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ICH E21 Guideline on inclusion of pregnant and breastfeeding individuals in clinical trials

Step 2b

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INTERNATIONAL COUNCIL FOR HARMONISATION OF TECHNICAL REQUIREMENTS FOR PHARMACEUTICALS FOR HUMAN USE

ICH HARMONISED GUIDELINE

INCLUSION OF PREGNANT AND BREASTFEEDING INDIVIDUALS IN CLINICAL TRIALS E21

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ICH HARMONISED GUIDELINE

INCLUSION OF PREGNANT AND BREASTFEEDING INDIVIDUALS IN CLINICAL TRIALS

E21

ICH Consensus Guideline

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1 1. INTRODUCTION

2 1.1 Objective

- 3 The objective of this guideline is to provide recommendations for the appropriate inclusion
- 4 and/or retention of pregnant and/or breastfeeding individuals in clinical trials and facilitate the
- 5 generation of robust clinical data that allow for evidence-based decision making on the safe
- and effective use of medicinal products by these individuals and their healthcare providers
- 7 (HCPs).

8 **1.2 Scope**

- 9 The scope of this guideline includes pre- and postmarketing clinical trials of investigational
- products (see ICH E6(R3)) for indications in the general population and indications specific to
- pregnant or breastfeeding individuals.
- 12 In principle, inclusion of pregnant and breastfeeding individuals in clinical trials should be
- 13 considered for all products where individuals of childbearing potential are among the
- anticipated user population. It is especially important for conditions where there is high unmet
- medical need for treatment in pregnancy or while breastfeeding; however, the scope of this
- 16 guideline is not limited to these scenarios.

1.3 Background

- Many individuals who are pregnant or breastfeeding have acute or chronic medical conditions
- 19 (including physical and/or mental health conditions that occur or may be exacerbated during
- pregnancy and the postpartum period) that require new, ongoing, or preventative treatment(s).
- 21 Physiological changes during pregnancy can also have an impact on the pharmacokinetics (PK)
- 22 and/or pharmacodynamics (PD) of a medicinal product and there may be a need to modify the
- 23 dosage of medicinal products in pregnant individuals.
- 24 Pregnant and breastfeeding individuals are often excluded from clinical trials and those who
- become pregnant while participating in a clinical trial are frequently discontinued from the
- 26 clinical trial. As a result, pregnancy- as well as breastfeeding-specific information in the
- 27 product labeling on benefits and risks of medicinal product use is, at best, sparse and treatment
- decisions need to be made in the absence of this information. This lack of data has the following
- 29 potential consequences for pregnant and breastfeeding individuals:

- HCPs and/or patients avoiding or discontinuing indicated treatments leading to exacerbation of the condition or harm to the patient, pregnancy, or the child;
- HCPs and/or patients inadvertently choosing treatments harmful to the patient, pregnancy, or the child;
- Use of a dose or treatment regimen that is sub- or supra-therapeutic, leading to increased risk for under-treatment and/or adverse reactions;
- Avoidance or premature discontinuation of breastfeeding, or discontinuation of indicated treatment to allow for breastfeeding.
- 38 The potential magnitude of the public health impact of these negative consequences is considerable.

2. GENERAL PRINCIPLES

- 41 This guideline recommends that medicinal product use in pregnancy and/or breastfeeding
- 42 receives careful consideration and is incorporated into planning throughout investigational
- 43 product development from nonclinical studies through post-approval use of the product.
- 44 Proactive planning for obtaining data related to use in pregnancy and/or breastfeeding through
- 45 nonclinical and clinical studies (or the rationale for not obtaining data) should be done from
- 46 the early stages of formulating the development strategy for the investigational product.
- 47 Sponsors of drug development programs and clinical trials are encouraged to consider
- strategies to generate data that support informed decision-making on the safety, dosing, and
- 49 efficacy of the medicinal product's use during pregnancy and breastfeeding. Sponsors are
- recommended to consult with regulatory authorities as early as possible and as needed
- 51 throughout the investigational product development process regarding the plans for the
- 52 participation of pregnant and/or breastfeeding individuals in clinical trials. Every effort should
- 53 be made to reduce the burden of study procedures on pregnant and breastfeeding study
- 54 participants and it is essential to avoid any undue influence or coercion when pregnant or
- breastfeeding individuals are included or planned to be included in clinical trials. Early
- engagement with appropriate stakeholders, including patients, provides opportunities to
- address all relevant aspects of these clinical trials.

- Assessing the safety in pregnant and breastfeeding individuals is complex as there are potential
- impacts on the fetus and breastfed child to consider. When considering including pregnant or
- 60 breastfeeding individuals in clinical trials, it is important to evaluate the risks and benefits
- based on all available data, ensure that risks have been appropriately mitigated, and plan studies
- 62 that can yield scientifically robust data (see Sections 4.1.2 and 5.1.1).
- 63 Collection of data pertinent to use of an investigational product in pregnant and breastfeeding
- 64 individuals should continue into the postmarketing period. Ongoing safety monitoring of
- product use in these populations in the postmarketing period contributes to the identification
- of safety signals, especially for rare or delayed outcomes, that are unlikely to be thoroughly
- addressed in pre-authorization clinical trials. Real-world data (RWD) used to generate
- real-world evidence (RWE) can be helpful in assessing the usage and potential benefits or risks
- of an investigational product in pregnant and breastfeeding individuals.
- 70 Ongoing assessment of an investigational product during pregnancy and breastfeeding may
- draw from a variety of data sources, such as pharmacovigilance-generated data, electronic
- health records, medical claims or health insurance databases, medicinal product or disease
- 73 registries, or other sources (such as digital health technologies). Because pregnancy and
- breastfeeding present unique issues when gathering RWD, such as mother-child linkage, it is
- encouraged to proactively prepare platforms for post-approval data collection and to collect
- 76 background information on population and disease-specific risks to assist with data
- 77 interpretation.

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- Available data and assessment of investigational product benefits and risks during pregnancy
- and breastfeeding are expected to be included and updated as necessary in labeling documents.
- 80 Any statements in the prescribing information regarding pregnancy outcomes should be based
- on and reflect the robustness and limitations of the data as well as consideration of baseline
- rates of the outcomes in the indicated population when known. Additional considerations for
- labeling are included in Appendix 1.

3. ETHICAL CONSIDERATIONS

- 85 Including pregnant and breastfeeding individuals in clinical trials to support safe and effective
- data-driven use of medicinal products is ethical and supported by the Declaration of Helsinki
- and ICH guidelines, specifically ICH E6(R3) and ICH E8(R1). In addition to the
- 88 responsibilities of the sponsor and regulatory authorities, Institutional Review Boards (IRBs)

or Ethics Committees (ECs) have responsibility for evaluating whether the risks of conducting the trial are reasonable in relation to anticipated benefits. Consideration should be given to the use of IRBs or ECs experienced in working with pregnant and breastfeeding participants. For protocols involving pregnant or breastfeeding individuals, this responsibility involves considerations for the participant, for their pregnancy, and the fetus or breastfed infant. Ensuring ethical conduct of the trial therefore requires additional considerations regarding any need for appropriate safeguards related to pregnancy or breastfeeding (including risk mitigation measures implemented in the protocol and stopping criteria), as well as additional considerations regarding informed consent (Sections 4.4 and 5.5).

