

Research Article

# Integrated Skin Sensitization Assessment Based on OECD Methods (II): Hazard and Potency by Combining Kinetic Peptide Reactivity and the “2 out of 3” Defined Approach

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## Abstract

Depending on regulatory requirements, three questions on the skin sensitization risk for new chemicals with potential skin contact by consumers must be answered by experimental testing: (i) Binary hazard assessment to identify sensitizers, (ii) subclassification of sensitizers according to the Global Harmonized System (GHS), and (iii) a Point of Departure (PoD) for risk assessment. The Organisation for Economic Co-operation and Development (OECD) published a test guideline incorporating the “2 out of 3” Defined Approach (2o3 DA) for skin sensitization hazard assessment. At the same time, the kinetic direct peptide reactivity assay (kDPRA) was added to the test guidelines as a stand-alone method for GHS subclassification. For the 2o3 DA, at least two *in vitro* tests are conducted. The cell-based tests and the kDPRA generate, next to a binary outcome with a fixed threshold, continuous dose-response data, which can be used in quantitative regression models to derive a PoD. The 2o3 DA is flexible for the sequence of testing. Here we show different testing sequences and how they can be combined with kDPRA data to provide a PoD in parallel to hazard identification (hazard ID) and GHS subclassification. Using these different testing sequences, a set of 188 chemicals with available *in vitro* data was evaluated for the final PoD. The results indicate that testing can start with DPRA / kDPRA and either of the cell-based assays, and that testing can stop with two congruent tests without major impact on the final PoD for chemicals within the applicability domain of the kDPRA.

## 1 Introduction

A significant effort underway is developing next-generation risk assessment approaches for skin sensitization that don't rely on the need for new animal test data. New Approach Methodologies (NAMs), non-animal test methods, have been developed to identify skin sensitization hazards and with a new focus on determining potency information for risk assessment purposes (Bernauer et al., 2021; Dent et al., 2018; Ezendam et al., 2016; Gilmour et al., 2020; Kleinstreuer et al., 2018). The ban on animal testing for new cosmetic ingredients was implemented in Europe within the Cosmetic legislation (Regulation (EC) No 1223/2009) in 2009, which led to the rapid development of NAMs by both academic and industrial laboratories (Ezendam et al., 2016). Three OECD guidelines have been published that cover mechanistic key events (covalent binding to protein, keratinocyte activation, and dendritic cell activation). These three mechanistic events map to key events 1-3 of the skin sensitization AOP (OECD, 2014). There are eight non-animal test methods approved in OECD TG (Direct Peptide Reactivity Assay, DPRA; Amino Acid Derivative Reactivity Assay, ADRA; kDPRA; ARE-Nrf2 luciferase assay KeratinoSens<sup>TM</sup>, KS; ARE-Nrf2 luciferase assay LuSens; human Cell Line Activation Test, h-CLAT; U937 cell line activation test, U-SENS<sup>TM</sup>; and Interleukin-8 Reporter Gene Assay, IL-8 Luc Assay)(OECD, 2018a,b, 2021b).

Recent work has been focused on finding ways to combine NAM data to generate integrated approaches to testing and assessment (IATA) or defined approaches (DA). Defined Approaches for skin sensitization contain fixed data interpretation procedures on how to combine data obtained from different *in chemico*, *in vitro* and *in silico* methods to conclude whether a substance is a skin sensitizer and, if so, what is the skin sensitization potency (Gilmour et al., 2020; Hoffmann et al., 2018; Kleinstreuer et al., 2018). Two simple DAs for assessing skin sensitization have been published in a new guideline (OECD, 2021a). The OECD Guideline 497 includes the 2o3 DA and integrated testing strategy (ITSv1 and ITS v2) DA. In the 2o3 DA, a hazard assessment is provided by two concordant, non-borderline (non-BL) results from DPRA, KS, and h-CLAT (Bauch et al., 2012; Natsch et al., 2021; Urbisch et al., 2015). The 2o3 DA does not provide information on the skin sensitization potency. While ITS v1 and v2 integrate an *in silico* prediction, the 2o3 is the sole DA based only on experimental data from OECD validated tests.

