

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

DRAFT ICH HARMONISED GUIDELINE

**DETECTION OF TOXICITY TO REPRODUCTION FOR HUMAN
PHARMACEUTICALS**

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1 **1 SCOPE OF THE GUIDELINE**

2 This guideline applies to pharmaceuticals, including biotechnology-derived pharmaceuticals,
3 vaccines (and their novel constitutive ingredients) for infectious diseases, and novel
4 excipients that are part of the final pharmaceutical product. It does not apply to cellular
5 therapies, gene therapies and tissue-engineered products. The methodological principles
6 (e.g., study design, dose selection and species selection) outlined in this guideline can also
7 apply to pharmaceuticals intended for the treatment of serious and life threatening diseases,
8 such as advanced malignancies (i.e., see ICH S9 (3)). This guideline should be read in
9 conjunction with ICH M3(R2) (1), ICH S6(R1) (2) and ICH S9 (3) regarding whether and
10 when non-clinical reproductive toxicity studies are warranted.

11

12 **2 INTRODUCTION & GENERAL PRINCIPLES**

13 The purpose of this guideline is to provide key considerations for developing a testing
14 strategy to identify hazard and characterize reproductive risk for human pharmaceuticals.
15 The guidance informs on the use of existing data and identifies potential study designs to
16 supplement available data to identify, assess, and convey risk. General concepts and
17 recommendations are provided that should be considered when interpreting study data and
18 making an assessment of reproductive risk in support of clinical development and marketing
19 approval.

20 To assess a human pharmaceutical’s effects on reproduction and development, the
21 information should generally include exposure of adult animals and the impact on all stages
22 of development from conception to sexual maturity. No guideline can provide sufficient
23 information to cover all possible cases, and flexibility in testing strategy is warranted.
24 Regardless of the pharmaceutical modality (see Glossary), key factors to consider when
25 developing an overall integrated testing strategy include:

- 26 • The anticipated pharmaceutical use in the target population (especially in relation to
27 reproductive potential and severity of disease);
- 28 • The formulation of the pharmaceutical and route(s) of administration intended for
29 humans;
- 30 • The use of any existing data on toxicity, pharmacodynamics, pharmacokinetics, and
31 similarity to other compounds in structure or activity;
- 32 • Selection of specific studies, test species/test system and dose levels.

33

34 These concepts are discussed in more detail throughout the guideline, which defines a
35 thoughtful approach for developing a testing strategy. This guideline recommends the use of
36 information about the pharmaceutical and the patient population in order to perform only
37 those studies essential to evaluate the stages (see below) for which there is insufficient
38 knowledge to inform about the risk to reproduction and development.

39 As appropriate, observations through one complete life cycle (i.e., from conception in one
40 generation through conception in the following generation) permit detection of immediate
41 and latent adverse effects. For the purposes of this guidance, gestation day 0 (GD 0; see
42 Glossary) is when positive evidence of mating is detected. The following stages of
43 reproduction are generally assessed:

44 A) Premating to conception (adult male and female reproductive functions, development
45 and maturation of gametes, mating behavior, fertilization).

46 B) Conception to implantation (adult female reproductive functions, preimplantation
47 development, implantation).

48 C) Implantation to closure of the hard palate (adult female reproductive functions,
49 embryonic development, major organ formation).

50 D) Closure of the hard palate to the end of pregnancy (adult female reproductive
51 functions, fetal development and growth, organ development and growth).

52 E) Birth to weaning (adult female reproductive functions, neonate adaptation to
53 extrauterine life, pre-weaning development and growth).

54 F) Weaning to sexual maturity (post-weaning development and growth, adaptation to
55 independent life, attainment of full sexual function).

56 The stages covered in individual studies are left to the discretion of the Sponsor, although
57 the timing of studies within the pharmaceutical development process is dependent on study
58 populations and phase of pharmaceutical development (see ICH M3(R2) (1), ICH S6(R1)
59 (2) and ICH S9 (3)).

60 This guideline also provides considerations for interpreting all available nonclinical
61 information as part of the risk characterization.

62 **3 STRATEGIES FOR REPRODUCTIVE TOXICITY ASSESSMENT**

63 **3.1 Considerations/Principles**

64 The initial step is to determine if reproductive toxicity testing for each of the various
65 reproductive stages is warranted and, if so, what are the most appropriate studies to conduct.
66 The considerations should include: a) the target patient population and duration of dosing, b)
67 the known pharmacology of the compound, c) the known toxicity of the compound, d) any
68 existing knowledge of the impact of the target(s) on reproductive risk (e.g., human and/or
69 animal genetics, or class effects), and e) data from *in vitro* and non-mammalian assays
70 (alternative assays, see Glossary) that could be relied upon to identify hazard and/or risk (see
71 Section 3.3.2). Approaches for qualifying and use of alternative assays in assessing
72 reproductive risk are discussed below (Sections 3.3.2 and 9.5). Generally, most alternative
73 assays being developed address endpoints related to Embryo-Fetal Development (EFD) and
74 are thus discussed in section 3.3.2. However, as new assays are developed for other
75 reproductive endpoints, they can be similarly deployed with appropriate qualification.

76 The experimental strategy to generate the data should consider minimizing the use of
77 animals. Alternative assays and/or *in vivo* studies with fewer animals can be used to identify
78 hazards in a tiered manner. Reductions in animal use can also be achieved by deferring
79 definitive EFD studies (see Section 9.4.3.3) until later in pharmaceutical development (see
80 below). Alternative assays can replace definitive assays in some circumstances where as in
81 others they can be used to defer traditional assays until later in development (see Section
82 3.3). An important component of the overall strategy is the timing for the additional
83 information to support ongoing clinical development (e.g., developmental toxicity (see
84 Glossary) data to support dosing women of childbearing potential).

85

86 Reproductive and developmental studies should in general be conducted according to Good
87 Laboratory Practice (GLP) as they will contribute to risk assessment. However, if a human
88 developmental or reproductive risk is defined during the conduct of a relevant non-GLP
89 study, repetition of the study to confirm the finding(s) under GLP conditions is not
90 warranted. Preliminary EmbryoFetal Development (pEFD; see Glossary) studies should be
91 conducted under high-quality scientific standards with data collection records readily
92 available or under GLP conditions. It is recognized that GLP compliance is not expected for
93 some study types, or aspects of some studies, employing specialized test systems or methods,
94 such as disease models or surrogate molecules (see Glossary), or literature. However, high
95 quality scientific standards should be applied, with data collection records readily available.
96 Areas of non-compliance should be identified and their significance evaluated relative to the
97 overall safety assessment.

98

99 **3.1.1 Target Patient Population/ Therapeutic Indication Considerations**

100 The patient population or therapeutic indication can influence the extent of reproductive
101 toxicity testing. For example:

- 102 • If the female patient population is post-menopausal there is no utility in evaluating
103 any of the reproduction stages;
- 104 • A pharmaceutical for use in an elderly male does not warrant conduct of studies to
105 evaluate stages E and F;
- 106 • If the disease indicates that reproductive toxicity will have minimal impact on the
107 usage of the pharmaceutical in the target population, studies evaluating only stages C
108 and D can be warranted;
- 109 • Short-term therapies under highly controlled settings.

110 **3.1.2 Pharmacology Considerations**

111 Before testing, it should be determined if the pharmacologic effects are incompatible with
112 fertility, normal EFD, or measurement of endpoints of the study being considered (e.g., a
113 general anesthetic and measurement of mating behavior). This assessment could be based
114 on data with other pharmaceuticals with similar pharmacology on the pathways affected, or

115 on knowledge of effects in humans with related genetic diseases. Based on these
116 considerations, sometimes no testing for a particular reproductive endpoint can be
117 warranted. In contrast, testing for only off-target effects can be warranted if the expected
118 pharmacologic effects on reproductive endpoints are non-adverse. Examples include
119 patients with a condition that mimics the target pharmacology who have normal
120 reproductive capability and healthy offspring; or when other pharmaceuticals have similar
121 pharmacology or pathways affected but have no demonstrated reproductive risk.

122 **3.1.3 Toxicity Considerations**

123 Repeat-dose toxicity studies with sexually mature animals can provide important
124 information on toxicity to reproductive organs. The existing toxicology data for the
125 compound should always be considered, taking into account the dose levels, toxicokinetic
126 profile, and dosing duration. For example, the evaluation of fertility effects for a
127 pharmaceutical that damages testicular tissue might warrant modifications to the standard
128 fertility study, if such a study would be appropriate.

129 Sometimes, toxicity in animals precludes attaining a systemic exposure relevant to the
130 human exposure under conditions of use and this should be addressed.

131 **3.1.4 Timing Considerations**

132 General guidance on the timing for conduct of reproductive toxicity studies covering Stages
133 A-F relative to clinical studies is described in the ICH M3(R2) and ICH S9 guidelines (1,3).
134 The timing for when to conduct specific reproductive toxicity assessments should take into
135 consideration the points discussed above. Based on these factors, it can sometimes be
136 appropriate to consider altering timing of the assessment of specific reproductive stages. For
137 example, if there is an equivocal observation from a preliminary study and other compounds
138 in the class are without risk, then consideration should be given to accelerating the definitive
139 studies. In contrast, there can be circumstances for deferring studies. For example, when
140 other studies have revealed a risk and appropriate precautions in clinical trials have been
141 taken, the conduct of definitive studies evaluating the relevant reproductive stages can be
142 deferred to later in development than is recommended in ICH M3(R2) (1). When conducting
143 enhanced Pre- and PostNatal Development (ePPND) studies in NonHuman Primates (NHP)
144 see ICH S6(R1) (2) for timing.

145 Additional options that include study deferral are discussed in Section 3.3.3.

146 **3.1.5 Other Considerations for Reproductive Toxicity Studies**

147 For some species and compounds, it can be more appropriate to test multiple reproductive
148 stages in a single study (e.g., monoclonal antibodies in NHPs; see ICH S6(R1) (2)).
149 Consideration can also be given to evaluation of reproductive toxicity endpoints as a
150 component of another study type (e.g., male fertility as part of a repeat-dose toxicity study,
151 see Section 3.2).

152 When designing a pre- and post-natal development (PPND) or ePPND study, thought should
153 be given to the value for juvenile animal endpoints for supporting the safety of pediatric use
154 (see Section 9.4.2.1).

155 Alternative assays are described as part of an integrated testing strategy for assessing
156 embryo-fetal developmental endpoints as described in the examples below (see Section
157 3.3.2.1).

158

159 **3.2 Strategy to Address Fertility and Early Embryonic Development**

160 The aim of the fertility study is to test for disturbances resulting from treatment from before
161 mating of males and/or females through mating and implantation. This comprises evaluation
162 of Stages A and B of the reproductive process (see Sections 6 and 9.4).

163 Fertility studies are generally only performed in rodents or rabbits. Mating evaluations are
164 not generally feasible in non-rodents such as dogs and NHPs. For example if NHPs are the
165 only pharmacologically relevant species (as for many monoclonal antibodies, see ICH
166 S6(R1) (2)), fertility evaluations can be based on the results of the repeat-dose toxicity
167 studies (e.g., histopathological examinations).

168 Histopathology of the reproductive organs from the repeat-dose toxicity studies is a sensitive
169 method of detecting the majority of effects on male and female fertility, provided animals are
170 sexually mature.

171 Dogs and minipigs used in long-term repeat-dose studies should have, in general, sexually
172 matured by the end of the study. If NHPs are to be used to assess effects on fertility, there
173 should be a sufficient number of sexually mature animals at study termination.

174 If repeat-dose toxicity studies are used to assess effects on fertility, a comprehensive
175 histopathological examination of the reproductive organs from both male and female animals
176 should be performed (Note 1).

177 When there is cause for concern based on mode of action or data from previous studies,
178 additional examinations can be included in repeat-dose toxicity studies, e.g., sperm
179 collection, or monitoring of the estrous or menstrual cycle. Studies of two to four weeks
180 treatment duration can be expected to provide an initial evaluation of effects on the
181 reproductive organs. This information will later be supplemented with similar evaluations in
182 the subchronic and chronic toxicity studies.

183 A dedicated fertility study includes a mating phase and serves to detect effects that cannot be
184 assessed by histopathology of the reproductive organs. However, if the drug has clinically
185 relevant adverse effects on male or female reproductive organs in the repeat-dose toxicity
186 studies, a routine fertility study in the affected sex will be of limited value and not warranted.
187 Likewise, a fertility study is not warranted for pharmaceuticals that will not be used in
188 subjects of reproductive age. Generally, the repeated-dose toxicity study results can be used
189 to design the fertility study without the need for further dose ranging studies.

190 If no adverse effects on fertility are anticipated, male and female rodents can be evaluated in
191 the same fertility study. However, if effects on fertility are identified, the affected sex should
192 then be determined. In addition, if it cannot be determined whether effects are reversible
193 based on the pathophysiological evaluation, then reversibility of induced effects should be
194 evaluated. These determinations can have an important impact on risk assessment.

195

196 **3.3 Strategies to Address Embryo Fetal Development (EFD)**

197 The aim of the EFD studies is to detect adverse effects on the pregnant female and
198 development of the embryo and fetus consequent to exposure of the female during the period
199 of major organogenesis (Stage C). EFD studies include full evaluation of fetal development
200 and survival. For most non-highly targeted pharmaceuticals (e.g., small molecules), effects
201 on EFD are typically evaluated in two species (i.e., rodent and non-rodent). There are cases
202 where testing for effects on EFD in a single species can suffice. General strategies to address
203 EFD studies are shown in Figure 3-1.

204 **3.3.1 Routine Approach for Addressing EFD Risk**

205 In situations where the use of rodent or rabbit species is appropriate, at least one of the test
206 species should exhibit the desired pharmacodynamic (PD) response (Section 4). If the
207 pharmaceutical is not pharmacodynamically active in any routinely used species (Section
208 9.3), genetically modified (GM) animals or use of a surrogate molecule can be considered.
209 If it is a highly-targeted pharmaceutical these data can be sufficient. If the pharmaceutical is
210 non-highly targeted, it can be appropriate to also administer it to a rodent or a rabbit to test
211 for off-target effects.

212 However, under some circumstances other approaches can be used to defer (Table 3-1) or
213 replace (Section 9.5.5) definitive studies. Alternatively, there can be adequate information
214 to communicate risk without conducting additional studies. Evidence suggesting an adverse
215 effect of the intended pharmacological mechanism on EFD (e.g., mechanism of action,
216 phenotypic data from genetically modified animals, class effects) can be sufficient to
217 communicate risk.

218 Non-routine animal models or a surrogate molecule can be considered in place of NHPs for
219 either small molecules or biotechnology-derived products, if appropriate scientific
220 justification indicates that results will inform the assessment of reproductive risk (Section
221 4.3).

222 In certain justified cases, testing for effects on embryo-fetal development in a single species
223 can suffice. One example is for highly targeted pharmaceuticals (e.g., for biotechnology-
224 derived products, see ICH S6(R1)) when there is only one relevant species that can be used
225 in reproductive testing (2). Another circumstance is for non-highly targeted pharmaceuticals
226 when it can be shown that a single species is a relevant model for the human, based on
227 pharmacodynamics, pharmacokinetics and metabolite profiles, as well as toxicology data. If
228 the result is clearly positive (teratogenic and/or embryofetal lethal; TEFL; see Glossary)
229 under relevant exposure, testing in a second species is not warranted.

230 When there are no pharmacologically relevant species (e.g., the pharmacological target only
231 exists in humans), EFD studies in two species can still be warranted to detect off-target
232 effects or secondary pharmacology as appropriate based on the therapeutic modality and the
233 indication.

234 For biotechnology-derived products, when no relevant species can be identified because the
235 biopharmaceutical agent does not interact with the orthologous target in any species relevant
236 to reproductive toxicity testing, use of surrogate molecules or transgenic models can be
237 considered, as described in detail in ICH S6(R1) (2). If there are no relevant species,
238 genetically modified animals, or surrogate, *in vivo* reproductive toxicity testing is not
239 meaningful; however, the approach used should be justified.

