efficacy against Omicron variants is less than against the previous dominant variants of concern.
- MC cases (39,368), NMC cases (11,660).
- Relevant lack of efficacy events\textsuperscript{158}: 51,028 (Vaccination failure [24,762] and Drug ineffective [26,266]).
- Country of incidence (≥2%): Austria (31,629), US (4734), UK (2316), Germany (1856), France (1478), Netherlands (1291); the remaining 7724 cases were distributed among 71 countries.
- Subjects’ gender: female (27,177), male (21,802) and unknown (2049).

\textsuperscript{157} LOE cases are assessed according to the definition provided in the EMA corePSUR19 guidance (EMA/362988/2021) and classified into confirmed vaccination failure, suspected vaccination failure, and not a vaccination failure.

\textsuperscript{158} LOE PTs recorded in the 51,028 cases were Vaccination failure (24,404) and Drug ineffective (26,624). Upon review after DLP, some cases were re-assessed: in 423 cases the PT Drug ineffective was reassessed to Vaccination failure, and in 65 cases the PT Vaccination failure was reassessed to Drug ineffective.
COVID-19 mRNA vaccine (nucleoside modified)
Periodic Safety Update Report (PSUR) 3
19 December 2021 through 18 June 2022

- Subjects' age in years (n = 48,297), range: 1.5 – 107.0 years, mean: 47.3 years, median: 47.0 years.
- Relevant event seriousness: all serious.\textsuperscript{159}

\textbf{Confirmed vaccination failure (24,077 cases)}

Vaccination failure was reported in 24,077 cases, indicative of appropriately and fully vaccinated subjects (appropriate series of 2 doses at the appropriate interval), who developed clinical, and laboratory confirmed (e.g., COVID-19 PCR test, antigen test) COVID-19 infection, on or after day 7 post second dose. In 5029 of these 24,077 cases, a booster dose was also administered (including 4735 cases with administration of the third dose and 294 cases with administration of the fourth dose).

- Age groups: Child (40), Adolescent (1053), Adult (18,337), Elderly (4475) and Unknown (172).
- Time to event onset was known for 23,013 cases; in the remaining 1064 cases, it was implied that vaccination failure was reported on or after day 7 post second dose, however, detailed information was not provided.
  - Time to onset reported after the second dose.
    - \( \geq 91 \) days to \( \leq 180 \) days: 13,650 subjects
    - \( \geq 181 \) days to \( \leq 270 \) days: 799 subjects
    - \( \geq 271 \) days to \( \leq 360 \) days: 270 subjects
    - \( \geq 361 \) days to \( \leq 450 \) days: 70 subjects
    - \( \geq 451 \) days to \( \leq 501 \) days: 9 subjects

  - Time to onset reported after the third dose.
    - \( \geq 91 \) days to \( \leq 180 \) days: 671 subjects
    - \( \geq 181 \) days to \( \leq 270 \) days: 264 subjects
    - \( \geq 271 \) days to \( \leq 293 \) days: 7 subjects

  - Time to onset reported after the fourth dose.
    - \( \geq 91 \) days to \( \leq 180 \) days: 129 subjects
    - \( \geq 181 \) days to \( \leq 293 \) days: 9 subjects
    - 213 days: 1 subject

\textsuperscript{159} Includes 5 cases where LOE was captured as non-serious and upgraded to serious after the PSUR DLP.
- Reported COVID-19 infection related events (>5 occurrences)\textsuperscript{160}: COVID-19 (23,679), COVID-19 pneumonia (285), SARS-CoV-2 test positive, Suspected COVID-19\textsuperscript{161} (107 each), Vaccine breakthrough infection (21), Breakthrough COVID-19 (8), and Post-acute COVID-19 syndrome (6).

- Outcome of COVID-19 infection related events: resolved/resolving (2187), resolved with sequelae (29), not resolved (673), unknown (21,115), and fatal (221).

- Of the 24,077 subjects with confirmed vaccination failure, in 880 cases, the COVID-19 events were severe, resulting in:
  - Hospitalisation (non-fatal/non-life threatening): 623
  - Disability: 13
  - Life threatening: 40
  - Death: 204.

\textbf{Suspected vaccination failure (1402 cases)}

Lack of efficacy (PTs Drug ineffective or Vaccination failure) was reported in 1402 cases, wherein the subjects received 2 doses of vaccine at appropriate interval and reported to develop COVID-19 infection on or after day 7 post second dose, but laboratory confirmation of the infection (e.g., COVID-19 PCR test, antigen test) was not reported or clinical disease was unconfirmed (i.e., asymptomatic COVID-19). In 307 of these 1402 cases, a booster dose was also administered (including 300 cases with administration of the third dose and 7 cases with administration of the fourth dose).

- Age groups: Child (3), Adolescent (46), Adult (991), Elderly (298) and Unknown (64).

- Time to event onset was known for 1036 cases; in the remaining 366 cases, it was implied that vaccination failure was reported on or after day 7 post second dose, however, detailed information was not provided.

  - Time to onset reported after the second dose.
    - day 7 to \( \leq \) 90 days: 138 subjects
    - $\geq$ 91 days to \( \leq \) 180 days: 413 subjects
    - $\geq$ 181 days to \( \leq \) 270 days: 272 subjects
    - $\geq$ 271 days to \( \leq \) 360 days: 32 subjects
    - 438 days: 1 subject

  - Time to onset reported after the third dose.
    - day 1 to \( \leq \) 90 days: 115 subjects

\textsuperscript{160} Some cases reported more than 1 PT referring to a SARS-CoV-2 infection related event.

\textsuperscript{161} In these cases reporting Suspected COVID-19, upon review, the infection was assessed to be confirmed.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

- ≥ 91 days to ≤ 180 days: 51 subjects
- ≥ 181 days to ≤ 234 days: 11 subjects

- Time to onset reported after the fourth dose.
  - day 1, 4 days and 29 days: 3 subjects

- Reported COVID-19 infection related events (≥ 32 occurrences)\textsuperscript{160}: Suspected COVID-19 (610), COVID-19 (527), Asymptomatic COVID-19 (239), and COVID-19 pneumonia (32).

- Outcome of COVID-19 infection related events: resolved/resolving (664), resolved with sequelae (4), not resolved (62), unknown (662), and fatal (24).

\textit{Not a vaccination failure cases (25,549 cases)}

There were 25,549 cases reporting Drug ineffective that were indicative of occurrence of COVID-19 infection:

- in subjects who experienced COVID-19 infection from day 14 after receiving the first dose to day 6 after receipt of the second dose;

- in subjects who have not received the appropriate series of two doses or for whom it was not possible to determine whether they received the appropriate series of 2 doses at the appropriate interval;

- in subjects for whom it was not possible to determine how many days have passed since the first or second dose administration.

- Age groups: Infant (1), Child (214), Adolescent (565), Adult (18,448), Elderly (4350) and Unknown (1971).

- Reported COVID-19 infection related events (>2 occurrences)\textsuperscript{160}: COVID-19 (22,973), Suspected COVID-19 (2075), Asymptomatic COVID-19 (266), COVID-19 pneumonia (207), SARS-CoV-2 test positive (51), Breakthrough COVID-19 (44), Vaccine breakthrough infection (42), Post-acute COVID-19 syndrome (28), Multisystem inflammatory syndrome in children (7), Coronavirus infection (6), Coronavirus test positive, Multisystem inflammatory syndrome (4 each), and Pneumonia viral (3).

- Outcome of COVID-19 infection related events: resolved/resolving (3346), resolved with sequelae (160), not resolved (1245), unknown (20,746), and fatal (221).

According to the RSI, subjects may not be protected until at least 7 days after their second dose of the vaccine, therefore for the above 25,549 cases where lack of efficacy was reported, the reported events may represent signs and symptoms of intercurrent or undiagnosed COVID-19 infection or infection in an individual who was not fully vaccinated, rather than vaccine ineffectiveness.
SARS-CoV-2 Variants (11,901 cases)

In 11,901 of the 51,028 cases, information on SARS-CoV-2 variants was provided.

- **Delta (India) variant**\(^{162}\) (11,274 cases)\(^{163}\)
  - Country of incidence (>3 occurrences): Austria (11,164), France (84), Germany (16), and US (4).
  - Lack of efficacy events: Vaccination failure (6591) and Drug ineffective (4683).
  - Outcome of COVID-19 infection related events\(^{160}\): resolved/resolving (50), resolved with sequelae (2), not resolved (23), unknown (11,156), and fatal (51).

- **Omicron variant**\(^{162}\) (606 cases)
  - Country/region of incidence (>2 occurrences): Hong Kong (391), France (79), Germany (40), US (39), Japan (12), Spain (6), Austria (4), Belgium, Brazil, Mexico, and Norway (3 each).
  - Lack of efficacy events: Vaccination failure (404) and Drug ineffective (202).
  - Outcome of COVID-19 infection related events\(^{160}\): resolved/resolving (81), not resolved (11), unknown (503), and fatal (18).

- **Alpha (UK) variant**\(^{162}\) (19 cases)
  - Country of incidence: Austria, Germany (5 each), France, Italy (4 each), and Poland (1).
  - Lack of efficacy events: Vaccination failure (16) and Drug ineffective (3).
  - Outcome of COVID-19 infection related events\(^{160}\): resolved/resolving (8), not resolved (1), unknown (10), and fatal (1).

- **Others** (2 cases)
  - In 2 other cases, variant was reported as Beta (South Africa\(^{162}\)) and South African or Brazilian (as reported), respectively.

**Literature**

Review of the literature identified significant new information with regards to the use of BNT162b2 and lack of therapeutic efficacy. Please refer to Section 11 Literature and

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\(^{162}\) As per WHO Nomenclature (Countries in which earliest samples were documented were additionally listed, when applicable).

\(^{163}\) Includes 30 cases reporting SARS-CoV-2 variant as Indian variant/lineage specified as B.1.617 and 6 cases reporting SARS-CoV-2 variant as AY lineages.
Section 17.2 Newly Identified Information on Efficacy and Effectiveness for the review of these articles.

Conclusion

No new safety signals have emerged based on a review of these cases.

16.3.4.6. Off-Label Use

Search criteria - PTs Contraindicated product administered; Contraindicated product prescribed; Drug effective for unapproved indication; Drug ineffective for unapproved indication; Intentional device use issue; Intentional product use issue; Intentional underdose; Off label use; Off label use of device; Prescribed underdose; Product administered to patient of inappropriate age; Product use in unapproved indication; Product use issue; Therapeutic product effective for unapproved indication; Therapeutic product ineffective for unapproved indication.

Please refer to Section 6.3.1.1.2.3 Third Dose/Booster Dose for the amendments made regarding booster doses of the BNT162b2 vaccine.

Of the 38,130 cases, 8325 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

- 7960 cases reporting the PT Product administered to patient of inappropriate age (7958) and Contraindicated product administered (2) were found to be indicative of a potential medication error. These cases are referenced in Section 9.2 Medication Errors.

- 352 cases reported exposure in utero; these cases are referenced in Section 16.3.5.3 Use in Pregnant/Lactating Women.

- 9 cases reported the event Intentional product use issue (6) and Intentional underdose (3). Three (3) of these cases were not reported with use of the BNT162b2 vaccine. In 1 case, only the intention of misuse was reported. These cases did not report any additional events potentially indicative of off label use. The remaining 5 cases are referenced in Section 16.3.4.3 Abuse, Misuse and Drug Dependency.

- 1 case reported the (fatal) PT Drug ineffective for unapproved indication with no additional events indicative of off-label use. This case is referenced in Section 16.3.4.7 Unexpected Therapeutic Effect.

- 3 additional cases did not report relevant off label use (i.e., off label use not reported with the BNT162b2 vaccine).

Clinical Trial Data

Not applicable.
Post-Authorisation Data

- Number of cases: 29,805 (5.9% of 507,683 cases, the total PM dataset), compared to: 22,533 (3.4%) cases retrieved in the PSUR #2. A general increase in cases reporting Interchange of vaccine products was noted (54.0% of PM cases from PSUR #2 versus 83.3% of PM cases retrieved during this reporting period).

- MC cases (6091), NMC cases (23,714).

- Country of incidence (≥2%): UK (10,172), Netherlands (6230), Germany (4368), France (1516), Poland (602)

- Subjects' gender: female (20,994), male (7831) and unknown (980).

- Subjects' age in years (n = 26,283), range: 0.01–104 years, mean: 45.8 years, median: 44.0 years.

- Medical history (n = 12,399): the most frequently (≥2%) reported medical conditions include PT Disease risk factor (1663), COVID-19 (1591), Suspected COVID-19 (1448), Hypertension (1089), Breast feeding (1061), Asthma (746), Immunodeficiency (581), Hypothyroidism (340), Diabetes mellitus (319), Hypersensitivity (296), Steroid therapy (293), Depression (281), Drug hypersensitivity (279), Seasonal allergy (271).

- COVID-19 Medical history (n = 3001): the most frequently (≥2%) reported medical conditions included COVID-19 (1591) and Suspected COVID-19 (1448).

- Co-suspects (n = 1745 cases): the most frequently (≥2%) reported co-suspect vaccines/medications included COVID-19 vaccine MRNA (MRNA 1273) (681), COVID-19 vaccine NRVV AD (CHADOX1 NCOV-19) (420), Influenza vaccine (188), Influenza vaccine inact SAG 4V (80), Influenza vaccine inact SPLIT 4V (64), JNJ 78436735 (51), COVID-19 vaccine (50).

- Number of events: 174,381 (of which 32,211 were events of interest).

- Relevant event seriousness:42 serious (10,382), non-serious (21,845).

- Most frequently reported relevant PTs (≥2%): Off label use (29,562) and Product use issue (2531). Of note, of the 29,805 cases, 696 did not report additional events. The majority of cases described off-label use as
  - intentionally used in unapproved populations such as those mentioned below:
    - It is unknown whether the BNT162b2 vaccine is excreted in human milk.
    - Administration of the vaccine in pregnancy should be considered when potential benefits outweigh any potential risks for the mother and foetus.
o Immunocompromised persons, including individuals receiving immunosuppressant therapy, may have a diminished immune response to the vaccine.

o The administration of the BNT162b2 vaccine should be postponed in individuals suffering from acute severe febrile illness.

o The safety and efficacy have not yet been established in individuals under 5 years of age. The safety and effectiveness of a booster dose of in individuals 16 through 17 years of age is based on safety and effectiveness data in adults at least 18 through 55 years of age.

  - alternative dosing or scheduling regimens (i.e., Full primary series not received, longer/shorter number of days between doses than recommended)

o The primary series of the BNT162b2 vaccine is administered as 2 doses at greater than or equal to 21 days (preferably 3 weeks) apart. Off label is currently considered when the 2nd dose of the vaccine is administered outside the 19-42 day range from the 1st dose.

  - co-administration with other vaccines (i.e., influenza)

o No interaction studies have been performed

  - administration of COVID-19 vaccines from different manufacturers and third/booster/extra doses.

  - administration of COVID-19 vaccine formulations indicated for a different age group.

  - usage of poor quality COVID-19 vaccines due to either preparation (i.e., dilution technique) and/or storage issues (i.e., used after the expiry or beyond use date).

Analysis by dose interval

Among these cases, 9 (all non-serious) reported administration of 3 doses of BNT162b2 with different time intervals than the recommended posology and included the relevant PTs\(^{164}\) Off label use (9) and Product use issue (1).

  • Upon review, there were no significant differences were identified in the occurrence of the most frequently relevant PTs and clinical co-reported AEs reported in those who received the 3 doses of vaccine at a different time interval than the recommended posology when compared to the population receiving BNT162b2 in unapproved

\(^{164}\) More than 1 reported in 1 case.
conditions Clinical events reported more than once in this population included Headache (4), Pain, Pyrexia, and Vaccination site pain (2 each).

Literature

Review of the literature did not identify any significant new information with regards to the off-label use of BNT162b2.

Conclusion

Review of these cases did not identify new safety information related to off-label use.

16.3.4.7. Unexpected Therapeutic Effect

Search criteria - PTs Device effect increased; Drug effect faster than expected; Drug effective for unapproved indication; Therapeutic product effective for unapproved indication; Therapeutic response changed; Therapeutic response increased; Therapeutic response prolonged; Therapeutic response unexpected; Therapeutic product effect increased; Therapeutic product effect prolonged.

Clinical Trial Data

- There were no serious clinical trial cases with the above PTs reported during the reporting period; no serious cases were retrieved in the PSUR #2.

Post-Authorisation Data

- Number of cases: 664 (0.1 % of 507,683 cases in the total PM dataset), compared to 844 cases (0.1%) retrieved in the PSUR #2.
- MC cases (76), NMC cases (588).
- Country of incidence (≥10 occurrences): Germany (297), US (71), Netherlands (64), UK (34), Australia (28), Canada (20), France (18), Japan (17) Belgium (14), Sweden (10); the remaining 91 cases were distributed among 30 countries.
- Subjects’ gender: female (379), male (224), unknown (61).
- Subjects’ age in years (n = 371), range: 7 – 96, mean: 53.2, median: 54.
- Medical history (n = 489): the most frequently (≥10 occurrences) reported medical conditions included the PTs Asthma (30), Psoriasis (29), Migraine, Seasonal allergy (24 each), Multiple sclerosis, Pain (21 each), Diabetes mellitus, Skin papilloma (15 each), Hypertension (13), Fatigue (12), Depression, Fibromyalgia, and Headache (10 each).
- Co-suspects (n = 28 cases): COVID-19 vaccine mRNA (mRNA 1273) (7), bupropion, COVID-19 vaccine (2 each), adalimumab, atogepant, bupropion/naltrexone, COVID-19
vaccine NRVV AD, hepatitis A vaccine, ibuprofen, influenza vaccine, JNJ 78436735, levonorgestrel/ethinyl estradiol, palbociclib, PF-07321332/ritonavir, pneumococcal 13-valent conjugate vaccine, sarilumab, senna alexandrina extract, sucralfate, tetanus vaccine, thyroid (1 each).

- Number of events: 1447 (of which 664 were events of interest).
- Relevant event seriousness: serious (11), non-serious (653).
- Relevant PTs: Therapeutic response unexpected (656), Therapeutic response changed (5), Therapeutic product effect increased (3).
- In most of the cases (when specified), the unexpected therapeutic effect included improvement in the following: pain, breathing, allergies, skin conditions (including warts and psoriasis), autoimmune/inflammatory diseases (e.g., arthritis, multiple sclerosis, ulcerative colitis), migraine/headache, infections (e.g., herpes and other viral infections, fungal infections), neoplasia (remission/regression of various cancers), movement/mobility, energy, hair growth/loss, menstruation, and general health/well-being (e.g., “felt better”).
- Time to event onset (n = 146), range: <24 hours - 368 days, median: 1.5 days.
  - <24 hours: 50 events;
  - 1 day: 23 events;
  - 2-7 days: 35 events;
  - 8-14 days: 14 events;
  - 15-30 days: 9 events;
  - 31-180 days: 12 events;
  - >180 days: 3 events.
- Relevant event outcome: resolved/resolving (95), resolved with sequelae (1), not resolved (73), unknown (495).

**Analysis by age group**

- PM: Paediatric (3 [1 Child, 2 Adolescent]), Adults (285), Elderly (109), Unknown (267).
  - There was no meaningful difference between different age groups.

**Literature**

Review of the literature did not identify any significant new information regarding the use of BNT162b2 and unexpected therapeutic effects.

**Conclusion**

In most of the cases (when specified), the unexpected therapeutic effect included improvement in the following: pain, breathing, allergies, skin conditions (including warts and psoriasis), autoimmune/inflammatory diseases (e.g., arthritis, multiple sclerosis, ulcerative colitis), migraine/headache, infections (e.g., herpes and other viral infections, fungal
infections), neoplasia (remission/regression of various cancers), movement/mobility, energy, hair growth/loss, menstruation, and general health/well-being (e.g., “felt better”). In the majority of the cases, the subject experienced the unexpected therapeutic effect following the first dose (when recorded).

No significant new information was identified with regards the use of BNT162b2 and unexpected therapeutic effects.

16.3.5. Update on Special Patient Populations

Any new data identified during the reporting interval for use of BNT162b2 by special patient populations is analysed below.

16.3.5.1. Use in Elderly Patients

Of the 56,799 cases, 4 post-marketing cases were determined to be non-contributory and were not included in the discussion for the following reason:

- upon review, these 4 cases reported the PT Maternal exposure during pregnancy (4) in subjects greater than 65 years of age (in these 4 cases, it is most likely that the subjects’ age was erroneously reported).

Clinical Trial Data

- Number of cases: 211 (BNT162b2 [180], blinded therapy [26], placebo [4], BNT162b2S01 [1]); (31.6 % of 668 cases in the total CT dataset), compared to 233 cases (32.3%) retrieved in the PSUR #2.
- Country of incidence: US (162), Argentina (26), Germany (9), Brazil (7), China (4), Israel (2), Dominican Republic (1).
- Subjects' gender: female (88), male (123).
- Subjects' age in years (n = 211), range: 65 – 87, mean: 73.1, median: 73.
- Medical history (n = 193): the most frequently (≥20 occurrences) reported medical conditions included the following HLGTs: Vascular hypertensive disorders (114), Lipid metabolism disorders (73), Joint disorders (63), Glucose metabolism disorders (incl diabetes mellitus) (51), Gastrointestinal motility and defaecation conditions (42), Appetite and general nutritional disorders (35), Allergic conditions (31), Depressed mood disorders and disturbances (28), Cardiac arrhythmias (26), Sleep disorders and disturbances (25), Coronary artery disorders, Thyroid gland disorders (24 each), Gastrointestinal therapeutic procedures (23), Bronchial disorders (excl neoplasms) (22), Anxiety disorders and symptoms, Prostatic disorders (excl infections and inflammations) (20 each).
- COVID-19 Medical history: None.
- Co-suspects (n = 4 cases): amiodarone, amlodipine, atenolol, diltiazem, etoricoxib, furosemide, hydrochlorothiazide/triamterene, hydroxyzine, losartan, metformin, sulfamethoxazole/trimethoprim, tamsulosin (1 each).
- Number of events: 274.
Most frequently (≥5 occurrences) reported PTs: Atrial fibrillation (11), Cerebrovascular accident, Osteoarthritis, Prostate cancer (9 each), Condition aggravated\textsuperscript{165} (8), Acute kidney injury, Acute respiratory failure, Dyspnœa (5 each).

Of the 274 events, the only related event was for BNT162b2 and coded to the PT Dehydration (1).

Time to event onset: n = 247, range: from <24 hours to 504 days, median: 125 days.
- <24 hours: 2 events;
- 1 day: 3 events;
- 2-7 days: no events;
- 8-14 days: 3 events;
- 15-30 days: 12 events;
- 31-180 days: 179 events (20 of which had a fatal outcome);
- >180 days: 48 events. (6 of which had a fatal outcome).

Event outcome: fatal (27), resolved/resolving (185), resolved with sequelae (19), not resolved (43).

Post-Authorisation Data
- Number of cases: 56,584 (11.1% of 507,683 cases in the total PM dataset), compared to 87,982 cases (13.4%) retrieved in the PSUR #2.
- MC cases (28,690), NMC cases (27,894).
- Country of incidence (≥500 occurrences): Germany (10,884), Austria (9277), France (7504), US (3203), UK (3119), Japan (2225), Netherlands (2140), Sweden (2091), Australia (1747), Italy (1596), Spain (1187), Malaysia (1025), Denmark (992), Belgium (984), Poland (804), Philippines (689), Slovenia (614), Norway (602), Finland (592), Canada (556); the remaining 4753 cases were distributed among 62 countries.
- Subjects' gender: female (33,348), male (22,179), unknown (1057).
- Subjects' age in years (n = 54,943), range: 65 – 120, mean: 73.9, median: 72.
- Medical history (n = 18,647): the most frequently (≥1000 occurrences) reported medical conditions included the following HLGTs: Vascular hypertensive disorders (6169), Glucose metabolism disorders (incl diabetes mellitus) (2721), Allergic conditions (2365), Bronchial disorders (excl neoplasms) (1789), Cardiac arrhythmias (1712), Lipid metabolism disorders (1683), Joint disorders (1580), Thyroid gland disorders (1409), Therapeutic procedures and supportive care NEC (1392), Lifestyle issues (1342), Coronary artery disorders (1226), Central nervous system vascular disorders (1081).

\textsuperscript{165} The aggravated condition were: Adenocarcinoma pancreas, Back pain, Benign prostatic hyperplasia, Cardiac failure congestive, End stage renal disease, Gastroesophageal reflux disease, Hypertrophic cardiomyopathy, Muscle rupture, and Supraventricular tachycardia (1 each).

- Co-suspects (n = 1956 cases) the most frequently (≥10 occurrences) reported co-suspect medications included: COVID-19 vaccine (422), COVID-19 vaccine mRNA (mRNA 1273) (274), COVID-19 vaccine NRVV AD (258), influenza vaccine (154), adalimumab (151), influenza vaccine INACT SAG 4V (100), influenza vaccine INACT SPLIT 4V (58), pneumococcal polysaccharide vaccine 23-valent (32), apixaban (29), upadacitinib (24), influenza vaccine INACT SAG 3V (23), JNJ 78436735 (18), prednisone (17), mepolizumab (16), rivaroxaban (13), rituximab (12), casirivimab, imdevimab, influenza vaccine INACT SPLIT 3V (11 each), atorvastatin, ibritinib, levodopa, risankizumab (10 each).

- Number of events: 167,970; the most frequently (>1000 occurrences) reported events were coded to the PTs were: COVID-19 (8394), Inappropriate schedule of product administration (5063), Fatigue (4864), Headache (4712), Drug ineffective (4627), Vaccination failure (4515), Pyrexia (4261), Off label use (3847), Myalgia (3431), Immunisation (3355), Arthralgia (3137), Interchange of vaccine products (2920), Dizziness (2815), Vaccination site pain (2798), Pain in extremity (2796), Malaise (2468), Dyspnoea (2455), Nausea (2155), Chills (2012), Asthenia (1666), Herpes zoster (1771), Pain (1706), Rash (1677), Pruritus (1194), Vomiting (1178), Diarrhoea (1175), Paraesthesia (1096), Chest pain (1089).

- Event seriousness: 42 serious (73,170), non-serious (94,882).

- Time to event onset (n = 119,721), 166 range: from <24 hours to 492 days, median: 2 days.
  - <24 hours: 37,098 events (733 of which had a fatal outcome);
  - 1 day: 19,235 events (440 of which had a fatal outcome);
  - 2-7 days: 21,444 events (674 of which had a fatal outcome);
  - 8-14 days: 9213 events (373 of which had a fatal outcome);
  - 15-30 days: 8260 events (395 of which had a fatal outcome);
  - 31-180 days: 21,670 events (991 of which had a fatal outcome);
  - >180 days: 2801 events (168 of which had a fatal outcome).

- Event outcome: 78 fatal (5367), resolved/resolving (52,311), resolved with sequelae (4003), not resolved (39,949), unknown (66,774).

166 This number does not include 48,801 events for which administration and/or event onset dates were not provided or were incomplete; or events without a meaningful time to onset value as per reported information. Please note, multiple episodes of the same PT event were reported with different latencies within some cases hence the sum of latencies exceeds the total number of PT events.
Analysis by presence of comorbidities

- Number of elderly subjects with reported comorbidities: 10,304 (18.2% of the 56,584 cases in the total elderly dataset).

- Of the cases that reported medical histories, the percentage of cases reporting an AE with a fatal outcome is higher in subjects with comorbid conditions (72.1%) when compared to the percentage of cases involving an AE with a fatal outcome in subjects without comorbidities (27.9%).

- Upon review of the most frequently (≥200 occurrences) reported AEs in cases that recorded medical histories, the PT COVID-19 Pneumonia was the only event that had a significant proportional reporting ratio of >3:1 in the elderly population with comorbidities compared to the elderly population without comorbidities.

