OECD GUIDELINE FOR THE TESTING OF CHEMICALS

Defined Approaches (DAs) for Serious Eye Damage and Eye Irritation for including a new DA for solids
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1. Introduction

1.1. General Introduction

1. The assessment of eye irritation/serious eye damage originally involved the use of albino rabbits according to the Draize eye test method (OECD Test Guideline 405) (1). The hazard potential of a test chemical was determined based on its effect on corneal opacity (CO), iritis (IR), conjunctival redness (CR), and conjunctival chemosis (CC). Based on the severity of effects and/or the timing of their reversibility, classifications are derived according to the serious eye damage/eye irritation classification criteria defined by the United Nations (UN) Globally Harmonized System of Classification and Labelling of Chemicals (GHS) (2). According to the UN GHS classification system, Category 1/serious damage (Cat. 1) is defined as causing irreversible effects (not fully reversible within 21 days) on the eye/serious damage to the eye. Category 2/irritation (Cat. 2) is defined as causing reversible effects (fully reversible within 21 days) on the eye/eye irritation. This category may be divided into the optional Categories 2A (effects fully reversible within 21 days) and 2B (effects fully reversible within 7 days). When none of the Cat. 1 or Cat. 2 classification criteria are met, the test chemical does not require classification which corresponds to No Category (No Cat.).

2. In 2022, a stand-alone in vitro method (OECD TG 492B was adopted for the identification of test chemicals not requiring classification (UN GHS No Cat), requiring classification for eye irritation (UN GHS Cat 2) and requiring classification for serious eye damage (UN GHS Cat 1) (3). Furthermore, several Test Guidelines (TGs) on in vitro methods have been adopted for the identification of test chemicals inducing serious eye damage (UN GHS Cat. 1) or for the identification of test chemicals not requiring classification for eye irritation and serious eye damage hazards (UN GHS No Cat.), notably OECD TG 437, TG 438, TG 460, TG 491, TG 492, TG 494, and TG 496 (4, 5, 6, 7, 8, 9, 10). Data generated with these in vitro methods are proposed to be used together, as well as with information sources such as physicochemical properties, in silico and read-across predictions from chemical analogues, within integrated approaches to testing and assessment (IATA) or defined approaches (DAs) (11). Results from the individual information sources cannot be used in DAs if the chemicals are known to clearly fall outside the applicability domains of the methods, as may be detailed in the respective assay TGs. The prediction from a DA may be used alone or along with further information as part of an IATA (11) or according to the applicable legal criteria.

3. The major difficulty for a single in vitro test method to fully replace the in vivo rabbit eye test (TG 405) is to predict the middle category (UN GHS Cat. 2) and it is therefore recommended to make use of testing strategies (e.g., Top-Down or Bottom-Up approach) that combine the strengths of individual in vitro test methods to address the required ranges of irritation potential (12). The determination of the most relevant in vivo endpoint(s), in particular the effects on cornea, iris or conjunctiva, is important for the development of adequate in vitro methods as it allows to better understand the relationship between the in vitro and the in vivo data (13, 14). For this reason, it is recommended to take into consideration the most important drivers for Cat. 1 and Cat. 2 classifications as well as the distribution of in vivo effects for chemicals not requiring classification when selecting
reference chemicals for the development, evaluation and/or validation of alternative methods and/or strategies for serious eye damage and eye irritation testing (11) (see “In vivo reference data (Draize eye test)” in the Supporting document to the Guideline (GL) on Defined Approaches (DAs) for Serious Eye Damage/Eye Irritation (15)).

4. Modes of action for eye irritation are unknowable for the majority of chemicals and do not provide additional insight in evaluation of the test methods and DAs, and thus they are not considered for the analysis of the test chemicals for the DAs in the current Guideline (see paragraph 27 of the SD for more information).

5. Results from multiple information sources can be used together in DAs to predict the eye hazard potential of test chemicals. A DA consists of a fixed data interpretation procedure (DIP) (i.e. a mathematical model, a rule-based approach) applied to data (e.g. in silico predictions, in chemico, in vitro data) generated with a defined set of information sources to derive a prediction without the need for expert judgment. The DAs use method combinations intended to overcome some of the limitations of the individual, stand-alone methods in order to provide increased confidence in the overall obtained result. The DAs provide information that can be used for eye hazard identification.

6. Testing laboratories should consider all relevant available information on the test chemical prior to conducting the studies according to a DA. Such information could include, for example, the identity and chemical structure of the test chemical and its physicochemical properties. Such information should be considered in order to determine whether the individual OECD TG methods under a specific DA are applicable for the test chemical.

7. When performing a hazard evaluation based on the output from the in vivo Draize eye test, from an in chemico test, from an in vitro test, from an in silico approach, from a DA, and any combination thereof, the same principles always apply, i.e. all available information relevant to the chemical in question should be taken into consideration as well as toxicological data on structurally related test chemicals, if available. However, specific regulatory requirements in the applicable legislation should be applied.