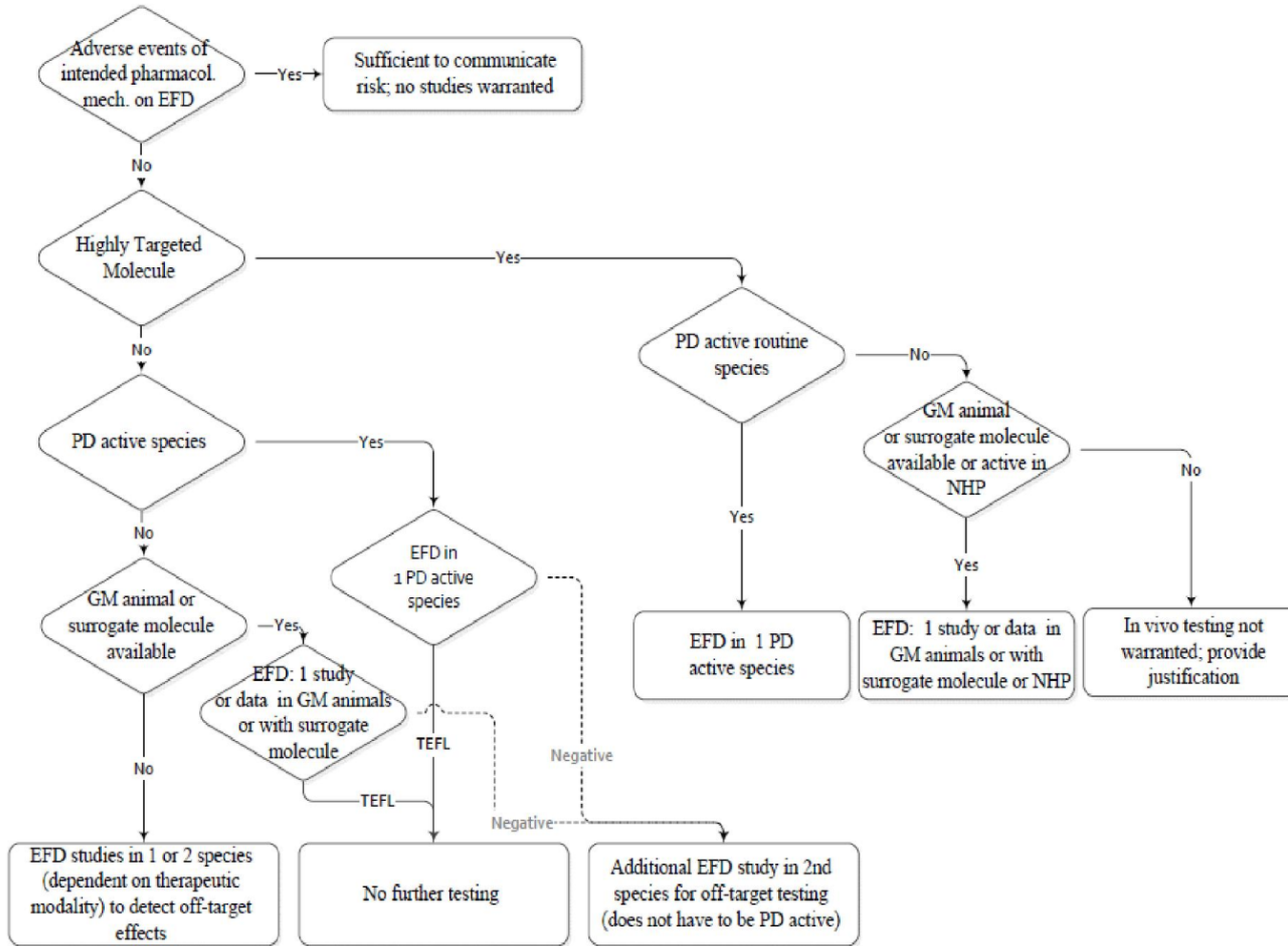
240 For other therapeutic modalities that lack orthologous target engagement in useful
241 reproductive toxicology species and also have anticipated off-target effects, use of surrogate
242 molecules or transgenic models can be considered.

243 Several scenarios of use for integrated testing strategies are described in Annex 9.5.5.

244

245

Figure 3-1: General Strategy to Address EFD



246

247 3.3.2 Optional Approaches for Addressing EFD Risk

248 3.3.2.1 Use of Alternative Assays

249 Use of alternative *in vitro*, ex vivo, and non-mammalian *in vivo* assays (alternative assays)
 250 can reduce animal use while preserving the ability to detect relevant reproductive risks. The
 251 use of qualified (Note 2) alternative assays can be an appropriate approach in lieu of the
 252 routine approach discussed above. Use of qualified alternative assays is appropriate for risk
 253 assessment under certain circumstances where they are interpreted in conjunction with *in*
 254 *vivo* reproductive testing. Although they are not a replacement for all *in vivo* reproductive
 255 testing, they can reduce *in vivo* mammalian animal studies and/or animal usage (Section
 256 3.3.2.1). Several scenarios of use for integrated testing strategies are described in Annex
 257 9.5.5. Furthermore, while a study in a second species could be conducted under the routine
 258 approach, the use of an alternative assay could be more informative in some circumstances,
 259 taking into consideration route of administration, exposure, and mechanism of action.

260 The circumstances justifying the incorporation of alternative assays in an integrated testing
261 strategy for assessing EFD risk will be dependent upon a number of factors. These could
262 include the severity of the disease, the characteristics of the patient population, or the
263 limitations of some traditional test systems for specific therapeutic targets. The
264 pharmacological or biological plausibility for developmental toxicity is a key consideration.

265
266 This guideline does not recommend specific assays, but basic principles are included to assist
267 in assay qualification for potential regulatory use (Section 9.5.2).

268 For appropriate use of alternative assays it is important to know the reliability and
269 predictivity for *in vivo* reproductive outcomes. The Annex provides information on various
270 reference compounds that can be used to assess alternative methods for embryo-fetal
271 development/deaths (Note 3). It is possible that a suite of assays/assessments will show
272 improved predictivity.

273
274 Where applicable, testing strategies can take into consideration data from qualified
275 alternative assays in combination with one or more *in vivo* mammalian EFD studies. Any
276 alternative assay integrated into a testing strategy should be qualified for its intended context
277 of use (Section 9.5). When alternative assays are used to contribute to the risk assessment
278 they should generally be conducted according to GLP, particularly when the assay results do
279 not identify a hazard. Contexts of use (see Glossary) could include, but are not limited to:

- 280 a. Being part of an integrated testing strategy for assessing embryo-fetal developmental
281 endpoints as described in the scenarios in Section 9.5.5;
- 282 b. Deferral of definitive studies as discussed in Section 3.3.3;
- 283 c. Complete replacement of one species when used in conjunction with an enhanced
284 pEFD study in one species (see Scenarios in Section 9.5.5);
- 285 d. There is evidence (e.g., a mechanism of action affecting fundamental pathways in
286 developmental biology, phenotypic data from genetically modified animals, class
287 effects) suggesting an adverse effect on EFD, or contributing to the weight of
288 evidence when animal data are equivocal;
- 289 e. Toxicity (on-target related and/or off-target) in a routine animal species precludes
290 attaining a systemic exposure relevant to the human exposure under conditions of use,
291 but higher exposures can be attained in an alternative assay;
- 292 f. Low systemic exposure (e.g., no embryo-fetal exposure) in humans such as following
293 ophthalmic administration.

294 The information from the alternative qualified test systems should be used with all available
295 *in vivo* nonclinical and human data as part of an integrated risk assessment approach (see
296 Principles of Risk assessment; Section 7).

297 **3.3.2.2 *In vitro* and Non-mammalian Exposure Information**

298 As stated in section 7 of this guideline, for the purposes of risk assessment, it is important to
 299 consider exposure in the interpretation of non-clinical studies assessing reproductive toxicity.
 300 This also applies to assays conducted using *in vitro* or non-mammalian systems. The
 301 pharmacokinetic parameter used is dependent upon how the assay was qualified in relation to
 302 the *in vivo* concentrations at which the EFD observations were made, considering any
 303 normalization factors used in the assay qualification. For example, the maximum
 304 concentration tested without an adverse effect in the *in vitro* system can be compared to the
 305 C_{max} in humans for the determination of potential human risk, applying the normalization
 306 factor used in the assay qualification.
 307

308 **3.3.3 Potential Approaches to Defer *in vivo* Testing as Part of an Integrated Testing**
 309 **Strategy**

310 Table 3-1 illustrates approaches to support inclusion of Women Of Child-Bearing Potential
 311 (WOCBP) in clinical studies while deferring conduct of definitive assays. This applies to
 312 circumstances where 2 definitive EFD studies are warranted for the pharmaceutical.
 313

314 One such approach is the use of an enhanced pEFD study for one of the species. In this case,
 315 the pEFD study (see ICH M3(R2)) should be conducted in accordance with GLP regulations,
 316 the number of pregnant animals should be increased from 6 to ≥ 8 per group, and include
 317 fetal skeletal examinations.
 318

319 **Table 3-1. Approaches for Deferral of Definitive EFD Studies in 2 Species**

Approach	Stage of Development			
	Limited inclusion of WOCBP ^a	Unlimited inclusion of WOCBP up to start of Phase 3 (supports Phase 2a/b) ^b	Unlimited inclusion of WOCBP up to marketing (supports Phase 3)	To support marketing ^c
A	1 st species EFD (enhanced pEFD or definitive) + Qualified alternative assay		2 nd species definitive EFD	1 st species definitive EFD if not conducted earlier
B	1 st species pEFD + 2 nd species EFD (enhanced pEFD or definitive)		1 st species definitive EFD	2 nd species definitive EFD if not conducted earlier
C ^d	2 species pEFD	2 species definitive EFD		

^a Up to 150 WOCBP receiving investigational treatment for a relatively short duration (up to 3 months).
^b All approaches include “where precautions to prevent pregnancy in clinical trials (see above) are used.”
^c For monoclonal antibodies, the ePPND is generally conducted before marketing approval (see ICH S6(R1)).
^d See ICH M3(R2) for regional differences.

320 **3.4 Strategy to Address Effects on PPND**

321 The aim of the PPND study is to detect adverse effects following exposure of the mother
322 from implantation through weaning on the pregnant or lactating female and development of
323 the offspring. Since manifestations of effects induced during this period can be delayed,
324 development of the offspring is monitored through sexual maturity (i.e., Stages C to F). The
325 usual species used for PPND is the rat; however, other species can be used as appropriate
326 with modifications of the endpoints assessed.

327 In most cases, a preliminary PPND study is optional because the appropriate information is
328 generally available from prior studies to design the definitive study. However, a preliminary
329 PPND study with termination of the pups before or at weaning can be used to select dose
330 levels or inform study design and to provide pup exposure data.

331 For pharmaceuticals that can only be tested in the NHP, the ePPND study can provide a
332 limited assessment of post-natal effects, but it is not feasible to follow the offspring through
333 maturity. For the timing of the ePPND study see ICH S6(R1) (2).

334 **3.5 Toxicokinetics (TK)**

335 TK investigations are generally expected and the use of the data is discussed throughout this
336 document. General concepts regarding TK data collection are discussed in ICH S3A.

337 Determination of the pharmaceutical's concentration in the fetus can be of interest to
338 facilitate interpretation of discordant or equivocal evidence of developmental hazard.
339 However, determination of placental transfer is generally not warranted because of limited
340 ability to translate data to human fetal exposures.

341

342 Many pharmaceuticals are excreted in milk, although lactational excretion data in animals are
343 of uncertain value for human risk assessment. Therefore, measurement of drug
344 concentrations in the milk of animals is generally not warranted. However, determination of
345 a pharmaceutical's concentrations in the offspring can support interpretation of findings
346 observed during the pre-weaning period.

347 **4 TEST SYSTEM SELECTION**

348 **4.1 Routine Test Species**

349 When a study is warranted, a mammalian species should be used. For the primary species, it
350 is generally desirable to use the same species and strain as in other toxicity studies to avoid
351 additional studies to characterize pharmacokinetics and metabolism, and/or for dose-range
352 finding. The species used should be well-characterized with respect to health, fertility,
353 fecundity, and background rates of malformation and embryo-fetal death. Generally, within
354 and between reproductive studies animals should be of comparable age, weight and parity at
355 the start. The easiest way to fulfil these factors is to use animals that are young, sexually
356 mature adults at the time of the start of dosing with the females being virgin, with the

357 exception of NHP where proven mothers can be an advantage for ePPND studies.

358 The species chosen for testing should be relevant and justified based on their advantages and
359 disadvantages (see Table 9-1 in Section 9.3). If the species selected differs considerably from
360 the human in regard to the considerations below, the impact should be considered when
361 interpreting the reproductive toxicity data (see Principles of Risk Assessment, Section 7).
362 Assessing all of the reproductive endpoints or parameters of interest in a single test species,
363 however, is not always possible.

364 Additional points to consider in selection of a species relate to the interaction of the
365 pharmaceutical with the species including:

- 366 a. The pharmacokinetic and metabolite profile (including adequate exposure to major
367 human metabolites, as discussed in ICH M3(R2) (1));
- 368 b. Whether the species expresses the pharmacologic target (e.g., is an endogenous or
369 exogenous target) and whether the pharmaceutical has adequate affinity for the target
370 in the species selected;
- 371 c. Whether the functional pharmacological activity of the pharmaceutical is exhibited in
372 the test species.

373 For highly targeted molecules, selection of a pharmacologically relevant species is
374 particularly important as described in more detail in ICH S6(R1) (2).

375 **4.1.1 Rat as the Primary Species for Reproductive Toxicity Testing**

376 The rat is the most often used rodent species for reasons of practicality, general knowledge
377 of pharmacology in this species, the extensive toxicology data usually available for
378 interpretation of nonclinical observations from development of the pharmaceutical, and the
379 large amount of historical background data. Thus, in many cases based on how species are
380 selected for general toxicity studies, the rat is generally appropriate for reproductive toxicity
381 testing.

382 **4.1.2 Rabbit as the Secondary Species for EFD studies**

383 For assessment of EFD only, a second mammalian non-rodent species is often warranted,
384 although there are exceptions (e.g., vaccines, therapeutic antibodies, etc., see Sections 4.1.3
385 and 4.2, respectively). The rabbit has proven to be useful in identifying human teratogens
386 that have not been detected in rodents; and the rabbit is routinely used as the non-rodent
387 species based on the extensive historical background data, availability of animals, and
388 practicality.

389 **4.1.3 Species Selection for Preventative and Therapeutic Vaccines**

390 The animal species selected for testing of vaccines (with or without adjuvants) should
391 demonstrate an immune response to the vaccine. Typically, rabbits, rats, and mice are used.
392 Nonhuman primates should be used only if no other relevant animal species is available,
393 even though quantitative and qualitative differences can exist in the responses (e.g., in

394 humoral and cellular endpoints). It is usually sufficient to conduct developmental toxicity
395 studies using only one animal model.

396 Rabbits are the most common species used for vaccine developmental toxicity studies, but
397 other species are also appropriate. In primates (as in humans), the transfer of maternal
398 antibodies across the placenta is limited, but generally increases over the course of gestation.
399 In other species routinely used in reproductive testing the time course of transfer differs.
400 The type of developmental toxicity study conducted and the choice of the animal model
401 should be justified based on the immune response observed and the ability to administer an
402 appropriate dose.

403 When there is a lack of an appropriate animal model (including NHP), a developmental
404 toxicity study in rabbits, rats, or mice can still provide important information regarding
405 potential embryo/fetal toxic effects of the vaccine components/formulation and safety of the
406 product during pregnancy.

407 **4.2 Non-routine Test Species**

408 There are cases where it can be appropriate to use strategies other than those involved using
409 the routine species discussed above. A commonly encountered example is where the rabbit is
410 unsuitable for EFD testing. In situations like this, one can consider alternative species or
411 approaches that can inform the risk assessment.

412 Many other species have been used to evaluate the effects of pharmaceuticals on the various
413 reproductive stages. The suitability of alternative species will depend on the reproductive
414 endpoints to be assessed (see Table 9-1 in Section 9.3).

415 NHPs can also be used for evaluating reproductive toxicity, especially for biotechnology-
416 derived products, as described in ICH S6(R1) (2). NHPs should be considered if they are the
417 only pharmacologically relevant species, provided that it is not already clear that the
418 pharmacology of the pharmaceutical is incompatible with normal development or
419 maintenance of pregnancy. There are additional factors that further limit the utility of
420 studies in NHPs for reproductive risk assessment (see Annex 9.3 and ICH S6(R1)). An
421 alternative animal model can be considered in place of NHPs for either small molecules or
422 biotechnology-derived products by using a surrogate molecule that elicits the appropriate
423 pharmacologic activity in the animal model, or data from genetically modified animals. The
424 results of the studies can inform the assessment of reproductive risk (see Sections 4.3 and 7).

425 For biotechnology-derived products, when no relevant species can be identified because the
426 biopharmaceutical agent does not interact with the orthologous target in any species relevant
427 to reproductive toxicity testing, use of surrogate molecules or genetically modified models
428 can be considered, as described in ICH S6(R1) (2) and Section 4.3.2. For some therapeutic
429 modalities that lack orthologous target engagement in useful reproductive toxicology species
430 and also have anticipated off-target effects, the testing strategy should address both of these
431 situations.

432 In lieu of, or in addition to, the use of an *in vivo* mammalian study for assessment of
433 reproductive toxicity, alternative approaches that can be considered include assessment of
434 pharmacologic or mechanistic information, non-mammalian *in vivo* studies, or *in vitro*
435 assays that predict reproductive toxicity (see Principles of Risk assessment Section 7).

436 4.3 Other Test Systems

437 4.3.1 Use of Disease Models

438 Disease animal models are not routinely used in reproductive toxicity testing; however, there
439 are some cases where they can be informative. Studies in disease models can be of value in
440 cases where the data obtained from healthy animals could be misleading or otherwise not
441 apply to the disease conditions in the clinical setting. Examples of situations where a
442 reproductive toxicity study in a disease model could contribute information to the risk
443 assessment include studies with pharmaceuticals that are replacement therapies, when the
444 target is only present in disease state, or when the pharmacologic activity of the test article
445 could yield confounding results in healthy animals (e.g., causes hypoglycemia or
446 hypotension).