4. PREGNANCY

4.1 Development Strategy

Sponsors should anticipate that the approach to include pregnant individuals in clinical trials will require careful assessment of benefits and risks that may evolve depending on multiple factors, including the stage of clinical development, the duration of treatment, the indication being sought, and the strength of the available evidence. In addition, the approach may differ based on the anticipated trimester of pregnancy of participants to be included in the clinical trial. This section of the guideline lays out considerations for incorporating these complexities into the development strategy of an investigational product.

4.1.1 Factors to Consider When Planning for Pregnancy Data Collection

- Incorporating evidence collection for pregnant individuals into the development strategy starts with considering the targeted condition, patient population, and existing treatments. In addition, sponsors should consider how pregnancy might affect the disease state (e.g., potential worsening of the disease/condition if under- or untreated), as well as how the patient's disease (and its treatment) could impact the pregnancy and its outcomes (e.g., the potential increase in risk of adverse pregnancy outcomes due to inadequate disease control). These considerations will influence the timing and the type of data to be collected (see Section 4.2).
- When the investigational product is likely to be used by individuals of child-bearing potential, collecting data on safety, efficacy, PK during pregnancy, and predicted exposure to the fetus is important to support informed decision-making. Data should be collected as early as possible and appropriately timed in product development. Sponsors are encouraged to evaluate and update the development strategy as new information or data become available.

120 121	Situations that represent an especially high medical need for such data collection include but are not limited to:
122	Public health emergencies;
123 124 125 126	• Diseases that, if left untreated, are likely to adversely affect the health of the pregnant individual, the outcome of the pregnancy, and/or the health of the fetus/child (e.g., certain autoimmune diseases such as systemic lupus erythematosus (SLE) or human immunodeficiency virus (HIV) infection);
127 128 129	 Diseases for which the available treatments are not satisfactory in pregnancy and/or are known to carry high risks for the pregnant individual and/or the fetus/child (e.g., known or suspected teratogenicity or increased risk of pregnancy loss).
130 131 132 133	In these scenarios, the development strategy should aim for early acquisition of data from pregnant individuals unless there exists justification for postponement. Sponsors should proceed proactively with activities to generate the data and evidence necessary to enable inclusion in clinical trials at a later stage.
134 135 136 137	Depending on the characteristics and pharmacology of the investigational product and/or the disease/condition and available data from other similar medicinal products, it may be considered appropriate to design studies that include participants for an entire pregnancy, any time during pregnancy, or certain pregnancy trimesters only (e.g., avoiding third trimester exposure for non-steroidal anti-inflammatory drugs).
139 140 141	Clinical trials of prenatal interventions intended to improve outcomes of the fetus/neonate are not the focus of this guideline, however the principles discussed in this guideline may still apply.
142	4.1.2 Evidence Needed to Support Inclusion of Pregnant Individuals in Clinical Trials
143 144 145 146 147	In alignment with the principles of ICH E8(R1), the approach to collecting data from pregnant individuals in clinical trials involves a systematic expansion of data collection across relevant sources and patient populations, guided by data-driven decisions to safeguard study participants. Development programs should aim to generate the nonclinical and clinical data necessary to enable the inclusion of pregnant participants in clinical trials at the appropriate
148	stage of clinical development.

- The data and evidence needed to support the decision to include pregnant individuals in a clinical trial or to enable ongoing participation of individuals who become pregnant will depend on a weight of evidence approach and consideration of the following:
- The indication and the intended population;
- Nonclinical data;

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- The prospect of benefit;
- The clinical pharmacology of the investigational product;
- Biological plausibility of harm due to pregnancy exposure;
- When during the pregnancy the investigational product would be administered;
- The novelty of the investigational product (i.e., the availability of data from molecular entities or treatments similar to the investigational product).
- In the development strategy, the plan for collection of clinical data should be informed by an integrated assessment of these factors.

Prior to proceeding to studies including pregnant individuals, the results from relevant nonclinical studies need to be evaluated. These studies may include the standard Developmental and Reproductive Toxicology (DART) studies (see ICH M3 and ICH S5), the standard battery of genotoxicity studies if relevant (see ICH S2), appropriately qualified/validated alternative tests, and any relevant modeling. It is necessary to assess the nonclinical studies on how informative these studies would be on the safety of the investigational product for the intended patient population and make necessary adjustments to the type of studies needed and/or the study design. For instance, the timing and/or necessity for DART studies may be influenced by the characteristics of the investigational product (such as biotechnology derived pharmaceuticals as outlined in ICH S6(R1)), the clinical indication (such as those covered by ICH S9), and/or the intended patient population (e.g., exposure during the third trimester or the first trimester). Nonclinical data evaluation should be further explored to understand any potential risk to a pregnancy. When risks are identified, further investigations may be warranted with modified reproductive toxicology studies to characterize

176	them further (e.g., studies that investigate risks to the embryonic period vs. fetal period,
177	duration of dosing).
178	In addition to gathering the nonclinical data needed to proceed to studies in pregnancy,
179	acquiring clinical data in non-pregnant individuals will also usually be necessary. Generally,
180	clinical data that support safety and prospect of benefit in non-pregnant study participants could
181	reasonably be expected to be applicable for pregnant individuals. The necessary quantity and
182	type of data from non-pregnant participants will typically be similar to the data needed for an
183	investigational product to proceed through clinical development.
184	When the necessary nonclinical and clinical data become available, the sponsor should perform
185	a benefit-risk assessment that incorporates all relevant information described above, using a
186	weight of evidence approach. The objective of this assessment should be to determine whether
187	the risks of proceeding with trials in pregnancy are reasonable given the anticipated benefits.
188	If the sponsor determines that proceeding with trials in pregnancy is not yet reasonable, they
189	should seek to obtain further data unless there is a rationale for not studying the investigational
190	product in pregnancy. If the sponsor determines that proceeding with trials in pregnancy is
191	appropriate, then the following approaches/actions (in no specific order) need to be considered
192	and/or incorporated into the development strategy:
193	• Recruitment of pregnant individuals into ongoing and/or subsequent clinical trials;
194	• Removal of mandatory contraception requirements in ongoing and/or subsequent
195	clinical trials;
196	• Ongoing participation of individuals who become pregnant during clinical trials;
197	• Implementation of study(ies) specifically designed to be conducted in pregnant
198	individuals if needed.
199 200	4.1.3 When All the Data Necessary to Support a Favorable Benefit-risk Assessment are Not Yet Available
201	Before reaching the point where it may be appropriate to incorporate pregnant individuals into
202	the clinical development program, clinical studies using the investigational product will
203	typically have mandatory contraception requirements. Sponsors should recognize and plan for
204	the fact that pregnancies can occur when the study population includes individuals of

- 205 childbearing potential even when rigorous approaches to mandatory contraception are 206 implemented. Implications for study design and implementation when an unintended 207 pregnancy occurs are discussed in Section 4.2.11.
- A decision will need to be made regarding potential continuation on the investigational product when pregnancies occur despite mandatory contraception. Such continuation may often be inappropriate, but there could be exceptions. Considerations in the decision making should include the following:
- Information obtained to date regarding the safety in pregnancy of the investigational product (nonclinical as well as any clinical findings);
 - The participant's current health status, including the pregnancy and the underlying health condition;
- Risks of suspending study treatment (e.g., possible exacerbation of the treated disease, suitability or teratogenicity of alternative treatments, or impact of the disease on pregnancy);
- Any potential loss of the possible benefit (effectiveness) that might be obtained from the study treatment (e.g., through improvements in the underlying condition).
- If the conclusion is for treatment with the investigational product to continue, then the participant should be reconsented as a pregnant participant.

