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主旨:國際醫藥法規協和會(ICH)S1B(R1)指引草案進入公開諮 詢階段,請協助轉知所屬,彙整該指引有關建議並請於 110年8月31日前惠復,請查照。

說明:

- -、S1B(R1)工作係藥品之齧齒動物致癌性檢測(Addendum to the Guideline on Testing for Carcinogenicity of Pharmaceuticals)指引之修訂,該指引草案現進入法規諮 詢階段公開徵求各界意見,請協助轉知所屬,並惠予彙整 所屬建議,依意見彙整表提供中英文建議惠復。
- 二、SIB(R1)指引草案及意見彙整表請至本署ICH草案公開諮詢 專區下載,路徑:首頁>業務專區>藥品>ICH專區>草案公開 諮詢專區。草案相關資料亦可參考ICH Public Consultations (https://www.ich.org/page/publicconsultations)。
- 正本:台灣藥學會、中華民國學名藥協會、中華民國西藥商業同業公會全國聯合會、中 華民國藥劑生公會全國聯合會、台灣年輕藥師協會、台灣醫藥品法規學會、台灣 研發型生技新藥發展協會、臺灣製藥工業同業公會、台灣藥品行銷暨管理協會、



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第2頁,共2頁



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

ICH HARMONISED GUIDELINE

ADDENDUM TO THE GUIDELINE ON TESTING FOR CARCINOGENICITY OF PHARMACEUTICALS S1B(R1)

Draft version Endorsed on 10 May 2021 Currently under public consultation

At Step 2 of the ICH Process, a consensus draft text or guideline, agreed by the appropriate ICH Expert Working Group, is transmitted by the ICH Assembly to the regulatory authorities of the ICH regions for internal and external consultation, according to national or regional procedures.

S1B(R1) Document History

Code	History	Date
S1B(R1)*	Endorsement by the Members of the ICH Assembly	10 May 2021
	under Step 2 and release for public consultation.	

*This addendum is complementary to the S1 Guidelines (S1A, S1B and S1C(R2)) and is not intended to replace the existing S1B Guideline. At Step 4 of the ICH process, this addendum will be integrated with the S1B Guideline.

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ICH HARMONISED GUIDELINE

ADDENDUM TO THE GUIDELINE ON TESTING FOR CARCINOGENICITY OF PHARMACEUTICALS

ICH S1B(R1)

ICH Consensus Guideline

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1 **PREAMBLE**

- 2 This Addendum is to be used in close conjunction with ICH *S1A Guideline on the Need for*
- 3 Carcinogenicity Studies for Pharmaceuticals, S1B Testing for Carcinogenicity of
- 4 *Pharmaceuticals, and S1C(R2) Dose Selection for Carcinogenicity Studies.* The Addendum is
- 5 complementary to the S1 Guidelines.

6 **1. INTRODUCTION**

7 **1.1 Scope of the Addendum**

8 This Addendum covers all small molecule pharmaceuticals where carcinogenicity evaluations 9 are recommended as described in S1A.

10 **1.2 Purpose of the Addendum**

11 This Addendum expands the testing scheme for assessing human carcinogenic risk of small

- 12 molecule pharmaceuticals by introducing an additional approach that is not described in the
- 13 original S1B Guideline. This is an integrative approach that provides specific weight of
- 14 evidence [WoE] criteria that inform whether or not a 2-year rat study adds value in completing
- 15 a human carcinogenicity risk assessment. The Addendum also adds a plasma exposure ratio-
- based approach for setting the high dose in the rasH2-Tg mouse model,¹ while all other aspects of the recommendations for high dose selection in S1C(R2) Guideline would still apply.
- The first definition of the first dose selection in DTC(R2) Guideline would still appry.
- 18 Application of this integrative approach would reduce the use of animals in accordance with the 2P
- 19 3Rs (reduce/refine/replace) principles, and shift resources to focus onto generating more
- 20 scientific mechanism-based carcinogenicity assessments, while promoting safe and ethical
- 21 development of new small molecule pharmaceuticals.