8. Two rule-based DAs for non-surfactant liquids and one rule-based DA for neat solids are included in this GL, and are described with respect to their intended regulatory purpose: hazard identification, i.e. discrimination between three UN GHS categories i.e., Category 1 (Cat. 1) on “serious eye damage”; Category 2 (Cat. 2) on “eye irritation” and No Category (No Cat.) for chemicals “not requiring classification and labelling” for eye irritation or serious eye damage (2). The evaluation and review of the DAs are described in detail in the Supporting Document for Evaluation and Review of TG 467 on DAs for Serious Eye Damage / Eye Irritation (15). For the non-surfactant liquids, a dataset of at least 86 chemicals with DA predictions, data on individual information sources, highly curated Draize eye test data, and physicochemical properties, was compiled and is attached as Annex B (spreadsheets) to the Supporting Document for Evaluation and Review of TG 467 on DAs for Serious Eye Damage / Eye Irritation (15). A list of 109 solids with DA predictions, data on individual information sources, and highly curated Draize eye test data was compiled and is attached as Annex A (spreadsheets) to the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (15). The list of chemicals was used to evaluate the performance of the DAs. The set of liquids and solids covers a broad range of uses and chemicals classes, with a wide range of organic functional groups (79 different OFGs for the liquids and 111 different OFG for the solids) defined according to OECD QSAR Toolbox analysis (version 3.2; https://www.oecd.org/chemicalsafety/risk-assessment/oecd-qsar-toolbox.htm).

9. The dataset is chemically diverse as shown by the physicochemical properties covered by these chemicals: it contains small and large molecules, as well as hydrophobic and hydrophilic substances. Further details on the chemical characterization of the
1.2. DAs and Use Scenarios included in the Guideline

11. The DAs currently described in this GL are:

- **Part I** - Defined Approaches 1 for Eye hazard identification based on physicochemical properties and *in vitro* data (DAL-1).

- **Part II** - Defined Approaches 2 for Eye hazard identification based on *in vitro* data (DAL-2).

- **Part III** - Defined Approach (DAS) for Eye hazard identification based on *in vitro* data for neat solids (DAS).

12. The DAs described in this GL can each be used to address countries' requirements for identifying chemicals causing serious eye damage (*i.e.* UN GHS Category 1), eye irritation (*i.e.* UN GHS Category 2), and test chemicals not requiring classification (*i.e.* UN GHS No Category), though they do so with different performance (detailed in the respective descriptions of each DA).

13. The DAs described in this GL are not designed to distinguish between Categories 2A and 2B.

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1 DIP with the BCOP OP-KIT test method did not meet the acceptance criteria for the current GL and thus is not used for DAL-1 or DAL-2 (see Annex A of the supporting document).

2 Other similar methods from OECD TG 492 were not used for the DAL-1 analysis due to insufficient availability in data for those methods.

3 DIP with the BCOP OP-KIT test method did not meet the acceptance criteria for the current GL and thus is not used for DAS.

4 DIP with the EpiOcular™ EIT test method did not meet the acceptance criteria for the current GL and thus is not used for DAS.
15. DAL-1 and DAL-2 are applicable to liquids (i.e., pipettable test substances) and DAS is applicable to solids (i.e., not pipettable test substances). For additional details see Section 2 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (15).

16. The performance of DAL-1 PCP/EpiOcular/LLBO described in this GL for discriminating between the three UN GHS categories was evaluated using 94 non-surfactant liquids (17 Cat. 1, 22 Cat. 2, and 55 No Cat.) for which physicochemical properties, EpiOcular™ EIT predictions, BCOP LLBO predictions (available for all in vivo classified results but missing for 14/55 in vivo No Cat. substances), and classifications based on Draize Eye test data are available (for additional details see Section 2 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (15)).

17. The performance of DAS PCP/SkinEthic/LLBO described in this GL for discriminating between the three UN GHS categories was evaluated using 86 non-surfactant liquids (17 Cat. 1, 23 Cat. 2, and 46 No Cat.) for which physicochemical properties, SkinEthic™ HCE EIT predictions, BCOP LLBO predictions (available for all in vivo classified results but missing for 11/46 in vivo No Cat. substances), and classifications based on Draize Eye test data are available (for additional details see Section 2 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (15)).

18. The performance of the DAL-2 STE/LLBO described in this GL for discriminating between the three UN GHS categories was evaluated using 164 non-surfactant liquids (17 Cat. 1, 24 Cat. 2, and 123 No Cat.) for which STE predictions, BCOP LLBO predictions (available for all in vivo classified results but missing for 67/123 in vivo No Cat. substances), and classifications based on Draize Eye test data are available (for additional details see Section 2 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (15)).

19. The performance of the DAS SkinEthic/LLBO described in this GL for discriminating between the three UN GHS categories was evaluated using 109 solids (31 Cat. 1, 18 Cat. 2, and 60 No Cat.) for which SkinEthic™ HCE EIT predictions, BCOP LLBO predictions (available for all in vivo classified results but missing for 5/60 in vivo No Cat. substances), and classifications based on Draize Eye test data are available (for additional details see Section 2 and Annex B.3 of the Supporting Document to the GL on DAs for Serious Eye Damage/Eye Irritation (15).

1.3. Performance and Applicability

1.3.1. Performance of the DAs

20. Table 1.1 provides an overview of the DAs included in this Guideline (GL), their information sources used, and summarises their performance against the Draize Eye reference data. More details are provided in Part I, Part II, and Part III of this GL, as well as in the Supporting documents to the GL on DAs for Serious Eye Damage and Eye Irritation (15).

