

DRAFT Guidance on the Biocidal Products Regulation

Volume III: Human health
Part A: Information requirements
DRAFT Version 2, 31 May 2021

ABC

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Title...**Reference:** xxxx-xxxx-xxxx**ISBN:** xxxx-xxxx-xxxx**Cat. Number:** ED-xxxxxx-xxxx-xxxx**DOI:** xx.xxxx/xxxxxx**Publ.date:** Month 20xx**Language:** EN

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List of abbreviations

| Standard term / Abbreviation | Explanation |
|------------------------------|--|
| °C | Degree(s) Celsius (centigrade) |
| ADME | Absorption, distribution, metabolism and excretion |
| ADI | Acceptable daily intake |
| ADS | Additional data set |
| AEL | Acceptable exposure level, overall systemic limit value for the human population |
| ARfD | Acute Reference Dose |
| AUC | Area under the curve |
| BCF | Bioconcentration factor |
| BPD | Biocidal Products Directive. Directive 98/8/EC of the European Parliament and of the Council on the placing on the market of biocidal products |
| BPR | Biocidal Products Regulation. Regulation (EU) No 528/2012 of the European Parliament and of the Council concerning the making available on the market and use of biocidal products |
| Cat | Category |
| CDS | Core data set |
| CLH | Harmonised classification and labelling |
| CLP (Regulation) | Classification, Labelling and Packaging Regulation. Regulation (EC) No 1272/2008 of the European Parliament and of the Council on Classification, Labelling and Packaging of substances and mixtures |
| CWM | Cincinnati water maze |
| DG | European Commission Directorate General |
| DG SANTE | European Commission Directorate-General for Health and Food Safety |
| DNA | Deoxyribonucleic acid |
| DNT | Developmental Neurotoxicity |
| EATS | Oestrogen, androgen, thyroid, steroidogenesis (ED modalities) |
| EC | European Communities or European Commission |
| EC method | Test Method as listed in the Test Methods Regulation |
| ECHA | European Chemicals Agency |
| ED | Endocrine disruption; endocrine disruptor |
| EFSA | European Food Safety Agency |
| EU | European Union |
| FISH | Fluorescence in-situ hybridisation |
| FOB | Functional observation battery |
| g | Gram(s) |

| | |
|--------|---|
| GC | Gas chromatography |
| GIVIMP | Good <i>in vitro</i> Method Practices |
| GLP | Good laboratory practice |
| h | Hour(s) |
| HPLC | High performance (or pressure) liquid chromatography |
| IATA | Integrated Approach on Testing and Assessment |
| IPCS | The WHO International Programme on Chemical Safety |
| ISBN | International standard book number |
| ITS | Integrated testing strategy |
| IUCLID | International Uniform Chemical Information Database |
| kg | Kilogram(s) |
| LD50 | Lethal dose for 50% of the group of tested animals |
| LLNA | Murine local lymph node assay |
| mg | Milligram(s) |
| MMAD | Mass median aerodynamic diameter |
| mol | Mole(s) |
| MWM | Morris water maze |
| MRL | Maximum residue limit |
| MS | Mass spectrometry |
| MSCA | Member State competent authority |
| NAFTA | North American Free Trade Agreement |
| nm | Nanometre(s) |
| NMR | Nuclear magnetic resonance |
| NOAEL | No observed adverse effect level |
| OECD | Organisation for Economic Cooperation and Development |
| OPPTS | Office of Prevention, Pesticides, and Toxic Substances (U.S.-EPA) |
| Pa | Pascal(s) |
| PBK | Physiologically based kinetics |
| PBPK | Physiologically based pharmaco(toxico)-kinetics |
| pH | pH-value, negative decadic logarithm of the hydrogen ion concentration |
| PND | Postnatal day |
| PPI | Pre-pulse inhibition |
| PPPR | Plant Protection Products Regulation. Regulation (EC) No 1107/2009 of the European Parliament and of the Council of concerning the placing of plant protection products on the market |
| PT | Product-type |

| | |
|-------------------------|--|
| (Q)SAR | (Quantitative) structure activity relationship |
| RAM | Radial arm maze |
| REACH | Regulation (EC) No 1907/2006 of the European Parliament and of the Council concerning the Registration, Evaluation, Authorisation and Restriction of Chemicals |
| s | Second(s) |
| Test Methods Regulation | Regulation (EC) No 440/2008 laying down test methods pursuant to the REACH Regulation |
| UV | Ultraviolet |
| WHO | World Health Organisation |

1. Dossier Requirements for Active Substances

Toxicological profile for human and animal including metabolism

Considerations before initiating testing

Before testing is initiated all available information should be scrutinised for evidence that may indicate severe effects, serious specific system or target organ toxicity (e.g. neurotoxicity or immunotoxicity), delayed effects or cumulative toxicity. Consideration should also be given to tests already performed/submitted for the purpose of other regulatory programmes. All available information on toxicity should be taken into account when choosing the dose range for a new study. If there is concern that an effect is not adequately covered by existing OECD Test Guidelines, specialised study protocols may be used. Whenever deviating from OECD Test Guidelines, a justification is required. These specialised study protocols should be designed on a case-by-case basis in order to enable an adequate characterisation of these hazards, including the dose-response, threshold for the toxic effect and an understanding of the nature of the toxic effects. Where a need is identified for a modification in the study protocol to cover specific needs, this will be done in consultation with the evaluating Member State.

The endpoints that need to be addressed for the purpose of the BPR are interlinked and in certain cases sequential testing strategy is needed to decide which tests need to be performed and in which order. This is due to the impact that the results from one study can have on the classification and labelling and the risk management measures, which can make the requirement for testing of other endpoints redundant.

For each toxicological endpoint and the respective information requirements, all available information has to be collected and evaluated before concluding on the need to conduct further testing using integrated testing strategies (ITS) where relevant.

The Test Methods Regulation is regularly updated to follow the approval of new OECD Test Guidelines. In accordance with Point 5 of BPR Annex II, the latest version of an adopted test guideline should always be used when generating new data, independently from whether it is published by the EU or OECD.

The Test Methods Regulation is regularly updated to follow the approval of new OECD Test Guidelines. In accordance with Point 5 of BPR Annex II, the latest version of an adopted test guideline should always be used when generating new data, independently from whether it is published by the EU or OECD. In addition to the test methods mentioned for each data requirement, new OECD validated tests for genotoxicity should be taken into account once available in deciding the test strategy.

General considerations for animal data reporting

Where submitted, historical control data should be from the same species and strain, maintained under similar conditions in the same laboratory and should be from contemporaneous studies. Additional historical control data not fulfilling these conditions, or from other laboratories may be reported separately as supplementary information.

The information on historical control data provided should include:

(a) identification of species and strain, name of the supplier, and specific identification if the supplier has more than one geographical location;

(b) name of the laboratory and the dates when the study was performed;

- 1 (c) description of the general conditions under which animals were maintained, including the
2 type or brand of diet and, where possible, the amount consumed;
- 3 (d) approximate age, in days, and weight of the control animals at the beginning of the study
4 and at the time of sacrifice or death;
- 5 (e) description of the control group mortality pattern observed during or at the end of the
6 study, and other pertinent observations (such as diseases, infections);
- 7 (f) name of the laboratory and the examining scientists responsible for gathering and
8 interpreting the pathological data from the study;
- 9 (g) for carcinogenicity studies: a statement of the nature of the tumours that may have been
10 combined to produce any of the incidence data.

11 The historical control data should be presented on a study by study basis giving absolute values
12 plus percentage and relative or transformed values where these are helpful in the evaluation. If
13 combined or summary data are submitted, these should contain information on the range of
14 values, the mean, median and, if applicable, standard deviation.

15 The doses tested should be selected on the basis of the results of short-term testing and, where
16 available at the time of planning the studies, on the basis of metabolism and toxicokinetic data.
17 Dose selection should consider toxicokinetic data such as saturation of absorption measured by
18 systemic availability of active substance and/or metabolites.

19 Doses causing excessive toxicity should not be considered relevant to evaluations. Determination
20 of blood concentration of the active substance (for example around Tmax) should be considered
21 in long-term repeated dose toxicity studies.

22 **1.1. Skin corrosion or irritation**

23 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.1 Skin corrosion or irritation</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) skin corrosion, <i>in vitro</i> testing;</p> <p>(c) skin irritation, <i>in vitro</i> testing;</p> <p>(d) skin corrosion or irritation, <i>in vivo</i> testing</p> | <p>The study/ies in column 1 do(es) not need to be conducted if:</p> <ul style="list-style-type: none"> - the available information indicates that the substance meets the criteria for classification for skin corrosion or irritation, - the substance is a strong acid (pH ≤ 2,0) or base (pH ≥ 11,5), - the substance is spontaneously flammable in air or in contact with water or moisture at room temperature, - the substance meets the classification criteria for acute toxicity (Category 1) by the dermal route, or - an acute toxicity study by the dermal route provides conclusive evidence on skin corrosion or irritation adequate for classification. <p>If results from one of the two studies listed in point (b) or point (c) in column 1 of this row already allow</p> |

| | |
|--|--|
| | <p>conclusive decision on the classification of a substance or on the absence of skin irritation potential, the second study does not need to be conducted</p> <p>An <i>in vivo</i> study for skin corrosion or irritation shall be considered only if the <i>in vitro</i> studies listed in points (b) and (c) in column 1 of this row are not applicable, or the results of these studies are not adequate for classification and risk assessment</p> <p><i>In vivo</i> studies for skin corrosion or irritation that were carried out or initiated before 15 April 2022 shall be considered appropriate to address this information requirement</p> |
|--|--|

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2 For skin corrosion/irritation, the information must be sufficient to conclude on the classification
3 of the substance, i.e. that the criteria are met for classifying as skin corrosion (Cat. 1 of CLP) or
4 as skin irritation (Cat. 2 of CLP), or that no classification is warranted.

5 The information below provides brief guidance for the assessment of skin corrosion or irritation.
6 To support this, please refer to chapter R.7.2.6 of *REACH Guidance on Information Requirements*
7 *and Chemical Safety Assessment Chapter R.7a* where detailed information is given on the
8 different steps/tiers, as well as on the OECD Guidance Document on an Integrated Approach on
9 Testing and Assessment (IATA) for skin corrosion/irritation (2014).

10 The testing and assessment strategy aims at identifying skin corrosion/irritation by using all the
11 information available. A basic principle of the strategy is that the results of one study or
12 information source are evaluated before another study is initiated. The strategy seeks to ensure
13 that the data requirements are met in the most efficient and humane manner so that animal
14 usage and costs are minimised.

15 **Tier a) assessment of the available human, animal and non-animal data**

16 In this Tier, all available information (including physico-chemical properties) should be evaluated
17 before undertaking any new testing and to avoid, as far as possible, *in vivo* testing of corrosive
18 and severely irritating substances. In case new testing is needed, *in vitro* tests must be
19 performed first, and it should be assessed whether *in vivo* testing can be completely avoided.

20 Further guidance regarding the assessment of existing information (physicochemical properties,
21 grouping, (Q)SARs and expert systems, *in vitro* data, human data and animal data) is available
22 within the *Guidance on the Application of the CLP Criteria, BPR Guidance Volume III Parts B+C*
23 *and REACH Guidance on Information Requirements and Chemical Safety Assessment Chapter*
24 *R.7a*.

25 In principle information requirements for skin irritation/corrosion do not apply in cases when:

26 1. The available information already indicates that the criteria are met for classification as
27 corrosive to the skin or as a skin irritant.

28 2. The substance is a strong acid (pH < 2) or base (pH > 11.5).

29 3. The substance is spontaneously flammable in air or in contact with water or moisture at
30 room temperature.

31 4. The substance meets the classification criteria for acute toxicity (Category 1) by the
32 dermal route.

1 5. An acute toxicity study by the dermal route provides conclusive evidence on skin
2 corrosion or irritation adequate for classification.

3 If a good quality *in vivo* skin irritation study is already available i.e. study was carried out or
4 initiated before 15 April 2022, it can be used to fulfil the standard information requirement.

5 For existing animal data, the use of methods other than those that are specified in the Annex to
6 the EU Test Methods Regulation or the corresponding OECD methods may be accepted on a
7 case-by-case basis. If the test was performed in other species than the rabbit, evaluation must
8 be made with caution. Such information may be available e.g. from dermal toxicity studies in
9 the rat or sensitisation studies in guinea pigs. One must note that the skin of the rat is less
10 sensitive compared to rabbit skin, and the guinea pig skin is even less sensitive. Much lower
11 exposures are employed in dermal toxicity testing and, in general, the scoring of dermal effects
12 is performed less accurately. The results of dermal toxicity testing in rats or skin sensitisation
13 tests in guinea pigs will not be adequate for classification for skin irritation/corrosion, unless the
14 results indicate skin corrosivity that warrants classification as Skin Corrosive Category 1. In any
15 other case, such information must be used in a Weight of Evidence assessment.

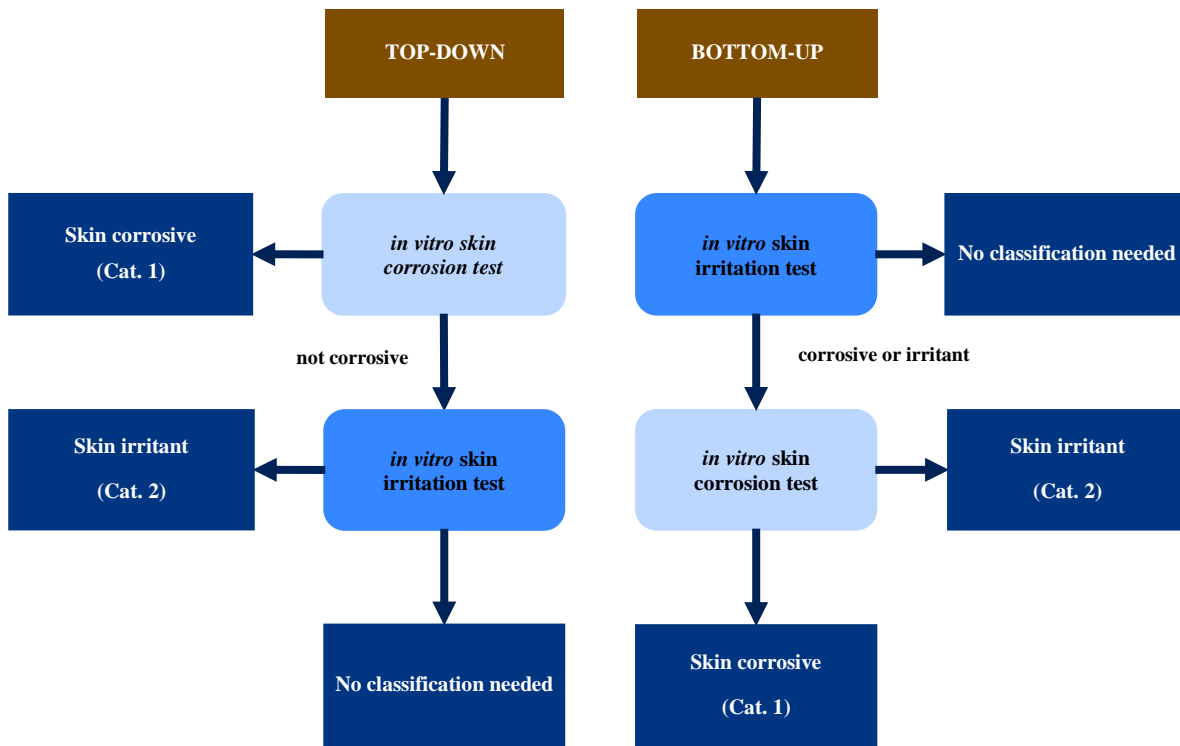
16 Existing human data include historical data that should be taken into account when evaluating
17 intrinsic hazards of substances. New testing in humans for hazard identification purposes is not
18 acceptable for ethical reasons. Existing data can be obtained from case reports, poison
19 information centres, medical clinics, occupational experience, epidemiological studies and
20 volunteer studies. Their quality and relevance for hazard assessment should be critically
21 reviewed. However, in general, human data can be used to determine a corrosive or irritating
22 potential of a substance. Good quality and relevant human data have precedence over other
23 data. However, absence of incidence in humans does not necessarily overrule positive, good
24 quality *in vitro* data or existing animal data.

25 **Considerations before performing further testing**

26 If after the analysis in Tier a) further testing is needed to assess the potential for skin irritation
27 or skin corrosion, the test methods mentioned below should be used. Where new testing is
28 needed, please see also the general information under *Considerations before initiating testing* in
29 chapter 1.

30 The tests will provide information on the degree and nature of the effects on skin especially with
31 regard to the reversibility of responses.

32 New *in vitro* testing should be performed following a top-down or bottom-up approach, based
33 on presumed properties (Figure 1). The top-down approach should be used when the available
34 information suggests that the substance may be irritant or corrosive to the skin. The bottom-up
35 approach should be followed when all available information suggests that the substance may not
36 be irritant to the skin.



1

2 **Figure 1. Schematic presentation of top-down and bottom-up approaches for skin**
 3 **corrosion/irritation.**

4 After following this scheme, no new *in vivo* testing is normally necessary unless:

- 5 a) the available *in vitro* methods cannot be used due to substance specific limitations, or
 6 b) the results of the *in vitro* test(s) performed do not enable a clear conclusion on
 7 classification and/or are insufficient for appropriate risk assessment.

8 Before performing any *in vivo* studies, it is necessary to identify any skin corrosion/irritation
 9 studies that may already be available, even if not fully equivalent to an OECD TG or an EU test
 10 method. If there are several studies and the results from such studies are consistent, they may
 11 together provide sufficient information on the skin corrosion/irritation potential of the substance.

12 Tier b) skin corrosion, *in vitro* testing

13 If after the analysis in Tier a) above, further testing is needed to assess the potential for skin
 14 corrosion, one of the test methods listed in Table 1 should be used. Before testing, consider
 15 whether corrosion or irritation would not be expected, in which case the bottom-up approach
 16 could be considered instead (see Figure 1).

17 **Table 1. In vitro test methods for skin corrosion:**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|--|--------------------------------------|--|
| Transcutaneous electrical resistance tests | B.40 / TG 430 | Cat. 1 or non-corrosive |

| | | |
|---------------------------|------------------|--|
| Human skin model test(s)* | B.40bis / TG 431 | Cat. 1, 1A, 1B/1C or non-corrosive |
| Membrane barrier test | B.65 / TG 435 | Cat. 1, 1A, 1B and 1C or non-corrosive |

1

2 The limitations and the scope of a given test method within a test guideline should be taken into
3 account when selecting the most appropriate *in vitro* method for a particular substance and when
4 interpreting the test results. Where new testing is needed, please see also the general
5 information under *Considerations before initiating testing* in chapter 1.

6 **Tier c) skin irritation, *in vitro* testing**

7 To examine the potential for skin irritation, the method(s) listed in the Table 2 below should be
8 used.

9 **Table 2. *In vitro* test methods for skin irritation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|--|--------------------------------------|--|
| Reconstructed human epidermis test(s)* | B.46 / TG 439 | Cat. 1/Cat. 2 or not classified |

10 * The test guideline contains multiple methods/protocols using reconstructed human epidermis.

11 The limitations and the scope of a given test method within a test guideline should be taken into
12 account when selecting the most appropriate *in vitro* method for a particular substance and when
13 interpreting the test results. Where new testing is needed, please see also the general
14 information under *Considerations before initiating testing* in chapter 1.

15 **Tier d) skin corrosion or irritation, *in vivo* testing**

16 *In vivo* testing in Tier d) is required only as a last resort if the information assessed in the Tiers
17 (a-c) above are not sufficient for concluding on the classification and/or for performing a risk
18 assessment. In such a case, an *in vivo* skin irritation study should be performed using the test
19 method listed in Table 3.

20 **Table 3. *In vivo* test methods for skin corrosion/irritation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|---|--------------------------------------|--|
| Acute Dermal Irritation/Corrosion test (<i>in vivo</i>) | B.4 / OECD TG 404 | Cat. 1, Cat. 2 or not classified |

21

22 In interpreting *in vivo* information, particular attention should be given to the persistence of
23 irritation effects, even those which do not lead to classification. Effects such as erythema,
24 oedema, fissuring, scaling, desquamation, hyperplasia and opacity, which do not reverse within
25 the test period may indicate that a substance will cause persistent damage to the human skin.

26 **1.2. Serious eye damage or eye irritation**

1 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|--|
| <p>8.2 Serious eye damage or eye irritation</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) serious eye damage or eye irritation, <i>in vitro</i> testing;</p> <p>(c) serious eye damage or eye irritation, <i>in vivo</i> testing</p> | <p>The study/ies in column 1 do(es) not need to be conducted if:</p> <ul style="list-style-type: none"> –the available information indicates that the substance meets the criteria for classification for eye irritation or causing serious damage to eyes, – the substance is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$), – the substance is spontaneously flammable in air or in contact with water or moisture at room temperature, or –the substance meets the classification criteria for skin corrosion leading to classification of the substance as “serious eye damage” (category 1). <p>If results from a first <i>in vitro</i> study do not allow a conclusive decision on the classification of the substance or on the absence of eye irritation potential (an)other(s) <i>in vitro</i> study(ies) for this endpoint shall be considered.</p> <p>An <i>in vivo</i> study for serious eye damage or eye irritation shall be considered only if the <i>in vitro</i> study(ies) listed in point (b) in column 1 of this row are not applicable, or the results obtained from these studies are not adequate for classification and risk assessment <i>In vivo</i> studies for serious eye damage or eye irritation that were carried out or initiated before 15 April 2022 shall be considered appropriate to address this information requirement</p> |

2
3 For serious eye damage or eye irritation, the information must be sufficient to conclude on the
4 classification of the substance, i.e. that the criteria are met for classifying as serious eye damage
5 (Cat 1 of CLP) or as eye irritation (Cat 2 of CLP), or that no classification is warranted.

6 The information below provides brief guidance for the assessment of serious eye damage or eye
7 irritation. To support this, please refer to chapter R.7.2.11 of REACH Guidance on Information
8 Requirements and Chemical Safety Assessment Chapter R.7a where detailed information is given
9 on the different steps/tiers, as well as in the OECD 2019 *Guidance Document on Integrated*
10 *Approaches to Testing and Assessment (IATA) for Serious Eye Damage and Eye Irritation*,
11 Second Edition.

12 The testing and assessment strategy is aimed at the identification of serious eye damage/eye
13 irritation by using different elements where appropriate, depending on the information available.
14 A basic principle of the strategy is that the results of one study or from an information source
15 are evaluated before another study is initiated. The strategy seeks to ensure that the data
16 requirements are met in the most efficient and humane manner so that animal usage and costs
17 are minimised.

18 **Tier a) Assessment of the available human, animal and non-animal data**

19 In this Tier, all available information (including physico-chemical properties) must be evaluated
20 before undertaking any new testing and to avoid, as far as possible, *in vivo* testing of corrosive

1 and severely irritating substances. In case new testing is needed, *in vitro* tests must be
2 performed first, and it should be assessed whether *in vivo* testing can be completely avoided.

3 Further guidance regarding the assessment of existing information (physicochemical properties,
4 grouping, (Q)SARs and expert systems, *in vitro* data; human data and animal data) is available
5 within the *Guidance on the Application of the CLP Criteria, BPR Guidance Volume III Parts B+C*
6 and *REACH Guidance on Information Requirements and Chemical Safety Assessment Chapter*
7 *R.7a*.

8 In principle information requirements for eye irritation do not apply in cases when:

- 9 1. The available information indicates that the substance meets the criteria for classification
10 for eye irritation or causing serious damage to eyes,
- 11 2. The substance is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$),
- 12 3. The substance is spontaneously flammable in air or in contact with water or moisture at
13 room temperature, or
- 14 4. The substance meets the classification criteria for skin corrosion leading to classification
15 of the substance as 'serious eye damage' (category 1).

16 If a good quality *in vivo* eye irritation study is already available i.e. study was carried out or
17 initiated before 15 April 2022, it can be used to fulfil the standard information requirement.

18 For existing animal data, the use of methods other than those specified in the Annex to the EU
19 Test Methods Regulation, or corresponding OECD methods may be accepted on a case-by-case
20 basis. To support this, please refer to the ECHA Guidance Vol III Parts B+C, and section 1.5.5.1.2
21 "Testing data for irritation/corrosion (skin and eye)" of REACH Guidance on Information
22 Requirements and Chemical Safety Assessment Chapter R.7a.

23 Existing human data include historical data that should be taken into account when evaluating
24 intrinsic hazards of substances. New testing in humans for hazard identification purposes is not
25 acceptable for ethical reasons. Existing data can be obtained from case reports, poison
26 information centres, medical clinics, occupational experience, epidemiological studies and
27 volunteer studies. Their quality and relevance for hazard assessment should be critically
28 reviewed. However, in general, human data can be used to determine a corrosive or irritating
29 potential of a substance. Good quality and relevant human data have precedence over other
30 data. However, absence of incidence in humans does not necessarily overrule positive, good
31 quality *in vitro* data or existing animal data.

32 **Considerations before further testing**

33 If after the analysis in Tier a) further testing is needed to assess the potential for serious eye
34 damage or eye irritation, the test methods listed in Tables 4 and 5 should be used. Where new
35 testing is needed, please see also the general information under *Considerations before initiating*
36 *testing* in chapter 1.

37 New *in vitro* testing should be performed following a top-down or bottom-up approach, based
38 on presumed properties. The top-down approach starts with an *in vitro* test able to identify
39 substances causing serious eye damage (Cat 1 of CLP). This approach should be used when all
40 available information and the Weight-of-Evidence assessment indicate a high *a-priori* probability
41 of the substance being seriously damaging to the eye. The bottom-up approach starts with an
42 *in vitro* test able to identify substances not requiring classification for serious eye damage/eye
43 irritation. This approach should be followed when all available information and the Weight-of-
44 Evidence assessment indicate a high *a-priori* probability of the substance being non-irritant to
45 the eyes.

1 After following this scheme, no new *in vivo* testing is normally necessary unless:

- 2 a) the available *in vitro* methods cannot be used due to substance specific limitations, or
3 b) the results of the *in vitro* test(s) performed do not enable a clear conclusion on
4 classification and/or are insufficient for appropriate risk assessment.

5 Before performing any *in vivo* studies, it is necessary to identify any serious eye damage/eye
6 irritation studies that may already be available, even if not fully equivalent to an OECD TG or an
7 EU test method. If there are several studies and the results from such studies are consistent,
8 they may together provide sufficient information on the serious eye damage/eye irritation
9 potential of the substance.

10 **Tier b) Serious eye damage or eye irritation, *in vitro* testing**

11 If after the analysis in Tier a) above further testing is needed to assess the potential for serious
12 eye damage or eye irritation, the test methods in Table 4 below should be used. Where new
13 testing is needed, please see also the general information under *Considerations before initiating*
14 *testing* in chapter 1.

15 **Table 4: *In vitro* test methods for serious eye damage/eye irritation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|----------------|--------------------------------------|--|
| BCOP | B.47 / OECD TG 437 | Cat. 1 or not classified |
| ICE | B.48 / OECD TG 438 | Cat. 1 or not classified |
| STE | B.68 / OECD TG 491 | Cat. 1 or not classified |
| Macromolecular | N.A. / OECD TG 496 | Cat. 1 or not classified |
| FL | B.61 / OECD TG 460 | Cat. 1 |
| RhCE | B. 69 / OECD TG 492 | Not classified |
| Vitrigel | N.A. / OECD TG 494 | Not classified |

16 **Abbreviations:** BCOP = Bovine Corneal Opacity and Permeability; FL = Fluorescein Leakage; ICE = Isolated Chicken
17 Eye; N.A. = not available; RhCE = Reconstructed human Cornea-like Epithelium Test Method; STE = Short-Time
18 Exposure.

19 The limitations and the scope of a given test method within a test guideline should be taken into
20 account when selecting the most appropriate method for a particular substance and when
21 interpreting the test results. The latest version of an adopted test guideline should always be
22 used when generating new data, independently of whether it is published by the EU or OECD.

23 The test methods mentioned above are suitable either for the direct identification of effects
24 leading to serious eye damage (Cat. 1 of CLP) or substances not requiring classification under
25 CLP. Currently there are no internationally adopted methods available for the direct identification
26 of effects leading to eye irritation (Cat. 2 of CLP).

27 If the results of one *in vitro* assay do not allow concluding on the classification of the substance
28 or on the absence of eye irritation potential, additional *in vitro* studies may need to be performed.

1 **Tier c) Serious eye damage or eye irritation, *in vivo* testing**

2 *In vivo* testing is required only as a last resort if the information assessed in the Tiers a) and b)
3 above are not sufficient for concluding on the classification and/or for performing a risk
4 assessment. In such a case, an *in vivo* eye irritation study should be performed using the test
5 method in Table 5.

6 **Table 5. *in vivo* test methods for serious eye damage/eye irritation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|--|--------------------------------------|--|
| Acute Eye Irritation/Corrosion test (<i>in vivo</i>) | B.5 / OECD TG 405 | Cat. 1, Cat. 2 or not classified |

7

8 **1.3. Skin sensitisation**

9 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|--|
| <p>8.3 Skin sensitisation</p> <p>The information shall allow to conclude whether the substance is a skin sensitizer and whether it can be presumed to have the potential to produce significant sensitisation in humans (Category 1A). The information should be sufficient to perform a risk assessment where required</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) skin sensitisation, <i>in vitro</i> testing. Information from <i>in vitro</i> or <i>in chemico</i> test method(s) referred to in point 5 of the introductory part of this Annex and addressing each of the following key events of skin sensitisation:</p> <p>(i) molecular interaction with skin proteins;</p> <p>(ii) inflammatory response in keratinocytes;</p> <p>(iii) activation of dendritic cells;</p> <p>(c) skin sensitisation <i>in vivo</i> testing. The Murine Local Lymph</p> | <p>The study/ies in column 1 do(es) not need to be conducted if:</p> <ul style="list-style-type: none"> - the available information indicates that the substance meets the criteria for classification for skin sensitisation or skin corrosion - the substance is a strong acid (pH ≤ 2,0) or base (pH ≥ 11,5), or - the substance is spontaneously flammable in air or in contact with water or moisture at room temperature. <p><i>In vitro</i> tests do not need to be conducted if:</p> <ul style="list-style-type: none"> - an <i>in vivo</i> study referred to in point (c) of column 1 of this row is available, or - the available <i>in vitro</i> or <i>in chemico</i> test methods are not applicable for the substance or the results obtained from those studies are not adequate for classification and risk assessment. <p>If information from test method(s) addressing one or two of the key events described under point (b) in column 1 of this row allows for classification of the substance and risk assessment, studies addressing the other key event(s) do not need to be conducted</p> <p>An <i>in vivo</i> study for skin sensitisation shall be conducted only if <i>in vitro</i> or <i>in chemico</i> test methods described under point (b) in column 1 of this row are not applicable, or the results obtained from those studies are not adequate for classification and risk assessment</p> <p><i>In vivo</i> skin sensitisation studies that were carried out or initiated before 15 April 2022 shall be considered</p> |

| | |
|---|--|
| Node Assay (LLNA) is the first-choice method for <i>in vivo</i> testing. Another skin sensitisation test may only be used in exceptional cases. If another skin sensitisation test is used, justification shall be provided | appropriate to address this information requirement' |
|---|--|

1
2 If the substance is a skin sensitiser based on *in vitro/in chemico* testing and the results of *in vitro/in chemico* testing allow a sufficiently reliable conclusion that the substance has the
3 potential to produce significant sensitisation in humans (Cat. 1A), no further testing is required.
4

5 If the substance is a skin sensitiser based on *in vitro/in chemico* testing, but the results of *in vitro/in chemico* testing allow a sufficiently reliable conclusion that the substance does not have
6 the potential to produce significant sensitisation in humans, the substance can be presumed to
7 be a moderate skin sensitiser (Cat. 1B). In this case, no further testing is needed. However, if
8 significant sensitisation (Cat. 1A) cannot be excluded with sufficient confidence based on *in vitro/in chemico* testing, additional information (*in silico/in vitro/in chemico*) would need to be
9 generated to strengthen the weight of evidence. If still no reliable conclusion can be reached, as
10 a last resort *in vivo* testing (LLNA) would need to be performed (Tier c).
12

13 According to data requirements, it is necessary to conclude whether the substance *can be*
14 *presumed to have the potential to produce significant sensitisation in humans (Category 1A)*.
15 However, in case there is already existing *in vivo* information (study initiated before 15 April
16 2022) that does not allow assessing the skin sensitisation potency, this information can still be
17 used to fulfil the information requirement and no additional testing is required. In such cases,
18 any information on skin sensitisation potency coming from such studies should be used together
19 with existing information from other sources or with additional non-animal test data to refine
20 the classification and risk assessment.

21 **Tier a) Assessment of the available human, animal and non-animal data**

22 In this Tier, all available information (including physico chemical properties) should be evaluated
23 before undertaking any new testing. In case new testing is needed, *in vitro* tests must be
24 performed first, and it should be assessed whether *in vivo* testing can be completely avoided.

25 Further guidance regarding the assessment of existing information (physicochemical properties,
26 grouping, (Q)SARs and expert systems, *in vitro* data, human data and animal data) is available
27 within the *Guidance on the Application of the CLP Criteria, BPR Guidance Volume III Parts B+C*
28 and *REACH Guidance on Information Requirements and Chemical Safety Assessment Chapter*
29 *R.7a*.

30 In principle information requirements for skin sensitisation do not apply if:

- 31 1. the available information indicates that the substance meets the criteria for classification
32 for skin sensitisation or skin corrosion,
- 33 2. the substance is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$), or
- 34 3. the substance is spontaneously flammable in air or in contact with water or moisture at
35 room temperature

36 The decision on the need to test a substance for skin sensitisation when it fulfils one or more of
37 the above conditions requires expert judgment. This is because the information on skin
38 sensitisation from the active substance will be used for the assessment of this property for
39 products containing the substance, and it needs to be taken into account e.g. whether sub-
40 corrosive concentrations of a substance may still have sensitising properties. For a substance
41 that is corrosive, strong acid or strong base, the decision-making process on testing needs to

1 take into account all the available information as specified in this tier.

2 *In vitro* skin sensitisation test does not need to be performed in cases when:

3 1. *in vivo* study for skin sensitisation is already available, or

4 2. the available *in vitro* or *in chemico* test methods are not applicable for the substance or
5 the results obtained from those studies are not adequate for classification and risk
6 assessment.

7 If a good quality *in vivo* skin sensitisation study is already available, i.e. study was carried out
8 or initiated before 15 April 2022, it can be used to fulfil the information requirement even if no
9 conclusion on the skin sensitisation potency (Cat 1A or 1B of CLP) can be made.

10 For existing animal data, the use of methods other than those that are specified in the Annex to
11 the EU Test Methods Regulation or the corresponding OECD methods may be accepted on a
12 case-by-case basis, considering the reliability of the information and the relevance for
13 classification and labelling.

14 When reliable and relevant human data are available, they can be useful for hazard identification
15 and are even preferable over animal data. However, the lack of positive findings in humans does
16 not necessarily overrule positive results in good quality animal data. When human studies have
17 been performed for safety assessment, the aim is to ensure that a specific concentration does
18 not induce skin sensitisation, however those studies do not determine whether a substance has
19 an intrinsic property to cause skin sensitisation. The situation is similar when diagnostic tests
20 are carried out to see if an individual is sensitised to a specific agent, and not to determine
21 whether the agent can cause sensitisation.

22 **Considerations before performing further testing**

23 If after the analysis in Tier a) further testing is needed to assess the potential for skin
24 sensitisation, the test methods mentioned below should be used. Where new testing is needed,
25 please see also the general information under *Considerations before initiating testing* in chapter
26 1.

27 The tests can provide information on i) whether the substance is a skin sensitiser or not, and ii)
28 how potent sensitiser the substance is.