22 **1.3 Background**

23While the S1B Guideline calls for flexibility in considering approaches to address pharmaceutical carcinogenicity testing, the basic scheme generally recommends a long-term 24rodent study which, in practice, is usually a 2-year study in rats, along with a second rodent 25carcinogenicity study in mice (2-year or short-term study). Since publication of the ICH S1B 26Guideline, scientific advances toward elucidation of mechanisms of tumorigenic action, greater 27understanding of the limitations of rodent models, and several retrospective analyses of 28pharmaceutical datasets indicate that 2-year rat carcinogenicity studies might not add value to 29human carcinogenicity risk assessment in some cases and the carcinogenic potential could have 30

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¹ The rasH2-Tg mouse was developed in the laboratory of Tatsuji Nomura of the Central Institute for Experimental Animals (1). The model is referred to in the S1B Guideline as the TgHras2 transgenic mouse. The official nomenclature for the model is CByB6F1-Tg(HRAS)2Jic which is maintained by intercrossing C57BL/6JJic-Tg(HRAS)2Jic hemizygous male mice with BALB/cByJJic female mice. The littermates derived from these intercrosses are the transgenic rasH2-Tg animals with the tg/wt genotype, and the wild type rasH2-Wt animals with a wt/wt genotype.

Since other short-term models mentioned in S1B have not gained significant use compared to rasH2-Tg over the past 20 years, pharmaceutical development experience with these models is far more limited. Therefore, other short-term carcinogenicity models referred to in S1B would not qualify for a plasma exposure ratio-based high dose selection.

It is appropriate to use wild-type rasH2-Wt littermates of rasH2-Tg mice for dose range-finding studies and for generating exposure data.

- been assessed adequately based on a comprehensive assessment of all available
 pharmacological, biological, and toxicological data (2-9).
- To determine whether the conclusions from these retrospective analyses could be confirmed in a real- world setting (i.e., prior to knowledge of the 2-year rat carcinogenicity study outcomes), an independent international prospective study was conducted under ICH SI(R1) RND *Proposed Change to Rodent Carcinogenicity Testing of Pharmaceuticals – Regulatory Notice Document.* The conclusion from this prospective evaluation confirmed that an integrated WoE approach could be used to adequately assess the human carcinogenic risk for certain pharmaceuticals in lieu of conducting a 2-year rat study.²
- In addition, an exposure ratio endpoint (based on animal to human plasma AUC) for high dose selection in 2-year rodent studies as per ICH S1C(R2) has not been globally accepted for use in the rasH2-Tg mouse study. Therefore, a comprehensive analysis was conducted to assess exposures and outcomes in rasH2-Tg studies from available information.³ As described in Section 3, the results of this analysis indicate that there is no value in exceeding a 50-fold exposure ratio for high dose selection in this model.

462. A WEIGHT OF EVIDENCE APPROACH TO ASSESS THE HUMAN47CARCINOGENIC POTENTIAL OF SMALL MOLECULE PHARMACEUTICALS

48 Over the course of drug development, it is important for sponsors to develop a scientifically 49 robust strategy for carcinogenicity assessment that considers key biologic, pharmacologic, and 50 toxicologic information. The integrative WoE assessment approach described in sections 2.1 51 and 2.2 may support a conclusion that the test compound is either:

- likely to be carcinogenic in humans such that the product would be labeled accordingly
 and any 2-year rat carcinogenicity studies would not add value; or
- likely not to be carcinogenic in humans such that a 2-year rat study would not add value (may also not be carcinogenic in rats, or may likely be carcinogenic in rats but through

² Conduct and results of the prospective study will be summarized; ICH Website of RND and PEP updates will be cited; and future DRA manuscript pointed to. These new citations will appear in the Step 4 Version and this footnote modified.