21. The performance of the DAs for UN GHS classification (Cat. 1, Cat. 2, and No Cat) when compared to the Draize Eye test reference data yielded balanced accuracies of 69.2% (DAL-1 PCP/EpiOcular/LLBO), 75.2% (DAL-1 PCP/SkinEthic/LLBO), 74.3% (DAL-2 STE/LLBO), and 66.7% (DAS SkinEthic/LLBO). Note that there is a raised degree of uncertainty relating to the derived Cat. 1 and Cat. 2 accuracies (correct predictions), as compared with the No Cat. accuracies due to the lower number of reference chemicals within these categories. It was however not possible to increase the number of chemicals because of the limited number of available Draize Eye test results with a Cat. 1 or Cat. 2
classification. Detailed performance statistics are reported in Part I, Part II, and Part III and in Section 5 of the Supporting documents to the GL on DAs for Serious Eye Damage and Eye Irritation (15).

Table 1.1 Summary of the DAs included in this Guideline – Eye hazard identification

<table>
<thead>
<tr>
<th>DA</th>
<th>DAL-1 PCP/EpiOcular/LLBO (N=94)</th>
<th>DAL-1 PCP/SkinEthic/LLBO (N=86)</th>
<th>DAL-2 STE/LLBO (N=164)</th>
<th>DAS SkinEthic/LLBO (N=109)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Information Sources</td>
<td>Physicochemical properties, EpiOcular™ EITL (TG 492), BCOP LLBO (TG 437)</td>
<td>Physicochemical properties, SkinEthic™ HCE EITL (TG 492), BCOP LLBO (TG 437)</td>
<td>STE (TG 491), BCOP LLBO (TG 437)</td>
<td>SkinEthic™ HCE EITS (TG 492), BCOP LLBO (TG 437)</td>
</tr>
<tr>
<td>Applicable</td>
<td>Non-surfactant neat liquids</td>
<td>Non-surfactant neat liquids, liquids and solids dissolved in water</td>
<td>Non-surfactant neat solids</td>
<td></td>
</tr>
<tr>
<td>Performance vs. Draize Eye test (Correct classification)</td>
<td>70.5% No Cat. (N=55) 59.1% Cat. 2 (N=22) 70.5% Cat. 1 (N=17)</td>
<td>79.7% No Cat. (N=46) 68.7% Cat. 2 (N=23) 76.5% Cat. 1 (N=17)</td>
<td>85.3% No Cat. (N=123) 56.3% Cat. 2 (N=24) 81.2% Cat. 1 (N=17)</td>
<td>70.0% No Cat. (N=60) 52.3% Cat. 2 (N=18) 77.4% Cat. 1 (N=31)</td>
</tr>
</tbody>
</table>

Note: For performance, accuracy reflects correct classification rate within each UN GHS category.

EITL: Eye Irritation Test protocol for liquids and EITS protocol for solids
Solid: non-pipettable neat substance

1.3.2. Applicability domain of the DAs and of the individual components of the DAs

22. DAL-1 PCP/EpiOcular/LLBO and DAL-1 PCP/SkinEthic/LLBO are not applicable for surfactants and solids. Both DAs are applicable to neat liquids, excluding mixtures, UVCBs and multi-constituent substances. For impurities with concentration > 5% and < 20%, the physicochemical properties of the impurities also need to be determined, and only when all components meet the exclusion criteria, the liquid is predicted No Cat., in all other cases, proceed with an RhCE test method.

23. DAL-2 STE/LLBO is not applicable for surfactants and solids dispersed in water. The DAL-2 STE/LLBO is applicable to non-surfactant neat liquids, liquids and solids dissolved in water.

24. DAS SkinEthic/LLBO is not applicable to liquids. The DAS SkinEthic/LLBO is applicable to non-surfactant neat solids (i.e., not pipettable test substances).

25. Users should refer to the limitations of the individual in vitro test methods as specified in their respective TGs, which are revised as new data become available and should be consulted regularly. The most up-to-date published version of the respective TGs should always be used. Users should also refer to the limitations of the individual methods for measuring the physicochemical properties as specified in their respective GLs.

1.3.3. Uncertainty of DAs

26. Details on accepting the results of individual information sources to determine confidence in DA predictions are provided in Sections 2.1.4, 3.1.4, and 4.1.4 and in the respective TGs (TG 437; TG 491; TG 492) (4, 7, 8).
1.4. References


2. PART I - Defined Approaches 1 (DAL-1), based on physicochemical properties and in vitro data, for neat non-surfactant liquids

27. Part I of this GL applies to DAL-1 that is intended for hazard identification, i.e. distinguishing between serious eye damage and eye irritation potential of test chemicals (or the absence thereof), specifically for neat non-surfactant liquids based on physicochemical properties and in vitro data. A summary of the DAL-1 for hazard identification is provided below; additional detailed information can be found in the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation.

2.1. DAL-1

2.1.1. Summary

28. The DAL-1 is intended for the identification of the eye irritation hazard of a test chemical without the use of animal testing, i.e. UN GHS Cat. 1 vs. UN GHS Cat. 2 vs. UN GHS No Cat. The data interpretation procedure (DIP) is not designed to provide information on sub-categorisation of Cat. 2 into 2A and 2B.

