#### ICH HARMONISED GUIDELINE

# NONCLINICAL SAFETY TESTING IN SUPPORT OF

## **DEVELOPMENT OF PAEDIATRIC MEDICINES**

## ICH S11 指引之意見彙整表

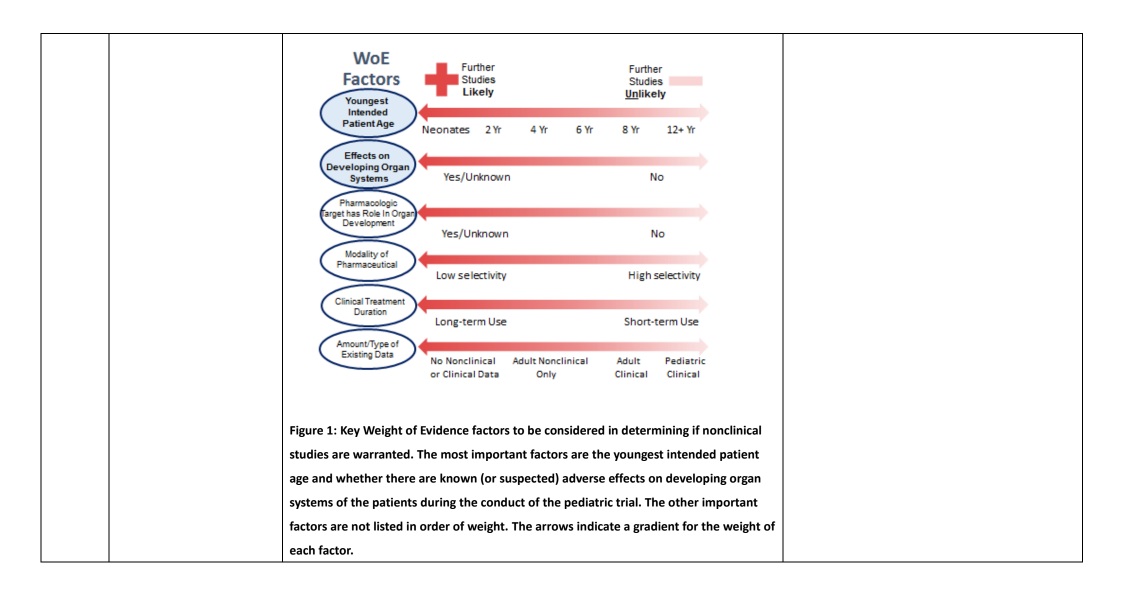
段落	標題	內文	相關建議及意見
			(請提供中英文內容)
22-29		1.1. Objectives of the Guideline	
		The purpose of this document is to recommend international standards	
		for, and promote harmonization of, the nonclinical safety studies	
		recommended to support the development of pediatric medicines.	
		Harmonization of the guidance for nonclinical safety studies will define the	
		current recommendations and reduce the likelihood that substantial	
		differences will exist among regions. It should facilitate the timely conduct	
		of pediatric clinical trials and reduce the use of animals in accordance with	
		the 3Rs (replace/reduce/refine) principles.	
30-38	<b>1. INTRODUCTION</b>	1.2. Background	
		Several regional guidelines have previously been issued by various	
		regulatory agencies and were not in complete agreement on the need for,	
		timing of, and design of juvenile animal studies (JAS).	
		There are ICH guidelines that refer to the need for and/or timing or study	
		design of JAS (e.g., ICH E11, M3, S5, S6, and S9); the current guideline is	
		intended to complement the existing 36 ICH guidelines. This guideline	
		reflects current thinking based on collations of examples by regulatory	
		agencies, by industry surveys, and literature.	

		1
39-48	1.3. Scope	
	This guideline recommends an approach for the nonclinical safety	
	evaluation of medicines intended for development in pediatric	
	populations. This can include products with prior adult use, as well as	
	products being considered for initial human use in pediatrics (see Section	
	4).	
	The ICH S9 guideline should be consulted for recommendations on	
	whether to conduct JAS for those pharmaceuticals included in the scope of	
	the ICH S9 guideline, i.e., anticancer pharmaceuticals. The ICH S11	
	guideline should be consulted for study design in all cases where a study is	
	considered to be warranted.	
	Tissue engineered products, gene and cellular therapies, and vaccines are	
	excluded from the 47 scope of this guideline.	
49-73	1.4. General Principles	
	Pediatric patients represent a population different from adults when	
	considering the rapid growth and postnatal development of several organ	
	systems. The continued development of these systems can affect drug	
	pharmacokinetics (PK), pharmacodynamics (PD), and/or off-target effects	
	of medicines, potentially leading to differences in toxicity and/or efficacy	
	profiles both between pediatric age groups and when compared to adults.	
	An early consideration of nonclinical support for pediatric medicine	
	development is recommended. In this respect, changing the design and/or	
	timing of the traditional nonclinical program is one way to address	

potential safety concerns for the pediatric patient. For example, dosing can	
be initiated at a younger age in a repeat-dose toxicity study to support the	
corresponding developmental stages in pediatric patients. Another	
approach could be to conduct the Pre- and Postnatal Development (PPND)	
study earlier than usual, with modifications that demonstrate adequate	
offspring exposure and incorporate additional endpoints (see ICH S5).	
These changes can obviate the need for, or limit the design of, a dedicated	
JAS.	
An understanding of the overall clinical development plan is needed to	
design an appropriate, efficient nonclinical plan. Prior to each pediatric	
trial, a weight of evidence (WoE; see Section 2) based decision should be	
made to determine whether additional nonclinical investigations are	
warranted. The outcome of such a WoE assessment can be different for	
each trial for the same pharmaceutical depending on pediatric age and	
indication.	
The conduct of additional nonclinical investigations should be undertaken	
only when previous animal and human data are judged to be insufficient	
to support pediatric studies. JAS are designed to address identified safety	
concerns that cannot be adequately addressed in other nonclinical studies	
or pediatric clinical trials, including potential long-term safety effects. This	
guideline recommends a customized JAS that comprises core design	
elements and potential additional elements driven by specific concerns.	

74-89		2.1 Clinical Context	
		The pediatric clinical development plan for a pharmaceutical is discussed	
		in the ICH E11 guideline, and needs to be understood before an	
		appropriate nonclinical plan can be designed. The pediatric clinical plan	
		includes the indication/condition, the intended pediatric age group(s), and	
		the treatment regimen (particularly, the duration of dosing during the	
		stages of development). The clinical development of a medicine for pediatric	
		patients usually follows initial adult clinical studies. If needed, the design and	
		timing of additional nonclinical investigations are dependent on the identified	
		safety concerns and the intended clinical use.	
	2. DETERMINING THE	In case of a severely debilitating or life-threatening disease, or one in	
	NEED FOR ADDITIONAL	which there is serious unmet medical need in a pediatric population, the	
	NONCLINICAL SAFETY	sponsor and regulatory agencies should consider the 85 timing impact of	
	INVESTIGATIONS	producing additional data to support patient access to a pharmaceutical.	
		This decision should be based upon a careful and cautious risk-benefit	
		evaluation. If a safety concern is identified for further clinical	
		development, appropriate nonclinical studies (e.g., JAS) should 88 be	
		considered, and could be conducted in parallel with clinical investigation.	
90-115		2.2 Weight of Evidence Approach	
		The nonclinical development plan for a pediatric pharmaceutical depends	
		on an integrated assessment based on the totality of the clinical context	
		together with the pharmacology, pharmacokinetic (ADME), and nonclinical	
		in vitro and in vivo animal and clinical safety data, i.e., a WoE approach. A	
		WoE approach considers multiple factors evaluated together and,	
		therefore, a single factor should not be considered in isolation. The	

importance of each factor should be weighted such that the final decision	
concludes whether available data adequately address safety concerns in	
the proposed pediatric population or whether additional nonclinical	
studies are warranted.	
The WoE evaluation should be conducted when designing the initial	
pediatric development plan, but revisited if there are changes in age	
ranges and/or indications. The WoE outcome can be different for each trial	
depending on the pediatric population and the disease to be treated.	
Figure 1 below shows some key factors that should be considered as part	
of the WoE evaluation to determine the need for further nonclinical	
investigations. The individual factors are presented below on the left of	
the figure. The most important factors are the youngest intended patient	
age and whether there are known (or suspected) adverse effects on	
developing organ systems of the patients during the conduct of the	
pediatric trial. The other important factors are not listed in order of weight	
in the figure. The list is not all inclusive for every situation, as there may be	
additional specific factors to consider (e.g., clinical management). The WoE	
factors are further described in the following sections.	



116-206	2.3 Factors to Inform the Weight of Evidence Evaluation
	2.3.1 Clinical Information
	The most relevant safety and efficacy data for pediatric patients come
	from other pediatric subpopulations and adults exposed to the
	pharmaceutical. This established efficacy and safety profile is usually the
	first point to consider when determining if additional nonclinical studies
	are warranted.
	The youngest intended patient age is one of the most important factors to
	be considered. The use of existing clinical data from older subgroups may
	not necessarily be sufficient (see ICH E11). Further nonclinical studies are
	more likely to be warranted at the lower end of the age range.
	The duration of clinical treatment is another factor in determining whether
	additional nonclinical studies are warranted. Longer durations of
	treatment are more likely to expose a pediatric subject during a
	developmentally sensitive window. Whereas short-term use of a
	pharmaceutical is less likely to affect some aspects of development such as
	growth, a long duration of use is more likely to warrant further nonclinical
	studies than short-term treatments.
	Additional nonclinical studies are not warranted when existing clinical data
	are considered sufficient to support pediatric use and/or if identified
	safety concerns can be clinically managed. A JAS is not warranted to
	confirm toxicity in target organs in which sensitivity to toxicity is not
	expected to differ between adults and pediatric patients. Developmental

differences in target or off-target tissue maturity do not, in isolation, necessarily mean a JAS is required, but are a concern that needs to be considered.

#### 2.3.2 Pharmacological Properties

Primary or secondary pharmacological properties of a pharmaceutical can be responsible for unwanted side effects. This may raise concerns for pediatric use if effects occur in systems/organs in development and/or if developing organs have a different sensitivity from mature organs. A review of the literature on the developmental expression and ontogeny of drug target(s) (e.g., receptor, enzyme, ion channels, protein), or the known/potential role of the target during development is recommended. Existing data from genetically modified animals (e.g., the knock-out of a receptor) may also identify developmental effects of potential concern for the pediatric population, which could be included in the WoE evaluation.

If the known pharmacology of a medicine has the potential to impact the development of the intended pediatric population, or the role of the pharmacology on development is not understood or reasonably predictable, further nonclinical investigations should be considered. Potential adverse effects of pharmaceuticals with high selectivity for their target (e.g., monoclonal antibodies) are more likely to be related to exaggerated pharmacology and therefore be more predictable than effects of pharmaceuticals with lower selectivity for their pharmacologic target. Pharmaceuticals with lower selectivity may have secondary pharmacodynamics effects and thus are more likely to warrant further

nonclinical investigations. Considerations should be given whether conducting in vitro or ex vivo investigations using juvenile (i.e., animal) or pediatric (i.e., human) tissues would be useful to determine potential age-related differences in sensitivity, density, and distribution of molecular pharmacological/toxicological targets.