³ The approach taken for determining an adequate exposure margin for high dose selection for the rasH2-Tg short-term model is similar to that described previously for the 2-year rat and mouse studies (10,11) and Hisada S, Tsubota K, et al (Manuscript in preparation) Survey of Available Data to Assess Tumorigenic Sensitivity of rasH2-Tg Mice and 2-year Rodent Models. Draft Summary: Results were analyzed from studies conducted for 50 drugs in the 6-month rasH2-Tg model and the 2-year rat, 15 of which were also evaluated in the 2-year mouse. For 13 studies concluded to be positive in rasH2-Tg, 6 genotoxic carcinogens were positive within 0.1 - 3-fold of the AUC exposure ratio or body surface area adjusted dose ratio (rodent:human), and 7 nongenotoxic carcinogens were positive all within 1 - 50-fold. Among those 7, three tested positive only at exposures evaluated that exceeded 25-fold. The rasH2-Tg model was 20-fold more sensitive to 10-fold less sensitive than the 2-yr rat or mouse among these 13 drugs that were tested in all 3 models, while 3 of the 13 drugs tested negative in the 2-year rat study. Eight of 37 drugs that tested negative in rasH2-Tg were evaluated at greater than 50-fold exposure ratios (60 to >200fold). For 11 compounds testing positive in 2-year rat studies at exposure ratios of <25-fold, and testing negative in rasH2-Tg, high dose selection in rasH2-Tg was limited by maximum tolerated dose (MTD) at exposure ratios of <50-fold for 9 drugs, and for the other 2 drugs, exposure margins exceeded 50-fold. Human relevance of the tumorigenic potential observed in rats for these 11 drugs has been questioned. In conclusion, when high exposures are tolerated in rasH2-Tg mice, there appears to be some value in exceeding 25-fold, but the overall evidence indicates no benefit to exceeding a 50-fold exposure margin. (Note: this summary paragraph may be deleted upon publication of Hisada et al).

- 56 well recognized mechanisms known to be human irrelevant); or
- uncertain with respect to the carcinogenic potential for humans, and a 2-year rat carcinogenicity study is likely to add value to human risk assessment.
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In cases where the WoE assessment leads to a conclusion of uncertainty regarding human carcinogenicity potential, the approach described in S1B of conducting a 2-year rat carcinogenicity study together with a carcinogenicity assessment in mice (short term or 2-year study) remains the most appropriate strategy.

64 **2.1 Factors to consider for a WoE assessment**

65 A WoE approach is based on a comprehensive assessment of the totality of data relevant to 66 carcinogenic potential available from public sources and from conventional drug development 67 studies. These factors include:

- data that inform carcinogenic potential based on drug target biology and the primary
 pharmacologic mechanism of the parent compound and active major human
 metabolites. This includes drug target distribution in rat and human; available
 information from genetically engineered models; human genetic association studies;
 cancer gene databases; and carcinogenicity information available on the drug class,
- results from secondary pharmacology screens for the parent compound and major
 metabolites that inform off-target potential, especially those that inform carcinogenic
 risk (e.g., binding to nuclear receptors),
 - histopathology data from repeated-dose toxicity studies completed with the test agent, with particular emphasis on the long term rat study, including exposure margin assessments of parent drug and major metabolites,⁴
- 4) evidence for hormonal perturbation, including knowledge of drug target and
 compensatory endocrine response mechanisms; weight, gross and microscopic changes
 in endocrine and reproductive organs from repeated-dose toxicity studies; and results
 from reproductive toxicology studies,⁵
- genetic toxicology study data using criteria from ICH S2(R1) Genotoxicity Testing and
 Data Interpretation for Pharmaceuticals Intended for Human Use; equivocal
 genotoxicity increases uncertainty with respect to the carcinogenic potential,
- 6) evidence of immune modulation in accordance with ICH S8 Immunotoxicity Studies
 for Human Pharmaceuticals; it is generally recognized (12,13) that standard rat and
 mouse carcinogenicity studies are not reliable for identifying this specific human risk.

⁴ Histopathology findings from long term rat toxicity studies of particular interest for identifying carcinogenic potential in a 2-year rat study include cellular hypertrophy, cellular hyperplasia, persistent tissue injury and/or chronic inflammation, foci of cellular alteration, preneoplastic changes, and tumors. It is important to provide an understanding of the likely pathogenesis, and/or address the human relevance of such findings. While long term rat toxicity study data are shown to be of highest value for assessing the likely outcome and value of conducting a 2-year rat study, short term rat studies can sometimes also provide histopathologic conclusions of value.