Assessing skin sensitization potency is needed for the binary subclassification of sensitizers into 1A (strong sensitizers) and 1B (other sensitizers) in the UN Global Harmonized System (GHS). The kDPRA assay, which has been recently added to OECD guideline 442c, is a standalone assay for the application of Sub-category 1A (Natsch et al., 2020; OECD, 2021b; Wareing et al.,

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2020). An assessment of potency on a more granular scale is needed for next generation risk assessment of new chemical entities. Thus, it is advantageous for risk assessors to have available approaches that can provide continuous PoD values so that more quantitative assessments can be made to help protect workers and consumers.

Linear regression models using KS and kinetic peptide reactivity data have been proposed to provide a PoD value in the form of a predicted EC3 value in the Local Lymph Node Assay (LLNA) (Natsch et al., 2015, 2018). Building on this previous approach using regression models, updated quantitative models using input data from the kDPRA, the KS and the h-CLAT were generated to calculate a PoD (Natsch and Gerberick, 2022). The predictive models were produced using a comprehensive database that included test data from the accepted OECD methods. All models were examined using a set of case studies selected based on multiple LLNA reference data in the OECD database. The robustness of the models was characterized by comparing a comprehensive historical database vs. the curated dataset provided by the OECD working group on DA. The predicted PoD were within or close to the variation of the historical LLNA data for most of the cases studies. Overall, the models predict the *in vivo* value with a median fold-misprediction factor of around 2.5. The various models offer risk assessors flexibility in the choice of tests and a PoD value can still be determined when there are compatibility issues or chemicals are outside the chemical domain of an individual assay.

In this paper, it is demonstrated how the kDPRA and these quantitative models can be combined in different testing sequences in the 2o3 DA to provide at the same time (i) hazard ID, (ii) GHS sub-classification, and (iii) PoD-determination based on the validated *in vitro* tests. The integrated assessment presented is solely based on *in vitro* data from the three OECD test guidelines. Thus this work further advances the 3R for skin sensitization testing as it gives practical guidance how to finally combine the methods and evaluates these proposed strategies on a large number of chemicals.

## 2 Materials and methods<sup>1</sup>

### 2.1. Database used

The analysis in this paper is based on a comprehensive database on 188 chemicals with data in the kDPRA, KS, h-CLAT and the LLNA and no new data were generated for this study (Table ESM1-1<sup>2</sup>; the data presented are a subset of the larger database presented in a parallel paper (Natsch and Gerberick, 2022). For 154 of these chemicals, data are also available in the OECD reference database (OECD DB) compiled by the OECD DA working group (OECD, 2021c). For all the 188 chemicals, LLNA data from published historical compilations are available. In parallel, for the subset in the OECD database, a curated LLNA value is available based on evaluating the original data with a set of fixed rules (OECD, 2021d). The analysis for accuracy of the PoD determination was made with the historical LLNA data and in parallel with the curated LLNA data, using the historical data only for the chemicals not in the OECD DB.

### 2.2. Regression models and statistics

The data normalizations and calculations are described in a parallel paper (Natsch and Gerberick, 2022). Based on the test data from the OECD tests, a Prediction Spreadsheet can be used to calculate a predicted EC3 as PoD based on regression equations. The following four regression models are further used here and are implemented in this prediction Spreadsheet:

For a PoD assessment based on KS and kDPRA data:

$$\text{EQ1} \quad \text{pEC3} = 0.42 + 0.40 \times \text{Log } k_{\text{max norm}} + 0.15 \times \text{Log EC1.5}_{\text{norm}} + 0.36 \times \text{Log IC50}_{\text{norm}} - 0.21 \times \text{Log VP}_{\text{norm}}$$

For a PoD assessment based on h-CLAT and kDPRA data:

$$\text{EQ4} \quad \text{pEC3} = 0.18 + 0.36 \times \text{Log } k_{\text{max norm}} + 0.21 \times \text{Log MIT}_{\text{norm}} + 0.35 \times \text{Log CV75}_{\text{norm}} - 0.19 \times \text{Log VP}_{\text{norm}}$$

For a PoD assessment based on KS, h-CLAT and kDPRA data:

$$\text{EQ5} \quad \text{pEC3} = 0.20 + 0.34 \times \text{Log } k_{\text{max norm}} + 0.20 \times \text{Log MIT}_{\text{norm}} + 0.09 \times \text{Log EC1.5}_{\text{norm}} + 0.21 \times \text{Log CV75}_{\text{norm}} + 0.11 \times \text{Log IC50}_{\text{norm}} - 0.19 \times \text{Log VP}_{\text{norm}}$$

For a PoD assessment based on KS and h-CLAT data:

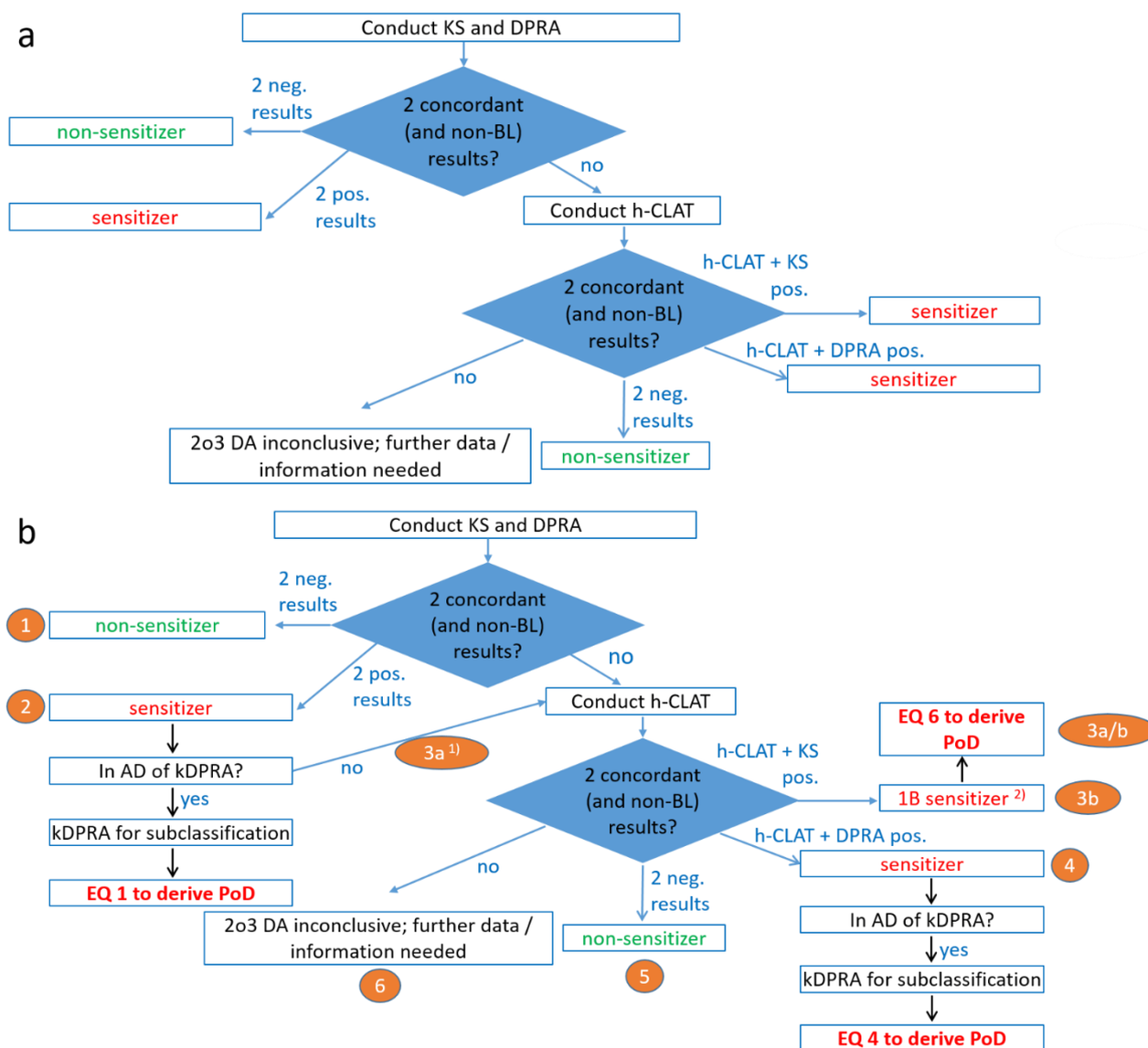
$$\text{EQ6} \quad \text{pEC3} = 0.09 + 0.276 \times \text{Log MIT}_{\text{norm}} + 0.22 \times \text{Log EC1.5}_{\text{norm}} + 0.34 \times \text{Log CV75}_{\text{norm}} + 0.06 \times \text{Log IC50}_{\text{norm}} - 0.12 \times \text{Log VP}_{\text{norm}}$$