Literature

Review of the literature did not identify any significant new information regarding the use of BNT162b2 in elderly patients.

Conclusion

No significant differences in the reporting proportion of the most frequently reported AEs were noted between the elderly dataset and the non-elderly dataset, apart from the PTs indicative of lack of therapeutic effect, for which the reporting proportion is higher in the elderly population: COVID-19 (14.8% versus 8.8%) and Vaccination failure (7.9% versus 4.4%). This is expected due to age-related decline in immunity that results not only in increased susceptibility to infection, but also reduces the prophylactic efficacy of vaccinations\(^\text{167}\).

The interval data reviewed did not identify any new safety information regarding the use of BNT162b2 in elderly patients.

16.3.5.2. Use in Paediatric Patients

Search criteria - Paediatric cases are identified as cases where the Age Range derived field value for the patient is “Less than or equal to 17 years”. Cases indicative of exposure to the vaccine during the mother’s pregnancy or through breastfeeding were excluded.

- Of the 31,930 cases, 3 cases were determined to be non-contributory and were not included in the discussion for the following reasons:

in 2 cases the data reported (e.g., height, weight, clinical data) were not consistent with paediatric subjects;

- in 1 case, exposure to the vaccine occurred during the mother’s pregnancy.

16.3.5.2.1. Paediatric Subjects <5 Years of Age\textsuperscript{168}

Clinical Trial Data

- Number of cases: 62 (blinded therapy [43], BNT162b2 [18] and pre-randomisation\textsuperscript{169} [1]), originated from Protocols C4591007, C4591007-OPENLABEL and C4591024 (9.3% of 668 cases, the total CT dataset), compared to 25 cases (3.5%) retrieved in the PSUR #2.

- Country of incidence of relevant cases: US (28), Poland (21), Brazil, Spain (4 each), Finland, and Germany (2 each).

- Subjects' gender: female (28), male (33).

- Subjects' age in years (n = 61), range: 0.58 – 4, mean: 2.4, median: 2.0.

- Medical history (n = 22): the most frequently reported (\geq 2): Chronic kidney disease, Renal transplant (4 each), Febrile convulsion, Gastrostomy, Urethral valves (3 each), Asthma Food allergy, Autism spectrum disorder, Biopsy kidney, Cough, Cystostomy, Eating disorder, Eczema, Heart transplant, Hypospadias, Orchidectomy, Orchidopexy, Pulmonary valves stenosis, Pyrexia, Reflux nephropathy, Respiratory disorder, Stem cell transplant, and Suture removal (2 each).

- COVID-19 Medical history: None.

- Co-suspects (n = 1): Paracetamol (1).

- PTs reported in the relevant cases (n=67): PTs reported in more than 1 case: Febrile convulsion, Gastroenteritis, Gastroenteritis rotavirus (4 each), Bronchiolitis (3), Adenovirus infection, Anaphylactic reaction, Appendicitis, Dehydration, Humerus fracture, Lower respiratory tract infection, Metapneumovirus infection, Pneumonia, and Pyrexia (2 each).

All events were assessed as unrelated to BNT162b2 or blinded therapy.

- Time to event onset: n =66,\textsuperscript{170} range: from 1 day to 298 days, median: 59 days.
  - 1 day: 2 events;

\textsuperscript{168} The paediatric vaccine for individuals aged between 6 months and 4 years was approved first in the US on 17 June 2022; the administration of BNT162b2 in subjects < 5 years was unapproved during the reporting period of this PSUR.

\textsuperscript{169} This case is not included in the analysis below.

\textsuperscript{170} This number does not include 1 event for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
- 2-7 days: 10 events;
- 8-14 days: 1 event;
- 15-30 days: 11 events;
- 31-298 days: 42 events.

- Duration of relevant events (n = 60 out of 61 occurrences with outcome of resolved/resolved with sequelae)\(^{171}\), range: >1 day to 36 days, median: 4 days.
  - <24 hours: 6 events
  - 1 day: 10 events
  - 2-7 days: 23 events
  - 8-14 days: 16 events
  - 15-36 days: 5 events.

- Event outcome: resolved/resolving (63), not resolved (3), resolved with sequelae (1).

**Post-Authorisation Data**

- Number of cases: 275\(^{172}\) (0.5% of 507,683 cases, the total PM dataset), compared to 83 cases (0.01%) retrieved in the PSUR #2.
- MC cases (78), NMC cases (197).
- Country of incidence (≥1%): Germany (179), Iraq (28), US (18), Australia (15), France (9), Austria (7), Ireland, UK (4 each), Italy, and Philippines (3 each).
- Subjects' gender: female (119), male (142) and unknown (14).
- Subjects' age in years (n = 267), range: 0.01-4.50, mean: 2.4, median: 2.33.
- Medical history (n = 12): the reported medical conditions included Breast feeding, Cardiac disorder (2 each), Abnormal behaviour, Arrhythmia, Asthma, Atelectasis, Cardiomyopathy, Cytogenetic abnormality, Feeding disorder, Food allergy, Gait inability, Gastrooesophageal reflux disease, Hypersensitivity, Hypertension, Infection, Lactose intolerance, Lung disorder, Mite allergy, Pyelonephritis, Ventricular septal defect, and Weight gain poor (1 each).
- Co-suspect vaccines (n = 2): influenza vaccine and influenza vaccine inactive split 3V (1 each).

\(^{171}\) This number does not include 1 event for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\(^{172}\) Cross-referenced with Section 16.3.4.6 Off-Label Use, since the administration of BNT162b2 was approved in subjects <5 years of age (≥ 6 months of age) only in the US since 17 June 2022.
• Number of events: 856. The most frequently reported PTs (>5): Product administered to patient of inappropriate age (202), Off label use (133), Vaccination site pain (88), Product use issue (75), Pyrexia (43), Fatigue (37), Headache, Rash, Vaccination site erythema (14 each), Arthralgia, Vaccination error (10 each), Diarrhoea (8), Malaise, Urticaria (7 each), Chest pain, Chills, Nausea, and Vomiting (6 each).

• Event seriousness\(^{173}\): (58), non-serious (799).

• Time to event onset: n = 735\(^{174}\), range: from <1 day to 28 days, median: <24 hours.
  - <1 day: 558 events;
  - 1 day: 87 events;
  - 2-7 days: 62 events;
  - 8-14 days: 8 events;
  - 15-28 days: 20 events.

• Duration of relevant events (n = 212 out of 268 occurrences with outcome of resolved/resolved with sequela)\(^{175}\), range: from <1 day to 62 days, median: 1 day.
  - <1 day: 53 events;
  - 1 day: 81 events;
  - 2-7 days: 72 events;
  - 8-14 days: 3 events;
  - 14-62 days: 3 events.

• Event outcome\(^{178}\): resolved/resolving (279), not resolved (87), resolved with sequela (12), unknown (480).

16.3.5.2.2. Paediatric Subjects ≥5 Years and ≤11 Years of Age\(^{176}\)

Clinical Trial Data

• Number of cases: 25 (blinded therapy [6] and BNT162b2 [19]), originated from Protocols C4591007, C4591007-OPENLABEL and C4591024 (3.7% of 668 cases, the total CT dataset), compared to 18 cases (2.5%) retrieved in the PSUR #2.

• Country of incidence: US (13), Brazil (6), Germany (3), Poland (2), and Spain (1).

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\(^{173}\) One case reported different seriousness for the same event.

\(^{174}\) This number does not include 124 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\(^{175}\) This number does not include 56 events with outcome resolved or resolved with sequela for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\(^{176}\) The administration of BNT162b2 in ≥ 5 years to ≤ 11 years population was approved by EMA on 25 November 2021.
Subjects' gender: female (8), male (17).

Subjects' age in years (n = 25), range: 5 – 11, mean: 7.7, median: 7.0

Medical history (n = 21): the most frequently (≥2) reported medical conditions included Renal transplant (4), Dermatomyositis (3), Anal stenosis, Anoplasty, Asthma, Attention deficit hyperactivity disorder, Colostomy, Constipation, Currarino syndrome, Epilepsy, Factor V Leiden mutation, Inguinal hernia repair, Juvenile idiopathic arthritis, Malnutrition, Spinal cord operation, Spondylolisthesis, and Tumour necrosis factor receptor-associated periodic syndrome (2 each).

COVID-19 Medical history (n = 1): COVID-19 (1).

Co-suspect vaccines/medications: none.

PTs (34): Gastroenteritis (3), Dermatomyositis, Intestinal obstruction, Pyrexia (2 each), Appendicitis, Asthma, Colitis, Condition aggravated, Constipation, Depression, Depression suicidal, Device related infection, Diarrhoea, Drug therapy, Febrile convulsion, Hypertension, Hyponatraemia, Influenza, Kidney transplant rejection, Large intestine benign neoplasm, Mental status changes, Myalgia, Myositis, Rhinovirus infection, Seizure, Small intestinal obstruction, Syncope, Tibia fracture, and Vomiting (1 each).

All events were assessed as unrelated to BNT162b2 or blinded therapy.

Time to event onset: n = 34, range: 1 day to 282 days, median: 83 days.
  - 1 day: 1 event;
  - 2-14 days: 0 events;
  - 15-30 days: 3 events;
  - 31-90 days: 15 events;
  - 91-282 days: 15 events.

Duration of relevant events (n = 30 out of 34 occurrences with outcome of resolved/resolved with sequelae)\(^{177}\), range: 1 day to 10 days, median 3 days.
  - 1 day: 9 events;
  - 2-7 days: 18 events;
  - 8-10 days: 3 events.

Event outcome: resolved/resolving (31), resolved with sequelae (3).

Post-Authorisation Data

Number of cases: 9605 (1.9% of 507,683 cases, the total PM dataset), compared to 1227 cases (0.2%) retrieved in the PSUR #2.

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\(^{177}\) This number does not include 4 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
COVID-19 mRNA vaccine (nucleoside modified)  
Reporting Period  
Periodic Safety Update Report (PSUR) 3  
19 December 2021 through 18 June 2022

- MC cases (6573), NMC cases (3032).
- Country of incidence (≥2%): US (2503), Australia (1428), Philippines (1264), Germany (1177), Japan (859), Italy (409), and Spain (386).
- Subjects' gender: female (3925), male (4133) and unknown (1547).
- Subjects' age in years (n = 8372), range: 5 – 11,25, mean: 8.4, median: 9.0.
- Medical history (n = 846): the most frequently (≥10) reported medical conditions included Asthma (158), Food allergy (62), Seasonal allergy (59), Attention deficit hyperactivity disorder (41), Hypersensitivity (34), Autism spectrum disorder, Epilepsy (30 each), Drug hypersensitivity (28), Rhinitis allergic (27), Eczema (26), Dermatitis atopic (23), Mite allergy (21), Allergy to animal (19), Constipation, Type 1 diabetes mellitus, Urticaria (15 each), Bronchospasm, Headache, Seizure (12 each), Obesity (11), Migraine (10).
- COVID-19 Medical history (n = 136): COVID-19 (121), Suspected COVID-19 (10), Asymptomatic COVID-19, Exposure to SARS-CoV-2 (2 each), and Post-acute COVID-19 syndrome (1).
- Co-suspects (n = 44): the most frequently (>1) reported co-suspect vaccines/medications included influenza vaccine (8), adalimumab, COVID-19 vaccine (7 each), measles vaccine live (Enders-Edmonston)/ mumps vaccine live (Jeryl Lynn)/ rubella vaccine live (Wistar RA 27/3), varicella zoster vaccine live (Oka/Merck) (3 each), diphtheria vaccine toxoid/ pertussis vaccine acellular/tetanus vaccine toxoid, influenza vaccine inact split 3V, meningococcal vaccine B RFHP, NADA, NHBA OMV, and sodium chloride (2 each).
- Number of events: 22,457.
- Event seriousness:42 (3735), non-serious (18,725).
- Most frequently reported PTs (>3% of cases): Product administered to patient of inappropriate age (1338), Pyrexia (1289), Vaccination site pain (1213), Poor quality product administered (1063), Headache (976), Product administration error (753), Vomiting (733), Rash (556), Overdose (516), Product preparation error (429), Fatigue (425), Nausea (410), Abdominal pain (371), Dizziness (366), Chest pain (331), COVID-19 (309), Pain in extremity (293), and Underdose (290).
- Time to event onset (n = 16,236)178, range: from <1 day to 385 days, median: <1 day.
  - <1 day: 8574 events;
  - 1 day: 3414 events;
  - 2 days: 1102 events;
  - 3-7 days: 1594 events;
  - 8-14 days: 593 events;
  - 15-30 days: 551 events;

178 This number does not include 6242 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
- 31-60 days: 273 events;
- 61-385 days: 135 events.

- Duration of relevant events (n = 3787 out of 7329 occurrences with outcome of resolved/resolved with sequelae)\(^{179}\), range: from <1 day to 109 days, median 1 day.
  - <1 day: 1258 events;
  - 1 day: 955 events;
  - 2-7 days: 1263 events;
  - 8-14 days: 171 events;
  - 15-109 days: 140 events.

- Relevant event outcome: resolved/resolving (9811), resolved with sequelae (73), not resolved (3274), fatal (58), unknown (9257).

- Fatal cases: 20
  - Age: 5 years (1), 6 years (3), 7 years (4), 8 years (2), 9 years (1), 10 years (2), 11 years (5), unknown (2).
  - MC cases (17), NMC cases (3).
  - Gender: females (9), males (9), unknown (2).
  - Country: Philippines (6), Australia (4), Germany, Spain (3 each), Albania, Japan, Portugal, UK (1 each).
  - Fatal PTs (58): the most frequently (≥2) reported AEs included Dyspnoea (4), Cardiac arrest, Cardio-respiratory arrest, Pyrexia (3 each), Abdominal pain, Cough, COVID-19, Death, Headache, Myocarditis, Seizure, and Vomiting (2 each).
  - Medical history (n = 7): Autoimmune thyroiditis, Asphyxiating thoracic dystrophy, Brain malformation, Bronchitis, Bronchospasm, Cerebral palsy, Cognitive disorder, COVID-19, Dependence on respirator, Developmental delay, Dysphagia, Epilepsy, Gastrostomy, Hypoxic-ischaemic encephalopathy, Immunodeficiency, Intellectual disability, Joint dislocation, Kidney transplant rejection, Motor dysfunction, Myoclonic epilepsy, Neonatal asphyxia, Obstructive sleep apnoea syndrome, Pneumonia, Renal impairment, Renal transplant, Rhinitis allergic, Scoliosis, Seizure, Severe myoclonic epilepsy of infancy, Type 1 diabetes mellitus, and Varicella zoster virus infection (1 each).

The 20 fatal cases are summarised below:

- In 2 cases (1 MC and 1 NMC) reporting only Death as the fatal AE, neither cause of death nor information on autopsy was provided. The limited information provided prevented any meaningful assessment.

\(^{179}\) This number does not include 3542 events with outcome resolved or resolved with sequelae for which time to event onset partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
- In 2 cases, the underlying medical conditions could have predisposed to the occurrence of the fatal AEs:

  - MC case; age: 11 years; gender: male; fatal PT: Acute respiratory failure, occurred 2 days after the 1st dose of BNT162b2; medical history: brain malformation, bronchitis, cognitive disorder, dysphagia, gastrostomy, joint dislocation, myoclonic epilepsy, obstructive sleep apnoea syndrome, pneumonia, scoliosis; autopsy: not performed.

  - MC case; age: 6 years; gender: female; fatal PTs: Renal impairment, Epilepsy, Apnoea, Varicella zoster virus infection, Seizure, Sudden death, Product administered to patient of inappropriate age, death occurred 7 days after the 1st dose of BNT162b2; medical history: developmental delay, epilepsy, immunodeficiency, renal impairment, seizure, severe myoclonic epilepsy of infancy, varicella zoster virus infection; autopsy: unknown if performed.

- In one case, the reporter concluded that the death "had nothing to do" with the administration of BNT162b2 and was due to natural causes:

  - MC case; age: 6 years; gender: male; fatal PTs: Myocarditis, Cardio-respiratory arrest, COVID-19, occurred 7 days after the 1st dose of BNT162b2; medical history: rhinitis allergic, autoimmune thyroiditis), type I diabetes mellitus); autopsy: performed, results are pending.

  - In the remaining 15 cases (13 MC and 2 NMC) reporting the following fatal PTs Dyspnoea (4), Cardiac arrest, Pyrexia (3 each), Abdominal pain, Cardio-respiratory arrest, Cough, Headache, Vomiting (2 each), Abdominal pain upper, Acute respiratory distress syndrome, Adverse event following immunisation, Arteriovenous malformation, Blood pressure decreased, Blood pressure immeasurable, Bradycardia, Cardiac failure acute, Cerebral haemorrhage, COVID-19, Cyanosis, Diarrhoea, Drug ineffective, haematemesis, Heart rate decreased, Immunisation, Influenza like illness, Multisystem inflammatory syndrome, Myocarditis, Nasopharyngitis, Nausea, Off label use, Pulmonary embolism, Respiratory failure, and Seizure (1 each), no confounding factors have been identified. In most cases (9) the limited information available does not allow a medically meaningful assessment; in the remaining cases (6) a causality between the vaccination and the occurrence of the fatalities cannot be ruled out, based on the temporal relationship, although no laboratory data or autopsy results provided evidence of a causal relationship.
16.3.5.2.3. Paediatric Subjects ≥12 Years of Age\textsuperscript{180}

Clinical Trial Data

- Number of cases: 15 (BNT162b2 [14] and blinded therapy [1]) originated from Protocol C4591001 (2), C4591001-OPEN LABEL (10), C4591007-OPEN LABEL (1), C4591024 (1), and C4591031-OPEN LABEL (1) (2.2% of 668 cases, the total CT dataset), compared to 24 cases (3.3%) retrieved in the PSUR #2.
- Country of incidence: US (14) and Germany (1).
- Subjects' gender: female (7) and male (8).
- Subjects' age in years (n = 15), range: 12 – 17, mean: 14.6, median: 15.0.
- Medical history (n = 11): the most frequently (≥2) reported medical conditions included Anxiety (6), Seasonal allergy (4), Depression (3), Attention deficit hyperactivity disorder, Insomnia, and Rhinitis allergic (2 each).
- COVID-19 Medical history: None.
- Co-suspects (n = 1): aripiprazole, duloxetine (1 each).
- PTs (17): Suicidal ideation, Suicide attempt, Toxic shock syndrome (2 each), Addison’s disease, Appendicitis, Constipation, Depression, Fractured skull depressed, Herpes zoster, Major depression, Mucocutaneous rash, Pectus excavatum, Subdural haematoma, and Syncope (1 each).
- All events were assessed as unrelated to BNT162b2 or blinded therapy.
- Time to event onset (n = 17), range: from 6 days to 284 days, median: 96 days.
  - 6 days: 1 event;
  - 7-30 days: no events;
  - 31-90 days: 7 events;
  - 91-284 days: 9 events.
- Duration of relevant events (n = 8 out of 8 occurrences with outcome of resolved/resolved with sequelae), range: from <1 day to 86 days, median 3.5 days.
  - <1 day: 1 event;
  - 1 day: 1 event;
  - 2-7 days: 5 events;
  - 86 days: 1 event.
- Event outcome: resolved/resolving (14), not resolved (2), resolved with sequelae (1).

\textsuperscript{180} The administration of BNT162b2 in this subpopulation was approved by EMA on 31 May 2021.
Post-Authorisation Data

- Number of cases: 21,945 (4.3% of 507,683 cases, the total PM dataset), compared to 18,451 cases (2.8%) retrieved in the PSUR #2.
- MC cases (13,478), NMC cases (8467).
- Country of incidence (>2%): Germany (3333), Philippines (3026), Australia (2220), UK (1656), Austria (1645), Malaysia (1189), Taiwan, Province of China (1186), France (1061), US (1025), Italy (628), Netherlands (582), Japan (484), and Mexico (454).
- Subjects' gender: female (11,656), male (9813) and unknown (476).
- Subjects' age in years (n = 21,661), range: 12 - 17, mean: 14.7, median: 15.0.
- Medical history (n = 2837): the most frequently (>1%) reported medical conditions included Asthma (286), Hypersensitivity (210), Seasonal allergy (194), Food allergy (108), Attention deficit hyperactivity disorder (90), Drug hypersensitivity (74), Mite allergy (73), Epilepsy (62), Depression (51), Autism spectrum disorder (48), Allergy to animal (45), Anxiety, Migraine (40 each), Immunodeficiency, Obesity (39 each), Rhinitis allergic (37), Eczema (35), Non-tobacco user (34), Acne (33), Headache (31), and Dermatitis atopic (29).
- Co-suspects (n = 148): the most frequently (>2%) reported co-suspect vaccines/medications included COVID-19 vaccine (33), adalimumab (18), influenza vaccine (15), COVID-19 vaccine mRNA (MRNA 1273) (13), influenza vaccine inactive split 4V, mestanol/norethisterone (8 each), HPV vaccine VLP R.L. 2V (yeast) (7), HPV vaccine VLP R.L. 2V (baculovirus) (5), HPV vaccine, infliximab (4 each), ibuprofen, and semaglutide (3 each).
- Number of events: 61,071.
- Relevant event seriousness:4 serious (19,558), non-serious (41,530).
- Most frequently reported PTs (>2%): Headache (3495), Pyrexia (3395), Dizziness (2376), Chest pain (1956), Fatigue (1919), Vaccination site pain (1804), Nausea (1669), COVID-19 (1600), and Dyspnoea (1267).
- Time to event onset: (n = 45,162)181, range: from <1 day to 476 days, median: 1 day.
  - <1 day: 18,167 events;
  - 1 day: 10,527 events;
  - 2-7 days: 7412 events;
  - 8-14 days: 2061 events;

181 This number does not include 16,067 events for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
- 15-30 days: 2003 events;
- 31-181 days: 4802 events;
- 182-476 days: 190 events.

- Duration of relevant events (n = 9201 out of 19,141 occurrences with outcome of resolved/resolved with sequelae)\(^{182}\), range: <1 day to 329 days, median 1 day.
  - <1 day: 2914 events;
  - 1 day: 1832 events;
  - 2-7 days: 3247 events;
  - 8-14 days: 530 events;
  - 15-30 days: 317 events;
  - 31-181 days: 341 events;
  - 182-329 days: 20 events.

- Relevant event outcome: fatal (169), resolved/resolving (28,719), not resolved (12,336), resolved with sequelae (332), unknown (19,645).

**Fatal cases (62)**

- Age: 12 years (12), 13 years (13), 14 years (5), 15 years (6), 16 years (11), 17 years (9), unknown (6).
- MC cases (45), NMC cases (17).
- Gender: females (28), males (32), unknown (2).
- Country (≥ 2): Philippines (19), US (8), Malaysia, Poland (6 each), Germany (4), Austria, Brazil, Japan, Taiwan (Province of China), UK (2 each).
- Fatal PTs (169): the most frequently (≥ 3) reported AEs included Death (16), Dyspnoea (8), Pyrexia (7), Cardiac arrest (6), Myocarditis (5), Cardiac failure, Headache (4 each), Asthenia, Seizure, Shock, and Vomiting (3 each).
- Medical history (n = 13): Attention deficit hyperactivity disorder, Obesity (2 each), Abdominal pain, Agitation, Amenorrhoea, Asthma, Bedridden, Chest pain, Colloid brain cyst, Cough, Cystic fibrosis, Cyst removal, Decreased appetite, Depression, Diabetes insipidus, Dizziness, Drug hypersensitivity, Dyspnoea, Dyssomnia, Exercise adequate, Fatigue, Feeling abnormal, Fracture, Headache, Hereditary cerebral degeneration, Hypertension, Kawasaki's disease, Lipoedema, Liver disorder, Lymphoedema, Lymphostasis, Oral contraception, Osteogenesis imperfecta, Ovarian enlargement, Palpitations, Physical deconditioning, Pulmonary embolism, Pulmonary veno-occlusive disease, Seasonal allergy, Somatic symptom disorder, Substance abuser, Substance use, and Weight decreased (1 each).

\(^{182}\) This number does not include 9940 events with outcome resolved or resolved with sequelae for which partial administration or event onset dates were reported or events without a meaningful time to onset value as per reported information.
The 62 fatal cases are summarised below:

- In 15 cases (9 MC and 6 NMC) reporting only Death as the fatal AE, neither cause of death nor information on autopsy was provided. The time to fatal event onset is available in 4 cases: 5 days, 7 days, 49 days, and 144 days (1 each). The limited information provided prevented any meaningful assessment.

- In 2 cases, the subjects did not die due to illness, but due to unfortunate accidents:
  - MC case; age: 17 years; gender: male; fatal PT: Fall, occurred 24 days after the vaccination; autopsy: unknown if performed.
  - MC case; age: 16 years; gender: male; fatal PT: Road traffic accident, occurred approximately 110 days after the 2nd dose of BNT162b2; autopsy: unknown if performed.

- In 6 cases, the underlying medical conditions could have predisposed to the occurrence of the fatal AEs:
  - MC case; age: 16 years; gender: female; fatal PT: Dyspnorea, occurred 3 days after the 1st dose of BNT162b2; medical history: bronchial asthma; autopsy: unknown if performed.
  - NMC case; age: 16 years; gender: female; fatal PTs: Dyspnorea (developed 6 days after the 1st dose of BNT162b2), Brain injury, Cardiac failure acute, Hypoxia, Cardiac failure (all developed 38 days after the 1st dose of BNT162b2), Sudden death, Pulmonary veno-occlusive disease, Pulmonary arterial hypertension (all developed 41 days after the 1st dose of BNT162b2), Brain oedema, Sudden cardiac death, Brain injury, Acute kidney injury, Pneumonitis, Epistaxis, Acute respiratory failure, Cardiac failure congestive (all unknown onset date); medical history: pulmonary veno-occlusive disease, amenorrhoea, cough, dyspnorea, fatigue, fracture, hypertension, lipoedema, lymphoedema, lymphostasis, osteogenesis imperfecta, ovarian enlargement; autopsy: not performed.
  - NMC case; age: 16 years; gender: female; fatal PTs: Pulmonary embolism, Cardiac arrest (all developed 2 days after the 3rd dose of BNT162b2); medical history: obesity, oral contraception, pulmonary embolism; autopsy: performed, results not provided.
  - MC case; age: 17 years; gender: male; fatal PTs Pneumococcal sepsis, Cardiac failure, Pneumonia pneumococcal (all occurred 92 days after the 2nd dose of BNT162b2); medical history: agitation, attention deficit hyperactivity disorder, depression, dyssomnia, regular exercise. Autopsy results: the subject died after consumption of from the beginning pneumonia and the influx of germs into the bloodstream as a result of cardiovascular failure. The concentration determined in the blood and brain does not justify in itself a fatal intoxication in view of a long-
term intake with a tolerance effect but may have favoured the onset of death due to a substance-typical respiratory and circulatory depressive effect, also increased in combination with the effect of [masked]. The findings obtained during the autopsy and the results of the chemical-toxicological examination can be reconciled with a protracted occurrence of death.

- MC case; age: 13 years; gender: female subject; fatal PTs: Malaise (developed 1 day after the 1st dose of BNT162b2, Lot number FJ1763), Palpitations, Chest pain (all occurred 5 days after the 1st dose of BNT162b2, Lot number FJ1763), Loss of consciousness, Pulseless electrical activity, Cardiac arrest (all occurred 64 days after the 1st dose of BNT162b2, Lot number FJ1763); medical history: Kawasaki's disease, palpitations, weight decreased, decreased appetite, feeling abnormal. Autopsy results showed that there was no possibility of myocarditis and angina pectoris, and there was no thrombus. Since symptoms such as episodes of palpitations had appeared before the vaccination, it was assessed that the vaccination was possibly related to the death, but the possibility of being the exacerbation factor could not be ruled out.

