

## **Avamys**

Procedural steps taken and scientific information after the authorisation

| Application number    | Scope   | Opinion/<br>Notification <sup>1</sup> issued on | Commission Decision Issued <sup>2</sup> / amended on | Product<br>Information<br>affected <sup>3</sup> | Summary                           |
|-----------------------|---|---|--|---|-----------------------------------|
| IB/0047               | B.II.e.3.z - Change in test procedure for the immediate packaging of the finished product - Other variation | 24/01/2022                                      | n/a  |   |                                   |
| PSUSA/9154/<br>202104 | Periodic Safety Update EU Single assessment - fluticasone furoate   | 02/12/2021                                      | n/a  |   | PRAC Recommendation - maintenance |

<sup>&</sup>lt;sup>1</sup> Notifications are issued for type I variations and Article 61(3) notifications (unless part of a group including a type II variation or extension application or a worksharing application). Opinions are issued for all other procedures.



<sup>&</sup>lt;sup>2</sup> A Commission decision (CD) is issued for procedures that affect the terms of the marketing authorisation (e.g. summary of product characteristics, annex II, labelling, package leaflet). The CD is issued within two months of the opinion for variations falling under the scope of Article 23.1a(a) of Regulation (EU) No. 712/2012, or within one year for other procedures.

<sup>3</sup> SmPC (Summary of Product Characteristics), Annex II, Labelling, PL (Package Leaflet).

| N/0045      | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)  | 10/09/2021 |            | PL                                     |  |
|-------------|---|------------|------------|--|--|
| IAIN/0044/G | This was an application for a group of variations.  B.I.a.1.a - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - The proposed manufacturer is part of the same pharmaceutical group as the currently approved manufacturer  B.I.a.1.f - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - Changes to quality control testing arrangements for the AS -replacement or addition of a site where batch control/testing takes place  B.I.a.4.z - Change to in-process tests or limits applied during the manufacture of the AS - Other variation | 17/05/2021 | n/a        |  |  |
| IA/0043     | A.7 - Administrative change - Deletion of manufacturing sites   | 10/02/2021 | 21/04/2021 | Annex II and<br>PL                     |  |
| IAIN/0042   | B.II.b.2.c.1 - Change to importer, batch release arrangements and quality control testing of the FP - Replacement or addition of a manufacturer responsible for importation and/or batch release - Not including batch control/testing  | 15/01/2021 | n/a        |  |  |
| IB/0041     | C.I.z - Changes (Safety/Efficacy) of Human and<br>Veterinary Medicinal Products - Other variation   | 26/08/2020 | 21/04/2021 | SmPC, Annex<br>II, Labelling<br>and PL |  |

| II/0040               | Update of section 4.8 of the SmPC in order to add bronchospasm with a frequency 'not known' and dyspnoea with a frequency 'common' to the list of adverse drug reactions based on post-marketing experience and clinical trials reports. The package leaflet is updated accordingly. In addition, the MAH took the opportunity to update the list of local representatives in the package leaflet.  C.I.3.b - Change(s) in the SPC, Labelling or PL intended to implement the outcome of a procedure concerning PSUR or PASS or the outcome of the assessment done under A 45/46 - Change(s) with new additional data submitted by the MAH | 02/04/2020 | 21/04/2021 | SmPC and PL                  | Based on clinical trials and post-marketing data, causal relationship between fluticasone furoate and bronchospasm and/or dyspnoea cannot be excluded. Therefore, bronchospasm and dyspnoea are included in the list of adverse drug reactions in the product information of Avamys. |
|-----------------------|--|------------|------------|------------------------------|--|
| IB/0039/G             | This was an application for a group of variations.  B.II.e.4.a - Change in shape or dimensions of the container or closure (immediate packaging) - Non-sterile medicinal products  | 16/01/2019 | n/a        |                              |  |
| T/0037                | Transfer of Marketing Authorisation  | 12/10/2018 | 06/12/2018 | SmPC,<br>Labelling and<br>PL |  |
| PSUSA/9154/<br>201804 | Periodic Safety Update EU Single assessment - fluticasone furoate  | 29/11/2018 | n/a        |                              | PRAC Recommendation - maintenance  |
| IA/0038/G             | This was an application for a group of variations.  A.4 - Administrative change - Change in the name and/or address of a manufacturer or an ASMF holder  | 23/10/2018 | n/a        |                              |  |