The parameters used in these equations are (i) from the kDPRA the  $\text{Log } k_{\text{max norm}}$ , the normalized, logarithmic rate constant, (ii) from KS  $\text{Log IC50}_{\text{norm}}$ , the normalized IC50 value (concentration for 50% reduction in cellular viability) and  $\text{Log EC1.5}_{\text{norm}}$ , the normalized EC1.5 value indicating the concentration for 1.5-fold induction of luciferase activity and (iii) from h-CLAT the normalized  $\text{Log MIT}_{\text{norm}}$ , indicating the lowest concentration for either 1.5-fold CD86 or 2-fold CD54 induction and the  $\text{Log CV75}_{\text{norm}}$  indicating concentration for 25% reduction in viability. In addition, the  $\text{Log VP}_{\text{norm}}$  describes volatility for chemicals evaporating significantly from the LLNA vehicle within 60 min.

To assess prediction accuracy of quantitative models, the ratio between the larger and the smaller values of the measured and predicted EC3 value was calculated in each case to give the fold-misprediction. Median and geometric means were calculated for this measure of the data fit and the number of chemicals mis-predicted by > 5-fold or > 10-fold in either direction is listed.

<sup>1</sup> **Abbreviations:** AD, Applicability Domain; AOP, Adverse Outcome Pathway; BL, Borderline; DA, Defined Approach; DPRA, Direct Peptide Reactivity Assay; GHS, Global Harmonized System; hazard ID, >Hazard Identification; h-CLAT; Human Cell Line Activation Test; IATA; Integrated Approaches to Testing and Assessment; ITS, Integrated Testing Strategy; kDPRA, kinetic Direct Peptide Reactivity Assay; KS, KeratinoSens™; LLNA, Local Lymph Node Assay, NAM, New Approach Methodologies; PoD, Point of Departure; OECD, The Organisation for Economic Co-operation and Development; OECD DB, OECD reference database on defined approaches; QRA, Quantitative Risk Assessment; TG; Test Guideline; VP, Vapor Pressure; 2o3 DA, "2 out of 3" Defined Approach.

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**Fig. 1: An economic testing sequence includes GHS sub-classification and PoD determination into the 2o3 DA**  
 (a) The decision tree in Figure 2.1 in OECD TG 497 with KS and DPRA conducted first. (b) The decision tree expanded integrating the kDPRA for GHS subclassification and PoD determination as proposed in the current work. The numbers in orange bubbles indicate the different scenarios discussed in the text.

<sup>1)</sup> Chemicals outside of the applicability domain of the kDPRA according to APPENDIX III, ANNEX 1 of OECD TG 442C can be assessed based on h-CLAT and KS data if potency information is required.

<sup>2)</sup> Chemicals negative in DPRA and kDPRA, but positive in h-CLAT and KS are normally not 1A sensitizers based on kDPRA (TG442C) and based on DA ITS (TG497). Chemicals assessed with EQ 6 based on being outside of AD of kDPRA (scenario 3a) are not considered 1B chemicals directly unless DPRA is negative.

For assessment of subclassification, sensitizers were discriminated from non-sensitizers with the 2o3 DA taking borderline (BL) outcomes in the individual tests into account as described in OECD TG 497 (OECD, 2021a). Data are presented as a three-way classification Table. For analysis of this prediction of three classes, only the OECD data were used as BL analysis could not be made on the additional published h-CLAT data.

### 3 Results

#### 3.1 An economic testing sequence to include GHS sub-classification and PoD determination into the 2o3 DA

In the 2o3 DA, a hazard assessment is provided by two concordant, non-BL results from DPRA, KS, and h-CLAT (OECD, 2021a). The testing sequence does not affect the outcome of this hazard assessment. Here, we provide the most economical testing sequence and indicate two alternative approaches (either starting with h-CLAT or conducting all assays by default). The goal of all these testing sequences, as described here, is to provide hazard ID, GHS sub-classification and PoD determination based on results from DPRA, kDPRA, KS and h-CLAT.