29. The DAL-1 presented in this GL describes the combination of one and/or three physicochemical properties with the results of two in vitro test methods (RhCE and BCOP LLBO) for the identification of the eye hazard potential of non-surfactant liquid substances primarily for the purposes of classification and labelling without the use of animal testing (1). The physicochemical properties can be retrieved from publicly available databases, can be determined by new experimental studies, or may be predicted using computational methods (e.g. Quantitative Structure-Activity Relationships ((Q)SAR)). The RhCE models that are part of DAL-1 are the EpiOcular™ EITL and the SkinEthic™ HCE EITL (OECD TG 492) (2). Furthermore, the Bovine Corneal Opacity and Permeability (BCOP) test method with the laser light-based opacimeter (LLBO) is used (OECD TG 437) (3).

30. The DAL-1 PCP/EpiOcular/LLBO was compared to 94 chemicals with curated Draize Eye test reference data and demonstrated a balanced accuracy of 68.7% (see Table 2.1). The DAL-1 PCP/SkinEthic/LLBO was compared to 86 chemicals with curated Draize Eye test reference data and demonstrated a balanced accuracy of 75.0% (see Table 2.2).

2.1.2. Data interpretation procedure

31. The data interpretation procedure (DIP) applied uses the readout of the prediction models of each of the individual test method as defined by the TGs and/or information on the physicochemical properties. A scheme of DAL-1 is presented in Figure 2.

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5 Surfactant, also called surface-active agent, this is a substance, such as a detergent, that can reduce the surface tension of a water and thus allow it to foam or penetrate solids; it is also known as a wetting agent.
property exclusion rules based on water solubility (WS) or a combination of octanol-water partition coefficient (LogP), vapour pressure (VP) and surface tension (ST) of the neat liquid are used in a first step to identify liquid chemicals with no serious eye damage or eye irritation potential (details are provided in section 5.1.2. of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation). Liquids that are not identified as No Cat. according to the physicochemical property-based exclusion rules, are then evaluated based on a RhCE test method (EpiOcular™ EIT or SkinEthic™ HCE EIT) in Step 2. Liquids that result in a tissue viability > 60% are classified No Cat. Liquids that result in a tissue viability ≤ 60% are then evaluated based on the BCOP LLBO test method in a third step. Liquids that result in an opacity > 145 are predicted Cat. 1 and the remaining liquids are assigned Cat. 2. Note that it is also possible to start with a RhCE method, followed by the physicochemical property exclusion rules in case the tissue viability measured with EpiOcular™ EIT or SkinEthic™ HCE EIT > 60% (Figure 2.2). Furthermore, when a RhCE method is used as a first step and if the tissue viability > 60%, the prediction is based on the stand-alone method.
Figure 2.1. Scheme of the DAL-1 option 1; step 1 physicochemical exclusion rules (WS: water solubility in mg/mL; or LogP: octanol-water partition coefficient / VP: vapour pressure in mm Hg / ST: surface tension of the neat liquid in dyne/cm) to identify No Cat., step 2 RhCE EITL test method used to identify No Cat., and step 3 BCOP LLBO used to identify Cat. 1.
Figure 2.2. Scheme of the DAL-1 option 2: step 1 RhCE EITL test method used to identify No Cat., step 2: physicochemical exclusion rules (WS: water solubility in mg/mL; or LogP: octanol-water partition coefficient / VP: vapour pressure in mm Hg / ST: surface tension of the neat liquid in dyne/cm) to identify No Cat., and step 3 BCOP LLBO used to identify Cat. 1.
2.1.3. Description and limitations of the individual information sources

32. The individual information sources in the DA are the physicochemical properties and test methods included in OECD TG for serious eye damage/eye irritation or the absence thereof (OECD TG 437, 492) (2, 3), and the protocols are detailed therein.

33. The following in vitro test methods from those TGs have been characterised and included in the DAL-1.

- The RhCE EITL test methods: the methods measure the ability to induce cytotoxicity. In case borderline results are obtained, additional testing should be conducted, as specified in OECD TG 492 (2).

- BCOP LLBO test method: the eye hazard potential of a test chemical is measured by its ability to induce opacity and permeability in an isolated bovine cornea. Note that only opacity measurement is considered in the DAs. In case borderline results are obtained for opacity measurements, additional testing should be conducted, as specified in OECD TG 437 (3).

34. Any restriction regarding the applicability domain identified in the respective test method TGs (TG 437, TG 492) and analytical methods for measuring the physicochemical properties (GL 104, GL 105, GL 107, GL 115, GL 117, GL 123) is applicable to this GL (2, 3).

35. Measurements of physicochemical properties should be performed according to the OECD Guidelines (GL) and test reports are required corresponding with the information requested on data and reporting in each specific OECD GL (see Annex E). Prediction of physicochemical properties should use models that are based on the 5 OECD principles for QSAR models and that have a QMRF (QSAR Model Reporting Format).

2.1.4. Procedure for dealing with borderline result in test guidelines relevant to DAL-1

36. The first decision on whether each information element can be used is dictated by the applicability domain as described in the TGs of the respective in vitro methods (TG 437, TG 492) (2, 3). Even for within-domain substances, test results are inherently subject to variation and these variations increase the uncertainty of a test result, especially when close to a (classification) cut-off threshold, i.e. in the borderline range. The following procedures are in place to control the degree of uncertainty are described within the TGs of the respective information sources.