Further nonclinical studies might not add value when the underlying pharmacology has already 157 identified a particular hazard.

#### 2.3.3 Pharmacokinetic Data

Important differences can exist in the ADME of pharmaceuticals depending on the age of both 160 patients and animals, leading to potential differences in efficacy and toxicity. These differences are usually most prominent in neonates and infants. Similarly, maturation of the gastrointestinal (GI), liver, and renal systems can result in rapidly changing systemic exposures, particularly in young animals.

The use of clinical PK modelling and simulation systems for the purpose of predicting PK/ADME characteristics in pediatric populations can be more relevant than conducting JAS. If the results of the PK modelling and simulation indicate that there will be significant differences between adult and pediatric patients, then nonclinical investigations (e.g., in vitro studies) can be helpful to determine the potential impact of these differences on toxicity.

### 2.3.4 Nonclinical Safety Data

Existing nonclinical toxicity study data should be evaluated for signals that could indicate potential effects in organs undergoing development in pediatric subjects. Findings occurring in animals at similar exposures as those likely to be achieved in pediatric subjects are of higher concern, particularly if the findings occur in organs/tissues that undergo significant postnatal development at the intended pediatric age (see Appendix A). Safety signals that occur in adult animals of more than one species are of increased concern. Depending on the age of the animals at study start and the endpoints included, some of these concerns may have been addressed in existing repeat-dose toxicity studies.

Genotoxicity testing and safety pharmacology investigations are normally conducted to support adult clinical trials and, therefore, should be available before pediatric clinical trials commence. If a safety pharmacology study shows an effect in an organ system known to be developing in the intended pediatric patient population, the possible impact of the effect should be carefully considered. Additional genotoxic and safety pharmacology assessments in juvenile animals are generally not needed to support pediatric indications.

Reproductive and developmental toxicity study data may also be available. If PPND study data are available and have shown clinically relevant and sustained systemic exposures in offspring during the relevant postnatal period, these data can contribute to the WoE evaluation. The review of such data should include the maternal tolerance of the drug during pregnancy and lactation, as this could have impacted on the findings in the offspring. Observations of adverse effects in offspring would not, on their own, indicate that a JAS is recommended. However, if there is an identified safety concern that could lead to effects on postnatal development, it should be considered in the WoE evaluation. These data in rodents are primarily relevant to preterm and term neonates if exposure is demonstrated.

In some cases modification of a rodent PPND study can obviate the need for a JAS, provided potential concerns for the pediatric population have been appropriately addressed in the study design (see ICH S5). For enhanced PPNDs (ePPND) studies conducted in the non-human primate (NHP), the data from the offspring can characterize toxicity during early postnatal development, provided relevant exposure and/or PD effects are confirmed in the offspring. When available, ePPND data should be evaluated in combination with data from the general toxicity studies in assessing the value of additional nonclinical investigations.

#### 2.3.5 Feasibility

The decision to conduct an additional animal study should also consider the technical and practical feasibility of the study design and endpoints (see Section 3). If a study in animals cannot be conducted with dose levels that provide acceptable systemic exposures in the range of those expected in pediatric patients, even with an alternative route of administration or frequency of dosing, the conduct the JAS may not be informative or warranted.

207-214		2.4 Application and Outcome of the Weight of Evidence Evaluation	
		All of the WoE factors described above should be considered when	
		determining whether additional nonclinical investigations are warranted.	
		Additional nonclinical studies are not warranted if identified safety	
		concerns can be clinically monitored and/or managed. When a study is	
		warranted, the specifics of the identified safety concerns will define the	
		objectives of the nonclinical investigation; this could be a JAS or another	
		study (e.g., in vitro or ex vivo investigations).	
		Examples of applying the WoE approach are in Appendix B.	
215-235		3.1 General Considerations/Study Objectives	
		Once it is decided that a JAS is warranted, Section 3 should be consulted to	
		design the appropriate study. This section contains recommendations on	
		study design considerations, core endpoints to be included in all studies,	
		and additional endpoints that can be included to address specific	
		concerns. A JAS design including all potential additional endpoints is not	
		recommended without rationale.	
	3. DESIGN OF		
	NONCLINICAL JUVENILE	If the reason to conduct a study is primarily driven by a specific, identified	
	ANIMAL STUDIES	safety concern for pediatric patients, the study design should be	
		customized to address particular aspects of function or development of a	
		target organ or system of concern. If the rationale to conduct a study is	
		based on a concern for patient safety due to lack of relevant knowledge of	
		the pharmacology, the study design would generally be broader and	
		include additional endpoints as appropriate.	
L		1	

The maturation of human and animal organ systems can influence
susceptibility to toxicity. Understanding the relative level of maturity and
function across species during development is needed not only to design
the appropriate JAS but also to aid the translation of nonclinical toxicity
findings to a specific human age range. This "age" or "stage" mapping can
be challenging and is not uniform across different organ systems or
species, as the relative maturity at birth, rate of postnatal maturation,
and/or regulation of maturation can be quite different between humans
and animals. While not comprehensive, Appendix A, Figures A1-A6 provide
an overview of age-dependent development of organ systems by species.
3.2 Preliminary/ Dose Range Finding Studies
Preliminary studies such as dose range-finding (DRF) or PK studies with
small group sizes of juvenile animals of relevant age are highly
recommended to perform tolerability and PK/TK (toxicokinetic)
assessments. This is particularly valuable when dosing starts prior to
weaning to avoid unexpected mortality, excessive toxicity, and/or
irrelevant exposures in a definitive JAS.
Dosing should be initiated at the youngest planned starting age of the
animals in the definitive JAS to evaluate the most critical period for
tolerability and exposure differences. The DRF dosing period generally
lasts a few weeks, e.g., typically until shortly after weaning in rodents. If
there are important age-related differences in tolerated dose levels
between adults and juveniles, a second DRF study may be needed to select
adequate dose levels or a dosing regimen for the definitive JAS. See
sections on route of administration (3.6) and dose selection (3.7) for more

information on the use of preliminary studies to prepare for anticipated
changes in dosing route and/or dose level adaptation during the course of
a definitive JAS.
In a preliminary or DRF JAS, lack of tolerability of a pharmaceutical at
clinically relevant systemic exposures can indicate a significant concern for
the corresponding clinical age range. When the reason for greater
sensitivity or significant differences in toxicity profiles between juvenile
and adult animals at similar systemic exposure is not understood,
additional investigations (e.g., assessment of protein-binding values or
blood-brain barrier penetration) can be useful for the interpretation of
these differences.
In certain circumstances, DRF studies can explore the usefulness of
particular endpoints, tissues, or biomarkers and thus refine the study
design of the definitive JAS.
3.3 Animal Test System Selection
When a JAS is warranted, in most cases a single species is considered
sufficient. In principle, the rat should initially be considered as the species
for a JAS. Other species have been used in JAS (e.g., mouse, rabbit, dog,
minipig, NHP). In all cases, the selected species should be justified, as
nonclinical studies in a non-relevant species can give rise to
misinterpretation and are not recommended.
The following factors should be considered when selecting an appropriate
species:

An understanding of the ontogeny of the pharmacological or
toxicological target (e.g., the receptor) in animals in comparison to
that in the intended pediatric population
Preference for a species and strain for which adult repeated-dose
toxicity data are available to allow a comparison of the toxicity and
systemic exposure profiles between juvenile and adult animals.
Toxicological target organs
<ul> <li>the relative stage of organ/system development in the juvenile</li> </ul>
animal as compared to the intended pediatric population (see
also Section 3.4)
the ability of the animal model to detect toxicity endpoints of
concern
The technical/practical feasibility to conduct the study in the selected
species
Similarity of ADME characteristics
Advantages and disadvantages of using different rodent or non-rodent
species are outlined in Appendix A, Table A1.
While for biopharmaceuticals NHPs are pharmacological responders in
many cases, the conduct of JAS in NHPs is challenging for both scientific
and practical reasons. There is limited added value of performing JAS in
younger NHP as compared to the 2-4 year old NHP used in general toxicity
studies and, therefore, alternative approaches to obtaining the necessary
data are encouraged. Only in rare cases is the value of JAS conducted in
NHP justifiable.

	Consistent with ICH S6, a homologous protein, when available, can be	
	considered for the purposes of hazard identification in the rodent or other	
	non-rodent species.	
	JAS in two species would be warranted only in a pediatric-first situation	
	(see Section 4) or where there are multiple specific concerns for postnatal	
	development and one species alone is not able to address them.	
	If a pediatric PD model of disease exists (e.g., enzyme replacement	
	therapy), appropriate safety endpoints can be incorporated in these	
	studies. This information could contribute to the WoE evaluation and/or	
	potentially obviate the need for a dedicated JAS.	
290-336	3.4 Age of Animals, Dosing Period, and Dosing Regimen	
	The age of dosing initiation in animals should developmentally correspond	
	to the youngest age of the intended pediatric population, which will	
	depend on a human-to-animal comparison of developmental periods of	
	organ system(s) of toxicological concern. As comparative organ system	
	correlations are not aligned for each organ across species, priority should	
	be given to any target organ/ system of potential concern or to particularly	
	vulnerable developing systems in the intended patient population. The	
	animal age at dosing initiation should be justified using relevant	
	information (see Appendix A).	
	When determining the duration of administration in JAS, it is important to	
	consider the age range and the shorter developmental period of animals	
	compared to humans, the duration of treatment for the intended pediatric	

population, the safety concern to be assessed, and the developmental
stage of target organs/functions of the intended pediatric population
relative to that of the animals used for toxicology studies.
The dosing period in JAS is not only defined by the pediatric age stages
(e.g., > 2 years) or the clinical dosing duration but also by the specific
stages of organ development for the organs of concern (see Appendix A).
To evaluate the impact on a pediatric developmental stage, a longer dosing
period in animals can be appropriate to address a concern of a certain
organ system that develops late (e.g., central nervous system [CNS])
compared to a system with shorter developmental window (e.g., kidney).
In contrast to nonclinical studies for adult populations (see ICH M3), a
short treatment duration in pediatric patients can require a longer dosing
duration in the JAS to capture the developmental age range of the
intended pediatric population. For example, to include the youngest
intended patients of 2 years old up to patients 12 years of age with a
clinical dosing duration of 14 days, the JAS can have a dosing period longer
than 14 days to incorporate exposure at all developmental stages
corresponding to human patients from 2 to 12 years old (e.g., in the rat
this would be approximately 6 weeks dosing duration, roughly postnatal
day (PND) 21 to 65, See Appendix A).
Dosing up to maturation can be feasible in non-rodent species like the dog,
minipig, and rabbit, as these species mature over a period of a few to
several months, and with relative consistency. In contrast, the interval
between birth and maturity for NHPs is several years, making dosing

during the entire developmental period not practical. Eurthermore, NHDc	
and maturity.	
When a DRE study demonstrates that a dose level or duration of dosing is	
approach may only be needed at the dose that is not tolerated. This	
approach is especially applicable in cases when the clinical dosing period is	
comparable to or shorter than the dosing period in the JAS subgroups; it	
may also have value to identify critical windows of susceptibility. The	
benefits of this approach should be considered with the drawbacks, such	
as substantially increasing the required number of animals and difficulties	
interpreting data at different ages. See Section 3.7 Dose Selection	
regarding dose adjustment as an alternate strategy to be considered in this	
situation.	
Dosing frequency in JAS may not be exactly the same as in the clinical	
regimen. For example, even though a clinical regimen is once a week, daily	
dosing in juvenile animals can be needed to achieve and maintain relevant	
systemic exposures to evaluate the effects on developing organ systems	
and/or to maintain systemic exposures at relevant levels during the entire	
developmental period of concern.	
	<ul> <li>comparable to or shorter than the dosing period in the JAS subgroups; it may also have value to identify critical windows of susceptibility. The benefits of this approach should be considered with the drawbacks, such as substantially increasing the required number of animals and difficulties interpreting data at different ages. See Section 3.7 Dose Selection regarding dose adjustment as an alternate strategy to be considered in this situation.</li> <li>Dosing frequency in JAS may not be exactly the same as in the clinical regimen. For example, even though a clinical regimen is once a week, daily dosing in juvenile animals can be needed to achieve and maintain relevant systemic exposures to evaluate the effects on developing organ systems</li> </ul>