Data from long term toxicity studies in non-rodents and mice may also be useful for providing additional context on the human relevance of rat study findings (e.g., species-specific mechanistic differences) and whether there is value in conducting a 2-yr rat study.

⁵ If microscopic changes in endocrine and reproductive tissues including atrophy, hypertrophy, hyperplasia are observed, or statistically and biologically significant test article associated endocrine or reproductive organ weight changes are observed this may be considered evidence of functional hormonal perturbation even when changes in hormone levels are not documented. Such findings may be suggestive of potential carcinogenic risk unless investigated for human relevance and demonstrated otherwise.

The above WoE factors may be sufficient to conclude whether or not a 2-year rat study would add value. However, where one or more WoE factors may be inconclusive or indicate a concern for carcinogenicity, the Sponsor can conduct investigations that could inform human

- 92 relevance of the potential risk. Possible approaches may include, but are not limited to:
- 93 1) additional investigational studies, or analyses of specimens collected from prior studies
- 94 (e.g., special histochemical stains, molecular biomarkers, serum hormone levels,
- 95 further characterization of immunomodulation, alternative *in vitro* or *in vivo* test
- 96 systems, data from emerging technologies, etc.), and
- 2) clinical data generated to inform human mechanistic relevance at therapeutic doses and
 exposures (e.g., urine drug concentrations and evidence of crystal formation; targeted
 measurements of clinical plasma hormonal alterations; human imaging data, etc.).
- 100 **2.2 Integration of WoE Factors for Assessing Human Carcinogenic Risk**

An integrated analysis of the WoE factors described above determines whether or not a 101 standard 2-year rat study would contribute to the human carcinogenic risk assessment. While 102all factors will contribute to the integrated analysis, the relative importance of each factor will 103 vary depending on the specific molecule being considered. A summary of key outcomes and 104 examples based on the experience accrued during the ICH S1 RND study (S1(R1) RND 105Proposed Change to Rodent Carcinogenicity Testing of Pharmaceuticals – Regulatory Notice 106 Document), are provided in Appendix 1 demonstrating how the WoE factors could be 107 integrated in determining the need for a 2-year rat study. 108

Experience from the ICH S1 RND study indicates that an established profile of other 109 compound(s) in a drug class contributes substantially to assessing human carcinogenic risk 110 111 associated with modulation of the pharmacologic target. Compounds with novel drug targets (i.e., first-in-class) are, nevertheless, considered eligible for an integrative WoE-based 112approach. For such candidates, a higher evidentiary standard is expected to establish that there 113114is no cause-for-concern in regard to target biology. Appendix 1 provides an example where a 115WoE assessment led to a conclusion that a 2-year rat study would not add value to human carcinogenic risk assessment for a drug inhibiting a novel target. 116

When the WoE assessment concludes that conduct of a 2-year rat study is not warranted, the Sponsor should seek alignment with the Drug Regulatory Agency [DRA] of each region where marketing approval is sought. When a sponsor decides to conduct a 2-year rat study in accordance with ICH S1B, there is no obligation to seek concurrence nor to document their rationale with each DRA.

122 **2.3 Mouse Carcinogenicity Studies**

123 A carcinogenicity study in mice, either 2-year or a short-term transgenic model as specified in

- 124 ICH S1B, remains a recommended component of a carcinogenicity assessment plan, even for
- 125 those compounds where the integrated WoE assessment indicates a 2-year rat study would not
- 126 contribute significant value.⁶ However, in some cases, for example, when the WoE evaluation

⁶ The WoE approach described for the rat is not appropriate for eliminating the mouse as a second rodent carcinogenicity species because: (1) 6-month chronic toxicity studies are not generally conducted with mice so the WoE approach cannot be implemented and no database is available to confirm this approach, (2) the results of carcinogenicity studies in mice will often provide different outcomes from the corresponding rat carcinogenicity study, so a direct extrapolation cannot be made, and (3) a 6-month rasH2-Tg mouse has been adopted as an acceptable carcinogenicity study model.