|           | or supplier of the AS, starting material, reagent or intermediate used in the manufacture of the AS or manufacturer of a novel excipient A.7 - Administrative change - Deletion of manufacturing sites A.7 - Administrative change - Deletion of manufacturing sites A.7 - Administrative change - Deletion of manufacturing sites  |            |            |             |  |
|-----------|---|------------|------------|-------------|--|
| WS/1263/G | This was an application for a group of variations following a worksharing procedure according to Article 20 of Commission Regulation (EC) No 1234/2008.  B.I.a.1.a - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - The proposed manufacturer is part of the same pharmaceutical group as the currently approved manufacturer  B.I.a.3.a - Change in batch size (including batch size ranges) of AS or intermediate - Up to 10-fold increase compared to the originally approved batch size | 16/11/2017 | n/a        |             |  |
| IAIN/0034 | C.I.3.a - Change(s) in the SPC, Labelling or PL intended to implement the outcome of a procedure concerning PSUR or PASS or the outcome of the assessment done under A 45/46 - Implementation of wording agreed by the competent authority  | 18/08/2017 | 30/04/2018 | SmPC and PL |  |

| IB/0032   | C.I.z - Changes (Safety/Efficacy) of Human and<br>Veterinary Medicinal Products - Other variation   | 24/05/2017 | 30/04/2018 | SmPC, Annex<br>II, Labelling<br>and PL |
|-----------|---|------------|------------|--|
| IB/0031/G | This was an application for a group of variations.  B.II.e.1.a.2 - Change in immediate packaging of the finished product - Qualitative and quantitative composition - Semi-solid and non-sterile liquid pharmaceutical forms  B.II.e.2.z - Change in the specification parameters and/or limits of the immediate packaging of the finished product - Other variation  | 10/03/2017 | n/a        |  |
| IB/0029/G | This was an application for a group of variations.  B.II.e.2.z - Change in the specification parameters and/or limits of the immediate packaging of the finished product - Other variation  B.II.d.1.h - Change in the specification parameters and/or limits of the finished product - Update of the dossier to comply with the provisions of an updated general monograph of the Ph. Eur. for the finished product  B.II.e.7.a - Change in supplier of packaging components or devices (when mentioned in the dossier) - Deletion of a supplier  B.II.e.1.a.2 - Change in immediate packaging of the finished product - Qualitative and quantitative composition - Semi-solid and non-sterile liquid pharmaceutical forms | 11/07/2016 | n/a        |  |

| II/0030/G | This was an application for a group of variations.   | 23/06/2016 | n/a        |             |   |
|-----------|--|------------|------------|-------------|---|
|           | Type II-Submission of a post authorisation safety study PASS 201077 Retrospective Case-Control Studies of Rare Adverse Events Associated with Intranasal Steroids  Type IB-Submission of a new RMP v.11 to include cataracts and glaucoma as identified risks, following the recent PRAC assessment (EMA PSUSA/00009154/201504)  In addition, the MAH took the opportunity to reflect in the RMP the current SmPC wording with regard to patient with hepatic impairment agreed in the previously approved variation  EMEA/H/C/000770/II/0027.  C.I.11.z - Introduction of, or change(s) to, the obligations and conditions of a marketing |            |            |             |   |
|           | authorisation, including the RMP - Other variation  C.I.13 - Other variations not specifically covered elsewhere in this Annex which involve the submission of studies to the competent authority  |            |            |             |   |
| II/0027   | Update of sections 4.2 and 4.4 of the SmPC to remove warnings associated with hepatic impairment. The Package Leaflet is updated accordingly. In addition, the MAH took the opportunity to update the contact details for the local representatives in the Package Leaflet.  | 17/12/2015 | 07/03/2016 | SmPC and PL | The MAH has provided safety data showing that moderate or severe hepatic impairment is not expected to result in a clinically relevant effect when the recommended intranasal fluticasone furoate dose is used. The corresponding wording in sections 4.2 and 4.4 of the SmPC and the Package Leaflet has been deleted accordingly. |
|           | C.I.4 - Change(s) in the SPC, Labelling or PL due to   |            |            |             |   |