- MC case; age: 13 years; gender: male; fatal PTs Brain death, Condition aggravated (all occurred within 1 month of unknown dose number of BNT162b2, Lot number FG9428); medical history: colloid brain cyst; autopsy: unknown if performed.

In the remaining 39 cases (30 MC and 9 NMC) reporting the following fatal PTs Pyrexia (7), Dyspnoea (6), Myocarditis (5), Cardiac arrest, Headache (4 each), Asthenia, Seizure, Shock, Vomiting (3 each), Cardiac failure, Cardiac infection, Cardiomegaly, Depressed level of consciousness, Diarrhoea, Dizziness, Hypoesthesia, Multiple organ dysfunction syndrome, Myocardial infarction, Myocardial injury, Pneumonia, Toxic cardiomyopathy (2 each), Abdominal pain upper, Adverse event following immunisation, Agranulocytosis, Aneurysm ruptured, Anisocoria, Anuria, Atrioventricular block, B-cell type acute leukaemia, Brain injury, Cardiogenic shock, Cerebral haemorrhage, Chest discomfort, Chills, Coma, Compartment syndrome, Completed suicide, Contusion, Cough, COVID-19, Death, Dehydration, Diabetic ketoacidosis, Enterovirus infection, Extensive swelling of vaccinated limb, Fallot's tetralogy, Gait inability, Haematemesis, Haemorrhage intracranial, Head banging, Hypertension, Immunisation, Loss of consciousness, Malaise, Meningitis meningococcal, Metabolic acidosis, Multi-organ disorder, Musculoskeletal stiffness, Nausea, Nervous system disorder, Off label use, Pain in extremity, Peripheral swelling, Pleural effusion, Pruritus, Pulse absent, Pulseless electrical activity, Rash, Rash pruritic, Renal failure, Respiratory arrest, Rhinovirus infection, Sepsis, Septic shock, Slow response to stimuli, Stress cardiomyopathy, Sudden death, Thrombosis, Unresponsive to stimuli, Vaccination failure, Vaccination site pain, and Ventricular tachycardia (1 each), no confounding factors have been identified. In 19 cases the limited information available does not allow a medically meaningful assessment, in the remaining 20 cases a causality between the vaccination and the occurrence of the fatalities cannot be ruled out, based on the temporal relationship, although no laboratory data or autopsy results provided evidence of a causal relationship.
Analysis by presence of comorbidities

- Number of subjects with reported comorbidities: 960 (3.0% of 31,927 cases, the total paediatric dataset).

- Upon review, there was no significant differences in the occurrence of the most frequently reported AEs in the subjects with comorbidities compared to the population without underlying diseases was identified.

Analysis of confounders and risk factors

- Among the 31,927 cases involving paediatric subjects, 4423 included one or more confounders that prevented a clear causality assessment: co-suspect and/or multiple concomitant drugs (1286 cases), underlying medical history and/or comorbidities (4037 cases) or predisposing factors (e.g., asthma, cardiac disorders, depression, diabetes, menstrual disorders, renal disease, respiratory disorders, seizures/epilepsy) (503 cases).

Literature

Review of the literature did not identify any significant new information regarding the use of BNT162b2 in paediatric subjects.

Conclusion

No relevant differences in the occurrence of the most frequently reported events, case seriousness and case outcomes were identified among the 3 paediatric age groups presented above. Additionally, no significant differences in the reporting proportion of the most frequently reported AEs were noted between the paediatric dataset and the non-paediatric dataset, apart from the Pts Vomiting (6.1% versus 2.0%) and Product administered to patient of inappropriate age (5.8% versus 1.3%).

No new significant safety information was identified based on the review of the cases reported in the overall paediatric population. The most frequently reported AEs\(^\text{183}\) were consistent with the known reactogenicity of the vaccine and listed in Section 4.4 Special warnings and precautions for use and/or in Section 4.8 Undesirable effects of the CDS.

16.3.5.3. Use in Pregnant/Lactating Women\(^\text{184}\)

*As part of the approval letter for the emergency use of Tozinameran - COVID-19 mRNA vaccine (nucleoside modified) - COMIRNATY®, the WHO requested the MAH to present the outcome of the cases of pregnancy observed in the clinical studies.*

\(^{183}\) For the CT cases, the analysis was focused on AEs assessed as related to BNT162b2 or blinded therapy.

\(^{184}\) Exposure *in utero* cases are included.
In the final AR of the 4th SMSR (01 March 2021 – 31 March 2021), the MAH was requested to present data according to annex 3 of the “Guideline on the exposure to medicinal products during pregnancy: need for post-authorisation data (EMEA/CHMP/313666/2005)”.

These requests are addressed within the section, providing a cumulative review of pregnancy and lactation cases originating from clinical trials along with incremental pregnancy and incremental lactation cases from CTs, and incremental pregnancy and lactation cases from PM and presenting the data according to annex 3 of the “Guideline on the exposure to medicinal products during pregnancy: need for post-authorisation data”.

Search criteria - "Selects Pregnancy cases from the data set. Pregnancy cases are identified as cases where:

- Patient Pregnant Flag is “Yes”;
- If there is a value for Pregnancy Outcome, Birth Outcome, or Congenital Anomaly;
- If Delivery Notes are available;
- If any of the valid events on the case contains one of the following:
  - SOC Pregnancy, puerperium and perinatal conditions, or
    HLT Exposures associated with pregnancy, delivery and lactation; Lactation disorders, or
    PT Exposure via body fluid.

Clinical Trial Data

Cumulative review (Pregnancy Cases)

- Number of pregnancy cases: 697 (28.7% of the total 2426 cases from the CT dataset). These 697 cases represent 669 unique pregnancies (2 cases [a mother case and a foetus/baby case] or 3 cases [a mother case and 2 foetus/baby cases for twins] were created for 28 pregnancies). Cases originated from clinical studies C4591001 (155), C4591015 (120), C4591001-OPENLABEL (91), C4591031-OPENLABEL (7), C4591031 (6), C4591020 (2), C4591017 (1), BNT162-01-OPENLABEL (1), BNT162-17 (2), and C4591006 (328) and study treatment was reported as BNT162B2 (466), blinded therapy (188), placebo (42) and BNT162C2 (1).
- Country of incidence: Japan (322), US (200), Brazil (49), Argentina (46), South Africa (44), Spain (19), UK (12), Germany (3) and Turkey (2).
- Of the 597 mother cases, 431 cases reported exposure to vaccine in utero without the occurrence of any clinical event. The frequently reported pregnancy related events (>1 occurrence) were coded to the PTs Maternal exposure before pregnancy (272), Maternal exposure during pregnancy (139), Maternal exposure timing unspecified (12), Exposure during pregnancy (6), Drug exposure before pregnancy (2).
- One hundred sixty-six (166) mother cases, 139 serious and 27 non-serious, reported additional clinical events, which occurred in the vaccinated mothers.
  - The frequently reported pregnancy related events (>1 occurrence) reported in these cases were coded to the PTs Maternal exposure during pregnancy (57), Abortion spontaneous (46), Maternal exposure before pregnancy (30), Pre-eclampsia (7), Cephalo-pelvic disproportion (6), Abortion missed, Foetal death, Postpartum haemorrhage, Premature separation of placenta (4 each), Abortion threatened, Delivery, Ectopic pregnancy, Gestational hypertension, Premature delivery, Premature labour (3 each), Abortion incomplete, Hyperemesis gravidarum, Maternal exposure via partner during pregnancy, Miscarriage of partner, Uterine disorder (2 each).
  - Other reported clinical events were coded to the PTs COVID-19 (9), Anaemia (2), Abdominal wall haematoma, Cholelithiasis, Dehydration, Drug eruption, Endometritis, Lower respiratory tract infection, Osteoarthritis, Pneumonia, Pruritis, Pyelonephritis, Urinary tract infection, Urinary tract procedural complication, Vascular pseudoaneurysm, Venous thrombosis limb (1 each).
  - Of the 58 cases reporting spontaneous abortion or abortion related events, in 25 cases the mother had a medical history of spontaneous abortion, alcohol/tobacco use during pregnancy, ectopic pregnancy, obesity, diabetes mellitus, uterine disorder, depression or anembryonic gestation, which might have contributed to the event and in 33 cases there was limited information regarding the mother’s obstetric history, which precluded meaningful assessment.
  - Of the 19 cases reporting elective termination, in 10 cases, the mother had a medical history of spontaneous abortion, induced abortion, alcohol/tobacco use and in the remaining 9 cases there was limited information regarding mother’s obstetric history which precluded meaningful assessment.
  - In 3 cases reporting foetal death/stillbirth the mother had a medical history of amniotic cavity infection, HIV infection and/or spontaneous abortion, which might have contributed to the event.
  - In 3 cases reporting ectopic pregnancy, in 1 case, the mother had a medical history of tobacco use which might have contributed to the event, and in the remaining 2 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful assessment.

- Hundred (100) baby/foetal cases, 98 serious and 2 non-serious. Cases are classified according to pregnancy outcome.
  - Pregnancy outcome: Live birth with congenital anomaly: Thirty-one (31) of these cases reported 39 congenital anomalies that were coded to the PTs Atrial septal defect (4), Ankyloglossia congenital, Hypoxic-ischaemic encephalopathy, Neonatal hypotension, Trisomy 21 (2 each), Cleft lip, Coma neonatal, Congenital rubella syndrome, Congenital skin dimples, Congenital skin disorder, Craniostenosis, DiGeorge's syndrome, Gnathoschisis, Microcephaly, Neonatal pneumothorax, Neonatal intestinal perforation, Neonatal seizure, Nervous system disorder, Newborn persistent pulmonary hypertension, Osteochondrodysplasia, Patent ductus arteriosus,
Polydactyly, Pyelonephritis acute, Renal failure neonatal, Renal tubular necrosis, Sepsis neonatal, Sex chromosome abnormality, Syndactyly, Thanatophoric dwarfism, Thrombocytopenia neonatal, Ventricular septal defect, Vesicoureteric reflux (1 each). Of these 31 cases, information regarding trimester of exposure was available in 17 cases. Of these 17 cases, in 12 cases foetus was exposed during the 3rd trimester, in 4 cases foetus was exposed during the 2nd trimester, and in 1 case exposure occurred during the 1st trimester. Of these 31 cases, in 5 cases the mother of the baby was on multiple concomitant medications, alcohol use, advanced age of the mother (i.e., 43 years) and/or had a medical history of in vitro fertilization which increases the chance of gene mutation. In the remaining 26 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

- Pregnancy outcome: Still birth without foetal defect: During the reporting period there was 1 case reporting stillbirth without foetal defect. The event reported in this case was coded to the PT Neonatal respiratory distress syndrome. The information regarding trimester of exposure was unknown. In this case the mother of the baby had underlying medical history of amniotic cavity infection, which might have led to the development of the reported event.

- Pregnancy outcome: Live birth without congenital anomaly: Sixty-eight (68) cases reported live birth babies without congenital anomaly. Of these 68 cases, information regarding trimester of exposure was available in 40 cases. Of these 40 cases, in 23 cases, foetus was exposed during the 3rd trimester, in 14 cases foetus was exposed during the 2nd trimester, and in 3 cases exposure occurred during the 1st trimester. The frequently reported events (>1 occurrence) in these 68 cases were coded to PTs Jaundice neonatal (11), Foetal distress syndrome (8), Premature baby (6), Neonatal pneumonia, Neonatal respiratory distress, Bronchiolitis, Neonatal respiratory distress syndrome, Hyperbilirubinaemia neonatal (3 each), Foetal hypokinesia, Neonatal tachypnoea, Dehydration, Gastroenteritis, Patent ductus arteriosus, Anaemia neonatal, Sepsis neonatal, Hypoglycaemia neonatal, Meconium aspiration syndrome (2 each). In all these 68 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

Of the 697 cases, 658 cases provided pregnancy outcomes, which are provided in Table 67 below. Pregnancy outcome was pending or not provided in the remaining 39 cases.
<table>
<thead>
<tr>
<th>Pregnancy outcome</th>
<th>Prospective cases</th>
<th>Retrospective cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>564 (80.9%)</td>
<td>94 (13.5%)</td>
</tr>
<tr>
<td></td>
<td>Timing of exposure in pregnancy</td>
<td>Timing of exposure in pregnancy</td>
</tr>
<tr>
<td></td>
<td>Before conception</td>
<td>1st trimester</td>
</tr>
<tr>
<td>Ectopic pregnancy</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Spontaneous abortion</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>Elective termination (foetal defects)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Elective termination (no foetal defects or unknown)</td>
<td>0</td>
<td>14</td>
</tr>
<tr>
<td>Stillbirth with foetal defects</td>
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<tr>
<td>Stillbirth without foetal defects</td>
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<tr>
<td>Live birth with congenital anomaly</td>
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<tr>
<td>Live birth without congenital anomaly</td>
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<td>97</td>
</tr>
<tr>
<td>Total</td>
<td>0</td>
<td>132</td>
</tr>
</tbody>
</table>
Cumulative review (Lactation cases)

- Number of lactation cases: 141 (5.8% of the total 2426 cases from the CT dataset). All these 141 cases were non-serious. Of these 141 cases, 140 cases reported only exposure to vaccine during breastfeeding (PT Exposure via breast milk) without the occurrence of any clinical events. In the remaining case the clinical event was coded to the PT Respiratory syncytial virus infection. In this case there was limited information regarding mother’s obstetric history, which precluded meaningful causality assessment.

Incremental review (CT cases)

- Number of pregnancy cases: 41 (6.1% of the total 668 cases from the CT dataset). These 41 cases represent 37 unique pregnancies (2 cases [a mother case and a foetus/baby case] for 4 pregnancies). Cases originated from clinical studies C4591015 (24), C4591001-OPENLABEL (10), C4591001, C4591031-OPENLABEL (3 each), C4591031 (1) and study treatment was reported as blinded therapy (27), and BNT162b2 (14).

- Country of incidence: South Africa (15), Brazil (11), US (6), Argentina (5), Spain (3), UK (1).

- Twenty-three (23) serious maternal cases reported additional clinical events, which occurred in the vaccinated pregnant females.
  - The frequently reported pregnancy related events (>1 occurrence) reported in these cases were coded to the PTs Maternal exposure during pregnancy (8), Abortion spontaneous (7), Cephalo-pelvic disproportion (3), Abortion missed, Maternal exposure before pregnancy (2 each).
  - Other reported clinical events were coded to the PTs Abdominal wall haematoma, COVID-19, Pneumonia, Urinary tract infection (1 each).
  - Of the 11 cases reporting spontaneous abortion or abortion related events, in 4 cases, the mother had a medical history of spontaneous abortion or had underlying condition of obesity, which might have contributed to the event and in 7 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

- Eighteen (18) serious baby/foetal cases are classified according to pregnancy outcome.
  - Pregnancy outcome: Live birth with congenital anomaly: Five (5) of these cases reported 5 congenital anomalies that coded to the PTs Congenital rubella syndrome, DiGeorge's syndrome, Pyelonephritis, Syndactyly, Trisomy 21 (1 each). Of these 5 cases, information regarding trimester of exposure was available in 2 cases and in these 2 cases foetus was exposed during the 2nd trimester in 1 case and the 3rd trimester in the remaining case. Of these 5 cases, in 1 case reporting Trisomy 21, the age of the mother was 43 years and advanced maternal age is a risk factor for Trisomy 21. In the remaining 4 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.
  - Pregnancy outcome: Live birth without congenital anomaly: Thirteen (13) cases reported live birth babies without congenital anomaly. Of these 13 cases, information
regarding trimester of exposure was available in 7 cases. Of these 7 cases, in 5 cases, foetus was exposed during the 2nd trimester and in 2 cases foetus was exposed during the 1st and the 3rd trimester each. The frequently reported clinical events (>1 occurrence) in these 13 cases were coded to the PTs Foetal distress syndrome (3), Meconium aspiration syndrome, Gastroenteritis, Jaundice neonatal (2 each). In all these 13 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

Of the 41 cases, 38 cases provided pregnancy outcomes, which are provided in Table 68 below. Pregnancy outcome was pending or not provided in the remaining 3 cases.
Table 68. Clinical Trial Data: Pregnancy Outcome during the Reporting Interval

<table>
<thead>
<tr>
<th>Pregnancy outcome</th>
<th>Prospective cases</th>
<th>Retrospective cases</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>38 (92.7% of pregnancy cases)</td>
<td>0 (0% of pregnancy cases)</td>
<td></td>
</tr>
<tr>
<td><strong>Timing of exposure in pregnancy</strong></td>
<td><strong>Before conception</strong></td>
<td><strong>1\textsuperscript{st} trimester</strong></td>
<td><strong>After 1\textsuperscript{st} trimester</strong></td>
</tr>
<tr>
<td>Ectopic pregnancy</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Spontaneous abortion</td>
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</tr>
<tr>
<td>Elective termination (foetal defects)</td>
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<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Elective termination (no foetal defects or unknown)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Stillbirth with foetal defects</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Stillbirth without foetal defects</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Live birth with congenital anomaly</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Live birth without congenital anomaly</td>
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<td>1</td>
<td>15</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>0</td>
<td>3</td>
<td>17</td>
</tr>
</tbody>
</table>

CONFIDENTIAL
Page 335
Post-Authorisation Data

Incremental review (Pregnancy cases)

- Number of pregnancy cases: 3642 (0.7% of 507,683 cases, the total PM dataset), compared to 5239 cases (0.8%) retrieved in the PSUR #2. These 3642 cases represent 3419 unique pregnancies (2 cases [a mother case and a foetus/baby case] or 3 cases [a mother case and 2 foetuses/baby cases for twins] were created for 223 pregnancies).

- Country of incidence (>100 occurrences): Germany (837), UK (475), Netherlands (461), Philippines (309), France (302), Sweden (162), Australia (110).

- Of the 3320 mother cases, 535 cases reported exposure to vaccine in utero without the occurrence of any clinical event. The pregnancy exposure events were coded to the PTs Maternal exposure during pregnancy (355), Maternal exposure timing unspecified (116), Maternal exposure before pregnancy (52), Exposure during pregnancy (7), Drug exposure before pregnancy (4), Foetal exposure during pregnancy (1).

- There were 2785 mother cases of which 1479 were serious and 1306 were non-serious reporting additional clinical events, which occurred in the vaccinated pregnant females. Additional pregnancy related events reported in these cases (>50 occurrences) were coded to the PTs Abortion spontaneous (566), Labour pain (151), Vaginal haemorrhage (78), Heavy menstrual bleeding (50). Other frequently reported (>100 occurrences) clinical events were coded to the PTs Headache (410), Vaccination site pain (407), Fatigue (363), Pyrexia (206), Malaise (194), Myalgia (192), Nausea (178), Chills (156), Pain in extremity (135). The distribution of clinical events that were not pregnancy related (>100 occurrences) was similar in the pregnant mothers when compared with non-pregnant women of childbearing age.

- Three hundred twenty-two (322) baby/foetal cases, 283 serious and 39 non-serious. Cases are classified according to pregnancy outcome.
  - Pregnancy outcome: Live birth with congenital anomaly: Thirty-nine (39) of these cases reported 72 congenital anomalies that were coded to the PTs Foetal malformation (4), Atrial septal defect, Congenital anomaly, Ventricular septal defect (3 each), Congenital cystic lung, Congenital hydronephrosis, Congenital skin dimples, Exomphalos, Foetal cardiac disorder, Foetal chromosome abnormality, Foetal growth restriction, Kidney malformation, Pulmonary valve stenosis congenital (2 each), Anal atresia, Ankyloglossia congenital, Arnold-Chiari malformation, Cleft lip, Cleft palate, Cloacal exstrophy, Congenital amputation, Congenital foot malformation, Congenital haematological disorder, Congenital hand malformation, Congenital heart valve disorder, Congenital musculoskeletal disorder, Congenital musculoskeletal disorder of limbs, Congenital musculoskeletal disorder of spine, Congenital oral malformation, Cryptorchism, Double outlet right ventricle, Dysmorphism, Enlarged foetal cisterna magna, Fallot's tetralogy, Foetal arrhythmia, Foetal growth abnormality, Growth retardation, Heart disease congenital, Heart valve incompetence, Hepatic cytolysis,

\[185\] Few additional events reported were coded to PTs Pre-eclampsia (20), Amniotic cavity infection (1).
Hypospadias, Meningomyelecele, Neonatal deafness, Neonatal infection, Polydactyly, Pulmonary artery stenosis congenital, Pulmonary sequestration, Renal aplasia, Renal disorder, Renal dysplasia, Renal failure, Renal fusion anomaly, Renal hypertrophy, Spina bifida, VACTERL syndrome (1 each). Of these 39 cases, information regarding trimester of exposure was available in 19 cases. Of these 19 cases, in 13 cases foetus was exposed during the 1st trimester, in 4 cases foetus was exposed during the 2nd trimester, and in 2 case exposure occurred during the 3rd trimester. Of these 39 cases, in 2 cases the mother of the baby was an asymptomatic gene carrier or had familial risk factors. In the remaining 37 cases, there was limited information regarding mother’s obstetric history, which precluded meaningful causality assessment.

- Pregnancy outcome: Spontaneous abortion: Thirty-seven (37) cases reported spontaneous abortion. Of these 37 cases, information regarding trimester of exposure was provided in 17 cases. Of these 17 cases, in 15 cases, foetus was exposed during the 1st trimester, in 2 cases foetus was exposed during the 2nd and the 3rd trimester each. The most frequently reported events (>1 occurrence) in these 37 cases other than exposure related events were coded to PTs Foetal growth restriction (18), Congenital anomaly (8), Foetal heart rate abnormal (3), Cytogenetic abnormality, Foetal vascular malperfusion (2 each). Of these 37 cases, in 4 cases mother had underlying medical history (i.e., spontaneous abortion, induced abortion and/or tobacco abuse), which might have contributed to the reported events. In the remaining 33 cases, there was limited information regarding obstetric history or co-suspect medications of the mother, which precluded meaningful causality assessment.

- Pregnancy outcome: Elective termination: Twenty-three (23) cases reported elective termination of pregnancy. Of these 23 cases, 22 cases reported elective termination due to foetal defects and 1 case reported elective termination without foetal defects or unknown. Of these 23 cases, information regarding trimester of exposure was provided in 8 cases. Of these 8 cases, in 7 cases foetus was exposed during the 1st trimester, in 1 case, foetus was exposed during the 2nd trimester. The most frequently reported events (>1 occurrence) in these 23 cases other than exposure related events were coded to the PTs Heart disease congenital (4), Foetal malformation (3), Congenital central nervous system anomaly, Abortion induced (2 each). Of these 23 cases, in 5 cases mother had underlying medical history (i.e., spontaneous abortion, and/or gestational diabetes), which might have contributed to the reasons for elective termination of foetus. In the remaining 18 cases, there was limited information regarding obstetric history or co-suspect medications of mother, which precluded meaningful assessment.

- Pregnancy outcome: Stillbirth: Twenty-one (21) cases reported foetal death/neonatal death. Of these 21 cases, 15 cases reported stillbirth with foetal defects and remaining 6 cases reported stillbirth without foetal defect. Of these 21 cases, information regarding trimester of exposure was provided in 6 cases. Of these 6 cases, in 3 cases foetus was exposed during the 1st trimester, in the remaining 3 cases, foetus was exposed during the 2nd trimester. The most frequently reported events (>1 occurrence) in these 21 cases other than exposure related events were coded to the
PTs Premature baby (7), Foetal hypokinesia (5), Foetal death, Foetal heart rate abnormal (4 each), Foetal growth restriction (3). Of these 21 cases, in 5 cases the mother had underlying medical history (i.e., spontaneous abortion, and/or obesity), which might have contributed to the reported event. In the remaining 16 cases, there was limited information regarding obstetric history or co-suspect medications of mother, which precluded meaningful causality assessment.

- Pregnancy outcome: Live birth without congenital anomaly: Two hundred two (202) cases reported live birth babies without congenital anomaly. Of these 202 cases, information regarding trimester of exposure was available in 58 cases. Of these 58 cases, in 26 cases, foetus was exposed during the 3rd trimester, in 20 cases foetus was exposed during the 2nd trimester, and in 12 cases exposure occurred during the 1st trimester. The frequently reported events (≥5 occurrence) in these 202 cases other than exposure related events were coded to PTs Premature baby (74), Foetal growth restriction (22), Foetal hypokinesia (12), Jaundice neonatal (9), Foetal heart rate abnormal, Congenital anomaly, Foetal distress syndrome (7 each), Immunisation (6), Neonatal respiratory distress syndrome, Breech presentation (5 each). Of these 202 cases, in 1 case reporting cerebral thrombosis and cerebral haemorrhage foetal the baby was delivered using vacuum extractor, which might have led to development of reported event. In the remaining 201 cases, there was limited information regarding the mother’s obstetric history, which precluded meaningful causality assessment.

Of the 3642 cases, 1898 cases provided pregnancy outcomes, which are provided in Table 69 below. Pregnancy outcome was pending or not provided in the remaining 1744 cases.
Table 69. Post-Authorisation Data: Pregnancy Outcome during the Reporting Interval

<table>
<thead>
<tr>
<th>Pregnancy outcome</th>
<th>Prospective cases</th>
<th>Retrospective cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1032 (28.3% of pregnancy cases)</td>
<td>866 (23.8% of pregnancy cases)</td>
</tr>
<tr>
<td></td>
<td>Timing of exposure in pregnancy</td>
<td>Timing of exposure in pregnancy</td>
</tr>
<tr>
<td></td>
<td>Before conception</td>
<td>1st trimester</td>
</tr>
<tr>
<td>Ectopic pregnancy</td>
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<td>Spontaneous abortion</td>
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<td>14</td>
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<td>Elective termination (no foetal defects or unknown)</td>
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<td>0</td>
</tr>
<tr>
<td>Stillbirth with foetal defects</td>
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<td>Stillbirth without foetal defects</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Live birth with congenital anomaly</td>
<td>0</td>
<td>14</td>
</tr>
<tr>
<td>Live birth without congenital anomaly</td>
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<td>100</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>0</td>
<td>130</td>
</tr>
</tbody>
</table>

a. 19 December 2021 through 18 June 2022.
**Incremental review (Lactation cases)**

- Number of lactation cases: 3771 (0.7% of 507,683 cases, the total PM dataset), compared to 2670 cases (0.4%) retrieved in the PSUR #2.
  - Breast feeding baby cases: 3119, of which:
    - Two thousand six hundred eighty-nine (2689) cases reported exposure to vaccine during breastfeeding (PT Breast feeding, Exposure via breast milk and Maternal exposure during breast feeding) without the occurrence of any clinical events.
    - Four hundred thirty (430) cases, 66 serious and 364 non-serious, reported clinical events that occurred in the infant/child exposed to vaccine via breastfeeding (PT Exposure via breast milk and Maternal exposure during breast feeding); the frequently reported clinical events (>10 occurrences) were coded to the PTs Pyrexia (76), Diarrhoea (56), Crying (38), Poor feeding infant (34), Immunisation (33), Fatigue (26), Somnolence (25), Infant irritability (22), Rash (21), Vomiting (19), Irritability (16), Abdominal pain, Malaise (15 each), Rhinorrhea (14), Restlessness (13), Body temperature increased (12), Faeces discoloured, Insomnia (11 each).
  - Breast feeding mother cases: 652, of which:
    - Sixty-nine (69) mother cases who were breast feeding their baby while taking the vaccine were reported without the occurrence of any clinical events.
    - Five hundred eighty-three (583) cases, 75 serious and 508 non-serious, reported clinical events in mothers who were breast feeding; the frequently reported clinical events (>20 occurrences) were coded to the PTs Headache (93), Fatigue (86), Pyrexia (68), Vaccination site pain (60), Myalgia (54), Malaise (46), Immunisation (44), Chills (42), Pain in extremity (34), Nausea (31), Lymphadenopathy (30), Arthralgia (24), Dizziness, Influenza like illness, Interchange of vaccine products, Pain (21 each).