29 **Tier b) Generation of new *in chemico/in vitro* test data**

30 If after the analysis in Tier a) above further testing is needed to assess the potential for skin
31 sensitisation, the test methods listed in Table 6 should be used. The limitations and the scope
32 of a given test method within a test guideline should be taken into account when selecting the
33 most appropriate *in vitro* method for a particular substance and when interpreting the test
34 results. Where new testing is needed, please see also the general information under
35 *Considerations before initiating testing* in chapter 1.

36 As specified in the data requirement, all three key events need to be addressed. In case the *in*
37 *chemico/in vitro* methods for one or more of the skin sensitisation key event(s) are not suitable
38 for the substance, a scientific justification of that needs to be provided.

39 **Table 6. *In chemico/in vitro* test methods for skin sensitisation**

| AOP KEY EVENT | TEST METHOD | EU TEST METHODS/ OECD TEST GUIDELINE | OUTCOME ACCORDING TO THE TEST METHOD/GUIDELINE |
|--|----------------|---|--|
| Key Event 1 Peptide/protein binding | DPRA | B.59/TG 442C | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| | ADRA | N.A/TG 442C | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| | kDPRA* | N.A/ TG 442C | Skin sensitiser (Cat 1A) or non-category 1A (cannot differentiate between Cat 1B and non-sensitiser) |
| Key Event 2 Keratinocyte response | KeratinoSens™ | B.60/TG 442D | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| | LuSens | N.A/N.A | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| Key Event 3 Monocytic /Dendritic cell response | h-CLAT | B.71/TG 442E | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| | U-SENS™ | B.71/TG 442E | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| | IL-8 Luc Assay | B.71/TG 442E | Skin sensitiser (Cat 1) or non-sensitiser with complementary information |
| Defined approaches | 2 out of 3* | N.A/xxx | Skin sensitiser (Cat 1) or non-sensitiser |
| | ITS v1 and v2* | N.A/xxx | Skin sensitiser (Cat 1, 1A and 1B) and non-sensitiser |

1 * **Note to PEG members:** adopted by WNT-33, pending approval from of the OECD Chemicals and Biotechnology
2 Committee (meeting 8-10 June 2021).

3 **Abbreviations:** DPRA: Direct Peptide Reactivity Assay, ADRA: Amino acid Derivative Reactivity Assay, kDPRA: kinetic
4 DPRA, h-CLAT: Human Cell Line Activation test, U-SENS™: U937 cell line activation Test, IL8-Luc assay: Interleukin-8
5 Reporter Gene Assay, ITS: Integrated testing strategy

6 **Tier c) Generation of new *in vivo* test data**

7 If after the analysis in Tiers a) and b) above further testing is needed to assess the potential for
8 skin sensitisation, the test methods listed in Table 7 should be used. Where new testing is
9 needed, please see also the general information under *Considerations before initiating testing* in
10 chapter 1.

1 **Table 7. *In vivo* Murine Local Lymph Node assay (LLNA) test methods for skin sensitisation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|-------------------------------|--------------------------------------|---|
| Local Lymph Node Assay (LLNA) | B.46 / TG 429 | Skin sensitiser (Cat. 1, 1A and 1B) or non-sensitiser |
| LLNA: DA. | B.50 / TG 442A | Skin sensitiser (Cat. 1) or non-sensitiser |
| LLNA: BrdU-ELISA | B.51 / TG 442B | Skin sensitiser (Cat. 1) or non-sensitiser |

2

3 The EU method B.46/OECD TG 429 is recommended because information provided by the LLNA
4 assay according to this method should be adequate for the assessment of the skin sensitisation
5 potency. For the two LLNA variants there are no CLP criteria available to predict the skin
6 sensitisation potency (Cat 1A or 1B).

7 Specific limitations that may be described within the Test Guideline protocol should be taken into
8 account before performing a test and when interpreting the test results.

9 If the LLNA assay is not considered suitable due to the properties of the substance to be tested,
10 other OECD Test Guideline protocols can be used for the assessment of skin sensitisation, such
11 as the methods in Table 8. If another *in vivo* method than LLNA is used, a scientific justification
12 shall be provided.

13 **Table 8. Other *in vivo* test methods for skin sensitisation**

| TEST METHOD | EU TEST METHOD / OECD TEST GUIDELINE | CLASSIFICATION ACCORDING TO CLP REGULATION |
|------------------------------|--------------------------------------|--|
| Guinea Pig Maximization test | B.6 / TG 406 | Skin sensitiser (Cat. 1, 1A and 1B) or non-sensitiser* |
| Buehler Assay | B.6 / TG 406 | Skin sensitiser (Cat. 1, 1A and 1B) or non-sensitiser* |

14 * Due to the study design, potency estimation for skin sensitising substances (Cat 1A or 1B
15 according to CLP) based on Guinea Pig Maximization study or Buehler study is rarely possible.

16 1.4. Respiratory sensitisation and irritation

17 1.4.1. Respiratory sensitisation (ADS)

18 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|-------------------------------|---|
| 8.4 Respiratory sensitisation | |

19

20 There are currently no standard tests and no OECD test guidelines available for respiratory

1 sensitisation. Since an active substance identified as a skin sensitizer can potentially induce a
2 hypersensitivity reaction, potential respiratory sensitisation and respiratory elicitation after
3 dermal sensitisation should be taken into account when appropriate tests are available or when
4 there are indications of respiratory sensitisation effects.

5 The assessment of the potential of a substance to induce respiratory sensitisation should include
6 the assessment of the available existing information including physico-chemical properties,
7 grouping, (Q)SARs and expert systems, *in vitro* data, human and animal data, and the outcome
8 of immunotoxicity assessment (see section 1.13.4 of this guidance). The assessment should also
9 consider *Guidance on the Application of the CLP Criteria* and *BPR Volume III Human health Parts*
10 *B+C*.

11 The following information should be provided where available, including any details necessary
12 for the evaluation of the information (please see also ECHA Guidance Vol III, Parts B+C):

- 13 • Information on respiratory sensitisation or any incidences of respiratory hypersensitivity
14 of workers or others exposed.
- 15 • Evidence that the substance can induce specific respiratory hypersensitivity will usually
16 be based on human experience data. The clinical history data including both medical and
17 occupational history, and reports from appropriate lung function tests related to exposure
18 to the substance should be submitted, if available.
- 19 • Reports of other supportive evidence, such as:
 - 20 ○ Information of a chemical structure within the active substance that is related to
21 substances known to cause respiratory hypersensitivity;
 - 22 ○ *In vivo* immunological tests;
 - 23 ○ *In vitro* immunological tests;
 - 24 ○ Studies indicating other specific but non-immunological mechanisms of action;
25 and
 - 26 ○ Data from a positive bronchial challenge test.

27 **1.4.2. Respiratory irritation (not in BPR Annex II)**

28 There is no testing requirement for respiratory irritation under the BPR, and there are currently
29 no standard tests or OECD TGs for respiratory irritation. Consequently, respiratory irritation is
30 not included in the testing strategies suggested in this Guidance. Nevertheless, account should
31 be taken of any existing and available data that provide evidence of the respiratory
32 corrosion/irritation potential of a substance. One should consider if the data on dermal or ocular
33 corrosion/irritation might contain information that is relevant for respiratory effects. Information
34 from cases where symptoms have been associated with occupational exposures can be used on
35 a case-by-case basis to characterise the respiratory irritation potency of a substance.
36 Information from acute and repeated dose inhalation toxicity studies may also be considered
37 sufficient to show that the substance causes respiratory irritation at a specific concentration level
38 or range. The data need to be carefully evaluated with regard to the exposure conditions and
39 sufficient documentation is required. Any confounding factors should be taken into account.

40 The exposure of atopic patients with bronchial asthma to some biocidal gases can result in so-
41 called acute, unspecific hyperreactivity, an exacerbation or airway hyperresponsiveness (AHR).
42 AHR is accompanied by adverse effects on human health and can constitute a serious health
43 impairment especially in infants. Experimental animal testing systems for AHR are not a data
44 requirement under BPR nor a part of an existing OECD TG, but any information on AHR should

1 be considered for the active substance if it has the irritation potency and exposure can take
2 place to the gas form.

3 Additional considerations for the evaluation of all available data with regard to respiratory
4 irritation are provided in BPR Volume III Human health Parts B+C, REACH Guidance on
5 Information Requirements and Chemical Safety Assessment Chapter R.7a and Appendix to
6 REACH Guidance Chapter R.8: Guidance for preparing a scientific report for health-based
7 exposure limits at the workplace (chapter A.8-17.2.2.2.1).

8 1.5. Mutagenicity

9 Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.5 Mutagenicity</p> <p>The assessment of this endpoint shall comprise the following consecutive steps:</p> <ul style="list-style-type: none">— an assessment of the available <i>in vivo</i> genotoxicity data— an <i>in vitro</i> test for gene mutations in bacteria, an <i>in vitro</i> cytogenicity test in mammalian cells and an <i>in vitro</i> gene mutation test in mammalian cells are required— appropriate <i>in vivo</i> genotoxicity studies shall be considered in case of a positive result in any of the <i>in vitro</i> genotoxicity studies | |

10 The testing of genotoxicity is intended to identify substances that might cause permanent
11 transmissible changes in the amount or structure of a single gene or gene segments, a block of
12 genes or chromosomes.
13

14 The aim of genotoxicity testing is to:

- 15 • predict genotoxic potential;
- 16 • identify genotoxic carcinogens at an early stage;
- 17 • elucidate the mechanism of action of active substances inducing germ-line mutations,
18 which may lead to inherited disorders.

19 Appropriate dose levels, depending on the test requirements, should be used in either *in vitro*
20 or *in vivo* assays. A tiered approach should be adopted, with selection of higher tier tests being
21 dependent upon interpretation of results at each stage.

22 At least one *in vitro* test for gene mutations in bacteria, one test for cytogenicity in mammalian
23 cells and one test for gene mutation in mammalian cells are required.

24 Collection and evaluation of available information

25 For the assessment of existing information (physicochemical properties, grouping, [Q]SARs and

expert systems, *in vitro* data, human data and animal data) further guidance is available within the *Guidance on the Application of the CLP Criteria* and BPR Volume III Human health, Evaluation and Assessment (Parts B+C). For further information, the following documents can be considered:

- Overview on Genetic Toxicology TGs (OECD 2017). OECD Series on Testing and Assessment, No. 238, OECD Publishing, Paris, <https://doi.org/10.1787/9789264274761-en>
- Clarification of some aspects related to genotoxicity assessment (EFSA 2017) <https://doi.org/10.2903/j.efsa.2017.5113>

Generation of new test data

If after the analysis above further testing is needed to assess the potential for genotoxicity *in vitro*, the test methods below should be used. Where new testing is needed, please see also the general information under *Considerations before initiating testing* in chapter 1.

Testing for genotoxicity (in vitro assays)

The test guideline protocols to follow for the investigation of *in vitro* genotoxicity are listed below (section 1.5.1 to 1.5.3 of this guidance). These should be used taking into account some considerations described here but also taking into account the existing information for this endpoint and its assessment.

If there are indications of micronucleus formation in an *in vitro* micronucleus assay, further testing with appropriate centromere labelling should be conducted to clarify if there is an aneugenic or clastogenic response. Further investigation of the aneugenic response may be considered to determine whether there is sufficient evidence for a threshold mechanism and threshold concentration for the aneugenic response (particularly for non-disjunction).

Active substances which display highly bacteriostatic properties as demonstrated in a range finding test do not need an Ames test. Such substances should be tested in at least one *in vitro* mammalian cell test for gene mutation, i.e. in either an In Vitro Mammalian Cell Gene Mutation Tests Using the Thymidine Kinase Gene (OECD 490) or an In Vitro Mammalian Cell Gene Mutation Tests using the Hprt and xpRT genes assay (OECD 476). If the Ames test is not performed, this should be justified.

For active substances bearing structural alerts that have given negative results in the standard test battery, additional testing may be required if the standard tests have not been optimised for these alerts. The choice of an additional study or study plan modifications depends on the chemical nature, the known reactivity and the metabolism data on the structurally alerting active substance.

1.5.1. *In vitro* gene mutation study in bacteria

Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.5.1 <i>In vitro</i> gene mutation study in bacteria | |

Test methods for *in vitro* gene mutation in bacteria:

- 1 • EC method B.13/14 Mutagenicity - reverse mutation test using bacteria.
2 • OECD Test Guideline 471: Bacterial Reverse Mutation Test.
3

4 **1.5.2. In vitro cytogenicity study in mammalian cells**

5 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.5.2 In vitro cytogenicity study in mammalian cells | |

6
7 Test methods for *in vitro* cytogenicity in mammalian cells:

- 8
9 • EC method B.10 Mutagenicity - *In vitro* mammalian chromosome aberration test.
10 • OECD Test Guideline 473: *In vitro* Mammalian Chromosome Aberration Test.
11 • OECD Test Guideline 487. *In vitro* Mammalian Cell Micronucleus Test.

12 With the current state of knowledge, the *in vitro* cell micronucleus test can be considered as the
13 preferred method for examining *in vitro* cytogenicity in mammalian cells due to its increased
14 sensitivity and ability to identify also the effect of aneugens provided that appropriate
15 centromere labelling is performed in case of positive results.
16

17 **1.5.3. in vitro gene mutation study in mammalian cells**

18 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.5.3 In vitro gene mutation study in mammalian cells | |

19
20 Test methods for *in vitro* gene mutation in mammalian cells:

- 21 • OECD Test Guideline 476: In Vitro Mammalian Cell Gene Mutation Tests using the Hprt
22 and xprt genes.
23 • OECD Test Guideline 490: In Vitro Mammalian Cell Gene Mutation Tests Using the
24 Thymidine Kinase Gene.
25

26 **1.6. In vivo genotoxicity study (ADS)**

27 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| 8.6 <i>In vivo</i> genotoxicity study The assessment shall comprise the following tiers: (a) If there is a positive result in any of the <i>in vitro</i> genotoxicity studies as listed in 8.5 and there | The study/ies in column 1 do(es) not need to be conducted if: — the results are negative for the three <i>in vitro</i> tests listed in 8.5 and no other concern has been identified (e.g. metabolites of concern formed in mammals), or — the substance meets the criteria to be classified as |

| | |
|---|--|
| <p>are no reliable results available from an appropriate <i>in vivo</i> somatic cell genotoxicity study, an appropriate <i>in vivo</i> somatic cell genotoxicity study shall be conducted;</p> <p>(b) A second <i>in vivo</i> somatic cell genotoxicity study may be necessary depending on the <i>in vitro</i> and <i>in vivo</i> results, type of effects, quality and relevance of all available data;</p> <p>(c) If there is a positive result from an <i>in vivo</i> somatic cell genotoxicity study available, the potential for germ cell mutagenicity should be considered based on all available data, including toxicokinetic evidence to demonstrate whether the substance has the capacity to reach germ cells. If no clear conclusions about germ cell mutagenicity can be made, additional investigations shall be considered</p> | <p>a germ cell mutagen category 1A or 1B.</p> <p>The germ cell genotoxicity test does not need to be conducted if the substance meets the criteria to be classified as a carcinogen, category 1A or 1B and a germ cell mutagen category 2'</p> |
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2 Collection and evaluation of available information

3 For the assessment of existing information (physicochemical properties, grouping, (Q)SARs and
4 expert systems, *in vitro* data, human data and animal data), further guidance is available within
5 the *Guidance on the Application of the CLP Criteria* and *BPR Volume III Human health Parts B+C*.

6 Generation of new test data

7 If after the analysis above further testing is needed to assess the potential for genotoxicity *in*
8 *vivo*, the test methods below should be used. Where new testing is needed, please see also the
9 general information under *Considerations before initiating testing* in chapter 1.

10 Testing for genotoxicity: *In vivo* studies in somatic cells (Tiers a-b)

11 Before any decisions are made on the need for *in vivo* testing, a review of the *in vitro* test results
12 and all available information on the toxicokinetic and toxicodynamic profile of the test substance
13 is needed. A particular *in vivo* test should be conducted only when it can be reasonably expected
14 from all the properties of the test substance and the proposed test protocol that the specific
15 target tissue will be adequately exposed to the test substance and/or its metabolites. If
16 necessary, a targeted investigation of toxicokinetics should be conducted before progressing to
17 *in vivo* testing (e.g. a preliminary toxicity test to confirm that absorption occurs and that an
18 appropriate dose route is used).

19 The comet assay and the *in vivo* micronucleus test can be combined into a single acute study
20 with appropriate modification of treatment and sampling times. These same endpoints can be
21 integrated into *in vivo* test as part of one of the short-term toxicity studies described under
22 section 1.9 of this guidance.

1 In the interest of ensuring that the number of animals used in genotoxicity tests is kept to a
2 minimum, using both males and females is not always necessary. In accordance with standard
3 guidelines, testing in one sex only is possible when the substance has been investigated for
4 general toxicity and no sex-specific differences in toxicity have been observed.

5 If the *in vitro* mammalian chromosome aberration test or the *in vitro* micronucleus test is positive
6 for clastogenicity, an *in vivo* test for clastogenicity using somatic cells such as metaphase
7 analysis in rodent bone marrow or micronucleus test in rodents should be conducted.

8 In case of a positive result in the *in vivo* micronucleus assay, appropriate staining procedure
9 such as fluorescence in-situ hybridisation (FISH) should be used to identify an aneugenic and/or
10 clastogenic response. For this purpose, two sets of slides should always be prepared before
11 scoring.

12 If any of the *in vitro* gene mutation tests is positive, an *in vivo* test to investigate the induction
13 of gene mutation should be conducted, such as the Transgenic Rodent Somatic and Germ Cell
14 Gene Mutation Assay.

15 When conducting *in vivo* genotoxicity studies, only relevant exposure routes and methods (*such*
16 *as* admixture to diet, drinking water, skin application, inhalation, gavage) should be used. There
17 should be convincing evidence that the relevant tissue will be reached by the chosen exposure
18 route and application method. Other exposure techniques (*such as* intraperitoneal or
19 subcutaneous injection) that are likely to result in abnormal kinetics, distribution and metabolism
20 should be justified.

21 The available test guideline protocols for assessing the *in vivo* genotoxic potential of a substance
22 are listed below and reflect current state of knowledge. The choice of the most appropriate test
23 to conduct should reflect the considerations described in this section and future
24 recommendations or changes within the OECD Test Guideline programme for this endpoint.

25 Test methods for *in vivo* genotoxicity:

- 26 • EC method B.12 - Mutagenicity - *In vivo* mammalian erythrocyte micronucleus test EC
27 method
- 28 • B.11 - Mutagenicity – *In vivo* mammalian bone-marrow chromosome aberration test
- 29 • OECD Test Guideline 474: Mammalian Erythrocyte Micronucleus Test
- 30 • OECD Test Guideline 475: Mammalian Bone Marrow Chromosome Aberration Test
- 31 • OECD Test Guideline 488: Transgenic Rodent Somatic and Germ Cell Gene Mutation
32 Assays
- 33 • OECD Test Guideline 489: In Vivo Mammalian Alkaline Comet Assay

34 **Testing for genotoxicity: *In vivo* studies in germ cells (Tier c)**

35 The potential to affect germ cells should always be considered for substances classified as
36 category 2 mutagens or giving positive results in *in vivo* tests for genotoxic effects in somatic
37 cells. The first step is to make an appraisal of all the available toxicokinetic and toxicodynamic
38 properties of the test substance. Expert judgment is needed at this stage to consider whether
39 there is sufficient information to conclude that the substance poses a mutagenic hazard to germ
40 cells. If this is the case, it can be concluded that the substance may cause heritable genetic
41 damage and no further testing is justified. Consequently, the substance is classified as a category
42 1B mutagen. If the appraisal of mutagenic potential in germ cells is inconclusive, additional
43 investigation will be necessary. In the event that additional information about the toxicokinetics

1 of the substance would resolve the problem, toxicokinetic investigation (i.e. not a full
2 toxicokinetic study) tailored to address this is required. The type of mutation produced in earlier
3 studies, namely gene, numerical chromosomal or structural chromosome changes, should be
4 considered when selecting the appropriate assay.

5 Alternatively, other methods can be used if deemed appropriate by expert judgment. These may
6 include the mammalian spermatogonial chromosome aberration test (OECD Test Guideline 483)
7 or gene mutation tests with transgenic animals (OECD Test Guideline 488). The comet assay as
8 described in the OECD Test Guideline 489 is, at present, not considered appropriate to measure
9 DNA strand breaks in mature germ cells.

10 The available test guideline protocols for assessing *in vivo germ cell mutagenicity* of a substance
11 are listed below according to the current state of knowledge. The choice of the most appropriate
12 test to conduct should reflect the considerations described in this section and future
13 recommendations or changes within the OECD Test Guideline programme for this endpoint.

14 Test methods for *in vivo* germ cell genotoxicity:

- 15 • EC method B.23 Mammalian spermatogonial chromosome aberration test.
- 16 • OECD Test Guideline [483: Mammalian Spermatogonial Chromosome Aberration Test](#).
- 17 • OECD Test Guideline 488: Transgenic Rodent Somatic and Germ Cell Gene Mutation
18 Assays.

19 **Specific considerations for *in vivo* genotoxicity testing**

20 For substances that are short-lived, reactive, *in vitro* mutagens, or for which no indications of
21 systemic availability have been presented, the analysis of tissues at initial sites of contact with
22 the body is a crucial element of the testing strategy. Expert judgment should be used on a case-
23 by-case basis to decide which tests are the most appropriate. The main options to investigate
24 local genotoxicity are the *in vivo* comet assay and the gene mutation test with transgenic
25 rodents. Both assays employ methods by which any tissue (containing nucleated cells) of an
26 animal can in theory be examined for effects on the genetic material. This gives the possibility
27 to examine site-of-contact tissues, i.e. epithelium of the respiratory or gastro-intestinal tract
28 (e.g. nasal epithelium and lungs for inhalation; glandular stomach and duodenum for oral route)
29 as target tissues of the assays. For any given substance, expert judgment, based on all the
30 available toxicological information, will indicate which of these tests are the most appropriate.
31 The route of exposure should be selected that best allows assessing the hazard posed to humans.
32 For poorly soluble or insoluble substances, the possibility of release of active molecules in the
33 gastrointestinal tract may indicate that a test involving the oral route of administration is
34 particularly appropriate.

35 Special testing requirements in relation to photogenotoxicity may be indicated by the structure
36 of a molecule for substances that absorb light within the range of natural sunlight (290-700 nm).
37 If the ultraviolet/visible molar extinction/absorption coefficient of the active substance and its
38 major metabolites is less than $1.000 \text{ L} \times \text{mol}^{-1} \times \text{cm}^{-1}$, photogenotoxicity testing is not required.
39 Please see also the ICH Guidance S10 on Photosafety Evaluation of Pharmaceuticals¹.

40 **1.7. Acute toxicity**

41 **Information requirement according to BPR Annex II:**

¹ Available at <https://www.ema.europa.eu/en/ich-s10-photosafety-evaluation-pharmaceuticals>.

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.7 Acute toxicity</p> <p>In addition to the oral route of administration (8.7.1), for substances other than gases, the information mentioned under 8.7.2 to 8.7.3 shall be provided for at least one other route of administration</p> <ul style="list-style-type: none"> — The choice for the second route will depend on the nature of the substance and the likely route of human exposure — Gases and volatile liquids should be administered by the inhalation route — If the only route of exposure is the oral route, then information for only that route need be provided. If either the dermal or inhalation route is the only route of exposure to humans then an oral test may be considered. Before a new dermal acute toxicity study is carried out, an <i>in vitro</i> dermal penetration study (OECD 428) should be conducted to assess the likely magnitude and rate of dermal bioavailability — There may be exceptional circumstances where all routes of administration are deemed necessary | <p>The study/ies do(es) not generally need to be conducted if:</p> <ul style="list-style-type: none"> — the substance is classified as corrosive to the skin |

1
2 Assessment of the acute toxic potential of a chemical is necessary to determine the adverse
3 health effects that might occur following accidental or deliberate short-term exposure.

4 Administration via different routes makes an overall assessment of relative acute hazard in
5 different exposure routes possible.

6 **Collection and evaluation of available information**

7 For the assessment of existing information (physicochemical properties, grouping and read-
8 across, (Q)SARs and expert systems, *in vitro* data, human data and animal data), further
9 guidance is available within the Guidance on the Application of the CLP Criteria, the BPR Volume
10 III Human health Guidance, Parts B+C and in the REACH Guidance on Information Requirements
11 and Chemical Safety Assessment Chapter R.7a: Endpoint specific guidance.

12 **1.7.1. By oral route**

13 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| 8.7.1 By oral route The Acute Toxic Class Method is the preferred method for the determination of this endpoint | The study need not be conducted if: – the substance is a gas or a highly volatile substance |

1

2 **Generation of new test data**

3 If after the analysis of all available information further testing is needed to assess the potential
 4 for acute toxicity by the oral route, the test methods below should be used. Where new testing
 5 is needed, please see also the general information under *Considerations before initiating testing*
 6 in chapter 1.

7 Test methods for Acute toxicity via oral route:

- 8 • EC method B.1 tris Acute oral toxicity - Acute toxic class method
- 9 • OECD Test Guideline 423: Acute oral toxicity: acute toxic class method
- 10 • EC method B.1 bis Acute oral toxicity - fixed dose procedure
- 11 • OECD Test Guideline 420: Acute oral toxicity: fixed dose procedure
- 12 • OECD Test Guideline 425: Acute oral toxicity: up-and-down procedure
- 13 • OECD Test Guideline 401: Acute oral toxicity (acceptable only if performed before
 14 December 2002)

15 According to the BPR data requirement, the acute toxic class method is the preferred study.
 16 However, taking into account animal welfare, in performing new studies the fixed dose procedure
 17 should be considered.

18 **1.7.2. By inhalation**

19 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.7.2 By inhalation Testing by the inhalation route is appropriate if exposure of humans via inhalation is likely taking into account: – the vapour pressure of the substance (a volatile substance has vapour pressure > 1×10^{-2} Pa at 20 °C) and/or – the active substance is a powder containing a significant proportion (e.g. 1 % on a weight basis) of particles with particle size MMAD < | |

| | |
|--|--|
| 50 micrometers or – the active substance is included in products that are powders or are applied in a manner that generates exposure to aerosols, particles or droplets of an inhalable size (MMAD < 50 micrometers) – the Acute Toxic Class Method is the preferred method for the determination of this endpoint | |
|--|--|

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Generation of new test data

3 If after the analysis of available information, and the considerations listed below, further testing
4 is needed to assess the potential for acute toxicity by inhalation, the test methods below should
5 be used. Where new testing is needed, please see also the general information under
6 *Considerations before initiating testing* in chapter 1.

7 If there is absence of information on particle/droplet size and where there is potential for
8 exposure via inhalation from the use of biocidal products containing the active substance, an
9 acute inhalation study should be performed.

10 Test methods for Acute toxicity via inhalation route:

- 11 • OECD Test Guideline 436: Acute Inhalation Toxicity – Acute Toxic Class Method
- 12 • OECD Test Guideline 433: Acute Inhalation Toxicity: Fixed Concentration Procedure
- 13 • EC method B.2 Acute toxicity (inhalation)
- 14 • OECD Test Guideline 403: Acute Inhalation Toxicity

15 When selecting an acute inhalation study, preference should be given to OECD TG 436 (according
16 to BPR Annex II requirements) and secondarily to OECD TG 433, as these methods have been
17 designed to use less animals than EU B.2/OECD TG 403. However, in some circumstances, e.g.
18 if a dose-response curve is needed for risk assessment purposes, testing according to EU B.2 /
19 OECD TG 403 may be considered appropriate (see also the OECD Guidance Document 39).

20 The full study using three dose levels may not be necessary if a substance at an exposure
21 concentration equal to the limit concentrations of the test guideline (limit test) or at the
22 maximum attainable concentration produces no compound-related mortalities.

23 The head/nose only exposure should be used, unless whole body exposure can be justified.

24 **1.7.3. By dermal route**

25 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.7.3 By dermal route Testing by the dermal route is necessary only if: – inhalation of the substance is | |

| | |
|--|--|
| unlikely, or – skin contact in production and/or use is likely, and either – the physicochemical and toxicological properties suggest potential for a significant rate of absorption through the skin, or – the results of an in vitro dermal penetration study (OECD 428) demonstrate high dermal absorption and bioavailability | |
|--|--|

1

2 **Generation of new test data**

3 Dermal toxicity must be reported for an active substance except for gases.

4 If after the analysis of all available information further testing is needed to assess the potential
 5 for acute toxicity by the dermal route, the following test methods should be used. Where new
 6 testing is needed, please see also the general information under *Considerations before initiating*
 7 *testing* in chapter 1.

8 Test methods for Acute toxicity via dermal route:

- 9 • EC method B.3 Acute toxicity (dermal)
- 10 • OECD Test Guideline 402: Acute Dermal Toxicity

11 For substances with low acute dermal toxicity, a limit test with 2000 mg/kg body weight may be
 12 sufficient.

13 **1.8. Toxicokinetics and metabolism studies in mammals**14 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.8 Toxicokinetics and metabolism studies in mammals The toxicokinetics and metabolism studies should provide basic data about the rate and extent of absorption, the tissue distribution and the relevant metabolic pathway including the degree of metabolism, the routes and rate of excretion and the relevant metabolites | |

15

16 The generation of toxicokinetic data should be considered in light of the generation of other
 17 toxicity data (e.g. on repeated dose toxicity, mutagenicity, reproductive toxicity) to assist in the
 18 estimation of systemic exposure to the active substance and/or its metabolites and the
 19 correlation of the effects observed with internal dose estimates. This is important in establishing

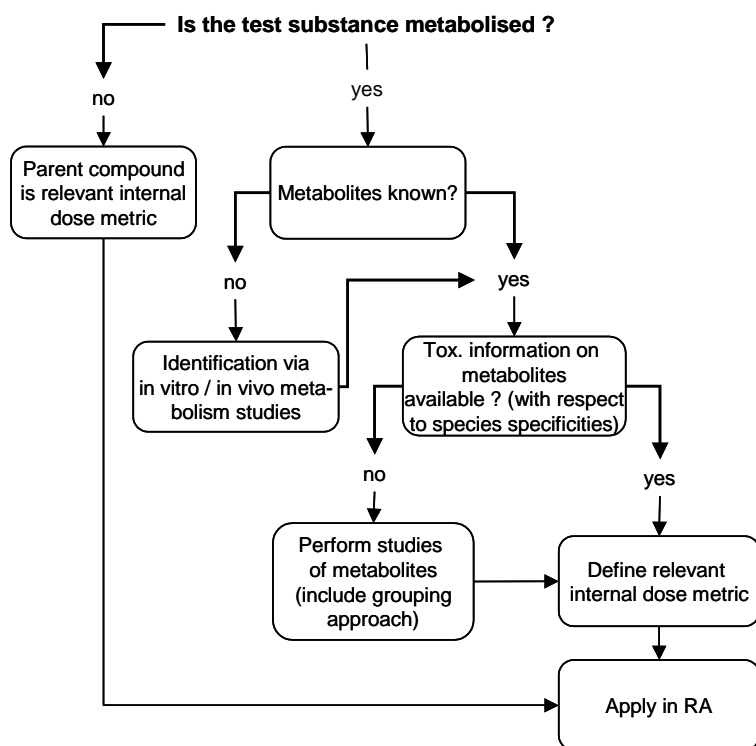
1 the mode of action of the active substance and whether administered doses cause non-linear
2 dose response due to saturation kinetics. Such information is valuable in the derivation of
3 assessment factors, route-to-route extrapolation and hazard characterisation, as well as in
4 considering the validity of read-across and grouping approaches.

5 **Collection and evaluation of available information**

6 For the assessment of existing information (physicochemical properties, grouping, (Q)SARs and
7 expert systems, *in vitro* data, human and animal data) further guidance is available within *BPR*
8 *Guidance Volume III Human health Parts B+C* and the REACH Guidance on Toxicokinetics within
9 the REACH CSA&IR, Chapter R.7c: Endpoint specific guidance.

10 **Generation of new test data**

11 Following the evaluation of all available data, a decision should be made on which type of kinetic
12 data and which test design is the most appropriate. It is preferred to generate kinetic data within
13 the toxicity studies such as repeated dose toxicity studies where possible. The sections below
14 describe the issues to consider when designing new tests for toxicokinetics and the available
15 techniques for the tests suitable for ADME (absorption, distribution, metabolism, elimination)
16 estimation. See Figure 2 below regarding the use of metabolism information, and also Figure 3
17 in Chapter 1.9, explaining how toxicokinetic data can be used in the design of repeated dose
18 toxicity studies.



19

20 **Figure 2. Use of increasing knowledge on substance metabolism**

21 The OECD Test Guideline 417 provides the protocol for the conduct of toxicokinetic studies either
22 as standalone test or in combination with repeated dose toxicity studies.

23 *In vivo* studies provide an integrated perspective on the relative importance of different
24 processes in the intact biological system for comparison with the results of the toxicity studies.
25 To ensure a valid set of toxicokinetic data, a toxicokinetic *in vivo* study has to consist of several
26 experiments that include blood/plasma kinetics, mass balances and excretion experiments as
27 well as tissue distribution experiments. Depending on the problem to be solved, selected

1 experiments (e.g. plasma kinetics) may be sufficient to provide data for further assessments
2 (e.g. bioavailability).

3 The high dose level administered in an ADME study should be linked to the dose levels that cause
4 adverse effects in toxicity studies. Ideally there should also be a dose without toxic effect, which
5 should be in the range of expected human exposure including consideration of limit of
6 quantification. A comparison between toxic dose levels and those that are likely to represent
7 human exposure values may provide valuable information for the interpretation of adverse
8 effects and is essential for extrapolation and risk assessment.

9 In an *in vivo* study the systemic bioavailability is usually estimated by the comparison of either
10 dose-corrected amounts excreted, or of dose-corrected areas under the curve (AUC) of plasma
11 (blood, serum) kinetic profiles, after extra- and intravascular administration. The systemic
12 bioavailability is the dose-corrected amount excreted, or AUC determined after an extravascular
13 substance administration divided by the dose-corrected amount excreted, or AUC determined
14 after an intravascular substance application, which corresponds by definition to a bioavailability
15 of 100%. This is only valid if the kinetics of the compound is linear, i.e. dose-proportional, and
16 relies upon the assumption that the clearance is constant between experiments. If the kinetics
17 is not linear, the experimental strategy has to be revised on a case-by-case basis, depending of
18 the type of non-linearity involved (e.g. saturable protein binding, saturable metabolism, etc).

19 Generally *in vitro* studies provide data on specific aspects of toxicokinetics such as metabolism.
20 A major advantage of *in vitro* studies is that it is possible to carry out parallel tests on samples
21 from the species used in toxicity tests and samples from humans, thus facilitating interspecies
22 comparisons (e.g., metabolite profile, metabolic rate constants).

23 In recent years, methods to integrate a number of *in vitro* and *in silico* information into a
24 prediction of ADME *in vivo* by the use of appropriate physiologically based kinetic (PBK) models
25 have been developed. Such methods allow both the prediction of *in vivo* kinetics at early stages
26 of development, and the progressive integration of all available data into a predictive model of
27 ADME. The uncertainty associated with the prediction depends largely on the amount of available
28 data. PBK models have become an important tool to facilitate the translation of doses that elicit
29 biological responses in cellular systems to exposure levels *in vivo* (OECD 2021).

30 Information on the concentration of the active substance and relevant metabolites in blood and
31 tissues, for example around the time to reach the maximum blood (serum/plasma) concentration
32 (T_{max}) or other relevant toxicokinetic parameter, should be generated in short and long-term
33 studies on relevant species to better use the toxicological data generated in terms of
34 understanding the toxicity studies. If such information is not considered essential for the
35 assessment, full justification should be provided.