337-368	3.5 Off-Treatment Period Assessments	
557-500		
	Inclusion of an evaluation period after treatment has stopped in a JAS can	
	help address two issues: 1) whether any effects observed during treatment	
	are reversible, persistent, or progressive and 2) whether any effects	
	emerge later in development as a result of early life exposure (i.e., delayed	
	onset of changes). The need for an off-treatment period is dependent on	
	the outcome of the WoE assessment and the endpoints to be evaluated in	
	the study.	
	In general, an off-treatment period should be included to understand	
	persistence, progression, or reversibility of a specific effect if this cannot	
	be predicted by scientific assessment (Note 1). The principles of	
	reversibility in ICH M3 apply to JAS endpoints that are similar to those in	
	repeat-dose toxicity studies in adults (e.g., histopathology, clinical	
	pathology). The duration of such an off-treatment period should be	
	sufficient to allow the potential recovery of the effect, and should take into	
	account the elimination of the pharmaceutical. The demonstration of full	
	reversibility is not considered essential. A trend towards reversibility	
	(decrease in incidence and/or severity) and a scientific assessment that	
	this would eventually progress to full reversibility could be sufficient. If	
	reversibility or irreversibility of a specific effect is well characterized in	
	adult animals, it is generally not necessary to confirm this in a JAS. There	
	are endpoints in a JAS that are not amenable to the classic approach of	
	reversibility assessment, such as the timing of onset of puberty and	
	neurobehavioral assessments (e.g., learning). Additionally, the timing of	
	the off-treatment period in relation to the life stage of the animals should	

be considered.	
Some alterations can only be identified following an appropriate	
off-treatment period to allow maturation of an organ system and	
expression of the alteration. Therefore, some assessments can only b	e
meaningfully performed after a certain level of maturity is expected t	o be
reached (e.g., behavioral assessment, immunological response in	
T-cell-dependent antibody response [TDAR]). These assessments can	be
conducted in off-treatment periods after dosing duration has covered	lall
critical developmental windows related to the clinical use. This is esp	ecially
relevant in cases in which the clinical population is only the very your	ng,
	-
In non-rodents, the addition of post-treatment groups for JAS can be	less
useful due to the more protracted development period, high	
inter-individual variability, and fewer and less well characterized	
assessments available to identify delayed or altered development.	
3.6 Route of Administration	
The intended clinical route of administration should be used when	
feasible, but obtaining adequate systemic exposure is paramount.	
Alternative administration routes should be considered in cases of	
practical difficulties; changing routes during the course of the study c	an
	off-treatment period to allow maturation of an organ system and expression of the alteration. Therefore, some assessments can only b meaningfully performed after a certain level of maturity is expected t reached (e.g., behavioral assessment, immunological response in T-cell-dependent antibody response [TDAR]). These assessments can conducted in off-treatment periods after dosing duration has covered critical developmental windows related to the clinical use. This is espi relevant in cases in which the clinical population is only the very your such that the JAS dosing duration would cease at an immature age ar animals will continue to mature during the off-treatment period to ar that an appropriate assessment can be conducted.In non-rodents, the addition of post-treatment groups for JAS can be useful due to the more protracted development period, high inter-individual variability, and fewer and less well characterized assessments available to identify delayed or altered development. <b>3.6 Route of Administration</b> The intended clinical route of administration should be used when feasible, but obtaining adequate systemic exposure is paramount.

	redents). The validity of using an alternative desing route should be	
	rodents). The validity of using an alternative dosing route should be	
	justified (e.g., supported by TK data in representative juvenile animals).	
	If the pharmaceutical is intended for use by two or more clinical routes of	
	administration, a JAS with a single route of administration is sufficient, but	
	should provide adequate exposure in juvenile animals for all intended	
	clinical routes of administration.	
379-398	3.7 Dose Selection	
	It is desirable to establish a dose-response relationship for adverse effects	
	and to determine a no-observed adverse effect level (NOAEL) in juvenile	
	animals. Dose levels should be selected to achieve some overlap in the	
	range of exposure in adult animals to enable comparison of effects	
	between adults and young animals. However, the high dose should not	
	result in marked toxicity that can confound the growth and development	
	endpoints and complicate the assessment. Body weight loss or lack of gain	
	during rapid growth periods has the potential to confound results, and is	
	therefore not desirable in a JAS. The low dose should preferably result in	
	exposure levels similar to the anticipated exposure in the intended clinical	
	population. For small molecules, selection of the high dose in accordance	
	with ICH M3 applies. For biotechnology-derived products, the principles	
	for dose selection described in ICH S6 apply.	
	There can be changes in systemic exposure due to maturation of the	
	ADME systems that can make it challenging to meet the dose selection	
	aims described above. In cases in which preliminary studies demonstrated	
	that juvenile animals are markedly more sensitive than adult animals, or	

	there are substantial changes in systemic exposure as the animals mature,
	dose adjusting should be considered. Dose adjustment (dose increase or
	decrease) during a JAS can be appropriate to evaluate endpoints when
	exposure separation between dose levels can otherwise not be maintained
	throughout the study. Adjusting doses during the study is intended to keep
	the exposures somewhat consistent; generally, not more than one or two
	adjustments during a JAS would be expected.
399-558	3.8 Endpoints
	Each JAS should include the core endpoints defined in Section 3.8.1 below,
	unless justified otherwise. Each additional endpoint (see Section 3.8.2)
	should be considered and justified to address an identified safety concern
	(Note 2).
	For the interpretation of study results in JAS it is important to have
	appropriate historical control data (HCD) at relevant ages of the
	species/strain/sex used (Note 3).
	3.8.1 Core Endpoints
	3.8.1.1 Mortality and Clinical Observations
	Mortality should be evaluated throughout the experimental period.
	Clinical observations, including physical examinations, should be
	conducted as they can identify overt behavioral effects both on and off
	treatment
	Clinical observations during the lactation period should include maternal
	nursing behavior, and should capture clinical observations unique to
· · · ·	

juvenile animals as much as possible. After weaning, clinical observations should be recorded as for adult animals.

#### 3.8.1.2 Growth

Growth should be assessed by body weights in conjunction with long bone length. As body weight increases dramatically during the early postnatal period, individual weight measurements should be frequently recorded to inform dose calculations. Generally, one long bone (e.g., femur) measured for length at necropsy is sufficient (Note 4).

### 3.8.1.3 Food Consumption

Food consumption during the post weaning period should be assessed as appropriate for the species.

### 3.8.1.4 Sexual Development

The physical indicators of onset of puberty (e.g., for rodents, the age of vaginal opening in females and balanopreputial separation in males) are recommended when the treatment period encompasses the relevant developmental window.

## 3.8.1.5 Clinical Pathology

Standard clinical pathology examinations (serum chemistry and hematology) should be assessed as a terminal endpoint at necropsy if evaluation is planned at an age in which expected clinical pathology ranges are known and can support interpretation of histopathology findings.

### 3.8.1.6 Anatomic Pathology

At the end of the treatment and/or off-treatment periods, gross pathology, organ weights (Note 5), and comprehensive collection and preservation of tissues should be conducted for animals allocated to necropsy. Histopathology should be performed on major organs (e.g., bone, brain, ovary, testis, heart, kidney, liver) and those with macroscopic lesions. Testicular histopathology should include a qualitative evaluation of spermatogenic progression in mature animals.

### 3.8.1.7 Toxicokinetics

TK sampling should be conducted near the beginning and end of the dosing period. If dosing is started pre weaning, interim TK assessment(s) should be considered. A preliminary or DRF JAS with TK assessment, which is recommended (see Section 3.2), will inform on the sampling day and the time points of sample collection.

When designing the TK component of a JAS, microsampling and sparse sampling (if justified) are strongly encouraged (see ICH S3) from the view of 3Rs.

For protein therapeutics, samples for anti-drug antibodies should be collected and evaluated if appropriate (see ICH S6).

## 3.8.2 Additional Endpoints to Address Identified Concerns

### 3.8.2.1 Growth

As appropriate for the species, crown rump length, body length (e.g., nose/tail), and/or withers height can be used as an indicator of growth. Serial non-invasive measurement of long bone length using ultrasonic echo or X-ray can be appropriate in non-rodents in addition to a direct measurement at necropsy.

## 3.8.2.2 Skeletal Examinations

When there is an identified concern about bone metabolism or structure, the measurements of bone-related biomarkers and/or expanded histopathology (e.g., histomorphometry) should be considered. Assessment of bone mineral density (e.g., micro densitometry, dual energy X-ray absorptiometry, peripheral quantitative computed tomography [CT]) or bone structure (e.g., micro CT) can also be conducted as appropriate.

# 3.8.2.3 Clinical Pathology

Additional haematology, serum chemistry, and/or biomarkers can be considered to further characterize identified concerns on target organs/tissues. Other parameters such as urinalysis or coagulation assessments can be added when warranted.

Samples collected throughout the study at different ages and/or a series of samples collected within a short time period (e.g., 24 to 48 hours) can also be useful.

Due to the limitation in obtaining adequate sample volumes from juvenile animals (especially rodents), any additional samples that may require additional animals therefore are only recommended when critical to address a concern. When sample volume constraints exist, the parameters to be measured should be selected according to a priority based on the identified concern(s).

#### 3.8.2.4 Anatomic Pathology

Additional tissues/organs can be evaluated to address specific concerns. Immunohistochemical or other special staining methods for tissue sections, electron microscopy, histomorphometry, or other imaging techniques can be warranted for interpretation of some findings.

#### 3.8.2.5 Ophthalmologic Examinations

When there is concern for ocular toxicity, including retina and optic nerve, assessment of ocular endpoints should be considered. Standard ophthalmological examinations (e.g., palpebral reflex, ophthalmoscopy, slit-lamp microscopy) are not a routine endpoint for JAS, because structural development of the eye is largely completed during the prenatal period in humans.