4.1.4 When Existing Data Suggest a Safety Concern for Pregnancy

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If nonclinical and/or clinical data suggest that the investigational product is potentially harmful to the pregnant individual and/or the fetus, the sponsor may conclude that inclusion of pregnant individuals in clinical trials is initially not warranted. However, for some investigational products, the benefits of use in pregnancy may still outweigh the potential risks. Examples include situations where the target disease has a serious negative impact (e.g., diseases such as malaria, which are known to have adverse effects on both the mother and the fetus) or where available treatment(s) have a safety concern in pregnancy (e.g., methotrexate for SLE). In such cases, including pregnant individuals in the trial may be considered on a case-by-case basis. In determining whether that is appropriate, it is essential to consider what additional data are needed to characterize the benefit-risk and to explore whether any potential risks can be

- mitigated. Additionally, consideration should be given to the fact that medical needs and potential risks associated with the product may differ depending on the trimester of exposure.
- 236 4.1.5 Strategies for Obstetric Conditions
- For the development of investigational products intended for obstetric conditions (e.g.,
- pre-eclampsia or preterm birth), clinical trials in pregnant individuals are necessary to evaluate
- 239 the investigational product's efficacy, safety, and dosing. In these scenarios, the data needed to
- proceed in clinical development and support a marketing application will be specific to the
- 241 condition.

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4.2 Inclusion of Pregnant Individuals in Clinical Trials

- 243 This section applies to trials that allow inclusion of pregnant individuals and those designed to
- be conducted as stand-alone trials in pregnant individuals. When a trial conducted in
- 245 individuals of childbearing potential has no requirement for contraception, such a trial
- essentially enables inclusion of pregnant individuals. Acquiring data on medicinal products
- during early pregnancy is only likely to occur in trials that have no requirement for
- contraception. These trials will be important to help characterize the product's safety profile in
- pregnancy unless there is a good rationale for not doing so.

250 4.2.1 Study Design and Implementation

- 251 While this guideline focuses mainly on the inclusion of pregnant individuals in interventional
- clinical trials, other trial types may be acceptable if they are appropriate for inclusion of
- 253 pregnant individuals. The sponsor should carefully consider which study design would be most
- 254 appropriate for the evaluation of an investigational product in pregnant individuals.
- Additionally, the safety impact on the pregnancy by all products used within the trial (i.e., test
- and comparator products) should be considered.

257 4.2.2 Expertise Considerations

- 258 Given the specialist knowledge required for investigational product and disease impacts on
- pregnancy, embryo-fetal development, and neonatology, consultation with relevant specialist
- 260 (e.g., obstetrician or maternal fetal medicine specialist) should be available for study design
- and safety monitoring (e.g., Data Monitoring Committee or other safety oversight body) to help
- interpret any adverse events (AEs) reported during pregnancy.

263	4.2.3 Sample Size
264	Study designs should consider the number and proportion of pregnant individuals expected to
265	be enrolled in trials, taking into consideration expected withdrawal rates based on the target
266	population and trial conditions.
267	For clinical trials with non-obstetric indications, estimating the number of pregnant participants
268	can help determine assessable endpoints. The PK data during pregnancy to enable appropriate
269	dose estimates may be obtained in most cases. However, low participant numbers may limit
270	safety conclusions, especially for rare adverse outcomes like specific birth defects.
271	The number of participants required to determine an efficacy endpoint should be achieved by
272	design for clinical trials of investigational products used for obstetric indications or in trials
273	designed for pregnant individuals only.
274	4.2.4 Pharmacokinetics and Dosing Considerations
275	There may be a need to modify the dose or frequency of investigational product administration
276	during pregnancy.
277	The physiological changes that occur during pregnancy may affect absorption, distribution,
278	metabolism, and elimination of the product potentially leading to an altered PK/PD profile of
279	the investigational product. In addition, the extent of these physiological changes can vary over
280	the course of pregnancy, so PK/PD should be assessed during the different trimesters and
281	postpartum. Depending on the duration of treatment, PK/PD measures should be assessed from
282	the same participant wherever possible. The postpartum assessment period should be
283	sufficiently long to understand PK/PD changes until the return to pre-pregnancy state.
284	For clinical trials that include pregnant participants, it is essential to include in the protocol
285	whether pregnant participants should receive the same dose as non-pregnant participants or a
286	different dose. Dose adjustments may be needed for pregnant participants in cases where
287	efficacy becomes suboptimal because of insufficient systemic exposure, or where the
288	therapeutic index or safety margins are narrow. To initially estimate the dosage/dosing regimen
289	for pregnant participants, clinical and dose-exposure data from non-pregnant participants could
290	be considered. Modeling approaches, such as physiologically based pharmacokinetics (PBPK)
291	modeling, which accounts for the PK alterations in pregnancy, may help to estimate the dosing
292	strategy. Any observed PK alterations in pregnant participants, exposure-response analysis, and

- population PK analysis, all provide important information for proper dose selection for
 pregnant participants.
- The dosing strategy for pregnant participants should be based on all the available evidence at
- 296 the stage of the clinical development program. The proposed dosing strategy should be
- 297 confirmed or further revised based on the findings of the clinical trial (e.g., safety concerns in
- 298 the trial and the clinical impact of overexposure or underexposure).

4.2.5 Fetal Exposure Assessment

- 300 Before including pregnant individuals, predicting the extent of fetal exposure may be helpful
- 301 for benefit-risk assessment. In the absence of data, risk assessments should assume a certain
- degree of fetal exposure. Currently, it is challenging to evaluate fetal exposure with available
- methods such as umbilical cord blood sampling. However, PBPK modeling could be a useful
- 304 option for estimating fetal exposure. Despite the limitations, fetal exposure data could
- 305 contribute to the overall pharmacologic and safety profile of the investigational product in
- fetuses and infants.

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4.2.6 Endpoints and Outcomes

- Pregnant participants should be evaluated with the same efficacy, safety, PK, and PD endpoints
- as those in the general study population, with the same frequency of evaluation whenever
- feasible (for information on analysis, see Section 4.2.10). Additional endpoints may also be
- needed for pregnant participants (e.g., PK/PD data). When the planned method to measure the
- endpoint may present a risk in pregnancy (e.g., CT scans), the participant should be followed
- 313 for safety or efficacy using alternative methods when available. Considerations regarding the
- 314 type of data to be collected are similar whether the participant is enrolled while pregnant or
- becomes pregnant during trial participation.