447 Recognizing that no animal model perfectly replicates human disease, there are several
448 factors to be considered in choosing to study toxicity to reproduction in a disease animal
449 model. The model should be pharmacologically relevant and appropriate for the reproductive
450 endpoints being assessed. The pathophysiology of the disease course in the model should be
451 characterized. Some differences from the human pathophysiology would not preclude its use
452 provided that these are unlikely to confound data interpretation. Animal to animal variability
453 should be characterized and appropriate within the context of the study. Reference data for
454 the study endpoints should be available or should be generated during the study to aid data
455 interpretation.

456 Although disease animal models can be used in definitive reproductive toxicity studies, they
457 are more likely to be used as supplementary approaches to understand the relevance of
458 adverse reproductive effects of the pharmaceutical in normal animals. The use of disease
459 animal models and the design of the study for reproductive toxicity testing should be
460 justified.

461

462 4.3.2 Use of Genetically Modified Models and Use of Surrogate Molecules

463 For both genetically modified models and for surrogate molecules the effect of the intended
464 pharmacology on reproduction is being investigated and thus informs the assessment of risk.
465 For example, if the pharmacology is linked to adverse effects on reproduction, it can
466 reasonably be concluded that the adverse effects would be experienced in some proportion of
467 pregnant women receiving the pharmaceutical. However, the actual proportion of individuals
468 affected (incidence) cannot be determined from animal studies, even if the actual
469 pharmaceutical and a pharmacologically relevant species are used.

470 Genetically modified models can be used to create disease models or to characterize the
471 on-target and off-target effects of a pharmaceutical on reproductive toxicity parameters.
472 Such models can inform on whether the pharmacology of the target is closely linked to
473 adverse effects on reproduction and development. When these models are used and
474 off-target effects are anticipated based on therapeutic modality, the clinical candidate should
475 be evaluated with this model to assess both on- and off-target effects.

476 When the clinical candidate does not have adequate activity against the target receptor in the
477 routine test species, surrogate molecules can be used for any modality to assess potential
478 adverse effects on reproductive toxicity. Using surrogate molecules is analogous to
479 identifying class-effects from structurally diverse molecules with similar pharmacology.
480 The overall approach is comparable to using a surrogate antibody that is pharmacologically
481 active in the species being tested rather than using the humanized antibody that is
482 pharmacologically active only in the NHP.

483 If there are no adverse effects on reproduction associated with the target pharmacology,
484 evaluation of off-target reproductive toxicity using the clinical candidate is warranted.

485

486 **5 DOSE LEVEL SELECTION, ROUTE OF ADMINISTRATION AND SCHEDULE**

487 As part of the dose selection process, route of administration and schedule are important
488 components in the design of reproductive toxicity studies. The dose selection should
489 optimize exposure relative to humans considering route, schedule, and pharmacokinetics
490 profile, to the extent that is practical.

491 The choice of dose levels, schedule and route of administration should be based on all
492 available information (e.g., pharmacology, repeated-dose toxicity, pharmaco-/toxicokinetics,
493 and Dose Range Finding studies) and a rationale should be provided. Guidance on the
494 principles of dose selection is given in ICH M3(R2) Q&A (1) and ICH S6(R1) (2), and all
495 available data should be used. Dose levels should be selected to investigate dose-response
496 relationships for the primary endpoints of the study. Using doses similar to those used in the
497 repeat dose toxicity studies of comparable duration permits interpretation of potential effects
498 on reproductive and/or developmental endpoints within the context of general systemic
499 toxicity and enables integration of data. When sufficient information on tolerability and
500 pharmaco-/toxicokinetics in the test system is not available, appropriately designed
501 exploratory studies are advisable.

502 Dosing schedules used in the toxicity studies influence the exposure profile which can be
503 important in the risk assessment. Usually mimicking the clinical schedule is sufficient, but is
504 not always warranted. A more frequent (e.g., twice a day) or a less frequent schedule can be
505 appropriate to provide an exposure profile more relevant to the clinical exposure. When a
506 more frequent schedule is contemplated, pragmatic factors (e.g., study logistics, stress on
507 animals) should be considered.

508 In general the route of administration should be similar to the clinical route, provided the
509 relevant human reproductive risk can be assessed. In circumstances where systemic exposure
510 cannot be achieved or only small multiples of the clinical systemic exposure are achieved in
511 the absence of maternal toxicity, a different route of administration should be considered.
512 Use of a route of administration other than the clinical route should be justified in the context
513 of the general toxicology program. When multiple routes of administration are being
514 evaluated in humans, a single route in the test species can be adequate provided sufficient
515 systemic exposure is achieved compared to that of the clinical routes.

516 It is not always warranted to use pregnant animals for dose selection, even if the reproductive
517 study assesses pregnant animals. However, when exposure-based endpoints are used as the
518 basis for selection of the dose levels (Section 5.1.3), it can be important to have TK from
519 pregnant animals. If the TK is derived from non-pregnant animals for dose selection, then the
520 achievement of the TK endpoint should be confirmed in pregnant animals.

521 **5.1 Dose Selection Common to all Pharmaceuticals, Including Biotechnology-** 522 **derived Pharmaceuticals**

523 There are a number of dose selection endpoints that can be used for reproductive toxicity
524 studies. All the endpoints discussed in this section are considered equally appropriate in
525 terms of study design. The high dose in the definitive study should be one that is predicted to
526 produce the anticipated change in the endpoint as described below in Sections 5.1.1 to 5.1.6.
527 The selected high dose should be based on the observations made in appropriately designed
528 studies, including the effects observed at higher dose levels in other studies (e.g., repeat-dose,
529 TK, pEFD).

530 Justification for high dose selection using other endpoints than specified below, can be made
531 on a case-by-case basis.

532 **5.1.1 Toxicity-based Endpoints**

533 This endpoint is based on the prediction of minimal toxicity in the parental animals at the
534 high dose. Minimal toxicity is defined as having an adverse effect on the parental animals
535 without having an anticipated direct effect on the reproductive outcome. Factors limiting the
536 high dose determined from previously conducted studies could include:

- 537 • Alterations in body weight (gain or absolute; either reductions or increases). Minor,
538 transient changes in body weight gain or in body weight are not considered dose
539 limiting. When assessing weight change effects, the entire dosing duration of the
540 study should be considered and the absolute change that is appropriate is dependent
541 on the parameter being measured, the species, strain, and the window of development
542 being evaluated.
- 543 • Specific target organ toxicity (e.g., ovarian, uterine) or clinical pathology
544 perturbations (e.g., changes in glucose) that would interfere with the study endpoints
545 within the duration of the planned reproductive or developmental toxicity study.
- 546 • Exaggerated pharmacological responses (e.g., excessive sedation or hypoglycemia)

547 • Toxicological responses (e.g., convulsions, increased TEFL).

548 **5.1.2 Absorption, Distribution, Metabolism and Excretion (ADME)-based Saturation**
549 **of Systemic Exposure Endpoint**

550 High dose selection based on saturation of systemic exposure measured by systemic
551 availability of pharmaceutical-related substances can be appropriate (see ICH M3(R2) (1)).
552 There is, however, little value in increasing the administered dose if it does not result in
553 increased plasma concentration. For the purposes of this guideline, saturation of exposure is
554 defined as substantial increases in dose that result in minimal increases in total exposure
555 (e.g., a doubling of the dose resulting in only an approximate 20% increase in exposure).

556 **5.1.3 Exposure-based Endpoint**

557 It can be appropriate to select doses based on exposure margins above the exposure at the
558 maximum recommended human dose (MRHD). For pharmaceuticals having primary and
559 secondary pharmacology (or off-target effects) in the test species (e.g., small molecules), a
560 systemic exposure representing a large multiple of the human AUC (area under the exposure
561 curve) or C_{max} can be an appropriate endpoint for high-dose selection. This dose selection
562 approach can be applied when there are qualitatively similar metabolite profiles between
563 humans and the test species. The rationale for the metric used should be provided. Doses
564 anticipated to provide an exposure > 25-fold of the clinical systemic exposure at the MRHD
565 are generally considered appropriate as the maximum dose for reproductive toxicity studies
566 (Note 4). Usually this is based on the parent moiety if it is the pharmacologically active
567 agent. There are other cases (e.g., prodrugs, pharmacologically active metabolites) for which
568 the Sponsor should provide a justification for the moieties included in the exposure multiple
569 calculations.

570 When evaluating a pharmaceutical against a human endogenous target using an exposure-
571 based endpoint, it is recommended to choose at least one species with pharmacodynamic
572 activity. For studies using a surrogate molecule a dose should be used that has adequate
573 pharmacodynamic activity in the test species. In addition to testing the surrogate, if the
574 clinical candidate is anticipated to have secondary pharmacology or off-target effects, the
575 clinical candidate should also be tested at doses anticipated to provide an exposure > 25-fold
576 at the MRHD in the routine species.

577 Alternatively, instead of using a surrogate, for clinical candidates that have some
578 demonstrated pharmacodynamic activity in the test species only at exposures > 25-fold,
579 doses that achieve pharmacodynamic activity in the routine test species can be used.
580 However, it should be noted that irrelevant off-target effects are likely to be observed.

581 If none of the routine test species are pharmacodynamically relevant, but the target is
582 endogenous and the clinical candidate is anticipated to have off-target effects, an alternative
583 endpoint rather than the exposure-based endpoints should be considered (e.g., limit dose,
584 maximum feasible dose, toxicity-based endpoints).

585 When there is no human endogenous target (e.g., viral target), a > 25-fold exposure multiple
586 of the MRHD is sufficient for high dose selection.

587 **5.1.3.1 Considerations for Total vs. Fraction Unbound Pharmaceutical Exposure**

588 The choice for the use of total vs. fraction unbound pharmaceutical exposures should be
589 justified. The total exposure can be used as the default, unless the fraction unbound results in
590 a lower exposure margin than that of the total; in this case the lower exposure multiple
591 should be used for the comparison of animal vs. human exposures. Alternatively, the fraction
592 unbound pharmaceutical exposure can be used regardless of whether it generates a lower or
593 greater exposure multiple than that of the total exposure provided the following applies:

- 594 • The fractions unbound can be calculated accurately from the total pharmaceutical
595 exposure, is reproducible at the effective concentrations in humans and at the
596 toxicological concentrations in animals, and the fractions unbound are statistically
597 significantly different.

598

599 Two examples of how this calculation might impact the exposure multiples are provided
600 below.

- 601 • 25 fold exposure multiple not met: If the total exposure is 25 $\mu\text{M}\cdot\text{hr}$ in animals and 1
602 $\mu\text{M}\cdot\text{hr}$ in humans and unbound protein fraction is 5% and the unbound fraction in
603 animals is 1%, then the margin would be 5.

- 604 • 25 fold exposure multiple exceeded: If the exposure is 10 $\mu\text{M}\cdot\text{hr}$ in animals and 5
605 $\mu\text{M}\cdot\text{hr}$ in humans and unbound protein fraction is 1% in human and 20% in animals,
606 then the unbound ratio would be 40 rather than the apparent ratio of 2 based on total.

607 **5.1.3.2 Exposure-based Approach for Highly Targeted Therapeutics**

608 Highly targeted therapies (e.g., monoclonal antibodies, therapeutic proteins) are those that
609 exhibit no or minimal off-target effect. For these therapeutics that exhibit pharmacodynamic
610 effects in the test species, high dose selection can be accomplished by either identifying a
611 dose which provides the maximum intended pharmacological effect in the preclinical species
612 or a dose which provides an approximately 10-fold exposure multiple over the maximum
613 exposure to be achieved in the clinic, whichever one is higher (ICH S6(R1)) (2). Corrections
614 for large differences in target binding affinity and *in vitro* pharmacological activity between
615 the nonclinical species and humans should be considered in dose selection such that a higher
616 dose can be appropriate to elicit pharmacodynamic effects, if not limited by toxicity or
617 feasibility. If the routine species do not exhibit pharmacological activity and a surrogate
618 molecule is used, a dose of the surrogate that is 10-fold that which elicits the intended
619 pharmacological activity in the test species can be appropriate.

620 **5.1.4** *Maximum Feasible Dose (MFD) Endpoint*

621 Use of the MFD should maximize exposure in the test species, rather than maximize the
622 administered dose (see also ICH M3(R2) (1)).

623 The MFD can be used for high dose selection when the physico-chemical properties of the
624 test substance (or formulation) associated with the route/frequency of administration and the
625 anatomical/physiological attributes of the test species limit the amount of test substance that
626 can be administered.

627 **5.1.5** *Limit Dose Endpoint*

628 A limit dose of 1 g/kg/day can be applied when other dose selection factors have not been
629 achieved with lower dose levels (see also ICH M3(R2) (1) for other considerations).

630 **5.1.6** *Selection of Lower Dose Levels*

631 It is generally desirable to establish a “no observed adverse effect level” for developmental
632 and reproductive toxicity. Having selected the high dose, lower doses should be selected
633 taking into account exposure, pharmacology, and toxicity, such that there is separation in
634 anticipated outcomes between groups. Any dose level that yields a sub-therapeutic exposure
635 is not generally informative to risk assessment, unless it is the highest dose that can be
636 achieved without toxicity in the parental animals. For some of the variables in reproductive
637 toxicity studies the ability to discriminate between background and treatment effects can be
638 difficult and the presence or absence of a dose-related trend can be informative. The low dose
639 should generally provide a low multiple (e.g., 1 to 5-fold) of the human exposure MRHD.
640 The exposure at the mid dose should be intermediate between the exposures at the low and
641 the high doses; however, dose spacing that results in less than 3-fold increase in exposure is
642 not generally recommended.

643 **5.2** **Dose Selection and Study Designs for Vaccines**

644 This guideline covers vaccines (adjuvanted or not) used in both preventative and therapeutic
645 indications against infectious diseases. The principles outlined can be applicable to the
646 nonclinical testing of vaccines for other indications as well (e.g., cancer). The types of
647 studies depend on the target population for the vaccine and the relevant reproductive risk.
648 Generally, reproductive studies are not warranted for vaccines being developed for neonates,
649 pre-pubertal children, or geriatric populations.

650 For reproductive toxicity studies of vaccines it is typically sufficient to assess a single dose
651 level capable of inducing an immune response in the animal model (Section 4.1.3). This
652 single dose level should be the maximum human dose without correcting for bodyweight
653 (i.e., 1 human dose = 1 animal dose). If it is not feasible to administer the maximum human
654 dose to the animal because of a limitation in total volume that can be administered or because
655 of dose-limiting toxicity (e.g., local, systemic), a dose that exceeds the human dose on a
656 mg/kg basis can be used. To use a reduced dose, justification as to why a full human dose
657 cannot be used in an animal model should be provided.

658 The vaccination regimen should maximize maternal antibody titers and /or immune response
659 throughout the embryonic, fetal, and early postnatal periods. Timing and number of doses
660 will depend on the onset and duration of the immune response of the particular vaccine.
661 When developing vaccines to be given during pregnancy, the sponsor should justify the
662 specific study design based upon its intended use (e.g., protecting the mother during
663 pregnancy or protecting the child early postnatally).

664 Daily dosing regimens can lead to overexposure to the vaccine constituents. Episodic dosing
665 of pregnant animals rather than daily dosing is recommended. Also, episodic dosing better
666 approximates the proposed clinical immunization schedule for most preventive and
667 therapeutic vaccines for infectious disease indications. Considering the short gestational
668 period of routine animal species, it is generally recommended to administer a priming dose(s)
669 to the animals several days or weeks prior to mating in order to elicit peak immune response
670 during the critical phases of pregnancy (i.e., the period of organogenesis). The dosing
671 regimen can be modified according to the intended vaccination schedule in humans.

672 At least one dose should be administered during early organogenesis to evaluate potential
673 direct embryotoxic effects of the components of the vaccine formulation and to maintain a
674 high antibody response throughout the remainder of gestation. If EFD toxicity is observed,
675 this can be further assessed using subgroups of animals that are dosed at certain time points.

676 In cases where a vaccine includes a novel, active constitutive ingredient (including novel
677 adjuvants) consideration of additional testing strategies similar to those for non-vaccine
678 products can be appropriate.

679 It is recommended that the route of administration be similar to the clinical route of
680 administration.

681 **6 DESIGN AND EVALUATION OF IN VIVO MAMMALIAN STUDIES**

682 The testing strategy to evaluate the potential reproductive risk of a pharmaceutical can
683 include one or more *in vivo* studies. Although three separate study designs have been
684 employed for the development of the majority of pharmaceuticals, various combinations of
685 these study designs can be conducted to reduce animal use. All available pharmacological,
686 kinetic, and toxicological data for the pharmaceutical should be considered in determining
687 which study design(s) should be used. Study details for fertility, EFD, and PPND studies, and
688 combinations thereof, can be found in Annex 9.4. Different approaches are listed below.

689 **6.1 Three separate studies to assess all stages (A□F)**

- 690 • Fertility and Early Embryo Development (FEED)
 - 691 ○ If effects on fertility are suspected, based on mode of action or on the results of
 - 692 repeat dose studies, it can be advisable to dose males and females in separate arms
 - 693 or separate studies comprising mating with untreated animals of the opposite sex.
- 694 • Embryo-Fetal Development (EFD)

695 • Pre- and PostNatal Development, including maternal function (PPND)

696 **6.2 Single study design**

697 A combination of fertility, gestation, and postnatal development (Stages A→F).

698

699 A single study design in rodents might be appropriate when reproductive toxicity is not
700 expected. If such a study provides clearly negative results at appropriately selected doses, no
701 further reproduction studies in that species are warranted. In this study, all newborns and
702 pups, including stillbirths and culled pups, should be examined for morphological
703 abnormalities. If reproductive and developmental toxicity is observed, these toxicity risks
704 should be assessed in detail.