- TG 492 (RhCE EITL): A single test composed of at least two tissue replicates should be sufficient for a test chemical when the result is unequivocal. However, in cases of borderline results, such as non-concordant replicate measurements and/or mean percent tissue viability equal to 60±5% a second test should be considered, as well as a third one in case of discordant results between the first two tests.

- TG 437 (BCOP LLBO): UN GHS Cat. 1 prediction based on opacity (Lux/7, mean opacity > 145), but 1 of 3 corneas with opacity (Lux/7) < 130; in cases of borderline results in the first testing run, a second testing run should be considered, as well as a third one in case of discordant predictions between the first two testing runs.
2.1.5. Predictive capacity of the DAL-1 PCP/EpiOcular/LLBO vs. the Draize Eye test

37. The predictive capacity of DAL-1 PCP/EpiOcular/LLBO is reported based on data generated by the Draize eye test (see Table 2.1) (see Section 2.1 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation). Performance statistics are reported for weighted predictions as compared to Draize eye test reference data. DA predictions for specific chemicals and further details are available in Section 5 and Annex B.2 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation for liquids (5).

Table 2.1. Performance of DAL-1 PCP/EpiOcular/LLBO in comparison to Draize Eye reference data

<table>
<thead>
<tr>
<th>UN GHS</th>
<th>Prediction using DAL-1 PCP/EpiOcular/LLBOa</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cat 1</td>
<td>Cat 2</td>
<td>No Cat</td>
</tr>
<tr>
<td>Cat. 1 (N=17)' % (n/N)</td>
<td>76.5% (13.0/17.0)</td>
<td>23.5% (4.0/17.0)</td>
<td>0.0% (0.0/17.0)</td>
</tr>
<tr>
<td>Cat. 2 (N=22)' % (n/N)</td>
<td>27.3% (6.0/22.0)</td>
<td>59.1% (13.0/22.0)</td>
<td>13.6% (3.0/22.0)</td>
</tr>
<tr>
<td>No Cat. (N=55)' % (n/N)</td>
<td>5.5% (3.0/55.0)</td>
<td>24.0% (13.2/55.0)</td>
<td>70.5% (38.8/55.0)</td>
</tr>
</tbody>
</table>

68.7% balanced accuracy overall

a The proportion given is based on a weighted calculation which takes into account (where they exist) multiple results from an individual information source for a given chemical, and applying a correction factor so that all chemicals have a weight of 1. To improve the readability of the numbers in the table, the numbers n/N have been rounded, so they may deviate slightly from the percentage corresponding to the weighted calculation.

b EpiOcular™ EITL protocol for liquids.

Note 1: The performance is the same for the two versions of the DIP (Fig. 2.1 and Fig 2.2).

2.1.6. Predictive capacity of the DAL-1 PCP/SkinEthic/LLBO vs. the Draize Eye test

38. The predictive capacity of DAL-1 PCP/SkinEthic/LLBO is reported based on data generated by the Draize eye test (see Table 2.2) (see Section 2.1 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation). Performance statistics are reported for weighted predictions as compared to Draize eye test reference data. DA predictions for specific chemicals and further details are available in Section 5 and Annex B.2 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation for liquids (5).
Table 2.2. Performance of DAL-1 PCP/SkinEthic/LLBO in comparison to Draize Eye reference data

<table>
<thead>
<tr>
<th>UN GHS</th>
<th>Prediction using DAL-1 PCP/SkinEthic/LLBO (^b)</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cat 1</td>
<td>Cat 2</td>
<td>No Cat</td>
</tr>
<tr>
<td>Cat. 1 (N=17), (^a)</td>
<td>76.5%(13.0/17.0)</td>
<td>23.5%(4.0/17.0)</td>
<td>0.0%(0.0/17.0)</td>
</tr>
<tr>
<td>Cat. 2 (N=23), (^a)</td>
<td>30.4%(7.0/23.0)</td>
<td>68.7%(15.8/23.0)</td>
<td>0.9%(0.2/23.0)</td>
</tr>
<tr>
<td>No Cat. (N=46), (^a)</td>
<td>3.1%(1.4/46.0)</td>
<td>17.2%(7.9/46.0)</td>
<td>79.7%(36.7/46.0)</td>
</tr>
</tbody>
</table>

75.0% balanced accuracy overall

\(^a\) The proportion given is based on a weighted calculation which takes into account (where they exist) multiple results from an individual information source for a given chemical, and applying a correction factor so that all chemicals have a weight of 1. To improve the readability of the numbers in the table, the numbers n/N have been rounded, so they may deviate slightly from the percentage corresponding to the weighted calculation.

\(^b\) SkinEthic™ HCE EITL protocol for liquids

Note 1: The performance is the same for the two versions of the DIP (Fig. 2.1 and Fig 2.2).

### 2.1.7. Demonstration of Proficiency

39. The DAL-1 relies on a simple, rule-based data interpretation procedure and requires no expert judgment. Proficiency chemicals for the individual information sources are defined in the respective TGs (2, 3). Proficiency for the individual information sources demonstrates proficiency for the DA.