3.8.2.6 CNS Assessments
There are different categories of CNS assessments, such as:
detailed clinical observations
behavioral tests
learning and memory tests, and
expanded neuropathology evaluations
Selection of any additional CNS assessments should be based only on the
particular concerns identified in the WOE evaluation. In addition, the
timing of these assessments should take into consideration whether the
results will be used to identify adverse effects due to an extension of
pharmacology, developmental neurotoxicity (i.e., effects that emerge or
are still present after cessation of treatment) or both.
Detailed CNS-related clinical observations document the severity and the
onset and duration of the clinical signs relative to dosing (e.g., hyper- or
hypoactivity, tremors). These parameters should be assessed when a CNS
concern has been identified by the WoE evaluation and should be
collected during on- and off-treatment periods as appropriate.
Behavioral testing can include a modified Irwin test, functional
observational battery (FOB), assessment of locomotor activity, evaluation
of coordination and reflexes, and/or acoustic startle response (e.g.,
habituation or prepulse inhibition). These tests should be appropriate for
the species being tested and the timing of these assessments should be
determined relative to the level of maturity in the test species.

In addition, learning and memory can be evaluated by a variety of methods. Different methods assess different aspects of learning and memory. When specific aspects of learning and memory have been identified as areas of concern based on the WoE evaluation, then tests capable of assessing those aspects should be selected. Learning and memory should be evaluated typically during the off-treatment period as this period is most relevant to assess potential persistent or delayed effects. If learning and memory testing is performed during the treatment period, the potential for confounding pharmacological effects (e.g., sedation, decreased motor coordination) should be considered and avoided.

Any CNS areas or components (e.g., hippocampus, myelin) that are identified by the WoE evaluation as potential targets of concern should be assessed with additional neuropathological examinations as appropriate (e.g., additional levels for sections, immunohistochemistry, special stains). These assessments are typically performed at times of scheduled necropsy, unless there is a specific concern related to timing to be investigated. Imaging technologies may also be useful in specific circumstances (e.g., magnetic resonance imaging).

Postnatal CNS assessments are most commonly conducted and characterized in the rat. For those pharmaceuticals where the rodent is an inappropriate species, some behavioral tests are also available in other species (e.g., dogs, minipigs). Learning and memory assessments are infrequently conducted in NHPs. In NHPs, behavioral observations can

	-	
	provide the primary assessment of potential CNS effects in a JAS or ePPND	
	study.	
	3.8.2.7 Reproductive Assessments	
	If there is an identified concern for effects on female and/or male	
	reproductive organs or function, histopathology examinations and organ	
	weights can be expanded to include reproductive and/or endocrine tissues	
	in addition to the gonads. Reproductive system effects identified as	
	irreversible in adult animals need not be confirmed in a JAS.	
	In rodents, for concerns relevant for females, assessment of estrous	
	cyclicity is recommended as an initial assessment of reproductive and	
	endocrine function. For concerns relevant for male rodents, sperm analysis	
	(e.g., counts, motility, morphology) and/or testicular	
	immunohistochemistry can be considered to further characterize effects if	
	they can add critical information not already captured elsewhere.	
	The timing of the treatment and assessments in relation to that of sexual	
	maturation in the species tested is critical. The timing of folliculogenesis	
	and spermatogenesis should be considered in the study design and timing	
	of reproductive assessments. Assessment of reproductive organs or	
	function (e.g., estrous cyclicity, sperm count, or qualitative histologic	
	assessment of spermatogenesis) can only be conducted in sexually mature	
	animals. If the clinical age range 530 is prepubertal, the concern is	
	whether treatment of a medicine with reproductive toxic potential would	
	cause any delayed effect on sexual maturation or reproductive function in	
	adulthood. In this situation, a study should be designed to treat only	

during immaturity, and then allow the animal to mature without treatment, and conduct assessments after maturation is reached.

Mating assessments are not generally recommended in JAS. In male rodents, mating assessments have low sensitivity due to a large functional reserve of the testis. In female rodents, assessment of estrous cyclicity and ovarian histology can identify many developmental reproductive liabilities. In non-rodent species mating assessments are not practical due to the protracted duration of development and high degree of individual variability.

The feasibility of other additional reproductive assessments is such that the large majority are conducted in rodents, although they can be considered for those nonrodent species that achieve maturity during the conduct of a JAS. In NHP, additional reproductive assessments are not typically included in JAS.

Hormonal assessments are only recommended in JAS if they can add critical information not already captured elsewhere as there is considerable hormonal variability during puberty. Any hormone assessment should be justified, and the timing and specific hormones assessed should be well characterized for the age the assessment occurs.

If the pharmacological class or data in animals or humans give cause for	
concern for the development of the immune system, assessments for	
immunotoxicity should be considered as outlined in ICH S8. Such concerns	
can include, but are not limited to, a transient, prolonged or permanent	
decrease or increase in the number or function of a lymphocyte subtype	
or a sustained increase or decrease in immunoglobulin class. Functional	
assays such as the TDAR should be performed after appropriate times of	
development (e.g., after PND 45 for the rat).	
3.8.2.9 Other Possible Assessments 555	
If there are additional tissues or endpoints for which concerns are	
identified and cannot be managed clinically, appropriate evaluations	
should be planned and performed when nonclinical investigations can add	
useful information.	
3.9 Allocation of Animals to Study Groups	
3.9.1 Preweaning Allocation	
In most species, initiation of a JAS during the preweaning phase presents a	
unique situation of dosing offspring within a litter. The maternal animal is	
a critical component of the study providing nutrition and care, but only the	
offspring are the test system. The study should be designed to reduce	
potential confounders of data from offspring related to genetics, maternal	
care, and littermates (i.e., nature and nurture confounders). Generally,	
genetic siblings and/or littermates should not be assigned to the same	
endpoints, especially for the core study endpoints. This can be achieved by	
the way the litters are constructed in combination with how they are	
	<ul> <li>immunotoxicity should be considered as outlined in ICH S8. Such concerns can include, but are not limited to, a transient, prolonged or permanent decrease or increase in the number or function of a lymphocyte subtype or a sustained increase or decrease in immunoglobulin class. Functional assays such as the TDAR should be performed after appropriate times of development (e.g., after PND 45 for the rat).</li> <li><b>3.8.2.9 Other Possible Assessments 555</b>         If there are additional tissues or endpoints for which concerns are identified and cannot be managed clinically, appropriate evaluations should be planned and performed when nonclinical investigations can add useful information.     </li> <li><b>3.9 Allocation of Animals to Study Groups 3.9.1 Preweaning Allocation</b>         In most species, initiation of a JAS during the preweaning phase presents a unique situation of dosing offspring within a litter. The maternal animal is a critical component of the study providing nutrition and care, but only the offspring are the test system. The study should be designed to reduce potential confounders of data from offspring related to genetics, maternal care, and littermates (i.e., nature and nurture confounders). Generally, genetic siblings and/or littermates should not be assigned to the same endpoints, especially for the core study endpoints. This can be achieved by</li></ul>

assigned to dose groups and subsets of endpoints.

It is advisable to utilize litter sizes and sex ratios reasonably similar to the natural mean litter sizes for that species and strain. As for the method of assigning dose groups, it is desirable to prevent animals in a control group from being exposed to the test pharmaceutical, thus is it preferred that all animals in a litter be assigned to the same treatment group.

JAS can become large and complex, therefore it is especially important that the study design balances scientific rigor against animal use. Investigators should know all the planned endpoints (core and additional) to design the littering and subset assignment strategy efficiently. Efficiency in study design is critical to reduce animal use as per the 3R principles, and should be measured by the number of maternal animals and litters needed to supply the study. For animal species with low and variable litter sizes or single offspring, the same approach for group allocation design as in general toxicity studies can be appropriate.

After the study has started, each litter size should remain comparable across and within dose groups, as much as possible, while in the preweaning phase because litter size affects pup growth rate. Litter handling, dose group and endpoint subset allocation methods, and specifics of the testing model (e.g., age when litters culled, litter size and sex distribution, fostering, assignment of groups and subsets for evaluation) should be clearly described in the study plan and report. For statistical analysis, data collected from offspring while part of a litter

r		1
	should not be o	onsidered an independent variable since an individual
	offspring is dep	endent on maternal and 587 littermate factors.
	There are differ	ent allocation methods for litter management in
	preweaning, m	ultiparous animals. Appendix C provides one example of an
	approach for ro	dents that controls for potential genetic, maternal care,
	and littermate	plases. Other methods are acceptable if they appropriately
	consider these	piases and the study objectives.
	3.9.2 Postwear	ing Allocation
	In multiparous	animal species, if possible, it is still recommended to
	allocate the litt	ers to minimize the genetic bias and maternal and
	littermate varia	bles. In particular when dosing starts in the early
	postweaning pl	ase, and, when offspring are supplied from a limited
	number of natu	ral mothers in the test facility, the study should be
	designed in cor	sideration of the potential confounders similar to those at
	preweaning all	ocation.
599-603	3.10 Animal Nu	mbers and Sex
	A JAS should us	e an adequate number of animals to evaluate the selected
	endpoints (e.g.	body weights, reversibility, behavioral assessments). To
	reduce the nun	ber of animals, combining assessment of endpoints in the
	same animals o	an be effective. It is recommended that JAS be performed
	in both female	and male animals.

604-647		Section 3 should be consulted to determine study designs needed to	
001017		address the points below.	
		A common clinical approach for non-oncology pediatric-only/first	
		pharmaceuticals starts with a First in Human (FIH) study in healthy adult	
		volunteers prior to any pediatric trial. As per ICH M3, this approach	
		generally includes nonclinical repeat-dose toxicity studies of appropriate	
		duration in rodent and non-rodent animals as well as safety pharmacology	
		and genetic toxicology studies before initiation of adult clinical trials.	
		Principles of ICH S6 can also apply. The repeat-dose toxicity studies to	
		support FIH in adults could be performed in several ways; in both species	
	4. CONSIDERATIONS	in adult animals or in one or both species by initiating dosing in juvenile	
	FOR	animals and continuing treatment into maturity including additional	
	PAEDIATRIC-FIRST/ONLY	endpoints (see Sections 2 and 3).	
	DEVELOPMENT		
		Alternatively, there are cases where pediatric patients are treated without	
		any prior adult patientor healthy volunteer data (e.g., for a life-threatening	
		or debilitating disease that only exists in children and when the	
		pharmaceutical cannot be given safely to adult volunteers). In these cases,	
		the FIH trial will be in pediatric patients and the nonclinical program would	
		generally include one JAS in a rodent and one JAS in a non-rodent species,	
		if feasible. Safety pharmacology and genotoxicity testing would be	
		conducted as appropriate for adult use; in vivo studies need not be	
		conducted in juvenile animals (see Section 2.3.4).	
		After initial clinical trials, JAS can be important to support continued	

clinical development in pediatric patients on a case-by-case basis, driven	
by cause for safety concern (see Section 2) and duration of clinical	
treatment. The principles of ICH M3 should also be considered. If the	
pharmaceutical is intended to treat a chronic disease, chronic toxicity	
studies should be conducted in one rodent and one non-rodent species. In	
at least one of these studies, dosing should start at an age	
developmentally matched to the lowest age of the intended patient	
population. In principle, a single set of chronic studies that start dosing	
from ages that developmentally correlate to the youngest pediatric patient	
age can provide nonclinical safety data sufficient to cover all ages and	
durations of pediatric development up to marketing, and can replace adult	
chronic and separate JAS. Further nonclinical assessments of reproductive	
toxicity and carcinogenic potential can be warranted.	
For biopharmaceuticals, studies in juvenile animals should be limited to	
relevant species, as per ICH S6. When the NHP is the only relevant species,	
a JAS in NHPs could support initial clinical use. Non-invasive safety	
pharmacology endpoints can be included in the juvenile or standard NHP	
repeated-dose studies. Genotoxic and carcinogenic potential should be	
addressed as outlined in ICH S6.	
JAS in NHP are typically conducted starting at 10-12 months of age, thus	
limiting the lowest pediatric age ranges. In cases where JAS is not feasible	
to support the youngest pediatric age, alternative approaches (e.g., in vitro	
assays, genetically-modified animals, surrogate molecules) should be	
considered if available and relevant.	