An efficient testing sequence is shown in Figure 1. Testing starts with DPRA and KS since these tests are more economical in most test laboratories and they lead to less inconclusive /borderline outcomes as compared to the h-CLAT (OECD, 2021c), thus

fewer instances will require the third test to be conducted. Also, during the validation of the 2o3 DA at the OECD it was clearly shown, that the sequence of testing does not affect the outcome of the 2o3 DA. Two non-BL negative results lead to a negative call (Scenario 1), while two non-BL positive results are sufficient for classification as a sensitizer (Scenario 2). If the chemical is within the applicability domain (AD) of the kDPRA, conducting the kDPRA provides the information on whether the chemical must be sub-classified as 1A. The combined dose-response information from a positive kDPRA and the positive KS is then applied in the regression model in the standardized prediction spreadsheet using EQ1 to derive the PoD. However, if the chemical is not within the AD of the kDPRA (Scenario 3a), it is recommended to perform the h-CLAT to gather more evidence on potency by applying EQ6. If either the DPRA or KS was negative or BL, the h-CLAT must be conducted according to the 2o3 scheme. Two negative, non-BL outcomes again indicate a non-sensitizer (Scenario 5), and a BL outcome leads to an inconclusive assessment (Scenario 6). A positive h-CLAT with a positive result from either KS or DPRA leads to classification. If the DPRA and the h-CLAT are positive (Scenario 4), chemicals within the AD of the kDPRA can then be sub-classified based on the kDPRA and assessed for PoD with the regression model EQ4.

If the DPRA is negative, two positives in KS and h-CLAT can lead to classification (Scenario 3b) and a PoD can be derived based on EQ6. In this case a sub-classification of 1B can directly be made: a negative call in the DPRA and hence a negative call in the kDPRA is sufficient for chemicals to be classified as 1B. (Note: This is also consistent with the outcome from the alternative validated DA, ITS, whereby a chemical negative in the DPRA is not classified as 1A sensitizer, as it cannot reach a score 6 or 7 (OECD, 2021a)). However, in scenario 3a (i.e., a chemical not in the AD of the kDPRA but the DPRA is positive), the GHS sub-classification is inconclusive. In this case, the outcome of the PoD with EQ6 may still be used for a WoE assessment to indicate whether the LLNA potency is predicted to be at an EC3 < 2%. However, according to the OECD guideline, this would not be sufficient for a conclusive 1B classification.

For scenario 6, i.e., an inconclusive outcome of the 2o3 due to borderline results, the result can be due to borderline negative results. In this case also no relevant PoD can be calculated as no EC1.5 or MIT or reaction rate is derived from the borderline tests. On the other hand, if the result is borderline positive, EC1.5 or MIT values are available and a PoD can be calculated, but it has a lower certainty. These values were still given in ESM1<sup>2</sup> (13 cases), as OECD guideline states that inconclusive outcomes could still be used in a weight-of-evidence.

This proposed testing sequence might be further simplified if chemical reactivity is expected, e.g., based on structural alerts. Then the kDPRA could be directly done instead of the DPRA. A positive result in the kDPRA (> 13.89% Cys peptide depletion) may then be used as a positive rating along with a positive from KS and/or h-CLAT. If the kDPRA were negative, the DPRA would still need to be conducted to confirm the negative result. This approach may further save tests if a chemical has a high likelihood of a positive outcome in the (k)DPRA.

## 3.2 Alternative testing sequences

### 3.2.1 All data is generated

The tiered economic testing strategy in Figure 1 with conditional testing in h-CLAT based on the outcome of the first two tests may be considered time-consuming by some users. An alternative option is to test a new chemical directly in KS, h-CLAT and DPRA by default. If two tests are positive, and one is the DPRA, the kDPRA is conducted. In this case the hazard ID and the GHS sub-classification can directly be made based on the data (unless BL results are obtained, or the chemical is outside of the AD), and the PoD can be calculated with EQ5 taking all evidence into account. If the chemical is outside of the AD of the kDPRA but positive in h-CLAT and KS, application of EQ6 is warranted (identical to Scenario 3a in Figure 1). As EQ5 is used for most chemicals in this approach, the derived PoD can differ from the approach in Figure 1, which relies on models based on data from two positive tests (EQ1, EQ4 and EQ6).

### 3.2.2 Testing starts with DPRA and h-CLAT

The testing sequence in Figure 1 can also be modified, and the testing starts with DPRA and h-CLAT. KS then is only conditionally used in the same way as h-CLAT is used in Figure 1 (Figure ESM2-1<sup>3</sup>). This alternative approach will not change the outcome for GHS sub-classification and hazard ID. However, in this case, the PoD is more frequently derived with EQ4 instead of EQ1, as all chemicals positive in the first two assays (i.e., h-CLAT and DPRA) will be assessed based on EQ4.