|                       | new quality, preclinical, clinical or pharmacovigilance data  |            |            |             |                                   |
|-----------------------|---|------------|------------|-------------|-----------------------------------|
| N/0028                | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)  | 15/12/2015 | 07/03/2016 | PL          |                                   |
| PSUSA/9154/<br>201504 | Periodic Safety Update EU Single assessment - fluticasone furoate   | 03/12/2015 | n/a        |             | PRAC Recommendation - maintenance |
| II/0026               | Update of section 4.8 of the SmPC to add the new ADR nasal septum perforation with a frequency classification of very rare. The Package leaflet has been updated accordingly. In addition, the MAH took the opportunity to update the contact details of the local representatives in Malta, France, Cyprus and the UK in the Package Leaflet.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data | 22/10/2015 | 07/03/2016 | SmPC and PL | N/A                               |
| IA/0024               | B.II.e.7.b - Change in supplier of packaging components or devices (when mentioned in the dossier) - Replacement or addition of a supplier  | 31/07/2015 | n/a        |             |                                   |
| IAIN/0023/G           | This was an application for a group of variations.  A.4 - Administrative change - Change in the name and/or address of a manufacturer or an ASMF holder or supplier of the AS, starting material, reagent or intermediate used in the manufacture of the AS or  | 08/05/2015 | n/a        |             |                                   |

|         | manufacturer of a novel excipient  B.I.a.1.a - Change in the manufacturer of AS or of a starting material/reagent/intermediate for AS - The proposed manufacturer is part of the same pharmaceutical group as the currently approved manufacturer  |            |            |                          |   |
|---------|--|------------|------------|--------------------------|---|
| II/0022 | Update of the sections 4.2 and 5.2 of the SmPC with modifications to hepatic impairment dose recommendations for fluticasone furoate nasal spray (FFNS).  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data  | 26/03/2015 | 07/03/2016 | SmPC                     |   |
| 11/0020 | Update of sections 4.8, 5.1, and 6.5 of the SmPC in order to incorporate the QRD comments received as part of the renewal assessment procedure (EMEA/H/C000770/R/0014). The Package Leaflet is updated accordingly.  C.I.4 - Change(s) in the SPC, Labelling or PL due to new quality, preclinical, clinical or pharmacovigilance data | 21/11/2013 | 28/04/2014 | SmPC, Annex<br>II and PL | Update the sections 4.8 and section 5.1 of Avamys SmPC, as well as the corresponding sections in Annexes II and III of Avamys, in order to incorporate the QRD comments received at the renewal assessment procedure (EMEA/H/C000770/R/0014).  In addition, MAH made slight correction to section 6.5 'Nature and contents of container' to clarify that '14.2ml' does not refer to the volume of Avamys suspension, but to the capacity of the glass container, and to update Czech Republic email address in Annex III of Avamys. |
| IA/0021 | A.7 - Administrative change - Deletion of manufacturing sites  | 28/10/2013 | n/a        |                          |   |
| N/0019  | Minor change in labelling or package leaflet not connected with the SPC (Art. 61.3 Notification)   | 22/08/2013 | 28/04/2014 | PL                       |   |