When the WoE evaluation indicates the 2-year rat study adds no value, a carcinogenicity study in mice (either 2-year or short-term) is also not recommended in the EU.

strongly indicates no carcinogenic risk to humans and data indicate that only subtherapeutic,
pharmacologically inactive drug exposures can be achieved in the mouse, it may not be
appropriate to conduct any mouse carcinogenicity study.

130 3. CLARIFICATION ON CRITERIA FOR SELECTION OF THE HIGH DOSE FOR RASH2-TG MOUSE CARCINOGENICITY STUDIES

132In practice, a plasma exposure (AUC) ratio for high dose selection in the absence of dose limiting toxicity or appropriate use of other dose setting criteria as outlined in ICH S1C(R2) in 133this model, has not been globally accepted as an endpoint. Therefore, available data from 134experience with 50 compounds evaluated in the rasH2-Tg mouse model were analyzed and the 135conclusion reached that there was no value in exceeding a 50-fold plasma AUC exposure ratio 136(rodent:human) to support carcinogenicity assessment. Therefore, all criteria for selection of 137the high dose for carcinogenicity studies as specified in S1C(R2) for 2-year rodent studies are 138applicable to rasH2-Tg, including an AUC plasma exposure ratio, except that the exposure 139ratio will be 50-fold in rasH2-Tg rather than 25-fold as for 2-year studies conducted in wild 140 type rodents. All other aspects of S1C(R2) remain applicable to rasH2-Tg. 141

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182 APPENDIX 1: CASE STUDIES APPLYING THE WEIGHT OF EVIDENCE 183 APPROACH

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185 **Preamble**

One outcome of the ICH S1 RND study was the recognition that programs with the following
WoE attributes are more likely to support a conclusion that the results of a 2-year rat study
would not contribute value to human carcinogenicity risk assessment.

- Target biology is well characterized and not associated with cellular pathways known to be involved with human cancer development. Often, the pharmaceutical target was non-mammalian and carcinogenicity data were available with the pharmacologic drug class.
- Results from chronic toxicity studies indicate no hyperplastic, hypertrophic, atypical cellular alterations, or degenerative/regenerative changes noted without adequate explanation of pathogenesis or human relevance, indicative of no on- or off-target potential of carcinogenic concern;
- No perturbation of endocrine and reproductive organs observed, or endocrine findings adequately explained with respect to potential human relevance;
- No identified concerns from secondary pharmacology screens intended to inform offtarget potential for the pharmaceutical
 - No evidence of immune modulation or immunotoxicity based on target biology and repeat dose toxicology studies
 - The overall assessment of genotoxic potential is concluded to be negative based on criteria from ICH S2(R1) Guidance.

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Although rasH2-Tg mouse study results were recommended when available as a WoE element in the initial RND, they did not significantly contribute to the prediction of the 2-year rat carcinogenicity study outcome. Therefore, a rasH2-Tg mouse study is not expected to be completed to support a WoE assessment. However, if rasH2-Tg mouse study results are available, they should be discussed in the assessment.

212

A series of case studies are provided to illustrate the application of the WoE approach. These 213cases are provided for illustrative purposes only and are not intended as guidance to indicate 214215the sufficiency of data to support a WoE assessment. Cases 1 and 2 describe the key WoE factors for that pharmaceutical and how the data were integrated to conclude that a 2-year rat 216study would not add value to the assessment of carcinogenic risk. In contrast to these cases, 217218Case 3 describes how data from the WoE factors were integrated to conclude that the carcinogenic potential for humans was uncertain, and a 2-year rat carcinogenicity study was 219likely to add value to human risk assessment. Case 4 describes a molecule for which a 2-year 220rat carcinogenicity study was concluded to not contribute value to human carcinogenicity 221assessment despite there being no data available for other molecules within the pharmacologic 222223class.