705 **6.3 Two study design**

706 • Combination of FEED and EFD (Stages A→D) + PPND (Stages C→F) studies.
707 This combination of the FEED and EFD, in addition to the PPND study provides all
708 the information obtained from conducting separate FEED and EFD and PPND
709 studies, but uses fewer animals.

710 • Combination of EFD (Stages C→D) + FEED and PPND (Stages A→C + D→F)
711 studies.

712 This combination study design does not include an assessment of external, soft
713 tissues, or skeletal morphology. It is most useful when no treatment-related TEFL
714 effects were observed in the EFD study. The fertility and PPND combined study
715 together with an EFD study, provide all the desired information for all stages of
716 development, but uses fewer animals than the three study design.

717

718 **6.4 Combination design of repeat-dose and fertility studies**

719 In cases where no effects on male or female fertility are expected, or where extending the
720 dosing period is appropriate due to observation of reproductive organ toxicity in long term
721 repeated dose toxicity study, a combination design of repeat-dose and fertility studies can be
722 considered. If effects on fertility are suspected, based on mode of action or on the results of
723 repeat dose studies, it can be advisable to dose males and females in separate studies
724 comprising mating with untreated animals of the opposite sex.

725

726 After a defined dosing period within the longer term repeat-dose toxicity study (e.g., 13- or
727 26-week repeat-dose study), males from the repeat dose study can be cohabited with sexually
728 mature females from a separate study arm (untreated sexually mature females or where the
729 female are treated for at least two weeks prior to mating). This combination study can reduce
730 the number of animals used; however, the number of male animals in the repeat-dose study
731 should be approximately 16 per group. Female animals and their fetuses will be examined
732 for endpoints described in the procedures of the fertility study (Annex Section 9.4.1).

733 The male dose duration period which precedes the period of cohabitation should be
734 determined based on the design principles of the fertility study described in Sections 3.2 and
735 9.4.1. The dosed males used for this assessment can come from any repeat-dose study
736 (e.g., 4-, 13-, or 26-week study) provided the dose duration is sufficient for the project aims,
737 the males are sexually mature, and the number of males available for cohabitation is
738 sufficient to assess effects on male fertility and implant survival. The group size selected to
739 assess male fertility should be justified based on species / strain characteristics. This
740 combination study can reduce the number of dosed males which can be particularly useful
741 with technically challenging exposure routes. It is also particularly useful where evaluation
742 of the long term effects on male reproductive performance is desired.

743 It is possible to assess both male and female fertility simultaneously using males from the
744 repeat-dose toxicity study by cohabiting the males with sexually mature females from a
745 separate study arm that have been treated with drug for at least two weeks. The females and
746 fetuses are assessed as described for the fertility study (Section 9.4.1). However, to detect
747 drug effects on the oestrus cycle, group size should be at least 16 unless justification for
748 smaller group sizes can be provided.

749

750 **6.5 Evaluation of Data**

751 **6.5.1 Data Handling/Data Presentation/Statistics for in vivo Studies**

752 The key to good reporting is the tabulation of individual values in a clear concise manner to
753 account for all animals that are being assessed. Because the data are derived from offspring
754 that are often not directly treated, clear and concise tabulation that permits any individual
755 animal from initiation to termination to be followed should be presented. This will enable
756 assessment of the contribution that the individual has made to any group summary values.
757 Group summary values should be presented with significant figures that avoid false precision
758 and that reflect the distribution of the variable.

759 For the presentation of data on structural changes (e.g., fetal abnormalities) the primary
760 listing (tabulation) should clearly identify the litters containing abnormal fetuses, identify the
761 affected fetuses in the litter and report all the changes observed in the affected fetus.
762 Secondary listings by type of change can be derived from this, as appropriate.

763 Graphical presentations that depict mean values for data collected on multiple days (e.g.,
764 mean body weights) are useful in visualizing a large amount of data. Annex or tabulations of
765 individual values such as bodyweight, food consumption, and litter values, should be
766 concise. While the presentation of absolute values should be the default, calculated values
767 such as bodyweight gain or litter survival indices can provide further support. Where data
768 from non-pregnant animals have been excluded from summary tables, this should be clearly
769 indicated.

770 Presentation of fetal abnormality findings should utilize terminology that is consistent and
771 easily understood.

772 Interpretation of study data should rely primarily on comparison with the concurrent control
773 group. Historical control/reference data are most useful when an interpretation of the data
774 relies on the knowledge of variability within the larger control population and specifically
775 among control groups in previous studies. For example, when trying to understand relevance
776 of malformations, historical control data are useful in interpreting the significance of rare
777 events. The individual laboratory's recent historical control database, if available, is
778 preferred over data compilations from other laboratories. Ideally, the historical data should
779 reflect data from contemporary studies (e.g., from years immediately preceding or following
780 the study conduct, if available) as genetic drift can be an issue.

781 Comparison of study data to the historical mean and standard deviation or range is often
782 performed. It can be important to take into consideration the frequency of the occurrence of
783 an event. If so, then the frequency should be presented.

784 **6.5.2** *Statistics*

785 Developmental and reproductive toxicity studies usually show a distribution of response that
786 does not follow a normal distribution, but can vary from any continuous to any discrete
787 distribution. As a result, this should inform the statistical method used. When employing
788 inferential statistics (determination of statistical significance) the basic unit of comparison
789 should be used. The experimental unit is a concept that is oftentimes misinterpreted but
790 refers to the units that have been randomized and treated. Therefore, cesarean and fetal data
791 should be calculated for the litter as the unit of measure; study result inferences are made
792 back to the mother, not to fetuses. This is because the pregnant females have been allocated
793 to different dose groups (not the fetuses or neonates) and the development of individual
794 offspring in a given litter is not independent. The responses of individual offspring in a given
795 litter are expected to be more alike than responses of offspring from different litters.
796 Similarly, for fertility studies the mating pair should be used as the basic unit of comparison.

797 In most cases, inferential statistics ("significance tests") will evaluate the relationship
798 between a response and treatment factor. The key outputs from a statistical model are then
799 the p-values and confidence intervals for assessing treatment effects – typically pairwise
800 comparisons back to vehicle and/or a trend test across all the groups. The output of such
801 significance tests should only be used as a support for the interpretation of results. Any
802 biologically meaningful difference in treated animals compared with concurrent controls
803 should be discussed. Statistical significance alone does not always constitute a positive
804 signal nor does lack of statistical significance constitute a lack of effect; historical controls,
805 biological plausibility, and reproducibility should be considered in this context. Use of
806 statistical significance alone for drawing inferences when dealing with studies with small
807 group sizes (e.g., NHP) should be approached with caution.

808 **7 PRINCIPLES OF RISK ASSESSMENT**

809 All available data on the pharmaceutical and any related compounds (e.g., surrogates or class
810 alerts), as well as information on human genetics, transgenic animals and the role of the
811 target in reproduction should be considered in this assessment. The amount of information
812 available can depend on the stage of pharmaceutical development, the nature of the
813 pharmaceutical and its intended use. The (projected) human exposure, comparative kinetics
814 between species and plausible mechanism of reproductive toxicity, if available, should be
815 considered.

816 Therapeutic benefit considerations can influence the appropriate level of human risk. For
817 instance, a higher degree of risk could be appropriate for a pharmaceutical intended to treat a
818 life-threatening disease for which all existing therapies have known adverse effects on
819 reproduction than for a life-style pharmaceutical. Human data (e.g., known effects of human
820 genetic variations, clinical trial experience) can greatly influence the overall assessment of
821 human risk of reproductive or developmental toxicity. Definitive human data will supersede
822 nonclinical data.

823 Any limitations (e.g., test system relevance, achieved exposure), uncertainties and data gaps
824 in the available nonclinical reproductive toxicity data package should be addressed and their
825 impact assessed.

826 Risk assessment should generate conclusions relevant for risk communication and
827 management for the intended patient population.

828 **7.1 Risk Assessment for Reproductive and Developmental Toxicities**

829 For human pharmaceuticals, an assessment should be conducted to identify potential risks on
830 human reproduction throughout pharmaceutical development.

831 Endpoints reflecting the full range of potential reproductive and developmental effects as
832 described in Section 2 should be addressed, if not otherwise justified.

833 Not all observations from nonclinical studies are considered to be adverse. An identified
834 effect of the pharmaceutical can also be considered as non-adverse if it is an adaptive change
835 (e.g., enzyme induction) which does not impact on reproductive or developmental function.

836 Adverse nonclinical effects should be evaluated to estimate the likelihood of increased
837 reproductive or developmental risk for humans under the proposed conditions of use of the
838 pharmaceutical. An analysis considering various factors that can increase or decrease the
839 level of concern is recommended. Such factors include animal-human exposure ratio, level of
840 maternal toxicity, dose-response relationship, type of observed effect(s), cross-species
841 concordance, or similarity between pharmacologic and toxicological mechanisms. For
842 example, concern for a reproductive or developmental risk would be increased in the event of
843 a finding observed under any of the following conditions: low relative exposure in animals,
844 cross-species concordance, absence of maternal toxicity, or similarity between
845 pharmacologic and reproductive/developmental toxicological mechanisms. Conversely,

846 concern can be decreased by high relative exposure in animals, absence of cross-species
847 concordance, excessive maternal toxicity or species-specific mechanisms.

848 When assessing effects on embryo-fetal development, one particular difficulty arises when
849 fetal toxicity is observed at dose levels that were also toxic for the mother. It cannot be
850 assumed that developmental toxicity was secondary to maternal toxicity unless such a
851 relationship can be demonstrated either de novo or from published precedence. One way of
852 doing this is to assess the degree of concordance between the severity of toxicity seen in the
853 individual dams and the effects on their litters.

854 Also, the consistency between studies can provide further evidence of an adverse effect of
855 the pharmaceutical (e.g., increased fetal lethality seen in a rodent EFD study consistent with
856 decreased live litter sizes in the PPND study). It is important to consider the exposure at
857 which specific effects were seen across studies and species. Knowledge of the mechanism of
858 reproductive or developmental effects identified in animal studies can help to explain
859 differences in response between species and provide information on the human relevance of
860 the effect (e.g., rodent-specific effects of prostaglandin synthetase inhibitors on
861 cardiovascular fetal development).

862 In general, TEFL are considered to be the critical endpoints in assessing prenatal
863 developmental toxicity. In contrast, reversible or minor manifestations of developmental
864 toxicity (e.g., changes in fetal weight, skeletal variations) by themselves are of minimal
865 concern from a risk assessment perspective. However, an increased incidence of variations
866 can influence the interpretation of an equivocal increase in related malformations. The extent
867 of concern will be influenced by other factors (e.g., exposure multiple at which the findings
868 occurred, cross-species concordance).

869 As in the case of developmental toxicity, reversible or minor manifestations of reproductive
870 toxicity (e.g., a transient inhibition of spermatogenesis) by themselves are of minimal
871 concern from a risk assessment perspective.

872 Comparison of pharmaceutical exposure at the No Observable Adverse Effect Level
873 (NOAEL) in the test species to that at the MRHD is a critical determination. This comparison
874 should be based on the most relevant metric (e.g., AUC, C_{max} , C_{min} , body surface area-
875 adjusted dose). In general, there is increased concern for reproductive or developmental
876 toxicity in humans when effects are seen in a relevant animal species and exposure at the
877 NOAEL is < 10-fold the human exposure at the MRHD. When exposure at the NOAEL is >
878 10-fold the human exposure at the MRHD, the concern is reduced. When the exposure in
879 animals at the NOAEL is > 25-fold the exposure at the MRHD, there is minimal concern for
880 the clinical use of the pharmaceutical (Note 4). If a significant difference in relative
881 exposures is observed between multiple test species, the appropriateness of the metric (e.g.,
882 AUC, C_{max}) being used for the interspecies exposure comparisons should be reassessed.
883 When an alternative metric fails to reduce the disparity between species, the assessment of
884 risk should be based on the most sensitive species. When applicable, the relative exposure
885 ratio should consider both the parent compound and its metabolites.

886 Generally, the results from definitive *in vivo* studies with adequate exposures compared to
887 the exposure at the MRHD carry more weight than those from alternative assays or
888 preliminary studies. Also, the exposure data obtained from *in vivo* studies can be used to
889 determine whether a positive signal identified in an alternative assay presents a risk at the
890 MRHD under the clinical conditions of use of the pharmaceutical.

891 7.2 Risk Assessment for Lactation

892 Generally, evaluations of a pharmaceutical's effects on lactation and its presence in milk in
893 animal studies have little relevance for human risk assessment. Pharmaceuticals can alter the
894 process of lactation in the nursing mother. While the outcome of the PPND (or ePPND) study
895 can inform the risk assessment and can inform as to whether there was extensive systemic
896 exposure in the suckling infant, information on the quantity of the pharmaceutical in milk
897 and production of milk is best derived from human experience, given that the composition of
898 milk varies significantly between rodents and humans. The risk for direct adverse effects on
899 the nursing infant depends on the concentrations of the pharmaceutical and its metabolites in
900 the milk, their absorption, and the age of the infant. Premature infants and neonates have a
901 different capacity to absorb, metabolize and excrete pharmaceuticals compared to older
902 infants.

903

904 8 ENDNOTES

905 **Note 1:** In particular, the testes and epididymides should be sampled and processed using
906 methods which preserve the tissue architecture and permits visualization of the spermatogenic
907 cycles. A detailed qualitative microscopic evaluation with awareness of the spermatogenic
908 cycle is sufficient to detect effects on spermatogenesis. A quantitative analysis of spermatogenic
909 stages (i.e., staging) is not generally recommended but can be useful to further characterize
910 any identified effects. In females, a detailed qualitative microscopic examination of the ovary
911 (including follicles, corpora lutea, stroma, interstitium, and vasculature), uterus and vagina
912 (rodents) should be conducted with special attention given to the qualitative assessment of
913 primordial and primary follicles.

914 **Note 2:** Qualified alternative assays within the context of this guideline can only be applied
915 under certain specific circumstances and have not been subject to formal validation. The EU
916 requires the use of non-animal approaches as soon as they are validated and accepted for
917 regulatory purposes (Directive 2010/63/EU, sector legislation and related guidance).
918 However, this EU directive does not apply to alternative assays qualified according to this
919 guideline.

920 **Note 3:** The ICH Reference Compound List in Annex 9.5.4 is not complete and as such we
921 are soliciting data for additional reference compounds (positive and negative) for potential
922 inclusion into the list, including relevant information as discussed below. These compounds
923 can be either pharmaceuticals or non-pharmaceuticals and should be commercially available.
924 Data to be submitted should include:

- 925 • Name, structure of the compound, suggested compound category, and CAS identifier
926 (if available);
- 927 • The specific TEFL observed in nonclinical test species;
- 928 • Exposures (C_{max} and AUC) at the Lowest Observed Adverse Effect Level (LOAEL) if
929 applicable and the NOAEL;
- 930 • References/sources for the specific data provided (will be made publicly available, if it
931 is not already):
- 932 See examples in Table 9-7 in Annex 9.5.4 for the type of data being requested, as
933 exemplified by four positive compounds (carbamazepine, fluconazole, 5-fluorouracil, and
934 topiramate) and one negative compound (saxagliptin). Data should be summarized using a
935 similar format as that shown in those examples.
- 936 This is not a request for data for the compounds listed in the Table 9-6 in Annex 9.5.4, nor is
937 this a request for examples of assays that could be used.
- 938 **Note 4:** An analysis of 20 known human teratogens showed that if malformations were
939 observed, exposure at the LOAEL in at least one species was < 25-fold the exposure at the
940 MRHD. This indicates that using a > 25-fold exposure ratio for high dose selection in the
941 development toxicity studies would have been sufficient to detect the teratogenic hazard for
942 all these therapeutics. The analysis also showed that for all human teratogens that were
943 detected in animal species the exposure at the NOAEL in at least one species was < 10-fold
944 the exposure at the MRHD.
- 945 In addition, a survey was conducted on EFD toxicity studies by the IQ DruSafe Leadership
946 Group. This survey identified 163 and 152 definitive rat and rabbit EFD studies,
947 respectively, that achieved ≥ 15 -fold animal to human parent drug exposure ratios (using
948 human exposure at the intended therapeutic dose) in the absence of confounding (i.e., dose-
949 limiting) maternal toxicity. An analysis showed that:
- 950 • Of the 163 rat studies, 51 (31%) achieved exposures ≥ 25 -fold human and only 6 (3.7%
951 of total cases) of these had TEFL findings. For all 6 rat cases, the LOAEL was
952 ≥ 50 -fold human exposure, one of which was predicted to be positive based on its
953 mechanism of action.
- 954 • Of 152 rabbit EFD studies, 35 (23%) achieved exposures ≥ 25 -fold human exposure
955 and only 2 (1.3%) of these had TEFL findings. For the 2 rabbit cases, the LOAEL was
956 ≥ 50 -fold human exposure.
- 957 These data show that dosing animals to achieve exposures ≥ 25 -fold human exposures when
958 there is no maternal toxicity (that would otherwise limit the high dose), only infrequently
959 detects a TEFL. In all these cases, TEFL findings were not observed until exposures
960 exceeded 50-fold and findings at such high exposures are not believed to be relevant to
961 human risk assessment. In the absence of confounding (i.e., dose-limiting maternal toxicity),

962 the selection of a high dose for EFD and PPND studies that represents a > 25-fold exposure
963 ratio to human plasma exposure of total parent compound at the intended maximal
964 therapeutic dose is therefore considered pragmatic and sufficient for detecting outcomes
965 relevant for human risk assessment.