### 2.1.8. Reporting of the DA

40. The reporting of the DA application should include at a minimum the following elements:

- Test chemical identification (e.g., chemical name, structural formula, composition, isomers, purity, chemical identity of impurities including their quantities as available, CAS number, batch and lot number, and other relevant identifiers).
- The DAL-1 option used, and the RhCE method used.
- Individual test reports performed per corresponding TGs (OECD TG 437, TG 492). Note that the chemical identity for each test report should match that above.
- Individual test reports on physicochemical properties corresponding with the information requested on data and reporting in each specific OECD GL (Annex E).
- Discussions on any uncertainties in the data with the in vitro methods and physicochemical properties applied in the DA that was used.
- Outcome of the DA application, including discussion of any uncertainties in the applied DA, as well as their predicted impact (e.g., over- or under-classification).
- Any deviation from or adaptation of the DA.
- Conclusion
2.2. References


3. PART II – Defined Approaches 2 (DAL-2), based on *in vitro* data, for non-surfactant neat liquids, liquids and solids dissolved in water

Part II of this GL applies to DAL-2 STE/LLBO that is intended for hazard identification, *i.e.* distinguishing between serious eye damage and eye irritation potential of test chemicals (or the absence thereof), specifically for non-surfactant neat liquids, liquids and solids dissolved in water based on *in vitro* data. A summary of the DAL-2 for hazard identification is provided below; additional detailed information can be found in the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation.

### 3.1. DAL-2

#### 3.1.1. Summary

The DAL-2 STE/LLBO is intended for the identification of the eye irritation hazard of a test chemical without the use of animal testing, *i.e.* UN GHS Cat. 1 vs. UN GHS Cat. 2 vs. UN GHS No Cat. The data interpretation procedure (DIP) is not designed to provide information on sub-categorisation of Cat. 2 into 2A and 2B.

The DAL-2 STE/LLBO presented in this GL describes the combination of two *in vitro* test methods (STE: OECD TG 491 and BCOP LLBO: OECD TG 437) for the identification of the eye hazard potential of non-surfactant neat liquids, liquids and solids dissolved in water primarily for the purposes of classification and labelling without the use of animal testing (1, 2, 3).

The DAL-2 STE/LLBO was compared to 164 chemicals with curated Draize Eye test reference data and demonstrated a balanced accuracy of 74.3% (see Table 3.1).

#### 3.1.2. Data interpretation procedure

The DIP applied uses the readout of the prediction models of each of the individual test methods as defined by the TGs (OECD 437, OECD 491) (1, 2). A scheme of DAL-2 STE/LLBO is presented in Figure 3.1. The STE test method is used to identify liquid chemicals with no serious eye damage or eye irritation potential (No Cat.: liquids that result in a mean cell viability > 70% at a 5% and 0.05% concentration) or to identify liquids that cause serious eye damage/eye irritation (Cat. 1: liquids that result in a mean cell viability ≤ 70% at a 5% and 0.05% concentration). For liquids that result in a mean cell viability ≤ 70% at 5% concentration but > 70% at 0.05%, the BCOP LLBO is needed. Liquids that result in an opacity > 145 are predicted as Cat. 1 and the remaining liquids are assigned to Cat. 2. Note that it is also possible to start with the BCOP LLBO followed by the STE test method, this scheme of DAL-2 is presented in Figure 3.2.

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6 Surfactant, also called surface-active agent, is a substance, such as a detergent, that can reduce the surface tension of a water and thus allow it to foam or penetrate solids; it is also known as a wetting agent.
Figure 3.1. Scheme of the DAL-2 STE/LLBO option 1: start with the STE test method followed by the BCOP LLBO test method.
Figure 3.2. Scheme of the DAL-2 STE/LLBO option 2: start with the BCOP LLBO test method followed by the STE test method.
3.1.3. Description and limitations of the individual information sources

46. The individual information sources in the DA are test methods included in OECD TGs (OECD TG 437, 491) for serious eye damage/eye irritation or the absence thereof (1, 2), and the protocols are detailed therein.

47. The following test methods from those TGs have been characterised and included in the DAL-2 STE/LLBO.

- BCOP LLBO test method: the eye hazard potential of a test chemical is measured by its ability to induce opacity and permeability in an isolated bovine cornea. Note that only opacity measurement is considered in the DAL-2. In case borderline results are obtained for opacity measurements, additional testing should be conducted, as specified in OECD TG 437 (1).

- STE test method: the eye hazard potential of a test chemical is assessed based on its ability to induce cytotoxicity on a confluent monolayer of Statens Seruminstitut Rabbit Cornea (SIRC) cells.

48. Any restrictions regarding the applicability domain identified in the respective TGs (TG 437, TG 491) are applicable to this GL (1, 2).

3.1.4. Procedures for dealing with borderline results in the test guidelines relevant to DAL-2 STE/LLBO predictions

49. The first decision on whether each information element can be used is dictated by the practical limitations as described in the TGs of the respective in vitro methods (TG 437, TG 491) (1, 2). Even for within-domain substances, test results are inherently subject to variation and these variations increase the uncertainty of a test result, especially when close to a (classification) cut-off threshold, i.e. in the borderline range. The following procedures to control the degree of uncertainty are described with the TG of the respective information sources.