224

225 Case 1: A small molecule inhibitor against a non-mammalian target 226

Prospective WoE Assessment: Concluded by all DRAs and Sponsor as likely not to be carcinogenic in both rats or humans such that a 2-year rat study would not add value

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230 Rationale

231 The WoE analysis supports the conclusion that the molecule was sufficiently studied at high

232	exposure margins, and cause-for-concern was not identified for any of the WoE factors.
233	
234	2-year Rat Study Results: No test article related neoplastic findings were present in the 2-year rat
235	study.
236	
237	WoE Criteria
238	
239	Knowledge of intended drug target and pathway pharmacology relative to carcinogenesis
240	Non-mammalian target excludes intentional alteration of potential mammalian
241	carcinogenic pathways.
242	• No evidence of carcinogenic outcome in 2-year rat studies conducted with other
243	compounds with the same non-mammalian pharmacological target
244	
245	Secondary Pharmacology Screen
246	 No evidence of off-target interactions at drug concentrations up to 10 μM, including
247	no interaction with estrogen, androgen, glucocorticoid receptors
248	
249	General Toxicology from Chronic Rat Study
250	• Chronic (6-month) toxicology study in Wistar rats dosed to saturation of absorption,
251	achieving up to a 31-fold margin to human exposure.
252	 No evidence of human specific major metabolites.
253	• No treatment-related histopathologic findings observed in standard battery of tissues
254	
255	General Toxicology from Chronic Non-rodent Study
256	• Chronic administration (9-month) to non-human primates identified bile duct
257	hyperplasia and hepatocellular hypertrophy, with reactive neutrophils and
258	regenerative hyperplasia. A No-Adverse-Effect-Level was identified which provided
259	a 5-fold margin to human exposure.
260	• Further evaluation in rats would not provide useful information, as similar findings
261	were not observed in the chronic rat study.
262	
263	Hormonal Perturbation
264	• No treatment-related findings on reproductive organ weights or histopathology
265	
266	Genetic Toxicology
267	• No evidence of genotoxic potential based on criteria from ICH S2(R1) Guidance
268	
269	Immune Toxicology
270	• No treatment-related changes in clinical pathology or histopathology of immune
271	tissues (e.g., lymphoid organs, spleen, thymus, bone marrow)
272	
273	Additional Special Investigations
274	• No data available
275	
276	
277	Case 2: A small molecule antagonist of a neuronal G-protein coupled receptor
278	· · · · · · · · · · · · · · · · · · ·
279	Prospective WoE Assessment: Unanimously concluded as likely to be carcinogenic in rats but not
280	in humans through well recognized mechanisms known to be human irrelevant, such that a 2-
281	year rat study would not add value

282

283 **Rationale**

The WoE analysis indicates the potential for rodent-specific liver and thyroid neoplasms based on the toxicology observed in the chronic rat study and on tumor outcome with the pharmacological class. Induction of hepatic cytochrome P450 was demonstrated. Evidence of hormonal perturbation is understood from target pharmacology, did not result in changes in reproductive organ weight or histopathology, and occurred at high multiples to human exposure.

2-year Rat Study Results: The 2-year rat study demonstrated hepatocellular hypertrophy but no neoplastic findings.

293294 WoE Criteria

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296 Knowledge of intended drug target and pathway pharmacology relative to carcinogenesis