966 9 GLOSSARY

967 **Alternative assay(s):** *In-vitro*, *ex-vivo* or non-mammalian *in-vivo* assay(s) intended to
968 evaluate a developmental endpoint (i.e., teratogenicity or embryo/fetal lethality; see TEFL).

969 **Applicability domain:** This describes the types of substances in terms of their physical
970 properties or specific types of substances for which the assay is appropriate. This applies to
971 what types of chemicals can meaningfully be tested in an assay, the applicable chemical
972 space. Examples of applicability could include physicochemical properties of the
973 pharmaceutical such as solubility, volatility, or assay interference by the molecule. The
974 applicability domain also refers to reasons why and conditions under which an assay can be
975 informative or cannot provide useful results. It could include the Training Set of the model
976 for which it is applicable to make predictions for new compounds.

977 **Assay qualification (for regulatory use):** Confirmation of the predictivity of an alternative
978 assay(s) to identify a defined adverse developmental outcome (i.e., TEFL), as outlined in this
979 guideline.

980 **Constitutive ingredients:** Chemicals or biologic substances used as excipients, diluents, or
981 adjuvants in a vaccine, including any diluent provided as an aid in the administration of the
982 product and supplied separately.

983 **Context of use:** For this guideline, context of use applies to regulatory conditions under
984 which the results of an assay can be relied upon. Examples could be: a stand-alone
985 replacement for an *in vivo* study under specified conditions, inclusion in a suite of
986 assays/assessments to replace *in vivo* studies, or to defer definitive studies to later in clinical
987 development.

988 **Developmental toxicity:** Any adverse effect induced prior to attainment of adult life. It
989 includes effects induced or manifested from conception to postnatal life.

990 **GD:** Gestation Day.

991 **GD 0:** The day on which positive evidence of mating is detected (e.g., sperm is found in the
992 vaginal smear / vaginal plug in rodents, or observed mating in rabbits).

993 **Highly targeted or highly selective pharmaceutical/therapeutic:** Therapeutics that exhibit
994 no or minimal off-target effects due to the nature of target binding (e.g., monoclonal
995 antibodies, therapeutic proteins).

996 **ICH Reference Compound List Categories Based on Intended Mechanism of Action:**

- 997 • **Channel modulator:** Compounds with a primary mode of action of targeting cellular
998 channels or transporters.
- 999 • **DNA modifiers:** Compounds with a primary mode of action of either DNA
1000 intercalation or DNA modification (direct [e.g., alkylation, methylation] or indirect
1001 [e.g., based on enzyme modulation]).
- 1002 • **Enzyme Modulator:** Inhibitor, activator, or inducer of enzymes not covered by other
1003 categories (e.g., Kinase Modulator).
- 1004 • **Hormone/Steroids:** Compounds with a primary mode of action of mimicking,
1005 modulating, or antagonizing paracrine, endocrine, or exocrine function.
- 1006 • **Kinase Modulator:** A specific subset of Enzyme Modulators specifically affecting
1007 kinases.
- 1008 • **Nucleoside Modulator/Nutrient Blocker/Central Metabolite Inhibitor:** Anti-
1009 metabolites of nucleosides, nutrients, or metabolic pathway intermediates.
- 1010 • **Oligonucleotide-based Modulators:** DNA or RNA-based oligonucleotides affecting
1011 transcription or translation.
- 1012 • **Receptor Modulator:** Compound that binds to a receptor, either nuclear- or
1013 membrane-based (non-kinase receptor modulators), to elicit a response.
- 1014 • **Secondary Messenger Modulator:** Binding to a target that directly alters cellular
1015 communications between intra- and extra-cellular compartments.
- 1016 • **Others:** Any other compounds that are not part of any of the above categories or for
1017 which there is no intended biological activity (e.g., industrial chemicals).
- 1018 **Malformation:** Permanent structural deviation that generally is incompatible with or
1019 severely detrimental to normal postnatal development or survival.
- 1020 **Modality:** Type of pharmaceutical such as small chemical entity, monoclonal antibody,
1021 oligonucleotide, nanobody, peptide, protein, vaccine.
- 1022 **Normalization Factor:** For the purposes of this guideline; a mathematical algorithm used to
1023 relate the alternative assay result and the *in vivo* observations to the exposures at which they
1024 occur.
- 1025 **Off-target or Secondary Pharmacological Activity:** Action or effect of a pharmaceutical
1026 not related to its intended therapeutic effect.
- 1027 **Pharmacologically Active or Primary Pharmacological Activity:** Eliciting the desired
1028 effects by either directly impacting the target (e.g., inhibition, activation, up regulation, or

- 1029 down regulation) or resulting in the intended physiological outcome (e.g., lower blood
1030 pressure).
- 1031 **PND:** Postnatal day.
- 1032 **PND 0:** Day last offspring of a litter is confirmed as delivered.
- 1033 **Preliminary EFD (pEFD):** A developmental toxicity study that includes exposure over the
1034 period of organogenesis, has adequate dose levels, uses a minimum of 6 pregnant animals per
1035 group, and includes assessments of fetal survival, fetal weight, and external and soft tissue
1036 alterations (see ICH M3(R2) (1)).
- 1037 **Enhanced pEFD:** A pEFD study that is GLP compliant, increases the number of pregnant
1038 animals to ≥ 8 per group, and includes fetal skeletal examinations.
- 1039 **Surrogate molecule:** A molecule showing similar pharmacologic activity in the test species
1040 as that shown by the human pharmaceutical in the human; for a biologic, it can also be
1041 referred to as a homologous protein.
- 1042 **TEFL:** Teratogenic and/or embryofetal lethal.
- 1043 **Teratogen:** For the purpose of this guideline; a pharmaceutical that causes malformations.
- 1044 **Training Set:** A set of data used to discover potentially predictive relationships.
- 1045 **Test Set:** A set of data used to assess the strength and utility of a predictive relationship.
- 1046 **Vaccine:** For the purpose of this guideline, this term refers to preventative or therapeutic
1047 vaccines for infectious diseases. Vaccine (inclusive of the term vaccine product) is defined as
1048 the complete formulation and includes antigen(s) (or immunogen(s)) and any additives such
1049 as adjuvants, excipients or preservatives. The vaccine is intended to stimulate the immune
1050 system and result in an immune response to the vaccine antigen(s). The primary
1051 pharmacological effect of the vaccine is the prevention and/or treatment of an infection or
1052 infectious disease.
- 1053 **Variation:** Structural change that does not impact viability, development, or function (e.g.,
1054 delays in ossification) which can be reversible, and are found in the normal population under
1055 investigation.

1056

1057 **10 REFERENCES**

- 1058 1. International Conference on Harmonisation M3(R2): Guidance on Nonclinical
1059 Safety Studies for the Conduct of Human Clinical Trials and Marketing
1060 Authorization for Pharmaceuticals (2009) together with ICH M3(R2) Questions &
1061 Answers (2012)

1062 2. International Conference on Harmonisation S6(R1): Preclinical Safety Evaluation
 1063 of Biotechnology-Derived Pharmaceuticals (2011)

1064 3. International Conference on Harmonisation (2009). S9: Nonclinical Evaluation for
 1065 Anticancer Pharmaceuticals.

1066

1067 **11 ANNEX**

1068 **11.1 Table of species advantages/disadvantages**

1069 **Table 9-1. Species for Developmental and Reproductive Toxicity Testing**

Species	Advantages	Disadvantages
Routine Species		
Rat	<ul style="list-style-type: none"> • Well-understood biology • Widely used for pharmacodynamics and drug discovery • Robust reproductive capacity with short gestation • Large group sizes and litter size • Suitable for all stages of testing • Widespread laboratory experience and high capacity • Extensive historical data 	<ul style="list-style-type: none"> • Different placentation (e.g., timing, inverted yolk sac) • Dependence on prolactin as the primary hormone for establishment and maintenance of early pregnancy, which makes them sensitive to some pharmaceuticals (e.g., dopamine agonists) <ul style="list-style-type: none"> • Highly sensitive to pharmaceuticals that disrupt parturition (e.g., Nonsteroidal anti-inflammatory drugs in late pregnancy) • Less sensitive than humans to fertility perturbations • Limited application for humanized monoclonal antibodies <ul style="list-style-type: none"> ○ Limited or no pharmacologic activity ○ Limited or no binding ○ Significant anti-drug immune response
Rabbit	<ul style="list-style-type: none"> • Similar advantages to rats plus • Non-rodent model • Readily amenable to semen collection • Placental transfer of antibodies more closely approximates primates than does rodents 	<ul style="list-style-type: none"> • Limitations similar to rat for biologics • Limited historical data for fertility and pre-/postnatal studies • Sensitive to gastrointestinal disturbances; (e.g., some antibiotics) • Prone to spontaneous abortion • Clinical signs difficult to interpret • Not generally used for general toxicology (except for vaccines), lack of kinetic or toxicity data • Limited use for pharmacodynamics

Mouse	<ul style="list-style-type: none"> • Similar advantages to rats • Genetically modified models available or readily generated • Amenable to surrogate approaches • Uses small amounts of test material 	<ul style="list-style-type: none"> • Similar limitations to rats • Small fetus size and tissue volumes • Stress sensitivity • Malformation clusters particularly evident • Less historical data with certain strains • Different placentation (e.g., timing, inverted yolk sac) • Less sensitive than humans to fertility perturbations
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Species	Advantages	Disadvantages
Non-routine Species		
NHP (Details are for Cyno)	<ul style="list-style-type: none"> • Phylogenetically and physiologically more similar to humans • More likely than rodents to show pharmacology and tissue reactivity to human proteins • Placentation similar to human • Larger size and tissue samples • Used in repeat-dose toxicity • Transfer of mAb across the placenta similar to humans 	<ul style="list-style-type: none"> • Low fecundity <ul style="list-style-type: none"> ◦ High background pregnancy loss ◦ Single offspring • Long menstrual cycle (30 days) and gestation (165 days) • Impractical for fertility (mating) studies • Sexual maturity occurs around 3 to 6 years of age • Separation of mother and neonate during postpartum bonding period can be detrimental to neonate • F1 reproduction function difficult to evaluate • Small group size (ethical considerations), hence low statistical power • Animal welfare considerations • Kinetics can differ from humans as much as other species • Limited historical control and laboratory experience/capability • Limited availability of breeding animals • Highly variable age, weight and parity at the start • Uses a large amount of test material

1071

Species	Advantages	Disadvantages
Mini-pigs	<ul style="list-style-type: none"> • Alternate non-rodent for general and reproductive toxicity testing • Susceptibility to some human teratogens • Short period of organogenesis (GD 11-35) • Defined genetic background and specific-pathogen-free animals • Short dose range-finding studies possible (mid-term) • Bred in and adapted to laboratory conditions • Sexual maturity at 3 to 5 months • Good litter size compared to NHP • Suitable for serial semen sampling and mating studies • Monitor pregnancy by ultrasound • Sufficient historical background data on reproductive endpoints 	<ul style="list-style-type: none"> • Limited number of experienced laboratories • Long gestation • Uses a large amount of test material • Large housing requirement • Minimal to no prenatal transfer of antibodies
Limited Use Species (primarily used for investigative purposes)		
Guinea pig	<ul style="list-style-type: none"> • Alternate rodent model that can demonstrate efficacy and cross-reactivity • Placental transfer of antibodies in the last part of gestation is at a similar level in humans 	<ul style="list-style-type: none"> • Historical control and laboratory experience limited to few laboratories • Sensitive to GI disturbances; susceptibility to some antibiotics • Validation of postnatal behavioral and functional tests is limited • Long fetal period • Lack of kinetic or toxicity data • Blood sampling more difficult

1072

Species	Advantages	Disadvantages
Hamster	<ul style="list-style-type: none"> • Alternate rodent model that can demonstrate efficacy and cross-reactivity 	<ul style="list-style-type: none"> • Higher postnatal loss due to cannibalization • Limited historical control and laboratory experience • Validation of postnatal behavioral and functional tests is limited • IV route difficult, can hide orally administered doses in cheek pouches • Aggressive • Sensitive to GI disturbances • Overly sensitive teratogenic response to many chemicals • Lack of kinetic or toxicity data • Blood sampling more difficult
Dog	<ul style="list-style-type: none"> • Usually have repeat-dose toxicity data • Large tissue volume • Readily amendable to semen collection 	<ul style="list-style-type: none"> • Twice yearly ovulators and long gestation (63 days) • Limited historical control and laboratory experience • Validation of postnatal behavioral and function tests is limited • Uses a large amount of test material • Immunogenicity/anaphylaxis concerns
Ferrets	<ul style="list-style-type: none"> • Alternate model that can demonstrate efficacy and cross-reactivity 	<ul style="list-style-type: none"> • Seasonal breeder unless special management system used (success highly dependent on human/animal interactions) • Minimal historical control data and laboratory experience

1073

1074 **11.2 *In vivo* Study Designs**

1075 The number of animals per group specified in individual studies is a balance based on
1076 scientific judgment from many years of experience with these study designs, and ethical
1077 considerations on the appropriate use of animals. Numbers group sizes can be adjusted when
1078 there is evidence either from the pharmacological action of the compound or from existing
1079 studies that the dosages used are expected to elicit an effect at a high frequency and therefore
1080 fewer animals are warranted to confirm the presence of an effect. The number of animals can
1081 differ according to the variable (endpoint) being considered, its prevalence in control
1082 populations (rare or categorical events) or dispersion around the central tendency (continuous
1083 or semi-continuous variables).

1084 For all but the rarest events (such as malformations, abortions, total litter loss), evaluation of
1085 16 to 20 litters for rodents and rabbits tends to provide a degree of consistency among

1086 studies. Below 16 litters per evaluation, between study results become inconsistent, and
1087 above 20 to 24 litters per group, consistency and precision is not greatly enhanced. These
1088 numbers relate litters available for evaluation. If groups are subdivided for different
1089 evaluations the number of animals starting the study should be adjusted accordingly.
1090 Similarly, in studies with 2 breeding generations, 16 to 20 litters should be available for the
1091 final evaluation of the litters of the F1 generation. To permit for natural attrition, starting
1092 group size of the F0 generation of at least 20 is recommended.

1093

1094 Provided below are representative study designs that could be utilized. However, parameters,
1095 timings, and assessments can be readily modified and still meet the study goals. Expert
1096 judgment should be used for adapting these framework designs for individual laboratories
1097 and purposes.

1098 **11.2.1 Fertility and Early Embryonic Development (FEED) Study**

1099 A fertility assessment in rodents is generally recommended (see Sections 3.2 and 4.1). The
1100 aim of the FEED study is to test for toxic effects/disturbances resulting from treatment from
1101 before mating (males/females) through mating and implantation. This comprises evaluation
1102 of stages A and B of the reproductive process (see Section 2). For females, this should detect
1103 effects on the estrous cycle, tubal transport, implantation, and development of
1104 preimplantation stages of the embryo. For males, it will permit detection of functional effects
1105 (e.g., epididymal sperm maturation) that cannot be detected by histological examinations of
1106 the male reproductive organs. The fertility study is designed to assess the maturation of
1107 gametes, mating behavior, fertility, preimplantation stages of the embryo, and implantation.

1108 A combined male/female FEED study is commonly used (See Table 9-2), but separate male
1109 only or female only options are possible by substituting the appropriate number of untreated
1110 males or females in the study designs and should be considered case-by-case.

1111 Table 9-2: FEED Study Design: Rats, combined male and female study

Parameter	Male and Female
Typical Group size	20 + 20
Number of dose groups	4
Administration period ^a	M: ≥ 2 weeks prior to cohabitation through at least confirmation of mating F: ≥ 2 weeks prior to cohabitation through implantation (GD6)
Mating ratio	1 male:1 female
Mating period ^b	≥ 2 weeks
Estrous cycle evaluation	Daily, commencing 2 weeks before cohabitation and until confirmation of mating
Clinical observations/mortality	At least once daily
Body weight	At least twice weekly
Food consumption	At least once weekly (except during mating)
Male euthanasia ^c	Perform macroscopic examination and preserve macroscopic findings, testes and epididymides for possible microscopic examination
Sperm analysis ^d	Optional
Mated female euthanasia ^e	Perform macroscopic examination and cesarean section; preserve macroscopic findings, ovaries and uteri for possible microscopic examination
Scheduled cesarean section: uterine implantation data	Corpora lutea counts, number of implantation sites, live and dead embryos

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- a: Available data (e.g., histopathology, weight of reproductive organs, in some cases hormone assays and genotoxicity data) from toxicity studies should be used to justify dosing duration, especially for detecting effects on spermatogenesis. Provided no effects have been found in repeated dose toxicity studies of at least 2 weeks duration that preclude this, a pre-mating treatment interval of 2 weeks for females and 2 weeks for males can be used. Treatment of males should continue throughout confirmation of mating, although termination following confirmation of female fertility can be valuable. Treatment of females should continue through at least implantation. This will permit evaluation of functional effects on fertility that cannot be detected by histopathological examination in repeated dose toxicity studies and effects on mating behaviour. If data from other studies show there are effects on weight or histology of reproductive organs in males or females, then a more comprehensive study should be considered.
- b: Most rats will mate within the first 5 days of cohabitation (i.e., at the first available estrus), but in some cases females can become pseudopregnant. Leaving the female with the male for up to 3 weeks permits these females to restart estrous cycles and become pregnant.
- c: It can be of value to delay sacrifice of the males until the outcome of mating is known. In the event of an effect on fertility, males could be mated with untreated females to ascertain any potential male mediation of the effect. The males can also be used for evaluation of toxicity to the male reproductive system if dosing is continued beyond mating and euthanasia delayed (e.g., histopathology, sperm analysis (see footnote d)).
- d: Sperm analysis (e.g., sperm counts, motility, and/or morphology) can be used as an optional method to confirm findings by other methods and to characterize effects further.
- e: Termination of females between days 13-15 of pregnancy in general is adequate to assess effects on fertility or reproductive function (e.g., to differentiate between implantation and resorption sites).