		A JAS in perinatal and preweaning NHP should only be conducted in the	
		situation of medicines with first and primarily neonatal clinical use, and	
		where alternative approaches to nonclinical safety assessment are not	
		feasible. Studies with direct dosing of offspring can require large numbers	
		of mature dams to populate even a relatively small JAS in NHP. Therefore	
		the design and endpoints should be clearly justified based on the clinical	
		concern. Design expectations should also be flexible; for example,	
		variability in gender distribution and starting weights are expected.	
648-658		5.1 Excipients	
		Dedicated JAS on excipients are generally not needed to qualify pediatric	
		formulations. To assess the safety of the pediatric clinical formulation,	
		available toxicity information on the excipients should be evaluated.	
		Pharmaceutical formulations used in pediatric indications can occasionally	
		contain novel excipients or excipients not previously used in pediatric	
		populations of a relevant age. If there are insufficient data to support the	
		use of the excipient in the intended pediatric population, a JAS can be	
	5. OTHER	warranted. Although JAS that are primarily intended to assess the safety of	
	CONSIDERATIONS	active ingredients need not always be conducted with the clinical	
		formulation, an excipient could be assessed in a JAS along with the active	
		ingredient, if such studies were being conducted.	
659-672		5.2 Combination Pharmaceuticals	
		The development of combination pharmaceuticals for pediatric use should	
		have a nonclinical evaluation consistent with the principles outlined in ICH	
		M3 (R2) for combination products in general together with the WoE	
		principles outlined in this guideline. For example, a combination JAS would	
		generally not be recommended for a combination of two late stage	

· · · · · · · · · · · · · · · · · · ·	[		
		entities for which there is adequate pediatric clinical experience with	
		co-administration. Whereas, a combination JAS might be warranted for a	
		combination of two early stage entities if a WoE evaluation suggests that a	
		JAS would address identified concerns. If additional nonclinical information	
		is needed, the study design should consider what assessment endpoints	
		are appropriate to address any concerns of administering the particular	
		combination. If a JAS is considered appropriate, assessment of the	
		combination as it is to be used clinically is generally sufficient and testing	
		of the individual active ingredients may not be critical. Alternatively, an	
		extra group with the combination could be added to a JAS that is already	
		being conducted with one of the single entities. This could eliminate the	
		need to do a separate study with the combination product.	
673-698		Enhanced Pre- and Postnatal Development Study (ePPND):	
		This study design is based on biopharmaceutical (NHP) experience and is a	
		PPND study which includes elements of the embryofetal development	
		(EFD) study in newborns and infants instead of the fetus.	
		Juvenile:	
		Any postnatal stage not fully matured in terms of morphology and	
	6. GLOSSARY	function	
		Pediatric First:	
		Pediatric-first development is when the pharmaceutical is developed for	
		pediatric patients before any clinical data are available in adults for any	
		indication.	
	I		

		Padiatuia Onlu	
		Pediatric Only:	
		Pediatric-only development describes development for an indication	
		requiring treatment 687 exclusively in pediatric ages (e.g., neonatal	
		respiratory distress syndrome).	
		Weight of Evidence:	
		An approach that evaluates a combination of information from several	
		independent sources to 691 determine if there is sufficient evidence to	
		support pediatric clinical trials or whether 692 additional nonclinical	
		assessments are recommended to address safety concerns that cannot be	
		managed clinically.	
		The weight given to the available evidence depends on factors such as the	
		quality of the data, consistency of results, nature and severity of effects,	
		and relevance of the information. The weight of evidence approach	
		requires use of scientific judgment and, therefore, should consider the	
		robustness and reliability of the different data sources.	
699-724		Note 1	
		If the off treatment period begins prior to maturity, the capacity and	
		character of the recovery can be influenced by the continued growth	
		and development of some organ systems, and should be carefully	
	7. NOTES	interpreted.	
		Note 2	
		The propensity for mortality to occur is generally higher in juvenile	
		animals compared to adult animals. Study-related procedures should be	

limited as much as possible before and at the time of weaning as they can contribute to mortality.

## Note 3

Assessments on immature animals should be done with reference to age-matched control data (e.g. body weights, clinical pathology, organ weights, histology) either from concurrent control animals or from other reference background data. This is especially important to consider in cases of unscheduled assessment of endpoints. JAS animals are generally not screened prior to initiation of treatment. Therefore, background rates of abnormalities in juveniles can differ from animals of the same age used in adult toxicity studies.

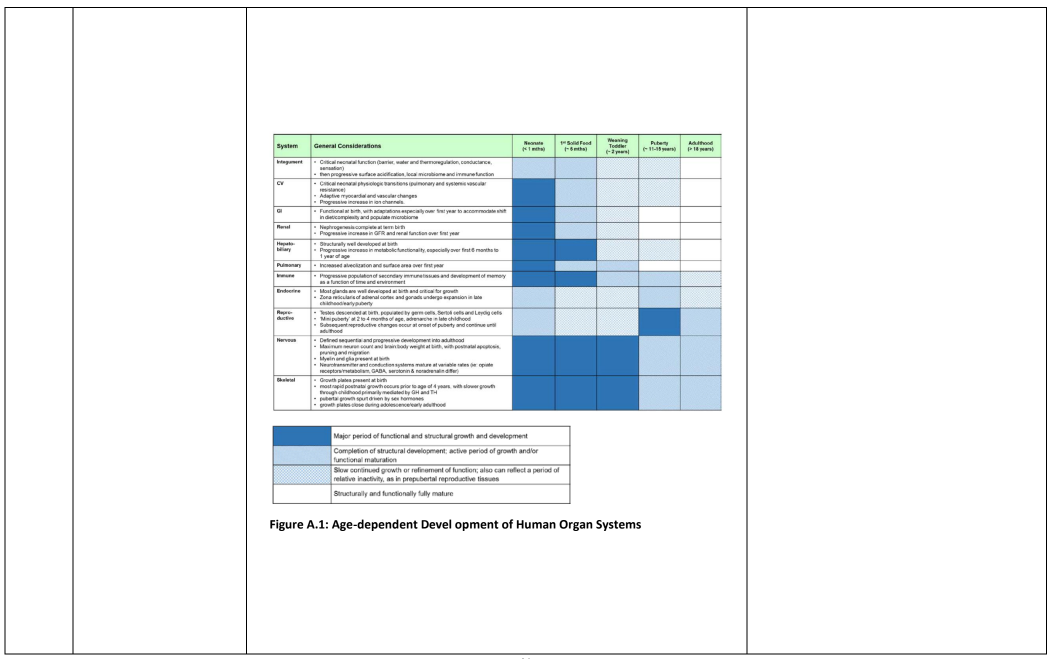
## Note 4

Since growth happens in spurts, frequent assessments of bone length for 'transient' effects on growth is challenging to appropriately power and offers limited value. An assessment using data from the end of treatment is more useful. An effect solely on decreased body weight gain is not necessarily an effect on growth.

## Note 5

Assessment of organ weight data should be done in the context of growth. For instance, if growth was restricted then absolute weights of most organs decrease in proportion to body weight; however, some organs have different sensitivity to growth effects.

725-735		1. ICH E11 Guideline: Clinical Investigation of Medicinal Products in the					
		Pediatric 726 Population; 2017					
		2. ICH M3 Guideline: Guidance on Nonclinical Safety Studies for the					
		Conduct of Human Clinical Trials and Marketing Authorization for					
		Pharmaceuticals; June 2009.					
	8. REFERENCES	3. ICH S5 Guideline: Detection of Toxicity to Reproduction for Medicinal					
		Products 731 and Toxicity to Male Fertility; 2000					
		4. ICH S6 Guideline: Preclinical Safety Evaluation of					
		Biotechnology-Derived 733 Pharmaceuticals; 2011					
		5. ICH S9 Guideline: Nonclinical Evaluation for Anticancer					
		Pharmaceuticals; 2009					
736-773		These tables reflect a high level overview of organ system development by					
		species to illustrate similarities and differences between the commonly					
	9. APPENDIX A:	used toxicology species, as compared to humans, for the timing and					
	OVERVIEW OF	relative duration of development. Specific milestones include birth,					
	AGE-DEPENDENT DEVELOPMENT OF	introduction of solid foods, weaning, puberty, and adulthood. The tables					
		are intended to aid in the assessment of the relevance of existing					
	ORGAN SYSTEMS BY	nonclinical data, as well as the selection of species, starting age, and					
	SPECIES	dosing duration for a JAS. These summary tables are based on a review of					
	JFLUILJ	current knowledge, but are not comprehensive. Species-specific and/or					
		organ system reviews in the literature can provide additional detail and					
		should be consulted for each specific situation.					