36 The main objective of the toxicokinetic data is to describe the systemic exposure achieved in
37 animals and its relationship to the dose levels and the time course of the toxicity studies. Other
38 objectives are:

39 (a) to relate the achieved exposure in toxicity studies to toxicological findings and contribute
40 to the assessment of the relevance of these findings to human health with a particular
41 regard to vulnerable groups;

42 (b) to support the design of a toxicity study (choice of species, treatment regimen, selection
43 of dose levels) with respect to kinetics and metabolism;

44 (c) to provide information which, in relation to the findings of toxicity studies, contributes to
45 the design of supplementary toxicity studies.

46 **Absorption, distribution, metabolism and excretion (ADME) after exposure by oral**
47 **route**

1 *Absorption*

2 Absorption is normally investigated by the determination of the test substance and/or its
3 metabolites in excreta, exhaled air and carcass (i.e. radioactivity balance). The biological
4 response between test and reference groups (e.g. oral versus i.v.) is compared and the
5 plasma/blood level of the test substance and/or its metabolites is determined.

6 *Distribution*

7 For determination of the distribution of a substance in the body, two approaches are available
8 at present for analysis of distribution patterns. Quantitative information can be obtained using
9 whole-body autoradiographic techniques, or by sacrificing animals at different times after
10 exposure and determination of the concentration and amount of the test substance and/or
11 metabolites in tissues and organs (EC method B.36: Toxicokinetics, OECD TG 417:
12 Toxicokinetics).

13 *Accumulative potential*

14 Information derived for the purpose of environmental risk assessment can be relevant for human
15 health risk assessment and the potential for a substance to accumulate. The static
16 bioconcentration factor (BCF) is the ratio of the concentration of a substance in an organism to
17 the concentration in water once a steady state has been achieved. The resulting fish BCF is
18 widely used as a surrogate measure for bioaccumulation potential. For further information, see
19 the Guidance on the BPR: Volume IV Environment (Part A; Parts B+C).

20 If single dose toxicity and tissue distribution data are not adequate to determine the potential
21 for accumulation, repeated dose administration may be needed to address the potential for
22 accumulation and/or persistence or changes in toxicokinetics.

23 Accumulating substances can also be measured in milk and therefore additionally allow an
24 estimation of transfer to the breast-fed pup.

25 *Metabolism*

26 *In vivo* toxicokinetic studies generally only determine the rates of total metabolic clearance (by
27 measuring radiolabelled products in blood/plasma, bile, and excrements) rather than the
28 contributions of individual tissues. It has to be taken into account that the total metabolic
29 clearance is the sum of the hepatic and potential extrahepatic metabolism.

30 *In vitro* tests can be performed using isolated enzymes, microsomes and microsomal fractions,
31 immortalised cell lines, primary cells and organ slices. Most frequently these materials originate
32 from the liver as this is the most relevant organ for metabolism, however, in some cases
33 preparations from other organs are used for investigation of potential organ-specific metabolic
34 pathways. In the absence of standardised *in vitro* methods, generation of novel *in vitro* ADME
35 data should be in accordance with the OECD guidance document on "Good *in vitro* Method
36 Practices" (GIVIMP) (OECD, 2018).

37 When using metabolically incompetent cells, an exogenous metabolic activation system is usually
38 added into the cultures. For this purpose, the post-mitochondrial 9000 g supernatant (S9
39 fraction) of whole liver tissue homogenate containing a high concentration of metabolising
40 enzymes is most commonly employed – the donor species needs to be considered in the context
41 of the study. In all cases metabolism may either be directly assessed by specific identification of
42 the metabolites or by subtractive calculation of the amount of parent substance lost in the
43 process.

44 *Excretion*

1 The major routes of excretion are in the urine and/or the faeces (via bile and directly from the
2 GI mucosa). For this purpose, urine, faeces and, in certain circumstances, bile are collected and
3 the amount of test substance and/or metabolites in these excreta is measured and those
4 accounting for 5% or more of the administered dose should be identified where possible (EC
5 method B.36: Toxicokinetics, OECD TG 417: Toxicokinetics).

6 The excretion of chemicals (metabolites) in other biological fluids such as saliva, milk, tears, and
7 sweat is usually negligible compared with renal or biliary excretion. However, in special cases
8 these fluids may be important to study either for monitoring purposes, or in the case of milk
9 allowing an assessment of the exposure of infants.

10 For volatile substances and metabolites, exhaled air has to be examined as it may be an
11 important route of elimination.

12 The use of *in silico* methods and physiologically based (pharmacokinetic (PBPK) modelling
13 should also be considered upfront in the assessment and toxicokinetic data generation.

14 Available data from human biological monitoring and biological marker measurement studies
15 should be part of the assessment. Further guidance on the use of these methods is provided in
16 *BPR Volume III Human health Parts B+C*.

17 **Aspects to consider in the design of tests for toxicokinetic data generation**

18 Limited data restricted to one *in vivo* test species (normally rat) may be all that is required as
19 regards absorption, distribution, metabolism and excretion after exposure by oral route. These
20 data can provide information useful in the design and interpretation of subsequent toxicity tests.
21 However, information on interspecies differences is crucial in extrapolation of animal data to
22 humans and information on metabolism following administration via other routes may be useful
23 in human risk assessments.

24 It is not possible to specify detailed information requirements in all areas, since the exact
25 requirements will depend on the results obtained for each particular test substance.

26 The studies should be designed on a case-by-case basis, considering generation of information
27 about the kinetics of the active substance and its metabolites in relevant species after being
28 exposed to the following conditions:

- 29 • a single oral dose (low and high dose levels);
- 30 • an intravenous dose (preferably), or if available, a single oral dose with assessment of
31 biliary excretion (low dose level); and
- 32 • a repeated dose.

33 When intravenous dosing is not feasible, a justification should be provided.

34 A key parameter is systemic bioavailability (F), obtained by comparison of the area under the
35 curve (AUC) after oral and intravenous dosing.

36 The information from the studies should include:

- 37 • rate and extent of oral absorption including maximal concentration in blood (C_{max}), AUC,
38 T_{max} and other appropriate parameters, such as bioavailability;
- 39 • potential for bioaccumulation;
- 40 • clearance and half-lives ($t_{1/2}$);

- 1 • distribution in major organs and tissues;
- 2 • information on the distribution in blood cells;
- 3 • chemical structure and quantification of metabolites in biological fluids and tissues;
- 4 • different metabolic pathways;
- 5 • route and time course of excretion of active substance and metabolites;
- 6 • information on enterohepatic circulation.

7 Comparative *in vitro* metabolism studies should be performed on animal species to be used in
8 pivotal studies and on human material (microsomes or intact cell systems) in order to determine
9 the relevance of the toxicological animal data and to guide in the interpretation of findings and
10 in further definition of the testing strategy.

11 An explanation must be given or further tests should be carried out where a metabolite is
12 detected *in vitro* in human material and not in the tested animal species.

13 **Absorption, distribution, metabolism and excretion after exposure by other routes**

14 Data on absorption, distribution, metabolism and excretion (ADME) following exposure by the
15 dermal route should be provided where toxicity following dermal exposure is of concern
16 compared to that following oral exposure. Before investigating ADME *in vivo* following dermal
17 exposure, the need to conduct an *in vitro* dermal penetration study should be considered in order
18 to assess the likely magnitude and rate of dermal bioavailability, also taking note of the
19 possibility of using default values for estimating dermal uptake and excretion as described in
20 *BPR Volume III Human health Parts B+C*.

21 Absorption, distribution, metabolism and excretion after exposure by the dermal route should
22 be considered on the basis of the above information, unless the active substance causes skin
23 irritation that would compromise the outcome of the study.

24 For volatile active substances (vapour pressure $>10^{-2}$ Pa at 20 °C) absorption, distribution,
25 metabolism and excretion after exposure by inhalation may be useful in human risk
26 assessments.

27 *Dermal absorption*

28 An appropriate dermal absorption assessment is needed. It is not always mandatory to submit
29 experimental data. If such data are not available, as a first step default values can be used
30 according to the EFSA Guidance Document on Dermal Absorption (EFSA, 2017).

31 The following Test Guidelines are available for skin absorption studies:

- 32 • EC method B.45 Skin Absorption: *In Vitro* Method (human tissue preferred over rat)
- 33 • OECD Test Guideline 428: Skin Absorption: *In Vitro* Method (human tissue preferred over
34 rat)
- 35 • EC method B.44 Skin Absorption: *In Vivo* Method
- 36 • OECD Test Guideline 427: Skin Absorption: *In Vivo* Method

37 If testing to assess the likely magnitude and rate of dermal bioavailability is necessary, the OECD
38 Test Guideline 428 for *in vitro* skin absorption should be considered first.

1 Percutaneous absorption depends on the partitioning of substances from the vehicle and
2 solubility in the vehicle. OECD TG 427 and TG 428 recommend conducting tests using test
3 preparations that are the same as (or a realistic surrogate to) those that humans may be exposed
4 to.

5 *In vitro* methods are designed to measure the penetration of chemicals into the skin and their
6 subsequent permeation through the skin into a fluid reservoir, as well as partition to the different
7 skin layers and possible deposition therein. Provided that the excised skin sample is intact and
8 its integrity has been proven by appropriate methods, it can reasonably be assumed that its
9 barrier function to what is generally a diffusional process has been maintained *in vitro* (also after
10 frozen storage [Harrison et al., 1984, Bronaugh 39 et al., 1986 and Steinling et al., 2001]).

11 Very lipophilic substances are difficult to examine *in vitro* because of their low solubility in most
12 receptor fluids. By including the amount retained in the skin *in vitro*, a more acceptable
13 estimation of skin absorption can be obtained. Water soluble substances can be tested more
14 accurately *in vitro* because they diffuse into the receptor fluid more readily (OECD, 2004a).

15 At present, results from *in vitro* methods seem to adequately reflect those from *in vivo*
16 experiments, supporting their use as a replacement test to measure percutaneous absorption
17 (Lehman et al. 2011).

18 Advantages of the *in vivo* method (EC method B.44, OECD TG 427) are that it uses a
19 physiologically and metabolically intact system and a species common to many toxicity studies,
20 and it can be modified for use with other species. The disadvantages are the use of animals, the
21 need for radiolabelled material to facilitate reliable results, difficulties in determining the early
22 absorption phase and the differences in permeability of the preferred species (rat) and human
23 skin. Animal skin is generally more permeable and therefore may overestimate human
24 percutaneous absorption. The experimental conditions should also be taken into account in
25 interpreting the results. For instance, dermal absorption studies in fur-bearing animals may not
26 accurately reflect dermal absorption in humans.

27 When valid (guideline-compliant and GLP) *in vitro* studies on human skin, *in vitro* studies in
28 animals and *in vivo* animal studies are available and conducted under the same experimental
29 conditions, and the results meet the quality criteria, in particular with respect to variability,
30 number of acceptable replicates and recovery, then the 'Triple Pack' approach can be used to
31 extrapolate the human dermal absorption values for risk assessment (OECD No. 156, draft) (see
32 also section 2.6 of this guidance).

33 *In silico* models might also provide information on dermal absorption, but currently they have
34 not gained regulatory acceptance. *In silico* models for prediction of dermal absorption for
35 pesticides have been evaluated and reported (Kneuer et al. 2018). Mathematical skin permeation
36 models are usually based on uptake from aqueous solution which may not be relevant for the
37 exposure scenario being assessed. In addition, the use of such models for quantitative risk
38 assessment purposes is often limited because these models have generally been validated by *in*
39 *vitro* data ignoring the fate of the skin residue levels. However, these models may prove useful
40 as a screening tool or for qualitative comparison of skin permeation potential. On a case-by-case
41 basis, and if scientifically justified, the use of (Q)SARs may prove useful, especially within a
42 group of closely related substances.

43 **Considerations for test substances and analytical methodology for toxicokinetic** 44 **studies**

45 Toxicokinetic and metabolism studies can be carried out using non-labelled compounds, stable
46 isotope-labelled compounds, radioactively labelled compounds or using dual (stable and radio-)
47 labelling. The labels should be placed in metabolically stable positions, avoiding the placing of
48 labels such as ¹⁴C in positions from which they can enter the carbon pool of the test animal. If
49 metabolic degradation of the test substance may occur, different labelling positions have to be

1 taken into account to be able to determine all relevant degradation pathways. The radiolabelled
2 compound must be of high radiochemical purity and of adequate specific activity to ensure
3 sufficient sensitivity in radio-assay methods.

4 Separation techniques are used in metabolism studies to purify and separate several radioactive
5 fractions in biota such as urine, plasma, bile and others. These techniques range from relatively
6 simple approaches such as liquid-liquid extraction and column chromatography to more
7 sophisticated techniques such as HPLC (high pressure liquid chromatography). These methods
8 also allow the establishment of a metabolite profile. Quantitative analytical methods are required
9 to follow concentrations of parent compound and metabolites in the body as a function of time.
10 The most common techniques used are LC/MS (liquid chromatography/ mass spectroscopy) and
11 high performance LC with UV-detection, or if ¹⁴C-labelled material is used, radioactivity detection
12 HPLC. It is worth mentioning that kinetic parameters generally cannot be calculated from
13 measurement of total radioactivity to receive an overall kinetic estimate. Nevertheless, to
14 generate exact values one has to address parent compound and metabolites separately. An
15 analytical step is required to define the radioactivity as chemical species. This is usually faster
16 than cold analytical methods. Dual labelling (e.g. ¹³C and ¹⁴C/¹²C) is the method of choice for
17 structural elucidation of metabolites (by MS and NMR spectroscopy). A cold analytical technique,
18 which incorporates stable isotope labelling (for GC/MS [gas chromatography/mass spectroscopy]
19 or LC/MS), is a useful combination. Unless this latter method has already been developed for
20 the test compound in various matrices (urine, faeces, blood, fat, liver, kidney, etc.), the use of
21 radiolabelled compound may be less costly than other methods.

22 In any toxicokinetic study, the identity and purity of the substance used in the test must be
23 assured. Analytical methods capable of detecting undesirable impurities will be required, as well
24 as methods to assure that the substance of interest is of uniform potency from batch to batch.
25 Additional methods will be required to monitor the stability and uniformity of the form in which
26 the test substance is administered to the organisms used in the toxicokinetic studies. Finally,
27 methods suitable to identify and quantify the test substance in toxicokinetic studies must be
28 employed.

29 In the context of analytical methods, *accuracy* refers to how closely the average value reported
30 for the assay of a sample corresponds to the actual amount of substance being assayed in the
31 sample, whereas *precision* refers to the amount of scatter in the measured values around the
32 average result. If the average assay result differs from the actual amount in the sample, the
33 assay is said to be *biased*, i.e., lacks specificity; bias can also be due to low recovery.

34 Assay *specificity* is perhaps the most serious problem encountered. Although *blanks* provide
35 some assurance that no instrument response will be obtained in the absence of the test chemical,
36 a better approach is to select an instrument or bioassay that responds to some biological,
37 chemical, or physical property of the test chemical that is not shared with many other
38 substances.

39 The assay method should be usable over a sufficiently wide range of concentrations for the
40 substance and its metabolites. The lower limit of reliability for an analytical method has been
41 perceived in different ways; frequently, the term sensitivity has been used to indicate the ability
42 of an analytical method to measure small amounts of a substance accurately and with requisite
43 precision. It is unlikely that a single analytical method will be of use for all these purposes.
44 Indeed, it is highly desirable to use more than one method. If two or more methods yield
45 essentially the same results, confidence in each method is increased.

46 **1.8.1. Further toxicokinetic and metabolism studies in mammals (ADS)**

47 **Information requirement according to BPR Annex II:**

INFORMATION REQUIRED

SPECIFIC RULES FOR ADAPTATION FROM STANDARD

| | INFORMATION |
|---|-------------|
| <p>8.8.1 Further toxicokinetic and metabolism studies in mammals</p> <p>Additional studies might be required based on the outcome of the toxicokinetic and metabolism study conducted in rat. These further studies shall be required if:</p> <ul style="list-style-type: none"> — there is evidence that metabolism in the rat is not relevant for human exposure — route-to-route extrapolation from oral to dermal/inhalation exposure is not feasible <p>Where it is considered appropriate to obtain information on dermal absorption, the assessment of this endpoint shall proceed using a tiered approach for assessment of dermal absorption</p> | |

1
2 With the core dataset, basic information about the rate and extent of absorption, the tissue
3 distribution and the relevant metabolic pathway including the degree of metabolism, the routes
4 and rate of excretion and the relevant metabolites should be provided by the toxicokinetic and
5 metabolism studies (BPR Annex II Section 8.8). Additional information might be needed based
6 on the outcome of the toxicokinetic and metabolism study conducted in rats (ADS according to
7 Annex II Section 8.8.1) or based on the evaluation of the toxicological and physicochemical
8 profile of the substance.

9 Further toxicokinetic/metabolism studies with repeated dose administration may be necessary
10 for example when there are indications for a potential of the active substance to accumulate, to
11 persist or to change the toxicokinetics e.g. by induction of metabolic enzymes. Section 1.8 of
12 this guidance provides guidance on the options available for the toxicokinetics study and its
13 integration with the repeated dose toxicity tests.

14 1.9. Repeated dose toxicity

15 Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.9 Repeated dose toxicity</p> <p>In general, only one route of administration is necessary and the oral route is the preferred route. However, in some cases it may be</p> | <p>The repeated dose toxicity study (28 or 90 days) does not need to be conducted if:</p> <ul style="list-style-type: none"> — a substance undergoes immediate disintegration and there are sufficient data on the cleavage products for systemic and local effects and no synergistic effects |

| | |
|---|---|
| <p>necessary to evaluate more than one route of exposure.</p> <p>For the evaluation of the safety of consumers in relation to active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route</p> <p>Testing by the dermal route shall be considered if:</p> <ul style="list-style-type: none">— skin contact in production and/or use is likely, and— inhalation of the substance is unlikely, and— one of the following conditions is met:<ul style="list-style-type: none">(i) toxicity is observed in an acute dermal toxicity test at lower doses than in the oral toxicity test, or(ii) information or test data indicate dermal absorption is comparable or higher than oral absorption, or(iii) dermal toxicity is recognised for structurally related substances and for example is observed at lower doses than in the oral toxicity test or dermal absorption is comparable or higher than oral absorption <p>Testing by the inhalation route shall be considered if:</p> <ul style="list-style-type: none">— exposure of humans via inhalation is likely taking into account the vapour pressure of the substance (volatile substances and gases have vapour pressure $> 1 \times 10^{-2}$ Pa at 20 °C), and/or— there is the possibility of exposure to aerosols, particles or droplets of an inhalable size (MMAD < 50 micrometers) | <p>are expected, or</p> <ul style="list-style-type: none">— relevant human exposure can be excluded in accordance with Section 3 of Annex IV <p>In order to reduce testing carried out on vertebrates and in particular the need for free-standing single-endpoint studies, the design of the repeated dose toxicity studies shall take account of the possibility to explore several endpoints within the framework of one study</p> |
|---|---|

1

2 Repeated dose toxicity testing provides information on adverse effects as a result of repeated or
3 prolonged exposure. The objectives of assessing repeated dose toxicity are to evaluate:

4 1. adverse effects based on human or non-human studies:

5 • whether exposure of humans to a substance is associated with adverse toxicological

- 1 effects occurring as a result of repeated daily exposure for a part of the expected
2 lifetime or for the major part of the lifetime; these human studies potentially may
3 also identify populations that have higher susceptibility;
- 4 • whether administration of a substance to experimental animals causes adverse
5 toxicological effects as a result of repeated daily exposure for a part or a major part
6 of the expected lifespan; effects that are predictive of possible adverse human health
7 effects;
- 8 2. the target organs, potential cumulative effects and the reversibility of the adverse
9 toxicological effects;
- 10 3. the dose-response relationship and threshold for any of the adverse toxicological effects
11 observed in the repeated dose toxicity studies;
- 12 4. the basis for risk characterisation and classification and labelling (C&L) of substances for
13 repeated dose toxicity;
- 14 5. the mode of action (MOA) and mechanism data.

15 Repeated dose toxicity tests may also provide information relevant for reproductive toxicity,
16 carcinogenicity, neurotoxicity, immunotoxicity and endocrine disruption. If new studies are
17 performed, including relevant investigations on these effects should be considered on the basis
18 of all the information on the substance.

19 For the assessment of existing information (physico-chemical properties, grouping and read-
20 across², [Q]SARs and expert systems, *in vitro* data, human data and animal data) further
21 guidance is available within the *Guidance on the Application of the CLP Criteria*, the *BPR Volume*
22 *III Human health Parts B+C* and the practical guides³ such as "How to use and report (Q)SARs".

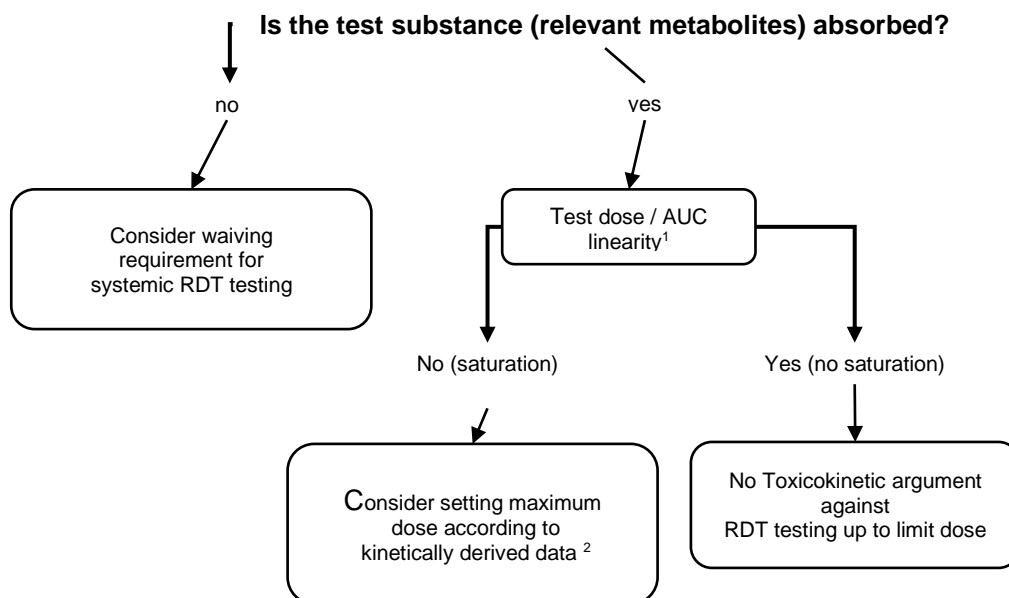
23 The most appropriate data on repeated dose toxicity are primarily obtained from studies in
24 experimental animals conforming to internationally agreed test guidelines.

25 Justification to replace the oral route by another significant route, or to require testing in addition
26 to the oral route needs to be provided.

27

² https://echa.europa.eu/documents/10162/13628/raaf_en.pdf

³ <https://echa.europa.eu/practical-guides>



¹ In the dose-range under consideration for RDT testing

² Meaning that the highest dose-level should not exceed the range of non-linear kinetics.

Figure 3. Use of toxicokinetic data in the design of repeated dose toxicity studies

1.9.1. Short-term repeated dose toxicity study (28 days), preferred species is rat

Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|--|
| 8.9.1 Short-term repeated dose toxicity study (28 days), preferred species is rat | The short-term toxicity study (28 days) does not need to be conducted if: <ul style="list-style-type: none"> (i) a reliable sub-chronic (90 day) study is available, provided that the most appropriate species, dosage, solvent and route of administration were used, (ii) the frequency and duration of human exposure indicates that a longer term study is appropriate and one of the following conditions is met: <ul style="list-style-type: none"> — other available data indicate that the substance may have a dangerous property that cannot be detected in a short-term toxicity study, or — appropriately designed toxicokinetic studies reveal accumulation of the substance or its metabolites in certain tissues or organs which would possibly remain undetected in a short term toxicity study but which are liable to result in adverse effects after prolonged exposure |

In principle, for substances where a 90-day repeated dose toxicity study needs to be performed, an additional 28-day repeated dose toxicity study will not be required.

If a 28-day repeated dose toxicity needs to be performed, the considerations described under

1 section 1.9.2 of this guidance regarding the generation of new test data should also be taken
2 into account.

3 **Generation of new test data**

4 If after evaluating the available information further testing is needed to assess repeated dose
5 toxicity, the test methods below should be used. Where new testing is needed, please see also
6 the general information under *Considerations before initiating testing* in chapter 1.

7 **Repeated dose toxicity (oral)**

8 Test methods for repeated dose toxicity via oral route:

- 9 • EC method B.7 Repeated dose (28 days) toxicity (oral).
- 10 • OECD Test Guideline 407: Repeated dose 28-day oral toxicity study in rodents.

11 **Repeated dose toxicity (dermal)**

12 If the substance is a severe irritant or corrosive, testing by the dermal route should be avoided
13 unless it can be performed at doses that do not cause irritation or corrosion and such doses are
14 still toxicologically relevant and the outcome can be used in risk assessment.

15 The following test methods for repeated dose toxicity via dermal route should be used:

- 16 • EC method B.9 Repeated dose (28 days) toxicity (dermal)
- 17 • OECD Test Guideline 410: Repeated dose dermal toxicity: 21/28-day study.

18 **Repeated dose toxicity (inhalation)**

19 The following test methods for repeated dose toxicity via inhalation route should be used:

- 20 • EC method B.8 Repeated dose (28 days) toxicity (inhalation)
- 21 • OECD Test Guideline 412: Subacute inhalation toxicity: 28-day study

22 **1.9.2. Sub-chronic repeated dose toxicity study (90-day), preferred species is** 23 **rat**

24 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| 8.9.2 Sub-chronic repeated dose toxicity study (90 days), preferred species is rat | <p>The sub-chronic toxicity study (90 days) does not need to be conducted if:</p> <ul style="list-style-type: none"> — a reliable short-term toxicity study (28 days) is available showing severe toxicity effects according to the criteria for classifying the substance as H372 and H373 (Regulation (EC) No 1272/2008), for which the observed NOAEL-28 days, with the application of an appropriate uncertainty factor allows the extrapolation towards the NOAEL-90 days for the same route of exposure, and — a reliable chronic toxicity study is available, |

| | |
|--|--|
| | provided that an appropriate species and route of administration were used, or — the substance is unreactive, insoluble, not bioaccumulative and not inhalable and there is no evidence of absorption and no evidence of toxicity in a 28-day 'limit test', particularly if such a pattern is coupled with limited human exposure |
|--|--|

1

2 **Generation of new test data**

3 If after evaluating the existing data further testing is needed to assess repeated dose toxicity,
4 the test methods described further below should be used. Where new testing is needed, please
5 see also the general information under *Considerations before initiating testing* in chapter 1.

6 **Considerations for the design of the repeated dose subchronic toxicity studies**

7 The study will be performed in a single rodent species, preferably the rat. The oral route will be
8 used unless one of the other routes is more appropriate based on either the most relevant route
9 of human exposure or the physico-chemical properties of the substance. The other routes should
10 be considered especially if route-to-route extrapolation is not appropriate and the predominant
11 human exposure occurs via dermal and/or inhalation route. In the 90-day study, potential
12 neurotoxic and immunotoxic effects (see also sections 1.13.2 and 1.13.4 of this guidance),
13 genotoxicity by way of micronuclei formation and effects potentially related to changes in the
14 endocrine system (see also section 1.13.3 of this guidance) must be carefully considered during
15 the conduct of the test and reported, taking into account potential limitations when modifying
16 test protocols in order to investigate specific effects.

17 Information on mode of action from structurally similar substances should also be considered in
18 the design of repeated dose toxicity tests.

19 Repeated dose toxicity studies should be designed to provide information as to the amount of
20 the active substance that can be tolerated without adverse effects under the conditions of the
21 study and to elucidate health hazards occurring at higher dose levels. Such studies provide useful
22 data on the risks for those handling and using biocidal products containing the active substance,
23 among other possible exposed groups. In particular, repeated dose toxicity studies provide an
24 essential insight into possible adverse effects of the active substance and the risks to humans
25 as a result of repeated exposure. In addition, repeated dose toxicity studies provide information
26 useful in the design of chronic toxicity studies.

27 The studies, data and information to be provided and evaluated should be sufficient to permit
28 the identification of effects following repeated exposure to the active substance, and in particular
29 to further establish or indicate:

- 30 (a) the relationship between dose and observed adverse effects;
- 31 (b) toxicity of the active substance including where possible the No Observed Adverse
32 Effect Level (NOAEL);
- 33 (c) target organs where relevant (including immune, nervous, reproductive and endocrine
34 systems);
- 35 (d) the time course and characteristics of adverse effects with full details of behavioural
36 changes and possible pathological findings at post-mortem;
- 37 (e) specific adverse effects and pathological changes produced;

- 1 (f) where relevant the persistence and reversibility of certain adverse effects observed,
2 following discontinuation of dosing;
- 3 (g) where possible, the mode of toxic action;
- 4 (h) the relative hazard associated with the different routes of exposure;
- 5 (i) relevant critical endpoints at appropriate time points for setting reference values,
6 where necessary.

7 Toxicokinetic data (e.g. concentration of the active substance and/or the main metabolites in
8 blood) should be included in repeated dose toxicity studies, unless it can be justified why this is
9 not necessary. To avoid increased animal use, the data may be derived in range finding studies.

10 If nervous system, immune system, reproductive system or endocrine system are specific
11 targets in repeated dose toxicity studies at dose levels not producing marked toxicity,
12 supplementary studies, including functional testing, need to be considered.

13 **Repeated dose toxicity (oral route)**

14 The following test methods should be used.

15 Test methods for sub-chronic repeated dose toxicity via oral route:

- 16 • EC method B.26 Sub-chronic oral toxicity test. Repeated dose 90-day oral toxicity study
17 in rodents.
- 18 • EC method B.27 Sub-chronic oral toxicity test. Repeated dose 90-day oral toxicity study
19 in non-rodents.
- 20 • OECD Test Guideline 408: Repeated dose 90-day oral toxicity study in rodents.
- 21 • OECD Test Guideline 409: Repeated dose 90-day oral toxicity study in non-rodents.

22 **Repeated dose toxicity (inhalation route)**

23 The following test methods for sub-chronic repeated dose toxicity via inhalation route should be
24 used:

- 25 • EC method B.29 Sub-chronic inhalation toxicity study 90-day repeated inhalation dose
26 study using rodent species.
- 27 • OECD Test Guideline 413: Subchronic inhalation toxicity: 90-day study.

28 **Repeated dose toxicity (dermal route)**

29 If the substance is a severe irritant or corrosive, testing by the dermal route should be avoided
30 unless it can be performed at doses that do not cause irritation or corrosion and such doses are
31 still toxicologically relevant and the outcome can be used in risk assessment.

32 The following test methods for sub-chronic repeated dose toxicity via dermal route should be
33 used:

- 34 • EC method B.28 Sub-chronic dermal toxicity test: 90-day repeated dermal dose study
35 using rodent species.
- 36 • OECD Test Guideline 411: Subchronic dermal toxicity test: 90-day study.

1 **1.9.3. Long-term repeated dose toxicity (≥ 12 months)**

2 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.9.3 Long-term repeated dose toxicity (≥ 12 months) | The long-term toxicity study (≥ 12 months) does not need to be conducted if: — Long-term exposure can be excluded and no effects have been seen at the limit dose in the 90-day study or — a combined long-term repeated dose/ carcinogenicity study (8.11.1) is undertaken |

3
4 Any new long-term toxicity study and carcinogenicity study (section 1.11 of this guidance) should
5 be combined. This section provides guidance covering both the long-term repeated dose toxicity
6 and the carcinogenicity study. The test is required for one rodent, the rat being the preferred
7 species. In exceptional cases and depending on the results obtained, testing in another
8 mammalian species (rodent or non-rodent, see also section 1.9.4 of this guidance for tests in
9 non-rodent species) may be considered.

10 **Generation of new test data**

11 If after the evaluation of available information further testing is needed to assess long-term
12 repeated dose toxicity, the test methods described further below should be used. Where new
13 testing is needed, please see also the general information under *Considerations before initiating*
14 *testing* in chapter 1.

15 The results of the long-term studies conducted and reported, taken together with other relevant
16 data and information on the active substance, should be sufficient to permit the identification of
17 effects, following repeated exposure to the active substance, and in particular should be
18 sufficient to:

- 19 • identify adverse effects resulting from long-term exposure to the active substance;
- 20 • identify target organs, where relevant;
- 21 • establish the dose-response relationship and mode of action;
- 22 • establish the NOAEL and, if necessary, other appropriate reference points.

23 Correspondingly, the results of the carcinogenicity studies taken together with other relevant
24 data and information on the active substance, should be sufficient to permit the evaluation of
25 hazards for humans to be assessed following repeated exposure to the active substance, and in
26 particular should be sufficient:

- 27 (a) to identify carcinogenic effects resulting from long-term exposure to the active
28 substance;
- 29 (b) to establish the species, sex, and organ specificity of tumours induced;
- 30 (c) to establish the dose-response relationship and mode of action;
- 31 (d) where possible, to identify the maximum dose eliciting no carcinogenic effect;

- 1 (e) where possible, to determine the mode of action and human relevance of any identified
2 carcinogenic response.
- 3 If comparative metabolism data indicate that either rat or mouse is an inappropriate model for
4 human cancer risk assessment, an alternative species should be considered.
- 5 Experimental data, including the elucidation of the possible mode of action involved and
6 relevance to humans, should be provided where the mode of action for carcinogenicity is
7 considered to be non-genotoxic. Suitable mode of action (MOA) studies can be considered to
8 confirm non-relevance of the non-genotoxic MOA to humans.
- 9 Investigation of toxicokinetic parameters generated within the combined long-term toxicity study
10 should also be considered as described also for short-term toxicity studies in section 1.9.2 of
11 this guidance.
- 12 The following test methods for long-term repeated dose toxicity should be used:
- 13 • EC method B.30 Chronic toxicity test.
 - 14 • EC method B.33 Combined chronic toxicity/carcinogenicity test.
 - 15 • OECD Test Guideline 452: Chronic Toxicity Studies.
 - 16 • OECD Test Guideline 453: Combined Chronic Toxicity/Carcinogenicity Studies.
17

18 **1.9.4. Further repeated dose studies (ADS)**

19 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.9.4 Further repeat dose studies</p> <p>Further repeat dose studies including testing on a second species (nonrodent), studies of longer duration or through a different route of administration shall be undertaken in case of:</p> <ul style="list-style-type: none"> — no other information on toxicity for a second non-rodent species is provided for, or — failure to identify a no observed adverse effect level (NOAEL) in the 28- or the 90-day study, unless the reason is that no effects have been observed at the limit dose, or — substances bearing positive structural alerts for effects for which the rat or mouse is an inappropriate or insensitive model, or — toxicity of particular concern | |

| | |
|--|--|
| <p>(e.g. serious/severe effects), or</p> <ul style="list-style-type: none">— indications of an effect for which the available data is inadequate for toxicological and/or risk characterisation. In such cases it may also be more appropriate to perform specific toxicological studies that are designed to investigate these effects (e.g. immunotoxicity, neurotoxicity, hormonal activity), or— concern regarding local effects for which a risk characterisation cannot be performed by route-to-route extrapolation, or— particular concern regarding exposure (e.g. use in biocidal products leading to exposure levels which are close to the toxicologically relevant dose levels), or— effects shown in substances with a clear relationship in molecular structure with the substance being studied were not detected in the 28- or the 90-day study, or— the route of administration used in the initial repeated dose study was inappropriate in relation to the expected route of human exposure and route-to-route extrapolation cannot be made. | |
|--|--|

1
2 When the available data are inadequate for hazard characterisation and risk assessment, further
3 repeated dose studies should be undertaken, including testing on a second species (non-rodent),
4 studies of longer duration than the studies already available or through a different route of
5 administration. However, testing should not be initiated before the evaluating competent
6 authority has indicated that further testing is necessary. The decision on further testing should
7 be based on expert judgement and on a case-by-case basis.