## 3.3 PoD outcome for chemicals with available KS, h-CLAT and kDPRA data

Table 1 summarizes the prediction accuracy for the three different testing sequences, namely (i) Prediction of the PoD according to Figure 1, (ii) prediction made based on EQ5 / EQ6 in cases when all data are generated and (iii) with h-CLAT done first (Figure ESM2-1<sup>3</sup>). The individual predictions and the scenario/equation used for each chemical with these three approaches are given in ESM1<sup>2</sup>, along with the correlation between the different assessments for each chemical (Figures ESM1<sup>2</sup>, 1-3). In all three cases, the prediction accuracy is quite similar and leads to a comparable number of > 5-fold or > 10 fold (i.e., a full potency class) mispredictions vs. the LLNA result. As is obvious from Table 1 and the scatter plots (Figure ESM1<sup>2</sup> 1 – 6), and the data on the individual chemicals in ESM1, for most chemicals, the predicted PoD are similar between the different testing sequences and there is no tendency that one testing sequence is, in general, less conservative. The number of overpredicted chemicals, however, is decreased when using all evidence, as the negative evidence for chemicals positive in only two assays is taken into account and this approach (EQ5) therefore also leads to a slightly better correlation to *in vivo* data (see Figures ESM1<sup>2</sup>, 4 - 6).

<sup>3</sup> doi:10.14573/altex.2201142s2

**Tab. 1: Chemicals rated positive by the 2o3 DA (n=116)<sup>a,b</sup> assessed for PoD using different testing sequences**

Approach	Fold-mispre-diction <sup>c</sup> (Geomean)	Fold-mispre-diction <sup>c</sup> (Median)	Chemicals > 5 – fold under-predicted <sup>d</sup> (n, %)	Chemicals > 10 – fold under-predicted (n, %)	Chemicals > 5 – fold over-predicted <sup>d</sup> (n, %)	Chemicals > 10 – fold over-predicted (n, %)
According to Figure 1	3.8	3.2	17 (15%)	7 (6%)	23 (20%)	10 (9%)
Performing h-CLAT and DPRA first	3.8	3.4	17 (15%)	6 (5%)	21 (18%)	11 (9%)
Using all evidence	3.4	2.6	19 (16%)	6 (5%)	16 (14%)	6 (5%)

<sup>a</sup> As compared to the parallel analysis (Natsch and Gerberick, 2022) comparing the different equations on all chemicals including negatives, this analysis is focused on the subset of chemicals rated positive in the 2o3 DA and assessed with different testing sequences

<sup>b</sup> For 21 chemicals, the published h-CLAT value could not be assessed whether the result is BL, and for 7 of those a h-CLAT and DPRA first call could lead to an inconclusive 2o3 assessment in theory. These data were treated as is, not taking potential BL results in h-CLAT into account for this analysis, which is focused on PoD, not hazard.

<sup>c</sup> The ratio between the higher and the lower values of the measured and predicted EC3 value. Predicted EC3 > 100% were set to 100%.

<sup>d</sup> Under-predicted chemicals: those for which the measured LLNA EC3 is < than the predicted EC3; over-predicted chemicals: Those with measured LLNA EC3 > than the predicted value.

### 3.4 Analysis of significant over- and under-predictions

To analyze individual mispredictions, we focused on the outcome of the testing sequence in Figure 1. Table ESM3-1<sup>4</sup>, see supplementary material, lists all the chemicals that are > 5-fold underpredicted, i.e., their potency as assessed by the LLNA is significantly higher than the predicted PoD. The chemicals in this Table are grouped and an individual discussion is given for each chemical. In summary, a set of 6 chemicals is underpredicted as weak sensitizers with predicted EC3 of 9.2 – 55%, while they are moderate sensitizers in the LLNA. These include *inter alia* primary amines/pro-haptens and amine-reactive chemicals, which are outside of the AD of the kDPRA (OECD, 2021b). For a larger group (n=12), the predicted PoD indeed indicates a significant sensitization potency (predicted EC3 0.05% – 5%), but the individual values are clearly below the strong to extreme potency observed in the LLNA. This indicates that the dynamic range for the exact potency assessment of some extreme sensitizers using the regression models is limited. However, a high sensitization potential is predicted for most chemicals in this group based on the *in vitro* data.