**Literature**

During the reporting period an article including new significant information regarding the use of BNT162b2 in pregnant/lactating women was identified. Please refer to Section 11 Literature for details.

**Conclusion**

There were no safety signals regarding use in pregnant/lactating women that emerged from the review of these cases or the medical literature.

**16.3.5.4. Use in Patients with Comorbidities**

Search criteria for immunocompromised patients: Patients with Medical history PTs included in SMQ Malignancy related conditions (Narrow and Broad Scope); SMQ Malignancy related therapeutic and diagnostic procedures (Narrow and Broad Scope); SMQ Malignant or unspecified tumours (Narrow and Broad Scope); HLGT (Primary Path): Immunodeficiency syndromes; HLT (Primary Path): Retroviral infections; PTs: Allogenic bone marrow
transplantation therapy, Allogenic stem cell transplantation, Autologous bone marrow transplantation therapy, Autologous haematopoietic stem cell transplant, Bone marrow transplant, Cord blood transplant therapy, Heart transplant, Liver transplant, Lung transplant, Pancreas islet cell transplant, Renal transplant, Small intestine transplant, Stem cell transplant.

Search criteria for patients with autoimmune or inflammatory disorders: Patients with Medical history PTs included in SMQ Immune-mediated/autoimmune disorders (Narrow and Broad scope); HLGTs (Primary Path) Autoimmune disorders; Immune disorders NEC; HLT (Primary Path) Neuromuscular junction dysfunction.

Search criteria for frail patients with comorbidities (e.g., COPD, Diabetes, Chronic Neurological Disease, Cardiovascular Disorders, active Tuberculosis): Patients with Medical history of PTs included in HLGTs (Primary Path) Bronchial disorders (excl neoplasms), Heart failures, Mental impairment disorders, Movement disorders (incl parkinsonism), Nephropathies, Pulmonary vascular disorders, Renal disorders (excl nephropathies); HLTs (Primary Path) Autonomic nervous system disorders, Diabetes mellitus (incl subtypes), Hepatic failure and associated disorders, Hepatic fibrosis and cirrhosis, Motor neurone diseases, Muscle tone abnormal, Neuromuscular disorders NEC, Pulmonary hypertensions, Respiratory failures (excl. neonatal); patients with Medical history of ongoing Tuberculosis.

Clinical Trial Data

- Number of cases: 259 (blinded therapy [36], BNT162b2, BNT162b2S01 [219, 1] and placebo [3]) (38.3% of 668 cases, the total CT dataset), compared to 286 cases (39.7%) retrieved in the PSUR #2.

- Country of incidence: US (188), Argentina (29), Germany (19), Brazil (13), China, South Africa, Spain (2 each), Dominican Republic, India, Israel, and Turkey (1 each).

- Subjects' gender: female (112), and male (147).

- Subjects' age (n = 259), range: 10 months – 87 years, mean: 57 years, median: 63 years.

- Medical history (n = 259): the most frequently (> 20 occurrences) reported medical conditions included Hypertension (117), Type 2 diabetes mellitus (71), Hypothyroidism (43), Obesity (42), Gastrooesophageal reflux disease, Osteoarthritis (41 each), Seasonal allergy (39), Anxiety, Depression (36 each), Hypercholesterolaemia (34), Asthma (33), Hyperlipidaemia (32), Insomnia (24), and Chronic obstructive pulmonary disease (23).


- Co-suspects (n = 7 cases): amlodipine, metformin (2 each), amiodarone, aripiprazole, atenolol, diltiazem, duloxetine, furosemide, hydrochlorothiazide/triamterene, hydroxyzine, losartan, metoprolol, semaglutide, tamsulosin, and warfarin (1 each).

- Number of relevant events: 341.

- Most frequently reported relevant PTs (≥4): Condition aggravated (13), Atrial fibrillation, Cerebrovascular accident, Pneumonia (8 each), Acute kidney injury, Dyspnoea,
Gastroenteritis, Osteoarthritis (5 each), Acute respiratory failure, Chest pain, Coronary artery disease, Pancreatic carcinoma, and Pulmonary embolism (4 each).

- Event outcome: fatal (18), resolved/resolving (251), resolved with sequelae (16), not resolved (55), and unknown (1).

Post-Authorisation Data

- Number of cases: 38,528 (7.6 % of 507,683 cases, the total PM dataset), compared to 66,813 cases (10.2%) retrieved in the PSUR #2.
- MC cases (13,011), NMC cases (25,517).
- Country of incidence (≥ 212 occurrences): France (7282), Germany (6733), UK (5557), US (3168), Sweden (1948), Italy (1560), Japan (1146), Austria (1066), Norway (1030), Netherlands (969), Spain (946), Denmark (670), Canada (603), Finland (543), Belgium (539), Czech Republic (460), Greece (393), Ireland (366), Estonia (275), Iraq, Portugal (272), Switzerland (260), Taiwan (246), Croatia (239), and Brazil (212).
- Subjects’ gender: female (26,999), male (10,838) and unknown (691).
- Subjects’ age in years (n = 36,076), range: 3 – 107 years, mean: 52.8 years, median: 53.0 years.
- Medical history (n = 38,528): the most frequently (≥1100 occurrences) reported medical conditions included: Asthma (7896), Hypertension (6254), Hypothyroidism (3732), Diabetes mellitus (3121), Seasonal allergy (2067), Type 2 diabetes mellitus (2001), Drug hypersensitivity (1971), Autoimmune thyroiditis (1752), Immunodeficiency (1647), Hypersensitivity (1633), Rheumatoid arthritis (1408), Food allergy (1239), Chronic obstructive pulmonary disease (1201), and Breast cancer (1121).
- Co-suspects (n = 976 cases): the most frequently (>10 occurrences) reported co-suspect vaccines/medications included: COVID-19 vaccine (471), COVID-19 Moderna vaccine (326), COVID-19 AstraZeneca vaccine (263), Influenza vaccine (107), adalimumab (90), ocrelizumab (62), quadrivalent influenza vaccine (39), COVID-19 JNC vaccine (34), levothyroixine, mycophenolate (20 each), apixaban, methotrexate (19 each), casirivimab, rituximab, tacrolimus (13 each), ibritinib (12), and pneumococcal vaccine (11).
- Number of events: 38,528.
- Event seriousness: serious (82, 607), non-serious (78,808).
- Most frequently reported relevant PTs (>10%): Headache (5518), Fatigue (5412), Pyrexia (3989), Off label use (3951).
- Reported event outcome: fatal (1081), resolved/resolving (19,899), resolved with sequelae (1540), not resolved (15,122), unknown (20,250).
Conclusion

The reporting proportion of not resolved cases (38.8%), cases resolved with sequelae (3.1%), and fatal cases (2.8%) in subjects with comorbidities is slightly higher than the reporting proportion observed in the overall population (31.6% for outcome of not resolved, 1.8% for outcome of resolved with sequelae, and 0.6% for fatal outcome). This is expected, considering the presence of the underlying diseases and/or poor intercurrent conditions.

No safety signals have emerged that would be considered specific to this population. Evaluation of cases in patients with comorbidities did not reveal any significant new safety information. Surveillance will continue. Data about each individual special sub-population are summarised in Section 16.3.5.5, Section 16.3.5.6 and Section 16.3.5.7.

16.3.5.5. Use in Immunocompromised Patients

Search criteria - Patients with Medical history of PTs included in Malignancy related conditions (SMQ Narrow); Malignancy related therapeutic and diagnostic procedures (SMQ Narrow); Malignant or unspecified tumours (SMQ Narrow); HLG: Immunodeficiency syndromes (Primary Path); HLT: Retroviral infections (Primary Path); PTs: Allogenic bone marrow transplantation therapy; Allogenic stem cell transplantation; Autologous bone marrow transplantation therapy; Autologous haematopoietic stem cell transplant; Bone marrow transplant; Cord blood transplant therapy; Heart transplant; Liver transplant; Lung transplant; Pancreas islet cell transplant; Renal transplant; Small intestine transplant; Stem cell transplant.

Clinical Trial Data

- Number of cases: 110 (BNT162b2 [90], blinded therapy [18], and BNT162B2S01, placebo [1 each]) (16.5% of 668 cases, the total CT dataset), compared to 110 cases (15.3%) retrieved in the PSUR #2.
- Country of incidence: US (78), Argentina (15), Germany (9), Brazil (5), South Africa (2), and Dominican Republic (1).
- Subjects' gender: female (58), and male (52).
- Subjects’ age in years (n = 110), range: 2–85 years, mean: 56.4 years, median: 64.5 years.
- Medical history (n = 110): the most frequently (>5 occurrences) reported relevant medical conditions included Hysterectomy (14), Cholecystectomy (10), Basal cell carcinoma (8), Colon cancer, HIV infection, Prostate cancer, Tonsillectomy (7 each), Benign prostatic hyperplasia, Breast cancer, Thyroidectomy (6 each), Breast conserving surgery (5).
- Co-suspects (n = 21): The reported co-suspect agents included amiodarone, amlodipine, diltiazem, hydroxyzine, losartan, metoprolol, tamsulosin (1 each).
- Number of events: 156.
Most frequently reported clinical PTs (>2%): Condition aggravated (8), Atrial fibrillation (4), Cerebrovascular accident (4), Gastroenteritis (4), Osteoarthritis (4), Pneumonia (4), Acute kidney injury (3), Peritonitis (3), Pyrexia (3).

BNT162b2 related events coded to the PT: None of the events were assessed as related to BNT162b2 and/or blinded therapy by the Sponsor or investigator.

Time to event onset: (n = 145 events),\textsuperscript{186} range: from <24 hours to ≤540 days, median: 113 days.
- <24 hours: 1 event;
- 1 day: 2 events;
- 2-7 days: 1 event;
- 8-14 days: 1 event;
- 15-30 days: 10 events;
- 31-181 days: 113 events;
- ≥182 days: 17 events.

Duration of event: (n = 95 of 102 events with outcome of resolved/resolved with sequelae), range: <24 hours to 122 days, median: 9 days
- <24 hours: 3 events;
- 1 day: 9 events;
- 2-7 days: 31 events;
- 8-14 days: 13 events;
- 15-30 days: 17 events;
- 31-122 days: 22 events.

Reported event outcome: fatal (10), resolved/resolving (119), resolved with sequelae (6), not resolved (20), and unknown (1).

Post-Authorisation Data

Number of cases: 8815 (1.7% of 507,683 cases, the total PM dataset), compared to 14,657 cases (2.2%) retrieved in the PSUR #2.

MC cases (3474), NMC cases (5341).

Country of incidence: France (2200), UK (2070), Germany (1085), US (726), Italy (314), Sweden (312), Japan (212), Austria (192), Spain (158), Netherlands (156), Denmark

\textsuperscript{186} This number does not include 2 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.
COVID-19 mRNA vaccine (nucleoside modified)
Periodic Safety Update Report (PSUR) 3
19 December 2021 through 18 June 2022

(131), Belgium (119), Canada (116), Norway (112); the remaining 912 cases were distributed among 53 countries.

- Subjects' gender: female (5967), male (2628) and unknown (220).
- Subjects’ age in years (n = 8073), range: 5 – 100, mean: 58.2, median: 60.0.
- Medical history (n = 8815). The most frequently (≥200 occurrences) reported relevant medical conditions included Immunodeficiency (1647), Breast cancer (1121), Thyroidectomy (566), Neoplasm malignant (466), Hysterectomy (407), Chemotherapy (377), Prostate cancer (330), Radiotherapy (272), Chronic lymphocytic leukaemia (243), Neoplasm (239).
- Co-suspects (n = 608): The most frequently (≥10 cases) reported co-suspect vaccines/medications included COVID-19 vaccine NVRV AD (113), COVID-19 vaccine (101), COVID-19 vaccine mRNA (95), Influenza vaccine (23), prednisone (20), mycophenolate mofetil (18), adalimumab (16), casirivimab/Imdevimab, tacrolimus (13 each), Influenza vaccine inact split 4V, JNJ 78436735, nivolumab, ocrelizumab (10 each).
- Number of events: 38,399.
- Event seriousness\textsuperscript{42}: serious (21,926), non-serious (16,507).
- Most frequently reported clinical PTs (≥3%): Immunisation\textsuperscript{43} (1248), Interchange of vaccine products (1223), Headache (1096), Fatigue (1030), Pyrexia (827), COVID-19 (740), Pain in extremity (686), Dyspnoea (605), Arthralgia (589), Myalgia (535), Dizziness (516), Pain (510), Nausea (488), Asthenia (478), Lymphadenopathy (456), Malaise (420), Chills (401), Chest pain (389), Vaccination site pain (374), Palpitations (326), Paraesthesia (313), Vomiting (292), Condition aggravated (254), Tachycardia (246).
- Time to event onset (n = 23,969 events),\textsuperscript{187} range: from <24 hours to ≤540 days, median: 1 day.
  - <24 hours: 8672 events;
  - 1 day: 4064 events;
  - 2-7 days: 4107 events;
  - 8-14 days: 1711 events;
  - 15-30 days: 1749 events;

\textsuperscript{187} This number does not include 103 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.

CONFIDENTIAL
Page 345
- 31-181 days: 2935 events.
- ≥182 days: 731 events.

- Duration of event (n = 3184 of 6987 events with outcome of resolved/resolved with sequelae)\textsuperscript{188}, range: <24 hours to 200 days, median: 3 days.
  - <24 hours: 396 events;
  - 1 day: 569 events;
  - 2-7 days: 1129 events;
  - 8-14 days: 383 events;
  - 15-30 days: 284 events;
  - 31-181 days: 401 events.
  - ≥182 days: 22 events.

- Event outcome\textsuperscript{78}: fatal (1006), resolved/resolving (10,930), resolved with sequelae (821), not resolved (8997), unknown (16,862).

**Analysis by age group**

- CT Data: Paediatric (16), Adults (39), and Elderly (55).
  - A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM Data: Paediatric (96), Adults (4828), Elderly (3198) and Unknown (693).
  - No significant difference was observed in the reporting proportion of frequently (≥3%) reported events between adult and elderly population except for the events coded to the PTs Headache, Lymphadenopathy, Palpitations and Tachycardia.
  - A higher reporting proportion of events coded to the PT Headache was observed in the adult population (16.6% [752 cases] in adults vs 8.2% [251 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Lymphadenopathy was observed in the adult population (7.4% [334 cases] in adults vs 2.6% [78 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Palpitations was observed in the adult population (5.2% [234 cases] in adults vs 2.1% [64 cases] in elderly) compared to the elderly population. A higher reporting proportion of events coded to the PT Tachycardia was observed in the adult population (4.1% [185 cases] in adults vs 1.5% [46 cases] in elderly) compared to the elderly population.

\textsuperscript{188} This number does not include 66 events for which partial administration and/or events without a meaningful time to onset/cessation value as per reported information.
○ No comparison was made to the paediatric population considering the limited number of cases.

Conclusion

No new significant safety information was identified based on a review of these cases.

16.3.5.6. Use in Patients with Autoimmune or Inflammatory Disorders

Search criteria for patients with autoimmune or inflammatory disorders: Patients with Medical history PTs included in SMQ Immune-mediated/autoimmune disorders (Narrow and Broad scope); HLGTS (Primary Path) Autoimmune disorders; Immune disorders NEC; HLT (Primary Path) Neuromuscular junction dysfunction.

Clinical Trial Data

- Number of cases: 102 (BNT162b2 [86], blinded therapy [14], and placebo [2]) (15.3% of 668 cases, the total CT dataset), compared to 101 cases (14.0%) retrieved in the PSUR #2.
- Of the 102 cases, the most frequently reported PTs (≥3%) included: Condition aggravated (6, 5.9%) and Atrial fibrillation (4, 3.9%).
- Event outcome: fatal (10), resolved/resolving (93), resolved with sequelae (3), and not resolved (24).
- In 6 cases (reporting 10 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Acute myeloid leukaemia, Adenocarcinoma pancreas, Cardiac failure congestive, Cardio-respiratory arrest, Completed suicide, Condition aggravated, COVID-19, Death, Pneumonia, and Sudden cardiac death (1 each). Of note, limited information regarding the cause of death was provided in 1 case (PT Death). Half (3 of 6 cases) of the fatal cases involved elderly subjects. The medical history reported included hypothyroidism, (3), colitis ulcerative, diabetes mellitus, narcolepsy, neuropathy peripheral (1 each).
- BNT162b2 related events coded to the PT Dehydration (1). Time to onset of event is 1 day and the event outcome is reported as resolved. None of the events were related to blinded therapy.

Post-Authorisation Data

- Number of cases: 21,000 (4.1% of 507,683, the total PM dataset), compared to 35,514 cases (5.4%) retrieved in the PSUR #2.
- MC cases (6424), NMC cases (14,576).
- Of the 21,000 cases, the most frequently reported clinical PTs (>3%) included: Fatigue (3103, 14.8%), Headache (3082, 14.7%), Pyrexia (2207, 10.5%), Immunization (1750, 8.3%), Pain in extremity (1680, 8.0%), Arthralgia (1675, 8.0%), Interchange of vaccine products (1568, 7.5%), Myalgia (1535, 7.3%), Dizziness (1478, 7.0%), Dyspnoea (1385,
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

6.6%), Vaccination site pain (1360, 6.5%), COVID-19 (1344, 6.4%), Nausea (1308, 6.2%), Pain (1226, 5.8%), Malaise (1180, 5.6%), Chills (1174, 5.6%), Asthenia (1083, 5.2%), Chest pain (932, 4.4%), Parasthesia (929, 4.4%), Lymphadenopathy (896, 4.3%), Condition aggravated (813, 3.9%), Palpitations (794, 3.8%), Tachycardia (646, 3.1%), and Hypoesthesia (640, 3.1%).

- Event seriousness: serious (39,651), non-serious (43,889).

- Event outcome: fatal (1295), resolved/resolving (27,683), resolved with sequelae (2277), not resolved (25,409), unknown (27,206).

- In 409 cases (reporting 1295 relevant events with a fatal outcome), the reported causes of death (≥ 20 occurrences) were coded to the PTs Death (63), Immunisation (44), Cardiac arrest, COVID-19 (36 each), COVID-19 pneumonia (33), Dyspnoea (23), Cardio-respiratory arrest (22), Interchange of vaccine products, Sudden death (21 each), and Cardiac failure (20). Of note, in 84 cases, limited information regarding the cause of death was provided (PTs Death and Sudden death). Immunisation and Interchange of vaccine products are discussed in the Section 16.3.4.6 Off Label Use. Most (326 of 409 cases) of the fatal cases involved elderly subjects. The most frequently (≥10 occurrences) reported medical history included diabetes mellitus (169), hypothyroidism (53), rheumatoid arthritis (36), type 1 diabetes mellitus (20), pulmonary fibrosis (15), rheumatic disorder (13), colitis ulcerative, psoriasis, and thyroid disorder (10 each).

- The most frequently reported clinical events observed in subjects with autoimmune or inflammatory disorders were consistent with those observed in the overall population.

**Exacerbation or Flare-up**

A focused analysis on exacerbation or flare of autoimmune or inflammatory disorders was conducted using PTs of interest (i.e., condition aggravated, disease progression), rather than all events.

- Of the 1117 cases that reported PTs indicative of exacerbation or flare, 345 cases were determined to be non-contributory and were not included in the discussion for the following reasons:
  - The events referred to an exacerbation/progression of an underlying condition that was not an autoimmune or inflammatory disorder (e.g., pain, arrhythmia, elevated blood pressure/hypertension, deep vein thrombosis, renal disease, migraine, fatigue/tiredness).

Therefore, 772 cases are included in the analysis below.

**Clinical Trial Data**

- Number of cases: 1 case (BNT162b2) (0.1% of 668 cases, the total CT dataset), compared to 1 (0.1%) retrieved in the PSUR #2.
In a case from [redacted], a 6-year-old male subject experienced a worsening of the dermatomyositis (PTs Condition aggravated and Dermatomyositis) approximately 87 days after receiving the second dose of the BNT162b2. During the hospitalisation, he was treated with methylprednisolone, albendazole, hydroxychloroquine, vitamin D, without any complications and recovered from the reported events. The events were considered unrelated to BNT162b2.

Post-Authorisation Data

- Number of cases: 771 (0.2% of 507,683 cases, the total PM dataset), compared to 750 (0.1%) retrieved in the PSUR #2.
- MC cases (274), NMC cases (497).
- Country of incidence: France (185), Germany (126), UK (118), Netherlands (54), Italy (51), US (35), Austria (23); the remaining 179 cases were distributed among 34 countries.
- Subjects' gender: female (584), male (180) and unknown (7).
- Subjects' age in years (n = 736), range: 9 – 90 years, mean: 50.7 years, median: 51 years.
- Relevant medical history: the most frequently (>20 occurrences) reported medical conditions included: Autoimmune thyroiditis (79), Hypothyroidism (53), Rheumatoid arthritis (49), Psoriasis (34), Pericarditis (29), Colitis ulcerative, Diabetes mellitus, Multiple sclerosis (28 each), Autoimmune disorder, Basedow's disease (27 each), Ankylosing spondylitis, Systemic lupus erythematosus (26 each), Immune thrombocytopenia (25), Sjogren's syndrome (22), Crohn's disease (21), Arthritis, and Psoriatic arthropathy (20 each).
- COVID-19 Medical history (n = 61): COVID-19 (43), Suspected COVID-19 (20), Post-acute COVID-19 syndrome (5), and SARS-CoV-2 test positive (1).
- Co-suspect vaccines/medications: Influenza vaccine (5), COVID-19 Vaccine MRNA (MRNA) 1273) (3), Adalimumab, COVID-19 Vaccine NRVV AD (CHADOXI NCOV-19) (2 each), acyclovir, colchicine, Hepatitis B vaccine, hydroxychloroquine, ocrelizumab, and pneumococcal vaccine polysacch 23V (1 each).
- Number of events: 4633 (of which 782 were events of interest ie, exacerbation/flare AEs).
- Relevant event seriousness: 42 serious (521), non-serious (266).
- Most frequently reported relevant PTs (>2%): Condition aggravated (548), Disease recurrence (200), and Concomitant disease aggravated (22).
- Time to event onset (n = 424)
  - <24 hours: 65 events (0 of which had a fatal outcome);
  - 1 day: 67 events;

189 This number does not include 7 events for which partial administration and/or events without a meaningful time to onset/cessation value as per reported information.
- 2-7 days: 129 events
- 8-14 days: 61 events;
- 15-30 days: 42 events;
- 31-180 days: 60 events;

- Duration of relevant events (n = 41 out of 112 occurrences with outcome of resolved/resolved with sequelae)\(^{190}\), range: 1 day to 160 days, median 17 days.
  - <24 hours: 2 events;
  - 1 day: 4 events;
  - 2 - 7 days: 6 events;
  - 8-14 days: 1 events;
  - 15-31 days: 11 events;
  - 32-181 days: 17 events;

- Relevant event outcome: fatal (4), resolved/resolving (224), resolved with sequelae (18), not resolved (332), unknown (208).

In 4 cases (reporting 4 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Disease recurrence (3), and Condition aggravated (1). Three of the 4 cases involved elderly subjects. The medical history reported included arthritis, autoimmune hepatitis, Miller Fisher syndrome, and thrombotic thrombocytopenic purpura.

**Analysis by age group**

- CT: Paediatric (1).

- PM: Paediatric (19), Adults (572), Elderly (155) and Unknown (25).
  - Exacerbation and/or flare of underlying autoimmune or inflammatory disorders occurred more frequently in the adult population, which is likely due to autoimmune disorders being more common in adults and the fact that adults are the largest group of vaccinated individuals reporting adverse events.

**Conclusion**

Overall, there were 772 cases (1 CT case and 771 PM cases [0.2% of the overall dataset]) that reported exacerbation/flares in subjects with autoimmune or inflammatory disorders following administration of BNT162b2. Exacerbation and/or flare of underlying autoimmune disease or inflammatory disorders occurred more frequently in the adult population which is likely due to the fact that the largest group of vaccinated individuals reporting AEs are in this age group. Also, the autoimmune or inflammatory disorders often occur during adulthood.

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\(^{190}\) This number does not include 1 event for which partial administration and/or events without a meaningful time to onset/cessation value as per reported information.
The most frequently reported clinical events observed in subjects with autoimmune or inflammatory disorders were consistent with those observed in the overall population.

16.3.5.7. Use in Frail Patients with Comorbidities (e.g. COPD, Diabetes, Chronic Neurological Disease, Cardiovascular Disorders, active Tuberculosis)

Search criteria - Patients with Medical history of PTs included in HLGTs (Primary Path) Bronchial disorders (excl neoplasms), Heart failures, Mental impairment disorders, Movement disorders (incl parkinsonism), Nephropathies, Pulmonary vascular disorders, Renal disorders (excl nephropathies); HLTS (Primary Path) Autonomic nervous system disorders, Diabetes mellitus (incl subtypes), Hepatic failure and associated disorders, Hepatic fibrosis and cirrhosis, Motor neurone diseases, Muscle tone abnormal, Neuromuscular disorders NEC, Pulmonary hypertensions, Respiratory failures (excl. neonatal); patients with Medical history of ongoing Tuberculosis.

Clinical Trial Data

- Number of cases: 153 (BNT162b2 [125], blinded therapy [25], and placebo [3]) (22.9% of 668 cases, the total CT dataset), compared to 176 cases (24.4%) retrieved in the PSUR #2.
- Country of incidence: US (123), Argentina (11), Germany (9), Brazil (3), China, Spain (2 each); the remaining 3 cases were distributed among 3 countries.
- Subjects' gender: female (56), male (97).
- Subjects' age in years (n = 153), range: 0.83 – 87 years, mean: 59.6 years, median: 64 years.
- Medical history (n = 153): the most frequently (≥5 occurrences) reported relevant medical conditions included Type 2 diabetes mellitus (71), Asthma (33), Chronic obstructive pulmonary disease (23), Diabetes mellitus (14), Cardiac failure congestive, Chronic kidney disease (10 each), Pulmonary embolism (7), and Bronchitis chronic (5).
- Co-suspects (n = 33 cases): The reported co-suspect agents included metformin (2), amiodarone, amlodipine, aripiprazole, atenolol, diltiazem, duloxetine, furosemide, hydrochlorothiazide/triamterene, hydroxyzine, losartan, semaglutide, tamsulosin, warfarin (1 each).
- Number of events: 187.
- Most frequently reported clinical PTs (>2%): Condition aggravated, Pneumonia (6 each), Cerebrovascular accident, Dyspnoea (5 each), and Coronary artery disease (4).
- BNT162b2 related events were coded to the PT: Dehydration (1). Time to onset of event is 1 day and the event outcome is reported as resolved. None of the events were related to blinded therapy.
• Time to event onset: \( n = 131 \),\(^{191}\) range: from 1 day to 178 days, median: 106 days.
  - <24 hours: 2 events (none had a fatal outcome)
  - 1 day: 1 event;
  - 2-7 days: 6 events;
  - 8-14 days: 0 events;
  - 15-30 days: 9 events;
  - 31-180 days: 113 events.

• Duration of relevant events \( n = 78 \) out of 103 occurrences with outcome of resolved/resolved with sequelae), range: 1 day to 78 days, median 5 days:
  - <24 hours: 4 events;
  - 1 day: 7 events;
  - 2-7 days: 43 events;
  - 8-14 days: 8 events;
  - 15-31 days: 10 events;
  - 32-181 days: 6 events.

• Reported event outcome: fatal (13), resolved/resolving (128), resolved with sequelae (10), not resolved (36), and unknown (0).

• In 9 cases (reporting 13 relevant events with a fatal outcome), the reported causes of death were coded to the PTs Death (2), Adenocarcinoma pancreas, Cardiac failure congestive, Cardio-respiratory arrest, Completed suicide, Condition aggravated, COVID-19, Drowning, Pneumonia, Pulmonary embolism, Respiratory failure, and Sudden cardiac death (1 each). Of note, in 2 cases, limited information regarding the cause of death was provided (PT Death). Most (5 of 9 cases) of the fatal cases involved elderly subjects. The most frequently (>1 occurrence) reported medical histories included type 2 diabetes mellitus (6) and Asthma (2).