8 **Requiring further repeated dose toxicity studies**

9 When all the toxicological data concern rodent species, an assessment of the data needs to be
10 performed to understand if testing with another species is likely to provide additional information
11 (e.g. potential of different mode of action within different species).

12 Further studies are not necessarily always needed when failing to identify a NOAEL. If the data
13 are sufficient for a robust hazard assessment and for classification and labelling, the LOAEL may
14 be used as the starting point for risk assessment.

15 Where the preferred animal species is an inappropriate or insensitive model, a study protocol
16 will be identified that can be reliably performed in a more suitable animal species. It is however
17 possible to conclude that e.g. a structural alert concerns an effect that is specific to humans
18 and/or none of the animal models is suitable for studying this specific effect. In this case all the

1 available information, including scientific literature and human data, will be taken into account
2 to judge whether the risk to humans can be concluded. The human data may consist of e.g.
3 records of worker/consumer experience, case reports, consumer tests or epidemiological studies.
4 Whether further testing will be required will depend on a case-by-case expert judgment.

5 If toxicity of particular concern is already established, the substance will be classified accordingly
6 and the appropriate risk management measures will be implemented, and therefore no further
7 testing is required.

8 In some cases, data derived by protocols designed for other endpoints, as for example the OECD
9 Test Guideline 443 (Extended One-Generation Reproductive Toxicity Study) may provide
10 valuable information on specific effects such as immunotoxicity, neurotoxicity or endocrine
11 disruption. Furthermore, where a need is identified for a modification in the study protocol to
12 cover specific needs, this will be done in consultation with the evaluating competent authority.
13 Non-standard protocols should be used only in exceptional cases, because the scientific value of
14 such results can be questioned.

15 A new repeated dose toxicity study for the purpose of performing quantitative risk
16 characterisation for local effects should not be performed by default due to the difficulty in
17 deriving threshold levels for local effects that are also relevant for humans. The benefit from the
18 generation of additional data for this purpose should be considered against the effectiveness of
19 qualitative risk characterisation as another option for ensuring safe use.

20 Further studies might be necessary e.g. when the biocidal product is used in one or more
21 consumer products and the (combined) exposure levels are close to toxicologically relevant dose
22 levels where effects on humans may be expected in the relevant timeframe. Any exposure-
23 triggered studies proposed or required should be considered on a case-by-case basis.

24 Effects may have been observed in substances with a clear relationship in molecular structure
25 with the active substance, where such effects were not detected in the 28- or the 90-day study.
26 The study protocol and the conditions in which the effects were seen in another substance will
27 be examined in detail in order to identify the conditions in which the effect would be expected
28 to occur for the substance to be studied. The study protocol will be selected to repeat and
29 possibly extend the conditions where the effect has been observed. However, where applicable,
30 mechanistic *in vitro* studies examining the specific mechanism of action of the related substances
31 should have preference over further animal studies.

32 If the route of administration used in the initial repeated dose study was inappropriate in relation
33 to the expected route of human exposure, the possibility of route-to-route extrapolation should
34 be carefully considered before concluding that it is not appropriate, taking into account the
35 toxicokinetic information available and the use of modelling approaches when performing route-
36 to-route extrapolation.

37 1.10. Reproductive toxicity

38 Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| <p>8.10 Reproductive toxicity</p> <p>For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route</p> | <p>The studies do not need to be conducted if:</p> <ul style="list-style-type: none"> – the substance meets the criteria to be classified as a genotoxic carcinogen (classified both as germ cell mutagen category 2, 1A or 1B and carcinogenic category 1A or 1B), and appropriate risk management measures are implemented including measures related |

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| | <p>to reproductive toxicity,</p> <ul style="list-style-type: none">— the substance meets the criteria to be classified as a germ cell mutagen category 1A or 1B and appropriate risk management measures are implemented including measures related to reproductive toxicity,—the substance is of low toxicological activity (no evidence of toxicity seen in any of the tests available provided that the dataset is sufficiently comprehensive and informative), it can be proven from toxicokinetic data that no systemic absorption occurs via relevant routes of exposure (e.g. plasma or blood concentrations below detection limit using a sensitive method and absence of the substance and of metabolites of the substance in urine, bile or exhaled air) and the pattern of use indicates that there is no or negligible human or animal exposure,— the substance meets the criteria to be classified as reproductive toxicity category 1A or 1B: May damage fertility (H360F), and the available data are adequate to support a robust risk assessment, then no further testing for sexual function and fertility will be necessary. A full justification must be provided and documented if investigations for developmental toxicity are not conducted, or— the substance is known to cause developmental toxicity, meeting the criteria for classification as reproductive toxicity category 1A or 1B: May damage the unborn child (H360D), and the available data are adequate to support a robust risk assessment, then no further testing for developmental toxicity will be necessary. A full justification must be provided and documented if investigations for sexual function and fertility is not conducted. <p>Notwithstanding the provisions of this column of this row, studies on reproductive toxicity may need to be conducted to obtain information on endocrine disrupting properties as laid down in 8.13.3.1.</p> |
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Terminology used

The terminology explained in the Regulation (EC) No 1272/2008 on classification, labelling and packaging of substances and mixtures (CLP Regulation⁴) is used in this guidance.

For the purpose of classification and labelling, reproductive toxicity is divided into three differentiations; (i) adverse effects on sexual function and fertility), (ii) adverse effects on development of the offspring, and (iii) effects on or via lactation.

Adverse effects on sexual function and fertility include any effect of a substance that has the

⁴ Regulation (EC) No 1272/2008 of the European Parliament and of the Council

1 potential to interfere with sexual function and fertility. This includes, but is not limited to,
2 alterations to the female and male reproductive system, adverse effects on onset of puberty,
3 gamete production and transport, reproductive (oestrus) cycle normality, sexual behaviour,
4 fertility, gestation length, parturition, pregnancy outcomes, premature reproductive senescence,
5 or modifications in other functions that are dependent on the integrity of the reproductive
6 system.

7 Developmental toxicity includes, in its widest sense, any effect interfering with normal
8 development of the organism, before or after birth and resulting from exposure of either parent
9 prior to conception, or exposure of the developing organism during prenatal development, or
10 postnatally to the time of sexual maturation. However, these effects can be manifested at any
11 point in the life span of the organism.

12 The major manifestations of developmental toxicity include (1) death of the developing
13 organism, (2) structural abnormality, (3) altered growth, and (4) functional deficiency.⁵

14 Developmental neurotoxicity and developmental immunotoxicity belong also under
15 developmental toxicity.

16 Adverse effects on sexual function and fertility of the offspring in adulthood can be of
17 developmental origin. Reproductive toxic effects that cannot be clearly assigned to either
18 impairment of sexual function and fertility or to developmental toxicity shall be classified as
19 reproductive toxicants (i.e. Repr. 1A; H360, Repr. 1B; H360 or Repr. 2; H361) without the
20 specification (F/f and or D/d) in the hazard statement (CLP 3.7.1.1).

21 Objectives

22
23 It is important that the hazardous properties and risks or lack of them with respect to
24 reproduction are concluded for active substances. The information requirements have three core
25 objectives:

- 26 • to have adequate information to conclude whether classification and labelling for adverse
27 effects on sexual function and fertility and on development is warranted or can be with
28 sufficient confidence excluded (e.g. by ensuring that sufficiently high dose levels have
29 been tested);
- 30 • to have sufficient information for the purpose of risk assessment;
- 31 • to obtain information on endocrine activity/endocrine disrupting properties.

32 The results from reproductive toxicity studies should allow identification of specific adverse
33 effects on reproduction for classification and labelling, identification of endocrine activity of the
34 active substance, and derivation of points of departure for both reproductive toxicity and non-
35 reproductive toxicity for risk assessment purposes.

36 In more detail, the results from required reproductive toxicity studies (and study summaries
37 with numerical results) should be sufficient to:

38 (a) To identify and assess any specific effect on sexual function and fertility in P0 and/or P1
39 generations

40 1) for classification and labelling

⁵ As written in 3.7.1.3 and 3.7.1.4 in Annex I to CLP (the definition for developmental toxicity is shortened here)

- 1 2) to establish NOAELs for sexual function and fertility (P0 and P1)
- 2 (b) To identify and assess any specific effect on development (observable during pre-, peri-
3 and postnatal periods, and including effects on developing nervous system) in F1 and/or
4 F2 generations
- 5 1) for classification and labelling
- 6 2) to establish NOAELs for development of offspring (F1 and F2)
- 7 (c) To identify and assess any non-reproductive toxicity in parental/maternal animals;
- 8 1) To assess the potential influence of other toxicity, i.e. non-reproductive toxicity
9 on reproductive toxicity, when reproductive toxicity co-occurs with other toxicity
10 in order to conclude on the specificity of observed effects on reproduction;
- 11 i. Effects on reproductive toxicity (sexual function and fertility and/or
12 development) which occur even in the presence of other toxicity are
13 considered evidence of reproductive toxicity unless it can be unequivocally
14 demonstrated or it is reasonable to assume that the reproductive effects
15 are solely secondary non-specific consequences of other toxicity (CLP).
- 16 2) To identify the lowest effective dose level and the NOAEL for non-reproductive
17 toxicity (some non-reproductive adverse effects may occur at lower doses than in
18 other repeated dose toxicity studies with similar exposure duration); e.g.
19 pregnant/lactating females may be more sensitive to certain effects as compared
20 to non-pregnant animals (different or enhanced effects).
- 21 3) To assess if such effects warrant or contribute to the classification for other hazard
22 class(es) such as STOT RE.
- 23 (d) To identify and assess effects related to endocrine activity in parental animals and
24 offspring that can contribute to identification of endocrine disrupters.

25 This guidance provides advice on how the applicant can address the reproductive toxicity of the
26 active substance and how the information requirements of BPR can be met, thereby providing
27 data on the hazardous properties for classification purposes and for the risk assessment and
28 endocrine activity.

29 **Fulfilling the data requirement**

30 Effects accentuated over generations should be reported.

31 **Steps 1 and 2 Collection and evaluation of available information**

32 For the assessment of existing information on the reproductive toxic properties of the substance
33 all the relevant information should be considered together (physicochemical properties,
34 grouping, (Q)SARs and expert systems, *in vitro* data, human data and animal data) please
35 consult the CLP Regulation Title II. Further guidance is available within the *BPR Volume III*
36 *Human health Parts B+C* and the *Guidance on the Application of the CLP Criteria*.

37 **Step 3 Generation of new test data**

38 If after the analysis in steps 1 and 2 above, further testing is needed to assess reproductive
39 toxicity, the test methods described in section 1.10 should be used. Core information
40 requirements include extended one-generation reproductive toxicity study (OECD TG 443) with
41 the extension of Cohort 1B to provide mainly information on effects on sexual function and

1 fertility, developmental toxicity observable peri- and postnatally and sometimes on effects on or
2 via lactation. Prenatal developmental toxicity studies (OECD TG 414) in two species provide
3 information mainly on effects interfering with normal development before birth. Furthermore,
4 information on developmental neurotoxicity (e.g. OECD TG 426) is required. If there are specific
5 concerns that are not addressed by the standard information requirements, additional testing
6 might be needed to produce necessary information for hazard identification (classification and
7 labelling) and risk management (including risk characterisation, other risk management
8 measures), or to conclude on the ED properties (see chapter 8.13.3).

9 Where new testing is needed, please see also the general information under *Considerations*
10 *before initiating testing* in chapter 1.

11 Information requirements can also be fulfilled by adaptations that reduce the requirement for
12 testing. Adaptation possibilities are specified in Column 3 of the information requirement or in
13 BPR Annex IV.

14 *Preliminary considerations*

15 When planning any reproductive toxicity studies, considerations such as the properties of the
16 test item, dose levels, vehicle, adequate study design, and animal species and strain, are
17 needed. Some of the most relevant considerations are presented below.

18 (i) Dose range-finding studies

19 The dose range-finding studies should be reported as separate study records (in IUCLID) to
20 provide sufficient information and justification for the doses selected for testing. The findings
21 from a range-finding study may also support the interpretation of the results from the main
22 study.

23 (ii) Selection of vehicle

24 Most of the test methods provide guidance on vehicle selection if that is needed. If other vehicles
25 than water is used, a justification is needed. The vehicle itself should not cause any adverse
26 effects, as that may interfere with the interpretation of the results and may invalidate the study.
27 The vehicle must not react with the substance or interfere with toxicokinetics of the substance
28 or affect significantly the nutritional status of the animals. The control group should receive the
29 same vehicle and at the same dosing volume as the treated groups.

30 (iii) Route of administration

31 BPR information requirements specify that for evaluation of consumer safety of active substances
32 that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route. The
33 selection of the route of administration focuses on identification of hazards (see the Introduction
34 to this Guidance and REACH Guidance R7a sub-section "Selection of the appropriate route of
35 administration for toxicity testing", under R.7.2 Human health properties or hazards) and
36 depends on the most appropriate route for identification of the intrinsic properties of the
37 substance.

38 According to the test methods for reproductive toxicity, the oral route (gavage, in diet, or in
39 drinking water) is the default route, except for gases. For the extended one-generation
40 reproductive toxicity study (EU B.56, OECD TG 443) dietary administration may be an
41 appropriate route to model human exposure. If another route of administration other than oral
42 is used, a robust justification is required. In practice, testing via the oral route is usually
43 performed with solids, liquids and dusts, while testing via inhalation route is usually performed
44 with gases and liquids with very high vapour pressure. Testing via dermal route might be
45 necessary under specific circumstances, for example for substances with high dermal penetration
46 and indications for a specific toxicity following dermal absorption. Dermal application or

1 inhalation by nose-only administration may need specific considerations to ensure that the
2 administration can be adequately conducted without causing confounding factors, such as
3 additional stress to the pregnant animals. During lactation, separating the dams from the pups
4 for 6 hours for whole body exposure might induce additional stress on the pups that might lead
5 to the observation of effects that are not necessarily test-item related. Deviations from the
6 default oral route of administration must be justified, such as having information on route-
7 specific toxicity or toxicokinetics indicating that oral administration would not be relevant for
8 assessing the human health hazards via inhalation, which would be the main route of foreseen
9 human exposure.

10 *In vivo* testing at concentration/dose levels causing corrosivity must be avoided. For irritating
11 substances, the vehicle should be chosen to minimise gastrointestinal irritation. For some
12 substances, dietary administration may allow adequate dosing without irritation compared with
13 administration via gavage. In certain cases, irritation/corrosivity may be avoided by testing of
14 neutral salts of alkaline or acidic substances in order to allow investigation of intrinsic properties
15 at adequate dose levels. If immediate hydrolysis of a substance occurs, it may be possible to
16 provide information on all the cleavage products. Such a read-across approach should be
17 adequately justified and documented according to BPR Annex IV, 1.5 and applying the principles
18 of Read-Across Assessment Framework, RAAF⁶. For corrosive or irritating vapours or gases for
19 which oral testing is not possible, the highest concentration for inhalation should be chosen
20 carefully maximising the toxicity while minimising the irritation.

21 Gavage dosing provides accurate information on dose levels, and the resulting toxicokinetics
22 follow generally daily bolus dosing with high maximum concentration in blood (C_{max}) and,
23 depending on the elimination rate, daily periods with essentially no exposure are possible.
24 Toxicity requiring high C_{max} values can be observed.

25 Using dietary or drinking water route of administration provides less accurate information on
26 dose levels due to loss of material due to spilling. On the other hand, the blood levels are steadier
27 for many hours due to distribution of feed and water consumption during the day. Toxicity
28 requiring longer effect levels per day are more easily observed. Studies involving routes of
29 administration that are not relevant exposure routes for active ingredients (e.g. intravenous or
30 intraperitoneal injection), and resulting in unrealistically high exposure levels or eliciting local
31 damage to the reproductive organs must be interpreted with extreme caution and on their own
32 are not normally the basis for hazard classification or risk assessment. However, they may
33 provide information on mechanisms/modes of action.

34 (iv) Selection of species

35 The most common species for reproductive toxicity testing is the rat. There is often good
36 historical background information for various rat strains that may be used to support the
37 interpretation of the results. The strain selected should have an adequate fecundity and not too
38 high incidence of spontaneous malformations or any other specific feature that may reduce the
39 adequacy of the strain to study reproductive toxicity of the active substance. To facilitate
40 integrated data interpretation together with other studies, it is recommended to use the same
41 (rat) strain in reproductive toxicity testing and repeated dose toxicity studies.

42 If there is information regarding the sensitivity of the species and strains, the most sensitive
43 species and strain should be used, taking into account human relevance. There is no need to
44 demonstrate the human relevance; human relevance is assumed unless demonstrated
45 otherwise. In choosing the appropriate species and strain, consideration must be given to the
46 suitability of the species and strain for the test protocol, and the availability of background
47 information on the species and strain for the test protocol. The species/strain selection should

⁶ https://echa.europa.eu/documents/10162/13628/raaf_en.pdf

- 1 be justified if the default species referred to in a test method is not used.
- 2 More information on species selection for prenatal developmental toxicity studies is given in
3 section 1.10.1.
- 4 (v) Dose level selection
- 5 The dose level selection should ensure data generation for classification and labelling, risk
6 assessment, and identification of endocrine disrupting properties.
- 7 The dose levels should be spaced to produce a gradation of toxic effects. Generally, at least
8 three dose levels and a concurrent control must be used, except where a limit test (1000 mg/kg
9 bw/day) does not produce observable toxicity. Expected human exposure may indicate the need
10 to use a dose level above 1000 mg/kg bw/day⁷. The conditions for applicability of a limit test are
11 provided in the individual test methods for reproductive toxicity. For inhalation exposure, OECD
12 guidance document 39 may be used.
- 13 In selecting dose levels, information should be considered from existing studies, as well as from
14 any dose range-finding studies that may need to be conducted. Toxicokinetic information may
15 provide reasons to adjust for example the dosing route and regime. Furthermore, toxicity and
16 toxicokinetics in pregnant animals may differ from those in non-pregnant animals. This may
17 cause challenges in selecting the highest dose level for the study, because the sensitivity of the
18 animals may differ at various phases of the study.
- 19 It is important to get information about the reproductive toxicity profile of a substance including
20 the spectrum of reproductive toxicity effects related to different dose levels as well as information
21 to allow evaluation of the severity of reproductive toxicity of a substance.
- 22 The highest dose level should be intended to produce sufficient toxicity to provide adequate
23 information on reproductive toxicity for the purpose of both classification and labelling (including
24 categorisation), risk assessment and identification of endocrine activity. For classification and
25 labelling it is important that the tested doses are sufficiently high to enable a conclusion on a
26 lack of reproductive toxic properties warranting a classification in Repr. 1B or Repr. 2 if clear
27 evidence warranting a category 1B on reproductive toxicity is not observed (see the CLP criteria).
28 Therefore, the top dose selection should demonstrate an aim to induce clear evidence of
29 reproductive toxicity (adverse effects on reproduction) without excessive toxicity and severe
30 suffering in parental animals (e.g. prostration, severe inappetence, excessive mortality) that
31 would compromise the interpretation of reproductive effects.
- 32 There are aspects to be considered in the dose level setting of OECD TG 414, 443 and 426.
33 Common to all these TGs is that the lowest dose should not produce any evidence of either
34 maternal or developmental toxicity (and allow to set the NOAEL). Dose level selection should

⁷ CLP, Annex I, Sections 3.7.2.5.7 –3.7.2.5.9 state on the limit dose and very high dose levels the following: "There is general agreement about the concept of a limit dose, above which the production of an adverse effect is considered to be outside the criteria which lead to classification, but not regarding the inclusion within the criteria of a specific dose as a limit dose. However, some guidelines for test methods, specify a limit dose, others qualify the limit dose with a statement that higher doses may be necessary if anticipated human exposure is sufficiently high that an adequate margin of exposure is not achieved. Also due to species differences in toxicokinetics, establishing a specific limit dose may not be adequate for situations where humans are more sensitive than the animal model." Section 3.7.2.5.8: "In principle, adverse effects on reproduction seen only at very high dose levels in animal studies (for example doses that induce prostration, severe inappetence, extensive mortality) would not normally lead to classification, unless other information is available, e.g. toxicokinetics information indicating that humans may be more susceptible than animals, to suggest that classification is appropriate. Please also refer to the section on maternal toxicity (3.7.2.4) for further criteria in this area." And section 3.7.2.5.9 continues: "However, specification of an actual 'limit dose' will depend upon test method that has been employed to provide the test results, e.g. in the OECD Test Guideline for repeated dose toxicity studies by oral route, an upper dose of 1000 mg/kg has been recommended as a limit dose, unless expected human response indicates the need for a higher dose level."

1 also demonstrate any dose response meaning that the mid dose level should produce minimal
2 observable toxic effects. However, there are some differences in the specifications for the top
3 dose level (see below). Irrespective of the specifications in OECD TGs regarding selection of the
4 top dose, for classification and labelling, as explained above, it is critical that the tested doses
5 are sufficiently high to enable a conclusion on a lack of reproductive toxic properties warranting
6 a classification in Repr. 1B or Repr. 2 if clear evidence on reproductive toxicity is not observed.

7 The OECD TGs 414 main specification for top dose:

- 8 • *“the highest dose should be chosen with the aim to induce some developmental*
9 *and/or maternal toxicity (clinical signs or a decrease in body weight) but not death or*
10 *severe suffering”*

11 The specifications in OECD TG 426 for top dose selection:

- 12 • *“the highest dose level should be chosen with the aim to induce some maternal*
13 *toxicity (e.g., clinical signs, decreased body weight gain [not more than 10%] and/or*
14 *evidence of dose-limiting toxicity in a target organ)”*

- 15 • *“the highest dose should be the maximum dose which will not induce excessive*
16 *offspring toxicity, or in utero or neonatal death or malformations, sufficient to*
17 *preclude a meaningful evaluation of neurotoxicity.”*

18 For the OECD TG 443, the highest dose level should be based on toxicity (adverse effects) and
19 selected with the aim to induce reproductive and/or other systemic toxicity, as stated in column
20 1 of the information requirement.

21 The top dose selection should not only follow the specifications in OECD TGs but also take into
22 account the applicability for classification and labelling purposes.

23 There is a need to study various aspects in parents and their offspring in OECD TG 443. The
24 study should be designed to ensure adequate assessment of the effects on sexual function and
25 fertility, i.e. the dose levels should not be reduced in order to get a sufficient number of offspring
26 for the assessment of developmental toxicity. Even if the amount of offspring is reduced due to
27 effects on sexual function and fertility, any offspring available at that those level should be
28 investigated for adverse effects on development. Also results at lower dose levels can still be
29 used to assess if showing adverse effects on development.

30 It is also important that toxicity in both female and male animals is seen, to ensure that
31 reproductive toxicity in either gender is not overlooked. If existing information, including results
32 from a dose-range finding study, show that the sensitivity between male and female animals
33 differs significantly, the dose setting should take these differences into account. The less
34 sensitive sex should be tested at higher doses than the more sensitive sex.

35 For all of the TGs, the aim to have appropriate dose level setting has to be demonstrated.

36 Dose level selection must be justified and documented to allow independent evaluation of the
37 choice made.

38 **Considerations on mechanisms or modes of action**

39 There is no requirement to investigate the mechanism or MoA and its relevance to humans in
40 order to classify for reproductive toxicity. Only if it is conclusively demonstrated that the clearly
41 identified mechanism or mode of action has no relevance for humans and other mechanisms or
42 MoAs can be excluded, a substance that produces the adverse effects on reproductive toxicity
43 only in experimental animals shall not be classified. Classification in category 2 may be more
44 appropriate than category 1B when *mechanistic information* raises doubt about relevance in

- 1 humans, as far as there is reassurance about the robustness and quality of the data.
- 2 Some reproductive effects may be mediated via specific maternally mediated mechanisms (e.g.,
3 reproductive effects due to chelating MoA) that may still be specific effects on reproduction and
4 shall not be dismissed from classification for reproductive toxicity due to specific maternally
5 mediated mechanism.
- 6 Information on mechanisms and modes of action are relevant for ED identification. Mechanistic
7 information may also indicate a specific concern that may help identifying the most specific tests
8 for e.g. associative learning and memory under DNT (see 1.10.3).

9 **1.10.1. Pre-natal development toxicity study (OECD TG 414) on two species**

10 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|--|
| 8.10.1 Pre-natal development toxicity study (OECD TG 414) on two species, preferred first species is rabbit (non- rodent) and preferred second species is rat (rodent); oral route of administration is the preferred route | The study on the second species shall not be conducted if the study performed on the first species or other available data indicate that the substance causes developmental toxicity meeting the criteria for classification as toxic for reproduction category 1A or 1B: May damage the unborn child (H360D), and the available data are adequate to support a robust risk assessment |

- 11
- 12 The prenatal developmental toxicity studies, taken together with other relevant data and
13 information on the active substance (e.g. the developmental parameters of the EOGRTS and
14 OECD TG 426), must be sufficient to permit the assessment of potential hazardous properties
15 and risks on the offspring following exposure to the active substance during the development.
- 16 The prenatal developmental toxicity study (EU B.31, OECD TG 414) provides a focused
17 evaluation of potential effects on prenatal development, although only effects that are
18 manifested before birth can be detected. Detailed information on external, skeletal and visceral
19 malformations and variations and other prenatal developmental effects are provided. Cesarean
20 section allows precise evaluation of the number of fetuses affected.
- 21 Prenatal developmental toxicity should be determined in two species by the oral route. The
22 information requirement indicates rabbit and rat as the preferred non-rodent and rodent species,
23 respectively (also in accordance with the test method EU B.31 / OECD TG 414). Information on
24 two species allows a comprehensive assessment of prenatal developmental toxicity. If there is
25 information regarding the sensitivity of the species and strains, the most sensitive species and
26 strain should be tested first, taking into account human relevance.
- 27 The prenatal developmental toxicity study in a second species can be omitted if the information
28 already warrants classification as toxic for reproduction category 1A or 1B for development and
29 the available data are adequate to support a robust risk assessment.
- 30 The rabbit is the preferred species for the first prenatal developmental toxicity study. Selecting
31 rabbit as the first species may be supported by arguments of being a more sensitive species
32 than the rat for the specific active substance.
- 33 On the other hand, most toxicity studies are conducted in the rat, and it may therefore be
34 considered that the first prenatal developmental toxicity study could also be conducted in this
35 species. Findings from previous studies can be used in dose selection, or the identification of

1 additional parameters for evaluation. In addition, the outcome of the prenatal developmental
2 toxicity study may be helpful in the interpretation of other reproductive toxicity studies, for which
3 the rat is generally the preferred species.

4 If one or both of the default species (rat and rabbit) are not suitable for prenatal developmental
5 toxicity testing, a more suitable species considering the human relevancy should be selected for
6 testing. An adequate justification must be provided for species other than the rat and the rabbit.
7 The results from prenatal developmental toxicity studies are considered relevant to humans
8 unless there is substance-specific toxicokinetic or toxicodynamic evidence showing otherwise.

9 Information on prenatal developmental toxicity coming from one- or multigeneration studies
10 (such as OECD TGs 443, 416, 426, 421, 422) is not equivalent to that from the prenatal
11 developmental toxicity study. The results from e.g. OECD TG 443 and 416 studies do not provide
12 confidence to conclude that there is no prenatal developmental toxicity. Structural malformations
13 and variations are not specifically investigated in one- and multigeneration studies. Therefore
14 information from one- or multigeneration studies do not cover the information on prenatal
15 developmental toxicity in rodent species. However, in addition to information on prenatal
16 developmental toxicity in two species, information on effects due to exposure during peri- and
17 postnatal developmental periods that is obtained from one- or multigeneration studies (e.g.
18 OECD TG 426, 443 and 426) is also relevant for developmental hazard identification and shall
19 be assessed to conclude on classification and labelling for developmental toxicity (CLP 3.7.1.4).

20 The latest update of the following test methods for pre-natal developmental toxicity should be
21 used:

- 22 • Prenatal developmental toxicity study (OECD TG 414, EU B.31).

23 Information on developmental toxicity observable during peri-postnatal period can be obtained
24 from:

- 25 • Developmental neurotoxicity study (OECD TG 426; EU B.53).
- 26 • Extended one-generation reproductive toxicity study (OECD TG 443, EU B.56).
- 27 • Two-generation reproductive toxicity study (OECD TG 416; EU B.35).

28 **Note regarding pre-natal developmental toxicity studies and assessment of endocrine** 29 **disruption:**

30 The studies for prenatal developmental toxicity may need to be conducted to clarify endocrine
31 activity of the substance. Conduct of the studies may be needed even if the classification
32 criteria for Repr 1B; H360D (adverse effects on development) are met.

33 OECD TG 414 has been updated with thyroid hormone and thyroid stimulating hormone
34 analysis in dams (T4, T3 and TSH) and anogenital distance (by sex and related to weight) in
35 foetuses to be measured in rats. Some findings, such as increased foetal weight or placental
36 weight, considered together with litter size, may indicate an endocrine disrupting mode of
37 action.

38 OECD GD 150 describes OECD TG 414 as a level 4 studies (*in vivo* assays providing data on
39 adverse effects on endocrine-relevant endpoints). Modes of action which may produce a
40 diagnostic response includes EAST modalities. Parameters are sensitive also to R modality but
41 are not diagnostic of R modality.

42 **1.10.2. Extended One-Generation Reproductive Toxicity Study**

43 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| <p>8.10.2 Extended One-Generation Reproductive Toxicity Study (OECD TG 443), with cohorts 1A and 1B and extension of cohort 1B to include the F2 generation with the aim to produce 20 litters per dose group, F2 pups must be followed to weaning and investigated similarly as F1 pups. Rat is the preferred species and oral route of administration is the preferred route.</p> <p>The highest dose level should be based on toxicity and selected with the aim to induce reproductive and/or other systemic toxicity</p> | <p>A two-generation reproductive toxicity study conducted in accordance with OECD TG 416 (adopted 2001 or later) or equivalent information shall be considered appropriate to address this information requirement, if the study is available and was initiated before 15 April 2022.</p> |

1

2 The extended one-generation reproductive toxicity study (EOGRTS, EU B.56, OECD TG 443),
3 taken together with other relevant data and information on the active substance, must be
4 sufficient to permit the assessment of potential hazardous properties and risks on sexual function
5 and fertility, and development, following repeated exposure to the active substance. The study
6 also includes certain parameters for endocrine disrupting modes of action.

7 Information on blood concentration of the active substance in parents and foetus/offspring may
8 be included and reported to enhance interpretation of the results. Furthermore, the
9 concentrations of active substance and its relevant metabolites should be measured in milk,
10 although not required in the OECD test guideline, where adverse effects are observed in the
11 offspring or are expected due to effects on or via lactation (for example from a range-finding
12 study).

13 OECD TG 443 is a modular study design with various investigational options. For BPR, OECD TG
14 443 with extension of Cohort 1B is the information requirement. The extension of Cohort 1B to
15 mate the Cohort 1B animals and produce the F2 generation is also recommended in OECD GD
16 150 for the identification of endocrine disruptors. This extension provides information on sexual
17 function and fertility of the offspring of the P0 parental animals and developmental toxicity of
18 the second filial generation, and is important for the identification of endocrine activity.

19 Developmental neurotoxicity is a separate information requirement (section 1.10.3) and can be
20 fulfilled with an OECD TG 443 with Cohorts 2A and 2B and with additional investigation of
21 cognitive functions, as specified by the minimum requirements for developmental neurotoxicity
22 under section 1.10.3.

23 Information on developmental immunotoxicity belongs to additional data set, and in section
24 1.13.4, a common recommendation for a test battery is described which should be used to
25 address a concern for developmental immunotoxicity. OECD TG 443 with Cohort 3 can be
26 considered as a screening level information on developmental neurotoxicity which may need to
27 be followed with confirmative investigations (see further details in section 1.13.4).

28 Important considerations regarding the study conduct are explained below. These are not clearly
29 expressed in OECD TG 443 or OECD GD 151 and/or need to be specified to ensure data applicable
30 to hazard classification, risk assessment and identification of endocrine activity.

1 Premating exposure duration

2 To ensure that sexual function and fertility are adequately studied, a ten-week pre-mating
3 exposure duration is required in P0 animals. The sexual function and fertility part of the
4 reproductive toxicity study should be capable of providing information that is adequate for both
5 risk assessment and classification and labelling, including categorisation. For the comprehensive
6 assessment of effects and for the classification and labelling purpose, it is important to produce
7 and evaluate the full spectrum of effects on sexual function and fertility. The pre-mating exposure
8 period must be sufficiently long to be able to provide full information on magnitudes, incidences,
9 severities and types of all effects (MIST information) to be assessed together, not only aiming
10 to detect the most sensitive adverse effects. The most conclusive outcome can be obtained when
11 mating is allowed after an exposure covering one full spermatogenic cycle (including sperm
12 maturation) and folliculogenesis, and an analysis of sperm parameters, organ weights and
13 histopathology of gonads and accessory sex organs are conducted around the same time after
14 the same exposure history. The full spermatogenesis, without sperm maturation, takes 48-53
15 days in rats, (e.g. Kerr *et al.*, 2006). After spermatogenesis, sperm maturation in rats takes
16 around two weeks in epididymides. When the pre-mating exposure is 10 weeks, it covers the
17 sperm development through all the stages. A ten-week pre-mating exposure duration covers the
18 full spermatogenesis and maturation meaning that the full cycle of development of sperm from
19 spermatogonia into mature sperm is exposed. Thus, a ten-week pre-mating exposure duration
20 allows an assessment of the adverse effects on male sexual function and fertility by combining
21 the information from all possible parameters in males evaluated at the same time.

22 Regarding females, fixed number of primordial follicles are endowed during early life and growth
23 of these dormant follicles is initiated before and throughout reproductive life. Duration of follicle
24 development from initial recruitment of a primordial follicle until cyclic recruitment into
25 preovulatory follicles takes 61 days in rats (e.g., McGee and Hsueh, 2000). This follicle
26 development is fully covered only after a sufficiently long exposure period, such as ten weeks.
27 Therefore, for both the P0 males and females, a ten-week pre-mating exposure duration is
28 required before mating.

29 The data on F1 generation provides the most conclusive information for sexual function and
30 fertility because the primordial germ cells develop, migrate and proliferate during embryonic
31 development and effects to these events can be investigated only when the animals are exposed
32 already *in utero*. Furthermore, the exposure period in F1 generation covers also the postnatal
33 period before sexual maturation. Therefore, information also on potential effects by exposure
34 during the developmental period on sexual function and fertility is obtained from F1 animals.
35 This full evaluation is possible as the mating and littering of the Cohort 1B animals is required in
36 an extended one-generation reproductive toxicity study (EU B.56, OECD TG 443).

37 It is important to expose all the developmental stages of the sperm and follicles before the
38 mating in order to be able to detect any potential adverse effect on sexual function and fertility.
39 Furthermore, a 10-week pre-mating exposure duration supports interpretation of results when
40 effects in P0/F1 generations and compared to those of P1/F2.

41 To allow the ten-week pre-mating period, the exposure can be started when the animals are
42 around 5 weeks old and mate them around 15 weeks of age.

43 *Number of litters produced*

44 The number of males and females mated should aim to produce 20 litters for both generations.
45 Typically, 24 or 25 males and females are used to aim at producing 20 litters.

46 *Investigating F1 and F2*

47 The F2 pups must be followed to weaning and investigated similarly as F1 pups. Termination
48 should take place at weaning (around post-natal day 20 or 21). By comparing effects and effect

1 levels between F1 and F2, it can be deduced if developmental effects are observed at lower
2 doses (indicating a higher sensitivity) in F2 compared to F1. Effects that are observed in filial
3 generations only and/or there is an increase in sensitivity in filial generation(s) is a strong
4 indication that the effects are developmental (see also CLP 3.7.1.4; developmental effects can
5 be manifested at any point in the life span of the organism).