4.2.7 Assessments and Data Collection for Pregnant Participants

- 317 Pregnancy-related assessments should be specified in the protocol and include those that are
- impacted by the disease.
- 319 Standard general recommendations on safety evaluation such as classification, assessment, and
- 320 reporting of AEs (i.e., ICH E2A, ICH E2F, ICH E6(R3), ICH E8(R1)) apply to studies
- 321 including pregnant participants. The safety assessment considerations in this section and in
- 322 Appendix 2 apply in addition to standard assessments. Furthermore, a plan to follow and collect

323 324	investigational product on maternal and fetal/infant/child health. How this is best achieved will
325	need to be considered on a study specific basis, and depends on several factors, including but
326	not limited to:
<i>5</i> 20	not mined to.
327	• The known properties of the investigational product;
328	• The known or potential safety risks of other investigational products in the same class,
329	including emerging data;
330	• The timing and extent of exposure during gestation (see also Section 4.2.5);
331	• Availability and appropriateness of additional methodologies focused on assessment of
332	gestational/fetal/infant/child health;
333	• The burden of additional assessments on the pregnant participant and the
334	newborn/infant/child.
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335	Where possible, additional information should be collected to aid in the interpretation of the
336	safety profile. These data may provide context where risks to pregnancy associated with the
337	underlying disease or other intrinsic or extrinsic factors are well-established (see Appendix 2).
338	Outcomes and data parameters reported should include precise definitions, as well as their
339	source(s).
340	Local routine pregnancy monitoring for trial participants may be part of study-specific
341	assessments. These may include prenatal and postpartum follow-up visits, neonatal
342	consultations, ultrasound scans, and blood and urine tests.
343	When feasible, appropriate, and allowed by local regulations, it may improve clinical
344	accessibility for the study participant to align and/or combine study visits with regular
345	pregnancy-related clinical visits, employ mobile study visits, or virtual (telemedicine) study
346	visits.
347	4.2.8 Assessments and Data Collection for Infants
348	The duration of follow-up should be considered on a case-by-case basis and will depend on the
349	investigational product's half-life, indication, nonclinical data, mechanism of action, timing
350	and duration of exposure, and time to manifestation of outcomes of interest, taking into

consideration that birth defects and functional or neurodevelopmental disorders may be diagnosed beyond birth. Infant characteristics at birth and outcomes in the neonatal period to be considered are included in Appendix 2. It is recognized that the follow-up may extend until past the clinical trial completion date. Sponsors should ensure a mechanism for such follow-up is in place. Options may include subgroup-specific safety follow-up studies, enrollment in existing programs such as pregnancy registries, or other appropriate methods to ensure longer-term data collection on infant outcomes.

4.2.9 Safety Monitoring

- Participants should be closely monitored for pregnancy-related AEs, with appropriate management plans if required. The impact of the investigational product on the health of the pregnancy and infant may not be fully revealed during a clinical trial. Depending on the investigational product and trial design, follow-up may be needed beyond the duration of the trial. Appropriate mechanisms for such follow-up should be considered.
 - Provision for suspending or discontinuing investigational product for pregnant participants should be considered in the event of an emerging pregnancy-related safety signal. Sources for the detection of a signal could include clinical trials and post-trial follow-up, from clinical use during pregnancy or pediatric use, or published data, if applicable.

4.2.10 Analysis and Interpretation

- Data on efficacy, PK, and safety for pregnant individuals can help inform conclusions regarding whether the efficacy, dosing, and safety of the investigational product in pregnant individuals are similar to the general population. Clinical trial data even from a small sample size may contribute important information for product labeling. In addition, PK data from a small set of pregnant participants can help to reinforce data from models approximating exposure in the pregnant population at large. However, care should be taken when analyzing clinical trial results in small subpopulations, such as pregnant individuals, as this may lead to difficulty with interpreting adverse pregnancy outcomes.
- Given that the indication for treatment (i.e., the underlying disease or condition) may be harmful to the pregnancy or embryo-fetal development, the pregnancy-related outcomes to be measured should be assessed in the context of known impacts of the disease on pregnancy and the fetus (e.g., congenital malformation in diabetes). Insight into the efficacy of the product in treating the underlying health condition in that case will be accompanied by insight into

382	whether and how treating the underlying health condition with the investigational product
383	benefits the pregnancy.
384 385	Interpretation of the causality of AEs in the infant exposed to investigational product <i>in utero</i> should be made with caution in instances where the sample size is small or if there is no control of the causality of AEs in the infant exposed to investigational product <i>in utero</i> should be made with caution in instances where the sample size is small or if there is no control of the causality of AEs in the infant exposed to investigational product <i>in utero</i> should be made with caution in instances where the sample size is small or if there is no control of the causality of AEs in the infant exposed to investigational product <i>in utero</i> should be made with caution in instances where the sample size is small or if there is no control of the causality of AEs in the infant exposed to investigational product <i>in utero</i> should be made with caution in instances where the sample size is small or if there is no control of the causality of AEs in the causality
386	arm. Possible confounders should also be considered. Additionally, the pregnancy trimester of
387 388	exposure should be considered when evaluating any associations between exposure and outcome, (e.g., neural tube defects are unlikely to result from third trimester exposures).
389	External reference rates of adverse pregnancy outcomes in the general population may be
390	helpful to provide context. However, disease-specific pregnancy registries or observational
391	studies may be more informative.
392 393	4.2.11 Considerations for Pregnancies Occurring During a Clinical Trial With Mandatory Contraception
394	In trials with mandatory contraception, as noted in Section 4.1.3, pregnancies do still occur. In
395	view of this, sponsors are encouraged to design protocols which:
396 397 398	 Allow as appropriate, the option of remaining in the trial with suspension of investigational product for the duration of the pregnancy, or earlier resumption once data to support resumption of investigational product are available;
399 400 401	2. In some cases where pregnancy occurred, allow the option of continuing on treatment after reconsenting (see Section 4.1.3 for considerations as to when this might be appropriate);
402 403	3. For both situations above, provide for additional data collection (e.g., PK, PD, and additional safety monitoring, see Appendix 2);
404	4. Specify whether and when unblinding would be expected. A participant becoming
405	pregnant should not automatically lead to the unblinding of the participant's treatment
406	assignment.
407	4.3 Recruitment and Retention of Pregnant Individuals in Clinical Trials
408	The general principles for recruitment outlined in ICH E6(R3) apply for clinical trials including
409	pregnant individuals.

410	Pregnancy is a time when social and/or family interests are enhanced compared to the health
411	of a non-pregnant individual. Such interests may influence a pregnant individual's autonomy
412	and either unduly encourage or deter their participation in a clinical trial.
413	Increasing wider awareness of opportunities and considerations around participating in clinical
414	trials while pregnant is recommended. Providing detailed information on the proposed study
415	and its potential impact on future pregnant individuals with the same condition can help address
416	concerns and improve recruitment for these trials.
417	Engaging with patients' advocacy groups, organizations managing disease specific registries
418	and clinicians experienced in conducting research in pregnant individuals before clinical trial
419	initiation may help reduce challenges to recruitment or barriers to participation for specific
420	disease areas and/or identify opportunities for reducing burden for pregnant participants. Early
421	engagement with relevant stakeholders may help recruitment in several ways:
422	Involving potential participants and other stakeholders such as relevant healthcare
423	teams (e.g., obstetric and maternal-fetal medicine professionals) early in the study
424	design stages, could provide input on patient-orientated outcomes of interest and/or
425	reducing burdens for inclusion of pregnant individuals in clinical trials (see
426	Section 4.3.2);
427	Consideration of cultural differences regarding aspects of the birth, cord blood, and
428	placenta (and use of placental samples) may identify important aspects;
429	• Engaging HCPs familiar with the community (e.g., midwives, community [home
430	health] nurses, or prenatal care providers) may help recruitment (e.g., introducing trial
431	information or asking for contact information to follow-up);
432	Involving healthcare teams relevant to pregnancy could enable education of HCPs
433	about the value of their patients participating in research on conditions which may affect
434	pregnancy and health of the future child, to address any concerns and to encourage
435	participation;
436	• Early consideration of how and when to engage with potential participants may enhance
437	the ability to recruit pregnant individuals (including those at a particular trimester of
438	pregnancy) to relevant clinical trials and may enable best use of sponsor resources.