Post-Authorisation Data

• Number of cases: 18,276 (3.6% of 507,683, the total PM dataset), compared to 33,889 cases (5.2%) retrieved in the PSUR #2.

• MC cases (6964), NMC cases (11,312).

• Country of incidence: France (3532), Germany (3124), UK (2189), US (1520), Sweden (1062), Japan (765), Italy (616), Austria (471), Norway (448), Spain (442), Denmark (408), Netherlands (396), Finland (305), Canada (260), Belgium (240), Czech Republic (234), Estonia (222), Iraq (220), Ireland (196), Greece (164), Taiwan, province of China

\(^{191}\) This number does not include 39 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.
COVID-19 mRNA vaccine (nucleoside modified) Reporting Period
Periodic Safety Update Report (PSUR) 3 19 December 2021 through 18 June 2022

(144), Portugal (143), Switzerland (136), Poland (102); the remaining 937 cases were distributed among 54 countries.

- Subject’s gender: female (11,576), male (6436), and unknown (264).
- Subject’s age in years (n = 17,342), range: 3 - 107 years, mean: 54.1 years, median: 55 years.
- Medical history (n = 18,276): the most frequently (≥75 occurrences) reported relevant medical conditions included Asthma (7896), Diabetes mellitus (3121), Type 2 diabetes mellitus (2004), Chronic obstructive pulmonary disease (1201), Type 1 diabetes mellitus (649), Cardiac failure (616), Chronic kidney disease (608), Pulmonary embolism (564), Renal failure (343), Parkinson's disease (247), Dementia (242), Hypokinesia (168), Cognitive disorder (166), Dementia Alzheimer's type (146), Bronchitis chronic (133), Renal disorder (117), Bronchiectasis (107), Asthma exercise induced (100), Cardiac failure chronic (81), Cardiac failure congestive (77), Bronchospasm, IgA nephropathy (76 each), and Hepatic cirrhosis (75).
- COVID-19 Medical history (n = 1226): COVID-19 (912), Suspected COVID-19 (268), COVID-19 pneumonia (38), Post-acute COVID-19 syndrome (36), SARS-CoV-2 test positive (13), Coronavirus infection (8), Asymptomatic COVID-19 (5), and Exposure to SARS-CoV-2 (1).
- Co-suspects (n = 929 cases): The most frequently (>5 occurrences) reported co-suspect vaccines/medications included COVID-19 vaccine (250), COVID-19 vaccine MRNA (MRNA 1273) (141), COVID-19 vaccine NRV V AD (CHADOXI NCOV-19) (129), influenza vaccine (58), influenza vaccine inact split 4V (24), ocrelizumab, prednisone (19 each), JNJ 78436735, mycophenolate mofetil (15 each), apixaban (14), influenza vaccine inact SAG 4V (13), tacrolimus (12), adalimumab (11), rituximab (9), prednisolone (8), atorvastatin, levophthyroxine, methotrexate (7 each), allopurinol, clopidogrel, influenza vaccine inact SAG 3V, and pregabalin (6 each).
- Number of events: 70,918
- Relevant event seriousness:42 serious (34,905), non-serious (36,098).
- Most frequently reported (≥3%) clinical PTs: Headache (2624, 15.0%), Fatigue (2570, 14.6%), Pyrexia (2012, 11.5%), Dyspnoea (1797, 10.2%), Immunisation (1533, 8.7%), COVID-19 (1446, 8.2%), Interchange of vaccine products (1383, 7.9%), Pain in extremity (1366, 7.8%), Dizziness (1255, 7.2%), Myalgia (1212, 6.9%), Arthralgia (1179, 6.7%), Vaccination site pain (1173, 6.7%), Nausea (1146, 6.5%), Malaise (1073, 6.1%), Asthenia (970, 5.5%), Chills (925, 5.3%), Pain (907, 5.2%), Chest pain (826, 4.7%), Palpitations (668, 3.8%), Lymphadenopathy (614, 3.5%), Paraesthesia (602, 3.4%), Cough (585, 3.3%), and Vomiting (561, 3.2%).
• Time to event onset (n = 46,814),\textsuperscript{192} range: from 1 day to 180 days, median: 2 days.
  - <24 hours: 16,088 events (334 of which had a fatal outcome);
  - 1 day: 9793 events;
  - 2-7 days: 9384 events;
  - 8-14 days: 3231 events;
  - 15-30 days: 3313 events;
  - 31-180 days: 5005 events.

• Duration of relevant events (n = 8391 out of 16,690 occurrences with outcome of resolved/resolved with sequelae)\textsuperscript{193}, range: 1 day to 181 days, median 3 days.
  - <24 hours: 1255 events;
  - 1 day: 1660 events;
  - 2 - 7 days: 3221 events;
  - 8-14 days: 760 events;
  - 15-31 days: 578 events;
  - 32-181 days: 917 events.

• Relevant event outcome\textsuperscript{78}: fatal (2258), resolved/resolving (24,735), resolved with sequelae (1867), not resolved (19,410), unknown (23,001).

• In 801 cases (reporting 2258 relevant events with a fatal outcome), the reported cause of death (≥26 occurrences) was coded to the PTs Death (144), Immunisation (92), COVID-19 (91), COVID-19 pneumonia (80), Cardiac arrest (62), Cardiac failure, Dyspnoea (50 each), Interchange of vaccine products (49), Sudden death (42), Cardio-respiratory arrest (40), Pulmonary embolism (38), Pneumonia (34), Respiratory failure (29), Pyrexia (28), and Myocardial infarction (26). Of note, in 186 cases, limited information regarding the cause of death was provided (PTs Death and Sudden death). Most (689 of 801 cases) of the fatal cases involved elderly subjects. The most frequently (≥20 occurrences) reported medical history included diabetes mellitus (169), type 2 diabetes mellitus (117), cardiac failure (113), chronic obstructive pulmonary disease (95), dementia (83), chronic kidney disease (72), asthma (55), cognitive disorder, pulmonary embolism (39 each), renal failure (38), Parkinson's disease (35), dementia Alzheimer's type (31), Cardiac failure chronic (27), and type 1 diabetes mellitus (20).

Analysis by age group

• CT Data: Paediatric (12), Adults (67), Elderly (74)).

\textsuperscript{192} This number does not include 1005 events for which partial administration and/or event onset dates were reported or events without a meaningful time to onset value as per reported information.

\textsuperscript{193} This number does not include 221 events for which partial administration and/or events without a meaningful time to onset/cessation value as per reported information.
A meaningful comparison between the different age groups is not possible due to the low number of cases.

- PM Data: Paediatric (625), Adults (11,157), Elderly (5906) and Unknown (588).
- No significant difference was observed in the reporting proportion of frequently (≥3%) reported events between adult and elderly population except for the event coded to PT Lymphadenopathy.
- A higher reporting proportion of events coded to PT Lymphadenopathy was observed in the adult population (4.7% [520 cases] in adults vs 1.0% [59 cases] in elderly) compared to the elderly population.
- No comparison was made to the paediatric population considering the limited number of cases.

Conclusion

The reporting proportion of not resolved cases (36.1%) and cases resolved with sequelae (3.1%) in frail subjects is similar to the reporting proportion observed in the overall population (31.7% for outcome of not resolved, 1.9% for outcome of resolved with sequelae). The reporting proportion of cases reporting fatal outcome (4.4%) in frail subjects is higher than the reporting proportion of cases reporting fatal outcome in the overall population (0.6%). This is expected, considering that most of the cases reporting a fatal outcome (64.4%) among the frail subjects involved subjects over 75 years of age who, due to their advanced age and underlying comorbidities, are more likely to die than younger individuals. Underlying comorbidities are likely to be contributory to their deaths.

The vaccine has been studied in individuals with stable chronic diseases (e.g., hypertension, obesity). It has not been systematically studied in frail individuals with severe comorbidities but there is much post-authorisation data in this population as they have generally been targeted as high priority for vaccination. No safety signals have emerged that would be considered specific to this population.

16.3.5.8. Interactions with other Vaccines

Search criteria - HLT Interactions.

- Of the 146 cases, 143 cases were determined to be non-contributory and were not included in the discussion for the following reasons:
  - in 1 case, the subject did not experience an interaction, but rather the reporter was inquiring about whether or not a drug interaction could potentially occur;
  - in 32 cases, the drug possibly interacting with BNT162b2 was not specified;
  - in 3 cases, alcohol (1), or herbal (2) interaction occurred;
  - in 1 case BNT162b2 was not involved in the interaction;
  - in 106 cases (of which 58 were serious), the subjects experienced drug interactions with the following medications rather than another vaccine (≥2): adalimumab (9), mycophenolate (6), upadacitinib (5), prednisone (4), budesonide, levetiracetam (3
each), botulinum toxin type A, capecitabine, ciclosporin, clozapine, corticosteroid nos, ethinylestradiol/ levonorgestrel, hyaluronic acid, infliximab, levothyroxine, methylphenidate, ocrolizumab, and venlafaxine (2 each).

Three of the 146 cases reported an interaction with another vaccine and are discussed below.

Clinical Trial Data

There were no relevant serious clinical trial cases reported during the reporting period, as in the PSUR #2.

Post-Authorisation Data

- Number of cases: 3 (0.0006% of 507,683 cases, the total PM dataset), compared to 18 (0.003%) retrieved in the PSUR #2.
- MC case (2), NMC case (1).
- Country of incidence: Finland, France, UK (1 each).
- Subjects' gender: female (3).
- Subjects' age in years (n =2), 61 and 68 years.
- Medical history (n = 3): Ankylosing spondylitis, Dermatitis, and Polychondritis (1 each).
- COVID-19 Medical history: none.
- Co-suspect vaccines (n = 3 cases): COVID-19 Vaccine Novavax, Pneumococcal vaccine Pneumovax and COVID-19 Vaccine AstraZeneca (1 each).
- Other co-suspects (n = 1 case): Methotrexate.
- Number of events: 26 (of which 3 were events of interest).
- Relevant event seriousness: serious (1), non-serious (2).
- Relevant PTs: Drug interaction (3).
- Co-reported AEs: Interchange of vaccine, Off label use, Pyrexia (2 each), Asthenia, Blood pressure decreased, Chills, COVID-19 Immunisation, Dermatitis, Dizziness, Eating disorder, Fall, Feeling hot, Hypersensitivity, Hypoesthesia, Illness, Nausea, Vaccination site induration, Vaccination site plaque, Vaccination site pruritus, and Vomiting (1 each). Of note, the following AEs may be associated with the interactions with other vaccines: Asthenia, Chills, Fall, Feeling hot, Hypersensitivity, Vomiting, Vaccination site induration, and Vaccination site plaque, Vaccination site pruritus. The outcome of these events was unknown or not resolved.
- Time to event onset: not available in the 3 cases.
- Relevant event outcome: resolving (1), unknown (2).
Analysis by age group comorbidities and dose

No comparison between the different age groups and presence of comorbidities was performed due to the limited number of cases.

Conclusion

Among the overall 146 cases, 143 were considered not relevant, as a drug interaction did not occur in 1 case, the interacting agents was not specified in 32 cases, BNT162b2 was not involved in 1 case and in the remaining 110 cases, the interaction occurred with alcohol, herbal or medications rather than another vaccine.

There were 3 cases in the overall post-marketing dataset that involved a vaccine interaction. The most frequently co-reported event (>2 occurrences) other than off label use and interchange of vaccines PTs was Pyrexia, which is consistent with the known reactogenicity of the vaccine and listed in Section 4.8 Undesirable effects of the CDS.

There is no indication of a safety signal noted based on the review of these cases.

16.4. Characterisation of Risks

As reported in Section 16.1 Summary of Safety Concerns, on 08 July 2022 (after the DLP of this PSUR), the MAH submitted the EU RMP version 5.1 to remove the important identified risk of anaphylaxis from the list of safety concerns in accordance with CHMP recommendation received on 10 March 2022 with the positive opinion for the Type II Variation 87 (EMEA/H/C/005735/II/0087) and based on the accumulation of post-authorisation safety information.

In line with this update to the EU-RMP, the MAH proposes to remove the important identified risk of anaphylaxis from the list of safety concerns for the next PSUR reporting period, because anaphylaxis is a known risk of vaccines that is understood by HCPs who administer vaccines and patients and does not considerably impact the benefit/risk profile of the vaccine. Product labeling and standards of medical care during the vaccination procedure provide adequate risk mitigation.

16.4.1. Characterisation of Important Identified and Potential Risks

A typical medicinal product has multiple risks associated with it and individual risks vary in terms of severity, effect on individual patients, and public health impact.

What constitutes an important risk depends upon several factors including the impact on the individual subject, the seriousness of the risk and its severity, and the impact on public health. In addition, risks, which, whilst not normally serious enough to require specific warnings or precautions but which occur in a significant proportion of the treated population, affect the quality of the treated person’s life, and which could lead to serious consequences if untreated should also be considered. Risks may be related to nonclinical or clinical safety or quality issues. The intended purpose and impact of the product e.g., whether it is intended to prevent disease, to prevent particular serious outcomes due to a condition or to reduce progression of a chronic disease must also be considered when characterising risks.
The following were considered in characterising risk(s) of this product:

- frequency of risk;
- public health impact (severity and seriousness/reversibility/outcomes);
- impact on the individual patient (effect on quality of life or could lead to serious consequences if left untreated);
- risk factors (including patient factors, dose, at risk period, additive or synergistic factors);
- preventability (i.e., predictability of a risk, whether risk factors have been identified, or possibility of detection at an early stage which could mitigate seriousness);
- potential mechanism;
- evidence source(s) and strength of the evidence.

Internal and external datasets were used to populate the table below with available data. In addition to literature searches for the drug itself and its class, external data sources were consulted.

Please see Appendix 8 for the characterisation of the important identified and important potential risks of BNT162b2, consistent with Part II, Module SVII of the BNT162b2 EU RMP version 5.0 adopted on 10 March 2022.

The threshold for investigating a safety concern further will depend upon the indication, the target population, and the likely impact on public health. For example, a safety concern with a vaccine might have a lower threshold for investigation than the same issue in a medicinal product used in the palliative treatment of metastatic cancer. The risks for BNT162b2 are well managed. No special investigations to further characterise any of these risks are necessary.

Summary information from clinical trials and post-marketing sources received by the MAH through 18 June 2022 is provided in Section 16.4.1.1 and Section 16.4.1.2.

16.4.1.1. Cumulative Characterisation of Important Identified Risks

<table>
<thead>
<tr>
<th>Table 70. Cumulative Characterisation of Important Identified Risks</th>
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<tbody>
<tr>
<td><strong>Risks</strong></td>
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<td>-----------</td>
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<tr>
<td>Anaphylaxis</td>
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</table>
**Table 70. Cumulative Characterisation of Important Identified Risks**

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Related SAEs: Anaphylactoid reaction (1) with an outcome of resolved. Related case gender: female Related case age: 17 years</td>
<td>(1924), Nausea (1125), Pruritus (1029), Dizziness (931), Erythema (870), Rash (819), Cough (795), Headache (782), Urticaria (779), Tachycardia (698), Throat tightness (615), Malaise (551), Blood pressure increased (537), Pyrexia (490), Vomiting (484), Blood pressure decreased (468), Feeling abnormal (460), Oropharyngeal discomfort (436), Palpitations (423), Chest discomfort (409), Hypoesthesia (398), Paraesthesia (392), Fatigue (382), Hypersensitivity (361), Wheezing (354), Loss of consciousness (330), Hypotension (329), Chills (325), Heart rate increased (305), Pallor (301), and Swollen tongue (301). Subjects' gender: female (6720), male (1395) and unknown (368). Subjects' age in years (n = 7763), range: 5 – 104 years, mean: 44.1 years, median: 43.0 years. Age group: Paediatric (325), Adults (6543), Elderly (909) and Unknown (706). Case source: Spontaneous (8285), Literature (166), Non-interventional study (25), Solicited (7). Event seriousness: serious (8779). Event outcome: Fatal (57), Not resolved (580), Resolved with sequelae (148), Resolved/resolving (6338), Unknown data (1667).</td>
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<tr>
<td></td>
<td>Based on the cumulative CT data, no new significant safety information was identified for BNT162b2 and anaphylaxis.</td>
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**Myocarditis and Pericarditis**

- **Myocarditis**
  - No. of cases: 4 of BNT162b2
  - No. of SAEs: 4
  - The relevant PTs: Myocarditis, Myopericarditis (2 each) Related SAEs: Myopericarditis (2), Myocarditis (1).

- **Pericarditis**
  - No. of cases: 3 of BNT162b2
  - No. of SAEs: 3

Cumulatively, there were 20,256 cases of Myocarditis and Pericarditis: 12,327 cases reported myocarditis and 9896 cases reported pericarditis (in 1967 of these 20,256 cases, the subjects developed both myocarditis and pericarditis).**Myocarditis**

- No. of cases: 12327
- Relevant PTs: Myocarditis (10390), Myopericarditis (1803), Carditis
Table 70. Cumulative Characterisation of Important Identified Risks

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
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<tbody>
<tr>
<td></td>
<td>• The most common PTs: Pericarditis (3)</td>
<td>(180), Eosinophilic myocarditis (8), Hypersensitivity myocarditis (5), Autoimmune</td>
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<td></td>
<td>• Related SAEs: None.</td>
<td>myocarditis (4), Giant cell myocarditis, Immune-mediated myocarditis (2 each).</td>
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<tr>
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<td>Based on the cumulative CT data, no new significant safety information was identified</td>
<td>• Frequently reported additional PTs (≥500 occurrences): Chest pain (4226), Dyspnoea</td>
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<td></td>
<td>for BNT162b2 and myocarditis/pericarditis.</td>
<td>(2659), Fatigue (2080), Palpitations (2009), Pericarditis (1964), Pyrexia (1838),</td>
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<td></td>
<td></td>
<td>Tachycardia (1375), Chest discomfort (1262), Headache (1046), Immunisation (886), Off</td>
</tr>
<tr>
<td></td>
<td></td>
<td>label use (846), Troponin increased (788), Dizziness (714), Interchange of vaccine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>products (691), Inappropriate schedule of product administration (585), Malaise (557),</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Asthenia (511), Nausea (502).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Subjects' gender: female (4203), male (7709) and unknown (415).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Subjects' age in years (n = 11,150), range: 6 – 102 years, mean: 35 years, median:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>31 years.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Age group: Paediatric (1836), Adults (8543), Elderly (866) and Unknown (1082).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Case source: Spontaneous (12,071), Literature (217), Clinical study (27), Solicited</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(12).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Event seriousness: serious (12,394)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Event outcome: Fatal (188), Not resolved (3639), Resolved with sequelae (296),</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Resolved/resolving (4780), Unknown data (3501).</td>
</tr>
<tr>
<td>Pericarditis</td>
<td>• No. of cases: 9896.</td>
<td>• Relevant PTs: Pericarditis (9824), Pleuropericarditis (75), Pericarditis constrictive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(16), Pericarditis adhesive (1).</td>
</tr>
<tr>
<td></td>
<td>• Frequently reported additional PTs (≥2%): Chest pain (4153), Dyspnoea (2535),</td>
<td>• Frequently reported additional PTs (≥2%): Chest pain (4153), Dyspnoea (2535),</td>
</tr>
<tr>
<td></td>
<td>Myocarditis (1848), Fatigue (1714), Palpitations (1610), Pyrexia (1150), Tachycardia</td>
<td>Myocarditis (1848), Fatigue (1714), Palpitations (1610), Pyrexia (1150), Tachycardia (1102), Chest discomfort (1027), Headache (776), Pericardial effusion (714), Immunisation (671), Off label use (603), Dizziness (555), Interchange of vaccine products (522), Malaise</td>
</tr>
</tbody>
</table>
Table 70. Cumulative Characterisation of Important Identified Risks

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>(441), Myalgia (386), Nausea (379), Pain (376), Arthralgia (361), Pain in extremity (357), Astenia (340), Inappropriate schedule of product administration (337), Paraesthesia (276), Syncope (262), Chills (238), Electrocardiogram abnormal (232), Cough (228), Heart rate increased (227), Angina pectoris (213), and Lethargy (208).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Subjects’ gender: female (4619), male (5062) and unknown (215).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Subjects’ age in years (n = 9188), range: 2 – 98 years, mean: 39.5 years, median: 37.0 years.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Age group: Paediatric (685), Adults (7657), Elderly (893), and Unknown (661).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Case source: Spontaneous (9806), Literature (45), Clinical study (41), Other solicited sources (4).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Event seriousness: serious (9916).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Event outcome: Fatal (32), Not resolved (3418), Resolved with sequelae (147), Resolved/resolving (3647), Unknown data (2676).</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Based on the accumulating data from post-authorisation use of the vaccine, including the consistent findings from passive and active surveillance databases of increased occurrences of myocarditis and pericarditis following vaccination with BNT162b2, myocarditis and pericarditis have been added as ADRs in section 4.8 Undesirable effects, in Appendix A and Appendix B of the CDS v. 14.0, made effective on 26 July 2022, after the DLP.

16.4.1.2. Cumulative Characterisation of Important Potential Risks

Table 71. Cumulative Characterisation of Important Potential Risks

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vaccine-Associated Enhanced Disease (VAED), including Vaccine-Associated Enhanced Respiratory Disease (VAERD)</td>
<td>There were no cases reporting COVID-19 infection associated with one of the PTs utilized to identify potential severe or atypical cases of COVID-19.</td>
<td>• No. of cases: 3472.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Relevant PTs most frequently reported (≥2%): Drug ineffective (1952), Vaccination failure (1520), COVID-19 pneumonia (1428), Dyspnoea (1062), Diarrhoea (498),</td>
</tr>
</tbody>
</table>

CONFIDENTIAL
Page 361
Table 71. Cumulative Characterisation of Important Potential Risks

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Based on the cumulative CT data, no new significant safety information was identified for BNT162b2 and VAED/VAERD.</td>
<td>Vomiting (239), Nausea (191), Respiratory failure (184), Myocarditis (174), Abdominal pain (130), Pulmonary embolism (129), Hypoxia (123), Acute respiratory distress syndrome (115), Cardiac failure (96), Acute kidney injury, Tachypnoea (94 each).</td>
<td>• Frequently reported additional PTs (&gt;100 occurrences): COVID-19 (2132), Pyrexia (683), Cough (547), Fatigue (385), Headache (379), Asthenia (330), Suspected COVID-19 (296), Malaise (182), Chest pain (175), Myalgia (172), Oxygen saturation decreased (163), Pain (159), Dizziness (144), Chills (141), Decreased appetite (124), Anosmia (121), Arthralgia (119), Oropharyngeal pain (118), Ageusia (111), Off label use (110), Pneumonia (106), and Immunisation (103).</td>
</tr>
<tr>
<td></td>
<td>• Subjects’ gender: female (1735), male (1662) and unknown (75).</td>
<td>• Subjects’ age in years (n = 3331), range: 5 – 104 years, mean: 65.8 years, median: 71.0 years.</td>
</tr>
<tr>
<td></td>
<td>• Age group: Paediatric (50), Adults (1292), Elderly (1994) and Unknown (136).</td>
<td>• Case source: Spontaneous (3393), Literature (26), Non-interventional study (53)</td>
</tr>
<tr>
<td></td>
<td>• Relevant event seriousness: serious (7324), non-serious (1017)</td>
<td>• Relevant event outcome: Fatal (1413), Not resolved (1261), Resolved with sequelae (80), Resolved/resolving (2784), Unknown data (2815).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Based on the cumulative PM data individual review of cases, no new significant safety information was identified for BNT162b2 and the potential risk of VAED/VAERD.</td>
</tr>
</tbody>
</table>

16.4.2. Description of Missing Information

Table 72 describes missing information associated with the use of BNT162b2.
Table 72. Description of Missing Information

<table>
<thead>
<tr>
<th>Topic</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use in pregnancy and while breastfeeding</td>
<td>The safety profile of the vaccine in pregnant and/or breastfeeding women was not studied in the pivotal clinical trial and the maternal clinical trial was terminated early due to participant recruitment difficulties. Many pregnant women have chosen to be vaccinated despite the lack of clinical trial safety data. It will be important to follow these women for pregnancy and birth outcomes. The timing of vaccination in a pregnant woman and the subsequent immune response may have varying favourable or unfavourable impacts on the embryo/foetus. The clinical consequences of SARS-CoV-2 infection to the woman and foetus during pregnancy is not yet fully understood and the pregnant woman’s baseline health status may affect both the clinical course of her pregnancy and the severity of COVID-19. These factors and the extent to which the pregnant woman may be at risk of exposure to SARS-CoV-2 will influence the benefit risk considerations for use of the vaccine. Cases indicative of use in pregnancy and while breastfeeding received during the reporting interval are summarised in Section 16.3.5.3 Use in Pregnant/Lactating Women.</td>
</tr>
<tr>
<td>Use in immunocompromised patients</td>
<td>The vaccine is being studied in ongoing clinical trials of individuals with immunocompromised conditions. Cases involving use of BNT162b2 in immunocompromised patients received during the reporting interval are summarised in Section 16.3.5.5 Use in Immunocompromised Patients.</td>
</tr>
<tr>
<td>Use in frail patients with co-morbidities (e.g. chronic obstructive pulmonary disease [COPD], diabetes, chronic neurological disease, cardiovascular disorders)</td>
<td>The vaccine has been studied in individuals with stable chronic diseases (e.g., hypertension, obesity), however, it has not been studied in frail individuals with severe comorbidities that may compromise immune function due to the condition or treatment of the condition. Therefore, further safety data will be sought in this population. Cases involving use of BNT162b2 in frail patients with comorbidities (e.g., COPD, diabetes, chronic neurological disease, cardiovascular disorders) received during the reporting interval are summarised in Section 16.3.5.7 Use in Frail Patients with Comorbidities (e.g. COPD, Diabetes, Chronic Neurological Disease, Cardiovascular Disorders, active Tuberculosis).</td>
</tr>
<tr>
<td>Use in patients with autoimmune or inflammatory disorders</td>
<td>There is limited clinical trial information on the safety of the vaccine in patients with autoimmune or inflammatory disorders. Cases involving use of BNT162b2 in patients with autoimmune or inflammatory disorders received during the reporting interval are summarised in Section 16.3.5.6 Use in Patients with Autoimmune or Inflammatory Disorders.</td>
</tr>
<tr>
<td>Interaction with other vaccines</td>
<td>There are no data on interaction of BNT162b2 mRNA vaccine with other vaccines at this time. Cases involving interactions with other vaccines received during the reporting interval are summarised in Section 16.3.5.8 Interactions with other Vaccines.</td>
</tr>
<tr>
<td>Long term safety data</td>
<td>At the time of the initial submission, 2-month post dose 2 safety data were available for approximately half of the patients who have received BNT162b2 mRNA vaccine in Study C4591001. The pivotal clinical study is ongoing and ongoing non-interventional safety studies will collect longer term post-marketing safety data.</td>
</tr>
</tbody>
</table>
17. BENEFIT EVALUATION

17.1. Important Baseline Efficacy and Effectiveness Information

BNT162b2 is indicated for active immunisation to prevent COVID-19 caused by SARS-CoV-2 virus in individuals 5 years of age and older.\textsuperscript{194}

Study C4591001 is a multicenter, placebo controlled- efficacy study in participants 12 years of age and older. Randomisation was stratified by age: 12 through 15 years of age, 16 through 55 years of age, or 56 years of age and older, with a minimum of 40% of participants in the \( \geq \)56-year stratum.\textsuperscript{195} The study excluded participants who were immunocompromised and those who had previous clinical or microbiological diagnosis of COVID-19.\textsuperscript{195} Participants with pre-existing stable disease, defined as disease not requiring significant change in therapy or hospitalisation for worsening disease during the 6 weeks before enrolment,\textsuperscript{196} were included as were participants with known stable infection with HIV, HCV, or HBV.\textsuperscript{195}

Efficacy analyses were performed with confirmed COVID-19 cases accrued during blinded placebo-controlled follow-up through 13 March 2021, representing up to 6 months of follow-up after Dose 2 for participants in the efficacy population, see table below.