6 All investigations required for F1 pups should be also performed for F2 pups until weaning. These
7 include:

- 8 • general observations (all signs of toxicity, morbidity, mortality),
- 9 • body weight,
- 10 • clinical observations (changes in skin, fur, eyes, mucous membranes, occurrence of
11 secretions and excretions, abnormalities of genital organs e.g. hypospadias or cleft
12 penis),
- 13 • clinical examination of the neonates, e.g. qualitative assessment of body temperature,
- 14 • state of activity and reaction to handling,
- 15 • litter examination/parameters including number and sex of pups, stillbirths and live
16 births,
- 17 • litter examination/parameters including presence of gross anomalies (externally visible
18 abnormalities, including cleft palate; subcutaneous haemorrhages; abnormal skin colour
19 or texture; presence of umbilical cord; lack of milk in stomach; presence of dried
20 secretions),
- 21 • anogenital distance in pups (preferred: relative to square root of body weight),
- 22 • presence and number of nipples/areolae in male pups (see OECD GD 151, Section 3).
- 23 • Macroscopic examination of all organs for abnormalities
- 24 • Retention for possible histopathology: mammary tissue and other organs as appropriate

25 Furthermore, from surplus F1 pups at weaning and from F2 pups, body weight is recorded and
26 macroscopic abnormalities investigated from all organs. The following organs are weighed: brain,
27 spleen, thymus and other organs as appropriate and these and mammary tissues are kept for
28 possible histopathology.

29 *(Developmental) neurotoxicity*

30 Required minimum investigations on developmental neurotoxicity are specified in section 1.10.3.
31 OECD TG 443 with Cohorts 2A and 2B and with additional investigation of cognitive functions
32 can fulfil these minimum requirements. However, even without the specific cohorts for
33 developmental neurotoxicity (Cohorts 2A and 2B), some parameters of (developmental)
34 neurotoxicity are investigated in P0, Cohort 1A, F1 pups, P1 (extension of Cohort 1B) as well as
35 F2 pups up to weaning and/or surplus pups. These comprise of:

- 36 • general observations on behavioural changes,
- 37 • clinical observations on autonomic activity (e.g., lacrimation, piloerection, pupil size,
38 unusual respiratory pattern),
- 39 • changes in gait, posture, response to handling,

- 1 • presence of clonic or tonic movements,
- 2 • stereotypy (e.g. excessive grooming, repetitive circling) or bizarre behaviour (e.g. self-
- 3 mutilation, walking backwards),
- 4 • clinical examination of the neonates, e.g. qualitative assessment of body temperature,
- 5 • state of activity and reaction to handling,
- 6 • brain weight and histopathology,
- 7 • histopathology of peripheral nerve, spinal cord and optic nerve,
- 8 • brain weight (F2 and surplus F1 pups)
- 9 • Thyroid hormones (T4 and TSH) (F2 and surplus F1 pups) (MoA).

10 Results on these parameters in the offspring should be assessed along with the information
11 described in 1.10.3 and the information in P0 shall be considered along with all other relevant
12 available information when considering the need for additional studies/investigations on adult
13 neurotoxicity (section yyyy).

14 *(Developmental) immunotoxicity*

15 Information on (developmental) immunotoxicity belongs to ADS. The developmental
16 immunotoxicity Cohort 3 in OECD TG 443 investigates primary IgM antibody response to a T cell
17 dependent antigen (immunization with antigen is part of the test). However, even without
18 specific cohort for developmental immunotoxicity (Cohort 3), some parameters of
19 (developmental) immunotoxicity are investigated in P0, Cohort 1A and F1/F2 pups up to weaning
20 and/or surplus pups. These comprise of:

- 21 • spleen weight and histopathology,
- 22 • thymus weight and histopathology,
- 23 • bone marrow histopathology,
- 24 • total and differential leukocyte count,
- 25 • splenic lymphocyte subpopulation analysis (CD4+ and CD8+ T lymphocytes, B
- 26 lymphocytes and NK cells) using one half of the spleen,
- 27 • weight of lymph nodes associated with and distant from the route of exposure,
- 28 • histopathology on the collected lymph nodes and bone marrow.

29 Results on these parameters should be carefully evaluated to inform on possible indications or
30 effects on (developmental) immunotoxicity. Possible concerns for (developmental)
31 immunotoxicity may need to be followed-up e.g., in investigations in adults or in a standalone
32 study for developmental immunotoxicity. Recommended parameters for a potential separate
33 developmental immunotoxicity study are presented in chapter 1.13.4.

34 In case the developmental immunotoxicity Cohort 3 is included to OECD TG 443 as a screening
35 investigation, it is important that this T cell dependent antibody response (TDAR) contains valid
36 positive and negative controls with sufficient number of reacting animals.

37 ***Two-generation reproduction toxicity study***

1 The two-generation reproductive toxicity study was a core information requirement for BPR until
2 the amendment of BPR Annex II⁸. Although the two-generation reproductive toxicity study
3 (OECD TG 416) lacks information on some parameters which are part of EU B.56 (OECD TG
4 443), it addresses the sexual function and fertility in two generations (P0 and F1). OECD TG 416
5 study or equivalent information is adequate instead of OECD TG 443 if the study is available and
6 was initiated before 15 April 2022 and is conducted in accordance with the version of OECD TG
7 416 adopted 2001 or later.

8 If the study is conducted, e.g., for other regulation, and was initiated after 15 April 2022, the
9 applicant may explore the possibilities to adapt the information requirement by substance
10 specific justifications according to BPR Annex IV. When considering the relevance of old
11 two(multi)-generation reproductive toxicity studies to address reproductive toxicity and ED,
12 these studies will be assessed in line with BPR Annex IV, 1.1.2 adaptation rules for existing
13 information. Thus, old existing non-guideline studies may fulfil the Column 1 core information
14 requirement or may serve as elements in a weight of evidence adaptation according to BPR
15 Annex IV, 1.2 to identify hazardous properties or support a category approach.

16 Where necessary for the assessment of the effects on reproduction and/or ED and as far as the
17 available information is not yet sufficient for concluding on classification and labelling for
18 reproductive toxicity, ED identification or NOAELs, supplementary studies/investigations may be
19 required to provide information on the lacking parameters and the possible mechanisms.

20 **Note regarding EOGRTS and assessment of endocrine disruption**

21 The EOGRTS is a Level 5 *in vivo* assay providing more comprehensive data on adverse effects
22 on endocrine-relevant endpoints over more extensive parts of the life cycle of the organism (see
23 OECD Guidance Document 150). OECD GD 150 recommends OECD TG 443 with extension of
24 Cohort 1B (to mate the Cohort 1B animals to produce F2 generation).

25 In particular, the EOGRTS includes investigations informing on oestrogenic, androgenic, thyroid-
26 related, and steroidogenesis-related activities. For example, the EOGRTS investigates endocrine-
27 sensitive parameters in parental animals and offspring, such as sexual function and fertility,
28 weights and histopathology of reproductive organs/ tissues (e.g. male and female reproductive
29 tissues/ organs, thyroid including thyroid hormone measurements, adrenals, pituitary),
30 anogenital distance, and developmental landmarks such as sexual maturation. Sexual
31 maturation should be investigated from 3 animals/sex/litter, from 20 litters per dose group.

32 **1.10.3. Developmental neurotoxicity**

33 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| 8.10.3 Developmental neurotoxicity Developmental Neurotoxicity Study in accordance with OECD TG 426, or any relevant study (set) providing equivalent information, or cohorts 2A and 2B of an Extended One-Generation Reproductive Toxicity study (OECD TG 443) with additional | The study shall not be conducted if the available data: <ul style="list-style-type: none"> — indicate that the substance causes developmental toxicity and meets the criteria to be classified as toxic for reproduction category 1A or 1B: May damage the unborn child (H360D), and — are adequate to support a robust risk assessment' |

⁸ Regulation (EU) 2021/525

| | |
|---------------------------------------|--|
| investigation for cognitive functions | |
|---------------------------------------|--|

1
2 The BPR data requirement describes three study options that can fulfil the information
3 requirement:

- 4 1. OECD TG 426,
5 2. Any relevant study (set) providing information equivalent to OECD TG 426, or
6 3. OECD TG 443 with Cohorts 2A and 2B and with additional investigation for cognitive
7 functions.

8 Investigations for developmental neurotoxicity in these three study options include tests for
9 clinical observations, motor activity, motor and sensory function and cognitive functions
10 (including associative learning and memory) as well as neuropathological examination and brain
11 weight. In this guidance, the tests or test types that are considered to constitute the minimum
12 requirements to fulfil the obligations to test developmental neurotoxicity (DNT) under BPR are
13 described.

14 The overview of the minimum requirements for DNT by performing OECD TG 426 or 443 is given
15 in Table 9 below. Fulfilling the information requirement by a study set equivalent to these
16 minimum requirements is also possible.

17 For fulfilling the minimum information requirements identified in Table 9, the following aspects
18 have been taken into account:

- 19 • The aim to investigate different nervous system functions in the most optimal manner
20 possible
21 • The minimum information requirements should be achievable with both OECD TG 426
22 and OECD TG 443 with additional investigations for cognitive functions and even if Cohort
23 3 (DIT) is included in OECD TG 443
24 • Examples of possible animal assignments described in OECD TG 426
25 • Different types of associative learning and memory tests should be performed in
26 adolescents and young adults (in OECD TG 426, OECD TG 443 or other study set), in
27 different animals at these two time points.

28 Alternative test methods (a battery of *in vitro* DNT assays) are not described because an OECD
29 guidance document for an integrated approach to testing and assessment (IATA) for DNT is still
30 under development. However, DNT *in vitro* testing battery is not considered as an option to fulfil
31 the minimum data requirements because it currently does not provide equivalent information to
32 the required minimum requirements in *in vivo* tests. Although results from *in vitro* studies
33 indicating DNT properties may strengthen the other available evidence on DNT, results from *in*
34 *vitro* studies showing no indication on DNT hazard do not allow concluding on DNT properties
35 due to limitations of *in vitro* studies as compared to information from *in vivo* studies. In addition,
36 *in vitro* information alone is currently not sufficient for classification and labelling in accordance
37 with the CLP Regulation. For further reading on DNT *in vitro* battery, see Sachana et al., 2021.

38 **Table 9. Minimum requirements of investigations and test types to detect DNT in OECD TG 426**
39 **and OECD TG 443.**

| INVESTIGATIONS IN F1 | OECD TG 426 | OECD TG 443 |
|----------------------|-------------|-------------|
|----------------------|-------------|-------------|

| GENERATION | | | | |
|--------------------------------|---|---|---|--|
| | Time point and minimum number of males and females per dose group* | Test method/test type | Time point, cohort and minimum number of males and females per dose group** | Test method/test type |
| Detailed clinical observations | Weekly during preweaning, at least every two weeks thereafter; (set 3: 20M+20F) | Reporting changes e.g. in autonomic activity (e.g., lacrimation, piloerection, pupil size, unusual respiratory pattern and/or mouth breathing, unusual urination or defecation), body position, activity level, gait, posture, reactivity to handling, placing or other environmental stimuli, clonic or tonic movements, convulsions, tremors, stereotypies, bizarre behaviour or aggression | Weekly, all F1 animals | Reporting occurrence of e.g. secretions and excretions and autonomic activity (e.g. lacrimation, piloerection, pupil size, unusual respiratory pattern), changes in gait, posture, response to handling, presence of clonic or tonic movements, stereotypy (e.g. excessive grooming, repetitive circling) or bizarre behaviour (e.g. self-mutilation, walking backwards) |
| FOB | - | - | PND 63- 75 (cohort 2A) | See Appendix A in OECD TG 443 |
| Brain weight | PND 22 (subset 1a: 10M+10F unfixed, and subset 1b: 10M+10F fixed) and at termination (PND 70) (at least subset 3a: 10M + 10F, subset 4a: 10M+10F) | | PND 21-22 (all surplus animals, cohort 2B, 10M+10F); PND 75-90 (cohort 2A, 10M+10F) | |
| Neuropathology and morphom | PND 22 (subset 1b: 10M+10F) and at | Staining of slices containing slices of olfactory bulbs, cerebral cortex, | PND 21-22 (cohort 2B, 10M+10F); PND 75-90 | 2A: Staining of slices containing slices of olfactory bulbs, cerebral cortex, hippocampus, |

| | | | | |
|--|--|---|---|---|
| etry | termination (PND 70) (subset 3a: 10M + 10F) | hippocampus, basal ganglia, thalamus, hypothalamus, midbrain (tectum, tegmentum, and cerebral peduncles), pons, medulla oblongata, cerebellum; spinal cord and the PNS at PND 70 only | (cohort 2A, 10M+10F) | basal ganglia, thalamus, hypothalamus, mid-brain (tectum, tegmentum, and cerebral peduncles), brain-stem and cerebellum, the eyes (retina and optic nerve), peripheral nerve, muscle and spinal cord*** |
| Motor activity (including habituation) | 1 or 3 (if tested for behavioural ontogeny) times during preweaning and Once on PND 60-70 (set 2: 20M + 20F) [in same animals at all time points] | Open-field test | Once on PND 63-75 (Cohort 2A, 10M + 10F) | Open-field test |
| Motor and sensory function | PND 25±2 (set 3: 20M+20F) and PND 60-70 (set 3: 20M+20F) | Acoustic startle with PPI and short-term habituation (PND 25±2) Grip strength and righting response (PND 60-70) | PND 24-25 (Cohort 2A, 10M+10F); PND63-75, (Cohort 2A,10M+10F) | Acoustic startle with PPI and short-term habituation (PND 24-25) Grip strength and righting response (part of FOB) (PND63-75) |
| Cognitive functions/ Learning and memory (L&M) | PND 25±2 (set 2: 20M+20F, same animals as in open-field test); PND 60-70 (set 4: 20M+20F); different animals at these two time points. | MWM or RAM (spatial explicit/allocentric L&M) at one time point and another type of associative L&M (implicit) test at the other time point, e.g., CWM | PND 25±2 (Cohort 1A [10M+10F] or Cohort 3 animals, if DIT investigations are not conducted, 10M+10F); PND 60-70 (Cohort 1A animals, 10M+10F) | MWM or RAM (spatial explicit/allocentric L&M) at one time point and another type of associative L&M (implicit) test at the other time point, e.g., CWM |
| Behavioural | At least two measures of at least 2 | Open-field as one test for behavioural ontogeny highly | - | - |

| | | | | |
|----------|--|---|--|--|
| ontogeny | behaviours during pre-weaning) (set 2: 20M+20F) | recommended. The other behaviour for behavioural ontogeny should not develop at the same age. | | |
|----------|--|---|--|--|

1 * The animal allocation for OECD TG 426 follows that of example 3 of the OECD TG 426 with 4 sets (divided to subset a
2 and b in some places) of 20 pups/sex/dose level (i.e. 1 male and 1 female per litter). Other animal allocations according
3 to OECD TG 426 are possible.

4 **In OECD TG 443 adverse effects on sexual function and fertility may limit the number of offspring available for
5 developmental investigations. However, the dosing should not be lowered in order to get a sufficient number of offspring.
6 The priority of the OECD TG 443 test is to identify potential effects on sexual function and fertility and if this effect leads
7 to an insufficient number of offspring, DNT should be investigated in OECD TG 426.

8 ***Histopathology of fixed peripheral nerve, spinal cord and eye (and optic nerve) is also performed in Cohort 1A(B)
9 offspring.

10 The sequence of tests should progress from the least invasive (e.g. observations in the home
11 cage and open field) to the most invasive (e.g. handling assessments) to minimize the influence
12 of stress on subsequent measures. Parameters that require descriptive measures should include
13 a clear description of what constitutes "other than normal", and ranking or scales describing
14 different severities of effects should be given. It is also recommended to include valid positive
15 controls if not already available for the laboratory and setup to ensure that the technical
16 personnel of the testing laboratory is able to correctly use the test procedures and animal model.
17 Those results should verify that the laboratory can effectively demonstrate effects that are
18 qualitatively and quantitatively consistent with those reported in other laboratories for the same
19 agent, at similar doses, and under comparable conditions. This outcome provides added
20 confidence that the absence of effects due to a treatment accurately reflects the situation rather
21 than being due to inadequate implementation of a valid test method (i.e., a false negative).
22 Positive control data also helps interpreting the results (Tyl et al., 2008).

23 The investigations are grouped below according to the main headings in OECD TG 426 into
24 physical and developmental landmarks and functional behavioural endpoints.

25 **Physical and developmental landmarks**

26 OECD TG 426 and 443 require the testing of physical and developmental landmarks, including
27 body weight, clinical observations, brain weight, neuropathology and sexual maturation, and
28 these should be investigated accordingly. Additional developmental landmarks (e.g. pinna
29 unfolding, eye opening and incisor eruption) can be optionally added as given in the OECD TG
30 426.

31 Effects on various parameters, even if not specific on the nervous system, may be relevant for
32 the interpretation of the effects on the nervous system. These include e.g. pup body weight,
33 morbidity, mortality, changes in skin, fur, eyes, mucous membranes, occurrence of some
34 secretions, unusual signs of urination or defecation, and sexual maturation.

35 Even when not specific for (developmental) neurotoxicity, physical and developmental landmarks
36 are relevant for the reproductive toxicity hazard assessment, both NOAEL/LOAEL determination
37 and classification and labelling. Adverse effects on development include death of the developing
38 organism, structural abnormalities, altered growth and functional deficiency, and adverse effects
39 on sexual function and fertility include effects on onset of puberty (sexual maturation) among
40 other effects (CLP 3.7.1.4 and 3.7.1.3).

41 **Clinical observations and FOB**

42 Clinical observations and functional observation battery (FOB) of the F1 generation should be
43 investigated according to OECD TG 426 or 443, depending on the selected TG.

1 Clinical observations required in OECD TG 426 and 443, and FOB required in OECD TG 443 are
2 often subjective evaluations, and therefore explicitly defined scores and criteria should be used.
3 Measures that are ranked provide more information than binary (all-or-nothing) measures. A
4 ranking or scale describing different levels of activity improve consistency across observers.
5 More details are given in NAFTA guidance (2016).

6 **Neuropathological examination**

7 Neuropathological evaluation and brain weight measurement of the offspring should be
8 conducted according OECD TG 426 or 443, depending on the selected TG.

9 All neuropathologic alterations should be assigned a subjective grade indicating severity and
10 their incidences should be reported. Cellular alterations (e.g., neuronal vacuolation,
11 degeneration, necrosis) and tissue changes (e.g. gliosis, leukocytic infiltration, cystic formation)
12 should be reported and assessed. Reporting should follow OECD GD 20 which requires that
13 ambiguous terminology should be avoided and the nomenclature used for describing lesions and
14 areas of the nervous system should follow standards and be as specific as possible. Further, the
15 cell types involved in the lesion should be described to the degree possible, and attention should
16 be paid to the distribution pattern of lesions, e.g. whether they are formed in bilateral and/or
17 symmetrical pattern.

18 A performance impairment detected in a behavioural test may not be reflected in outcomes from
19 brain pathology or brain morphometry, and vice versa. Behavioural effects may reflect e.g.
20 effects on specific ion channels or neurotransmitters affecting nerve cell communication and
21 such effects are not observed via standard histopathological staining procedures or
22 morphometry. However, when planning the test set for developmental neurotoxicity testing, if
23 there are already some neurohistopathological investigations (or any other information)
24 indicating effects on certain areas of the nervous system, one should ensure that the function of
25 these areas is specifically examined by selecting such behavioural test(s) from the possible
26 alternative tests that target the function of these structures.

27 **Functional/behavioural endpoints**

28 ***Behavioural ontogeny***

29 Behavioural ontogeny of the F1 generation should be investigated according OECD TG 426. In
30 OECD TG 443, testing behavioural ontogeny is not required, but it is recommended to test
31 behavioural ontogeny as in OECD TG 426.

32 According to OECD TG 426, ontogeny of at least two selected behaviours should be measured in
33 at least one pup/sex/litter during the appropriate age period (twice during pre-weaning), with
34 the same pups being used on all test days for all behaviours assessed. OECD TG 426 gives
35 righting reflex, negative geotaxis and motor activity as examples of behaviours for which their
36 ontogeny could be assessed, and the TG strongly recommends the use of motor activity to assess
37 behavioural ontogeny. If motor activity by open field is selected as one of the ontogeny
38 behaviours, it will be investigated three times during pre-weaning (see below the requirements
39 for motor activity). Generally, 20 males and 20 females per dose group should be selected for
40 investigations (1 pup per sex per litter).

41 ***Motor activity (including habituation)***

42 Motor activity of the F1 generation should be investigated according OECD TG 426 or 443,
43 depending on the selected TG and with the specifications below.

44 Motor activity should be monitored once (on PND 63-75 in Cohort 2A) according to OECD TG
45 443, and at least once during the pre-weaning and once on PND 60-70 according to OECD TG
46 426. If motor activity is tested for behavioural ontogeny, the test should be performed at least

1 three times during the pre-weaning period. In normal conditions locomotor capacity starts to
2 develop in rodents at around PND 13 and appears to be fully developed around PND 21 (NAFTA
3 guidance, 2016).

4 The OECD TG 426 or 443 does not specify the type of test arena for assessing motor activity,
5 other than that the motor activity testing must be conducted in automated test chambers. The
6 open-field test is the most suitable and therefore the required test for motor activity; it is widely
7 used to investigate hyper- or hypoactivity and habituation. As the open-field test may provide
8 also information on the anxiety-like behaviour, movements in central and peripheral parts should
9 be recorded and included in the analysis.

10 As an example, Qian et al. (2010) describe a methodology for measuring open field spontaneous
11 activity. The length of the test session should allow the detection of potential effects on motor
12 activity and on its habituation. It is necessary to determine precisely what measures are recorded
13 and reported. Since fine motor movements (e.g. sniffing, scratching, grooming) do not provide
14 a measure of locomotor or ambulatory activity, the activity test data should clearly distinguish
15 various activity measures and their types. Software settings for defining the type and threshold
16 for activity units can be critical for computing measures of ambulatory activity, and therefore
17 clear reporting on data collection and computation is necessary (e.g. recording instrumentation,
18 software versions and settings at each age). At least the distance traveled in the center,
19 periphery, and total (entire box), latency to enter the central area, as well as the number of
20 rearing activity should be recorded. Activity measures should be described broken down by dose
21 group, sex and 10-minute time blocks. Please see the NAFTA guidance (2016) for further
22 methodological and reporting aspects as well as the normal developmental stages in the
23 development of locomotor activity.

24 ***Motor and sensory function***

25 Minimum requirements for motor and sensory function of the F1 generation include testing for
26 auditory startle response with pre-pulse inhibition (PPI) and short-term habituation at least once
27 during adolescent period (PND 25±2 in OECD TG 426; or PND 24-25 in Cohort 2A in OECD TG
28 443), and grip strength and righting reflex in young adults (PND 60-70 in OECD TG 426; or as
29 part of FOB on PND 63-70 in Cohort 2A in OECD TG 443). Any other test set providing equivalent
30 information may also be used.

31 In the OECD TG 443 there is no heading for "motor and sensory function", but the tests
32 investigating these functions include the auditory startle test and grip strength and righting
33 reflex tests that are tested as part of the functional observational battery. The tests required in
34 this section for motor and sensory function in OECD TG 426 are based on the requirements
35 specified in OECD TG 443. The OECD TG 426 does not specify the required tests to ensure
36 "adequate quantitative sampling of sensory modalities and motor functions" but rather provides
37 a list of examples of tests (extensor thrust response, righting reflex, auditory startle habituation
38 and evoked potentials).

39 Additional tests for motor and sensory functions are recommended especially if there is a specific
40 concern for effects on some motor or sensory components that would not be adequately
41 addressed by this minimum set of tests. For example, cerebellar dysfunction often correlates
42 with abnormalities of gait synchronisation that can be sensitively measured by a rotating rod
43 (Lane and Dunnet, 2011, vol II).

44 Below is a short overview of the required specific tests that investigate different modalities of
45 sensory and motor functions and closely associated other key functions. The rotating rod test is
46 also summarised as it is recommended as an additional test.

47 *Acoustic startle test with pre-pulse inhibition (PPI) and short-term habituation*

48 Habituation (short- and/or long-term) and PPI of acoustic startle response can be measured

1 within one protocol. As it is a valuable predictive model for cognitive impairment, it is important
2 that PPI is included in the testing protocol for acoustic startle response in both OECD TG 426
3 and 443, although PPI is not specifically mentioned in OECD TG 443 as part of auditory startle
4 test, and it is reviewed only in the reference of OECD TG 426 (Koch, 1999). The PPI is a simple
5 addition to the acoustic startle and its short-term habituation test method. Acoustic startle and
6 its short-term habituation is a sensory-motor test involving only a short neural pathway, whereas
7 PPI adds a cognitive dimension to the test by predicting cognitive impairment involving a certain
8 limbic circuitry (cortico-striato-pallido-pontine), that converges with the primary startle circuit
9 in humans and rodents (Valsamis and Schmid, 2011).

10 Before measuring PPI, animals should always undergo startle habituation, so that startle
11 attenuations due to habituation do not interfere with PPI measurements. Before running the
12 habituation and PPI, the animal must adapt to the animal holder, startle box and background
13 noise via an acclimation period. A protocol design and data analysis is described in detail in
14 Valsamis and Schmid (2011).

15 *Grip strength*

16 The grip strength test measures the strength of limb flexor muscles in fore and hindlimbs that
17 are innervated by peripheral motor nerves. The test is a specified requirement as part of the
18 FOB in OECD TG 443, and it should be measured also as part of the sensory and motor testing
19 in OECD TG 426. The peak of the grip strength is measured by a grip strength meter for each
20 rat during five trials, separated by approximately 1 min between each trial, and the average is
21 used as the grip strength for each rat. Methodologies for measuring grip strength are well
22 established and protocols can be found e.g. in Torii *et al.*, 2010; Jeyasingham *et al* 2001.

23 *Righting reflex (postural reflex)*

24 Righting reflex is required as part of the FOB in OECD TG 443 and should be measured also as
25 part of the sensory and motor testing in OECD TG 426. Rats are momentarily held supine by the
26 shoulders and hip-girdle on a flat surface and released. Normal animals will immediately turn
27 over to recover their normal prone quadruped stance. The presence or absence of the reflex,
28 time taken and direction of response are noted and reported. The test should be performed three
29 times a day with an upper time limit of 3 min for each test. For each animal, the data for the
30 three tests are averaged. Normal animals will turn in either direction with equal frequency, but
31 they often turn away from the tester or a bright light source. It is therefore necessary to
32 randomly change the orientation in which the animals are held: head to the left for one test and
33 head to the right for another (Lane and Dunnet, 2011, vol II).

34 *Rotating rod*

35 Rotating rod test is not specifically required to fulfil the minimum requirements, but due to its
36 potential to detect e.g. basal ganglia and cerebellum dysfunctions, it is highly recommended to
37 be performed as part of the OECD TG 426 or OECD TG 443 (or any other study set providing
38 equivalent information). The rotarod is a horizontal cylinder that rotates about its long axis at
39 either constant or accelerating speeds. The animal is placed on the rotating cylinder
40 perpendicular to the direction of rotation facing away from the tester by allowing the animal to
41 walk off the open palm onto the rotating rod. In order to maintain position on top of the rod and
42 not fall off, it has to walk forwards synchronising stepping frequency and stride length to the
43 speed of rotation. For each trial, the parameters recorded are total time on the rod, time walking,
44 time spent in error (clinging or walking backwards) and time to first error (fall or cling). The trial
45 ends either when the animal falls or 180 s is reached (Lane and Dunnet, 2011, vol II).

46 Automatic time-to-fall is a useful measure for general motor ability, but it is not a sensitive
47 indicator of cerebellar function, which requires that the gait is synchronised to the speed of
48 rotation. Rodents undertake all possible alternative strategies they can to avoid falling, such as
49 clinging to the rod and being passively rotated or turning around, lying with their abdomen in

1 contact with the rod and shuffling backwards. All these alternative strategies may indicate
2 incorrect cerebellar function but will not be detected as "error" by automatic time-to-fall devices.
3 Therefore, it is important that the experimenter records also the additional parameters specified
4 above. An accelerating rotarod is quick and simple, but it is a less sensitive assessor of cerebellar
5 dysfunction than is the constant speed protocol (Lane and Dunnet, 2011, vol II).

6 ***Learning and memory (cognitive functions)***

7 Learning and memory of the F1 generation should be investigated with the specifications below.

8 The minimum information requirements for learning and memory, a component of cognitive
9 functions, include two different tests for associative learning and memory at two different time
10 points. Different test types of associative learning and memory should be performed at
11 adolescence (PND 25±2 days) and young adulthood (PND 60 and older). Different set of animals
12 is recommended to be used. In OECD TG 443, Cohort 1A animals can be allocated to two sets
13 of animals, 10 males and 10 females in both; the first set of animals to be tested at adolescence
14 and the other set of animals at young adulthood. If necessary, animals from other Cohorts (such
15 as Cohort 3 if not included to investigate developmental immunotoxicity) may be used also,
16 taking into account the integrity of the study. For OECD TG 426, the examples of the alternative
17 animal allocations can be followed. It is recommended to use more than 10 animals per sex if
18 possible, and e.g. example 3 of the OECD TG 426 may be used as the basis for animal allocation.

19 Two criteria for associative learning and memory tests are presented in paragraph 37 of OECD
20 TG 426 and these should be fulfilled also if the DNT is tested as part of OECD TG 443 or by other
21 means:

- 22 1) Learning should be assessed either as a change across several repeated learning trials or
23 sessions, or, in tests involving a single trial, with reference to a condition that controls
24 for non-associative effects of the training experience; and
- 25 2) The test(s) should include some measure of memory (short-term or long-term) in
26 addition to original learning (acquisition), in the presence of a measure of acquisition
27 obtained from the same test.

28 Different test types of associative learning and memory engage different brain regions,
29 combinations of regions and neural pathways. Different tests can have also different sensitivities
30 for observing effects on learning and memory. One of the required tests should investigate
31 explicit associative learning and memory and the other test should investigate implicit
32 associative learning and memory. Explicit memory (or declarative memory) is recalled
33 consciously whereas implicit memory (or nondeclarative memory) is recalled unconsciously
34 (Kandel, 2000).

35 Two examples of explicit associative learning and memory tests are the Morris water maze
36 (MWM) test and Radial arm maze (RAM) test, both investigating allocentric spatial learning and
37 memory. An example of one type of implicit associative learning and memory test is Cincinnati
38 water maze (CWM) which investigates egocentric navigational learning and memory. Allocentric
39 learning and memory in rodents is homologous to the same brain networks that in people
40 mediate memory for people, places, facts, and events. Egocentric navigation in rodents is
41 homologous to path finding and procedural learning and memory including skilled behaviours
42 such as driving a car and other highly trained behaviours that become semiautomatic in people.
43 However, the neural networks mediating egocentric and spatial navigation overlap despite partial
44 dissociations of the two systems. (Vorhees and Williams, 2015 and 2016).

45 Examples of other types of implicit associative learning and memory tests are classical and
46 operant/instrumental conditioning tests such as olfactory conditioning test, and acquisition and
47 retention of schedule-controlled behaviour. If there is any prior information indicating a need for
48 a specific test subtype, this should be used to select the most appropriate test. Although in OECD

1 TG 426 also the T-maze, Biel water maze and passive avoidance test are given as examples of
2 possible tests, these should not be selected because based on practical experience in regulatory
3 use, they have been suspected to be insensitive for detecting developmental neurotoxicants
4 (Levin, 2014; Vorhees and Williams, 2014; Vorhees and Makris, 2015). Below is an overview of
5 associative learning and memory tests that may be selected to fulfil the minimum information
6 requirements.

7 *Morris water maze (MWM)*

8 The MWM test studies allocentric spatial learning and memory that is a type of explicit learning
9 and memory. MWM test involves hippocampus, entorhinal cortex and surrounding structures.
10 The most basic MWM procedure tests allocentric learning and reference memory, but by an
11 appropriate modification of the basic protocol it is possible to study allocentric learning and
12 memory in more depth or with higher sensitivity or assess also other forms of learning and
13 memory. These variants of protocols are presented and the basic protocol is described (with
14 troubleshooting) in detail in Vorhees and Williams (2006).

15 The concept behind the basic MWM is that the animal must learn to use distal cues, such as
16 landmarks, to navigate a direct path to the hidden platform when started from different, random
17 locations around the perimeter of the tank. MWM is an open circular pool that is filled
18 approximately half-way with water. The interior is as featureless as possible, and the maze is
19 divided into four equal quadrants, and a relatively small hidden platform is positioned in the
20 middle of one of the quadrants below the water surface in a fixed location. The animal must
21 search in order to locate the hidden platform. The pool must be professionally constructed for
22 MWM to ensure that there are no proximal cues undermining the goal of the test. The correct
23 size of the tank is also one critical factor for obtaining valid spatial learning curve. Spatial learning
24 (spatial acquisition) is assessed across repeated trials (normally 4 trials per day, inter-trial
25 interval 15 s, repeated for 5-6 days) and reference memory (memory/probe trial) is determined
26 by the preference for the platform area when the platform is absent (animal placed in a novel
27 starting position to ensure that its spatial preference is a reflection of the memory of the goal
28 location rather than for a specific swim path, tested at least 24 h after the last learning trial,
29 trial length of 30 s recommended). Escape from water is relatively immune from motor activity
30 (e.g. on open field) or body mass differences, making it ideal for many experimental models. In
31 addition, the MWM has proven to be a robust and reliable test (Vorhees and Williams, 2006 and
32 2015).

33 *Radial arm maze (RAM)*

34 Similar to the MWM, the RAM test studies allocentric spatial learning and memory that is a type
35 of explicit learning and memory. However, the RAM test involves brain areas partly different
36 from the MWM (hippocampus, frontal cortex, mediodorsal thalamic nucleus, septum, amygdala
37 and mammillary bodies). RAM can be used with a variety of different procedures (reviewed e.g.
38 in Levin, 2014 and Vorhees and Williams, 2016). Typically, the RAM is used as an appetitive test.
39 In an eight arm RAM eight equally spaced arms extend from a central circular platform and four
40 of the eight arms are baited with a food reward. Over a course of successive daily test trials, the
41 rat is expected to learn which arms are baited (or never baited) and will efficiently retrieve the
42 food rewards at the ends of four baited arms by using visuospatial cues in the room. The
43 performance of the rat is measured by the time and distance to complete each trial, and by the
44 number the animal goes down a never baited arm between trials (reference memory error) or
45 re-entries into an arm it already visited within that trial (working memory error). RAM can be
46 also run with aversive (water escape) motivators. (Levin, 2014; Vorhees and Williams, 2016).

47 *Cincinnati water maze (CWM)*

48 The CWM test investigates egocentric navigational learning and memory that is a type of implicit
49 learning and memory. Dorsal striatum is considered as the key component in mediating
50 egocentric navigation. The CWM is an asymmetric 9-unit multiple-T labyrinthine maze that can

1 be used to test either egocentric (body-centered) navigation if tested under infrared lighting, or
2 combined allocentric and egocentric navigation if tested under standard light. In egocentric
3 navigation the animal uses internal and/or near (proximal) cues. Internal cues include
4 proprioceptive feedback from limb/joint receptors and stretch receptors in muscles and tendons
5 that provide a sense of speed of motion that, when combined with heading or directional
6 information and signposts about which way to turn, produce a pathway or route to and from
7 different locations. Signs or signposts are different from landmarks; a signpost is close whereas
8 a landmark is farther away from the organism. Although the CWM test run under the infrared
9 light provides the most stringent test of egocentric learning and memory and is more sensitive
10 than CWM test performed under standard light, the dark variant is more challenging and it takes
11 rats many trials over multiple days with multiple trial failures before learning the CWM to a
12 proficient level of performance. The extended length of the test, when used under infrared light,
13 may limit its applicability in a regulatory study (Vorhees and Williams, 2015 and 2016).

14 A day before the actual CWM test, whether using the standard light or infrared light procedure,
15 rats must be given acclimation trials consisting of a separate straight water channel under
16 standard light with a submerged platform at one end located in a different room than the maze.
17 If these acclimation trials are not given, rats will find the task too difficult, give up, and never
18 find the escape. The detailed test protocol is given e.g. in Vorhees and Williams (2016).