Table ESM3-2<sup>4</sup> provides data and discussion on the chemicals with a predicted PoD below the LLNA EC3, i.e., a higher potency is predicted *in vitro*. This group contains six false positives in the 2o3 vs. LLNA outcome. For four of those, positive human sensitization evidence or a strong alkylating potency indicate that the LLNA actually underpredicts the sensitization potential. In contrast, for two others, the reported human sensitization potential is rather weak (propyl paraben and benzocaine) and clearly over-rated by the *in vitro* approach. A further group (n=5) are very reactive and volatile chemicals. Although EQ1 corrects for high volatility, it does not fully predict the weak sensitization in LLNA observed for these highly reactive chemicals that evaporate rapidly under LLNA conditions (see Supplementary data file 1 in Natsch et al., 2015). However, these chemicals may be significantly more potent under (partial) occlusion or when present in a product limiting evaporation. Hence, this conservative assessment by the *in vitro* derived PoD may be appropriate. Another set of chemicals (n=5) are clearly over-predicted when assessed vs. LLNA data, but either clinical data or human repeat insult patch test indicate that these are very relevant human sensitizers, and the *in vitro* prediction is more conservative than the LLNA but could better reflect the human sensitization potency. No human data are available for the remaining seven chemicals, but they still include several highly reactive chemicals.

The analysis in Table 1 and in ESM3<sup>4</sup> is based on the comparison to the comprehensive historical LLNA database. An additional analysis was conducted based on the OECD curated EC3 values, taking the historical database values only where no curated EC3 was available. This analysis is shown and compared to above analysis in ESM2<sup>3</sup>. The outcome of both analysis is almost congruent.

**Tab. 2: GHS sub-classification of the chemicals in the OECD database by the 2o3 DA combined with kDPRA**

Prediction 2o3 DA with kDPRA	LLNA Result		
	NC (n =26)	1B (n = 85)	1A (n = 38)
NC	21	16	0
1B	3	34	7 (4) <sup>a</sup>
1A	1	14	26 (29)
Correct	84%	53%	79% (88%)
Under	NA	25%	21% (12%)
Over	16%	22%	NA
Inconclusive	n = 8	n =21	n = 5

<sup>a</sup> The values are based on applying only the prediction model of the kDPRA and 2o3 DA. In brackets are given the values if taking the AD of the kDPRA into account and applying Scenario 3a in Figure 1 (Using EQ6 for chemicals outside of AD of kDPRA).

<sup>4</sup> doi:10.14573/altex.2201142s3

### 3.5 Hazard ID and GHS sub-classification outcome for chemicals in the OECD database

If the kDPRA is combined with the 2o3 DA in a testing strategy, chemicals can be rated both for hazard and for GHS potency classes. As indicated above, this is independent of the testing sequence with all three testing proposals leading to the same outcome. In Table 2, we show the outcome for the classification rating on the chemicals in the OECD database for which an LLNA sub-classification is available in the database ( $n = 156$ ) compared to LLNA reference data. Chemicals in our dataset but not in the OECD database are excluded from this analysis since the published h-CLAT data could not be analyzed for BL outcomes for these chemicals.

Table ESM4-1<sup>5</sup>, see supplementary material, lists all chemicals which were not correctly predicted by this three-way classification using the 2o3 DA and the kDPRA. For each chemical, background information on what is known on the human sensitization potential or sensitization as reported from clinical studies is added. This analysis is partly overlapping with the analysis in ESM3<sup>4</sup>, as several chemicals for which the PoD is > 5-fold mispredicted as compared to the LLNA EC3 value by the integrated data from the cell-based assays and the kDPRA are also miss-classified by the prediction threshold of the kDPRA used for sub-classification and by the hazard models of the individual tests.

## 4 Discussion

The 2o3 DA has been accepted as an OECD standard for hazard ID. At the same time, the kDPRA can be used as a stand-alone test for GHS sub-classification once a chemical is identified as a skin sensitizer. Thus, combining these two approaches for classification and sub-classification, as illustrated here, is a straightforward strategy. This combination will not require further validation for both the hazard and the sub-classification decision as both prediction models were validated and implemented in the two OECD TG 497 and 442C for chemicals considered within the AD (OECD, 2021a,b).

For this classification approach, only the positive/negative answers from the validated prediction models in KS/h-CLAT/DPRA or the validated binary classification according to a quantitative threshold ( $\text{Log } k_{\text{max}} = -2$ ) in the kDPRA are used. However, the data generated are more granular (quantitative kinetic rate constant over several orders of magnitude in kDPRA and dose-response data over three orders of magnitude in the cell-based assays). As we showed in the parallel analysis (Natsch and Gerberick, 2022), this dose-response data can be used to estimate a PoD. Thus, the same test results generated for the GHS (sub)classification can be used for the potency assessment and to derive a PoD in the integrated testing and assessment sequences provided here.