\textsuperscript{194} As per information reported in the CDS version 13.0 dated 10 May 2022, in effect at the end of the reporting period. Since 17 June 2022, BNT162b2 is approved in individuals 6 months of age and older, as the paediatric Tris/Sucrose presentation - maroon cap was approved in the US.

\textsuperscript{195} Ref #12 of the CDS. Global Emergency Use Authorisation Application, Section 6.2.1.2.

\textsuperscript{196} Ref #21 of the CDS. Global Emergency Use Authorisation, Section 6.2.1.1 Phase 1 First-in-Human BNT162-01.
Table 73. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Age Subgroup – Participants without Evidence of Infection and Participants with or without Evidence of Infection prior to 7 Days after Dose 2 – Evaluatable Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=28,998 Cases n1&lt;sup&gt;b&lt;/sup&gt; Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n2&lt;sup&gt;d&lt;/sup&gt;)</th>
<th>Placebo N=21,096 Cases n1&lt;sup&gt;b&lt;/sup&gt; Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n2&lt;sup&gt;d&lt;/sup&gt;)</th>
<th>Vaccine Efficacy % (95% CI)&lt;sup&gt;f&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>All participants&lt;sup&gt;f&lt;/sup&gt;</td>
<td>77 6.247 (20,712)</td>
<td>850 6.003 (20,713)</td>
<td>91.3 (89.0, 93.2)</td>
</tr>
<tr>
<td>16 through 64 years</td>
<td>70 4.859 (15,519)</td>
<td>710 4.654 (15,515)</td>
<td>90.6 (87.9, 92.7)</td>
</tr>
<tr>
<td>65 years and older</td>
<td>7 1.233 (4192)</td>
<td>124 1.202 (4226)</td>
<td>94.5 (88.3, 97.8)</td>
</tr>
<tr>
<td>65 through 74 years</td>
<td>6 0.994 (3350)</td>
<td>98 0.966 (3379)</td>
<td>94.1 (86.6, 97.9)</td>
</tr>
<tr>
<td>75 years and older</td>
<td>1 0.239 (842)</td>
<td>26 0.237 (847)</td>
<td>96.2 (76.9, 99.9)</td>
</tr>
</tbody>
</table>

First COVID-19 occurrence from 7 days after Dose 2 in participants with or without evidence of prior SARS-CoV-2 infection<sup>198</sup>

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=22,166 Cases n1&lt;sup&gt;b&lt;/sup&gt; Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n2&lt;sup&gt;d&lt;/sup&gt;)</th>
<th>Placebo N=22,320 Cases n1&lt;sup&gt;b&lt;/sup&gt; Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n2&lt;sup&gt;d&lt;/sup&gt;)</th>
<th>Vaccine Efficacy % (95% CI)&lt;sup&gt;f&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>All participants&lt;sup&gt;f&lt;/sup&gt;</td>
<td>81 6.509 (21,642)</td>
<td>873 6.274 (21,689)</td>
<td>91.1 (88.8, 93.0)</td>
</tr>
<tr>
<td>16 through 64 years</td>
<td>74 5.073 (16,218)</td>
<td>727 4.879 (16,269)</td>
<td>90.2 (87.6, 92.4)</td>
</tr>
<tr>
<td>65 years and older</td>
<td>7 1.267 (4315)</td>
<td>128 1.232 (4326)</td>
<td>94.7 (88.7, 97.9)</td>
</tr>
<tr>
<td>65 through 74 years</td>
<td>6 1.021 (3450)</td>
<td>102 0.992 (3468)</td>
<td>94.3 (87.1, 98.0)</td>
</tr>
<tr>
<td>75 years and older</td>
<td>1 0.246 (865)</td>
<td>26 0.240 (858)</td>
<td>96.2 (77.2, 99.9)</td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by Reverse Transcription-Polymerase Chain Reaction (RT-PCR) and at least 1 symptom consistent with COVID-19 (symptoms included: fever; new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).

<sup>197</sup> Ref #53 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 19. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Subgroup – Blinded Placebo-Controlled Follow-up Period – Subjects Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluatable Efficacy (7 Days) Population.

<sup>198</sup> Ref #54 of the CDS. Interim Report – 6 Month Update (13 March 2021), Supplemental Table 14.59. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Subgroup – Blinded Placebo-Controlled Follow-up Period – Subjects With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluatable Efficacy (7 Days) Population.
Table 73. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Age Subgroup – Participants without Evidence of Infection and Participants with or without Evidence of Infection prior to 7 Days after Dose 2 – Evaluative Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period

* Participants who had no evidence of past SARS-CoV-2 infection (i.e., N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.

a. N = Number of participants in the specified group.
b. n1 = Number of participants meeting the endpoint definition.
c. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from 7 days after Dose 2 to the end of the surveillance period.
d. n2 = Number of participants at risk for the endpoint.
e. Two-sided CI for vaccine efficacy is derived based on the Clopper and Pearson method adjusted to the surveillance time.
f. Included confirmed cases in participants 12 through 15 years of age; 0 in the TRADENAME group (both with and with or without evidence of prior SARS-CoV-2 infection); 16 and 18 in the placebo group (with and with or without evidence of prior SARS-CoV-2 infection, respectively).

Subgroup analyses of vaccine efficacy by risk status in participants followed up to 6 months after Dose 2 (with a cut-off date of 13 March 2021) are presented in Table 74 and Table 75.

Table 74. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Risk Status – Participants Without Evidence of Infection* Prior to 7 Days After Dose 2 – Evaluative Efficacy (7 Days) Population During the Placebo-Controlled Follow-up Period

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=20,998 Cases</th>
<th>Placebo N=21,096 Cases</th>
<th>Vaccine Efficacy % (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n1b Surveillance Time (n2b)</td>
<td>n1b Surveillance Time (n2b)</td>
<td></td>
</tr>
<tr>
<td>First COVID-19 occurrence from 7 days after Dose 2</td>
<td>77</td>
<td>6.247 (20,712)</td>
<td>850</td>
</tr>
<tr>
<td>At risk 5</td>
<td>35</td>
<td>2.797 (9167)</td>
<td>401</td>
</tr>
<tr>
<td>No</td>
<td>42</td>
<td>3.450 (11,545)</td>
<td>449</td>
</tr>
<tr>
<td>Age group (years) and risk status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 through 64 and not at risk</td>
<td>41</td>
<td>2.776 (8887)</td>
<td>385</td>
</tr>
<tr>
<td>16 through 64 and at risk</td>
<td>29</td>
<td>2.083 (6632)</td>
<td>325</td>
</tr>
<tr>
<td>65 and older and not at risk</td>
<td>1</td>
<td>0.553 (1870)</td>
<td>53</td>
</tr>
</tbody>
</table>

199 Ref #55 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 20. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Risk Status – Blinded Placebo-Controlled Follow-up Period – Subjects Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluative Efficacy (7 Days) Population.
Table 74. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, by Risk Status – Participants Without Evidence of Infection* Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population During the Placebo-Controlled Follow-up Period

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=20,998 Cases n1&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Placebo N=21,096 Cases n1&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Vaccine Efficacy % (95% CI)&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n&lt;sup&gt;2&lt;/sup&gt;)</td>
<td>Surveillance Time&lt;sup&gt;e&lt;/sup&gt; (n&lt;sup&gt;2&lt;/sup&gt;)</td>
<td></td>
</tr>
<tr>
<td>65 and older and at risk</td>
<td>0.680 (2322)</td>
<td>0.656 (2304)</td>
<td>91.8 (81.4, 97.1)</td>
</tr>
<tr>
<td>Obese&lt;sup&gt;h&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>27</td>
<td>314</td>
<td>91.6 (87.6, 94.6)</td>
</tr>
<tr>
<td></td>
<td>2.103 (6796)</td>
<td>2.050 (6875)</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>50</td>
<td>536</td>
<td>91.1 (88.1, 93.5)</td>
</tr>
<tr>
<td></td>
<td>4.143 (13,911)</td>
<td>3.952 (13,833)</td>
<td></td>
</tr>
<tr>
<td>Age group (years) and obesity status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 through 64 and not obese</td>
<td>46</td>
<td>444</td>
<td>90.1 (86.6, 92.9)</td>
</tr>
<tr>
<td></td>
<td>3.178 (10,212)</td>
<td>3.028 (10,166)</td>
<td></td>
</tr>
<tr>
<td>16 through 64 and obese</td>
<td>24</td>
<td>266</td>
<td>91.3 (86.7, 94.5)</td>
</tr>
<tr>
<td></td>
<td>1.680 (5303)</td>
<td>1.624 (5344)</td>
<td></td>
</tr>
<tr>
<td>65 and older and not obese</td>
<td>4</td>
<td>79</td>
<td>95.2 (87.1, 98.7)</td>
</tr>
<tr>
<td></td>
<td>0.829 (2821)</td>
<td>0.793 (2800)</td>
<td></td>
</tr>
<tr>
<td>65 and older and obese</td>
<td>3</td>
<td>45</td>
<td>93.2 (78.9, 98.7)</td>
</tr>
<tr>
<td></td>
<td>0.404 (1370)</td>
<td>0.410 (1426)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by RT-PCR and at least 1 symptom consistent with COVID-19 (symptoms included: fever, new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).

* Participants who had no evidence of past SARS-CoV-2 infection (i.e., N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.

a. N = Number of participants in the specified group.
b. n1 = Number of participants meeting the endpoint definition.
c. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case ascertainment is from 7 days after Dose 2 to the end of the surveillance period.
d. n2 = Number of participants at risk for the endpoint.
e. Two-sided CI for vaccine efficacy is derived based on the Clopper and Pearson method adjusted for surveillance time.
f. Included confirmed cases in participants 12 through 15 years of age: 0 in the TRADENAME group; 16 in the placebo group.
g. At risk is defined as having at least 1 of the Charlson Comorbidity Index (CCI) category or obesity (BMI ≥ 30 kg/m<sup>2</sup> or BMI ≥ 95th percentile [12 through 15 Years of age]).
h. Obese is defined as BMI ≥ 30 kg/m<sup>2</sup>. For 12 through 15 years age group, obesity is defined as a BMI at or above the 95th percentile. Refer to the CDC growth charts at https://www.cdc.gov/growthcharts/html_charts/bmiagerev.htm.
Table 75. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Risk Status – Participants with or without Evidence of Infection prior to 7 Days after Dose 2 – Evaluable Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=22,166 Cases n1b Surveillance Time (n2b)</th>
<th>Placebo N=22,320 Cases n1b Surveillance Time (n2b)</th>
<th>Vaccine Efficacy % (95% CI)α</th>
</tr>
</thead>
<tbody>
<tr>
<td>First COVID-19 occurrence from 7 days after Dose 2 †</td>
<td>81 6.509 (21,642)</td>
<td>873 6.274 (21,689)</td>
<td>91.1 (88.8, 93.0)</td>
</tr>
<tr>
<td>At risk §</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>36 2.925 (9601)</td>
<td>410 2.807 (9570)</td>
<td>91.6 (88.1, 94.2)</td>
</tr>
<tr>
<td>No</td>
<td>45 3.584 (12,041)</td>
<td>463 3.466 (12,119)</td>
<td>90.6 (87.2, 93.2)</td>
</tr>
<tr>
<td>Age group (years) and risk status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 through 64 and not at risk</td>
<td>44 2.887 (9254)</td>
<td>397 2.779 (9289)</td>
<td>89.3 (85.4, 92.4)</td>
</tr>
<tr>
<td>16 through 64 and at risk</td>
<td>30 2.186 (6964)</td>
<td>330 2.100 (6980)</td>
<td>91.3 (87.3, 94.2)</td>
</tr>
<tr>
<td>65 and older and not at risk</td>
<td>1 0.566 (1920)</td>
<td>55 0.559 (1966)</td>
<td>98.2 (89.6, 100.0)</td>
</tr>
<tr>
<td>65 and older and at risk</td>
<td>6 0.701 (2395)</td>
<td>73 0.672 (2360)</td>
<td>92.1 (82.0, 97.2)</td>
</tr>
<tr>
<td>Obese §</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>28 2.207 (7139)</td>
<td>319 2.158 (7235)</td>
<td>91.4 (87.4, 94.4)</td>
</tr>
<tr>
<td>No</td>
<td>53 4.301 (14,497)</td>
<td>554 4.114 (14,448)</td>
<td>90.8 (87.9, 93.2)</td>
</tr>
<tr>
<td>Age group (years) and obesity status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16 through 64 and not obese</td>
<td>49 3.303 (10,629)</td>
<td>458 3.158 (10,614)</td>
<td>89.8 (86.2, 92.5)</td>
</tr>
<tr>
<td>16 through 64 and obese</td>
<td>25 1.766 (5584)</td>
<td>269 1.719 (5649)</td>
<td>91.0 (86.4, 94.3)</td>
</tr>
<tr>
<td>65 and older and not obese</td>
<td>4 0.850 (2899)</td>
<td>82 0.811 (2864)</td>
<td>95.3 (87.6, 98.8)</td>
</tr>
<tr>
<td>65 and older and obese</td>
<td>3 0.417 (1415)</td>
<td>46 0.420 (1462)</td>
<td>93.4 (79.5, 98.7)</td>
</tr>
</tbody>
</table>
Table 75. Vaccine Efficacy – First COVID-19 Occurrence from 7 Days after Dose 2, by Risk Status – Participants with or without* Evidence of Infection prior to 7 Days after Dose 2 – Evaluative Efficacy (7 Days) Population during the Placebo-Controlled Follow-up Period

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>TRADENAME N=22,166 Cases n1a</th>
<th>Placebo N=22,320 Cases n1b</th>
<th>Vaccine Efficacy % (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surveillance Time (n2b)</td>
<td>Surveillance Time (n2b)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by RT-PCR and at least 1 symptom consistent with COVID-19 (symptoms included: fever; new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).

* Participants who had no evidence of past SARS-CoV-2 infection (ie, N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.

a. N = Number of participants in the specified group.
b. n1 = Number of participants meeting the endpoint definition.
c. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from 7 days after Dose 2 to the end of the surveillance period.
d. n2 = Number of participants at risk for the endpoint.
e. Two-sided CI for vaccine efficacy is derived based on the Clopper and Pearson method adjusted for surveillance time.
f. Included confirmed cases in participants 12 to 15 years of age: 0 in the TRADENAME group; 18 in the placebo group.
g. Atrisk is defined as having at least 1 of the Charlson Comorbidity Index (CMI) category or obesity (BMI ≥30 kg/m² or BMI ≥95th percentile [12 through 15 years of age]).
h. Obese is defined as BMI ≥30 kg/m². For the 12 through 15 years of age group, obesity is defined as a BMI at or above the 95th percentile. Refer to the CDC growth charts at https://www.cdc.gov/growthcharts/html_charts/hmiagerev.htm.

Efficacy against severe COVID-19

As of 13 March 2021, vaccine efficacy against severe COVID-19 is presented only for participants with or without prior SARS-CoV-2 infection (Table 76) as the COVID-19 case counts in participants without prior SARS-CoV-2 infection were the same as those in participants with or without prior SARS-CoV-2 infection in both the TRADENAME and placebo groups.
Table 76. Vaccine Efficacy – First Severe COVID-19 Occurrence in Participants with or without* prior SARS-CoV-2 Infection Based on FDA† or Centers for Disease Control and Prevention (CDC)‡ Definition after Dose 1 or from 7 Days after Dose 2 in the Placebo-Controlled Follow-up

<table>
<thead>
<tr>
<th></th>
<th>TRADENAME Cases n1* Surveillance Time (n2†)</th>
<th>Placebo Cases n1* Surveillance Time (n2†)</th>
<th>Vaccine Efficacy % (95% CI‡)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>After Dose 1 d</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>30</td>
<td>96.7 (80.3, 99.9)</td>
</tr>
<tr>
<td></td>
<td>8.439e (22,505)</td>
<td>8.288e (22,435)</td>
<td></td>
</tr>
<tr>
<td><strong>7 days after Dose 2 f</strong></td>
<td>6.522e (21,649)</td>
<td>6.404e (21,730)</td>
<td>95.3 (70.9, 99.9)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>TRADENAME Cases n1* Surveillance Time (n2†)</th>
<th>Placebo Cases n1* Surveillance Time (n2†)</th>
<th>Vaccine Efficacy % (95% CI‡)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>After Dose 1 d</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>8.427e (22,473)</td>
<td>8.269e (22,394)</td>
<td>97.8 (87.2, 99.9)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>7 days after Dose 2 f</strong></td>
<td>6.514e (21,620)</td>
<td>6.391e (21,693)</td>
<td>100 (88.0, 100.0)</td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by RT-PCR and at least 1 symptom consistent with COVID-19 (symptoms included: fever; new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).

* Participants who had no evidence of past SARS-CoV-2 infection (ie, N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.

† Severe illness from COVID-19 as defined by FDA is confirmed COVID-19 and presence of at least 1 of the following:

- Clinical signs at rest indicative of severe systemic illness (respiratory rate ≥30 breaths per minute, heart rate ≥125 beats per minute, saturation of oxygen ≤93% on room air at sea level, or ratio of arterial oxygen partial pressure to fractional inspired oxygen <300 mm Hg);

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209 Ref #57 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 25. Vaccine Efficacy – First Severe COVID-19 Occurrence From 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period – Subjects With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population.

210 Ref #58 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 26. Vaccine Efficacy – First Severe COVID-19 Occurrence After Dose 1 – Blinded Placebo-Controlled Follow-up Period – Dose 1 All-Available Efficacy Population.

211 Ref #59 of the CDS. Interim Report – 6 Month Update (13 March 2021), Supplemental Table 14.61. Vaccine Efficacy – First Severe COVID-19 Occurrence Based on CDC-Definition After Dose 1 – Blinded Placebo-Controlled Follow-up Period – Dose 1 All-Available Efficacy Population.

212 Ref #60 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 28. Vaccine Efficacy – First Severe COVID-19 Occurrence Based on CDC-Definition From 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period – Subjects With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population.

213 Ref #61 of the CDS. EUA Amendment for Pfizer-BioNTech COVID-19 Vaccine, 6-Month Follow-Up Data from Participants 16 Years of Age and Older (13 March 2021), Section 6.2.1.2.1.2 Efficacy.
Table 76. Vaccine Efficacy – First Severe COVID-19 Occurrence in Participants with or without* prior SARS-CoV-2 Infection Based on FDA† or Centers for Disease Control and Prevention (CDC)‡ Definition after Dose 1 or from 7 Days after Dose 2 in the Placebo-Controlled Follow-up

- Respiratory failure [defined as needing high-flow oxygen, noninvasive ventilation, mechanical ventilation or extracorporeal membrane oxygenation (ECMO)];
- Evidence of shock (systolic blood pressure <90 mm Hg, diastolic blood pressure <60 mm Hg, or requiring vasopressors);
- Significant acute renal, hepatic, or neurologic dysfunction;
- Admission to an Intensive Care Unit;
- Death.

† Severe illness from COVID-19 as defined by CDC is confirmed COVID-19 and presence of at least 1 of the following:204
- Hospitalisation;
- Admission to the ICU;
- Intubation or mechanical ventilation;
- Death.

a. \( n_1 \) = Number of participants meeting the endpoint definition.
b. \( n_2 \) = Number of participants at risk for the endpoint.
c. Two-side CI for vaccine efficacy is derived based on the Clopper and Pearson method adjusted to the surveillance time.
d. Efficacy assessed based on the Dose 1 all available efficacy (modified intention-to-treat) population that included all randomised participants who received at least 1 dose of study intervention.205
e. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from Dose 1 to the end of the surveillance period.
f. Efficacy assessed based on the evaluable efficacy (7 Days) population that included all eligible randomised participants who received all dose(s) of study intervention as randomised within the predefined window, have no other important protocol deviations as determined by the clinician.205
g. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from 7 days after Dose 2 to the end of the surveillance period.

Efficacy and immunogenicity in adolescents 12 to 15 years of age

Vaccine efficacy in adolescents 12 to 15 years of age is presented in Table 77.

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205 Ref #62 of the CDS. Interim Report – 6 Month Update (13 March 2021), Table 4. Analysis Populations.
### Table 77. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2, Without Evidence of Infection and With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period, Adolescents 12 to 15 Years of Age Evaluable Efficacy (7 Days) Population

<table>
<thead>
<tr>
<th>First COVID-19 occurrence from 7 days after Dose 2 in adolescents 12 to 15 years of age without evidence of prior SARS-CoV-2 infection&lt;sup&gt;206&lt;/sup&gt;</th>
<th>Placebo</th>
<th>Vaccine Efficacy % (95% CI)&lt;sup&gt;207&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TRADENAME</strong>&lt;br&gt;N&lt;sup&gt;a&lt;/sup&gt;=1005 Cases n&lt;sup&gt;b&lt;/sup&gt;</td>
<td><strong>TRADENAME</strong>&lt;br&gt;N&lt;sup&gt;a&lt;/sup&gt;=978 Cases n&lt;sup&gt;b&lt;/sup&gt;</td>
<td><strong>TRADENAME</strong>&lt;br&gt;N&lt;sup&gt;a&lt;/sup&gt;=1119 Cases n&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Surveillance Time&lt;sup&gt;c&lt;/sup&gt; (n&lt;sup&gt;d&lt;/sup&gt;)</td>
<td>Surveillance Time&lt;sup&gt;c&lt;/sup&gt; (n&lt;sup&gt;d&lt;/sup&gt;)</td>
<td>Surveillance Time&lt;sup&gt;c&lt;/sup&gt; (n&lt;sup&gt;d&lt;/sup&gt;)</td>
</tr>
<tr>
<td>Adolescents 12 to 15 Years of Age</td>
<td>0</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>0.154 (1001)</td>
<td>0.147 (972)</td>
</tr>
<tr>
<td></td>
<td>100.0 (75.3, 100.0)</td>
<td></td>
</tr>
<tr>
<td>First COVID-19 occurrence from 7 days after Dose 2 in adolescents 12 to 15 years of age with or without&lt;sup&gt;*&lt;/sup&gt; evidence of prior SARS-CoV-2 infection&lt;sup&gt;207&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>TRADENAME</strong>&lt;br&gt;N&lt;sup&gt;a&lt;/sup&gt;=1119 Cases n&lt;sup&gt;b&lt;/sup&gt;</td>
<td><strong>TRADENAME</strong>&lt;br&gt;N&lt;sup&gt;a&lt;/sup&gt;=1110 Cases n&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Vaccine Efficacy % (95% CI)&lt;sup&gt;207&lt;/sup&gt;</td>
</tr>
<tr>
<td>Surveillance Time&lt;sup&gt;c&lt;/sup&gt; (n&lt;sup&gt;d&lt;/sup&gt;)</td>
<td>Surveillance Time&lt;sup&gt;c&lt;/sup&gt; (n&lt;sup&gt;d&lt;/sup&gt;)</td>
<td></td>
</tr>
<tr>
<td>Adolescents 12 to 15 Years of Age</td>
<td>0</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>0.170 (1109)</td>
<td>0.163 (1094)</td>
</tr>
<tr>
<td></td>
<td>100.0 (78.1, 100.0)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by RT-PCR and at least 1 symptom consistent with COVID-19 (symptoms included: fever; new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).

<sup>a</sup> Number of participants in the specified group.

<sup>b</sup> n<sup>1</sup> = Number of participants meeting the endpoint definition.

<sup>c</sup> Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from 7 days after Dose 2 to the end of the surveillance period.

<sup>d</sup> n<sup>2</sup> = Number of participants at risk for the endpoint.

<sup>*</sup> Participants who had no evidence of past SARS-CoV-2 infection (ie, N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.

In C4591001 an analysis of SARS-CoV-2 neutralising titers in a randomly selected subset of participants was performed to demonstrate non-inferior immune responses comparing adolescents 12 to 15 years of age to participants 16 through 25 years of age who had no serological or virological evidence of past SARS-CoV-2 infection. The immune response to TRADENAME in adolescents 12 to 15 years of age (n = 190) was non-inferior to the

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<sup>206</sup> Ref #46 of the CDS. Module 2.7.4 Summary of Clinical Safety, MAA Type II Variation (12-15 Years) Table: Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period – Subjects 12 Through 15 Years of Age and Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population.

<sup>207</sup> Ref #47 of the CDS. Module 2.7.4 Summary of Clinical Safety, MAA Type II Variation (12-15 Years) Table: Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2 – Blinded Placebo-Controlled Follow-up Period – Subjects 12 Through 15 Years of Age and With or Without Evidence of Infection Prior to 7 Days After Dose 2 – Evaluable Efficacy (7 Days) Population.
immune response in participants 16 through 25 years of age (n = 170), based on results for SARS-CoV-2 neutralising titers at 1 month after Dose 2. The GMT ratio of the adolescents 12 to 15 years of age group to the participants 16 through 25 years of age group was 1.76, with a 2-sided 95% CI of 1.47 to 2.10, meeting the 1.5-fold non-inferiority criterion (the lower bound of the 2-sided 95% CI for the GMR >0.67) which indicates a statistically greater response in the adolescents 12 to 15 years of age than that of participants 16 through 25 years of age.  

Efficacy and immunogenicity in participants ≥16 years of age after booster dose

Neutralizing SARS-CoV-2 antibody titers and S1-binding IgG antibodies were evaluated at 6 months after Dose 2 for Study C4591001. The data noted the persistence of a robust immune response elicited by BNT162b2 30 μg vaccination in adults for up to 6 months; and also suggest, based on the modest decline in GMTs and GMCs from 1 month to 6 months after receiving Dose 2, that vaccinees may benefit from a booster dose at 6 months or thereafter. Study C4591031 was designed to assess a booster dose in this participant population.

Study C4591031 is a Phase 3 randomised, placebo-controlled, observer-blind substudy to evaluate the safety, tolerability, and efficacy of a booster dose of BNT162b2. Participants ≥16 years of age who have completed a 2-dose primary series of BNT162b2 in Study C4591001, at least 6 months prior to randomisation, were enrolled and participants were randomised at a ratio of 1:1 to receive either BNT162b2 or placebo. Randomisation was stratified by age, such that approximately 60% of participants enrolled were to be ≥16 to 55 years of age and approximately 40% of participants >55 years of age.

Considering the observation of waning effectiveness and recommendation for booster doses in some countries, per the protocol, participants could be unblinded from 24 September 2021 onwards and those randomised to receive a booster dose of placebo were offered a dose of BNT162b2 30 μg to receive a booster of active vaccine. In Section 17.1 Important Baseline Efficacy and Effectiveness Information, the information on efficacy and effectiveness of the 2-month interim analysis of study C4591031 is presented.

17.2. Newly Identified Information on Efficacy and Effectiveness

Efficacy and effectiveness from the 6-month interim analysis of study C4591031 (16 years and older participants)

Study C4591031 Substudy A evaluated BNT162b2 boosting strategies across different population of participants (e.g., age groups). In the 6-month interim report for Substudy A efficacy analysis of a single booster dose of BNT162b2 30 μg from 7 days after booster dose during the blinded placebo-controlled follow-up period was evaluated; also incidence of

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208 Ref#48 of the CDS. Module 2.7.4 Summary of Clinical Safety, MAA Type II Variation (12-15 Years) Table: Summary of Geometric Mean Ratio – NT50 – Comparison of Subjects 12 Through 15 Years of Age to Subjects 16 Through 25 Years of Age (Immunogenicity Subset) – Subjects Without Evidence of Infection up to 1 Month After Dose 2 – Dose 2 Evaluable Immunogenicity Population.
COVID-19 cases through the entire study follow-up period in participants who received BNT162b2 initially or subsequently after unblinding was analysed.