| IB/0018/G | This was an application for a group of variations.  B.II.e.1.a.2 - Change in immediate packaging of the finished product - Qualitative and quantitative composition - Semi-solid and non-sterile liquid pharmaceutical forms  B.II.e.7.b - Change in supplier of packaging components or devices (when mentioned in the dossier) - Replacement or addition of a supplier | 18/06/2013 | 28/04/2014 | SmPC                                   |  |
|-----------|--|------------|------------|--|--|
| IG/0279   | A.1 - Administrative change - Change in the name and/or address of the MAH   | 18/04/2013 | 28/04/2014 | SmPC,<br>Labelling and<br>PL           |  |
| IG/0275   | C.I.z - Changes (Safety/Efficacy) of Human and<br>Veterinary Medicinal Products - Other variation  | 15/03/2013 | n/a        |  |  |
| R/0014    | Renewal of the marketing authorisation.  | 18/10/2012 | 17/12/2012 | SmPC, Annex<br>II, Labelling<br>and PL |  |
| IA/0015   | B.II.e.7.b - Change in supplier of packaging components or devices (when mentioned in the dossier) - Replacement or addition of a supplier   | 01/08/2012 | n/a        |  |  |
| II/0012   | The variation relates to an update of sections 4.4 (Warnings and Precautions), 4.8 (Undesirable effects) and 5.1 (Pharmacodynamic properties) of the Summary of Product Characteristics and section 4 'Possible Side Effects' of the Package Leaflet to include information on 'ocular changes' in the treated   | 21/06/2012 | 20/07/2012 | SmPC and PL                            | The assessment of the study FFR110537 has shown that once-daily Fluticasone Furoate Nasal Spray 110 mcg induces ocular changes. Information regarding 'ocular changes' in treated population was therefore added to the product information. |

|           | population following the CHMP assessment of the results from study FFR110537.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data   |            |            |             |  |
|-----------|---|------------|------------|-------------|--|
| II/0011   | The variation relates to an update by the MAH of sections 4.4 (Warnings and Precautions), 4.8 (Undesirable Effects) and 5.1 (Pharmacodynamic properties) of the SmPC (Summary of Product Characteristics) and section 4 'Possible Side Effects' of the Package Leaflet (PL) to include information on 'growth retardation' in the paediatric use following the CHMP assessment of the results of study FFR101782.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data | 21/06/2012 | 20/07/2012 | SmPC and PL | The assessment of the study FFR101782 have shown a one-year course of FFNS spray 110 mcg QD influences growth in pre-pubescent, paediatric subjects 5-8.5 years of age with periannial allergic rhinitis. Information regarding 'growth retardation' in paediatric use was therefore added to the product information. |
| IG/0150/G | This was an application for a group of variations.  C.I.9.c - Changes to an existing pharmacovigilance system as described in the DDPS - Change of the back-up procedure of the QPPV  C.I.9.h - Changes to an existing pharmacovigilance system as described in the DDPS - Other change(s) to the DDPS that does not impact on the operation of the pharmacovigilance system  | 05/04/2012 | n/a        |             |  |

| II/0010   | The variation relates to an update of section 4.8  (Undesirable Effects) of the SmPC (Summary of Product Characteristics) and section 4 'Possible Side Effects' of the Package Leaflet (PL) to include 'headache' following the assessment of the latest PSUR. In addition, minor changes have been made in accordance with the QRD template and for consistency throughout the product information.  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data  | 16/02/2012 | 15/03/2012 | SmPC and PL              | Following the assessment of the latest PSUR of fluticasone furoate, a number of reports on headache following intranasal fluticasone furoate were received from the MAH's Safety Database. A review of the spontaneously received reports, clinical trial data and literature supported the possibility that these events could be due to fluticasone furoate nasal spray. 'Headache' with a 'common' incidence was therefore added to the fluticasone furoate nasal spray product information. |
|-----------|---|------------|------------|--------------------------|---|
| II/0007/G | This was an application for a group of variations.  This was an application for a group of variations.  Update of Summary of Product Characteristics, Annex II and Package Leaflet.  Update of SmPC section 4.8 to add the adverse drug reactions rhinalgia, nasal discomfort and nasal dryness. Update of SmPC sections 4.4 and 4.8 regarding the long-term use of nasal corticosteroids, as requested by the CHMP. The Package Leaflet has been updated accordingly. In addition, minor linguistic and typographical amendments have been made in the SmPC and PL, and the SmPC, Annex II and PL was aligned with the latest version of the QRD template. | 22/09/2011 | 10/11/2011 | SmPC, Annex<br>II and PL | Based on results of review of safety data on nasal disorders from the literature, spontaneous reports and clinical trials, rhinalgia, nasal discomfort and nasal dryness were identified as potential adverse reactions in the frequency group 'uncommon'. Furthermore, the labelling was updated with recommended wording following a review of the long-term safety related to nasal corticosteroids.   |