19 *Olfactory conditioning*

20 Olfactory fear conditioning test involves amygdala, the key structure for initiating and controlling
21 fear reactions, but also playing a role in coding for the biological significance, intensity, or
22 salience of sensory stimuli (Buettner [ed], 2017). In humans, dysregulation of function of
23 amygdala is associated with abnormally heightened fear such as in anxiety disorders (Hakim et
24 al., 2019; Buettner [ed], 2017). Examples of methods for olfactory fear conditioning are given
25 in Kucharski and Spear (1984) and Crofton *et al.* (1993).

26 Aversive olfactory conditioning is a specific form of classical conditioning, also known as
27 Pavlovian learning, that is a fundamental form of learning and expressed between and within
28 species. The principle of Pavlovian fear learning is that an unpleasant unconditioned stimulus
29 (US), such as foot shock, that produces a strong negative response, irrespective of training, gets
30 associated with a neutral cue, odor in olfactory conditioning, that acts as a conditioned stimulus
31 (CS). Before this association the CS is a stimulus that at first induces only a minor orienting
32 response, but following contingent associations with the US (such that the CS predicts the
33 occurrence of the US), the CS acquires aversive properties itself and evokes an aversive
34 conditioned response (CR). Thereby after a certain number of pairings between the odor and
35 foot shocks, the sole presentation of the odor will trigger a freezing reaction in the rat (Buettner
36 [ed], 2017).

37 *Acquisition and retention of schedule-controlled behaviour*

38 Acquisition and retention of schedule-controlled behaviour involves dopaminergic projections to
39 nucleus accumbens, amygdala and prefrontal cortex. Examples of protocols for fixed interval
40 (FI) schedule of reinforcement can be found in Campbell and Haroutunian (1981) and Cory-
41 Slechta *et al.* (1983).

42 Schedule-controlled behaviour is an example of operant conditioning test. Ratio schedules of
43 reinforcement specify the number of responses that the animal must perform in order to obtain
44 a reinforcer. In fixed ratio schedules, this number is an unchanging feature of the schedule,
45 whereas in variable ratio schedules, it changes unpredictably from one reinforcer to the next. In
46 progressive ratio schedules, the required number of responses is systematically increased, from
47 one reinforcer to the next, between sessions or otherwise. Responding on progressive ratio
48 schedules is normally well maintained under lower ratios, but the rate of responding declines
49 with progressive increases in the ratio requirement. The ratio at which the subject stops
50 responding is known as the breaking point (Bradshaw and Killeen, 2012).

1 Dopamine is considered necessary for e.g. positive reinforcement and expression of learned
2 appetitive behaviours (reviewed for example in Fields et al., 2007), and reduced reward learning
3 might contribute e.g. to the onset and maintenance of major depressive disorder in humans
4 (Vrieze et al., 2014). For example, systemically administered dopamine antagonists have been
5 shown to reduce previously learned responses in simple operant tasks such as fixed ratio 1 for
6 food reward.

7 **Note regarding developmental neurotoxicity studies and assessment of endocrine**
8 **disruption**

9 In OECD TG 426, ED related investigations include parameters such as open field activity, spatial
10 learning and memory, AGD, sex distribution and results in tests with expected gender-dependent
11 reactions that may indicate and support endocrine activity of an active ingredient together with
12 other data. OECD TG 426 is a Level 4 study in the OECD Conceptual Framework for Testing and
13 Assessment of Endocrine Disrupting Chemicals (according to OECD GD 150), and it provides
14 data on adverse effects on endocrine-relevant endpoints. OECD TG 426 may produce responses
15 to EATS-modalities (oestrogen/androgen/thyroid/steroidogenesis), and non-diagnostic
16 responses to R (retinoid-related) modalities.

17 **1.10.4. Further studies**

18 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.10.4 Further studies</p> <p>A decision on the need to perform additional studies including those informing on the mechanisms should be based on the outcomes of the studies listed in 8.10.1, 8.10.2 and 8.10.3 and all other relevant available data</p> | |

19 A decision on the need to perform additional studies on additional species or strain or mechanistic
20 studies should be based on the outcome of the studies already conducted and all other relevant
21 information. If there is a specific concern that is not sufficiently addressed by the minimum study
22 requirements and there is a concern that the risks associated with such hazards would not be
23 sufficiently managed, a need for additional studies expected to provide answers to the identified
24 concerns may be decided. The decision on additional species/strain to be tested primarily
25 depends on consideration of all available information including the type of substance to be tested
26 (see above in preliminary considerations of 1.10). Mechanistic studies may strengthen the WoE
27 for reproductive toxicity when the *in vivo* evidence alone is not e.g. sufficiently convincing.
28

29 **1.11. Carcinogenicity**

30 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| <p>8.11 Carcinogenicity</p> <p>See 8.11.1 for new study requirements</p> | <p>A carcinogenicity study does not need to be conducted if:</p> <ul style="list-style-type: none"> — the substance is classified as mutagen category 1A or 1B. The default presumption would be that a |

| | |
|--|---|
| | genotoxic mechanism for carcinogenicity is likely. In these cases, a carcinogenicity test will normally not be required |
|--|---|

1
2 Carcinogenicity means the induction of cancer or an increase in the incidence of cancer occurring
3 after exposure to a substance or mixture. Substances and mixtures which have induced benign
4 and malignant tumours in well performed experimental studies on animals are considered as
5 known or presumed (Category 1) or suspected (Category 2) human carcinogens, unless there is
6 strong evidence that the mechanism of tumour formation is not relevant for humans.

7 Carcinogenicity testing under BPR is intended to provide information for classification and
8 labelling and for risk assessment. To conclude on appropriate classification and labelling, the
9 available data should be considered using the criteria and guidance associated with the CLP
10 regulation. For an appropriate risk assessment, the information on dose response has to be
11 sufficient and should allow concluding on the existence of a threshold (see Guidance *BPR Volume*
12 *III Human health Parts B+C*).

13 **Collection and evaluation of available information**

14 For the assessment of existing information (physicochemical properties, grouping and read-
15 across, (Q)SARs and expert systems, *in vitro* data, human data and animal data) further
16 guidance is available within the BPR Volume III Human health Parts B+C, Guidance on the
17 Application of the CLP Criteria and the practical guides⁹ such as "How to use and report (Q)SARs".

18 In addition to the waiving conditions indicated in the data requirement and in BPR Annex IV, the
19 study does not need to be conducted if:

- 20
- No genotoxic potential for humans is identified in genotoxicity tests, and
 - Possible mechanisms of toxicological effects observed in subchronic toxicity studies are without any indications of non-genotoxic carcinogenicity and there are no structural alerts for carcinogenicity, and
 - The subchronic studies in rodents and/or non-rodents are without indication of substance related adverse effects at the limit dose level.
- 21
22
23
24
25

26 **Generation of new test data**

27 If further testing is needed to assess carcinogenicity, the test methods below should be used.

- 28
- OECD Test Guideline 453: Combined chronic toxicity/carcinogenicity study.
 - EU: B.33. Combined chronic toxicity/carcinogenicity test
 - OECD Test Guideline 451: Carcinogenicity study
 - EU. B.32. Carcinogenicity test
- 29
30
31

32 Where new testing is needed, please see also the general information under *Considerations*
33 *before initiating testing* in chapter 1.

34 Other tests may contribute to a weight of evidence evaluation, e.g. by providing supporting

⁹ <https://echa.europa.eu/practical-guides>

1 information or mechanistic data.

2 For guidance on reporting historical control data see Section 1.

3 **Mode of action (MoA) and human relevance**

4 When carcinogenicity is observed, it may be necessary to further investigate the MoA and the
5 relevance of the effect for humans. All available data must be carefully considered to assess if it
6 can be concluded that the tumours are induced by a specific mechanism.

7 For the purpose of elucidating a non-genotoxic mode of action and human relevance, the need
8 for further investigations should be considered on a case-by-case basis, focusing first on
9 mechanistic studies (see also 1.13.5). The IPCS Framework for Analyzing the Relevance of a
10 Cancer Mode of Action for Humans (2007)¹⁰ may be useful in considering the testing/assessment
11 strategy.

12 **1.11.1. Combined carcinogenicity study and long-term repeated dose toxicity**

13 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.11.1 Combined carcinogenicity study and long-term repeated dose toxicity Rat, oral route of administration is the preferred route. If an alternative route is proposed a justification must be provided. For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route | |

14

15 See also section 1.9.3 of this guidance.

16

17 **1.11.2. Carcinogenicity testing in a second species**

18 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| 8.11.2 Carcinogenicity testing in a second species (a) A second carcinogenicity study | The second carcinogenicity study does not need to be conducted if the applicant can justify on the basis of scientific grounds that it is not necessary' |

¹⁰ IPCS (2007) Boobis A.R., Cohen S.M., Dellarco V., McGregor D., Meek M.E., Vickers C., Willcocks D., Farland W.: IPCS Framework for Analyzing the Relevance of a Cancer Mode of Action for Humans in IPCS Harmonization Project Document No. 4, Part 1, IPCS framework for analysing the relevance of a cancer mode of action for humans and case-studies. http://www.who.int/ipcs/methods/harmonization/areas/cancer_mode.pdf

| | |
|--|--|
| <p>should be conducted using the mouse as test species;</p> <p>(b) For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route</p> | |
|--|--|

1

2 If comparative metabolism data indicate that mouse is an inappropriate model for human cancer
3 risk assessment, an alternative species shall be considered.

4 **1.12. Relevant health data, observations and treatments**

5 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.12 Relevant health data, observations and treatments</p> <p>Justification should be provided if data is not available</p> | |

6

7 When no human studies/data are available, new studies on human volunteers should not be
8 conducted.

9 Data and information on any effects observed in humans may provide valuable information on
10 the validity of extrapolations from animal data to expected effects in humans, and to identify
11 any unexpected adverse effect that could be specific to humans.

12 All available data and information of adequate quality following accidental or occupational
13 exposure have to be submitted.

14 **1.12.1. Information on signs of poisoning, clinical tests, first aid measures, 15 antidotes, medical treatment and prognosis following poisoning**

16 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| <p>8.12.1 Information on signs of poisoning, clinical tests, first aid measures, antidotes, medical treatment and prognosis following poisoning</p> | |

17

18 Observations and information relevant to the recognition of the symptoms of poisoning, as well
19 as on the effectiveness of first aid and therapeutic measures must be included.

20 The reports should include a complete description of the exposure situations, clinical symptoms
21 observed, therapeutic measures and clinical follow-up.

- 1 A detailed description of clinical signs and details of clinical tests (such as biomonitoring and
2 patch tests) useful for diagnostic purposes must be included.
- 3 Symptoms of poisoning must be described, including full details of the time courses involved for
4 all exposure routes.
- 5 First aid measures in the event of poisoning and eye contamination must be provided.
- 6 Therapeutic regimes and the use of antidotes must be described. Information based on practical
7 experience must be provided where available, and otherwise, information must be provided
8 based on theoretical grounds as to the effectiveness of any treatment regimes. Contraindications
9 associated with particular regimes, particularly those relating to 'general medical problems' and
10 conditions, must be described. The expected effects and the duration of these effects following
11 poisoning must be described.

12 **1.12.2. Epidemiological studies**

13 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--------------------------------|---|
| 8.12.2 Epidemiological studies | |

14

15 Four major types of epidemiological studies may be submitted: (1) analytical epidemiology
16 studies on exposed populations, (2) descriptive or correlation epidemiology studies, (3) case
17 reports and (4) in very rare, justified cases, controlled studies in human volunteers.

18 Analytical epidemiology studies are useful for identifying a relationship between human exposure
19 and effects such as biological effect markers, early signs of chronic effects, disease occurrence,
20 or mortality. Such studies may provide the best data for risk assessment.

21 Descriptive epidemiology studies examine differences in disease rates among human populations
22 in relation to e.g. age and gender, and differences in temporal or environmental conditions.
23 Typically, these studies can only identify patterns or trends in disease occurrence over time or
24 in different geographical locations but cannot ascertain the causal agent or degree of human
25 exposure.

26 Case reports describe a particular health condition in an individual or a group of individuals who
27 were exposed to a substance. They may be particularly relevant when demonstrating effects
28 that cannot be observed in experimental animal studies. In many such studies, information is
29 lacking on critical aspects such as substance identity and purity, exposure, health status of the
30 persons exposed and even the symptoms reported; thorough assessment of the reliability and
31 relevance of case reports is therefore necessary.

32 For further information, please refer to REACH Guidance on information requirements and
33 chemical safety assessment, Chapter R.4: Evaluation of available information.

34 **1.12.3. Medical surveillance data, health records and case reports**

35 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|-----------------------------------|---|
| 8.12.3 Medical surveillance data, | |

| | |
|----------------------------------|--|
| health records and case reports' | |
|----------------------------------|--|

1

2 The reports should include detailed information on the design of the occupational surveillance
3 programme and exposure to the active substance and to other chemicals. Data relevant to the
4 mechanism of the action of substance should also be included where feasible. The data may
5 consist of published articles or unpublished medical surveys.

6 The following information on sensitisation should be provided where available, including any
7 details necessary for the evaluation of the information (please see also ECHA Guidance Vol III,
8 Parts B+C):

- 9 • Information on (respiratory) sensitisation or any incidences of (respiratory)
10 hypersensitivity of workers or others exposed.
- 11 • Evidence that the substance can induce specific respiratory hypersensitivity will usually
12 be based on human experience data. The clinical history data including both medical and
13 occupational history, and reports from appropriate lung function tests related to exposure
14 to the substance should be submitted, if available.
- 15 • Reports of other supportive evidence, such as:
- 16 ○ Information of a chemical structure within the active substance that is related to
17 substances known to cause respiratory hypersensitivity;
 - 18 ○ *In vivo* immunological tests;
 - 19 ○ *In vitro* immunological tests;
 - 20 ○ Studies indicating other specific but non-immunological mechanisms of action;
 - 21 ○ Data from a positive bronchial challenge test.

22 1.13. Additional studies (ADS)

23 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| <p>8.13 Additional studies</p> <p>Additional data which may be required depending on the characteristics and intended use of the active substance</p> <p>Other available data: Available data from emerging methods and models, including toxicity pathway-based risk assessment, <i>in vitro</i> and 'omic' (genomic, proteomic, metabolomic, etc.) studies, systems biology, computational toxicology, bioinformatics, and high-throughput screening shall be submitted in parallel</p> | |

24

1 Toxicity studies of metabolites

2 Supplementary studies, where they relate to substances other than the active substance, are
3 not a routine requirement. Decisions as to the need for supplementary studies should be made
4 on a case-by-case basis.

5 Where as a result of metabolism or other processes, metabolites from plants or in animal
6 products, soil, groundwater or open air differ from those in animals used for the toxicology
7 studies or are detected in low proportions in animals, further testing should be carried out on a
8 case-by-case basis, taking into account the amount of metabolite and the chemical structure of
9 the metabolite compared to the parent.

10 Supplementary studies on the active substance

11 Supplementary studies should be carried out where they are necessary to further clarify the
12 observed effects, taking into account the results of the available toxicological and metabolism
13 studies and the most important exposure routes. Such studies may include:

- 14 (a) studies on absorption, distribution, excretion and metabolism, in a second species;
- 15 (b) studies on the immunotoxicological potential;
- 16 (c) a targeted single dose study to derive appropriate acute reference values (ARfD, AEL);
- 17 (d) studies on other routes of administration;
- 18 (e) studies on the carcinogenic potential;
- 19 (f) studies on mixture effects.

20 The studies required should be designed on an individual basis, in the light of the particular
21 parameters to be investigated and the objectives to be achieved.

22 1.13.1. Phototoxicity - additional study (ADS)

23 Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|----------------------|---|
| 8.13.1 Phototoxicity | |

24 There is possible concern of phototoxicity if the active substance:
25

- 26 • Absorbs light within the range of natural sunlight (290-700 nm); and
- 27 • Is liable to reach the eyes or light-exposed areas of skin, either by direct contact or
28 through systemic distribution.

29 If the molar/extinction/absorption coefficient (MEC) of the active substance is less than 1000 L
30 x mol⁻¹ x cm⁻¹ (measured in methanol), the active substance is not considered to be
31 photoreactive enough to result in phototoxicity. MEC is also called molar absorptivity and it
32 reflects the efficiency with which a molecule can absorb a photon at a particular wavelength
33 (typically expressed as L mol⁻¹ cm⁻¹) and is influenced by several factors, such as solvent.
34 Detailed guidance on the use of the coefficient and the assessment of phototoxicity is provided

1 in the ICH Guidance S10 on Photosafety Evaluation of Pharmaceuticals¹¹.

2 The following test method for phototoxicity should be used:

- 3 • EC method B.41.
- 4 • OECD Test Guideline 432: *In vitro* 3T3 NRU phototoxicity test.

5 Where new testing is needed, please see also the general information under *Considerations*
6 *before initiating testing* in chapter 1.

7 The study should provide information on the potential of certain active substances to induce
8 cytotoxicity in combination with light.

9 Examples of phototoxic active substances:

- 10 • active substances that are phototoxic *in vivo* after systemic exposure and distribution to
11 skin;
- 12 • active substances that act as photoirritants/photosensitisers after dermal application to
13 skin.

14 A positive result should be taken into account when considering potential human exposure. For
15 photogenotoxicity see section 1.6 of this guidance.

16 **1.13.2. Neurotoxicity (ADS)**

17 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| <p>8.13.2 Neurotoxicity</p> <p>If the active substance is an organophosphorus compound or if there is an indication, knowledge of the mechanism of action or knowledge from acute or repeated dose studies that the active substance may have neurotoxic properties, additional information or specific studies (such as OECD TG 424 or OECD TG 418 or 419 or equivalent) will be required</p> <p>If anticholinesterase activity is detected a test for response to reactivating agents should be considered</p> <p>For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral</p> | |

¹¹ Available at <https://www.ema.europa.eu/en/ich-s10-photosafety-evaluation-pharmaceuticals>.

| | |
|-------|--|
| route | |
|-------|--|

- 1
2 Specific studies or additional specific investigations on neurotoxicity should be performed for
3 active substances that:
- 4 - are organophosphorus compounds;
 - 5 - have structural or other similarity to substance(s) capable of inducing neurotoxicity (e.g.
6 carbamate compounds);
 - 7 - induce specific indications of potential neurotoxicity such as clinical signs or effects in
8 functional tests indicating neurotoxicity or neuropathological lesions in toxicity studies;
 - 9 - have a neurotoxic mode of action unless the MoA has been demonstrated to be irrelevant
10 to humans and other MoAs can be excluded,

11 and these indications of neurotoxicity are not as such sufficient for classification and labelling for
12 neurotoxicity and/or risk management.

13 Indications or evidence of neurotoxicity can be acquired from the standard systemic toxicity
14 studies, but only when neurotoxicity is so pronounced that it is visible as clinical signs (e.g.
15 sedation, coma, convulsions) or by histopathological investigations. Lack of such effects does
16 not indicate lack of neurotoxicity as standard repeated dose studies do not include specific and
17 sensitive tests for neurotoxicity. Thereby also the potential effects seen in these standard
18 systemic toxicity studies normally represent high-dose effects and when testing further by
19 sensitive and specific methods it may be possible to detect also more subtle effects at lower
20 doses.

21 If additional information or specific studies are warranted for neurotoxicity, they should provide
22 adequate data to sufficiently investigate the neurotoxic potential of the active substance after
23 single and repeated exposure. The data should also be useful for classification and labelling in
24 accordance with CLP; therefore please consult the CLP criteria for STOT SE (CLP 3.8) and STOT
25 RE (CLP 3.9) under which neurotoxicity is assessed (note that developmental neurotoxicity is
26 part of reproductive [developmental] toxicity and discussed in chapter 1.10.3).

27 Specific neurotoxicity studies often investigate the function of different components of the
28 nervous system by specific and sensitive neurobehavioural tests and the histopathological effects
29 in the central and peripheral nervous systems. Other possible investigations may comprise of
30 neurophysiological (e.g. electroencephalography, electrophysiology) or biochemical studies
31 (investigating e.g. neurotransmitter levels, receptor expression and binding).

32 **Collection and evaluation of available information**

33 For the assessment of existing information (physicochemical properties, grouping, (Q)SARs and
34 expert systems, *in vitro* data, human data and animal data) further guidance is available within
35 the *Guidance on the Application of the CLP Criteria* and *BPR Volume III Human health Parts B+C*.

36 **Generation of new test data**

37 When it is considered necessary to conduct a neurotoxicity study, it is important that the study
38 design is discussed by the contractor/laboratory and the assessor before initiating the study,
39 paying particular attention to the specificity and sensitivity of the protocol to be used.

40 If further standard 28- or 90-day studies are to be conducted, additional neurotoxicity
41 parameters could be added if expected to be able to provide the missing information.

42 Neurotoxicity testing to conclude on classification and labelling and to establish a NOAEL for

1 neurotoxicity, is required when data from standard toxicity studies or any other available
2 information are indicative but not conclusive for neurotoxicity.

3 Test method for Neurotoxicity study in rodents:

- 4 • EC method B.43
- 5 • OECD Test Guideline 424

6 The OECD TG 424 is intended for confirmation or further characterisation of potential
7 neurotoxicity identified in previous studies or by other available information. It allows a flexible
8 approach where comprehensive investigations of specific neurotoxicity endpoints by sensitive
9 tests can be included. The dose levels should be adjusted to avoid confounding effects by general
10 toxicity, but they should be sufficiently high to allow to conclude on potential absence of effects
11 on the tested parameters. The procedures set out by OECD TG 424 can be used to investigate
12 both repeated dose and acute neurotoxicity. Possible inclusion of a satellite group for assessment
13 of reversibility of effects could be considered. For STOT SE and STOT RE both reversible and
14 irreversible effects are relevant.

15 The timing of the peak effect caused by the substance needs to be considered for the timing of
16 testing different neurotoxicity parameters. The duration of exposure and time after
17 administration needed to induce specific neurotoxic effects will depend on toxicokinetics of the
18 substance and the underlying mechanism(s) of action.

19 Testing during short-term peak exposures is important for revealing acute neurotoxic effects
20 that are often transient and to which tolerance may develop after repeated exposure. When the
21 test compound is administered as a bolus via the intravenous, subcutaneous or oral route and
22 causes acute neurotoxicity, it is essential to determine the time-effect course of the acute effect,
23 and to perform measurements of acute neurotoxicity parameters at the time of the peak effect.

24 Where cumulative toxicity or repeated-dose effects are the primary focus, testing should precede
25 the daily dose to rule out acute (less than 24 hour) effects. For delayed neurotoxicity a
26 sufficiently long period between the last dose and neurotoxicity testing is required.

27 For example, the acute and chronic neurotoxicity associated with exposure to specific volatile
28 organic solvents has been well identified based on human experience. The acute neurotoxic
29 effects are investigated with acute inhalation studies designed to detect findings such as
30 transient narcotic effects. However, long-term exposure to acute neurotoxicants may cause
31 additional neurotoxic effects of different nature and at lower doses than the acute neurotoxic
32 effects. To reveal these effects, repeated dose neurotoxicity studies should be performed by
33 using sensitive and specific tests. For some neurotoxic substances only a long exposure period
34 will elicit neurotoxic effects.

35 The most appropriate methods for further investigation of neurotoxicity should be determined
36 on a case-by-case basis, guided by the effects seen in the standard systemic toxicity tests, any
37 other available data. Methods which may be used are given in Table 10 below.

38 **Table 10 Methods for investigation of neurotoxicity**

| EFFECT | METHODS |
|-----------------------|--|
| Morphological changes | Neuropathology Gross anatomical techniques Immunocytochemistry Special stains |

| EFFECT | METHODS |
|-----------------------|---|
| Physiological changes | Electrophysiology Electroencephalogram (EEG) Evoked potentials |
| Behavioural changes | Functional observations Sensory function tests Motor function tests Cognitive function tests |
| Biochemical changes | Neurotransmitter analyses Enzyme/protein activity Measures of cell integrity |

1

2 Several MoAs, such as acetylcholine esterase (AChE) inhibition, have been associated with
3 neurotoxic effects. AChE may be inhibited to varying extents depending on animal or cell model,
4 dose, duration of exposure, and specific compound (Voorhees et al., 2016). Organophosphorus
5 compounds and carbamates are examples of compounds that can inhibit acetylcholinesterase,
6 but they have also other targets causing neurotoxicity (Voorhees et al., 2016; Lotti and Moretto,
7 2006). Exposure to high levels of organophosphorus compounds may cause cholinergic crisis in
8 humans and animals characterised by via overstimulation of the nervous system leading to
9 respiratory failure, flaccid paralysis, decreased blood pressure, parasympathetic discharge, and
10 even death. Lower (repeated dose) exposure levels have been associated with
11 neurodegenerative disease, psychiatric illness, and sensorimotor deficits in humans whereas in
12 rodent models deficits in learning and memory, attention and impulsive behaviour and some
13 other cognitive functions have been reported after repeated exposure to certain
14 organophosphorous compounds (see also delayed neuropathy below) (Voorhees et al., 2016).

15 There are many other neurotoxic MoAs as well. Based on the MoA, it needs to be carefully
16 considered which neurotoxicity test(s) is (are) most appropriate (specific and sensitive) to
17 investigate the adverse effects caused by the identified MoA. For example, in rats pyrethroids
18 may produce marked behavioural arousal, aggressive sparring, increased startle response, and
19 fine body tremor progressing to whole-body tremor, and prostration (T syndrome) and/or
20 profuse salivation, coarse tremor progressing to choreoatetosis, and clonic seizure (CS
21 syndrome) by affecting the function of sodium channels, GABA_A receptors and voltage-
22 dependent chloride channels. Degeneration of dopaminergic neurons in the substantia nigra may
23 result in Parkinson's disease-like symptoms manifested in rodents as e.g. impairments in
24 movement initiation, weight shifting, and in postural stability, whereas a substance targeting
25 hippocampal, amygdala and pyriform cortex neurons may cause cognitive impairment (Costa et
26 al., 2008).

27 **Delayed polyneuropathy studies**

28 Delayed polyneuropathy studies should provide sufficient data to evaluate if the active substance
29 may provoke delayed polyneuropathy after acute and/or repeated exposure by inhibition of
30 neuropathy target esterase (NTE). Organophosphate-induced delayed polyneuropathy (OPIDN)
31 results from exposure to certain organophosphorus compounds. It is characterised by distal
32 degeneration of some axons of both the peripheral and central nervous systems occurring 1-4
33 weeks after single or short-term exposures (Lotti and Moretto, 2005). The condition is
34 characterized by motor weakness, fatigue and paralysis and sensory numbness, tingling, and
35 pain. OPIDN has been attributable to inhibition of neuropathy target esterase (NTE), rather than
36 AChE, as inhibition of AChE is not necessary for the development of OPIDN (Woltje, 2015). Also,
37 some carbamates are known to inhibit neuropathy target esterase (NTE) (Lotti and Moretto,
38 2006). A repeated exposure study for delayed neuropathy may be waived unless there are

1 indications that the compound accumulates and significant inhibition of NTE or
2 clinical/histopathological signs of OPIDN occur at around the hen LD₅₀ as determined in the single
3 dose test.

4 Delayed neurotoxicity tests in the laying hen after acute and repeated exposure (OECD TG 418
5 and OECD TG 419) should be performed for active substances of similar or related structures to
6 those capable of inducing delayed polyneuropathy such as organophosphorus compounds, unless
7 there is already sufficient information to conclude on neurotoxicity.

8 Test methods for delayed neuropathy:

9 • OECD Test Guideline 418: Delayed Neurotoxicity of Organophosphorus Substances
10 Following Acute Exposure. (EC method B.37 Delayed neurotoxicity of organophosphorus
11 substances after acute exposure)

12

13 • OECD Test Guideline 419: Delayed Neurotoxicity of Organophosphorus Substances: 28-
14 day Repeated Dose Study (EC method B.38 Delayed neurotoxicity of organophosphorus
15 substances 28-day repeated dose study)

16 In OECD TG 418, a single dose of the test substance is administered orally to domestic hens,
17 NTE (and potentially AChE) activity is assayed 24 and 48 h after dosing, the animals are observed
18 for 21 days for ataxia, paralysis and other behavioural abnormalities, and 21-days after exposure
19 histopathological examination of selected neural tissues is performed. In OECD TG 419, the
20 exposure period is 28 days, NTE (and potentially AChE) activity is assayed 24 and 48 h after the
21 last dosing, the animals are observed for 14 days after the last dose and after which the
22 histopathological examination is performed.

23 1.13.3. Endocrine disruption

24 Information requirement according to BPR Annex II:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|--|
| <p>8.13.3 Endocrine disruption</p> <p>The assessment of endocrine disruption shall comprise the following tiers:</p> <p>(a) An assessment of the available information from the following studies and any other relevant information, including <i>in vitro</i> and <i>in silico</i> methods:</p> <p>(i) 8.9.1 A 28-day oral toxicity study in rodents (OECD TG 407);</p> <p>(ii) 8.9.2 A 90-day oral toxicity study in rodents (OECD TG 408);</p> <p>(iii) 8.9.4 A repeated dose oral toxicity study in non-rodents (OECD TG 409);</p> | <p>Where sufficient weight of evidence to conclude on the presence or absence of a particular endocrine disrupting mode of action is available:</p> <p>—further testing on vertebrate animals for that effect shall be omitted for that mode of action,</p> <p>—further testing not involving vertebrate animals may be omitted for that mode of action.</p> <p>In all cases, adequate and reliable documentation shall be provided'</p> |

| | |
|--|--|
| <p>(iv)8.10.1 A prenatal developmental toxicity study (OECD TG 414);</p> <p>(v) 8.10.2 An extended one-generation reproductive toxicity study (OECD TG 443) or two-generation reproductive toxicity study (OECD TG 416);</p> <p>(vi)8.10.3 A developmental neurotoxicity study (OECD TG 426);</p> <p>(vii) 8.11.1 A combined carcinogenicity study and long-term repeated dose toxicity study (OECD TG 451-3);</p> <p>(viii) A systematic review of the literature including studies on mammals and non-mammalian organisms;</p> <p>(b)If there is any information suggesting that the active substance may have endocrine disrupting properties, or if there is incomplete information on key parameters relevant for concluding on endocrine disruption, then additional information or specific studies shall be required to elucidate:</p> <p>(1) the mode or the mechanism of action; and/or</p> <p>(2)potentially relevant adverse effects in humans or animals</p> <p>For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to consider the oral route and conduct animal studies by the oral route</p> | |
|--|--|

1
2 This data requirement (8.13.3 Endocrine disruption) is a core data requirement although it is
3 placed under 8.13 Additional studies (ADS). This discrepancy is due to the change in the
4 legislation, as Regulation (EU) 2021/525 changed this data requirement from ADS to CDS.

5 This guidance should be read in conjunction with the ECHA/EFSA Guidance for the identification
6 of endocrine disruptors in the context of Regulations (EU) No 528/2012 and (EC) No 1107/2009
7 where the testing strategy is further elaborated.

8 Objectives

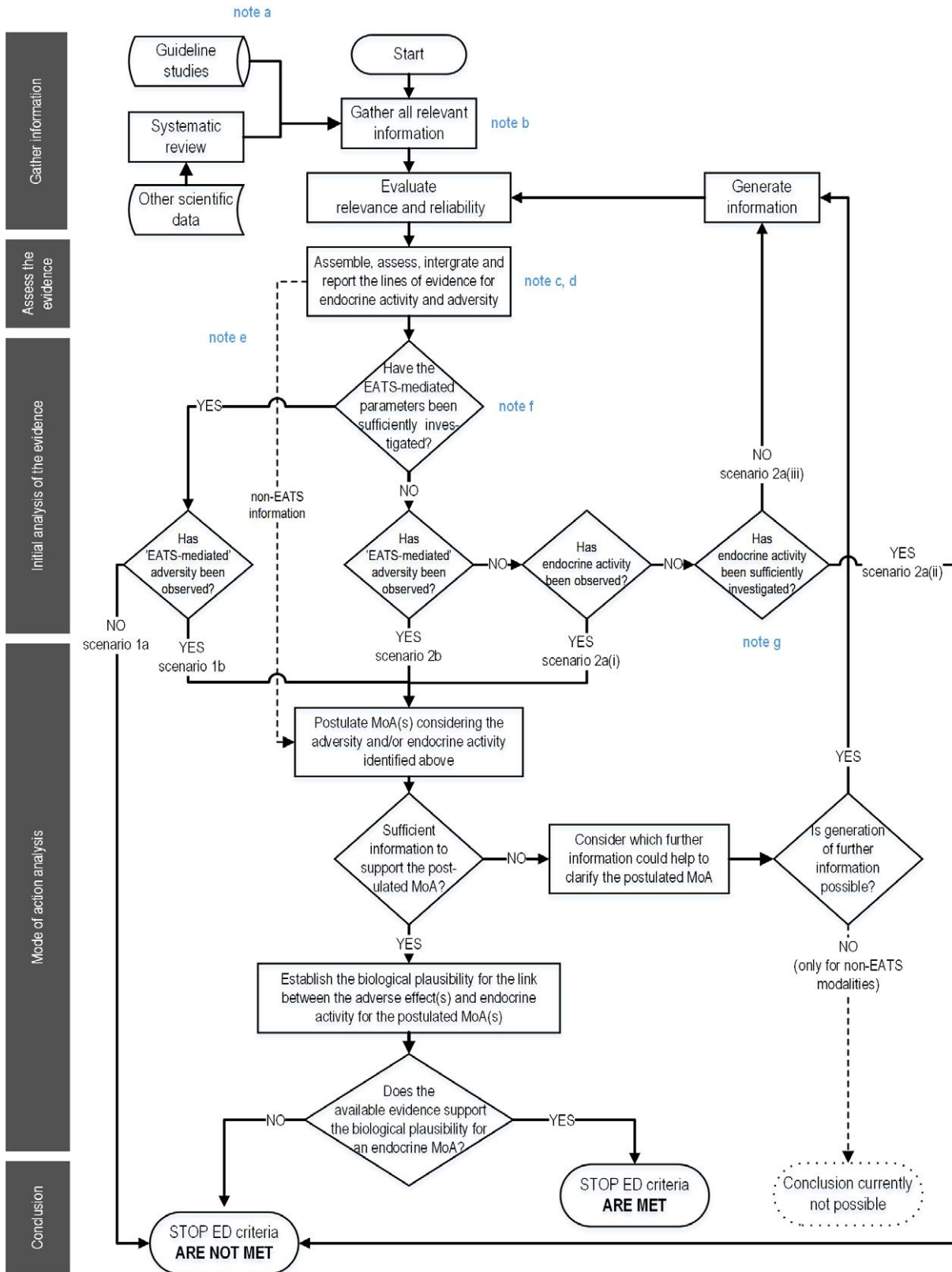
1 For each biocidal active substance, a conclusion is needed whether the substance meets the
2 criteria to be considered as an endocrine disruptor. To this end, there needs to be sufficient
3 information available to conclude whether the active substance interferes with the endocrine
4 system in a way that leads to adverse effects, in particular via any of the four modalities
5 addressed in the ED guidance¹². The information requirements for endocrine disruption have
6 three core objectives:

- 7 - to have sufficient information to conclude whether adverse effects occur that are
8 indicative of an endocrine mode of action
- 9 - to have sufficient information on endocrine activity
- 10 - to obtain sufficient information to perform a MoA analysis on endocrine activity/endocrine
11 disrupting properties if adversity is observed.

12 This guidance provides advice on the tests that an applicant can and should perform to address
13 the endocrine disrupting properties of the active substance and to conclude whether the ED
14 criteria are met or not.