439	The additional time required for follow-up of pregnancy and infant outcomes, may mean that
440	additional efforts are needed to support retention of participants such as: maintaining contact
441	information, discussing potential barriers and facilitators to study participation at every visit
442	(e.g., time constraints, financial burden, or availability of study personnel to answer questions).
443	4.3.1 Recruitment of Pregnant Individuals for Clinical Trials
444	Where available, local clinical research networks for obstetric care may help identify potential
445	study centers with expertise in the conditions under investigation, including ongoing care
446	during pregnancy. Appropriate use of electronic health records may help to identify patients,
447	but sponsors/investigators may need to consider possible issues regarding confidentiality (see
448	ICH E6(R3)) and misidentification (e.g., due to pregnancy loss). If recruited through obstetric
449	clinics or electronic healthcare records, consideration should be given to local privacy laws
450	regarding disclosing pregnancy status.
451	Recruitment at earlier timepoints of pregnancy may require different approaches as first
452	trimester pregnancies may be difficult to identify through electronic health records or
453	obstetric/antenatal care units. Reaching out to specialized care physicians with educational
454	material about a potential clinical trial in this target population may help recruitment of
455	participants early in pregnancy. Studies in early pregnancy could include individuals who have
456	been exposed to an investigational product in routine clinical care or who become pregnant in
457	a trial (see Section 4.1.3).
458	4.3.2 Reducing Burden and Harm on Pregnant Individuals in Clinical Trials
459	Every effort should be made to assess the potential impact of study procedures to reduce burden
460	on pregnant participants, which supports retention in the clinical trial and may minimize
461	missing data. The impact of study procedures on the birth plan and delivery should be
462	minimized.
463	Early identification of study procedures that are not applicable or could pose unacceptable risks
464	during pregnancy may enable use of alternative monitoring procedures and/or flexibility in trial
465	protocols. For instance, the protocol may need to allow for pregnant individuals to reduce or
466	suspend study assessments that are not necessary (e.g., pregnancy testing), or assessments
467	associated with additional risks to the fetus (e.g., X-rays, teratogenic rescue medications used
468	in the protocol, or medication adjustments) until their pregnancy outcome has occurred.

469 470	Allowing some flexibility in timing of trial procedures may help address additional considerations specific to pregnancy (e.g., nausea and vomiting in early pregnancy, additional
471	monitoring requirements with high-risk pregnancies) and may enhance adherence to protocols.
472	The rationale for any extra visits in the context of the study should be explained to the
473	participant along with how the investigator and their other medical care specialists will work
474	together to deliver the participant's care plan.
475	4.4 Informed Consent for Studies with Pregnant Participants
476	Informed consent of all participants should follow the usual process (see ICH E6(R3)), with
477	appropriate adaptations for pregnant participants. The primary consent for participation in
478	clinical trials should clearly state whether ongoing participation will be allowed during
479	pregnancy and, if so, under what conditions.
480	Depending on the study design, informed consent could include focusing on the pregnancy
481	aspects in the form of supplemental informed consent for participants who:
482	Are already pregnant;
483	• Could become pregnant during clinical trials in which contraception is not mandated;
184	• Have a pregnancy during a trial requiring mandatory contraception and need to
485	reconsent regarding pregnancy-related information if they wish to remain in the trial on
486	treatment during the pregnancy.
487	The consent form should reflect the potential benefits and risks of the investigational product
488	as applicable in the intended pregnancy trimester(s) of exposure. This may be especially
489	pertinent if recruitment of participants at various stages of pregnancy is part of the study design.
490	Information should be provided to participants in terms of the potential benefits and risks to
491	the individual and the fetus/infant/child of taking or not taking study medication and
492	assessments performed during the study. Local guidance on any additional consent
493	requirements should be followed as well as requirements for informed consent for pregnant
494	minors. IRBs and ECs experienced in this patient population may also advise regarding the
495	appropriateness of any proposed compensation for study participants.

The consent process should seek consent on follow-up of the pregnancy/infant/child. This may include information on the planned duration of follow-up and any additional data sources that may be used. The information provided to the patient and HCPs should make it clear how study procedures will be handled in the case of uncomplicated and complicated deliveries and that clinical care takes precedence over the study protocol. The informed consent should also include release of medical records to obtain relevant information on the course of the medical condition, the pregnancy, obstetric history, and follow-up information on the infant. It should also explain confidentiality of the study data and possible implications of participation (e.g., revealing of underlying genetic conditions that otherwise would not have been identified or follow-up of the exposed child may disclose underlying maternal conditions).

Participants who have a confirmed pregnancy while enrolled in a clinical trial should be provided with information to make an informed decision for both themselves and their fetus regarding options as per protocol for (1) staying on study investigational product, (2) suspending investigational product until later in or after pregnancy (3) discontinuing the investigational product and moving to pregnancy follow-up or (4) withdrawing from the study. The information provided to participants should clearly explain any changes to the protocol that are needed to allow for these individuals to reduce or suspend relevant study assessments until their pregnancy outcome occurs. Participants who withdraw from the study should understand the importance of follow-up of their pregnancy outcome and be encouraged to consent to collection of this data.

- Additional circumstances related to clinical trials in pregnancy where participants should be reconsented include:
- When mandatory contraceptive requirements of the trial have been removed while the trial is ongoing (see Sections 4.1.2 and 4.2.11);
 - When new information changes the assessment between benefits and risks for the pregnant participant or their fetus.