Demographics of participants in the evaluable efficacy populations without evidence of infection prior to 7 days after booster vaccination were similar in the BNT162b2 and placebo groups. This analysis population had similar demographics compared to the overall safety population, as did the evaluable efficacy population participants with or without evidence of infection prior to 7 days after booster vaccination and the all-available efficacy population.

For participants without evidence of infection prior to 7 days after booster vaccination in the evaluable efficacy population, the median duration of blinded follow-up after booster vaccination was 2.8 months as of the data cutoff date and was similar to the safety population. Of these participants originally randomized to the BNT162b2 group, the total exposure from booster vaccination to the data cutoff date was ≥6 months for most participants (99.0%).

Follow-up times after booster vaccination for participants with or without evidence of infection prior to 7 days after booster vaccination in the evaluable efficacy population were similar to the evaluable efficacy population.

After unblinding, in the all-available efficacy population there were 7 cases meeting severe criteria; all occurred after 20 December 2021, when the Omicron variant was the predominant strain, in participants who were baseline SARS-CoV-2 negative. In original BNT162b2 participants, there were 5 severe cases: 3 met the FDA definition, 1 met the CDC definition, and 1 met both definitions. In placebo participants who later received BNT162b2, there were 2 severe cases that met the FDA definition.

These results indicate that a booster dose of BNT162b2 30 µg given ≥6 months after the primary 2-dose series of BNT162b2 30 µg vaccination provided protection against COVID-19, and protection was strongest during the Delta variant wave, and sustained up to 4 months after vaccination; longer term protection against Delta variant relative to placebo cannot be estimated from this study due to unblinding and crossover of placebo control participants. For the same reason, RVE of boosted to non-boosted participants during the Omicron variant wave cannot be estimated in this study. Although the IR during Omicron wave is much higher than that of Delta wave, the IR in those participants that were ‘later’ vaccinated is lower than those participants that were ‘early’ vaccinated, which implies better protection against Omicron with recent vaccination.

**Efficacy and immunogenicity in children 5 through <12 years of age – after 2 doses**

Study C4591007 (Study 3) is a Phase 1/2/3 study comprised of an open-label vaccine dose-finding portion (Phase 1) and a multicenter, multinational, randomised, saline placebo-controlled, observer-blind efficacy portion (Phase 2/3) that has enrolled participants 5 through <12 years of age.

A descriptive efficacy analysis of Study 3 has been performed in 1968 children 5 through 11 years of age without evidence of infection prior to 7 days after Dose 2. This analysis
evaluated confirmed symptomatic COVID-19 cases accrued up to a data cut-off date of 08 October 2021.209

The descriptive vaccine efficacy results in children 5 through 11 years of age without evidence of prior SARS-CoV-2 infection are presented in Table 78. None of the cases accrued met criteria for severe COVID-19 or multisystem inflammatory syndrome in children (MIS-C). No cases of COVID-19 were observed in either the vaccine group or the placebo group in participants with evidence of prior SARS-CoV-2 infection.209

Table 78. Vaccine Efficacy – First COVID-19 Occurrence From 7 Days After Dose 2: Without Evidence of Infection Prior to 7 Days After Dose 2 – Phase 2/3 – Children 5 Through 11 Years of Age Evaluable Efficacy Population

<table>
<thead>
<tr>
<th>First COVID-19 occurrence from 7 days after Dose 2 in children 5 through 11 years of age without evidence of prior SARS-CoV-2 infection*</th>
<th>TRADENAME²</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10 mcg/dose</td>
<td></td>
</tr>
<tr>
<td></td>
<td>N=1305</td>
<td>N=663</td>
</tr>
<tr>
<td>Cases</td>
<td>n1b</td>
<td>n1b</td>
</tr>
<tr>
<td>Surveillance Time (n26)</td>
<td>3 (1273)</td>
<td>16 (637)</td>
</tr>
<tr>
<td>Vaccine Efficacy % (95% CI)</td>
<td>90.7 (67.7, 98.3)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Confirmed cases were determined by Reverse Transcription-Polymerase Chain Reaction (RT-PCR) and at least 1 symptom consistent with COVID-19 (symptoms included: fever; new or increased cough; new or increased shortness of breath; chills; new or increased muscle pain; new loss of taste or smell; sore throat; diarrhea; vomiting).
* Participants who had no evidence of past SARS-CoV-2 infection (i.e., N-binding antibody [serum] negative at Visit 1 and SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2), and had negative NAAT (nasal swab) at any unscheduled visit prior to 7 days after Dose 2 were included in the analysis.
² Pfizer-BioNTech COVID-19 Vaccine (10 mcg nucleotide).
 a. N = Number of participants in the specified group.
b. n1 = Number of participants meeting the endpoint definition.
c. Total surveillance time in 1000 person-years for the given endpoint across all participants within each group at risk for the endpoint. Time period for COVID-19 case accrual is from 7 days after Dose 2 to the end of the surveillance period.
d. n2 = Number of participants at risk for the endpoint.

In Study 3, an analysis of SARS-CoV-2 50% neutralising titers (NT50) 1 month after Dose 2 in a randomly selected subset of participants demonstrated effectiveness by immunobridging of immune responses comparing children 5 through less <12 years of age in the Phase 2/3 part of Study 3 to participants 16 through 25 years of age in the Phase 2/3 part of Study 2 who had no serological or virological evidence of past SARS-CoV-2 infection up to 1 month after Dose 2, meeting the prespecified immunobridging criteria for both the GMR and the seroresponse difference with seroresponse defined as achieving at least 4-fold rise in SARS-CoV-2 NT50 from baseline (before Dose 1).

209 Ref #82 of the CDS. Clinical Information Amendment - COVID-19 Vaccine C4591007 (5 to <12 Years) Efficacy Data in Phase 2/3 Study C4591007, October 2021.
The ratio of the SARS-CoV-2 NT50 in children 5 through <12 years of age to that of young adults 16 to 25 years of age was 1.04 (2-sided 95% CI: 0.93, 1.18), as presented in Table 79.\textsuperscript{210}

\textsuperscript{210} Ref.\#73 of the CDS Interim Report - Children 5 to <12 Years of Age: A Phase 1, Open-Label Dose-Finding Study to Evaluate Safety, Tolerability, and Immunogenicity and Phase 2/3 Placebo-Controlled, Observer-Blinded Safety, Tolerability, and Immunogenicity Study of a SARS-CoV-2 RNA Vaccine Candidate against COVID-19 in Healthy Children and Young Adults.
Table 79. Summary of Geometric Mean Ratio for 50% Neutralizing Titer – Comparison of Children 5 Through Less Than 12 Years of Age (Study 3) to Participants 16 Through 25 Years of Age (Study 2) – Participants Without* Evidence of Infection up to 1 Month After Dose 2 – Dose 2 Evaluable Immunogenicity Population

<table>
<thead>
<tr>
<th>Assay</th>
<th>Time Point</th>
<th>GMT(^a) (95% CI(^b))</th>
<th>GMT(^c) (95% CI(^c))</th>
<th>GMR(^d) (95% CI(^e))</th>
<th>Met Immunobridging Objective(^f) (Y/N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SARS-CoV-2 neutralization</td>
<td>1 month after</td>
<td>1197.6 (1106.1, 1296.6)</td>
<td>1146.5 (1045.5, 1257.2)</td>
<td>1.04 (0.93, 1.18)</td>
<td>Y</td>
</tr>
<tr>
<td>assay - NT50</td>
<td>Dose 2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(titer)(^f)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Abbreviations:** CI = confidence interval; GMR = geometric mean ratio; GMT = geometric mean titer; LLOQ = lower limit of quantitation; NAAT = nucleic acid amplification test; NT50 = 50% neutralizing titer; SARS-CoV-2 = severe acute respiratory syndrome coronavirus 2.

* Participants who had no serological or virological evidence (up to 1 month post-Dose 2 blood sample collection) of past SARS-CoV-2 infection (i.e., N-binding antibody [serum] negative at Visit 1 and 1 month after Dose 2, SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2, and negative NAAT [nasal swab] at any unscheduled visit up to 1 month after Dose 2 blood collection) and had no medical history of COVID-19 were included in the analysis.

a. \( n = \) Number of participants with valid and determinate assay results for the specified assay at the given dose/sampling time point.

b. Protocol-specified timing for blood sample collection.

c. GMTs and 2-sided 95% CIs were calculated by exponentiating the mean logarithm of the titers and the corresponding CIs (based on the Student t distribution). Assay results below the LLOQ were set to 0.5 \( \times \) LLOQ.

d. GMRs and 2-sided 95% CIs were calculated by exponentiating the mean difference of the logarithms of the titers (Group 1 [5 through <12 years of age] - Group 2 [16 through 25 years of age]) and the corresponding CI (based on the Student t distribution).

e. Immunobridging is declared if the lower bound of the 2-sided 95% CI for the GMR is greater than 0.67 and the point estimate of the GMR is \( \geq 0.8 \).

f. SARS-CoV-2 NT50 were determined using the SARS-CoV-2 mNeonGreen Virus Microneutralization Assay. The assay uses a fluorescent reporter virus derived from the USA_WA1/2020 strain and virus neutralization is read on Vero cell monolayers. The sample NT50 is defined as the reciprocal serum dilution at which 50% of the virus is neutralized.

Among participants without prior evidence of SARS-CoV-2 infection up to 1 month after Dose 2, 99.2% of children 5 through less than 12 years of age and 99.2% of participants 16 through 25 years of age had a seroresponse from before vaccination to 1 month after Dose 2. The difference in proportions of participants who had seroresponse between the 2 age groups (children - young adult) was 0.0% (2-sided 95% CI: -2.0%, 2.2%), as presented in Table 80.\(^{210}\)
Table 80. Difference in Percentages of Participants With Seroreponse – Participants Without* Evidence of Infection up to 1 Month After Dose 2 – Immunobridging Subset – Phase 2/3 – Comparison of 5 Through <12 Years of Age to Study 2 Phase 2/3 16 Through 25 Years of Age – Evaluative Immunogenicity Population

<table>
<thead>
<tr>
<th>Assay</th>
<th>Time Point&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Pfizer-BioNTech COVID-19 Vaccine</th>
<th>Study 3</th>
<th>Study 2</th>
<th>5 Through &lt;12 Years / 16 Through 25 Years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Study 3</td>
<td>Study 2</td>
<td>5 Through &lt;12 Years / 16 Through 25 Years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>10 mcg/Dose</td>
<td>30 mcg/Dose</td>
<td>5 Through &lt;12 Years / 16 Through 25 Years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5 Through &lt;12 Years</td>
<td>16 Through 25 Years</td>
<td>16 Through 25 Years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>N=264</td>
<td>N=253</td>
<td>N=253</td>
</tr>
</tbody>
</table>

| SARS-CoV-2 neutralization assay - NT50<sup>b</sup> (titer)<sup>b</sup> | 1 month after Dose 2 | 262 (99.2) (97.3, 99.9) | 251 (99.2) (97.2, 99.9) | -0.1 (95% CI: -1.2, 0.9) |

Abbreviations: LLOQ = lower limit of quantitation; NAAT = nucleic acid amplification test; N-binding = SARS-CoV-2 nucleoprotein-binding; NT50 = 50% neutralizing titer 50; SARS-CoV-2 = severe acute respiratory syndrome coronavirus 2.

Note: Seroreponse is defined as achieving a ≥4-fold rise from baseline (before Dose 1). If the baseline measurement is below the LLOQ, a postvaccination assay result ≥4 × LLOQ is considered a seroreponse.

* Participants who had no serological or virological evidence (up to 1 month post-Dose 2 blood sample collection) of past SARS-CoV-2 infection (i.e., N-binding antibody [serum] negative at Visit 1 and 1 month after Dose 2, SARS-CoV-2 not detected by NAAT [nasal swab] at Visits 1 and 2, and negative NAAT (nasal swab) at any unscheduled visit up to 1 month after Dose 2 blood collection) and had no medical history of COVID-19 were included in the analysis.

a. N = number of participants with valid and determinate assay results both before vaccination and at 1 month after Dose 2. These values are the denominators for the percentage calculations.
b. Protocol-specified timing for blood sample collection.
c. n = number of participants with seroreponse for the given assay at the given dose/sampling time point.
d. Exact 2-sided CI based on the Clopper and Pearson method.
e. Difference in proportions, expressed as a percentage (Group 1 [5 through <12 years of age] – Group 2 [16 through 25 years of age]).
f. 2-Sided CI, based on the Miettinen and Nurminen method for the difference in proportions, expressed as a percentage.
g. Immunobridging is declared if the lower bound of the 2-sided 95% CI for the difference in proportions is greater than -10.0%.
h. SARS-CoV-2 NT50 were determined using the SARS-CoV-2 mNeoGreen Virus Microneutralization Assay.

Efficacy and immunogenicity in children 5 through <12 years of age – after booster dose

Administration of a booster (third) dose of BNT162b2 10-μg to children 5 through <12 years of age in Study C4591007 elicited robust neutralizing titers against the wild-type variant of SARS-CoV-2 in an evaluable immunogenicity population of 67 children 5 to <12 years of age who were without evidence of SARS-CoV-2 infection.

Observed GMTs at 1-month post-Dose 3 were substantially increased (2720.9) compared with those at 1-month post-Dose 2 (1253.9) and prior to booster (Dose 3) vaccination (271.0).
The GMR for participants with available titers at 1-month post-Dose 3 compared to those with available titers at 1-month post-Dose 2 was 2.17 (2-sided 95% CI: 1.76, 2.68).

The observed proportion of participants who achieved seroresponse (ie, ≥4-fold rise in SARS-CoV-2 neutralizing titers from pre-Dose 1, or ≥4 × LLOQ for a pre-Dose 1 measurement <LLOQ) was high (100.0%) at 1-month post-Dose 2, waned by pre-Dose 3 (77.6%), and was increased at 1 month after Dose 3 (98.5%). The difference in seroresponse rates at 1-month post-Dose 3 compared with at 1-month post-Dose 2 was -1.5% (2-sided 95% CI: -8.0%, 2.4%).

Additionally, based on the FFRNT (a supportive assay), a third (booster) dose of BNT162b2 10-µg elicited neutralizing titers against a recombinant SARS-CoV-2 Omicron variant and recombinant wild-type (reference) strain of SARS-CoV-2 in an evaluable immunogenicity population of 29 children 5 to <12 years of age who were without evidence of SARS-CoV-2 infection.

The observed 1-month post-Dose 2, neutralizing GMTs for the Omicron variant and reference strain were 27.6 and 323.8, respectively which increased at 1-month post-Dose 3 to 614.4 and 1702.8 and, respectively, representing an increase from post-two-dose primary series to post-booster vaccination of 22-fold for Omicron and 5-fold for the reference strain.

The GMR of neutralizing titers against Omicron versus the reference strain at 1-month post-Dose 2 was 0.09 (2-sided 95% CI: 0.07, 0.10) and increased to 0.36 (2-sided 95% CI: 0.28, 0.47) at 1-month post-Dose 3, representing a fold-rise from 1-month post-Dose 2 to 1-month post-Dose 3 that was 4-times higher for the Omicron titers than for the reference strain titers obtained in the FFRNT assay.

The immune response associated with a booster (third) dose of BNT162b2 10-µg administered approximately 6 months after the second dose to children 5 to <12 years of age is expected to confer protection against COVID-19 including disease caused by Omicron. This is in the context of previously observed immunogenicity and efficacy results across pediatric, adolescent, and adult populations in the clinical development program and available real-world data, which have collectively shown that a booster (third) dose of BNT162b2 substantially increases the magnitude and breadth of neutralization and provides protection against symptomatic SARS-CoV-2 infection caused by variants including Omicron.

**Efficacy and immunogenicity in children 6 months to <5 years of age – after 3 doses**

Study C4591007 is the ongoing, randomized, placebo-controlled, Phase 1/2/3 pediatric study in healthy children from 6 months to <12 years of age. The pediatric vaccination series for children 6 months to <5 years of age was initially planned as a two-dose series given 3 weeks apart; however, based on emerging clinical and real-world data, the protocol was amended to add a third dose given at least 8 weeks after the second dose at the age-appropriate dose level.

Immunobridging (i.e., effectiveness) data were analyzed from approximately 4500 children across the 6 months to <5 years of age groups who were randomized 2:1 to receive three
doses of BNT162b2 3 \( \mu \)g or placebo with median follow-up of approximately 2 months after Dose 3 (inclusive of blinded and open-label periods).

**Immunobridging Results**

Immunobridging success criteria were met for both age groups, comparing the GMR and seroresponse for each C4591007 group who received three doses of BNT162b2 3-\( \mu \)g to adults 16 to 25 years of age in C4591001 who received two doses of BNT162b2 30-\( \mu \)g. Note, the CI lower bounds of the GMRs were \( \geq 1 \), indicating statistical significance.

- For children 2 to <5 years of age, the GMR for titers at 1-month post-Dose 3 of BNT162b2 3-\( \mu \)g compared to young adults 16 to 25 years of age at 1-month post-Dose 2 of BNT162b2 30-\( \mu \)g, all of whom were without evidence of prior SARS-CoV-2 infection, was 1.30 (2-sided 95% CI: 1.13, 1.50) and the difference in proportions who achieved seroresponse was 1.2\% (2-sided 95\% CI: -1.5\%, 4.2\%).

- For children 6 months to <2 years of age, the GMR for titers at 1-month post-Dose 3 of BNT162b2 3-\( \mu \)g compared to young adults 16 to 25 years of age at 1-month post-Dose 2 of BNT162b2 30-\( \mu \)g, all of whom were without evidence of prior SARS-CoV-2 infection, was 1.19 (2-sided 95\% CI: 1.00, 1.42) and the difference in proportions who achieved seroresponse was 1.2\% (2-sided 95\% CI: -3.4\%, 4.2\%).

**Wild-type Strain SARS-CoV-2 Neutralization**

Three doses of BNT162b2 elicited robust immune responses to wild-type SARS-CoV-2 in children who received 3-\( \mu \)g doses and in young adults who received 30-\( \mu \)g doses.

- For children 2 to <5 years of age without evidence of prior SARS-CoV-2 infection, the observed GMT before vaccination (20.7) was increased prior to Dose 3 (401.1) and then substantially increased at 1-month post-Dose 3 (1535.2). The GMFR at 1-month post-Dose 3 was 73.3 and the seroresponse rate was 100\%.

- For children 6 months to <2 years of age without evidence of prior SARS-CoV-2 infection, the observed GMT before vaccination (20.8) was increased prior to Dose 3 (317.0) and was substantially increased at 1-month post-Dose 3 (1406.5). The GMFR at 1-month post-Dose 3 was 68.4 and the seroresponse rate was 100\%.

Patterns observed for children in wild-type SARS-CoV-2 neutralization at 1-month post-Dose 3 were generally comparable to young adults 16 to 25 years of age at 1-month post-Dose 2.

**Omicron Variant SARS-CoV-2 Neutralization**

Three doses of BNT162b2 increased neutralizing titers to Omicron and Delta variants of SARS-CoV-2 in children who received 3-\( \mu \)g doses and in adults who received 30-\( \mu \)g doses.

- For children 2 to <5 years of age without evidence of prior SARS-CoV-2 infection who received three doses of BNT162b2 3-\( \mu \)g, FFRNT assay results showed neutralizing titers
against a recombinant Omicron variant increased from before Dose 3 (14.0) to 1-month post-Dose 3 (82.5). This represents a 5.9-fold increase in Omicron neutralizing titers from before Dose 3 to 1-month post-Dose 3.

- For children 6 months to <2 years of age without evidence of prior SARS-CoV-2 infection, FFRNT assay results showed neutralizing titers against a recombinant Omicron variant increased from before Dose 3 (14.3) to 1-month post-Dose 3 (127.5). This represents a 7.8-fold increase in Omicron neutralizing titers from before Dose 3 to 1-month post-Dose 3.

- Substantial increases in titers against a recombinant Delta variant and a wild-type reference strain were also observed after the second and third doses in both pediatric age groups.

Efficacy

Descriptive efficacy analyses for Phase 2/3 Study C4591007 populations of children 6 months to <2 years of age were based on symptomatic COVID-19 cases accrued from Dose 1 to a data cutoff date of 29 April 2022. These represent available data for a still-actively enrolling study.

Observed Vaccine Efficacy Across Total Population of Children 6 Months to <5 Years

VE was estimated across the total population of participants 6 months to <5 years of age randomized 2:1 to receive BNT162b2 3-μg vs placebo, which included 992 BNT162b2 recipients and 464 placebo recipients who received three doses of study intervention.

Based on COVID-19 cases confirmed from at least 7 days post-Dose 3 to the cutoff date, observed VE was 80.3% (2-sided 95% CI: 13.9%, 96.7%). Based on cases from Dose 1 onwards, observed VE was 25.5% (2-sided 95% CI: 7.7%, 39.6%).

RVE based on 2 cases reported at least 7 days after Dose 3 (original BNT162b2 group) vs 4 cases reported at least 7 days after Dose 2 (original placebo group unblinded to receive BNT162b2) during the calendar interval of 07 February 2022 to 29 April 2022 was 76.2% (2-sided 95% CI: -0.5%, 95.1%). Note that this time period corresponds to when Omicron was the dominant SARS-CoV-2 variant.

Observed Vaccine Efficacy in Each Age Group

VE was estimated for each age group based on COVID-19 cases confirmed at least 7 days post-Dose 3 or from Dose 1 to the data cutoff date. Observed VE from Dose 1 onwards, or from at least 7 days after Dose 3, was not meaningfully impacted by excluding cases involving coinfection with other respiratory pathogens.

In the 2 to <5 years of age group, VE was estimated from a population of 1835 BNT162b2 recipients and 915 placebo recipients of whom 606 and 280, respectively, received three doses. Based on cases confirmed from at least 7 days post-Dose 3 to the cutoff date, observed
VE of 82.3% (2-sided 95% CI: -8.0%, 98.3%). From Dose 1 onwards, observed VE of 32.6% (2-sided 95% CI: 10.8%, 48.8%)

In the 6 months to <2 years of age group, VE was estimated from a population of 1178 BNT162b2 recipients and 598 placebo recipients of whom 386 and 184, respectively, received three doses. Based on cases confirmed from at least 7 days post-Dose 3 to the cutoff date, observed VE of 75.5% (2-sided 95% CI: -370.1%, 99.6%). From Dose 1 onwards, observed VE of 14.0% (2-sided 95% CI: -21.2%, 38.4%).

Relative Vaccine Efficacy of Three Doses vs Two Doses in Each Age Group

For children 2 to <5 years of age, RVE during the calendar interval of 07 February 2022 to 29 April 2022 based on 2 cases reported at least 7 days after Dose 3 (original BNT162b2 group) vs 4 cases reported at least 7 days after Dose 2 (original placebo group unblinded to receive BNT162b2) was 84.0% (2-sided 95% CI: -11.8%, 98.6%).

For children 6 months to <2 years of age, RVE during the calendar interval of 07 February 2022 to 29 April 2022 based on 2 cases reported at least 7 days after Dose 3 (original BNT162b2 group) vs 2 cases reported at least 7 days after Dose 2 (original placebo group unblinded to receive BNT162b2) was 59.4% (2-sided 95% CI: -459.5%, 97.1%).

**Omicron-specific VE for the time period 01 December 2021 to 06 February 2022**

Given that the US Food and Drug Administration initially authorized a third dose of the vaccine for individuals aged 65 years and older and individuals at high risk of severe COVID-19 on 22 September 2021, early estimates of VE against Omicron are likely enriched for high-risk populations, including patients who are immunocompromised. Indeed, the analysis using data from the early portion of the Omicron wave showed early signs of waning effectiveness of the BNT162b2 mRNA COVID-19 vaccine against Omicron variant-related hospital and emergency department admission at 3 months or longer after receipt of a third dose in US adults aged 18 years and older.

Updated findings primarily show two things. First, waning effectiveness against Omicron-related hospitalisation observed at ≥3 months after a third dose of vaccine during the initial study period (data cutoff of 06 February 2022) was less pronounced after excluding individuals who were immunocompromised; original VE ≥3 months after a third dose of 55% (95% CI: 28–71) against hospitalisation vs 74% (95% CI: 52–86) after excluding individuals who were immune-compromised. Second, extending the analysis period through 18 March 2022, which captures the entire Omicron wave and results in the inclusion of more individuals who became eligible for booster doses on 29 November 2021, diminished the evidence of waning vaccine protection after a third dose. Specifically, after extending the analysis period, waning of VE against Omicron-related outcomes was no longer apparent, particularly in the immunocompetent population.

Thus, patients who were immunocompromised likely drove much of the observed waning seen in our initial report. Another explanation may be differences in severity of illness among patients admitted to the hospital or emergency department over time, which could result from
increasing levels of immunity due to natural infection and/or increased at-home COVID-19 testing during the updated study period\textsuperscript{211}

A more recent study by the same group, found that three doses of BNT162b2 conferred high protection against hospital and emergency department admission due to both the delta and omicron variants in the first 3 months after vaccination. However, 3 months after receipt of a third dose, waning was apparent against SARS-CoV-2 outcomes due to the omicron variant, including hospital admission. Additional doses of current, adapted, or novel COVID-19 vaccines might be needed to maintain high levels of protection against subsequent waves of SARS-CoV-2 caused by the omicron variant or future variants with similar escape potential. (2) \textsuperscript{212}

The UK health security agency released the COVID-19 vaccine surveillance report up to 16 June 2022, showing that BNT162b2 vaccine efficacy against symptomatic COVID-19 is lower and wanes faster for Omicron.\textsuperscript{213}

Tarof et al. also highlights vaccines have been effective against severe Omicron illness\textsuperscript{211,212} however waning against Omicron hospitalisation is observed $>$9m after the second vaccination dose and duration of protection $>$6m post-boost is unknown and could trigger higher rates of lack of efficacy, defined by breakthroughs and re-infections by the current Omicron subvariants BA.4 and BA.5.

A recent publication from Israel\textsuperscript{214} reports a low neutralisation efficiency against BA.4 and BA.5 even in sera obtained from BA.1-recovered from health care workers who previously received three or four vaccine doses. These findings suggest that an Omicron-specific vaccination might be indicated.

Hansen et al. evaluated the risk of reinfection, vaccine protection, and severity of infection with the BA.5 Omicron subvariant and they found a high protection against BA.5 from prior Omicron infection in triple-vaccinated individuals, and similar vaccine effectiveness for BA.5

\textsuperscript{211} Tarof SY, Slezak JM, Puzniak L. Effectiveness of a third dose of BNT162b2 mRNA COVID-19 vaccine in a large US health system: a retrospective cohort study. The Lancet Regional Health – Americas 2022;9: 100198 Published on line 14 February 2022.

\textsuperscript{212} Tarof SY, Slezak JM, Puzniak L. Durability of BNT162b2 vaccine against hospital and emergency department admission due to the omicron and delta variants in a large health system in the USA: a test-negative case-control study. Lancet Respir Med 2022; 10:689-99.


\textsuperscript{214} Kliker L, Zuckerman N, Atari N et al. COVID-19 vaccination and BA.1 breakthrough infection induce neutralising antibodies which are less efficient against BA.4 and BA.5 Omicron variants, Israel, March to June 2022. www.eurosurveillance.org submitted on 12 Jul 2022 / accepted on 28 Jul 2022 / published on 28 Jul 2022
infection as currently for BA.2. BA.5 infection was associated with an increased risk of hospitalisation which needs confirmation and continued surveillance as hospitalisations were low and stable during the study period.\textsuperscript{215} Adapted vaccines can help slow virus circulation and emergence of variants of concern.