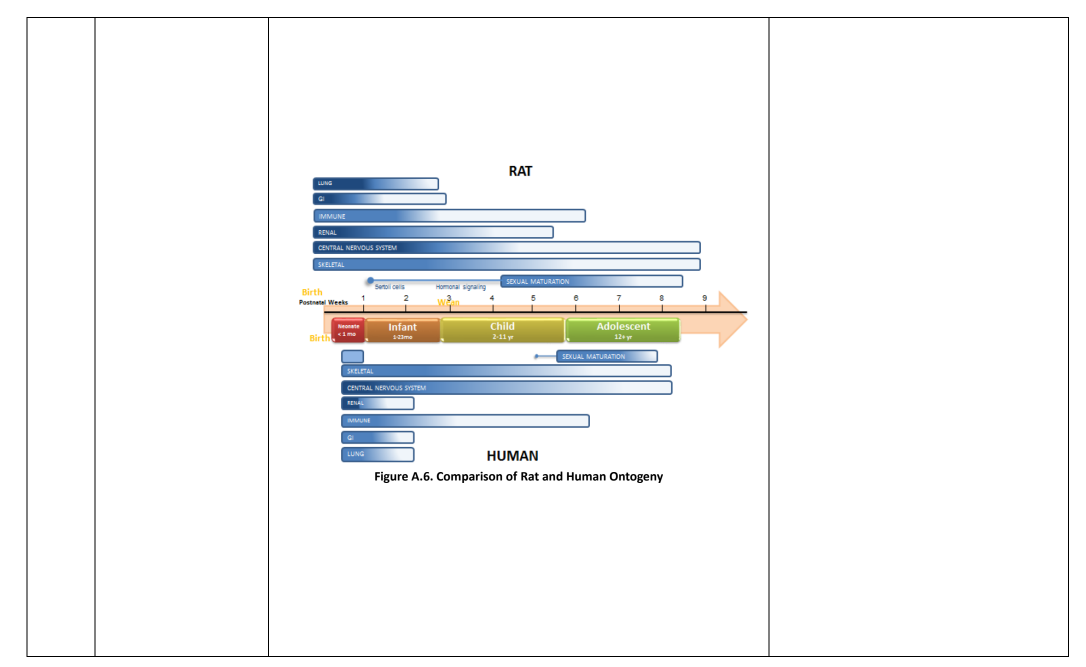


Integrated       • Circle in constal function (pairwine), water and thermoregulation, conductance, estatation), • These regulations in all 2 weeks of age       • Circle and constal functions (pairwine), water and thermoregulation, conductance, estatation), • These regulations is add to pPO_21         CV       • Social and constal functions (pairwine), water and thermoregulation, conductance, estatation, • Propression (monstale of social charges)       • Social and constal functions (pairwine), water and thermoregulation, conductance, estatation, • Propression (monstale of social charges)         GL       • Formation and social charges       • Social and constal functions (pairwine), water and thermoregulation, conductance, estatation, • Propression (monstale to PPO 21         Propression (monstale of social charges)       • Social and constal function confirms: 3 to 5 weeks of age         Rend       • Propression (monstale of social charges)       • Social and constal function confirms: 3 to 5 weeks of age         Network       • Social and charges of pairs (pairwine)       • Social and charges of pairs (pairwine)         Network       • Social and charges of pairs (pairwine)       • Social and charges of pairs (pairwine)         Network       • Social and charges of pairs (pairwine)       • Social and charges of pairs (pairwine)         Network       • Social and charges of pairs (pairwine)       • Social and charges of pairs (pairwine)       • Social and charges of pairs (pairwine)         Network       • Social and charges of pairs (pairwine)       • Social and charges of	System	General Considerations	Neonate (~ PND 1-10)	1 <sup>st</sup> Solid Food (~ PND 15)	Weaning (~ PND 21-25)	Puberty (M ~ PND 42, F ~ PND 35)	Adulthood (~ PND 70)	
CV <ul> <li>Critical monostrati drybinologic transitions (pulmonary and systemic vascular relations)         <ul> <li>Adaption in processing in consense in cardiomy costs and in on humanitis to PND 21</li> <li>Clinical and vascular charges</li> <li>Progressine increases in cardiomy costs and in on humanitis to PND 21</li> </ul> </li> <li>Clinical intervention small interactive engines and participation until after products         <ul> <li>Progressine increases in cardiomy costs and participation and interact robusins</li> <li>Progressine increases in GPR and recent function over first 3 to 5 weeks of age</li> <li>Progressine increases in GPR and recent function over first 3 to 5 weeks of age</li> </ul> </li> <li>Progressine increases in GPR and recent function over first 3 to 5 weeks of age</li> <li>Progressine diversity on thirst varies and a device and plates, with increase in metabolic increases of age</li> <li>Progressine diversity on thirst varies and device promoter of many and equivalence of age of the progressine diversity on thirst varies and device promoter of many and equivalence of age of the progressine production of the progressine diversity on the progressine diversity of the progressine diversity of the promoter of the progressine diversity of the progressine diversity of the progressine production of the progressine of the progressine production of the progressine of the progressine diversity of the progressine diversity of the progressine diversity of the progressine diversity of the progressine of the progressine diversity of the progressine of the progressine diversity of the progressine of the progressine of the progressine diversity of the progressine of the progressine diversity of the progressine of the progressine diversity of the progressine of the progressine dintervene din the progresis of the progressine of thep</li></ul>	Integument	<ul> <li>sensation)</li> <li>Thicker epidermis first 2 weeks of age</li> </ul>						
GI <ul> <li>Immuture at birth; bick gashte acid and poor pancreatic enzyme production unit after PND 14</li> <li>Highly parmable proximal small intestime allows absorption of mater proteins Adaptations in Sind evek postatatial to accommodate at birth Progressive incomplete at birth and cillcal for growth Repor- Progressive incomplete at birth and cillcal for growth Report Progressive incomplete at birth Completion of discretized and opport discretized and incomplements Report Progressive incomplete at birth Completion of structural growth through adulthood Progressive incomplete at birth Completion of structural development; active period of growth and/or</li>         Progressive incoma</ul>	cv	Critical neonatal physiologic transitions (pulmonary and systemic vascular						
Renal       • Neptrogenesis incompties at bith • Progressive incomes in GER and renal function over first: 3 to 5 weeks of age       Image: Comparison of Comparison over first: 2 to 5 weeks of age         Hepato- bility       • Structurally immature at bith • Progressive development of organized hepatic cords and plates, with increase in metabolic functionality own first 4 weeks of age       Image: Comparison over first 2 to 3 weeks of age         Pulmonary       • Secular at bith • Andorization occurs over first 2 to 3 weeks of age       Image: Comparison occurs over first 2 to 3 weeks of age         Immune       • Progressive population of secondary immune issues and development of memory as a function of the mand environment • TDAR typically assessed after PND 45       Image: Comparison over first 2 to 3 weeks of age         Repre- ductive       • Period of docreased androgen production by Leydig cells during 3rd week postnatal meets any for expansion of germ cells and Serbil cells • conset of pathety (constatil week 5 to 7)       Image: Comparison of the comparison of secure dimension plates and development, enset of pathety (constatil week 5 to 7)         Nervous       • Structural mutuation of officitory bubs, relinately, exceebellum, hippocampus, and ceetion of opticity (constatil week 5 to 7)       Image: Comparison of the comparison of an advelopment of advelopment at different tabs         Skeletal       • Regio constatil growth hough adutitood • Long bone growth plate structure not evident until PND 14 to 21, and remain open into aduthood       Image: Completion of structural growth and development; active period of functional and structural growth and/or       Image: Completion of st	GI	Immature at birth; lack gastric acid and poor pancreatic enzyme production until after PND 14     Highly permeable proximal small intestine allows absorption of intact proteins						
Hepsite- bilisity       Statusturally immature at birth Progressite development of and plates, with increase in advelocition occurs over first 2 to 3 weeks of age       Immature       Progressite development of manual evolution occurs over first 2 to 3 weeks of age         Pulmonary       - Saccular at birth advelocition occurs over first 2 to 3 weeks of age       Immune       Immune       Progressite population of secondary immune issues and development of memory as a function of time and environment to TDAR typically assessed after FND 45       Immune       Immune         Endocrine       - Most glandar as well developed at birth and critical for growth revised are well developed at birth and critical for growth environment of for and environment to mass of public be charges and appearance of sexual dimorphism occur at onset of public ty (postnatil week 5 to 7)       Immune         Nervous       - Structural maturation of olfactory buls, relinality or evolution, weight at PND 7, with extensive postnatal appoints, round and migration . Meelin not present at birth pathways mature at different rates       Immune at the evolution of the structural or evolution of a structural growth and development.         Steletal       - Rapid post and prosting in DS and pathhood . Orapidation of functional and structural growth and development.       Immune at the evolution of a structural growth and development.	Renal							
Immune        • Progressive population of secondary immune tissues and development of memory as a function of time and environment       • TORK typically assessed after PDIA 5         ■       ■       ■       ■       ■		Structurally immature at birth						
as a function of time and environment     as a function of time and environment       Endecrine     • Most glands are well deviceed at birth and critical for growth       Repro- ductive     • Period of decreased androgon production by Legidg colls during 3rd week postnatal necessary for expansion of germ cells and Serbil cells     •       Repro- ductive     • Period of decreased androgon production by Legidg colls during 3rd week postnatal necessary for expansion of germ cells and Serbil cells     •       • Structural maturation of ollscore y builds, reinaleys, cerebellum, hippocampus, and cerebral crobes occurs postnatally     •       • Structural maturation of ollscore y builds, reinaleys, cerebellum, hippocampus, and cerebral crobes occurs postnatal apoptoles, pruning and migration • Heydin not present at birth • Theydin not present at birth • Angid postnatal growth through adulthood • Long bone growth plate structure not evident until PND 14 to 21, and remain open into adulthood       Skeletal     • Rajid postnatal growth through adulthood • Long bone growth plate structure not evident until PND 14 to 21, and remain open into adulthood	Pulmonary	Saccular at birth     alveolization occurs over first 2 to 3 weeks of age						
Repro- ductive <ul> <li>Period of decreased androgen production by Leydig cells during 3rd week postnatal necessary for expansion of germ cells and Bestolicells</li> <li>Remaining reportative changes and appearance of sexual dimorphism occur at onsel of publicity (postnatal week 5 to 7)</li> <li>Nerveus</li> <li>Structural matrixets on of lactory bubs, refinaleye, cerebellum, hippocampus, and cerebral crotex occurs postnatali "Mainrum meunication of olactory bubs, refinaleye, cerebellum, hippocampus, and cerebral crotex occurs postnatali "Mein on present at birth</li> <li>Conduction systems, opiate receptors/metabolism, GABA, serotonin &amp; noradrenalin pathways mature at different rates</li> <li>Skeletal</li> <li>Ragio postnatal growth hiptel structure not evident until PND 14 to 21, and remain open into adulthood</li> <li>Long bone growth piate structure not evident until PND 14 to 21, and remain open into adulthood</li> <li>Completion of structural development; active period of growth and development</li> <li>Completion of structural development; active period of growth and/or</li> <li>Major period of functional and structural growth and development</li> <li>Completion of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of structural development; active period of growth and/or</li> <li>Major period of st</li></ul>	Immune	as a function of time and environment						
cerebral croke occurs posthalair         Maintum neuron ocuri and brain body weight at PND7, with extensive posthatal apoptosis, pruning and migration         Median chrosenant abit         • Megian chrosenant abit         • Orduction systems, optake receptors/metabolism, GABA, serotonin & noradrenalin pathways muter at different rates         Skeletal         • Ragid posthatal growth pitcupi adulthood         • Long bone growth pitce structure not evident until PND 14 to 21, and remain open into adulthood         • Completion of functional and structural growth and development         Completion of structural development; active period of growth and/or	Repro- ductive	<ul> <li>Period of decreased androgen production by Leydig cells during 3rd week postnatal necessary for expansion of germ cells and Sertoli cells</li> <li>Remaining reproductive changes and appearance of sexual dimorphism occur at onset of puberty (postnatal week 5 to 7)</li> </ul>						
Major period of functional and structural growth and development Completion of structural development; active period of growth and/or	Nervous	<ul> <li>Structural maturation of olfactory bulbs, relinaleye, cerebellum, hippocampus, and cerebral crotex occurs postnatally</li> <li>Maximum neuron count and brain body weight at PND7, with extensive postnatal apoptosis, pruning and migration</li> <li>Media not creasent at birth</li> </ul>						
Completion of structural development; active period of growth and/or	Skeletal	Rapid postnatal growth through adulthood     Long bone growth plate structure not evident until PND 14 to 21, and remain open into adulthood						
Completion of structural development; active period of growth and/or		Major period of functional and structural growth and develop	oment					
		Completion of structural development; active period of grow functional maturation	th and/or					
Slow continued growth or refinement of function; also can reflect a period of relative inactivity, as in prepubertal reproductive tissues		Slow continued growth or refinement of function; also can re	eflect a period	of				
Structurally and functionally fully mature				_				