5. BREASTFEEDING

5.1 Development Strategy

The benefit-risk considerations for medicinal product use during breastfeeding involve multiple factors, such as the amount of investigational product present in breastmilk, the extent

526	of absorption by the child, the potential benefits and risks of the medicine for the patient and
527	the breastfed child, available treatment alternatives, the benefits of breastfeeding, and available
528	alternatives to breastfeeding.
529	Sections 5.2 and 5.3 of this guideline discuss the following:
530	Obtaining information on the transfer of investigational product into breastmilk (either
531	without or with investigational product exposure to the infant as discussed in
532	Sections 5.2.1 and 5.2.2, respectively);
533	Subsequently, inclusion of breastfeeding individuals in clinical trials in the general
534	population after the investigational product's characteristics related to breastfeeding
535	have been determined (as discussed in Section 5.3).
536	The clinical development strategy for investigational product use in breastfeeding should be
537	tailored to the stage of development and existing knowledge about the investigational product.
538	Since investigational product exposure to the infant can be avoided by replacing breastmilk
539	with formula or other supplemental nutrition, whether and, if so, when to allow such exposure
540	during development must be carefully considered.
541	Sponsors should anticipate if, and when, clinical trials involving breastfeeding individuals may
542	be initiated and plan to conduct studies to gather information on exposure levels and effects on
543	a breastfed child if needed as early as possible in development. Early planning for when and
544	how to obtain the relevant data may enable optimizing the clinical development strategy of the
545	investigational product. Of note, there may still be a need to understand how the product may
546	affect lactation or the breastfed infant, even if the medicinal product is not to be used in
547	pregnancy.
548	The approach to collecting data related to breastfeeding should consider the level of
549	information available on the investigational product (e.g., physicochemical characteristics,
550	mechanism of entry into breastmilk, data from nonclinical studies such as pre- and postnatal
551	development and juvenile toxicology studies, and infant factors, such as differences due to
552	infant metabolic pathways). In addition, there could be other data sources to consider such as
553	use of the investigational product in pediatric patients. Early identification of available data
554	and knowledge gaps should be addressed to establish the safe and effective use of medicinal

products for breastfeeding individuals.

Individuals participating in efficacy clinical trials of the investigational product during pregnancy may be willing to participate in lactation studies. Data from such participants can provide important information for breastfeeding in the immediate postpartum period. Participants who are not intending to breastfeed could participate in lactation studies with no planned infant exposure.

5.1.1 Evidence Generation Planning Related to Investigational Product Use and Breastfeeding

- Developing a strategy to collect data relevant to breastfeeding can be broadly categorized into the following steps: (1) determine the concentration of investigational product in breastmilk (relative to maternal therapeutic blood levels), (2) use breastmilk concentration data for estimation of the daily infant dose and relative infant dose, and (3) collect infant exposure, safety, and benefit data, as applicable. Together this information is important in determining the appropriate breastfeeding and/or treatment advice.
- Lactation studies (see Section 5.2) which evaluate investigational product levels in breastmilk can contribute to an understanding of any potential effects on the breastfed infant and may be appropriate to be conducted as a clinical pharmacology trial. Studies which allow exposure of the child to the investigational product through breastmilk enable evaluation of whether the presence of the investigational product in milk has any impact on the breastfed infant.
- Milk composition and quantity may vary during lactation, with different patterns of breastfeeding and age of the child, which may affect the amount of investigational product to which the infant is exposed. Therefore, inclusion of individuals at different stages of breastfeeding is encouraged. Additionally, colostrum, foremilk, and hindmilk vary in composition, which should be considered when PK analysis of breastmilk is being planned.

5.1.2 Nonclinical Considerations

Nonclinical studies may be used to generate data on lactational exposure to an investigational product. The standard pre- and postnatal development (PPND) study (see ICH S5) exposes the pups both during gestation and lactation. This study provides information on the effects of the investigational product on both the pups (e.g., adverse effects on pups) and lactation (e.g., milk quality and quantity) that can characterize the potential risk(s) to a neonate. A challenge of this study is understanding whether any neonatal effects observed were related to the gestational or lactational exposure. To distinguish this, a juvenile toxicology study with direct dosing of juvenile animals can be used to further characterize potential risks (see ICH S11).

587	Qualified/validated alternative assays (ICH S5) may also be used to generate lactational
588	exposure data. In addition, appropriate use of modeling techniques, such as PBPK modeling,
589	may provide insights into likely levels of an investigational product in breast milk, and
590	subsequent infant exposure, absorption, and metabolism (see ICH M15).
591	5.2 Lactation Studies
592	5.2.1 Lactation Studies Assessing Investigational Product Levels in Maternal Milk
593	This section discusses lactation studies that assess product levels in maternal milk with no
594	infant exposure to investigational product through breastmilk (i.e., maternal-only studies).
595	These studies are usually conducted in breastfeeding patients but, when necessary, can be
596	conducted in breastfeeding healthy volunteers. In both cases, the participant must pump and
597	discard the breastmilk. The data collected from these studies are considered a prerequisite for
598	the planning of the studies described in Section 5.3.
599	Individuals could be enrolled once they have decided to stop breastfeeding their child or are
600	willing to interrupt breastfeeding during the study and until all investigational product would
601	be expected to be cleared from the breastmilk and maternal blood.
602	Lactation studies evaluating investigational product levels in breastmilk provide detailed
603	information about the amount/concentration and duration of an investigational product in
604	breastmilk. The data can also be used to model the likely exposure levels in the infant (e.g.,
605	amount of investigational product in milk and predicted absorption in the infant). As they are
606	usually short in duration, these studies could be designed as stand-alone studies or as an initial
607	sub-study of a larger trial that at some later point intends to enroll or include breastfeeding
608	participants.
609	Lactation studies that assess product levels in maternal milk only can also be conducted in
610	breastfeeding individuals who are taking a medicinal product as part of clinical care.
611	5.2.2 Lactation Studies Assessing Exposure in Breastfed Infants
612	This section discusses lactation studies that assess investigational product levels in the maternal
613	milk as well as in the infant exposed through breastmilk. These studies include both mother
614	and infant as part of the study population (i.e., mother-infant pair studies). This scenario
615	includes opportunistic studies which recruit patients who are already on a marketed medication
616	based on clinical need and choose to continue treatment during breastfeeding, stand-alone

lactation studies, and lactation studies conducted within clinical trials where breastfeeding

618	individuals are enrolled along with the general population.
619	For lactation studies in which the infant is exposed to the investigational product, that are not
620	opportunistic in design, data are needed to support a favorable benefit-risk profile in the infant.
621	Such data may include nonclinical data, lactation data on the amount of investigational product
622	in milk, and modeling to predict absorption in the infant. Uptake of the investigational product
623	in the infant needs to be evaluated, using paired sampling from mothers and their breastfed
624	infant. The study should evaluate whether the amount absorbed may have short and/or

long-term implications for the infant as appropriate.

5.3 Inclusion of Breastfeeding Individuals in Clinical Trials

The inclusion of breastfeeding individuals in clinical trials for indications in the general population may be permissible with the appropriate data available and considerations for benefit-risk for both the mother and the child. Lactation studies can support the benefit-risk profile of breastfeeding to the infant while participants are in the trial if they demonstrate no clinically relevant transfer of the investigational product into breastmilk or when there is no clinically relevant absorption in the infant. Inclusion of breastfeeding individuals in clinical trials may also be permissible when the infant has a potential benefit from investigational product exposure that outweighs the potential risks.

Depending on the numbers of participants, the inclusion of breastfeeding individuals in clinical trials may allow for evaluations of whether dose, efficacy, and safety are similar to the non-breastfeeding population. Additionally, it could be evaluated whether the investigational product affects breastfeeding.

5.3.1 Study Design

Clinical trials that enroll breastfeeding individuals should minimize the potential risks to the breastfed infant and assess safety in exposed infants. When there is reasonable scientific assumption that the investigational product may not be meaningfully absorbed from breastmilk or the potential benefits for mother and infant outweigh any potential risk to the infant, the protocol could allow a choice for participants to keep breastfeeding. Data collection should be planned such that the burden of trial participation remains manageable for trial participants (see Section 5.4.2).