|           | This application concerns grouping of variations of type IB (C.I.3.a) and type II (C.I.4).  C.I.4 - Variations related to significant modifications of the SPC due in particular to new quality, preclinical, clinical or pharmacovigilance data  C.I.3.a - Implementation of change(s) requested following the assessment of an USR, class labelling, a PSUR, RMP, FUM/SO, data submitted under A 45/46, or amendments to reflect a Core SPC - Changes with NO new additional data are submitted by the MAH   |            |     |          |
|-----------|--|------------|-----|----------|
| IB/0008   | B.II.d.2.d - Change in test procedure for the finished product - Other changes to a test procedure (including replacement or addition)   | 20/10/2011 | n/a |          |
| IG/0034/G | This was an application for a group of variations.  C.I.9.b - Changes to an existing pharmacovigilance system as described in the DDPS - Change in the contact details of the QPPV  C.I.9.c - Changes to an existing pharmacovigilance system as described in the DDPS - Change of the back-up procedure of the QPPV  C.I.9.e - Changes to an existing pharmacovigilance system as described in the DDPS - Changes in the major contractual arrangements with other persons or organisations involved in the fulfilment of pharmacovigilance obligations and described in the DD  C.I.9.g - Changes to an existing pharmacovigilance | 06/01/2011 | n/a | Annex II |

|         | of the Qualified Person for Pharmacovigilance (QPPV). Consequently, Annex II has been updated with the new version number.  Changes to QPPV Update of DDPS (Pharmacovigilance)  |            |            |                          | the DDPS performed since the last approved version.  Consequently, Annex II has been updated using the standard text including the new version number of the agreed DDPS. The CHMP considers that the Pharmacovigilance System as described by the MAH fulfils the requirements.  |
|---------|---|------------|------------|--------------------------|---|
| II/0003 | Inclusion of 'hypersensitivity' in section 4.8  'Undesirable Effects' of the Summary of Product Characteristics (SPC) and section 4 'Possible Side Effects' of the Package Leaflet (PL) accordingly. In addition, the MAH took the opportunity to make minor changes in sections 5.1 of the SPC and sections 3, 4 and 5 of the PL. Finally, the instructions on the use of Avamys in the PL were added.  Update of Summary of Product Characteristics and Package Leaflet | 19/11/2009 | 21/12/2009 | SmPC and PL              | Reports consistent with hypersensitivity reactions to intranasal fluticasone furoate were received from the MAH's Safety Database. A review of the spontaneously received reports, clinical trial data and literature supported the possibility that these events could be due to fluticasone furoate nasal spray. 'Hypersensitivity' with a 'rare' incidence was therefore added to the fluticasone furoate nasal spray product information. |
| II/0001 | Update the sections 4.2, 4.4 and 4.8 of the Summary of Product Characteristics (SPC) with safety information following the assessment of the first PSUR. Relevant sections of the Package Leaflet (PL) were amended in line with the SPC. The details of the Danish local representative were also amended in the PL. The Annex II was updated following the assessment of the RMP (version 05).  Update of Summary of Product Characteristics and Package Leaflet        | 23/04/2009 | 02/06/2009 | SmPC, Annex<br>II and PL | Following the assessment of the 1st PSUR of fluticasone furoate, an increased number of adverse drug reactions related to eye disorders was reported, particularly for cataracts and glaucoma. On this basis, the CHMP recommended an update of the product information and the information was subsequently included in the SPC.   |

| II/0002 | Update of Detailed Description of the | 19/02/2009 | 07/04/2009 | Annex II | The DDPS has been updated (version 6.2) to reflect the   |
|---------|---------------------------------------|------------|------------|----------|--|
|         | Pharmacovigilance System (DDPS).      |            |            |          | change of the Qualified Person for Pharmacovigilance     |
|         |                                       |            |            |          | (QPPV) as well as to notify other changes to the DDPS    |
|         | Changes to QPPV                       |            |            |          | performed since the last approved version. Consequently, |
|         | Update of DDPS (Pharmacovigilance)    |            |            |          | Annex II has been updated using the standard text        |
|         |                                       |            |            |          | including the new version number of the agreed DDPS.     |