	1					
System	General Considerations	Neonate (< 3 wks)	1st Solid Food (~ 3 wks)	Weaning (~ 8 wks)	Puberty (M ~ 5-8 mths) (F ~ 6-12 mths)	Adulthood (> ~ 12 mths)
cv	Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance)     Adoptime propagation and upper log shopped					
	<ul> <li>Adaptive myocardial and vascular changes</li> <li>Cardiac innervation continues development during approx. 2 to 4 months of age</li> <li>Significant increase in blood pressure and decrease in heart rate from week 1 to months of age</li> </ul>					
GI	Similarities in the stomach to that seen in human     At birth pastrointestinal tract is fully formed (functional development primarily between birth and weaning)					
Renai	Kidney is structurally and functionally immature at birth     Completion of nephrogenesis at approx. 2 weeks of age     Acid-base homeostasis develops postnatally     Concentrating ability develops prenatally					
Hepato- biliary	Hepatic structural maturation reached at approx. one week of age     Bile secretory function not fully mature at birth (at 4 to 6 weeks of age: 30 to 70 % adult values)	of				
Pulmonary	Considered acceptable model for the study of pulmonary toxicity in juvenile     population					
Immune	Immunologic tissues are largely structurally mature at or shortly after birth     Development of the immune system very similar to that seen in human, but plact     transfer of [go is poor     IgG transfer from dam primarily occurs during first 24 h postnatally via the colost     thymus undergoes rapid postnatal growth and reaches maximum size at 1 to 2					
Repro-	months of age					
Repro- ductive	Testicular descent incomplete at birth: occurring at 5 to 6 weeks of age     Males reach sexual maturity at approx. 7 to 8 months of age     Females reach sexual maturity at approx. 8 to 12 months of age					
Nervous	Rapid cognitive development through 12 to 18 weeks of age with critical developmental period for learning at approx. PND 18 to 28      Neonatal (minitiw) reflexes also disappear at approx. PND 28      Functional locomotor development occurs postnatally (standing approx. 3 weeks age with rapid progression through first month)	of				
Skeletal	<ul> <li>Long bone ossification primarily occurs postnatally with appearance of ossificatic centers between 1 to 10 weeks of age</li> <li>Most rapid long bone growth is complete by 5 months of age, with slower continu growth through puberty</li> </ul>	d				
	Major period of functional and structural growth and de	elopment				
	Completion of structural development; active period of functional maturation					
		an reflect a perio	d of			

System 0	General Considerations	Neonate (< 2 wks)	1= Solid Food (~ 2 wks)	Weaning (~ 4 wks)	Puberty (M ~ 3/4 mths, F ~ 4/5 mths)	Adulthood (> ~ 6 mths)
Integument •	<ul> <li>Critical neonatal function (barrier, water and thermoregulation, conductance, sensation)</li> </ul>					
cv ·	similarities in development to human     Critical neonatal physiologic transitions (pulmonary and systemic vascular					
1 C C C C C C C C C C C C C C C C C C C	<ul> <li>Adaptive myocardial and vascular changes</li> </ul>					
	Similarities in development to human					
	Maturity reached by approx. 4 weeks of age     Model for human stomach development					
Renal .	Nephron formation up to approx. 3 weeks after birth     Functional mature at approx. 3 months of age					
	Structurally and functionally immature at birth     Adult appearance at approx. 4 weeks of age and full function at 3 to 4 months of age					
· ·	Lungs are well developed at birth     Alveolization occurs over first 1 to 2 weeks of age and completed within 2 weeks of     age					
:	Very little function at birth     Anatomically full developed at approx. 4 weeks of age     Model for human immune development					
Repro- ductive	<ul> <li>Sexual maturity in males with approx. 3 to 4 months of age and in females with approx. 4 to 5 months of age</li> </ul>					
Nervous .	<ul> <li>Growth mainly in the late prenatal to early postnatal period</li> <li>Nenrous system complete by 6 months of age</li> <li>Brain development of the neonatal pig is similar to the human term neonate</li> <li>Neuromuscular system is more functionally mature at birth than in human</li> </ul>					
	<ul> <li>Rapid postnatal growth through adulthood; closure of the epiphysial growth plates at 18 months of age</li> <li>Full grown adults at approx. 24 months</li> </ul>					
Figure A	Major period of functional and structural growth and develop Completion of structural development; active period of grow functional maturation Slow continued growth or refinement of function; also can re relative inactivity, as in prepubertal reproductive tissues Structurally and functionally fully mature A.4: Age-dependent Development of	n and/or lect a period o		pig Org	an Syste	ems

Integrament       • Functional (harrier, water and thermoregulation, conductance, sensation) with hair       Image: CV       • Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance), resistance), resistance)       Image: CV       • Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance), resistance)       Image: CV       • Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance), resistance)       Image: CV       • Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance)       Image: CV
CV <ul> <li>Critical neonatal physiologic transitions (pulmonary and systemic vascular resistance)</li> <li>Adaptive myocardial and vascular changes</li> <li>Adaptive myocardial and vascular changes</li> <li>Myocardiocyte sepansion Through 3 months of age, then progressive growth</li> </ul> <li>GI</li> <li>Functional at birth, with adaptations especially over first year to accommodate shift in deliciomplexity and populate microborne</li> <li>Renal</li> <li>Nepbrogenesics complete at term birth Progressive increase in metabolic functionality, especially over first 3 to 8 months of age</li> <li>Hepato-</li> <li>Structurally well developed at birth, progressive increase in metabolic functionality, especially over first 3 to 8 months of age</li> <li>Pulmonary</li> <li>Structurally mature at birth with progressive growth</li> <li>Immune</li> <li>Progressive population of secondary immune tissues and development of memory as a function of time and environment</li>
Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Image: Adaptive myocardial and vascular changes       Image: Adaptive myocardial and vascular changes         Renal       • Nephrogenesis complete at term bith • Progressive increase in metabolic functionality, especially over first 3 to 6 months       Image: Adaptive myocardial changes         Pulmonary       • Structurally mature at bith with progressive growth       Image: Adaptive myocardial changes       Image: Adaptive myocardial changes         Immune       • Progressive population of secondary immune tissues and development of memory as a function of time and environment       Image: Adaptive myocardial changes       Image: Adaptive myocardial changes
GI <ul> <li>Functional at birth, with adoptations especially over first year to accommodate shift in diet/complexity and populate microbione</li> <li>Renal</li> <li>Nophrogenesis complete at term birth - Progressive increase in GPR and renal function over first 0 months of age</li> <li>Structurally work diventioned at the introduction over first 0 months of age</li> <li>Fundamentary over first 3 to 6 months</li> <li>Structurally mature at birth with progressive increase in metabolic functionality.</li> <li>Pulmonary</li> <li>Structurally mature at birth with progressive growth</li> <li>Immune</li> <li>Progressive population of secondary immune tissues and development of memory as a function of them and environment.</li> <li>Immune</li> <li>Structurally mature at birth with progressive and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth with progressive and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth with progressive and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth with progressive and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth and progressive and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth and progressive progression and development of memory</li> <li>A function of them and environment.</li> <li>Structurally mature at birth and environment.</li> <li>Struct</li></ul>
In distriction plexity and populate microbiome     Image: Constraint of the second secon
encode stream in the stream
biliary         especially over first 3 to 6 months           Pulmonary         • Structurally mature at birth with progressive growth         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and development of memory as a function of time and environment         Image: Constraint of the secondary immune tissues and the secondary immune tissues an
Pulmonary     • Structurally mature at birth with progressive growth     Immune       Immune     • Progressive population of secondary immune tissues and development of memory as a function of time and environment
Immune Progressive population of secondary immune tissues and development of memory as a function of time and environment
Endocrine   Most glands are well developed at birth and critical for growth
Zona reticularis of adrenal cortex expands at 3 to 6 months of age (adrenarche)     endocrine function of gonads expands at puberty
Repro- ductive         • Testes descended at birth, populated by germ cells, Serbic cells and Leydig cells           • Follicular development and artists begins at 3 to 6 months         • Solitosequent reproductive changes (menanche and spermarche) occur in at onset of puberty and continue unit advantioned
Nervous              • Defined sequential and progressive development into adulthood             • Postnatial apoptoris, pruning and migration most prominent before weaning             • Myelin and glip sceent at brit             • Neurotransmitter and conduction systems mature at variable rates (i.e.: serotonin             and noradrenain adfler)
Skeletal         • Growth plates present at birth           • Most rapid postnatal growth occurs prior to weaning, followed by slower growth until growth plates close during adulthood
Major period of functional and structural growth and development         Completion of structural development; active period of growth and/or functional maturation         Slow continued growth or refinement of function; also can reflect a period of relative inactivity, as in prepubertal reproductive tissues         Structurally and functionally fully mature

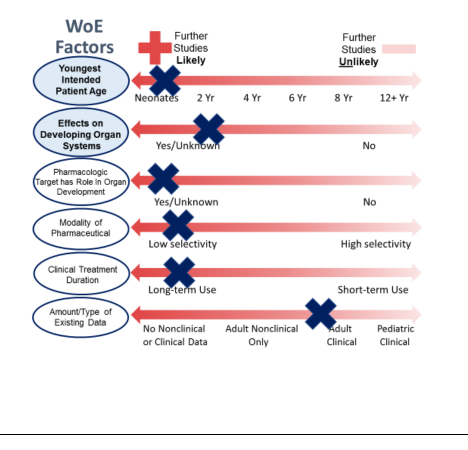


Species Advantages	Disadvantages	
Dog       Often used in general (adult) toxicology         Relatively large at birth       Relatively easy to handle         Litter size allows allocation of pups to different endpoin       Puppies can be separated from dams for a few hours         Breeding can be planned in advance       Breeding can be planned in advance	<ul> <li>Protracted development (~7-14 months to sexual maturity, ~18-24 months to skeletal maturity) with variable developmental milestones</li> <li>Altricial at birth (i.e. eyes do not open until ~2 weeks postnatally)</li> <li>Variable litter sizes and sex distribution can make it difficult to populate study with minimal bias (genetic/litter, sex distribution) across groups</li> <li>Limited historical background data, especially for nonstandard endpoints</li> <li>Substantial inter-individual variability in growth and development</li> <li>Seasonal breeder (supply &amp; study start over weeks or months)</li> <li>Not amenable to fostering</li> <li>Large body size requires comparably large amounts of test compound compared to rodents</li> </ul>	
Minipig/       • Many similar developmental milestones as humans         Pig       • Relatively large at birth         • Relatively easy to handle       • Breeding can be planned in advance         • Litter size allows allocation of piglets to different endpoints       • Amenable to cross fostering         • Relatively large litters usually allow balanced sex distribution       • Neonatal GI tract similar to human for orally administered drugs         • All routes of administration feasible (except inhalation); best model for dermal studies       • Short development (~6-9 months), relatively easy transport and housing compared to other large non- rodents	Less well established historical control data than dog or NHP toxicology species     Require colostrum for passive transfer of maternal Ig in perinatal period     Large body size requires comparably large amounts of test compound compared to rodents     IV and gavage administration can be challenging in very young piglets ed	
Species Advantages	Disadvantages	
NHP (cynomo lgus; rhesus and marmos       • Many similar developmental milestones as humans         • Neonates/infants similar to human for GI tract, immune system, cardiovascular, renal and special sense (eye, ear) development         • Macaque infants are relatively large at birth         • Extensive reference data from birth available         • Often used for (adult) general and reproductive toxicology (e.g., ePPND), especially for biopharmaceuticals         • Often the most pharmacologically relevant animal mode for highly targeted therapies         Rabbit       • Compressed development (~5-6 months) and small boo	<ul> <li>study to cover all developmental phases not practical</li> <li>Single offspring for macaques with high inter-individual variability in growth and development</li> <li>Marmosets typically have twins and require both maternal and paternal care in preveaning phase; offspring are relatively small</li> <li>Offspring highly dependent on maternal care over first month (minimal procedural intervention recommended; pre-weaning manipulation &amp; dosing feasible with risk of maternal rejection), and are cohoused with dam for first 3-6 months; with shipping and quarantine requirements it is rarely feasible to initiate studies in juvenile monkeys &lt; 9 months of age</li> <li>Neonatal NHP are precocious relative to human neonates in terms of musculoskeletal, CNS, endocrine and respiratory system</li> <li>Cannot synchronize breeding (supply &amp; study start over weeks or months for seasonal breeders such as rhesus)</li> <li>Ethical reservations (need strong rationale to justify use of juvenile NHP for toxicity testing)</li> <li>Developmental milestones less well established than other</li> </ul>	
Compressed development (~>-o months) and small box size requiring comparably low amount of test material Relatively easy to handle     Often used for reproductive toxicology; also can be use for ocular administration, evaluation of bone growth Litter size allows allocation of kits to different endpoints	nonrodent species     Not routinely used / well accepted in (adult) general toxicology     Handling young offspring can provoke cannibalism or rejection by     the mother	