1134 **11.2.2 Pre- and Postnatal Developmental (PPND) toxicity study**

1135 A PPND study in rodents is generally warranted (see Sections 3.4 and 4.1). The aim of the
 1136 PPND is to detect adverse effects on the pregnant/lactating female and on development of
 1137 the conceptus and the offspring following exposure of the female from implantation through
 1138 weaning. Since manifestations of effects induced during this period can be delayed,
 1139 observations should be continued through sexual maturity (i.e., stages C through F of the
 1140 reproductive process, see Section 2). The PPND toxicity study is designed to assess
 1141 enhanced toxicity relative to that in non-pregnant females, pre- and postnatal death of
 1142 offspring, altered growth and development, and functional deficits in offspring, including
 1143 maturation (puberty), reproductive capacity at maturity, sensory functions, motor activity,
 1144 and learning and memory.

1146 The females are permitted to deliver and rear their offspring to weaning at which time at least
 1147 one male and one female offspring per litter should be selected for rearing to adulthood and
 1148 mating to assess reproductive competence (see Table 9-3).

1149 Table 9-3: PPND Toxicity Study Design: Rats

Parameter

Typical Group size ^a	Approximately 20 females
Number of dose groups	4
Administration period	From implantation (GD 6/7) through weaning (PND 20/21)

F0 Females

Clinical observations/mortality	At least once daily
Body weight	At least twice weekly
Food consumption	At least once weekly at least until delivery
Parturition observations	GD 21 until complete
Necropsy	PND 21 At necropsy, preserve and retain tissues with macroscopic findings and corresponding control tissues for possible histological evaluation

F1 Pre-weaning

Clinical observations/mortality	Daily from PND 0
Litter size, live and dead	Daily from PND 0
Body weights and sex	PND 1, 4, 7, 14, and 21
Optional Standardization of litter size	≥ PND 4, to 4 or 5 pups per sex
Physical development and reflex ontogeny ^b	Depending on landmark

1150

F1 Post-weaning

Selection for post-weaning evaluation and group size ^c	PND 21, at least 1 male and 1 female/litter where possible to achieve 20 animals per group/sex
Clinical observations/mortality	Daily
Body weight	Weekly
Optional Food consumption	Weekly
Maturation (puberty) ^d	Females: vaginal opening, from PND 30 until complete Males: preputial separation, from Day 40 until complete

Other functional tests ^e	According to standard procedures
Reproductive performance	At least 10 weeks old, paired for mating (1M:1F) within the same group (not siblings)
Terminal procedures of males and females	Preserve organs with macroscopic findings for possible histological evaluation; keep corresponding organs of sufficient controls for comparison Cesarean section: uterine implantation data, corpora lutea counts, number of implantation sites, live and dead embryos

- 1151 a: In studies with 2 breeding generations, 16-20 litters should be available for the final evaluation of the litters of the F1
1152 generation. To permit for natural wastage, the starting group size of the F0 generation should be approximately 20.
1153 b: The best indicator of physical development is bodyweight. Achievement of preweaning landmarks of development such
1154 as eye opening and pinna unfolding as well as others is highly correlated with pup bodyweight. Reflexes, surface
1155 righting, auditory startle, air righting, and response to light are also dependent on physical development. Therefore,
1156 attention should be paid to differences in these parameters when observed in the absence of effects on bodyweight.
1157 c: One animal per sex per litter are retained to conduct behavioral and other functional tests, and to assess reproductive
1158 function. There can be circumstances where more animals per litter can be retained for independent functional
1159 assessments.
1160 d: Bodyweight should be recorded at the time of attainment to determine whether any differences from control are specific
1161 or related to general growth.
1162 e: Investigators are encouraged to adopt methods that would assess sensory functions, motor activity, and learning and
1163 memory. Learning and memory should be evaluated in a complex learning task. Assessments of locomotor activity and
1164 startle reflex with prepulse inhibition (if conducted) should be evaluated over a sufficient period of time to demonstrate
1165 habituation.
1166

1167 **11.2.2.1 Optional Modification of Rodent PPND Study to Assess Juvenile Toxicity**
1168 **Endpoints**

1169 In certain cases when a juvenile animal study is warranted, a PPND study can be modified to
1170 add juvenile toxicity endpoints to potentially reduce animal use and address a specific issue
1171 of concern (1). The following should be considered to support this approach:

- 1172 • Determine the period of exposure appropriate to support the pediatric use.
- 1173 • Demonstrate adequate exposure in the pups *via* the milk and/or consider direct dosing
1174 of pups for the period of developmental interest (TK sampling of the F1 generation
1175 using culled animals during the early post-partum period or study animals shortly
1176 before weaning can provide exposure data and can avoid pre-weaning dosing).

1177 Endpoints included in this modified PPND study should be based on the principles
1178 appropriate for juvenile animal study designs supporting pediatric uses and are not discussed
1179 in this (S5) guidance.

1180

1181 **11.2.2.2 Enhanced Pre- and Postnatal Developmental toxicity study (ePPND) in**
1182 **NHP**

1183 The ePPND toxicity study (Table 9-4) is a study in NHP that combines the endpoints from
1184 both the EFD and PPND studies in which dosing is extended throughout the gestation period
1185 to parturition (i.e., GD20 to parturition). See ICH S6(R1) for information on timing and
1186 additional parameters to be evaluated.

1187 Table 9-4: ePPND Toxicity Study Design: for cynomolgus monkey^a

Parameter	
Group size ^b	Generally ≥ 16 presumed pregnant
Number of dose groups	At least one treatment group plus a control group
Administration period	Initiates upon detection of pregnancy (approximately GD 20) to parturition
F0 Females	
Clinical observations/mortality	At least once daily
Body weight	At least weekly
Parturition observations	Document day of completion
Ultrasound evaluations	Only to track pregnancy status
Necropsy and tissue evaluation	Only as warranted
F1	
Clinical observations/mortality	Daily from PND 0
Body weights	Weekly
Morphometry/Physical development	After PND 0 and at regular intervals
Mother-infant interaction	Minimally in early postnatal period to confirm nursing; as appropriate thereafter
External evaluation	After PND 0 and at regular intervals
Skeletal evaluation	Month 1 and/or later
Visceral evaluation	At necropsy
Necropsy	Variable timing, depends on aim of the evaluations Preserve and retain tissues for possible histological evaluation

1188

1189 a: If an NHP other than the cynomolgus monkey is used, the study design should be adapted accordingly and a rationale
1190 provided.

1191 b: Group sizes in ePPND studies should yield a sufficient number of infants (6-8 per group at postnatal day 7) in order to
1192 assess postnatal development and provide the opportunity for specialist evaluation if warranted (e.g., immune system). Most
1193 ePPND studies accrue pregnant animals over several months. See ICH S6(R1) regarding accrual of animals.

1194 **11.2.3 Embryo-Fetal Developmental (EFD) Toxicity Study**

1195 The aim of the EFD toxicity study is to detect adverse effects on the pregnant female and
1196 development of the embryo and fetus consequent to exposure of the female from
1197 implantation to closure of the hard palate (Table 9-5). This comprises evaluation of stages C
1198 through D of the reproductive process (see Section 2). The embryo-fetal developmental
1199 toxicity study is designed to assess enhanced maternal toxicity relative to that in non-
1200 pregnant females, embryo-fetal death, altered growth, and structural changes.
1201

1202 **11.2.3.1 Dose Range Finding (DRF) Study**

1203 DRF studies in mated females are most often used to select appropriate dose levels, or dose
1204 schedules, for the definitive EFD studies but tolerability and TK data from existing repeat-
1205 dose toxicity can be sufficient for this purpose.

1206 **11.2.3.2 pEFD Study**

1207 The preliminary embryo-fetal developmental toxicity study (Table 9-5) is similar in design
 1208 to the definitive embryo-fetal developmental toxicity study. A typical pEFD study design
 1209 includes dosing over the period of organogenesis, has adequate dose levels, evaluates a
 1210 minimum of 6 pregnant females per group, and includes assessments of fetal survival and
 1211 weight, as well as external and soft tissue examinations (see ICH M3(R2)).

1212 **11.2.3.3 Definitive Embryo-fetal Developmental Toxicity Study**

1213 The females are cesarean sectioned near term and includes assessments of fetal survival and
 1214 weight, as well as external, soft tissue and skeletal examinations (Table 9-5). The timing
 1215 given in Table 9-5 is for rat and rabbit. For other species appropriate timing should be used.

1216 Table 9-5: Embryo-Fetal Developmental Toxicity Study Designs for Rat and Rabbit

Parameter	EFD			pEFD ^a
	Rat	Rabbit		
GLP Status	Yes	Yes		No
Minimum number of litters	16	16		6 (pregnant animal) ^g
Number of dose groups	4	4		4
Administration period ^b	GD6-17	GD7-19		Species appropriate
Antemortem endpoints				
Clinical observations/mortality	At least once daily	At least once daily		At least once daily
Body weight ^c	At least twice weekly	At least twice weekly		At least twice weekly
Food consumption	At least once weekly	At least once weekly		At least once weekly
Toxicokinetics	Yes	Yes		Optional
Postmortem endpoints				
Cesarean section ^d	GD20/21	GD28/29		Species appropriate
Macroscopic examination	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Uterine weight	Optional	Optional		Optional
Corpora lutea	Optional	Optional		Optional
Implant sites	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Live and dead conceptuses	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Early and Late resorptions	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Gross evaluation of placenta	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Fetal body weight	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Fetal sex	<input type="checkbox"/>	<input type="checkbox"/>		<input type="checkbox"/>
Fetal external evaluations ^{e,f}	Yes	Yes		Yes
Fetal soft tissue evaluations ^{e,f}	Yes	Yes		Yes
Fetal skeletal evaluations ^{e,f}	Yes	Yes		No

1217

- 1218 a: In an enhanced pEFD study the number of pregnant animals should be increased from 6 to ≥ 8 per group, include fetal
 1219 skeletal examinations, and it should be conducted in accordance with GLP regulations.
 1220 b: Females are dosed with the test substance from implantation to closure of the hard palate (i.e., stage C of the reproductive
 1221 process, see Section 2).
 1222 c: Daily weighing of pregnant females during treatment can provide useful information.
- 1223 d: Cesarean sections should be conducted approximately one day prior to parturition. Preserve organs with macroscopic
 1224 findings for possible histological evaluation; keep corresponding organs of sufficient controls for comparison.

1225 e: All fetuses should be examined for viability and abnormalities. To permit subsequent assessment of the relationship
1226 between observations made by different techniques fetuses should be individually identified. It is critical to be able to
1227 relate all findings by different examination techniques (i.e., body weight, external inspection, soft tissue and/or skeletal
1228 examinations) to a single specimen in order to detect patterns of abnormalities.

1229 f: It is preferable to examine all fetuses for both soft tissue and skeletal alterations, if permitted by the methods employed
1230 (e.g. fresh dissection or μ CT, MRI, etc.). When using techniques precluding evaluation of both soft tissue and skeletal
1231 changes in the same fetus, 50% of fetuses from each litter should be allocated to each examination. The internal soft
1232 tissues of the head should be examined in at least 50% of the fetuses.

1233
1234 g: Minimum number of litters equals the number of pregnant animals per group, not the number of litters for pEFD studies.

1235 11.2.4 **Combination Studies**

1236 11.2.4.1 **Fertility and Embryonic Development (FEFD)**

1237 The aim of the combined FEFD study is to test for toxic effects/disturbances resulting from
1238 treatment from before mating (males/females) through mating, implantation and until the
1239 end of organogenesis. This comprises evaluation of stages A to C of the reproductive
1240 process (see Section 2).

1241 A combined male/female FEFD is commonly used, but a separate female only option is
1242 possible where male fertility is assessed in a separate study such as a repeat dose study of
1243 suitable duration. The study would then use untreated males for mating purposes only. For
1244 specific study design and observational parameters see Sections 9.4.1 and 9.4.3 (FEED and
1245 EFD).

1246 11.2.4.2 **Fertility and PPND (FPPND)**

1247 The aim of the combined Fertility and Pre- and Postnatal Development study (FPPND) study
1248 is to test for toxic effects/disturbances resulting from treatment from before mating
1249 (males/females) and to detect adverse effects on the pregnant/lactating female and on
1250 development of the conceptus and the offspring following exposure of the female from
1251 implantation through weaning. Since manifestations of effects induced during this period
1252 can be delayed, observations should be continued through sexual maturity. This comprises
1253 evaluation of stages A to F of the reproductive process (see Section 2). The pre- and
1254 postnatal developmental toxicity study is designed to assess enhanced toxicity relative to
1255 that in non-pregnant females, pre- and postnatal death of offspring, altered growth and
1256 development, and functional deficits in offspring, including behavior, maturation (puberty)
1257 and reproductive capacity at maturity.

1258 The study design features should encompass those of the individual studies in terms of the
1259 number of animals used and the parameters assessed. For specific study design and
1260 observational parameters see Sections 9.4.1 and 9.4.2 (FEED and PPND, respectively).

1261 A combined male/female FPPND can be used, but a separate female only option is possible
1262 where male fertility is assessed in a separate study such as a repeat dose study of suitable
1263 duration. The study would then use untreated males for mating purposes only.
1264

1265 **11.3 Qualification of Alternative Test Systems for Regulatory Acceptance**

1266 A framework and testing scheme to facilitate the qualification of alternative assays,
1267 including a list of test compounds (ICH Reference Compound List), is provided in this
1268 section. The ICH Reference Compound List provides information on embryo-fetal toxicity
1269 for various reference compounds, organized by overarching categories. This list is generated
1270 recognizing that the context of use will inform on acceptability of particular alternative
1271 assessments. Performance factors for assay acceptance are also outlined. The ICH Reference
1272 Compound List is intended to be periodically updated.

1273 The applicability domain (see Glossary) together with the intended regulatory context of use
1274 influences the factors for assay qualification and the rigor for achieving regulatory
1275 acceptance.

1276 **11.3.1 Selection Factors for the ICH Reference Compound List**

1277 The ICH Reference Compound List aims to cover reference compounds known for their
1278 TEFL effects in animals or humans, even if the mode of action is uncertain.

1279 Availability of data showing clear TEFL effects in rats and/or rabbits in the absence of
1280 maternal toxicity represents an essential inclusion criterion for the selected positive
1281 compounds. This includes, when available, the multiples comparing human exposure to
1282 animal exposures where effects were seen.

1283 Availability of pharmacokinetic and toxicokinetic data in the test species is an important
1284 criterion for the selection of reference compounds. Thus, all compounds used should have
1285 non-clinical exposure data (C_{max} and/or AUC) under the approximate conditions tested
1286 yielding either negative or positive results in the *in vivo* studies for the species being
1287 predicted. While pharmaceuticals are preferred, other chemicals can be considered. The
1288 ICH Reference Compound List does not currently include biotechnology-derived
1289 pharmaceuticals. The list favors compounds with direct effects on the fetus; however, a few
1290 are known to depend on cytochrome P450 metabolic activation to cause TEFL. Cytotoxic
1291 and/or genotoxic compounds are included to a limited extent because they are expected to
1292 induce TEFL through their intrinsic property of preferentially damaging rapidly dividing
1293 cells.

1294 The performance of alternative assay(s) to detect species-specific differences can be
1295 evaluated by testing reference compounds known to cause TEFL in a single species;
1296 however, the number of such compounds available in the public domain is limited.

1297 Compounds not causing TEFL (negative compounds) are also included in the ICH
1298 Reference Compound List to permit assessment of assay specificity. These compounds can
1299 be negative at all *in vivo* doses tested, or can be positive (TEFL observed) at higher
1300 doses/exposures, provided the alternative assay predicts the transition from negative to
1301 positive. The alternative assay should predict a negative result at some extrapolated multiple
1302 under the conditions for which the *in vivo* study yielded a negative result (no TEFL).

1303 Further, the ICH Reference Compound List includes compounds from different
1304 chemical/pharmacologic classes with overlap with both negative and positive compounds to
1305 enable adequate coverage of the alternative assay for pharmaceuticals and diverse chemical
1306 structures and mode of action.