647	Given the specialist knowledge required for investigational product and disease impacts on
648	breastfeeding, postpartum physiology, and child health, consultation with relevant specialists
649	(e.g., specialists in breastfeeding and breastfeeding support) should be available for study
650	design and safety monitoring (e.g., Data Monitoring Committee or other safety oversight body)
651	to help interpret any AEs reported during the study.
652	As evaluation of the child's well-being and adequate development is crucial in these situations,
653	the presence of neonatologist/pediatricians in the study teams is also recommended.
654	5.3.2 Pharmacokinetics and Dosing Considerations
655	As there are physiological changes in the postpartum period (e.g., reduced plasma volume
656	during lactation), albeit to a lesser extent than during pregnancy and which progressively
657	normalize over time, the collection of PK data from the breastfeeding participant at various
658	stages of breastfeeding should be considered at least until return to pre-pregnancy status.
659	In general, changes in dosing regimen during breastfeeding are not expected to be necessary.
660	However, if dosages have been adjusted due to pregnancy, time to readjust to pre-pregnancy
661	doses may need to be considered. In addition, studies to assess alterations to the breastfeeding
662	strategy (e.g., timing of breastfeeding the child), in relation to dose regimen should be
663	considered, if applicable.
664	5.3.3 General Outcomes Related to Breastfeeding
665	When enrolled in clinical trials along with the general population, study participants who are
666	breastfeeding should, wherever possible, be evaluated with the same efficacy outcomes as those
667	in the general study population, with the same endpoints and frequency of evaluation.
668	If the planned assessment may expose a breastfed child to a specific risk (e.g., effect of
669	radiological contrast dye on the milk) alternative assessments or endpoints should be
670	considered or the breastmilk could be temporarily discarded for the required time to avoid
671	exposing the child to a specific risk.
672	Outcomes of interest related to breastfeeding should be selected with relevance for
673	investigational product labeling and health outcomes of mother and infant. Impact on lactation
674	itself should be evaluated (e.g., effects on breastmilk production). Data on lactation stage or
675	the schedule of breastfeeding, child age, other medical conditions of the mother or infant, and

676	concomitant therapies that could affect breastfeeding or have an impact on the infant should be
377	recorded.
270	Sparse PK sampling approaches can be useful to supplement detailed PK data to enlarge the
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379	patient population studied. Even when some trial data are available on the effects of the
380	investigational product on breastmilk production, the levels in the breastmilk, and the
381	absorption by the breastfed infant (when appropriate), it may be useful to collect data from
382	other breastfeeding study participants to enhance the dataset.
383	5.3.4 Safety Monitoring Related to Breastfeeding
384	Standard general recommendations on safety evaluation such as classification, assessment, and
385	reporting of AEs (i.e., ICH E2A, ICH E2F, ICH E6(R3), ICH E8(R1)) apply to studies
386	including breastfeeding individuals. In addition, the safety assessment considerations in this
387	section apply. When both the mother and the infant are exposed to the investigational product,
388	uptake of the product in the infant needs to be understood (or evaluated, if necessary), at
389	relevant timepoints. Where present, the study should evaluate whether the amount absorbed
390	may have short and/or long-term implications for the breastfed child (e.g., severity/frequency
391	of AEs or impact on growth and/or development, as appropriate). Depending on the specific
392	impact, a safety follow-up plan should be implemented.
393	The planned follow-up assessments should consider the general well-being of the child, as well
394	as any outcomes predicted from the pharmacologic effects and the safety profile of the
395	investigational product. Information from investigational products within the same class or
396	experience with use of the investigational product in pediatric populations may be helpful for
397	setting the safety follow-up plan. It should be considered whether monitoring of the effect on
398	lactation and the child may be needed beyond the duration of the trial.
399	Interpretation of the causality of AEs in the infant exposed to investigational product during
700	breastfeeding should be made with caution and take into consideration any medical condition
701	of the infant and other confounding factors (e.g., maternal diet, concomitant medicinal products
702	or need for supplemental nutrition with formula or other supplement), and any prior in utero

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exposure.

5.3.5 Discontinuation and Suspension of Treatment
The protocol should outline criteria for discontinuing breastfeeding in case of emerging safety
concerns to the breastfed child. Additionally, consideration should be given whether
adjustments to the breastfeeding strategy (e.g., timing or pump and discard) could serve as
effective measures to ensure infant safety, allowing the mother to continue participating in the
trial.
For studies involving breastfeeding participants, in addition to standard sources, any new safety
signal emerging from pediatric exposures should be considered (e.g., other or ongoing clinical
trials with the study investigational product(s)) as these might provide information relevant for
the exposed child.
5.4 Recruitment and Retention of Study Participants
5.4.1 Recruitment of Study Participants
Recruitment strategies for inclusion of breastfeeding participants may differ depending on
whether enrollment is for lactation studies or for clinical trials. Early consideration of how and
when to engage with potential participants may enhance the ability to recruit participants to
relevant studies to obtain clinically relevant information on investigational products in a timely
manner.
The following points should also be considered:
• Engaging patients and stakeholders in advance of recruitment to provide accurate,
relevant information on a specific trial may reduce concerns of potential participants
and their close family and/or social group, if applicable, about participating in research;
• Involving patients and other stakeholders such as relevant healthcare teams early in the
study design stages, could provide insights into how to better monitor and collect timely
information to enable any risk mitigation during the study to support recruitment and
retention of participants during the study;
• Providing education to HCPs about study participation for their patients and address
any concerns in order to encourage participation;

• Cultural differences regarding breastfeeding.

When an investigational product is to be used from the very early postpartum period, it could

733	be preferable to start screening procedures for patient enrollment during the pregnancy period
734	to be ready to potentially include the patient in the trial immediately after delivery. If screening
735	is started during pregnancy, some screening procedures may need to be repeated to confirm
736	eligibility before enrollment.
737	For clinical trials in which infants are exposed to investigational product through breastmilk,
738	recruitment efforts will need to include facilitating the understanding of benefits and risks
739	through educational materials for the mother and their families when appropriate and the
740	impact of trial participation on breastfeeding intentions. The purpose and types of study
741	procedures should be clearly explained to participants.
742	5.4.2 Reducing Burden on Participants
743	Flexibility can be incorporated into several aspects of the study to reduce the burden on
744	participants.
745	Early and avoidable discontinuation of participants can be mitigated by recognition and support
746	of the challenges of this period. To lessen the burden for participants, assessments required as
747	part of a study protocol may be integrated with information contained in records from standard
748	pediatric care visits where appropriate and feasible. Additional considerations to reduce burden
749	to study participation include:
750	• Quantities of breastmilk required for sample analysis should be minimized;
751	• Where appropriate, interventions for sampling infant blood should be minimized;
752	• Consideration should be given to providing breastmilk pumps for efficient milk
75 3	expression or use of alternative methods for sampling;
754	• Provision of care/activities for the child;
755	• If possible, and without compromising study integrity, provide real-time results to
756	participants in lactation studies evaluating investigational product levels in breastmilk,
757	to allow restarting of breastfeeding (if appropriate);
758	• It is recommended that participants collect and store samples or utilize home health
759	nurses, when appropriate;

- Encourage participants to pump and store breastmilk prior to dosing such that the infant can be fed for several hours to a day or more with pre-study milk;
- Lactation consultants (or their equivalent) can be used to help the participants continue to express sufficient quantities of milk during the clinical trial.