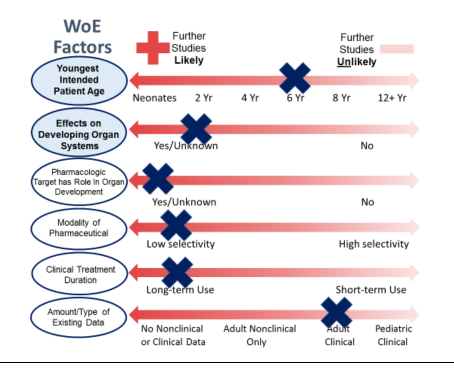
Other Species         Other species could be considered for cause when pharmacologically and toxicologically relevant. Examples of alternative mammalian test systems include the hamster, guinea pig, tree shrew, ferret, cat, sheep and goat. Advantages tend to be species and program specific, but often reflect use of that species in genetic or disease models, or when there is data supporting interpretation and translability of specific endpoints. <ul></ul>	

	clinical and nonclinical data including repeated dose toxicity data.	
	None of these data suggest a safety concern in a developing organ for	
	the intended pediatric population of adolescents (12 years and	
	above), for a one-month duration of clinical treatment. The WoE	
	analysis indicates that no additional nonclinical investigations are	
	needed.	
10. APPENDIX B: CASE STUDIES APPLYING THE WEIGHT OF EVIDENCE APPROACH	Were Factors       Further Studies       Further Studies         Vounget Intende       Likely       Unlikely         Intende       No       No         Patient Age       Neonates       2 Yr       4 Yr       6 Yr       8 Yr       124 Yr         Plarmacologic Trate has Role in Organ Development       Yes/Unknown       No       No       No         Modality of Pharmaceutical       Low selectivity       High selectivity       High selectivity         Clinical Treatment Duration       No Nonclinical Adult Nonclinical Only       Adult Pediatric Clinical Clinical	
	STUDIES APPLYING THE WEIGHT OF EVIDENCE	10. APPENDIX B: CASE STUDIES APPLYING THE WEIGHT OF EVIDENCE APPROACH  H H H H H H H H H H H H H H H H H H

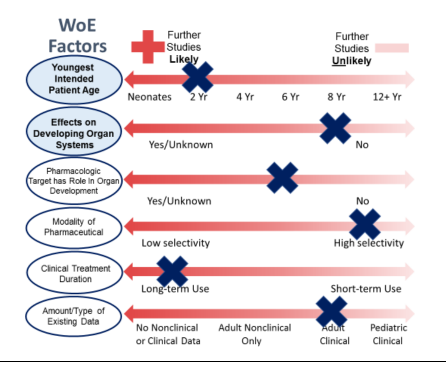
B. A small molecule with a novel mode of action intended for chronic use starting in neonates or infants has limited Phase 1 clinical and nonclinical safety data with no significant safety concerns identified. There are potential pharmacologic effects on developing organ systems. The WoE analysis indicates further nonclinical investigation, such as a core JAS with additional endpoints based on the targeted developing organ systems, would be useful.



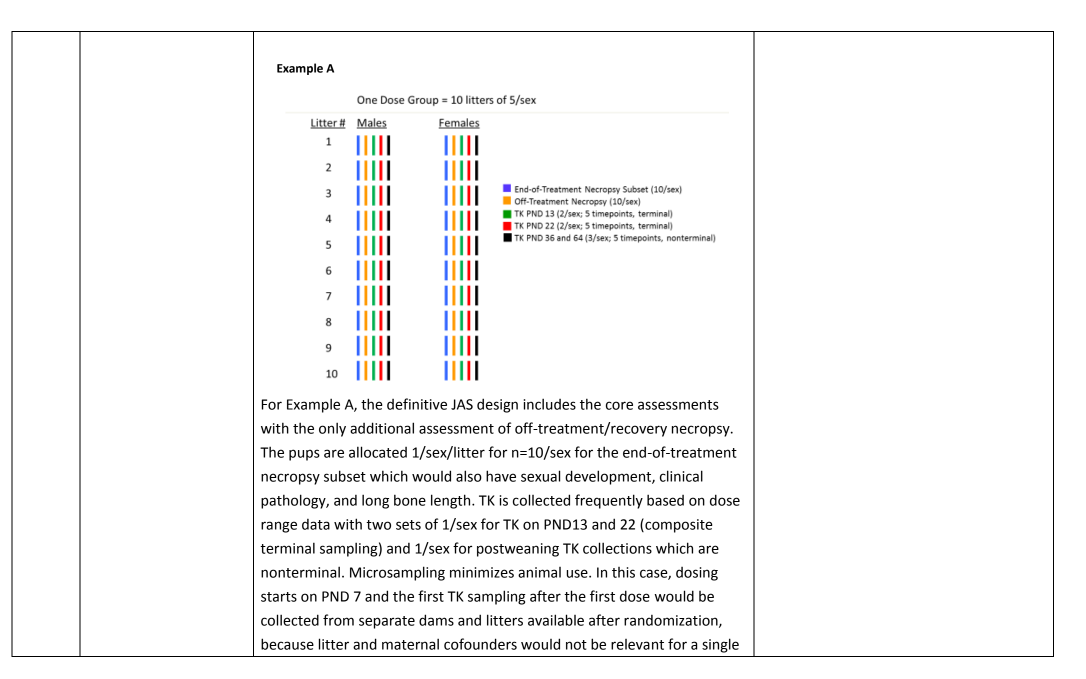
C. A small molecule with known pharmacology with a well characterized critical role in CNS development intended for chronic use in children (6 years and above) has nonclinical and adult clinical data. The concern for a potential effect on the developing CNS cannot be addressed clinically by monitoring and management. Existing data adequately addresses other developing systems. The WoE analysis warrants a post-weaning JAS study design that includes core endpoints and additional endpoints limited to CNS, including detailed clinical observations, behavioral assessments, a learning and memory evaluation, and expanded neuropathological examinations.

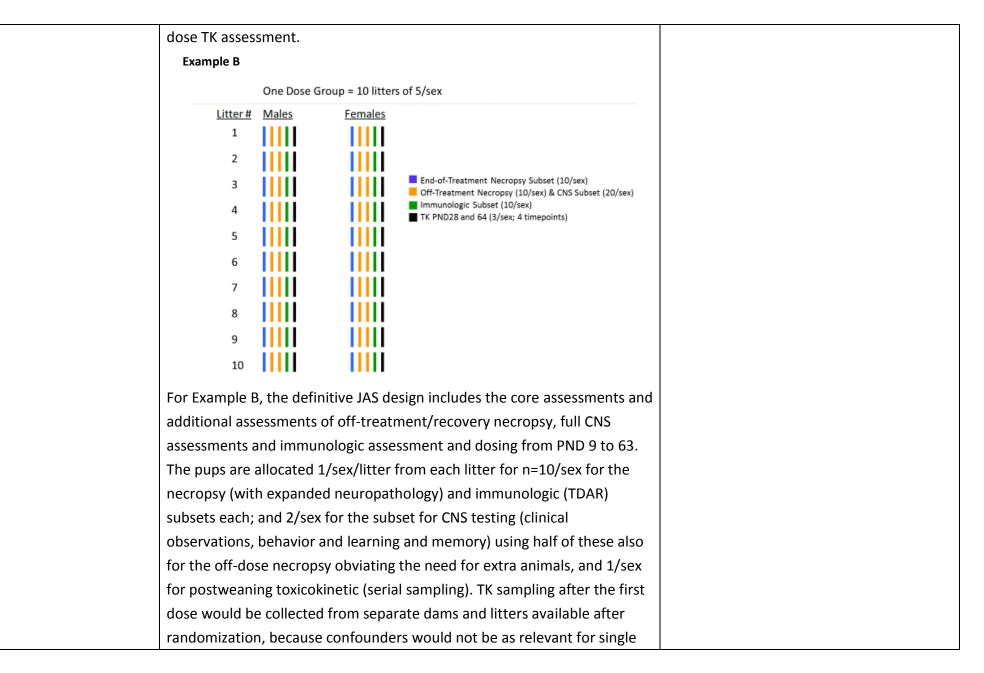


D. A monoclonal antibody targets a soluble cytokine and is intended for chronic pediatric use in rheumatologic and allergic diseases (>2 years old). The only findings are reversible decreased serum Ig and occasional injection site reactions (in both animals and adult patients). In a monkey ePPND study, offspring exposure was comparable to dams through PND 28 and decreased pharmaceutical Ig levels was detected on PND 28 and 56 postnatally.
 T-cell-dependent antibody response (TDAR) results were similar to controls (between 3-6 months postnatally). The WoE analysis does not warrant a JAS.



806-850		Natural Litters + Whole Litter Group Assignment + Inter-Litter
		Endpoint Subset Assignment
		Initiation of a JAS during the preweaning phase presents a unique
		situation and should be designed to reduce potential confounders related
		to genetics, maternal care, and littermates. This is achieved by how the
		litters are constructed in combination with how they are assigned to dose
		groups, and then to subsets of endpoints. In this approach, the offspring
		stay with their natural mother and are culled to the desired litter size with
		a balanced sex ratio. When necessary to minimize the required number of
		litters to supply the study, a very small percentage of pups are fostered to
	11. APPENDIX C:	other litters. Here, Wistar Han rat litters are culled to 10 offspring per
	EXAMPLE OF AN	litter composed of 5 males and 5 females (the mean natural litter size is
	APPROACH TO RODENT	~11). The whole litter is then assigned to the same dose group with 10
	PREWEANING LITTER	litters each assigned to each dose group. Offspring are arbitrarily assigned
	ALLOCATION	to subsets for specific endpoints in an inter-litter fashion, i.e., as one male
		or female from each litter in a dose group to the specific endpoints. The
		advantage of the whole litter group assignment is the littermates receive
		the same dose level so there is a low risk of cross contamination and
		confounding variables of high dose and control offspring competing for
		suckling position and time. Also, keeping the pups with genetic dams and
		assigning the endpoints in an inter-litter fashion ensures genetic, maternal
		care and littermate influences are distributed evenly.





dose TK assessment.		
---------------------	--	--