- TG 437 (BCOP LLBO): UN GHS Cat. 1 prediction based on opacity (Lux/7, mean opacity > 145), but 1 of 3 corneas with opacity (Lux/7) < 130; in cases of borderline results in the first testing run, a second testing run should be considered, as well as a third one in case of discordant predictions between the first two testing runs.

- TG 491 (STE): Standard deviation of the final cell viability derived from three independent repetitions should be less than 15% for both 5% and 0.05% concentrations of the test chemical. If the standard deviation is greater than or equal to 15%, the results should not be used and three more repetitions should be performed.

3.1.5. Predictive capacity of the DAL-2 STE/LLBO vs. the Draize Eye test

50. The predictive capacity of DAL-2 STE/LLBO is reported based on data generated by the Draize eye test (see Table 3.1) (see Section 2.1 and Annex B.3 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation). Performance statistics are reported for weighted predictions as compared to Draize eye test reference data. DA predictions for specific chemicals and further details are available in Section 5 and Annex B.2 of the Supporting document to the GL on DAs for Serious Eye Damage/Eye Irritation.
Table 3.1. Performance of DAL-2 STE/LLBO in comparison to Draize Eye reference data

<table>
<thead>
<tr>
<th>UN GHS</th>
<th>Prediction using DAL-2 STE/LLBO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cat 1</td>
</tr>
<tr>
<td>Cat 1 (N=17), % (n/N)</td>
<td>81.2%</td>
</tr>
<tr>
<td></td>
<td>(13.8/17.0)</td>
</tr>
<tr>
<td>Cat 2 (N=24), % (n/N)</td>
<td>30.2%</td>
</tr>
<tr>
<td></td>
<td>(7.2/24.0)</td>
</tr>
<tr>
<td>No Cat. (N=123), % (n/N)</td>
<td>4.1%</td>
</tr>
<tr>
<td></td>
<td>(5.1/123.0)</td>
</tr>
</tbody>
</table>

74.3% balanced accuracy overall

*a The proportion given is based on a weighted calculation which takes into account (where they exist) multiple results from an individual information source for a given chemical, and applying a correction factor so that all chemicals have a weight of 1. To improve the readability of the numbers in the table, the numbers n/N have been rounded, so they may deviate slightly from the percentage corresponding to the weighted calculation.

Note 1: The performance was obtained using the version of the DIP provided in Fig 3.1.

### 3.1.6. Proficiency chemicals

51. The DAL-2 STE/LLBO relies on a simple, rule-based data interpretation procedure and requires no expert judgment. Proficiency chemicals for the individual information sources are defined in the respective TGs (1, 2). Proficiency for the individual information sources demonstrates proficiency for the DAL-2 STE/LLBO.

### 3.1.7. Reporting of the DA

52. The reporting of the DA application should include at a minimum the following elements:

- Test chemical identification (e.g., chemical name, structural formula, composition, isomers, purity, chemical identity of impurities including their quantities as available, CAS number, batch and lot number, and other relevant identifiers).

- Describe the DAL-2 option used.

- Individual test reports performed per corresponding TGs (OECD TG 437, TG 491). Note that the chemical identity for each test report should match that above.

- Discussions on any uncertainties in the data with the *in vitro* methods applied in the DA that was used.

- Outcome of the DA application, including discussion of any uncertainties in the applied DA, as well as their predicted impact (e.g., over- or under-classification)

- Any deviation from or adaptation of the DA.

- Conclusion.

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3.2. References


4. PART III – Defined Approaches (DAS), based on in vitro data, for neat solids

54. Part III of this GL applies to DAS SkinEthic/LLBO that is intended for hazard identification, i.e., distinguishing between serious eye damage and eye irritation potential of test chemicals (or the absence thereof), specifically for neat solids based on in vitro data. A summary of the DAS for hazard identification is provided below; additional detailed information can be found in the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation (3).

4.1. DAS SkinEthic/LLBO

4.1.1. Summary

55. The DAS SkinEthic/LLBO is intended for the identification of the eye irritation hazard of a test chemical without the use of animal testing, i.e., UN GHS Cat. 1 vs. UN GHS Cat. 2 vs. UN GHS No Cat. The data interpretation procedure (DIP) is not designed to provide information on sub-categorisation of Cat. 2 into 2A and 2B.

56. The DAS SkinEthic/LLBO presented in this GL describes the combination of two in vitro test methods (SkinEthic™ HCE EITS: OECD TG 492 and BCOP LLBO: OECD TG 437) for the identification of the eye hazard potential of neat solids primarily for the purposes of classification and labelling without the use of animal testing (1, 2).

57. The DAS SkinEthic/LLBO was compared to 109 chemicals with curated Draize Eye test reference data and demonstrated a balanced accuracy of 66.7% (see Table 4.1).

4.1.2. Data interpretation procedure

58. The DIP uses the readout of the prediction models of each of the individual test methods as defined by the TGs (OECD 437, OECD 492) (1, 2). A scheme of DAS SkinEthic/LLBO is presented in Figure 4.1. The SkinEthic™ HCE EITS test method is used to identify solid chemicals with no serious eye damage or eye irritation potential (No Cat.: solids that result in a mean tissue viability > 50%). For solids that result in a mean tissue viability ≤ 50%, the BCOP LLBO is needed. Solids that result in an opacity > 145 or OD > 2.5, or both opacity > 145 and OD > 2.5 are predicted as Cat. 1 and the remaining solids are assigned to Cat. 2.