- Predominate receptor expression in brain with lower expression in some peripheral tissues, similar across species
- Receptor activation increases ACTH release from pituitary secondary to hypothalamic
 production of adrenocorticotropin-releasing hormone.
- Hypothalamic receptor ligand levels associated with LH surge and gonadotropin
 release in rats.
- Target knock-out mice showed no findings related to carcinogenicity.
- Long-term studies with other compound with same pharmacological target associated
 with thyroid follicular cell adenoma/carcinoma in rats, consistent with elevated
 thyroid stimulating hormone following off-target cytochrome P450 induction.
- Antagonist binding interaction identified for one off-target receptor with Ki 8-fold
 higher than Cmax at maximum clinical dose. Known target pharmacology of off target receptor not associated with tumorigenesis.
- 311 General Toxicology from Chronic Rat Study
 - Increased liver hypertrophy and organ weight at 50x to 74x margin to human exposure.
 - Increased thyroid follicular hypertrophy at 170x to 670x margin to human exposure.
 - No evidence of human specific metabolites.
 - An active major human metabolite in humans was also present in rats
- 318 General Toxicology from Chronic Non-rodent Study
- Increased liver hypertrophy and organ weight at ~230-fold human exposure.
- 321 Hormonal Perturbation
 - Reduced adrenal weight without histopathological correlates and reduced ACTH level at >74x human exposure in the chronic rat study, consistent with inhibition of drug target. Response noted to be growth suppressive.
- Irregular estrous cycles and decreased pregnancy rate were observed at 60-fold human exposure, and decreased numbers of corpora lutea, implantations, and live embryos were observed at >500-fold human exposure in a fertility study in rats. Considered consistent with inhibition of drug target.
- No treatment-related changes observed in reproductive organ weight or
 histopathology in chronic rat study.
- 331

332 Genetic Toxicology

No evidence of genotoxic potential of parent or major human metabolite based on
 criteria from ICH S2(R1) Guidance

336 Immune Toxicology

No treatment-related changes in clinical pathology, lymphocyte subsets, or
 histopathology of immune tissues (e.g., lymphoid organs, spleen, thymus, bone
 marrow)

340341 Additional Special Investigations

- Increased induction of CYP1A2 and CYP3A1 demonstrated
- Bone and teeth fluorosis related to defluorination of compound, demonstrated not to occur in humans
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347Case 3: A first-in-class small molecule inhibitor of a ubiquitously expressed348serine/threonine kinase

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Prospective WoE Assessment: Unanimously concluded to be uncertain with respect to the
 carcinogenic potential for humans, and a 2-year rat carcinogenicity study is likely to add value to
 human carcinogenicity assessment

354 Rationale

Significant carcinogenic uncertainty is based on a complex target pharmacology, the lack of precedent with the drug target, and histopathological changes of concern with inadequate mechanistic explanation from the chronic rat study which are supported by similar findings in cynomolgus monkeys. The immune toxicology observed in monkey will contribute to the overall assessment of risk but is not expected to be further informed by a rat carcinogenicity study.

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362 2-year Rat Study Results: The 2-year rat study demonstrated an increased incidence, lethality, 363 and reduced latency of pituitary tumors in both sexes. This carcinogenic outcome in rats would 364 contribute to the overall assessment of human carcinogenic potential.

365366 WoE Criteria

368 Knowledge of intended drug target and pathway pharmacology relative to carcinogenesis

- Target activation by inflammation-related oxidative stress promotes cellular apoptosis
 and is linked to control of cell proliferation; target inhibition suppresses apoptotic
 signaling and impacts cell proliferation, theoretically promoting cancer growth.
- Drug target displays tissue-dependent roles in cancer development, both promotion and suppression, in animal models.
 - No data available on tumor outcome from target inhibition in long term rodent or short term transgenic mouse studies
- 376377 General Toxicology from Chronic Rat Study
- Increased incidence and severity of renal basophilic tubules, eosinophilic droplets,
 and brown pigment in renal cortex starting at 14-fold human exposure. Etiology of
 lesions not empirically addressed.