Three doses of BNT162b2 conferred high protection against hospital and emergency department admission due to both the Delta and Omicron variants in the first 3 months after vaccination. However, 3 months after receipt of a third dose, waning was apparent against SARS-CoV-2 outcomes due to the omicron variant, including hospital admission.\textsuperscript{212}

Additional doses of current, adapted, or novel COVID-19 vaccines might be needed to maintain high levels of protection against subsequent waves of SARS-CoV-2 caused by the omicron variant or future variants with similar escape potential.\textsuperscript{212}

17.3. Characterisation of Benefits

Data in Section 17.1 demonstrates a high degree of efficacy against symptomatic and severe COVID-19 in non-immunocompromised people over 12 years of age, during the period at least 7 days following the second dose of vaccine. Efficacy is evident separately and at a similar level in people 12-15 years of age, 16-64 years of age, 65 to 74 years of age and 75 years of age and older. Efficacy also appears largely independent of risk factors (having at least 1 of the CMI categories) and obesity. Efficacy is also high against severe disease after the first dose. This is anticipated to deliver effective prevention of COVID-19 in the community and reduced hospitalisation, severe morbidity and death from COVID-19. Section 17.2 describes the newly identified information on efficacy and effectiveness of the 2-month and 6-month interim analysis of study C4591031 and on efficacy in children 6-month through <12 years of age. Additionally, the variation of VE against SARS-CoV-2 infection between 1 and 6 months after full vaccination and after booster dose and data concerning VE against Delta and Omicron variants are presented.

18. INTEGRATED BENEFIT-RISK ANALYSIS FOR APPROVED INDICATIONS

18.1. Benefit-Risk Context – Medical Need and Important Alternatives

BNT162b2 indications are provided in Section 1.

Incidence

COVID-19 is caused by a novel coronavirus labelled as SARS-CoV-2. The disease first emerged in December 2019, when a cluster of patients with pneumonia of unknown cause was recognised in Wuhan City, Hubei Province, China.\textsuperscript{216} The number of infected cases


rapidly increased and spread beyond China throughout the world. On 30 January 2020, the WHO declared COVID-19 a Public Health Emergency of International Concern and thus a pandemic.\textsuperscript{217}

Estimates of SARS-CoV-2 incidence change rapidly. The MAH obtained incidence and prevalence estimates using data from Worldometer, a trusted independent organisation that collects COVID-19 data from official reports and publishes current global and country-specific statistics online\textsuperscript{218}.

As of 05 April 2022, the overall number of people who had been infected with SARS-CoV-2 was over 475 million worldwide\textsuperscript{219}, an increase of 268 million since 15 August 2021.\textsuperscript{220}

Table 77 shows the incidence and prevalence as of 05 April 2022 for the US, UK, and EU-27 countries. In the EU and the UK, by 05 April 2022 the total number of confirmed cases had accumulated to over 148 million people, or 28,895 per 100,000 people (from 41 million, or 8074 per 100,000 by 15 August 2021). Across countries in the EU, the number of confirmed cases ranged from 13,859 to 50,185 cases per 100,000 people. Cyprus and Romania reported the lowest incidence rates while Netherlands, Slovenia, and Denmark reported the highest.\textsuperscript{219}

In the US, the number of confirmed cases had reached over 81 million (24,482 per 100,000 people) by 05 April 2022.\textsuperscript{219} This is an increase from 37 million (11,236 per 100,000) by 15 August 2021.\textsuperscript{220}

\begin{itemize}
\end{itemize}
Table 81. Incidence, Prevalence, and Mortality of COVID-19 as of 05 April 2022

<table>
<thead>
<tr>
<th></th>
<th>Total Cases</th>
<th>Incidence: Total Cases/100,000</th>
<th>Active Cases</th>
<th>Prevalence: Active Cases/100,000</th>
<th>Total Deaths</th>
<th>Mortality: Deaths/100,000</th>
<th>Population</th>
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<td>Global</td>
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<td>58,426,026</td>
<td>736</td>
<td>6,124,782</td>
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<td>7,938,283,964*</td>
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<td>1,052,505</td>
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<td>31,177</td>
<td>1,931,030</td>
<td>2,819</td>
<td>1,657,800</td>
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<td>15,155,839</td>
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<td>1,218,285</td>
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<td>514,219,196</td>
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<td>302</td>
<td>334,405,890</td>
</tr>
</tbody>
</table>

EU-27 Countries

<table>
<thead>
<tr>
<th>Country</th>
<th>Total Cases</th>
<th>Incidence: Total Cases/100,000</th>
<th>Active Cases</th>
<th>Prevalence: Active Cases/100,000</th>
<th>Total Deaths</th>
<th>Mortality: Deaths/100,000</th>
<th>Population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austria</td>
<td>3,912,606</td>
<td>43,012</td>
<td>252,955</td>
<td>2,781</td>
<td>16,051</td>
<td>177</td>
<td>9,096,500</td>
</tr>
<tr>
<td>Belgium</td>
<td>3,881,523</td>
<td>33,238</td>
<td>289,217</td>
<td>2,477</td>
<td>30,908</td>
<td>265</td>
<td>11,678,061</td>
</tr>
<tr>
<td>Bulgaria</td>
<td>1,141,859</td>
<td>16,653</td>
<td>173,569</td>
<td>2,531</td>
<td>36,608</td>
<td>534</td>
<td>6,856,744</td>
</tr>
<tr>
<td>Croatia</td>
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<td>27,178</td>
<td>7,321</td>
<td>180</td>
<td>15,634</td>
<td>385</td>
<td>4,060,882</td>
</tr>
<tr>
<td>Cyprus</td>
<td>444,174</td>
<td>13,859</td>
<td>318,853</td>
<td>3,617</td>
<td>951</td>
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<td>1,222,769</td>
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<tr>
<td>Denmark</td>
<td>2,924,746</td>
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<td>51,848</td>
<td>890</td>
<td>5,798</td>
<td>99</td>
<td>5,827,966</td>
</tr>
<tr>
<td>Estonia</td>
<td>560,233</td>
<td>42,183</td>
<td>96,637</td>
<td>5,243</td>
<td>2,475</td>
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<tr>
<td>Finland</td>
<td>907,786</td>
<td>16,339</td>
<td>858,532</td>
<td>15,453</td>
<td>3,254</td>
<td>59</td>
<td>5,555,812</td>
</tr>
<tr>
<td>France</td>
<td>26,025,500</td>
<td>39,717</td>
<td>2,267,053</td>
<td>3,460</td>
<td>142,655</td>
<td>218</td>
<td>65,526,762</td>
</tr>
<tr>
<td>Germany</td>
<td>21,908,379</td>
<td>26,003</td>
<td>4,369,710</td>
<td>5,186</td>
<td>130,969</td>
<td>155</td>
<td>84,253,677</td>
</tr>
<tr>
<td>Greece</td>
<td>3,096,135</td>
<td>29,961</td>
<td>257,121</td>
<td>2,488</td>
<td>27,746</td>
<td>268</td>
<td>10,333,792</td>
</tr>
<tr>
<td>Hungary</td>
<td>1,860,159</td>
<td>19,342</td>
<td>97,280</td>
<td>1,012</td>
<td>45,611</td>
<td>474</td>
<td>9,617,343</td>
</tr>
<tr>
<td>Iceland</td>
<td>1,474,374</td>
<td>29,286</td>
<td>154,166</td>
<td>3,062</td>
<td>6,799</td>
<td>135</td>
<td>5,034,484</td>
</tr>
<tr>
<td>Italy</td>
<td>14,877,144</td>
<td>24,669</td>
<td>1,274,305</td>
<td>2,113</td>
<td>159,909</td>
<td>265</td>
<td>60,305,943</td>
</tr>
<tr>
<td>Latvia</td>
<td>804,288</td>
<td>43,483</td>
<td>31,740</td>
<td>1,716</td>
<td>5,657</td>
<td>306</td>
<td>1,849,641</td>
</tr>
<tr>
<td>Lithuania</td>
<td>1,033,547</td>
<td>38,918</td>
<td>51,071</td>
<td>1,923</td>
<td>8,925</td>
<td>336</td>
<td>2,655,708</td>
</tr>
<tr>
<td>Luxembourg</td>
<td>219,390</td>
<td>34,076</td>
<td>18,113</td>
<td>2,813</td>
<td>1,041</td>
<td>162</td>
<td>643,829</td>
</tr>
<tr>
<td>Malta</td>
<td>82,845</td>
<td>18,675</td>
<td>7,969</td>
<td>1,796</td>
<td>649</td>
<td>146</td>
<td>443,605</td>
</tr>
<tr>
<td>Netherlands</td>
<td>7,935,106</td>
<td>46,131</td>
<td>1,227,890</td>
<td>7,138</td>
<td>22,037</td>
<td>128</td>
<td>17,201,349</td>
</tr>
<tr>
<td>Poland</td>
<td>5,971,998</td>
<td>15,810</td>
<td>522,946</td>
<td>1,384</td>
<td>115,395</td>
<td>305</td>
<td>37,773,933</td>
</tr>
<tr>
<td>Portugal</td>
<td>3,604,114</td>
<td>35,527</td>
<td>21,693</td>
<td>214</td>
<td>10,144,581</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Romania</td>
<td>2,864,475</td>
<td>15,066</td>
<td>192,684</td>
<td>1,013</td>
<td>65,129</td>
<td>343</td>
<td>19,012,700</td>
</tr>
<tr>
<td>Slovakia</td>
<td>17,702,096</td>
<td>21,699</td>
<td>85,628</td>
<td>1,567</td>
<td>19,462</td>
<td>356</td>
<td>5,464,279</td>
</tr>
<tr>
<td>Slovenia</td>
<td>978,154</td>
<td>47,038</td>
<td>34,930</td>
<td>1,680</td>
<td>6,512</td>
<td>313</td>
<td>2,079,439</td>
</tr>
<tr>
<td>Spain</td>
<td>11,551,574</td>
<td>24,690</td>
<td>530,130</td>
<td>1,133</td>
<td>102,541</td>
<td>219</td>
<td>46,786,531</td>
</tr>
<tr>
<td>Sweden</td>
<td>2,487,852</td>
<td>24,368</td>
<td>28,835</td>
<td>282</td>
<td>18,331</td>
<td>180</td>
<td>10,209,679</td>
</tr>
</tbody>
</table>

* World population based on https://www.worldometers.info/worldpopulation/

The reported numbers refer to cases that have been tested and confirmed to be carrying the virus and sometimes, depending upon the country, also presumptive, suspect, or probable cases of detected infection. There are large geographic variations in the proportion of the population tested, as well as in the quality of reporting across countries. People who carry the virus but remain asymptomatic are less likely to be tested and therefore mild cases are likely underreported. Further, as at-home rapid testing kits have become more readily

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available\textsuperscript{222} and formal testing resources reach capacity due to the Omicron variant, the true estimate of cases is estimated to be larger than formally reported counts, the true estimate of cases is estimated to be larger than formally reported counts. The numbers should therefore be interpreted with caution. While there is limited information on number of cases attributable to specific variants, recent case counts are likely to reflect the Omicron variant, which is currently the predominant strain in many countries, including the US\textsuperscript{223}. Omicron BA.1.1 was responsible for 57.3\%, BA.2 is responsible for 34.9\%, and B.1.1.529 was responsible for 7.9\% of all SARS-CoV-2 specimens sequenced by the CDC during the week ending 19 March 2022.\textsuperscript{223}

The main existing treatment options:

Through 18 June 2022, other COVID-19 vaccines were authorised\textsuperscript{224} in the European Union including COVID-19 Vaccine (inactivated, adjuvant), Spikevax (EU/1/20/1507), JCOVDEN (EU/1/20/1525), Vaxzevria (EU/1/21/1529), Nuvaxovid (EU/1/21/1618).

Natural history of the indicated condition in the untreated population, including mortality and morbidity:

Symptoms of COVID-19

The clinical manifestations of COVID-19 vary widely, from asymptomatic infection in 17–45\% of patients, across age groups\textsuperscript{225,226,227,228} to critical illness and death. The rate of asymptomatic infection decreases with increasing age and long-term care facilities are associated with a lower rate of asymptomatic infection when compared to household transmission or other healthcare facilities.\textsuperscript{228} One recent meta-analysis has estimated that 46.7\% of infections in children are asymptomatic.\textsuperscript{228} The most common symptoms of


COVID-19 are fever, cough, and shortness of breath for both children and adults.\textsuperscript{229,230} Confirming these observations in a recent systematic review, researchers examined 1,140 cases of COVID-19 in children from 23 published studies. They reported that 11% of cases were asymptomatic. Among symptoms, fever was reported in 48%, cough in 37%, and any nasopharyngeal symptom in 22%.\textsuperscript{231}

**Progression and Timeline of Mild to Moderate Disease**

Mild to moderate disease is defined as the absence of viral pneumonia and hypoxia. For those who develop symptoms, the incubation period is usually 4 to 5 days, with 97.5% experiencing symptoms within 11 days of exposure.\textsuperscript{232,233} Those with mild COVID-19 recover at home with supportive care and guidance to self-isolate. Those with moderate disease are monitored at home and are sometimes recommended to be hospitalised if conditions worsen.\textsuperscript{233} Data on rates of re-infection are limited but variants that are not neutralised by immune antiserum, such as the beta (South African), Delta, and Omicron variants, may lead to increased risk of re-infection in the future.\textsuperscript{233,234}

**Progression and Timeline of Severe Disease Requiring Hospitalisation**

Those with severe disease will require hospitalisation to manage their illness. Based on data that have been systematically collected for the US by the CDC between 01 August 2020 and 22 March 2022, there were 4,580,996 total hospital admissions for patients with confirmed COVID-19 in the US.\textsuperscript{235} For the week ending 20 March 2022, 9.3 per 100 000 population (country range: 2.7–42.8) were hospitalised due to COVID-19 in 17 countries of the EU/EEA with available data.\textsuperscript{236} As of 24 March 2022, 0.1% -1.5% of children who tested positive for


COVID-19 have been hospitalised (for any diagnosis) based on data reported from 25 states and New York City reporting, and 0.00%-0.01% of children with COVID-19 have died based on data reported from 46 states, New York City, Puerto Rico and Guam.\(^{237}\)

The most common symptoms in patients are fever (42-80%), shortness of breath (35-71%), fatigue (33-62%), cough (77-84%), chills (63%), myalgias (63%), headache (59%), and diarrhea (33%).\(^{238,239,240,241}\) COVID-19 patients also commonly experience gustatory disorders (44%) and olfactory disorders (53%).\(^{242}\) Among unhospitalised children < 18 years of age, 89% experienced one or more typical symptoms of COVID, including fever, cough, shortness of breath, and 22% experienced all three.\(^{240}\) Approximately 17% to 40% of those hospitalised with COVID-19 experience severe symptoms necessitating intensive care\(^{243,244,239}\) with 31% of children hospitalised experiencing severe COVID-19 that necessitates intensive care or invasive ventilation or ends in death. Risk factors for severe COVID-19 in hospitalised children include presence of a comorbid condition, younger age, and male sex.\(^{245}\) More than 75% of patients hospitalised with COVID-19 require supplemental oxygen.\(^{246}\)

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Studies early in the pandemic demonstrated that time from onset of illness to ARDS was 8-12 days and time from onset of illness to ICU admission was 9.5–12 days. In 12 countries of the EU/EEA with available data, 0.7 per 100,000 population (country range 0.1-4.1) were in the ICU due to COVID-19 for the week ending 20 March 2022. A recent meta-analysis found that, of patients <19 years of age, 11% went to the ICU, non-invasive ventilation was administered among 12%, and 4% required mechanical ventilation. A study of 82 cases in three pediatric hospitals noted that older children and those with higher body mass index or multiple comorbidities were more likely to receive respiratory support.

**Mortality**

As of 27 March 2022, there were 974,277 deaths reported in the US for all age groups among 79,766,087 individuals positive for COVID-19 (1.2% of cases) As of the week ending on 20 March 2022, the mortality rate was 29.6 per million population (country range: 7.1–119.0) in the EU. As of 27 March 2022, the UK has seen 165,046 deaths from COVID-19 in all age groups among 20,848,913 cases (0.8% of cases). According to a recent meta-analysis of paediatric studies published through October 2020, the mortality for pediatric patients is 0.1–2%. In a study from January through June 2020 using the National Child Mortality Database (NCMD) in England, 5.7% of 437 children 0-17 years of age who died were SARS-CoV-2 PCR positive and those who died of COVID-19 were older and were more likely to be non-White ethnicity.

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Mortality data are also presented from Worldometers, an independent organisation that publishes current, reliable COVID-19 statistics online.218 The mortality of SARS-CoV-2 infection is defined as the cumulative number of deaths among detected cases.

As of 05 April 2022, the overall SARS-CoV-2 mortality for the EU + UK was 1,218,285 deaths, or 237 per 100,000 people. Reported mortality among EU countries and the UK ranged from 52 to 534 deaths per 100,000. Cyprus and Finland reported the lowest mortality; Croatia, Hungary, and Bulgaria reported the highest.219

In the US, as of 05 April 2022, the mortality was 1,008,697 deaths (2302 per 100,000 people). Mortality in the US was higher than that of the UK (242 per 100,000).219

Overall reported mortality among hospitalised COVID-19 patients varies from 12.8% to 26% in the EU and UK and US244,255,256,257

**Complications of COVID-19 and Post-acute COVID**

Recent evidence has shown that a range of persistent symptoms can remain long after the acute SARS-CoV-2 infection. This condition has been called long COVID or post-acute COVID by some recognised research institutes; a universally accepted definition of long COVID has yet to be established.

Studies have shown that long COVID can affect the whole spectrum of people with COVID-19, from those with very mild acute disease to the most severe forms.

Studies around the world have reported various incidence rates for long COVID with different follow-up examination times after the acute infection, including 76% of people at 6 months, 32.6% at 60 days, 87% at 60 days, and 96% at 90 days. These finding are not fully corroborative, but they show that a substantial proportion of people who have had COVID-19 may develop long COVID.258

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Assuming at least 10% of COVID-19 survivors develop long COVID, it is estimated that 5 million people are facing long COVID globally.\textsuperscript{259}

This illness is poorly understood as it affects COVID-19 survivors at all levels of disease severity, even younger adults, children, and those not hospitalised. While the precise definition of long COVID may be lacking, the most common symptoms reported in many studies are fatigue and dyspnoea that last for months after acute COVID-19. Other persistent symptoms may include cognitive and mental impairments, chest and joint pains, palpitations, myalgia, smell and taste dysfunctions, cough, headache, and gastrointestinal and cardiac issues.

Presently, there is limited literature discussing the possible pathophysiology, risk factors, and treatments in long COVID, which the current review aims to address. In brief, long COVID may be driven by long-term tissue damage (e.g., lung, brain, and heart) and pathological inflammation (e.g., from viral persistence, immune dysregulation, and autoimmunity). The associated risk factors may include female sex, more than five early symptoms, early dyspnoea, prior psychiatric disorders, and specific biomarkers (e.g., D-dimer, CRP, and lymphocyte count), although more research is required to substantiate such risk factors.\textsuperscript{259}

Several studies are evaluating a potential impact of SARS Cov-2 vaccination on long COVID:

Ayoubkhani et al. described that a first dose of COVID-19 vaccine was associated with a reduction in long COVID symptoms 12.8% decrease (95% confidence interval $-18.6\%$ to $-6.6\%$, $P<0.001$), and evidence suggested a sustained improvement after a second dose, with an initial 8.8% decrease (95% confidence interval $-14.1\%$ to $-3.1\%$, $P=0.003$) in the odds of long COVID, with a subsequent decrease by 0.8% per week ($-1.2\%$ to $-0.4\%$ per week, $P<0.001$), at least over the median follow-up of 67 days in this study.

No evidence was found of differences in this relationship by sociodemographic characteristics, health related factors, vaccine type, or duration from infection to vaccination.

Although causality cannot be inferred from this observational evidence, vaccination may contribute to a reduction in the population health burden of long COVID.\textsuperscript{260}

Furthermore, Kuodzi et al.\textsuperscript{261} showed that two doses of BNT162b2 vaccine reduced the risk of the most common long COVID symptoms after COVID-19 infection, in a cross-sectional study preformed between 15 March 2020–15 November 2021. They found that patients who

\textsuperscript{259} Yung SJ. Long COVID or post-COVID-19 syndrome?putative pathophysiology, risk factors, and treatments. Infectious diseases 2021; VOL 0, No. 0, 1-18.


\textsuperscript{261} Kuodzi P, et al. medRxiv. Published online 17 January 2022. doi:10.1101/2022.01.05.22268800.
received 2 doses of BNT162b2 were 54% to 82% less likely to report 7 of the 10 most commonly reported symptoms compared with unvaccinated patients (all P<0.04).

Post COVID has also been described in children, a national survey in the UK found 7-8% of children with COVID-19 reported continued symptoms >12 weeks.262

Long COVID can appear after mild to severe infections, and after MIS-C. Most common symptoms: similar to adults and include fatigue, headache, insomnia, trouble concentrating, muscle and joint pain, and cough. Impact on quality of life: limitations of physical activity, feeling distressed about symptoms, mental health challenges, decreased school attendance/participation.

Post-COVID conditions may be less likely to occur after vaccine breakthrough in adolescents.263,264

Persons who were previously vaccinated were less likely to have symptoms between 12 and 20 weeks after infection compared to persons who were unvaccinated (OR 0.22; 95% 0.20, 0.25) with a lower occurrence of post-COVID conditions after infection compared to persons who were unvaccinated.263,264

Further research is needed, but vaccination may contribute to a reduction in the population health burden of long COVID.

18.2. Benefit-Risk Analysis Evaluation

Based on the safety data presented in Section 16 and the benefits presented in Section 17, this section presents an overall qualitative evaluation of the benefit risk analysis of BNT162b2 in prevention of COVID-19 infection. With respect to benefit, the nature, clinical importance, duration, efficacy profile, and pharmacokinetic benefits of BNT162b2 were considered. With respect to the risks, data from clinical trials, post-marketing, and literature sources were considered as well as important potential and identified risks, if applicable.

Limitations

Some limitations of the benefit-risk analysis may include missing information in certain special populations and the inherent limitations of the various data sources, as summarised below.


These limitations were considered when evaluating the overall benefit-risk profile of BNT162b2.

**Clinical trials:**

a) The participants in clinical trials are a relatively homogeneous group as they all meet study inclusion criteria. Importantly, certain populations may be excluded.

b) Close monitoring required as part of study participation likely identifies relatively common events. Events that are dose-related and pharmacologically predictable events may be distinguished. However, clinical studies may not be powered to pick up rare safety issues.

**Non-interventional (observational) study data:**

a) There is limited control over patient assessment as patient monitoring and diagnostics are per standard of care; no additional clinical monitoring is generally conducted.

b) Patient specific methodological challenges such as potential biases from patient selection, loss of patients through study attrition, and overall patient recall are also inherent limitations.

**Post-marketing data:**

a) Reports originate from multiple sources (consumer and healthcare professional) and they can be poorly characterised from a medical perspective.

b) Limited or incomplete information is common, including indication, medical history, concomitant medication use, and reason for reporting as an AE, making it difficult to fully characterise events and associated risk factors.

c) Difficult to contextualize quantitatively, as voluntary and sporadic reporting do not allow complete knowledge of total exposure or total number of events ever experienced in the exposed population. These data are generally not suitable to make between-drug comparisons.

**18.2.1. Benefits**

Please refer to Section 17.

**18.2.2. Risks**

An assessment of the important risks, identified and potential, was performed using the following data sources: pre-clinical studies, clinical studies, post-marketing experience, and literature as applicable. Interval findings are summarised in Table 82.

Based on pharmacovigilance monitoring activities, there has been no new safety information contributing importantly to the risks of BNT162b2.
Table 82. Summary of Important Risks

<table>
<thead>
<tr>
<th>Risks</th>
<th>Clinical Study Data</th>
<th>Post-Marketing Data</th>
<th>Literature Sources</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Important Identified Risks</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anaphylaxis</td>
<td>No new data from clinical studies were identified during the reporting interval.</td>
<td>Based on the review of post-marketing data, no new safety information was identified for BNT162b2 and anaphylaxis.</td>
<td>No new significant data received from literature sources.</td>
<td>The risk is communicated through the CDS, Sections 4.4 Special warnings and precautions for use, 4.8 Undesirable effects, Appendix A and Appendix B and in the EU SmPC, Sections 4.4 Special warnings and precautions for use and 4.8 Undesirable effects. It is being monitored through routine pharmacovigilance. Based upon review of the available information, no additional change to the RSI is warranted at this time.</td>
</tr>
<tr>
<td>Myocarditis and Pericarditis</td>
<td>No new data from clinical studies were identified during the reporting interval.</td>
<td>Based on the review of post-marketing data, no new safety information was identified for BNT162b2 and myocarditis and pericarditis.</td>
<td>During the reporting period an unpublished presentation including significant information on myocarditis was reviewed. Please refer to Section 11 Literature for details.</td>
<td>The risk is communicated through the CDS in the Section 4.4 Special warnings and precautions for use and EU SmPC in the Section 4.8 Undesirable effects. It is also included as an Important identified risk in the EU RMP and in the US PVP. Considering the accumulating data from post-authorisation use of the vaccine, myocarditis and pericarditis have been added as ADRs in the Section 4.8 Undesirable effects, in Appendix A and Appendix B of the CDS v. 14.0, made effective on 26 July 2022, after the DLP.</td>
</tr>
<tr>
<td><strong>Important Potential Risks</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VAED VAERD</td>
<td>No new data from clinical studies were identified during the reporting interval.</td>
<td>Based on the review of post-marketing data, no new safety information was identified for BNT162b2 and VAED-VAERD.</td>
<td>No new significant data received from literature sources.</td>
<td>VAED-VAERD is theoretical because it has not been described in association with the COVID-19 mRNA vaccine or it has not been reported from any other late phase clinical trial of other human vaccine. It is included as an Important Potential Risk in the EU-RMP and in the US-PVP. Based upon review of the available information, no additional change to the RSI is warranted at this time.</td>
</tr>
</tbody>
</table>
18.2.3. Overall Benefit-Risk

The important risks associated with the use of BNT162b2 are minimised through provision of relevant product information in the RSI to support safe use of the product. Risks have been evaluated in the context of the enumerated benefits of the product. Based on the available safety and efficacy/effectiveness data for BNT162b2, the overall benefit-risk profile of BNT162b2 remains favourable for all age groups in which it is authorised.

Table 83. Overall Benefit-Risk for BNT162b2

<table>
<thead>
<tr>
<th>Consideration</th>
<th>Favourable Benefit-Risk</th>
<th>Non Contributory</th>
<th>Unfavourable Benefit-Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severity of condition</td>
<td>The severity of the condition being treated, as well as comorbidities and outcomes in the population to be treated were considered. (See Section 18.1)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Unmet medical need</td>
<td>BNT162b2 meets an unmet medical need because there is - lack of alternative therapies, or - although alternative products are available in this class, this product may be the preferred therapeutic option or preferred in a select group of patients. (See Section 18.1)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Clinical benefit</td>
<td>The nature, clinical importance, duration, and generalizability of benefits were considered. (See Section 18.1)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Risk associated with treatment</td>
<td>The nature, seriousness, frequency, predictability, reversibility, impact on patients and public health of the product's risks were considered. (See Section 18.2.2)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Risk management</td>
<td>Risk minimisation measures currently in place for this product support a favourable benefit-risk balance. (See Section 18.2.2)</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Table was adapted from European Medicines Agency. Benefit-risk Methodology Project – Working package 2 report: Applicability of current tools and processes for regulatory benefit-risk assessment. 31 August 2010.

19. CONCLUSION AND ACTIONS

Risks have been evaluated in the context of the benefits of the vaccine. Based on the available safety and efficacy/effectiveness and immunogenicity data from the reporting interval for BNT162b2, the benefit-risk profile of BNT162b2 remains favourable. No additional changes to the BNT162b2 RSI or additional risk minimisation activities in addition to those in place are warranted at this time.

The MAH will continue to review the safety of BNT162b2, including all reports of adverse experiences and will revise the product documents if an evaluation of the safety data yields significant new information.