15 The ED criteria, the data requirements and the ED guidance all aim at assessing the endocrine
16 disruption hazard and are not intended to assess the presence or absence of a threshold for the
17 endocrine disrupting effect. The scientific discussion on the possibility to identify a threshold for
18 endocrine disrupting substances goes beyond the scope of this document. However, pending this
19 discussion, whenever a threshold is considered to exist, this has to be demonstrated on a case-
20 by-case basis. The analysis of whether or not a threshold can be set is the responsibility of the
21 applicant, and a proposal on a threshold should be based on appropriate data in the application
22 dossier. This means that it is possible that additional data might need to be generated after a
23 substance is identified as an ED in order to assess the risk.

¹² Note that while the ED guidance focuses on these four modalities, the ED criteria are not limited to these modalities. There may be cases where sufficient information is available on disruption of other part of the endocrine axes, which could lead to a substance meeting the ED criteria.



1
2

1 **Figure 4: Flow chart illustrating the ED assessment strategy.** The figure is from the
2 ECHA/EFSA Guidance (2018) for the identification of endocrine disruptors – for notes and
3 scenarios, please see this guidance. The assessment strategy illustrated in the flow chart is
4 applicable both for humans and non-target organisms and is driven by the availability of 'EATS-
5 mediated' parameters as these provide evidence for both endocrine activity and the resulting
6 potentially adverse effects. However, there may be situations where the 'EATS-mediated'
7 parameters are insufficiently investigated. In such cases, it may be possible to follow the
8 assessment strategy using the 'sensitive to, but not diagnostic of, EATS' parameters, without
9 the need to generate additional information on EATS-mediated parameters i.e. in case of
10 scenarios 2a(i) or 2b. If the required data are available, it is in principle possible to establish
11 endocrine disrupting MoA(s) on the basis of parameters indicating 'sensitive to, but not
12 diagnostic of, EATS' potential adversity and EATS endocrine activity.

13 **General overview of the assessment strategy**

14 This section contains an overview of the assessment strategy to determine whether a substance
15 meets the definition of an endocrine disruptor according to the ED criteria. More detailed
16 information on the assessment strategy and relevant test methods can be found in *ECHA/EFSA*
17 *Guidance for the identification of endocrine disruptors in the context of Regulations (EU) No*
18 *528/2012 and (EC) No 1107/2009.*

19 The ED criteria (Regulation (EU) 2017/2100) cover all endocrine-disrupting modes of action
20 (MoAs), i.e. adverse effects which may be caused by any endocrine modality. The ECHA/EFSA
21 Guidance focuses mainly on effects caused by the EATS (estrogenic, androgenic, thyroidal and
22 steroidogenic) modalities. This is because these pathways are currently the best understood, i.e.
23 with a relatively good mechanistic understanding of how substance-induced perturbations may
24 lead to adverse effects via an endocrine-disrupting MoA. In addition, standardised test methods
25 for *in vivo* and *in vitro* testing are currently available only for these modalities. However, there
26 may be situations where it is possible to conclude on ED properties also for non-EATS modalities.

27 To facilitate the assessment, the ECHA/EFSA Guidance has grouped the parameters investigated
28 in the standard test methods depending on the type of information they provide. The groups
29 are:

- 30 • ***In vitro* mechanistic** – parameters measured *in vitro* that provide information on
31 endocrine activity (e.g. by binding to and activating a receptor or interfering with
32 hormone production).
- 33 • ***In vivo* mechanistic** – parameters measured *in vivo* that provide information on
34 endocrine activity (e.g. changes in hormone levels or effects in a specific tissue known
35 to be mainly under endocrine control).
- 36 • **EATS-mediated** – parameters measured *in vivo* that may contribute to the evaluation
37 of adversity, while at the same time (due to the nature of the effect and the existing
38 knowledge) they are also considered indicative of an EATS MoA and thus (in the absence
39 of other explanations) also imply underlying *in vivo* mechanistic information.
- 40 • **Sensitive to, but not diagnostic of, EATS** – parameters measured *in vivo* that may
41 contribute to the evaluation of adversity, however, due to the nature of the effect and
42 the existing knowledge, these effects cannot be considered diagnostic on their own of
43 any one of the EATS modalities.

44 **Steps in the assessment strategy**

45 The starting point for the ED assessment strategy is that the other regulatory requirements for
46 the (active) substance are met and that the information is available. It is recognised that there
47 may be situations where the available information has not sufficiently investigated certain

1 parameters which are crucial for a robust conclusion on potential ED properties. In such cases,
2 additional data generation may need to be required.

3 The assessment strategy is based on the three conditions stipulated in the ED criteria – adversity,
4 endocrine activity and a biologically plausible link between the two – and on the grouping of the
5 parameters as described above. The ‘EATS-mediated’ parameters drive the assessment strategy
6 because, by providing evidence for both endocrine activity and the resulting potentially adverse
7 effects, they are considered indicative of an endocrine MoA. Parameters which are considered as
8 ‘sensitive to, but not diagnostic of, EATS’ and ‘EATS-mediated’ parameters are normally
9 investigated in the same tests. If there is no adversity seen in the ‘EATS-mediated’ parameters,
10 but adversity is observed in the same study in parameters considered ‘sensitive to, but not
11 diagnostic of, EATS’, then this adversity is not likely to be caused by alterations of the EATS
12 modalities. There may be situations where the ‘EATS-mediated’ parameters are not sufficiently
13 investigated (e.g. tests carried out according to outdated guidelines), and in such cases, any
14 adversity observed in parameters considered ‘sensitive to, but not diagnostic of, EATS’, cannot
15 be dismissed.

16 The assessment strategy is applicable both for humans and non-target organisms and in both
17 cases, Figure 4 illustrates the steps of the assessment. Each of the steps outlined in the figure
18 are described below.

19 **Gather information.** In this step, all available relevant information (including *in vitro* and *in*
20 *silico* methods) is gathered, evaluated for relevance and reliability, and extracted and reported
21 in the competent authority report. Relevant information is expected to be provided based on the
22 existing data requirements, including (but not limited to) the following:

23 (i) 8.9.1 A 28-day oral toxicity study in rodents (OECD TG 407)

24 (ii) 8.9.2 A 90-day oral toxicity study in rodents (OECD TG 408)

25 (iii) 8.9.4 A repeated dose oral toxicity study in non-rodents (OECD TG 409)

26 (iv) 8.10.1 A prenatal developmental toxicity study (OECD TG 414)

27 (v) 8.10.2 An extended one-generation reproductive toxicity study (OECD TG 443) or two-
28 generation reproductive toxicity study (OECD TG 416)

29 (vi) 8.10.3 A developmental neurotoxicity study (OECD TG 426)

30 (vii) 8.11.1 A combined carcinogenicity study and long-term repeated dose toxicity
31 study (OECD TG 451-3)

32 (viii) A systematic review of the literature including studies on mammals and non-
33 mammalian organisms

34 In addition to the studies listed in the data requirements, additional information must be
35 identified by performing a systematic literature review. The systematic review should focus on
36 information relevant for the ED assessment coming from *in vivo*, *in vitro* and *in silico* studies.
37 More information is provided in Appendix F of the ECHA/EFSA ED Guidance and EFSA (2010)
38 *Application of systematic review methodology to food and feed safety assessments to support*
39 *decision making.*

40 **Assess the evidence.** The information is assembled into lines of evidence, integrating
41 information for both adversity and endocrine activity for each of the EATS modalities. The lines
42 of evidence are assessed and reported in the dossier/CAR. If there is indication of non-EATS-
43 related endocrine activity and/or effects, this should be taken forward to the MoA analysis step
44 because the questions asked in the next step are tailored to the EATS modalities.

1 **Initial analysis of the evidence.** This step includes a decision tree. The decisions are driven
2 by the availability of 'EATS-mediated' parameters and/or evidence of endocrine activity. This
3 first step is to assess whether the available evidence already allows concluding that a substance
4 does not meet the ED criteria, or whether a more detailed analysis and/or additional information
5 is needed to conclude on the ED properties.

6 **MoA analysis.** This step aims to establish if there is a biologically plausible link between
7 observed adverse effects and endocrine activity. Different situations are outlined. Depending on
8 the available evidence, the applicant and the assessor need to identify the information that may
9 need to be generated to further investigate the adversity or the endocrine activity, or any
10 potential alternative MoA(s). In this step, it should be further investigated whether it is possible
11 to establish a plausible link between non-EATS endocrine activity and observed adversity, or
12 whether further information could be generated to clarify whether there is a non-EATS
13 endocrine-disrupting MoA.

14 **Conclusion on the ED criteria.** In this step, the conclusion is made whether the ED criteria
15 are met with respect to humans. The conclusion is transparently documented, including the
16 remaining uncertainties.

17 If a conclusion cannot be made whether the substance meets the ED criteria, then additional
18 information or specific studies shall be required. These are specified in the chapter below.

19 **1.13.3.1. Specific additional studies to investigate potential endocrine** 20 **disrupting properties (ADS)**

21 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.13.3.1 Specific additional studies to investigate potential endocrine disrupting properties may include, but are not limited to the following:</p> <ul style="list-style-type: none"> (a) the mammalian toxicity studies listed in 8.13.3(a); (b) the <i>in vitro</i> assays: <ul style="list-style-type: none"> (i) Estrogen receptor transactivation assay (OECD TG 455); (ii) Androgen receptor transactivation assay, (OECD TG 458); (iii) H295R steroidogenesis assay (OECD TG 456); (iv) the Aromatase assay (human recombinant) OPPTS 890.1200; (c) Uterotrophic bioassay in rodents (OECD TG 440) and Hershberger bioassay in rats (OECD TG 441); (d) Pubertal development and | |

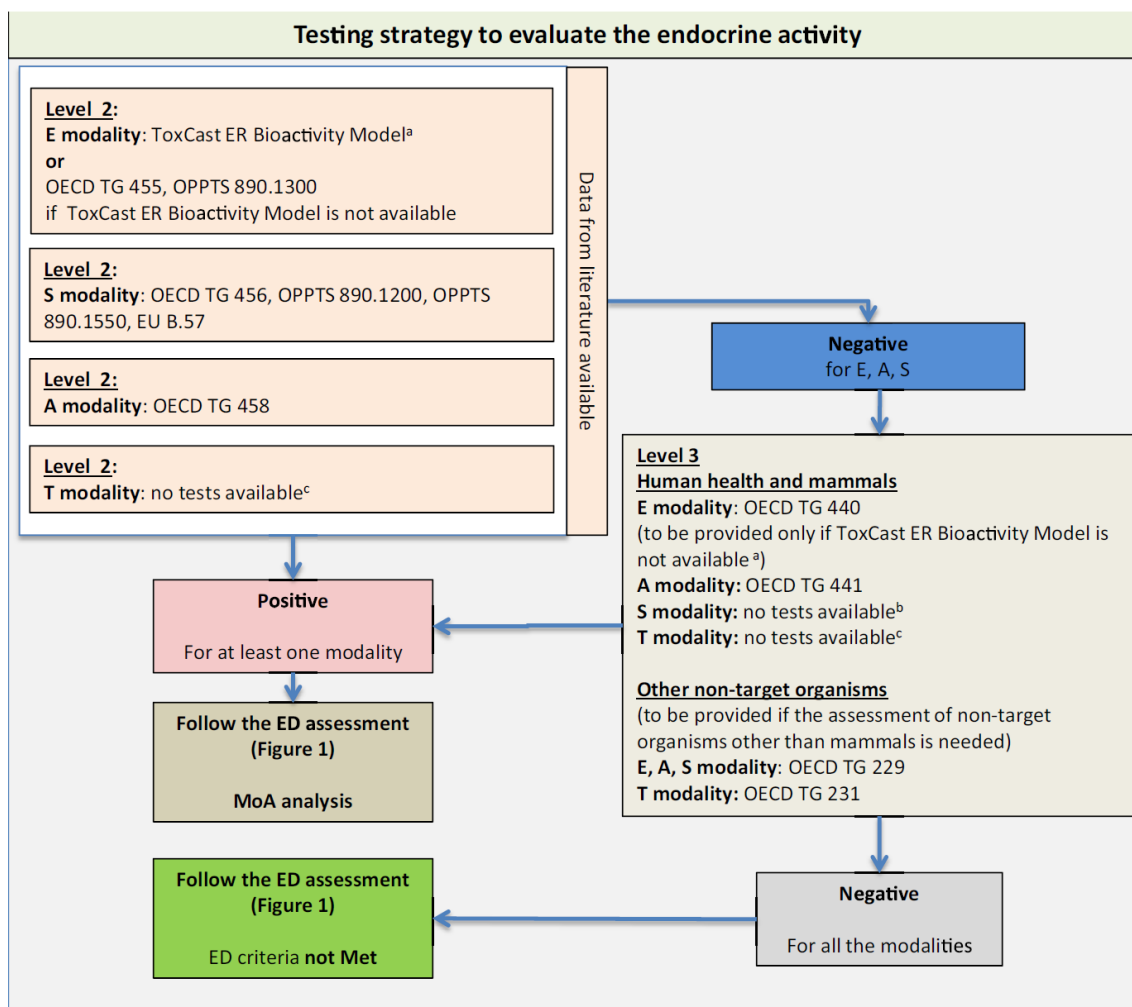
| | |
|--|--|
| <p>Thyroid Function in Intact Juvenile or Peripubertal Male Rats (OPPTS 890.1500).</p> <p>The decision to carry out studies in mammals shall be taken based on all available information, including a systematic review of the literature (including information on endocrine disrupting effects in non-target organisms) and the availability of suitable <i>in silico</i> or <i>in vitro</i> methods</p> | |
|--|--|

1
2 Point 8.13.3 of Annex II to the BPR states that *if there is any information suggesting that the*
3 *active substance may have endocrine disrupting properties* (or if available information is
4 incomplete), *then additional information or specific studies shall be required to elucidate: (1)*
5 *the mode or the mechanism of action and/or; (2) potentially relevant adverse effects in humans*
6 *or animals.* Point 8.13.3.1 of Annex II to the BPR specifies which additional studies to consider.

7 If additional data needs to be generated, there are several test methods available that
8 investigate specific endocrine modalities and/or further investigate potentially endocrine related
9 adverse effects. The decision on which additional studies to carry out depends on what
10 information is missing for a robust conclusion on ED properties.

11 Note that the methods mentioned under 8.13.3.1 (a) generally provide information on adversity,
12 while the systematic literature review from 8.13.3(a)(viii) can provide information on both the
13 adversity and endocrine activity. The studies listed under 8.13.3.1(b), 8.13.3.1(c) and
14 8.13.3.1(d) generally provide information on endocrine activity only, though exceptions may
15 apply.

16 Point 8.13.3.1 of Annex II to the BPR further specifies that, in all cases, *the decision to carry out*
17 *studies in mammals shall be taken based on all available information, including a systematic*
18 *review of the literature (including information on endocrine disrupting effects in non-target*
19 *organisms) and the availability of suitable in silico or in vitro methods.* The ECHA/EFSA guidance
20 recommends to first explore the modality with the strongest positive evidence. However, to
21 exclude ED properties, all EATS modalities must be sufficiently investigated in terms of endocrine
22 activity or endocrine related adversity. In case additional data needs to be generated, and in
23 line with the general desire to limit animal testing as much as possible, it is recommended to
24 investigate endocrine activity first. A general strategy for investigating endocrine activity is given
25 in Figure 5.



1

2 **Figure 5. Strategy to investigate EATS-related endocrine activity in the context of the**
 3 **ED assessment.** From ECHA/EFSA Guidance (2018). Note that the testing strategy also includes
 4 non-mammalian tests: since both mammalian and non-mammalian tests can inform on
 5 endocrine activity, all are included the assessment as discussed in the ED guidance.

6 **Point 8.13.3.1(a).** The existing information might give important information on endocrine
 7 activity and/or disruption *in vivo*, based on the mammalian toxicity studies listed in 8.13.3 (a).
 8 This will most likely be based on parameters measured *in vivo* that may contribute to the
 9 evaluation of adversity, while at the same time (due to the nature of the effect and the existing
 10 knowledge) are also considered indicative of an EATS MoA. Therefore, these endpoints would
 11 imply an underlying endocrine mode of action (in the absence of other explanations). In addition,
 12 some parameters measured *in vivo* may contribute only to the evaluation of adversity, because
 13 on their own, these effects cannot be considered diagnostic of any of the EATS modalities.

14 The existing mammalian *in vivo* information might also provide important information on
 15 endocrine activity. This would be based on parameters that are measured *in vivo* and, while
 16 providing information on endocrine activity, are usually not considered adverse. For example,
 17 changes in hormone levels are considered indicative of perturbation of the endocrine system,
 18 while not necessarily leading to an adverse effect.

19 **Point 8.13.3.1(b).** If further testing is needed for a robust conclusion on ED properties, the
 20 first step in generation new data shall be focused on investigating endocrine activity. The *in vitro*
 21 test methods listed below should be used. All assays are in principle required, unless it is possible
 22 to conclude that the substance meets the ED criteria. This is because each assay investigates a

1 different aspect of endocrine activity. Where new testing is needed, please see also the general
2 information under *Considerations before initiating testing* in chapter 1.

- 3 • *In vitro* assays investigating the E modality
 - 4 ○ OECD TG 455: Performance-based test guideline for stably transfected
 - 5 transactivation *in vitro* assays to detect estrogen receptor agonists and
 - 6 antagonists
- 7 • *In vitro* assays investigating the A modality
 - 8 ○ OECD TG 458: Stably Transfected Human Androgen Receptor Transcriptional
 - 9 Activation Assay for Detection of Androgenic Agonist and Antagonist Activity of
 - 10 Chemicals
- 11 • *In vitro* assays investigating the S modality
 - 12 ○ OECD TG 456: H295R steroidogenesis assay
 - 13 ○ OPPTS 890.1200: The Aromatase assay (human recombinant)
- 14 • *In vitro* assays investigating the T modality
 - 15 ○ Currently there are no validated OECD TGs to investigate the T modality
 - 16 specifically. However, several assays are described in the scientific literature. In
 - 17 addition, repeated dose toxicity studies inform on potential interference with the
 - 18 T modality, i.e. thyroid hormones and HDL/LDL cholesterol levels and the weight
 - 19 and histopathology of the thyroid gland.

20 **Point 8.13.3.1(c).** If the *in vitro* information is positive and sufficient to complete a MoA
21 analysis, additional data might not be needed. However, if the available *in vitro* (and *in silico*)
22 information is negative, the endocrine activity still needs to be further investigated using OECD
23 CF level 3 *in vivo* assays (see Figure 5). Specifically, the following assays should be considered:

- 24 • OECD TG 440: Uterotrophic Bioassay in Rodents: A short-term screening test for
25 oestrogenic properties, see also OECD GD 71 for how to investigate anti-estrogenic
26 effects.
- 27 • OECD TG 441: Hershberger Bioassay in Rats: A Short-term Screening Assay for
28 (Anti)Androgenic Properties, see also OECD GD 115 for how to investigate anti-
29 androgenic effects.

30 Before deciding on the need for *in vivo* testing, a review of the *in vitro* test results and all
31 available information on the toxicokinetic and toxicodynamic profile of the test substance is
32 needed.

33 In some cases, sufficient information might already be available from *in silico* or *in vitro* models
34 to allow a negative conclusion for a specific *in vivo* assay. Currently, this is described in the ED
35 guidance for the ToxCast ER pathway model¹³; whenever a reliable prediction is available for the
36 substance under investigation, the OECD TG 440 does not need to be conducted.

37 **Point 8.13.3.1(d).** The hazard identification of thyroid disruptors is currently hampered by a
38 lack of internationally validated test methods to investigate substance that alter thyroid

¹³ See: <https://comptox.epa.gov/dashboard/dsstoxdb/results?search=DTXSID0020446#bioactivity-toxcast-models>

1 homeostasis. Nevertheless, a data package that fulfils the information requirements should in
 2 most cases be sufficient to conclude on the T mediated adversity for a biocidal active substance.
 3 This is because the data is expected to include an assessment of thyroid histopathology, which
 4 is generally considered to be among the most sensitive and reliable means to detect thyroid
 5 disruption. In 8.13.3.1 (d) *Pubertal development and Thyroid Function in Intact Juvenile or*
 6 *Peripubertal Male Rats (OPPTS 890.1500)* is listed to investigate T mediated effects in more
 7 detail. The male assay is designed to detect interference with both the HPG and HPT axes. As a
 8 result, it will detect substances that interfere with the androgen and thyroid pathways, as well
 9 as effects on steroidogenesis. While the male assays can also detect estrogen receptor mediated
 10 effects, its accuracy on this is currently unknown. Note that while this assay is listed in the
 11 ECHA/EFSA ED guidance, it is not included in the testing strategy for endocrine activity. For a
 12 more detailed discussion on consideration on how to assess the potential for thyroid disruption
 13 for human health, see Appendix A of the ED guidance.

14 **1.13.4. Immunotoxicity and developmental immunotoxicity (ADS)**

15 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.13.4 Immunotoxicity and developmental immunotoxicity</p> <p>If there is any evidence from repeat dose or reproductive toxicity studies that the active substance may have immunotoxic properties, then additional information or specific studies shall be required to elucidate:</p> <p>(1) the mode or the mechanism of action; and/or</p> <p>(2) potentially relevant adverse effects in humans or animals.</p> <p>For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to consider the oral route and conduct animal studies by the oral route</p> | |

16

17 Immunotoxicity investigations should focus on:

18

- The potential to induce adverse effects involving the immune system;

19

20

- with special attention to the adverse immunotoxic outcome among susceptible and vulnerable groups;

21

- clarifying the type of the adverse immunotoxic outcomes when possible;

22

23

- hypersensitivity, immunosuppression, autoimmunity, or unintended stimulation of immune responses;

24

- impact on the developing immune system.

1 **Collection and evaluation of available information**

2 For the assessment of existing information (non-human data: physicochemical properties,
3 grouping, (Q)SARs and expert systems, *in vitro* data; human data and animal data) further
4 guidance is available within the *BPR Volume III Human health Parts B+C* and the *Guidance on*
5 *the Application of the CLP Criteria*).

6 The guidance for the evaluation of all available information before conducting new tests is
7 available in *BPR Volume III Human health Parts B+C* and is largely based on the WHO/IPCS
8 *Guidance on Immunotoxicity for Risk Assessment* (WHO, 2012).

9 Current standard animal studies provide information from an unchallenged immune system,
10 without functional tests, which can give only indications of immunotoxicity. Inclusion of
11 functional tests is needed to adequately assess the immunotoxic potential of active ingredients
12 (WHO/IPCS guidance for Immunotoxicity risk assessment for chemicals (WHO, 2012)).

13 **Generation of new test data**

14 If immunotoxicity potential is identified tests consisting of a more specific confirmatory set of
15 studies or in-depth mechanistic studies, is carried out to confirm and further characterize the
16 endpoint. It is worth noting that further testing to investigate immune function should be
17 conducted only if the outcomes of such studies can be interpreted in relation to the risk
18 assessment for the substance of interest. In addition, the need for further testing to characterise
19 effects of concern for immunotoxicity has to be considered on a case-by-case basis.

20 It should be considered that the conduct of the repeated dose toxicity tests and the reproductive
21 toxicity tests should be performed in a way that allows evaluation of immunotoxicity potential
22 (e.g. Repeated dose toxicity according to US EPA OPPTS 870.7800 [Health Effects Test
23 Guidelines Immunotoxicity] including parameters for immunotoxicity and OECD TG 443 -
24 extended one generation toxicity test- may be conducted with the developmental immunotoxicity
25 cohort). However, a separate study may be needed for confirmatory results of developmental
26 immunotoxicity.

27 Whether the immunotoxic properties should be investigated in adults or in the developing
28 organisms, or both, should be considered on a case by case basis taking into account the various
29 aspects affecting the decision, for example, the target population, toxicokinetics and mode of
30 action. Generally, a study in developing organisms is recommended as a more conservative
31 approach.

32 Immunotoxicity observed in animals exposed during adulthood only may trigger the need to
33 investigate also potential for developmental immunotoxicity unless substance specific
34 information is provided why these effects or mode of action would not be relevant in developing
35 organism. In addition, if the classification criteria for STOT are met, based on studies in adults,
36 this is not an adaptation rule allowing the omission of investigations on developmental
37 immunotoxicity but rather a trigger for a concern. This is due to expected higher sensitivity of
38 the developing organisms (see e.g. Dietert, 2014), which may lead to a lower point of departure
39 and/or to hazard classification for development.

40 A classification to Repr. 1B or 2 may be necessary if the effects are considered to be of
41 developmental origin, i.e. due to exposure during development. Sensitivity has been evaluated
42 in animal studies for nine reviewed (immuno)toxicants and, according to the authors, the
43 developing immune system was found to be at least as sensitive or more sensitive than the
44 general (developmental) toxicity parameters (Hessel et al., 2015).

45 The test methods to be used for further immunotoxicity studies will depend also on the weight
46 of evidence analysis. Different test methods can be employed for assessing immune suppression,
47 immune stimulation and autoimmunity as well as developmental immunotoxicity.

1 Reviews of principles and methods for immunotoxicity are available from WHO/IPCS:

- 2 • WHO/IPCS Environmental Health Criteria (EHC) 180, Principles and Methods for Assessing
3 Direct Immunotoxicity Associated with Exposure to Chemicals (WHO, 1996)
- 4 • WHO/IPCS Environmental Health Criteria (EHC) 212, Principles and Methods for Assessing
5 Allergic Hypersensitization Associated with Exposure to Chemicals (WHO, 1999)
- 6 • WHO/IPCS Environmental Health Criteria (EHC) 236, Principles and Methods for Assessing
7 Autoimmunity Associated with Exposure to Chemicals (WHO, 2007)
- 8 • WHO/IPCS Guidance for immunotoxicity risk assessment for chemicals, Harmonisation
9 project document No 10 (WHO, 2012)

10 Below a list of methods that can be considered for further immunotoxicity testing is provided.
11 This list is not exhaustive but provides the methodological aspects to consider on a case-by-case
12 basis.

13 **Immune Suppression**

- 14 • US EPA OPPTS 870.7800 Health Effects Test Guidelines Immunotoxicity
- 15 • Functional investigations as described under Additional Immunotoxicity Testing below

16 **Immune stimulation including hypersensitivity (skin and respiratory sensitisation)**

- 17 • LLNA assay (see sensitisation section)
- 18 • Functional investigations as described under Additional Immunotoxicity Testing below

19 **Autoimmunity**

- 20 • Functional investigations as described under Additional Immunotoxicity Testing below

21 **Additional Immunotoxicity Testing (adopted from ICH S8)**

- 22 • T-cell Dependent Antibody Response (TDAR)
- 23 • Immunophenotyping
- 24 • Natural Killer Cell Activity Assays
- 25 • Host Resistance Studies
- 26 • Macrophage/Neutrophil Function
- 27 • Assays to Measure Cell-Mediated Immunity

28 **Developmental Immunotoxicity**

- 29 • Protocols for independent developmental immunotoxicity studies with exposure during
30 development and functional investigations (such as described under Additional
31 Immunotoxicity Testing above) during development and/or adulthood
- 32 • Developmental immunotoxicity cohort in an OECD Test Guideline 443: Extended One-
33 Generation Reproductive Toxicity Study

1 Developmental immunotoxicity studies are designed to provide information on the potential
2 functional and morphological hazards to the immune system arising in the offspring from
3 exposure of the mother during pregnancy and lactation. For an independent developmental
4 immunotoxicity study there is currently no available internationally accepted protocol, such as
5 an OECD TG. However, protocols, considerations and recommendations for independent
6 developmental immunotoxicity studies have been published e.g. by Ogungbesan et al., (2019),
7 Boverhof et al., (2014), Collinge et al., (2012), WHO (2012), DeWitt et al., (2012a and 2012b),
8 Gupta (2011, page 219-225), Dietert and DeWitt (2010), Rooney et al., (2009), De Jong and
9 Van Loveren (2007), Dietert and Holsapple (2007), Holsapple et al., (2005). These studies
10 investigate changes in immune response due to effects on the innate or acquired immune
11 system. As immune response may also be affected by the function of other organs such as liver,
12 kidneys and the endocrine system, toxic effects on these organs in offspring may also be
13 reflected in changes in immune response. No single immune parameter is able to reflect the
14 entire complex and intricate function of immune system and so, integration of findings of
15 different tests is relevant to evaluate the relevance of the results on substance exposure.

16 The selection between the choices should be based on scientific and substance specific
17 considerations taking into account which method adequately addresses the scientific concern
18 with least amount of animals and investigations.

19 Some examples of aspects of these considerations are presented below. The nature and/or
20 severity of the identified concern may provide guidance to select between a separate study or
21 inclusion of parameters to other studies or a Cohort 3 in an OECD TG 443. Other aspects to
22 consider may include statistical power and the investigations included. It should be considered
23 whether the parameters/Cohort 3 or a separate study best address the particular concern
24 identified. The outcome of a separate developmental immunotoxicity study may differ from that
25 of the developmental immunotoxicity Cohort 3 in an OECD TG 443, if the exposure scenarios
26 and set ups are different.

27 Important aspects to be considered for study designs are 1) sufficient statistical power, 2)
28 separate analysis for males and females to assess potential sex differences, 3) selection of
29 sensitive parameters, and 4) selection of representative time points for each investigation, and
30 continuous exposure starting from implantation until investigations of immune parameters.

31 Although it is possible to combine the investigations for developmental immunotoxicity with
32 reproduction toxicity studies, this approach may limit the statistical power (number of animals
33 available; e.g. OECD TG 443) and investigations for sex differences. Furthermore, dose level
34 setting of a study for sexual function and fertility may not be optimal for investigating
35 developmental immunotoxicity.

36 As a common recommendation the test battery should include the following investigations:

- 37 a) Humoral immunity / antibody formation: T-cell dependent antibody response (TDAR) –
38 PND 45 or older;
- 39 b) Cell-mediated (antigen-specific) immune responses: Delayed type hypersensitivity assay
40 (DTH); *AND/OR* Cytotoxic T-lymphocyte (CTL) response; *AND/OR* NK cell assay;
- 41 c) Lymphoid organ weights (considered important to characterize effects; to be assessed
42 together with a) and b));
- 43 d) Histopathology of immune organs;
- 44 e) Supporting information: haematology, cytokine production, flow cytometric
45 immunophenotyping of lymphocyte sub-populations.

46 Developmental immunotoxicity investigations in an OECD TG 443 with DIT cohort (Cohort 3)

1 investigates less parameters with limited statistical power. The parameters investigated are
2 TDAR in Cohort 3 (10 males and 10 females), and lymphoid organ weights, histopathology, and
3 splenic lymphocyte subpopulation analysis in Cohort 1A (CD4+ and CD8+ T lymphocytes, B
4 lymphocytes, and natural killer cells). Cohort 3 contains 10 males and 10 females from different
5 litters where possible per group and TDAR (IgM) is investigated at PND 56±3. For lymph node,
6 bone marrow and splenic lymphocyte analysis the statistical power is 10 animals/sex /group in
7 Cohort 1A, for other lymphoid organs (thymus, spleen and the adrenal glands) the statistical
8 power is 20 animals/sex/group. Investigation from Cohort 1A are done at postnatal week 13.

9 Due to limited parameters and statistical power, the results from Cohort 3 in an OECD TG 443
10 cannot be considered as definitive but rather as screening results which may lead to confirmative
11 investigations. Therefore, where a concern for developmental immunotoxicity is identified, it is
12 recommended to investigate this using a testing battery described above with a sufficient
13 statistical power such as 20 animals/sex/group (representing 20 litters). Due to lack of OECD
14 TG for DIT, a detailed description of the test method used should be given with justifications for
15 the selected investigations and conditions.

16 Effects considered as adverse will be relevant to hazard classification and the human health risk
17 assessment, providing an N(L)OAEL, unless there is information to show that effects seen in
18 these studies could not occur in humans. Due to a complexity of the endpoint, adversity should
19 preferably be based on a holistic analysis of data by grouping similar parameters.

20 **Note regarding developmental immunotoxicity and assessment of endocrine**
21 **disruption**

22 Sex differences in effects may indicate hormonal co-influence to the parameter measured.

23 **1.13.5. Further mechanistic studies (ADS)**

24 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.13.5 Further mechanistic studies A decision on the need to perform additional studies should be based on all relevant data | |

25 This data may be relevant in the weight of evidence assessment with other information to assess
26 the toxicological properties of a substance, as it can provide information on the mode of action
27 (MoA) of the chemical. It can also provide information that can be used for refinement in the
28 evaluation of mixtures.
29

30 Studies of the mechanisms of toxicity/mode of action may provide useful supplementary
31 information when there are indications that the active substance may have effects on e.g.
32 carcinogenicity (genotoxic and non-genotoxic MoAs are relevant for the classification of
33 carcinogens in accordance with CLP), reproduction, neurotoxicity or immunotoxicity. Such
34 studies may in some cases be used in concluding that the effects observed in experimental
35 animals are not relevant to humans. For ED identification mechanistic studies may be needed
36 (see section 1.13.3). In addition, information on the MoA/mechanisms may clarify the observed
37 sex differences in effects (potential information on endocrine activity), differences between
38 toxicity in different life stages (e.g., sensitivity during development or elderly animals), or an
39 underlying cause (e.g., immunotoxicity) for other effects.

40 As a general principle, the effects observed in animal studies are considered relevant for humans
41 unless there is sufficient information to prove the contrary. In order to conclude that the adverse

1 effects are not relevant for humans, it is necessary to establish that the adverse effects are
2 caused by a MoA that is not relevant for humans and it must be also possible to exclude other
3 MoAs for the adverse effects seen.

4 **1.14. Studies related to the exposure of humans to the active** 5 **substance (ADS)**

6 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.14 Studies related to the exposure of humans to the active substance | |

7
8 Toxicity of degradation products, by-products and reaction products related to human exposure.

9 Information is required on the toxic effects of substances generated from an active substance,
10 other than mammalian metabolites, in normal use of biocidal product.

11 The decision as to the need for these data should be made on a case-by-case basis by expert
12 judgment. Where human exposure is significant, toxicity testing may be needed.

13 These data may be relevant for many product-types for example: product-types 1 and 2
14 (reaction products with water when the substance is used for human hygiene purposes or
15 reaction products with water or other materials released in water or air when the substance is
16 used for the treatment of bathing waters), product-type 5 (substances produced in a reaction
17 with drinking water), product-types 6, 7, 9 and 10 (residuals in treated materials), product-type
18 8 (irritating and sensitising effects of chemical compounds, such as metal salts, developed on
19 the surface of the treated wood) and product-type 18 (products, which may produce harmful
20 substances with water during gassing).

21 **1.15. Toxic effects on livestock and pets (ADS)**

22 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.15 Toxic effects on livestock and pets | |

23
24 For livestock and pets, an estimation of toxic effects and exposure via different exposure routes
25 (e.g. inhalation, licking, skin contact and ingestion of poisoned bait) is required. In exceptional
26 cases, toxicity testing in livestock and pets may be required. Toxic effects for livestock and pets
27 should be estimated or studied if the substance is to be used in spaces in which animals are
28 housed, kept or transported or exposure is possible via drinking water or feeding stuffs.
29 Information on lethal doses for different species, symptoms of poisoning, details of the time
30 courses in case of poisoning and antidotes should also be submitted, if available.

31
32 These data may be relevant e.g. for the following product-types:

- 33
34
35
36
- 3 (substances used for veterinary hygiene purposes)
 - 4 (disinfection of surfaces and equipment)
 - 5 (drinking water)

- 1 • 8, 10 (treated materials in areas in which animals are housed, kept or transported)
- 2 • 14, 15, 23 (ingestion of baits)
- 3 • 16, 17 (contaminated drinking water)
- 4 • 18, 19 (repellents to be used for veterinary hygiene purposes, residential indoor use).