7 A solid is a non-pipettable test substance.
Figure 4.1. Scheme of the DAS SkinEthic/LLBO: start with the SkinEthic™ HCE EITS test method followed by the BCOP LLBO test.
4.1.3. Description and limitations of the individual information sources

59. The individual information sources in the DA are test methods included in OECD TGs (OECD TG 492, 437) for serious eye damage/eye irritation or the absence thereof (1, 2), and the protocols are detailed therein.

60. The following test methods from those TGs have been characterised and included in the DAS SkinEthic/LLBO.

   • SkinEthic™ HCE EITS test method: the method measures the ability to induce cytotoxicity. In case borderline results are obtained, additional testing should be conducted, as specified in OECD TG 492 (1).
   
   • BCOP LLBO test method: the eye hazard potential of a test chemical is measured by its ability to induce opacity and permeability in an isolated bovine cornea. In case borderline results are obtained for opacity or permeability measurements, additional testing should be conducted, as specified in OECD TG 437 (2).

61. Any restrictions regarding the applicability domain identified in the respective TGs (TG 437, TG 492) are applicable to this GL (1, 2).

4.1.4. Procedures for dealing with borderline results in the test guidelines relevant to DAS SkinEthic/LLBO predictions

62. The first decision on whether each information element can be used is dictated by the practical limitations as described in the TGs of the respective in vitro methods (TG 437, TG 492) (1, 2). Even for within-domain substances, test results are inherently subject to variation and these variations increase the uncertainty of a test result, especially when close to a (classification) cut-off threshold, i.e., in the borderline range. The following procedures to control the degree of uncertainty are described within the respective TG.

   • TG 492 (SkinEthic™ HCE EITS): A single test composed of at least two tissue replicates should be sufficient for a test chemical when the result is unequivocal. However, in cases of borderline results, such as non-concordant replicate measurements and/or mean percent tissue viability equal to 50±5% a second test should be considered, as well as a third one in case of discordant results between the first two tests.
   
   • TG 437 (BCOP LLBO): UN GHS Cat. 1 prediction based on (i) opacity (Lux/7, mean opacity > 145), but 1 of 3 corneas with opacity (Lux/7) < 130 or (ii) OD (mean OD > 2.5), but 1 of 3 corneas with OD < 2.0; in cases of borderline results in the first testing run, a second testing run should be considered, as well as a third one in case of discordant predictions between the first two testing runs.

4.1.5. Predictive capacity of the DAS SkinEthic/LLBO vs. the Draize Eye test

63. The predictive capacity of DAS SkinEthic/LLBO is reported based on data generated by the Draize eye test (see Table 4.1) (see Section 5.1 and Annex A.2 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation for neat solids (3)). Performance statistics are reported for weighted predictions as compared to Draize eye test reference data. DA predictions for specific chemicals and further details are available in Section 7 and Annex A.2 of the Supporting document to the GL on DAs for Serious Eye Damage and Eye Irritation for neat solids (3).
Table 4.1. Performance of DAS SkinEthic/LLBO in comparison to Draize Eye reference data

<table>
<thead>
<tr>
<th>UN GHS</th>
<th>Prediction using DAS SkinEthic/LLBO</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cat 1</td>
</tr>
<tr>
<td>Cat. 1 (N=31), % (n/N)</td>
<td>77.4%</td>
</tr>
<tr>
<td></td>
<td>(24.0/31.0)</td>
</tr>
<tr>
<td>Cat. 2 (N=18), % (n/N)</td>
<td>29.5%</td>
</tr>
<tr>
<td></td>
<td>(5.3/18.0)</td>
</tr>
<tr>
<td>No Cat. (N=60), % (n/N)</td>
<td>1.7%</td>
</tr>
<tr>
<td></td>
<td>(1.0/60.0)</td>
</tr>
</tbody>
</table>

66.7% balanced accuracy overall

a The proportion given is based on a weighted calculation which takes into account (where they exist) multiple results from an individual information source for a given chemical, and applying a correction factor so that all chemicals have a weight of 1. To improve the readability of the numbers in the table, the numbers n/N have been rounded, so they may deviate slightly from the percentage corresponding to the weighted calculation.

4.1.6. Proficiency chemicals

64. The DAS SkinEthic/LLBO relies on a simple, rule-based data interpretation procedure and requires no expert judgment. Proficiency chemicals for the individual information sources are defined in the respective TGs (1, 2). Proficiency for the individual information sources demonstrates proficiency for the DAS.

4.1.7. Reporting of the DA

65. The reporting of the DA application should include at a minimum the following elements:

- Test chemical identification (e.g., chemical name, structural formula, composition, isomers, purity, chemical identity of impurities including their quantities as available, CAS number, batch and lot number, and other relevant identifiers).
- Individual test reports performed per corresponding TGs (OECD TG 437, TG 492). Note that the chemical identity for each test report should match that above.
- Discussions on any uncertainties in the data with the in vitro methods applied in the DA that was used.
- Outcome of the DA application, including discussion of any uncertainties in the applied DA, as well as their predicted impact (e.g., over- or under-classification).
- Any deviation from or adaptation of the DA.
- Conclusion.
4.2. References