1307 It is not critical for assay qualification purposes that the exposures achieved in animals that
1308 resulted in negative or positive TEFL outcome exceed the human exposures. This is in
1309 contrast to application of assay results for risk extrapolation where preferably the highest
1310 doses/exposures tested are at or above MRHD.

1311 Finally, the commercial availability of the selected compounds of appropriate quality was
1312 considered in the generation of the list.

1313 **11.3.2 Performance Factors**

1314 To be appropriate for regulatory use, the alternative assay(s) should be characterized using
1315 the ICH Reference Compound List. The list is not exhaustive and the recommendations
1316 provided are based on available information and pragmatic considerations. At least 45
1317 compounds in total should be tested. Other compounds can substitute for the non-core
1318 compounds, but their use should be justified according to the inclusion factors mentioned
1319 above.

1320 The compounds are distributed into multiple classes, covering a wide range of biological and
1321 chemical activities. All classes should be tested (at least 2 or 3 compounds from each class).
1322 An approximate 2:1 ratio of positive to negative compounds should be tested because it is
1323 important to identify positive compounds, but this ratio also ensures selectivity with the
1324 limited number of compounds available. For safety assessment purposes, and for some
1325 contexts of use, the false negative rate can be more important than the false positive rate.

1326 The sensitivity to detect a positive signal in an assay(s), should be at least 80%, with
1327 evidence of selectivity (i.e., differentiating between true positives and true negatives).

1328 The evaluation should identify the applicability domain and any limitations of the assay(s),
1329 and include assessments of accuracy, and reproducibility over time. Inter-laboratory
1330 reproducibility and transferability should be established if a particular assay is to be used in
1331 more than one laboratory.

1332 Individual assays or combinations of assays can be used to predict TEFL. The performance
1333 characteristics of each individual assay as well as the performance of the combined battery, if
1334 used, should be specified. Various statistical methods are available for determining which
1335 combination of assessments will give the best predictivity.

1336 **11.3.3 Assay Qualification Information to be Provided to Health Authorities**

1337 To enable evaluation of an alternative assay(s) for use in risk assessment for regulatory
1338 purposes, the following information should be provided.

1339 A detailed description should be presented concerning what the predictive model is, what
1340 species (e.g., rat, rabbit, and/or human outcomes) it is trying to predict, and what
1341 reproductive endpoint it assesses. The predictive model can consist of a single assay or a
1342 battery of assays used together to predict the endpoint of interest (e.g., TEFL) in the
1343 respective species such as rat. If a battery of assays is used, each should be fully described.
1344 The specific endpoint(s) used (e.g., gene signature, morphology) should be described and
1345 how the assessment is made, including how the endpoints were selected and the specific
1346 factors for positive and negative determinations, should be discussed.

1347
1348 The details of the algorithm employed for determining positive and negative outcomes from
1349 assay observations should also be presented. The predictive model should correlate
1350 concentrations tested in the alternative assay(s) to the *in vivo* exposure that results in an
1351 adverse outcome in the species being predicted. For example, concentrations associated with
1352 positive effects on the endpoint should take into consideration *in vivo* exposure such as C_{max}
1353 or AUC. This permits the model to be used for exposure-based risk assessment. The
1354 pharmacokinetic parameter used including any normalization factors employed to correlate
1355 with *in vivo* results should be presented (Section 3.5.3).

1356
1357 The compound list used to qualify the assay performance should be presented.
1358 Documentation should include a clear identification of the compound list used as the
1359 Training Set (see Glossary) to develop the assay, and the compound list used as the Test Set
1360 (see Glossary) to evaluate the assay's performance. The assay Training Set can include
1361 compounds of the sponsor's choice not on the ICH Reference Compound List. Additional
1362 compounds not in the ICH Reference Compound list can be used as part of the Training Set
1363 or the Test set, but not both. No more than 15% compounds from the ICH Reference
1364 Compound List can be used for the Training Set. This permits an adequate number of
1365 compounds from the ICH Reference Compound List to be used as part of the Test Set for
1366 qualification purposes. Reserving $\geq 85\%$ of compounds from the ICH Reference Compound
1367 List for the Test Set permits a sufficiently robust evaluation of the assay's predictivity.

1368
1369 The performance of the Training and Test sets should be evaluated separately and together
1370 and the results of each analysis presented. The performance summary should list the
1371 sensitivity, specificity, positive predictive value, and negative predictive value. If more than
1372 one assay is used, the performance of each assay should be provided separately in addition to
1373 the integrated assessment used for the predictive model. In the case of integration of more
1374 than one assay in the model, a clear description should be presented of how the integration of
1375 the individual assays is conducted to arrive at the integrated predictive model.

1376
1377 As part of the assay qualification and predictive model use, the category of compounds the
1378 assay can and cannot predict (e.g., a component of the applicability domain) should be
1379 defined from the following list of categories included in the ICH Compound Reference List
1380 (see Glossary): Channel modulator, DNA modifiers, Enzyme modulator, Hormone/steroids,
1381 Kinase modulator, Nucleoside modulator/nutrient blocker/central metabolite inhibitor,
1382 Receptor modulator, Oligonucleotide-based modulators, secondary messenger modulator,
1383 and Others. Additionally, human teratogens not detected *in vivo* by rat and/or rabbit should

1384 also be evaluated to understand if the assay can detect them, even if the assay(s) intended use
 1385 is to predict rat or rabbit outcomes. These results should be presented separately and the
 1386 sponsor should justify whether or not and if so, how, to include these results in their
 1387 predictivity assessment.

1388
 1389 Demonstration of assay reproducibility should be assessed and can be accomplished by
 1390 inclusion of at least one positive control and one negative control in either each assay run or
 1391 interspersed over time between test compound runs. The sponsor should justify their
 1392 approach to inclusion of positive and negative controls. The approach used to demonstrate
 1393 assay reproducibility should be described in the information provided. Additionally, several
 1394 of the compounds from the ICH Reference Compound List should be periodically reassessed
 1395 and the data provided along with compounds being evaluated for therapeutic development.
 1396 The source of reagents, biologic materials, and compounds tested should be provided.
 1397 Likewise, the source/reference of all *in vivo* exposure data used for compounds in the
 1398 qualification data set should also be presented, except for those compounds in the ICH
 1399 Reference Compound List since that would be the source (reference) information. Assays
 1400 should be developed with the understanding there is an expectation that regulatory studies
 1401 should generally be conducted in compliance with GLP.

1402
 1403 The sponsor of the alternative assay should state whether the assay qualification has been
 1404 previously submitted to any health authority in support of reproductive toxicity assessments
 1405 and, if so, to which one(s).
 1406

1407 **11.3.4 ICH Reference Compound List**

1408 The ICH Reference Compound List (Table 9-6) is not intended to cover tailored approaches
 1409 studying specific pharmaceutical targets or chemistry of structurally related analogs. For
 1410 particular pharmaceuticals and contexts of use, justification for use of particular
 1411 assays/assessments should be given (e.g., the Sponsor has *in vivo* information on other
 1412 pharmaceuticals in the class). Table 9-7 provides examples of data records for including
 1413 compounds in the ICH Reference Compound List for qualifying alternative assays.

1414 **Table 9-6. ICH Reference Compounds for Qualifying Alternative Assays**
 1415

Category	Positive Controls	Negative Controls
Channel Modulator	Sotalol	Hydrochlorothiazide
	Almokalant	Chlorthalidone
	Diltiazem	
	Topiramate	
	Trimethadione	
	Phenytoin (Diphenylhydantoin)	
	Carbamazepine	
DNA Modifiers	Cyclophosphamide	

Category	Positive Controls	Negative Controls
	Busulfan	
	Cisplatin	
	Thiotepa	
Enzyme Modulator	Aspirin	
	Captopril	Saxagliptin
	Enalapril	Vildagliptin
	Methimazole (Thiamazole)	
Hormone/Steroid	Dexamethasone	Progesterone
	Fluticasone	
Kinase Modulator	Afatinib	
	Ceritinib	
	Dabrafenib	
	Dasatinib	
	Ibrutinib	
	Pazopanib	
	Tacrolimus	
Nucleoside Modulator/ Central metabolite inhibitor	Imatinib	
	Cytarabine	
	5-Fluorouracil	
	Hydroxyurea	
	Methotrexate	
	Ribavirin	
	Teriflunomide	
Other	Warfarin	
	Artesunate / amodiaquine	Amoxicillin
	Clarithromycin	Clindamycin
	Doxycycline	Cyclobenzaprine
	Fluconazole	Erythromycin
	Pomalidomide	Sulfasalazine
	Tafamidis	
	Telavancin	
	Thalidomide	
Receptor Modulator	Valproic acid	
		Cetirizine
	Bosentan	Cyproheptadine
	Clobazam	Doxylamine
	Fingolimod	Maraviroc
	Metoclopramide	
	Plerixafor	

Category	Positive Controls	Negative Controls
	Sumatriptan	Nizatidine
Second Messenger Modulator	Theophylline	
Transcription Modulator	Acitretin	
	Isotretinoin (13- <i>cis</i> -retinoic acid)	
	Vismodegib	

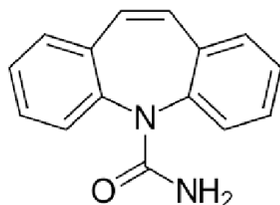
1416 **Table 9-7. Examples of Data Records for Including Compounds in Reference List for Qualifying**
 1417 **Alternative Assays**

1418 **Carbamazepine**

1419 **Proposed Class:** Other

1420 **CAS No.:** 298-46-4

1421 **Structure:**



1422

Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
250 mg/kg/day Fasted 200 mg/kg single PO dose: C _{max} = 32.7 µg/mL [3] (extrapolates to 41 µg/mL at 250 mg/kg) AUC _(0-24h) = 32.8 mg•min/mL = 547 µg•h/mL (extrapolates to 684 µg•h/mL at 250 mg/kg)	400 mg/kg Fasted 200 mg/kg single PO dose: C _{max} = 32.7 µg/mL [3] (extrapolates to 65 µg/mL at 400 mg/kg) AUC _(0-24h) = 32.8 mg•min/mL = 547 µg•h/mL (extrapolates to 1094 µg•h/mL at 400 mg/kg)	<u>650 mg/kg [2]</u> Maternal toxicity increased resorptions, increased skeletal and visceral abnormalities (4/119 offspring showed cleft palate, talipes, or anophthalmos) <u>600 mg/kg [4]</u> increased resorptions, increased skeletal and visceral abnormalities (edema and kinked tails)	NOAEL was not identified	225 mg/kg/day Exposure data available for 80 mg/kg [5]: C _{max} = 10.4 µg/mL (extrapolates to 29 µg/mL at 225 mg/kg) AUC _(0-24h) = 94.8 µg•h/mL (extrapolates to 267 µg•h/mL at 225 mg/kg)	Dosed 225 – 450 mg/kg [1] No malformations Decreased numbers of fetuses, increased resorptions in all groups Maternal toxicity at 450 mg/kg	Carbamazepine 10,11-epoxide metabolite present

Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
		<p><u>400 mg/kg [1, 2, 4]</u> Reduced maternal weight gain; increased visceral abnormalities; abortions</p> <p><u>250 mg/kg [1, 2]</u> kinked ribs in 2/119 fetuses (not considered a TEFL finding)</p>				
<ol style="list-style-type: none"> 1. Published Pharm/tox review of NDA 16-608 (December 19, 1967), 16608/S-000 Part 02. 2. Equetro (carbamazepine) extended-release capsules Label, Carbamazepine FDA approval package, Label 021710/S-011, S-012. 3. Shi L, Dang XL, Liu XY, Wei HM, Yang MM, Zhang Y. Effect of <i>Sophora flavescens</i> on the pharmacokinetics of carbamazepine in rats. Arch Pharm Res. 2014;37:1617-23. 4. Vorhees CV, Acuff KD, Weisenburger WP, Minck DR. Teratogenicity of carbamazepine in rats. Teratology. 1990;41:311-17. 5. Koumaravelou K, Adithan C, Shashindran CH, Asad M, Abraham BK. Effect of honey on carbamazepine kinetics in rabbits. Indian J Exp Biol. 2002;40(5):560-3 						

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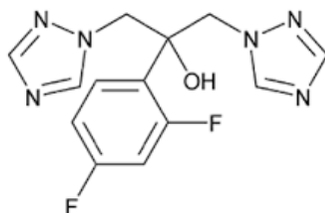
1424 **FLUCONAZOLE**

1425 **Proposed Class:** Other

1426 **CAS No.:** 86386-73-4

1427 **Structure:**

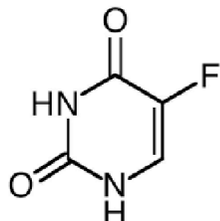
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1429

Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
50 mg/kg Following 20 mg/kg single oral dose: C _{max} [2] = 13.5 µg/mL (extrapolates to 34 µg/mL at 50 mg/kg) AUC [1] = 152 µg•hr/mL (extrapolates to 380 µg•h/mL at 50 mg/kg)	80 mg/kg 20 mg/kg single oral dose: C _{max} = 13.5 µg/mL [3] (extrapolates to 54 µg/mL at 80 mg/kg) AUC = 152 µg•h/mL [1] (extrapolates to 608 µg•h/mL at 80 mg/kg)	<u>80 –320 mg/kg [2, 3]</u> Increased embryolethality and fetal abnormalities (wavy ribs, cleft palate, and abnormal cranio-facial ossification) <u>≥25 mg/kg</u> Increases in fetal anatomical variants (supernumerary ribs, renal pelvis dilation) and delays in ossification were observed at 25 and 50 mg/kg and higher doses <u><10 mg/kg</u> No fetal effects	≤ 25 mg/kg 10 mg/kg single oral dose: C _{max} = 10.8 µg/mL (extrapolates to 27 µg/mL at 25 mg/kg)	75 mg/kg [2, 3] 10 mg/kg single oral dose: C _{max} = 10.8 µg/mL (extrapolates to 81 µg/mL at 75 mg/kg)	<u>75 mg/kg</u> Abortions	
<ol style="list-style-type: none"> Humphrey MJ, Jevons S, Tarbit MH. Pharmacokinetic evaluation of UK-49,858, a metabolically stable triazole antifungal drug, in animals and humans. Antimicrob Agents Chemother. 1985 Nov;28(5):648-53. Published Pharm/tox review of NDA 20322 (June 30, 1994), Part 01 Diflucan (Fluconazole) FDA Prescribing Information 						

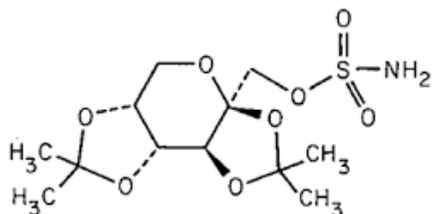
1430 **5-FLUOROURACIL**
 1431 **Proposed Class:** Nucleoside modulator
 1432 **CAS No.:** 51-21-8
 1433 **Structure:**
 1434



Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
15 mg/kg single dose IP (Ku wagata) 30 mg/kg , IP (Zhang) C _{max} = 7.74 µg/mL (extrapolates to 3.87 at 15 mg/kg) AUC = 11.66 µg•h/mL (extrapolates to 5.83 at 15 mg/kg)	12 – 37 mg/kg single IP dose on GD11 or 12 (Chaube) 17 mg/kg single dose IP on GD 9 (Ku wagata) 30 mg/kg , IP (Zhang) C _{max} = 7.74 µg/mL (extrapolates to 4.4 at 17 mg/kg) AUC = 11.66 µg•h/mL (extrapolates to 6.6 at 17 mg/kg)	<u>12 – 37 mg/kg</u> (Chaube) Cleft palate and deformed appendages <u>≥17 mg/kg</u> (Ku wagata) micro-anophthalmos, craniofacial defects, hydrocephaly, brain hernia, edema; embryo lethality at 30 mg/kg <u>≥15 mg/kg</u> decreased fetal weight	Not determined, <40 mg/kg	40 mg/kg SC GD12 (480 mg/m ²) PK: 20 mg/kg IV (Kar) C _{max} = 427 nmol/mL =55 µg/mL (extrapolates to 110 at 40 mg/kg) AUC = 2535 nmol•min/mL = 5.5 µg•h/mL (extrapolates to 11 at 40 mg/kg)	<u>40 mg/kg</u> (DeSesso) 2/5 females died, with fetuses of surviving females exhibiting anomalies of the limb in 85% of cases	5FU is a pro-drug: thymidylate synthetase inhibitor is 5FdUMP MW = 130.077 g/mol
Chaube S, Murphy ML. The teratogenic effects of the recent drugs active in cancer chemotherapy. In: Advances in Teratology. ed. DHM Woolham. Academic Press, New York. 1968 DeSesso, JM, Scialli AR, Goeringer GC. Teratology. 1995;51:172 (abstract) Kar R, Cohen RA, Terem TM, Nahabedian MY, Wile AG. Pharmacokinetics of 5-fluorouracil in rabbits in experimental regional chemotherapy. Cancer Res. 1986;46(9):4491-5.						

Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
Ku wagata M, Takashima H, Nagao T. A comparison of the <i>in vivo</i> and <i>in vitro</i> response of rat embryos to 5-fluorouracil. J Vet Med Sci. 1998;60(1):93-9.						
Zhang C, Li G, Wang Y, Cui F, Zhang J, Huang Q. Preparation and characterization of 5-fluorouracil-loaded PLLA-PEG/PEG nanoparticles by a novel supercritical CO ₂ technique. Int J Pharm. 2012;436(1-2):272-81.						

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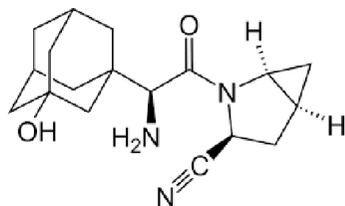
1436 **TOPIRAMATE**1437 **Proposed Class:** Channel Modulator1438 **CAS No.:** 97240-79-41439 **Structure:**

Rat NOAEL Dose AUC C _{max}	Rat LOAEL Dose AUC C _{max}	Rat Findings	Rabbit NOAEL Dose AUC C _{max}	Rabbit LOAEL Dose AUC C _{max}	Rabbit Findings	Notes
100 mg/kg Exposure (FDA pharmtox review) 30 mg/kg, female SD, 8 doses C _{max} = 22.2 µg/mL (extrapolates to 74 at 100 mg/kg)	400 mg/kg Exposure (FDA pharmtox review) 30 mg/kg, female SD, 8 doses C _{max} = 22.2 µg/mL (extrapolates to 296 µg/mL at 400 mg/kg)	<u>≥400 mg/kg</u> (FDA pharmtox review and/or topamax label) limb defects (ectrodactyly, micromelia, and amelia) <u>≥20 mg/kg</u>	10 mg/kg Exposure (FDA pharmtox review) 60 mg/kg, females, 14 doses C _{max} = 39 µg/mL (extrapolates to 6.5 at 10 mg/kg) AUC = 201	35 mg/kg Exposure (FDA pharmtox review) 60 mg/kg, females, 14 doses C _{max} = 39 µg/mL (extrapolates to 23 at 35 mg/kg) AUC = 201 µg•h/mL	<u>≥35 mg/kg</u> (FDA pharmtox review and/or topamax label) Embryofetal mortality increased at ≥35 mg/kg; Teratogenic effects (primarily rib/vertebral malformations) were observed at 120 mg/kg	In rats: maternal toxicity were seen at ≥400 mg/kg and maternal body weight gain was reduced at ≥100 mg/kg In rabbits: maternal toxicity (decreased body weight gain, clinical signs, and/or mortality)

Rat NOAEL Dose AUC C_{max}	Rat LOAEL Dose AUC C_{max}	Rat Findings	Rabbit NOAEL Dose AUC C_{max}	Rabbit LOAEL Dose AUC C_{max}	Rabbit Findings	Notes
AUC = 268 µg•h/mL (extrapolates to 893 at 100 mg/kg) In pregnant rats dosed w/ 200 mg/kg, at GD12-15, C _{1.5h} = 97 µg/mL (extrapolates to 49 at 100)	AUC = 268 µg•h/mL (extrapolates to 3573 at 400 mg/kg) In pregnant rats dosed w/ 400 mg/kg, at GD12-15, C _{1.5h} = 169 µg/mL	reduced fetal body weights and increased incidence of structural variations	µg•h/mL (extrapolates to 33.5 at 10 mg/kg)	(extrapolates to 117 at 35 mg/kg)		was seen at ≥35 mg/kg Rabbit LOAEL margins all <10
Topamax label (US): rat: oral doses of 20, 100, and 500 mg/kg or 0.2, 2.5, 30, and 400 mg/kg; rabbit: oral doses of 20, 60, and 180 mg/kg or 10, 35, and 120 mg/kg Published Pharm/tox review of NDA 20505/S000 (August 1, 1995)						

1440

1441 **SAXAGLIPTIN**
 1442 **Proposed Class:** Enzyme modulator
 1443 **CAS No.:** 361442-04-8
 1444 **Structure:**



1445
 1446

Rat NOAEL (Highest Dose Tested) Dose, AUC, C_{max}	Rat LOAEL	Rat Findings	Rabbit NOAEL (Highest Dose Tested) Dose, AUC, C_{max}	Rabbit LOAEL	Rabbit Findings	Notes
900 mg/kg C _{max} = 62 µg/mL AUC = 647 µg•h/mL	Not relevant	No malformations or embryofetal lethality noted. ≥240 mg/kg delayed ossification	200 mg/kg C _{max} = 34 µg/mL AUC = 111 µg•h/mL	Not relevant	No malformations or embryofetal lethality 200 mg/kg increased ossification	
Published FDA Pharm/tox review of NDA 022350/S000, Parts 2, 3, and 5 (March 3, 2009). Rat: oral dosages of 64, 240 and 900 mg/kg; rabbit: oral dosages of 8, 40 and 200 mg/kg						

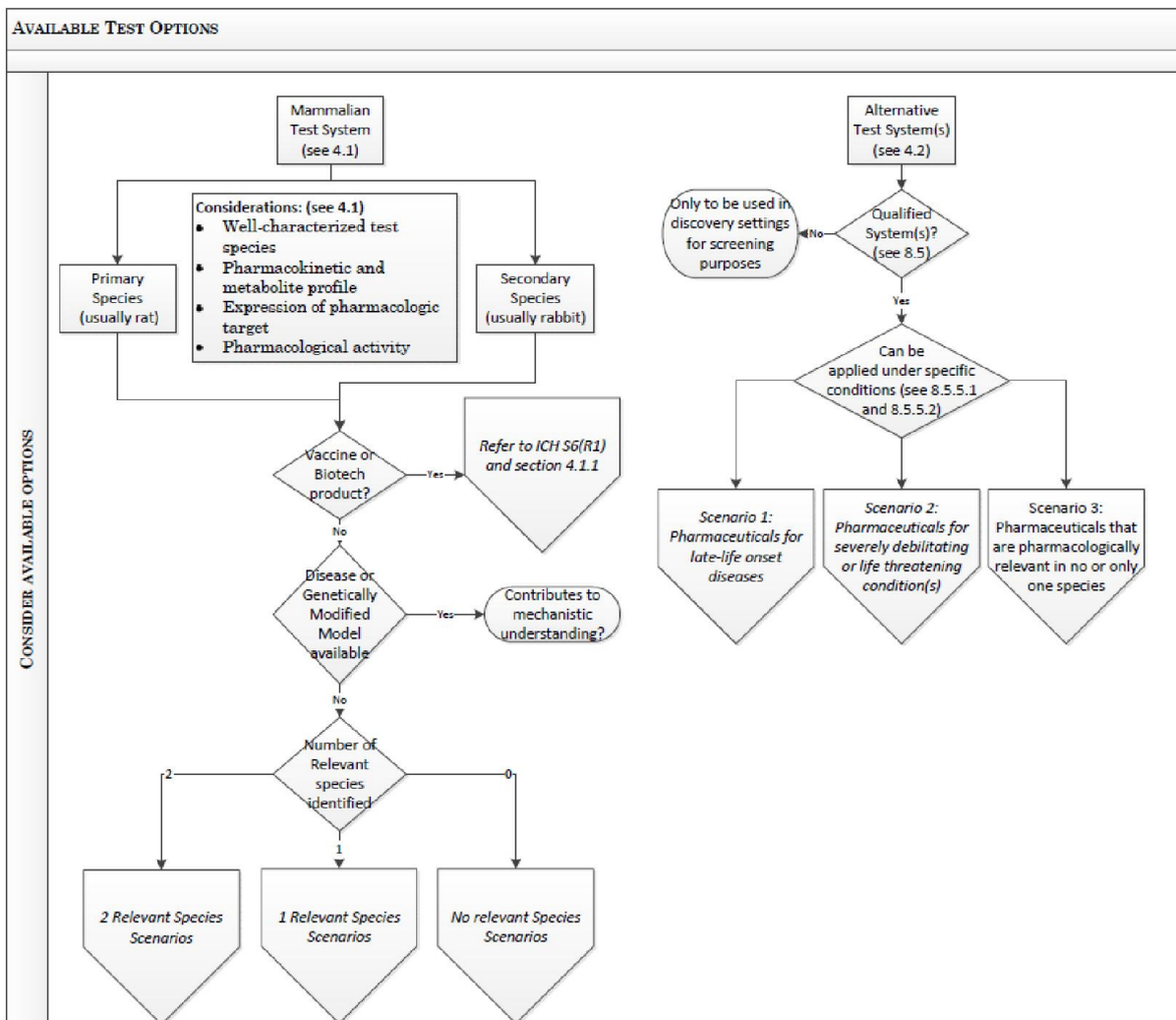
1447

1448 **11.3.5 Examples of EFD Testing Strategies**

1449 This section describes optional integrated testing strategies that can be used to detect adverse
 1450 effects on EFD. The use of a particular scenario needs to be justified.

1451 In circumstances other than those described in 9.5.5.1 and 9.5.5.2 below and elsewhere in
 1452 this guideline where use of alternative assays is proposed, positive results in alternative
 1453 assays can also reduce mammalian *in vivo* testing. In contrast, negative results in alternative
 1454 assays in most of these other circumstances would not be anticipated to reduce *in vivo*
 1455 testing. See Figure 9-1.

1456 Figure 9-1: Summary of Available Test Options



1457

1458 **11.3.5.1 Scenarios applicable when there are at least 2 relevant mammalian species**
1459 **(crf. Species selection)**

1460 This section describes optional integrated testing strategies that can be used to detect adverse
1461 effects on embryo-fetal development. The use of a particular testing strategy should be
1462 justified.

1463 **a) Scenario 1: Pharmaceuticals for late-life onset diseases (Figure 9-2)**

1464 1. When a qualified alternative assay predicts TEFL in one species (e.g., rat) or is
1465 equivocal, an EFD assessment (e.g., pEFD, enhanced pEFD) in another species (e.g.,
1466 rabbit) should be conducted to evaluate the multi-species risk and assess the finding
1467 *in vivo*.

1468 a. If TEFL is observed in the *in vivo* study (e.g., rabbit), the pharmaceutical will be
1469 considered to induce TEFL in multiple species based on the alternative assay and
1470 *in vivo* results.

1471 b. If no TEFL is detected in the *in vivo* study, a definitive EFD should be conducted
1472 in the species corresponding to the alternative assay to further assess the TEFL
1473 potential *in vivo*. If TEFL is observed in this definitive *in vivo* EFD study, the
1474 pharmaceutical will be considered positive in animal studies based on the
1475 positive alternative assay and *in vivo* for the same species. No further EFD
1476 studies are warranted, as a hazard has been identified and the risk assessment can
1477 be made based on the totality of the information. If no TEFL is observed in both
1478 *in vivo* EFD studies, the results from the alternative assay represent a false
1479 positive and the pharmaceutical will be considered not likely to induce TEFL,
1480 provided adequate exposure was achieved in the *in vivo* testing (e.g., exposures
1481 *in vivo* exceed the human exposure).

1482 2. When an alternative assay predicts a negative outcome (i.e., no TEFL) in one species
1483 (e.g., rat), an EFD study in another species (e.g., rabbit) should be conducted to
1484 determine if the pharmaceutical is positive for TEFL *in vivo*.

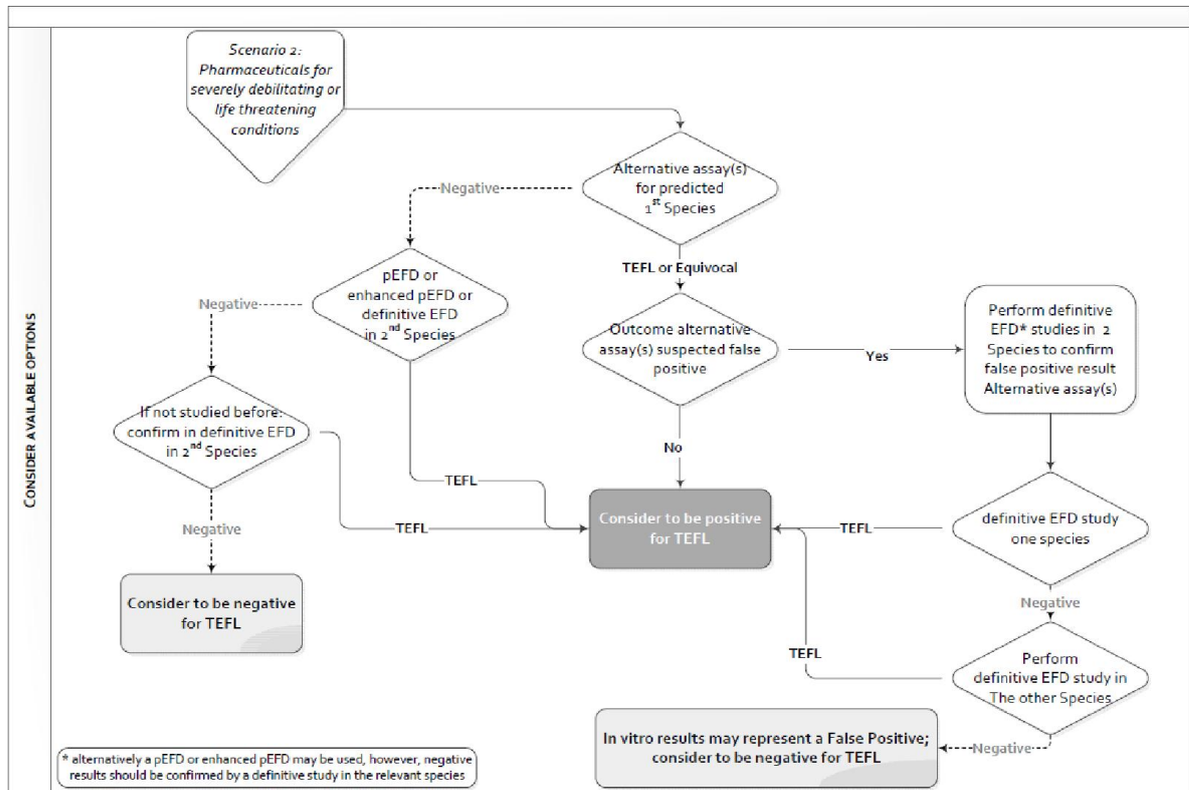
1485 a. If a TEFL outcome is observed in the second species EFD study, the
1486 pharmaceutical will be considered positive in animals. Further EFD studies
1487 would be warranted only if they would significantly alter the risk assessment
1488 (e.g., positive only at high multiples of the clinical exposure and thus another
1489 species could indicate a relevant risk at low exposures).

1490 b. If no TEFL is detected in the second species definitive EFD study, the
1491 pharmaceutical will be considered not likely to induce TEFL in animal studies
1492 (*in vitro* and *in vivo*) and no further EFD studies would be warranted.

1493 For the scenarios above where a rat EFD study is not conducted, an additional opportunity to
1494 confirm *in vitro* positive outcomes is presented in either rat fertility or pre-and postnatal

- 1512 i. If no TEFL is observed in both species *in vivo*, results from the alternative
1513 *in vitro* assay represent a false positive and the pharmaceutical will be
1514 considered negative *in vivo* and this information will be used in the risk
1515 assessment.
- 1516 ii. If one or more of these *in vivo* studies has positive TEFL outcome, the
1517 pharmaceutical will be considered positive *in vivo* and this will be factored
1518 into the risk assessment.
- 1519 2. If the alternative assay predicts a negative outcome (i.e., no TEFL), an EFD study in
1520 the other species (e.g., rabbit) should be conducted to determine if the pharmaceutical
1521 is positive *in vivo*.
- 1522 a. If a TEFL outcome is observed in the second species EFD study, the
1523 pharmaceutical will be considered positive in animals. Further EFD studies
1524 would be warranted only if they would significantly alter the risk assessment
1525 (e.g., positive only at high multiples of the clinical exposure and thus another
1526 species could indicate a relevant risk at low exposures).
- 1527 b. If no TEFL is observed in the second species definitive EFD study, the
1528 pharmaceutical will be considered negative in animals and no further EFD
1529 studies would be warranted.
1530

1531 **Figure 9-3: Scenario 2 Showing the Integrated Testing Strategies for EFD for**
 1532 **Pharmaceuticals for Severely Debilitating or Life Threatening Diseases**



1533
 1534

1535 **11.3.5.2 Scenarios applicable in case there is no or only 1 relevant mammalian**
 1536 **species (crf. Species selection)**

1537 **a) Scenario 3: Non-highly Targeted pharmaceuticals that are pharmacolo-gically active**
 1538 **in only one or no species**

1539 If there is evidence (e.g., mechanism of action, phenotypic data from genetically modified
 1540 animals, class effects) that there will be an adverse effect on pregnancy outcome, these data
 1541 can provide adequate information to communicate risk to reproduction and nonclinical *in*
 1542 *vivo* studies are not warranted. Similar approaches are discussed in other guidelines (ICH
 1543 S6(R1)(2) and ICH S9 (3)).

1544
 1545 If the evidence is lacking, inconclusive or negative for TEFL effects, an EFD study in a
 1546 single species should be conducted. If that study is positive for TEFL, an EFD study in a
 1547 second species is not warranted provided the observations occurred at relevant margins of
 1548 exposure and interpretation is not confounded by maternal toxicity.