381	Chronic irritation of limiting ridge in non-glandular stomach at 39-fold human
382	exposure. Etiology of lesions not empirically addressed.
383	 Increased liver weight without microscopic correlates.
384	• No evidence of human specific metabolites.
385	• An inactive major human metabolite in humans was also present in rats
386	Concerned Transiende our from Channie New and ent Study
387	General Toxicology from Chronic Non-rodent <u>Study</u>
388	• In monkeys, gastrointestinal epithelial degeneration, necrosis, reactive hyperplasia,
389	ectasia, inflammation, and ulceration, at doses ~12-fold human exposure
390	• Increased incidence of renal tubule degeneration /regeneration, necrosis, dilation, and
391	vacuolation at ~12-fold human exposure
392	
393	Hormonal Perturbation
394	• Increased adrenal weight and cortical hypertrophy in rats at 17-fold human exposure.
395	Etiology not empirically addressed.
396	
397	Immune Toxicology
398	• In monkeys, suppression of TDAR with no effect on NK cytotoxicity or granulocyte
399	function, and decreased lymphoid cellularity in spleen, thymus, lymph nodes at 12-
400	fold human exposure.
401	
402	Genetic Toxicology
403	• No evidence of genotoxic potential of parent or major human metabolite based on
404	criteria from ICH S2(R1) Guidance
405	
406	Additional Special investigations
407	• Increases in hepatic enzymes CYPs 1A, 3A, and 2B demonstrated.
408	F
409	
410	Case 4: A first-in-class small molecule inhibitor of a prostaglandin receptor
411	
412	Prospective WoE Assessment: Unanimously concluded as likely not to be carcinogenic in both rats
413	or humans such that a 2-year rat study would not add value
414	Detionale
415	Rationale When compared with the test egent discussed in Case 2, which is also first in along the drug
416	When compared with the test agent discussed in Case 3, which is also first-in-class, the drug
417	target in Case 4 is not associated with a role in cancer development, histopathological findings
418	were not observed in the chronic rat study, and a large margin of exposure was calculated at the high dage $(>50x)$. The accordance along across along indicated the test accord
419	the high dose (>50x). The secondary pharmacology screen also indicated the test agent
420	demonstrates target selectivity.
421	2 way Dat Study Daryltz, The 2 way not apprint a privity study did not domenstrate a dara valated
$\begin{array}{c} 422 \\ 423 \end{array}$	2-year Rat Study Results: The 2-year rat carcinogenicity study did not demonstrate a dose-related increase in tumors.
423 424	increase in tumors.
$\frac{424}{425}$	WoE Criteria
$425 \\ 426$	
420 427	Knowledge of intended drug target biology and pharmacologic mechanism relative to
428	carcinogenesis

429	• Receptor activation associated with allergic inflammatory response and currently
$420 \\ 430$	available data do not suggest a role in tumor initiation or progression.
$430 \\ 431$	 Knock-out mice of drug target showed no histological abnormalities or effects on
$431 \\ 432$	• Knock-out line of drug target showed no instological abiomanties of effects of immune function during one year of observation.
433	• No data available on tumor outcome in 2-year rat studies conducted with other
434	compounds with the same pharmacological target.
435	• No data available from a rasH2-Tg carcinogenicity study conducted with the test
436	agent.
437	
438	Secondary pharmacology screen
439	• Test agent was at least 300-fold more selective for drug target when compared with
440	other receptors in the same class as well as a sub-set of other assessed receptors
441	involved in the inflammatory response.
442	• Test agent was at least 2000-fold more selective for the drug target in a secondary
443	pharmacology screen of various receptors, ion channels, transporters and enzymes.
444	
445	General Toxicology from Chronic Rat Study
446	• Histopathological assessments conducted as part of repeated-dose toxicity studies up
447	to 26-weeks indicated no proliferative changes in any organ or tissue at the highest
448	dose tested (~ 54-fold human exposure based on AUC).
449	• No evidence of human specific metabolites.
450	
451	General Toxicology from Chronic Non-rodent Study
452	• Histopathological assessments conducted as part of repeated-dose toxicity studies up
453	to 39-weeks indicated no proliferative changes in any organ or tissue at the highest
454	dose tested (~ 45-fold human exposure based on AUC).
455	
456	Hormonal Perturbation
457	• No treatment-related findings on reproductive organ weights or histopathology.
458	
459	Genetic Toxicology
460	• No evidence of genotoxic potential based on criteria from ICH S2(R1) Guidance.
461	
462	mmune Toxicology
463	• In the 26-week rat toxicity study, there were no effects on immune function (including
464	the TDAR assay evaluating primary and secondary antibody responses) or adverse
465	effects on lymphocyte subsets at the highest dose tested (~54-fold human exposure
466	based on AUC).
467	
468	Additional Special Investigations
469	• Not performed.