5 **1.16. Food and feeding stuffs studies including for food producing** 6 **animals and their products (milk, eggs and honey) (ADS)**

7 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.16 Food and feeding stuffs studies including for food-producing animals and their products (milk, eggs and honey) Additional information related to the exposure of humans to the active substance contained in biocidal products | |

8

9 Evaluation of residues in food and feed from biocidal uses requires information on the nature of
 10 residues as well as quantification of residues, which is covered by data requirements listed under
 11 this endpoint in Annex II of the BPR (and the endpoint 8.10 in Annex III of the BPR). Normally
 12 standard residue study with radiolabeled compounds or other study providing equivalent
 13 information, designed to reflect the realistic use conditions of the biocidal product, would be
 14 necessary to identify residue composition. Chapter 5 of the BPR Guidance Volume III Human
 15 Health - Assessment & Evaluation (Parts B+C) and Guideline on risk characterisation and
 16 assessment of maximum residue limits (MRL) for biocides (EMA/CVMP/SWP/90250/2010),
 17 provides indications on the guidelines that would support the identification of the residue
 18 composition. The guidance recommends applying OECD TG 507, "Nature of the Pesticide
 19 Residues in Processed Commodities – High Temperature Hydrolysis".

20 Dietary Risk Assessment (DRA) follows a stepwise approach with each step leading to a more
 21 realistic estimate of residue amounts in foods. Lower-level steps generally involve calculation
 22 models populated with default values in the first tier, with the possibility of including additional
 23 data in higher tiers. With few exceptions, data from product- and use-specific residue studies
 24 with foods are only necessary if lower tiers fail to exclude a consumer risk. In addition, Maximum
 25 Residue Limits (MRLs) must be set according to the criteria outlined in the Commission Note¹⁴.

26 The basic use categories for DRA are "animal husbandry", "biocide-food contact (professional
 27 use)" and "biocide-food contact (non-professional use)". Depending on the use category,
 28 different calculation models and residue study designs apply. While some required information,
 29 e.g. metabolism in livestock and degradation during food processing is related to the active
 30 substance itself, other data are connected to the intended use of the respective biocidal product
 31 (e.g. supervised residue trials). The former can be submitted at the stage of the evaluation for

¹⁴ CA-March17-Doc.7.6.c-Final: *An interim approach for the establishment of maximum residue limits for residues of active substances contained in biocidal products for food and feed and specific migration limits in food contact materials.* See link under "related links" in <https://echa.europa.eu/guidance-documents/guidance-on-biocides-legislation>.

Direct link:

https://ec.europa.eu/health/sites/health/files/biocides/docs/2017_interimapproach_maximumresiduelimits_en.pdf.

1 active substance approval, while the latter must be generated at the product authorisation stage.
2 Guidance to address the non-professional uses (Chapter 5) and animal husbandry (Chapter 6)
3 are included in the BPR Guidance Volume III Human Health - Assessment & Evaluation (Parts
4 B+C). Guidance on professional use is under development and will be included as an additional
5 chapter in the same guidance. These guidance documents provide the methodology to estimate
6 the transfer of biocidal active residues into food and indications on the studies to be performed
7 to allow the identification of the residues and to support refinement options (for example, studies
8 to allow the quantification of transfer factor or to estimate the rinsing efficiency). Apart from the
9 above, there is currently no guidance for dietary risk assessment specifically for biocides.
10 Methodologies developed by other Agencies may be used to perform dietary risk assessment. In
11 addition, guidance documents developed by other Agencies, e.g. on metabolism in livestock and
12 degradation during food processing, may be used to support the assessment for biocides.

13 **1.16.1. Proposed acceptable residue levels i.e. maximum residue limits (MRL)**
14 **and the justification of their acceptability (ADS)**

15 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.16.1 Proposed acceptable residue levels i.e. maximum residue limits (MRL) and the justification of their acceptability | |

16
17 For product-type 5, any relevant regulations relating to acceptable or unacceptable residues in
18 drinking water must be taken into consideration in the justification.

19
20 For product-type 21, any directions or restrictions at the Community or national level related to
21 residues in fish and shellfish intended to be used as food or feeding stuffs must be taken into
22 consideration in the justification.

23
24 Please refer also to the Commission Note¹⁷ above.

25
26 **1.16.2. Behaviour of the residue of the active substance, its degradation**
27 **products and, where relevant, its metabolites on the treated or contaminated**
28 **food or feeding stuffs including the kinetics of disappearance (ADS)**

29 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.16.2 Behaviour of the residue of the active substance on the treated or contaminated food or feeding stuffs including the kinetics of disappearance Residue definitions should be provided where relevant. It is also important to compare residues found in toxicity studies with residues formed in food-producing animals and their products, as well | |

| | |
|------------------|--|
| as food and feed | |
|------------------|--|

1
2 *Residue definitions should be provided where relevant. It is also important to compare residues*
3 *found in toxicity studies with residues formed in food-producing animals, their product as well*
4 *as food and feed.*

5
6 *Residue definition should be provided when indirect exposure via food cannot be excluded.*

7 **1.16.3. Overall material balance for the active substance (ADS)**

8 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.16.3 Overall material balance for the active substance Sufficient residue data from supervised trials on food-producing animals and their products, as well as food and feed, to demonstrate that residues likely to arise from the proposed use would not be of concern for human or animal health | |

10
11 Point 8.16.3 of Annex II to the BPR states that *sufficient residue data from supervised trials on*
12 *food producing species and their products as well as food and feed to demonstrate that residues*
13 *likely to arise from the proposed use would not be of concern for human or animal health.*

14 **1.16.4. Estimation of potential or actual exposure of the active substance to** 15 **humans through diet and other means (ADS)**

16 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.16.4 Estimation of potential or actual exposure of humans to the active substance and residues through diet and other means | |

17
18
19 Expected consumer exposure via diet should be studied taking into account the average
20 consumption of different food types and drinking water.

21 **1.16.5. If residues of the active substance remain on feeding stuffs for a** 22 **significant period of time or also residues found in food of animal origin after** 23 **treatment on or around food producing animals (ADS)**

24 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|----------------------|---|
| | |

| | |
|---|--|
| <p>8.16.5 If residues of the active substance occur in or on feeding stuffs for a significant period of time or are found in food of animal origin after treatment on or around food-producing animals (e.g. direct treatment on animals or indirect treatment of animal houses or surroundings) then feeding and metabolism studies in livestock shall be required to permit evaluation of residues in food of animal origin</p> | |
|---|--|

1
2 Point 8.16.5 of Annex II to the BPR states that [...] (e.g. direct treatment on animals or indirect
3 treatment of animal houses or surroundings) then feeding and metabolism studies in livestock
4 shall be required to permit evaluation of residues in food of animal origin.
5

6 **1.16.6. Effects of industrial processing and/or domestic preparation on the
7 nature and magnitude of residues of the active substance**

8 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.16.6 Effects of industrial processing and/or domestic preparation on the nature and magnitude of residues of the active substance</p> | |

9
10 Provide information as implied by the title.

11
12 **1.16.7. Any other available information that is relevant (ADS)**

13 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.16.7 Any other available information that is relevant</p> <p>It may be appropriate to include information on migration into food, especially in the case of treatment of food contact materials</p> | |

14
15 Point 8.16.3 of Annex II to the BPR states that *it may be appropriate to include information on
16 migration into food, especially in the case of treatment of food contact materials.*

17
18 For instance information from other chemical programmes on ADI, MRL or relevant residues.

19
20 **1.16.8. Summary and evaluation of data submitted under 8.16.1. to 8.16.7.
21 (ADS)**

1 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.16.8 Summary and evaluation of data submitted under 8.16.1 to 8.16.8</p> <p>It is important to establish whether the metabolites found in food (from animals or plants) are the same as those tested in toxicity studies.</p> <p>Otherwise values for risk assessment (e.g. ADI) are not valid for the residues found</p> | |

2
3 Please follow the guidance in section 1.16 of this guidance.

4
5 **1.17. Tests to assess toxic effects of metabolites from treated plants**
6 **(ADS)**

7 **Information requirement according to BPR Annex II:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| <p>8.17 If the active substance is to be used in products for action against plants including algae then tests shall be required to assess toxic effects of metabolites from treated plants, if any, where different from those identified in animals</p> | |

8
9 This point on action against plants is considered as covered sufficiently by Regulation (EC) No
10 1107/2009 (PPPR) together with Regulations (EU) 283/2013 and (EU) 284/2013.

2. Dossier Requirements for Biocidal Products BPR Annex III, Title 1, 8 Toxicological Profile for humans and animals

Toxicological profile for humans and animals

This section describes the information requirements for biocidal products for the assessment of the toxicological profile for humans and animals.

Where new testing is needed, please see also the general information under *Considerations before initiating testing* in chapter 1.

2.1. Skin corrosion or irritation

Information requirement according to BPR Annex III:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| <p>8.1 Skin corrosion or irritation</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) skin corrosion, <i>in vitro</i> testing;</p> <p>(c) skin irritation, <i>in vitro</i> testing;</p> <p>(d) skin corrosion or irritation, <i>in vivo</i> testing</p> | <p>Testing of the product or mixture does not need to be conducted if:</p> <ul style="list-style-type: none"> – there are sufficient valid data on each component of the product or mixture to allow its classification in accordance with Regulation (EC) No 1272/2008, and synergistic effects between any of the components are not expected, – the product or mixture is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$), – the product or mixture is spontaneously flammable in air or in contact with water or moisture at room temperature, – the product or mixture meets the classification criteria for acute toxicity category 1 by the dermal route, or – an acute toxicity study by the dermal route provides conclusive evidence on skin corrosion or irritation adequate for classification. <p>If results from one of the two studies listed in points (b) or (c) in column 1 of this row already allow conclusive decision on the classification of product or mixture or on the absence of skin irritation potential, the second study does not need to be conducted</p> <p>An <i>in vivo</i> study for skin corrosion or irritation shall be considered only if the <i>in vitro</i> studies listed in points (b) and (c) in column 1 of this row are not applicable, or the results of these studies are not adequate for classification and risk assessment and the calculation method or bridging principles laid down in Regulation (EC) No 1272/2008 are not applicable</p> <p><i>In vivo</i> studies for skin corrosion or irritation that were carried out or initiated before 15 April 2022 shall be considered appropriate to address this information</p> |

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| | requirement |
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1
2 Please follow section 1.1 of this guidance.

3 4 2.2. Serious eye damage or eye irritation

5 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| <p>8.2 Serious eye damage or eye irritation</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) serious eye damage or eye irritation, <i>in vitro</i> testing;</p> <p>(c) serious eye damage or eye irritation, <i>in vivo</i> testing</p> | <p>Testing on the product or mixture does not need to be conducted if:</p> <ul style="list-style-type: none"> —there are sufficient valid data available on each component of the product or mixture to allow its classification in accordance with Regulation (EC) No 1272/2008, and synergistic effects between any of the components are not expected, — the product or mixture is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$), —the product or mixture is spontaneously flammable in air or in contact with water or moisture at room temperature, or — the product or mixture meets the classification criteria for skin corrosion leading to its classification as “serious eye damage” category 1 <p>If results from a first <i>in vitro</i> study do not allow a conclusive decision on the classification of the product or mixture or on the absence of eye irritation potential (an)other(s) <i>in vitro</i> study(ies) for this endpoint shall be considered</p> <p>An <i>in vivo</i> study for serious eye damage or eye irritation shall be considered only if the <i>in vitro</i> study(ies) under point (b) in column 1 of this row are not applicable, or the results obtained from these studies are not adequate for classification and risk assessment and the calculation method or bridging principles laid down in Regulation (EC) No 1272/2008 are not applicable</p> <p><i>In vivo</i> studies for serious eye damage or eye irritation that were carried out or initiated before 15 April 2022 shall be considered appropriate to address this information requirement</p> |

6
7 Please follow section 1.2 of this guidance.

8 9 2.3. Skin sensitisation

10 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| <p>8.3 Skin sensitisation</p> <p>The information shall allow to conclude whether the substance is a skin sensitizer and whether it can be presumed to have the potential to produce significant sensitisation in humans (Category 1A). The information should be sufficient to perform a risk assessment where required</p> <p>The assessment shall comprise the following tiers:</p> <p>(a) assessment of the available human, animal and non-animal data;</p> <p>(b) skin sensitisation, <i>in vitro</i> testing. Information from <i>in vitro</i> or <i>in chemico</i> test method(s) conducted in accordance with point 5 of the introductory part of this Annex and addressing each of the following key events of skin sensitisation:</p> <p>(i) molecular interaction with skin proteins;</p> <p>(ii) inflammatory response in keratinocytes;</p> <p>(iii) activation of dendritic cells.</p> <p>(c) skin sensitisation <i>in vivo</i> testing. The Murine Local Lymph Node Assay (LLNA) is the first-choice method for <i>in vivo</i> testing. Another skin sensitisation test may only be used in exceptional circumstances. If another skin sensitisation test is used, scientific justification shall be provided.</p> | <p>Testing on the product or mixture does not need to be conducted if:</p> <ul style="list-style-type: none"> – there are sufficient valid data available on each component of the product or mixture to allow its classification in accordance with Regulation (EC) No 1272/2008, and synergistic effects between any of the components are not expected, – the available information indicates that the product or mixture should be classified for skin sensitisation or skin corrosion, – the product or mixture is a strong acid ($\text{pH} \leq 2,0$) or base ($\text{pH} \geq 11,5$), or – the product or mixture is spontaneously flammable in air or in contact with water or moisture at room temperature. <p><i>In vitro</i> tests do not need to be conducted if:</p> <ul style="list-style-type: none"> – an <i>in vivo</i> study referred to in point (c) in column 1 of this row is available, or – the available <i>in vitro</i> or <i>in chemico</i> test methods are not applicable for the product or mixture or the results obtained from these studies are not adequate for classification and risk assessment. <p>If information from test method(s) addressing one or two of the key events described in point (b) in column 1 of this row already allows for classification of the substance and risk assessment, studies addressing the other key event(s) do not need to be conducted</p> <p>An <i>in vivo</i> study for skin sensitisation shall be considered only if <i>in vitro</i> or <i>in chemico</i> studies referred to in point (b) in column 1 of this row are not applicable, or the results obtained from these studies are not adequate for classification and risk assessment and the calculation method or bridging principles laid down in Regulation (EC) No 1272/2008 are not applicable</p> <p><i>In vivo</i> studies for skin sensitisation that were carried out or initiated before 15 April 2022 shall be considered appropriate to address this information requirement'</p> |
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2 Please follow section 1.3 of this guidance.

3 Any limitation of the additivity method specified in the Guidance on the Application of the CLP
4 Criteria (ECHA) for sensitisation with regard to addressing sub-corrosive concentrations with
5 sensitising potential should also be considered (see also section 1.3 of this guidance).

6 2.4. Respiratory sensitisation and irritation

7 2.4.1. Respiratory sensitisation (ADS)

8 Information requirement according to BPR Annex III:

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| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD |
|----------------------|---|

| | INFORMATION |
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| 8.4 Respiratory sensitisation | <p>Testing on the product/mixture does not need to be conducted if:</p> <ul style="list-style-type: none"> — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected |

Please follow section 1.4.1 of this guidance.

2.4.2. Respiratory irritation (not in BPR Annex III)

Please follow section 1.4.2 of this guidance.

2.5. Acute toxicity

Information requirement according to BPR Annex III:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| <p>8.5 Acute toxicity</p> <ul style="list-style-type: none"> — Classification using the tiered approach to classification of mixtures for acute toxicity in Regulation (EC) No 1272/2008 is the default approach | <p>Testing on the product/mixture does not need to be conducted if:</p> <ul style="list-style-type: none"> — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected |

Please follow section 1.7 of this guidance.

2.5.1. By oral route

Information requirement according to BPR Annex III:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|----------------------|---|
| 8.5.1 By oral route | |

Please follow section 1.7.1 of this guidance.

2.5.2. By inhalation

Information requirement according to BPR Annex III:

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|----------------------|---|
| 8.5.2 By inhalation | |

1 Please follow section 1.7.2 of this guidance.
2

3 **2.5.3. By dermal route**

4 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| 8.5.3 By dermal route | |

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6 Please follow section 1.7.3 of this guidance.
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8 **2.5.4. Biocidal products that are intended to be authorised for use with other biocidal products**

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10 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|---|---|
| 8.5.4 For biocidal products that are intended to be authorised for use with other biocidal products, the risks to human health, animal health and the environment arising from the use of these product combinations shall be assessed. As an alternative to acute toxicity studies, calculations can be used. In some cases, for example where there are no valid data available of the kind set out in column 3, this may require a limited number of acute toxicity studies to be carried out using combinations of the products | |

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12 **2.6. Information on dermal absorption**

13 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| 8.6 Information on dermal absorption Information on dermal absorption when exposure occurs to the biocidal product. The assessment of this endpoint shall proceed using a tiered approach | |

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15 It is not always mandatory to submit experimental data. If such data are not available, as a first

1 step default values can be used according to the EFSA Guidance Document on Dermal Absorption
2 (EFSA, 2017). The OECD Guidance Document on Percutaneous absorption/penetration (OECD,
3 2004a) and the EFSA Guidance on dermal absorption (EFSA, 2017) should be followed where
4 applicable for the estimation of dermal absorption both for the biocidal product and the active
5 substance (section 1.8 of this guidance).

6 The following Test Guidelines are available for the conduct of skin absorption studies:

- 7 • EC method B.45 Skin Absorption: *In Vitro* Method.
- 8 • OECD Test Guideline 428: Skin Absorption: *In Vitro* Method.
- 9 • EC method B.44 Skin Absorption: *In Vivo* Method.
- 10 • OECD Test Guideline 427: Skin Absorption: *In Vivo* Method.

11 If testing to assess the likely magnitude and rate of dermal bioavailability is necessary, the OECD
12 Test Guideline 428 for *in vitro* skin absorption should be considered first.

13 Before new studies are commenced, it should be checked whether the intended use is safe when
14 the appropriate default value is applied. If no experimental data are available, studies with
15 similar formulations should be looked for. If valid studies have been performed with the same
16 formulation for which authorisation is to be granted, these results should be used with a
17 preference for an *in vitro* study on human skin.

18 Dermal absorption can be measured *in vitro* and/or *in vivo*. If valid studies with the relevant
19 formulation are available, their results should be directly used for risk assessment. However,
20 any deviations from OECD TG 427 and OECD TG 428 require justification, including an
21 assessment of the impact of the deviation. Acceptable studies should be in full compliance with
22 OECD test guidelines 427 (*in vivo*) or 428 (*in vitro*) or at least similar to them in all main aspects,
23 based on expert judgement. The applicant should ensure that all relevant information is provided
24 in the study report, e.g. regarding the use of tape stripping. It must be acknowledged that both
25 guidelines leave a certain degree of freedom to modify the study design.

26 When valid (guideline-compliant and GLP) *in vitro* studies on human skin, *in vitro* studies in
27 animals and *in vivo* animal studies are available and conducted under the same experimental
28 conditions, and the results meet the quality criteria, in particular with respect to variability,
29 number of acceptable replicates and recovery, then the 'Triple Pack' approach can be used to
30 extrapolate the human dermal absorption values for risk assessment (OECD No. 156, draft). *In*
31 *vitro* studies on human skin are however considered sufficiently predictive and conservative and
32 should be normally used for the risk assessment – a complete "triple pack" including testing in
33 living animals is not required but available triple pack data may be used to refine the assessment.
34 *In vivo* studies on rats or *in vitro* studies on rat skin as "stand alone" information may also be
35 used, acknowledging however that this will result in clear overestimation of dermal absorption
36 in humans in the vast majority of cases.

37 Percutaneous absorption depends on the partitioning of substances from the vehicle and
38 solubility in the vehicle. OECD TG 427 and TG 428 recommend conducting tests using test
39 preparations that are the same as (or a realistic surrogate to) those that humans may be exposed
40 to.

41 Other types of studies (e.g. in human volunteers) could be taken into consideration in
42 exceptional cases but in general their use is not recommended.

43 In some cases, it may also be possible to estimate dermal absorption on the basis of existing
44 information that comes from other sources. Mostly, this will be extrapolation of experimental
45 data obtained with a similar formulation, but in this case strict and transparent rules should be

1 followed as to when another formulation or product can be considered similar. Expert judgment
2 will always be needed in these cases. A detailed justification and expert judgment is necessary
3 if less frequently used approaches are used, such as the application of QSARs or a comparison
4 of the results obtained in oral and dermal toxicity studies.

5 Dermal absorption can vary depending on the formulation, as well as due to other products that
6 are present on the skin. This is most relevant for biocidal products that are applied on the skin.
7 Any information of such interactions should be included in the assessment. This would normally
8 be considered in the need of risk management measures to avoid increased systemic exposure
9 due to other products that enhance dermal absorption.

10 **2.7. Available toxicological data relating to non-active substances (i.e.**
11 **substances of concern) and a mixture that a substance of concern is a**
12 **component of**

13 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|--|
| <p>8.7 Available toxicological data relating to:</p> <p>(a) non-active substance(s) (i.e. substance(s) of concern); and</p> <p>(b) a mixture that a substance(s) of concern is a component of</p> <p>Tests listed in Section 8 of the table in Title 1 of Annex II shall be carried out for the substance(s) of concern or a mixture that a substance(s) of concern is a component of if insufficient data are available and cannot be inferred through read-across, <i>in silico</i> or other accepted non-testing approaches</p> | <p>Testing on the product or mixture does not need to be conducted if all of the following conditions are met:</p> <ul style="list-style-type: none"> – there are valid data available on each of the components in the mixture to allow classification of the mixture in accordance with the rules laid down in Regulation (EC) No 1272/2008, – a conclusion can be made whether the biocidal product can be considered as having endocrine disrupting properties, – synergistic effects between any of the components are not expected' |

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15 **2.8. Food and feedingstuffs studies (ADS)**

16 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|------------------------------------|---|
| 8.8 Food and feedingstuffs studies | |

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18 **2.8.1. Feeding and metabolism studies in livestock (ADS)**

19 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| 8.8.1 If residues of the biocidal | |

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| product remain in or on feedingstuffs for a significant period of time, then feeding and metabolism studies in livestock shall be required to permit evaluation of residues in food of animal origin | |
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2 Please follow section 1.16 of this guidance.

3 **2.8.2. Residues in food (not in BPR Annex III)**4 If intended use of the biocidal product may lead to transfer of residues into foods, studies on the
5 nature of residues and studies on residue levels may be required.

6 Please follow section 1.16 of this guidance.

7 **2.9. Effects of industrial processing and/or domestic preparation on**
8 **the nature and magnitude of residues of the biocidal product (ADS)**9 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
|--|---|
| 8.9 Effects of industrial processing and/or domestic preparation on the nature and magnitude of residues of the biocidal product | |

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11 The objective of these studies is to establish whether breakdown or reaction products arise from
12 residues in the raw products during processing which may require a separate risk assessment.13 Depending on the level and chemical nature of the residue in the raw commodity, a set of
14 representative hydrolysis situations (simulating the relevant processing operations) should be
15 investigated, where appropriate. The effects of process other than hydrolysis may also have to
16 be investigated, where the properties of the active substance or metabolites indicate that
17 toxicologically significant degradation products may occur as a result of these processes. The
18 studies are normally conducted with a radio-labelled form of the active substance.

19 Please follow section 1.16 of this guidance.

20 **2.10. Other test(s) related to the exposure to humans (ADS)**21 **Information requirement according to BPR Annex III:**

| INFORMATION REQUIRED | SPECIFIC RULES FOR ADAPTATION FROM STANDARD INFORMATION |
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| 8.10 Other test(s) related to the exposure to humans Suitable test(s) and a reasoned case will be required for the biocidal product | |

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| In addition, for certain biocides which are applied directly or around livestock (including horses) residue studies might be needed | |
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2 Please follow section 1.16 of this guidance.

1 REFERENCES AND BACKGROUND DOCUMENTS

- 2 Billington, R., Lewis, W. R., Mehta, M. J., Dewhurst, I., et al. (2010). The mouse carcinogenicity
3 study is no longer a scientifically justifiable core data requirement for the safety assessment of
4 pesticides. *Critical Reviews in Toxicology*, 40(1), 35–49.
- 5 Boverhof DR, Ladics G, Luebke B, Botham J, Corsini E, Evans E, Germolec D, Holsapple M,
6 Loveless S, Lu H, van der Laan JW, White Jr KI and Yang Y (2014) Approaches and considerations
7 for the assessment of immunotoxicity for environmental chemicals: A workshop summary. *Regul*
8 *Toxicol Pharmacol* 68:96-107.
- 9 Boobis A.R., Cohen S.M., Dellarco V., McGregor D., Meek M.E., Vickers C., Willcocks D., Farland
10 W. (2007) IPCS Framework for Analyzing the Relevance of a Cancer Mode of Action for Humans
11 in IPCS Harmonization Project Document No. 4, Part 1, IPCS framework for analysing the
12 relevance of a cancer mode of action for humans and case-studies.
13 http://www.who.int/ipcs/methods/harmonization/areas/cancer_mode.pdf
14
- 15 Bronaugh, R. L., & Maibach, H. I. (1987). *In vitro* percutaneous absorption. In F. N. Marzulli, &
16 H. I. Maibach, *Dermatotoxicology* (pp. 121-34). Washington DC: Hemisphere Publishing.
- 17 Buettner [ed] (2017), *Springer Handbook of Odor*. First Edition, Springer International
18 Publishing Switzerland.
- 19 Collinge M, Thorn M, Peachee V and White K, Jr (2012) Developmental immunotoxicity (DIT)
20 testing of pharmaceuticals: Current practices, state of the science, knowledge gaps, and
21 recommendations. *J Immunotoxicol* 9, 210-230.
- 22 Costa LG, Giordano G, Guizzetti M, Vitalone A (2008), *Neurotoxicity of pesticides: a brief review*,
23 [Frontiers in Bioscience 13, 1240-1249, January 1, 2008].
24
- 25 De Jong W, Van Loveren H, (Eds) (2007) *Animal Models in Immunotoxicology*. *Methods Special*
26 *Issue* 41:1-142.
- 27 DeWitt JC, Peden-Adams MM, Keil DE and Dietert RR (2012a) Current status of developmental
28 immunotoxicity: early-life patterns and testing. *Toxicol pathol* 40:230-36.
- 29 DeWitt JC, Peden-Adams MM, Keil DE and Dietert RR (2012b) Developmental immunotoxicity
30 (DIT): assays for evaluating effects of exogenous agents on development on the immune
31 system. *Curr Protoc Toxicol* Chapter 18: Unit 18.15.
- 32 Dietert RR (2014) Developmental immunotoxicity, perinatal programming, and
33 noncommunicable disease: Focus on human studies. *Advances in Medicine* Vol 2014.
- 34 Dietert RR and DeWitt J (2010) Developmental immunotoxicity (DIT): the why, when, and how
35 of DIT testing. *Methods Mol Biol* 598:17-25.
- 36 Dietert RR and Holsapple MP (2007) methodologies for developmental immunotoxicity (DIT
37 testing. *Methods* 41:123-131.
- 38 ECETOC. (1992). *Evaluation of the neurotoxic potential of chemicals*. Monograph No. 18.
39 Brussels.
- 40 ECETOC. (1993). *Percutaneous Absorption*. Monograph 20. Brussels: ECETOC.
- 41 ECHA (European Chemicals Agency) and EFSA (European Food Safety Authority)
42 with the technical support of the Joint Research Centre (JRC) (2018), Andersson N, Arena M,
43 Auteri D, Barmaz S, Grignard E, Kienzler A, Lepper P, Lostia AM, Munn S, Parra Morte JM,

- 1 Pellizzato F, Tarazona J, Terron A and Van der Linden S. Guidance for the identification of
2 endocrine disruptors in the context of Regulations (EU) No 528/2012 and (EC) No 1107/2009.
3 EFSA Journal 2018;16(6):5311, 135 pp. <https://doi.org/10.2903/j.efsa.2018.5311> ECHA-18-G-
4 01-EN.
- 5 EFSA (European Food Safety Authority) (2017). Guidance on dermal absorption. EFSA Journal
6 2017;15(6):4873. DOI: <https://doi.org/10.2903/j.efsa.2017.4873>.
- 7 EFSA (European Food Safety Authority) (2010). Application of systematic review methodology
8 to food and feed safety assessments to support decision making. EFSA Journal 2010;8(6):1637,
9 90 pp. <https://doi.org/10.2903/j.efsa.2010.1637>
- 10 FAO. (2010). Manual on the Development and Use of FAO and WHO Specifications for Pesticides,
11 second revision.
- 12 Gupta RC (Ed) (2011) Reproductive and Developmental Toxicology. Elsevier Inc Academic Press,
13 Amsterdam, The Netherlands.
- 14 Hessel EV, Tonk ECM, Bos PM, van Loveren H and Piersma AH (2015) Developmental
15 immunotoxicity of chemicals in rodents and its possible regulatory impact. Crit Rev Toxicol
16 45:68-82.
- 17 Holsapple MP, Burns-Naas LA, Hastings KL, Ladics GS, Lavin AL, Makris SL, Yang Y and Luster
18 MI (2005) A proposed testing framework for developmental immunotoxicity (DIT). Toxicol Sci
19 83:18-24.
- 20 Howes, D., Guy, R. H., Hadgraft, J., Heylings, J., Hoeck, U., Kemper, F., et al. (1996). Method
21 for assessing percutaneous absorption – Report and Recommendations of ECVAM Workshop 13.
22 ATLA, 24, 81-106.
- 23 Janer, G., Slob, W., Hakkert, C. B., Vermeire, T., Piersman, H. A., __, et al. (2008). A
24 retrospective analysis of developmental toxicity studies in rat and rabbit: What is the added
25 value of the rabbit as an additional test species? Regulatory Toxicology and Pharmacology 50,
26 206-217.
- 27 Kerr JB, Loveland KL, O'Bryan MK, de Kretser DM (2006) Chapter 18 - Cytology of the Testis
28 and Intrinsic Control Mechanisms. In: Neill JD, Plant TM, Pfaff DW, Challis JRG, de Kretser DM,
29 Richards JS and Wassarman PM (eds) Knobil and Neill's Physiology of Reproduction, Third
30 Edition, pp. 827-947. Elsevier Inc Academic Press, Amsterdam, The Netherlands.
- 31 Kneuer C, Charistou A., Craig P, Eleftheriadou D, Engel N, Kjaerstad M, Krishnan S, Laskari V,
32 Machera K, Nikolopoulou D, Pieper C, Schoen E, Spilioti E and Buist H (2018). Applicability of in
33 silico tools for the prediction of dermal absorption for pesticides. EFSA supporting publication
34 2018:EN-1493. 156 pp. doi:10.2903/sp.efsa.2018.EN-1493
- 35 Lehman PA, Raney SG, Franz TJ (2011). Percutaneous absorption in man: in vitro-in vivo
36 correlation. 6 Skin Pharmacol. Physiol., 24:224-230.
- 37 Lotti M, Moretto A (2005). Organophosphate-induced delayed polyneuropathy. Toxicol Rev.
38 2005;24(1):37-49. doi: [10.2165/00139709-200524010-00003](https://doi.org/10.2165/00139709-200524010-00003)
- 39 McGee EA, Hsueh AJ (2000), Initial and cyclic recruitment of ovarian follicles. Endocr Rev
40 21:200-14.
- 41
- 42 NAFTA (2016). Developmental Neurotoxicity Study Guidance Document, North American Free
43 Trade Agreement (NAFTA) Technical Working Group on Pesticides (TWG).

- 1 OECD. (2000a). Guidance Document on Aquatic Toxicity Testing of Difficult Substances and
2 Mixtures. Series on Testing and Assessment, No 23.
- 3 OECD. (2004a). Guidance Document for the conduct of skin absorption studies, Series on Testing
4 and Assessment No 28. ENV/JM/MONO(2004)2.
- 5 OECD. (2004b). Guidance Document for Neurotoxicity Testing. Series on Testing and
6 Assessment No 20, Paris. ENV/JM/MONO(2004)25.
- 7 OECD. (2007). Guidance Document on the uterotrophic bioassay – procedure to test for
8 antioestrogenicity, Series on Testing and Assessment No 71, Paris. ENV/JM/MONO(2007)15.
- 9 OECD. (2008b). Guidance Documents on mammalian reproductive toxicity testing and
10 assessment. Series on Testing and Assessment No 43. ENV/JM/MONO(2008)16.
- 11 OECD. (2009). Guidance Document on the weanling Hershberger bioassay in rats: a short-term
12 screening assay for (anti)androgenic properties, Series on Testing and Assessment No 115.
13 ENV/JM/MONO(2009)41.
- 14 OECD. (2011). Guidance notes on dermal absorption, Series on Testing and Assessment No 156.
15 ENV/JM/MONO(2011)36
- 16 OECD (2018a), Guidance Document on Good In Vitro Method Practices (GIVIMP), OECD Series
17 on Testing and Assessment, No. 286, OECD Publishing, Paris.
- 18 OECD (2018b). Guidance Document on Inhalation Toxicity Studies, OECD Series on Testing and
19 Assessment, No.39, (Second Edition). ENV/JM/MONO(2009)28/REV1.
- 20 OECD. (2019a). Guidance on dermal absorption, Draft Second Edition, Series on Testing and
21 Assessment No 156.
- 22 OECD (2019b), Second Edition - Guidance Document on Integrated Approaches to Testing and
23 Assessment (IATA) for Serious Eye Damage and Eye Irritation, OECD Series on Testing and
24 Assessment, No. 263, OECD Publishing, Paris, [<https://doi.org/10.1787/84b83321-en>].
- 25 Ogungbesan A, Neal-Kluever A and Rice P (2019) Exploring the use of current immunological
26 assays for the developmental immunotoxicity assessment of food contact materials. Food Chem
27 Toxicol 133, 110801.
- 28 Rozman, K. K. (1986). Faecal excretion of toxic substances. In K. K. Rozman, & O. Hanninen,
29 Gastrointestinal Toxicology. Amsterdam: Elsevier.
- 30 Rooney AA, Yang Y and Makris SL (2009) Recent progress and diverge effects in developmental
31 immunotoxicology: overview of a symposium at the 46th Annual SOT Meeting, Charlotte, NC. J
32 Immunotoxicol 5:395-400. Erratum in J Immunotox 6:74.
- 33 Sachana M, Shafer TJ and Terron A (2021): Toward a Better Testing Paradigm for Developmental
34 Neurotoxicity: OECD Efforts and Regulatory Considerations. Biology 10, 86.UN. (2009).
35 Recommendations on the Transport of Dangerous Goods. Manual of Tests and Criteria.
36 ST/SG/AC.10/11/Rev.5. (UN-MTC). New York and Geneva.
- 37 US EPA. (1992). Dermal exposure assessment: Principles and Applications. EPA/600/8-91.001B.
38 Washington DC.Voorhees JR, Rohlman DS, Lein PJ, Pieper AA (2017), Neurotoxicity in Preclinical
39 Models of Occupational Exposure to Organophosphorus Compounds. Front Neurosci. 2016; 10:
40 590. doi: [10.3389/fnins.2016.00590](https://doi.org/10.3389/fnins.2016.00590)
41
- 42 WHO. (1986). WHO/IPCS Environmental Health Criteria (EHC) 60. Principles and Test Methods

- 1 for the Assessment of Neurotoxicity Associated with Exposure to Chemicals.
- 2 WHO. (1996). WHO/IPCS Environmental Health Criteria (EHC) 180, Principles and Methods for
3 Assessing Direct Immunotoxicity Associated with Exposure to Chemicals .
- 4 WHO. (1999). WHO/IPCS Environmental Health Criteria (EHC) 212, Principles and Methods for
5 Assessing Allergic Hypersensitization Associated with Exposure to Chemicals
- 6 WHO. (2007). WHO/IPCS Environmental Health Criteria (EHC) 236, Principles and Methods for
7 Assessing Autoimmunity Associated with Exposure to Chemicals.
- 8 WHO. (2012). Guidance for immunotoxicity risk assessment for chemicals, Harmonization
9 Project Document No. 10.
- 10 WHO/IPCS (2012) Guidance for Immunotoxicity Risk Assessment for Chemicals, IPCS
11 Harmonisation Project Document No 10. Available at:
12 <http://www.who.int/ipcs/methods/harmonization/areas/immunotoxicity/en/>