5.5 Informed Consent for Studies with Breastfeeding Participants

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- For informed consent the principles of ICH E6(R3) apply, and additional considerations for breastfeeding and lactation are outlined below.
- Depending on the study design, informed consent may need to consider the potential benefit and exposure risk to the mother and the infant, and risks related to study procedures for the mother and the infant (e.g., breastmilk sampling or blood draws). Consent should follow regional guidance related to parental consent. The consent should also include information on how clinical trial processes and procedures may impact breastfeeding and prioritizing participant and infant safety.
 - Participants enrolling in a lactation study should be informed that the primary purpose is to investigate the investigational product levels in the blood (i.e., maternal and may include infant) and breastmilk and the correlation between them. In a lactation study where the infant is not exposed to the investigational product, the participant should be advised about the duration that the investigational product will be present in breastmilk to avoid inadvertently exposing the breastfed child to the investigational product. The following should also be considered: timing of sampling and testing, duration of interruption of breastfeeding, the availability of nutritional alternatives to mother's milk, and conditions of their infant (e.g., prematurity) that may affect prioritizing breastmilk provision vs. research participation.
 - Additionally, depending on the study design, for studies that permit breastfeeding during exposure to the investigational product:
 - Up-to-date information about the investigational product and its clinical and nonclinical development should be made available, to support decisions regarding breastfeeding, especially in relation to investigational product transfer through breastmilk.
 - Local guidance on any additional consent requirements should be followed if an infant would be exposed to the investigational product through breastmilk.

789	• The informed consent should include follow-up plans for the infant, including the
790	frequency and type of safety assessments conducted, and access to infant medical
791	records, if appropriate.
792	• It may be appropriate for the informed consent to include release of information from
793	maternal medical records to obtain relevant information on the course of the medical
794	condition and the pregnancy.
795	There may be circumstances where participants should be reconsented (e.g., new information
796	that changes the assessment of benefits and/or risks of the investigational product for the
797	breastfeeding participant or the breastfed child).
798	IRBs and ECs experienced in this patient population may also advise regarding the
799	appropriateness of any proposed compensation for study participants.

6. APPENDICES

801	APPENDIX 1: CONSIDERATIONS FOR LABELING
802	Sources for information in product labeling include nonclinical data and clinical data such as
803	PK, PD, and dose data obtained through relevant studies and/or modeling and simulations,
804	clinical efficacy and safety trials, epidemiological studies, pregnancy registries, and
805	pharmacovigilance pertaining to pregnant and breastfeeding individuals.
806	When available, and depending on regional labeling guidances and subject to regulatory
807	review, the following information should be considered for inclusion in labeling:
808	Recommended dose during pregnancy and any dosage adjustments during pregnancy,
809	breastfeeding, and/or the postpartum period;
810	• The product's effects on the pregnancy (such as risk of miscarriage or pregnancy
811	complications);
812	• Risks of disease progression during pregnancy (e.g., potential worsening of the
813	disease/condition if under- or untreated);
814	• The potential for the product to cross the placenta;
815	• Effects on the fetus (such as risks of congenital malformation, effect on fetal growth,
816	and potential for long-term effects on the infant and the child);
817	• Extent of the product's presence in breastmilk and exposure of the breastfed infant;
818	• Effects of the product on lactation and on the breastfed child;
819	• Any adverse drug reactions or withdrawal symptoms in the neonate;
820	Any recommended measures to minimize a product's risk to pregnant and breastfeeding
821	individuals and to the fetus or the infant;
822	Any monitoring recommendations for pregnant and breastfeeding individuals and the
823	fetus or the infant;
824	• Any differences identified for the above items based on demographic, disease state, or
825	other subpopulations.

826 827	APPENDIX 2: ADDITIONAL OUTCOMES TO BE CONSIDERED IN CLINICAL TRIALS INCLUDING PREGNANT PARTICIPANTS
828	In addition to standard reporting requirements and Good Clinical Practice (GCP) (see
829	ICH E6(R3)), the following outcome parameters are to be considered, with attention to the
830	disease/condition being treated by the investigational product, investigational product
831	properties, duration of use, and therapeutic context.
832	Maternal and Gestational Outcomes of Interest:
833	Standard maternal and gestational measures of interest include pregnancy outcome, including
834	timing and underlying circumstances of pregnancy losses, (particularly if due to congenital
835	malformation), characteristics and gestational age at birth (e.g., cesarean section delivery or
836	preterm), and infant measurements at birth (e.g., weight).
837 838	In addition to these standard measures and where relevant, consideration should be given to the following:
839	• Identification of congenital malformation prenatally (e.g., fetal cardiac ultrasound);
840	Gestational/prenatal assessments and findings, including complications of pregnancy
841	(e.g., chorioamnionitis or intrauterine growth restriction);
842	Maternal conditions affecting gestational health (e.g., gestational diabetes, disease)
843	flares, or opportunistic infections);
844	• Obstetric history (e.g., miscarriages along with previous history of
845	preeclampsia/eclampsia, postpartum hemorrhage, caesarean section, or allergies to
846	specific medicinal products);
847	• Characteristics of childbirth including complications of labor (e.g., premature rupture
848	of membranes, method of delivery, stillbirth, or asphyxia);
849	• Placental pathology or notable placental abnormalities;
850	• Endpoints specific to multiple pregnancies, including chorionicity, zygosity, loss of one
851	or more fetuses in a higher-order multiple pregnancy, and conditions such as twin-twin
852	transfusion syndrome;

853	• Other relevant factors, e.g., use of folic acid, relevant paternal health factors, access to
854	and quality of prenatal care, or use of assisted reproduction (including donor
855	gametes/embryos).
856	Infant Characteristics at Birth:
857	Infant outcomes should include sex, gestational age at birth, infant weight at birth (e.g., small
858	for gestational age) and congenital malformations or other functional or morphological
859	abnormalities apparent at or immediately following birth.
860	Additional postnatal infant outcomes to be considered when relevant include:
861	Cardiovascular and respiratory examinations, including need for supplemental oxygen
862	or resuscitation;
863	• Developmental and functional assessments (e.g., APGAR or neurological assessment
864	(muscle tone, spontaneous activity)).
865	Outcomes in the Neonatal Period and Infant Follow-up:
866	Neonatal outcomes to consider when relevant within the first 28 days after birth include:
867	• Size- and growth-related assessments;
868	• Developmental (including neurologic) assessments;
869	• Feeding characteristics including use of breastmilk and/or formula, occurrence of
870	feeding difficulties, and gastrointestinal intolerances;
871	• Congenital malformations diagnosed in the neonatal period;
872	• Health of major organ systems (e.g., kidney or liver function);
873	• Postnatal infections or other health issues arising in the neonatal period including
874	hospitalizations.
875	Infant follow-up outcomes of interest will differ based on the maternal disease or disorder,
876	investigational product type, and gestational exposure. It should be considered that some

neurological and physical developmental delays or conditions may not be visible until later in
life.