The different sequences can start with either of the two cell-based assays or generate data with all three tests as a default. Different predictive equations can be applied for PoD determination depending on the generated data. The analysis of the outcome for the individual chemicals indicates that the different testing sequences using other predictive equations overall lead to surprisingly similar predictions (Figure S2 – S4). This confirms previous observations on data redundancy esp. between quantitative data from h-CLAT and KS (Natsch et al., 2015; Natsch and Gerberick, 2022). Still, it also indicates that the different testing sequences are all valid approaches and neither of them leads to an overall less conservative risk assessment. Since the various *in vitro* assessments correlate better to each other than to the *in vivo* data (see Figures S2-S7 in ESM 2), the key open question is whether other *in vitro* assays will provide further, more orthogonal information for a further improved PoD determination, or whether this asymptotic fit to *in vivo* data when adding more *in vitro* information also partly reflects the limitations of the *in vivo* data source.

In any case, it is of utmost importance to understand the sources of uncertainty between the *in vitro* and *in vivo* datasets. Part of the uncertainty comes from both the biological variabilities of the LLNA and the *in vitro* data. For the LLNA, analysis of repeated studies (Dumont et al., 2016; Hoffmann, 2015) indicates that the typical standard deviation of EC3 values is 1.8-fold to either direction, but larger discrepancies were noted in some cases. This will lead to uncertainty of the *in vivo* comparable esp. in instances where only one LLNA study is available. Biological variability in the *in vitro* data (Gabbert et al., 2020; Leontaridou et al., 2017) will further increase uncertainty and thus, variability in both datasets will always limit the fit between them. However, this data variability can only explain part of the prediction inaccuracy. A further part of the uncertainty is certainly because the *in vitro* tests are not yet a perfect reflection of the sensitization process, as they all only measure surrogates of some key events (e.g., no T cell activation).

On the other hand, as illustrated by the detailed analysis of the individual chemicals with > 5-fold misprediction, part of the inaccuracy may also be because the LLNA is not a perfect model of potency for all chemicals, reminding us that the LLNA itself measures only part of the sensitization process (antigen-presentation triggered cell proliferation in the lymph node). Thus, for some negatives in the LLNA but positive by the *in vitro* assessment, data from human studies and/or the alkylating potential observed in peptide reactivity studies indicate that the LLNA may be false-negative and the *in vitro* result may give a more accurate estimation of the sensitization risk. Similarly, several of those chemicals for which the potency is overestimated by the *in vitro* PoD are critical skin sensitizers from human clinical studies, especially some preservatives and glove allergens. When analyzing the under-predictions, on the other hand, the *in vitro* PoD appears not to perfectly cover the dynamic range for the very potent sensitizers. Thus, it is noteworthy that some of the extreme sensitizers are predicted as strong sensitizers based on the PoD, but the predictive models do not yet reflect their full potency in the LLNA.

Turning to the GHS classification and sub-classification outcome, the predictivity is better for predicting non-sensitizers and strong (1A) sensitizers in the LLNA, and the predictivity for the LLNA 1B sensitizers is less accurate with around 22-25% mispredictions in either direction. While a more limited predictivity for the intermediate class (where misprediction to either side is possible) is an intrinsic property for any three-way classification scheme, the absolute number of correct classifications may be considered relatively low. Here we thus provided a detailed analysis for the individual mispredicted chemicals regarding the GHS classification (ESM4<sup>5</sup>). Next to general limitations of prediction accuracy based on data variability discussed above, some of the predictive limitations for correct classifications can be attributed to (i) limitations of the applicability domain (AD) of the *in vitro* assays and partial coverage of key events, (ii) only partial coverage of the human sensitization potential and potency by the LLNA model, (iii) the fact that some *in vivo* and *in vitro* results are very close to the decision threshold (LLNA EC3 of 2% / kDPRA threshold of  $\text{log}k_{\text{max}} = -2$ ).

<sup>5</sup> doi:10.14573/altex.2201142s4

The kDPRA has an important weight in the potency determination (Natsch and Gerberick, 2022). Thus, it is critical to assess whether a chemical is in the AD of the kDPRA. OECD TG indicates that test chemicals with exclusive lysine-reactivity as observed in DPRA or ADRA are outside of the AD of the kDPRA as the kinetic reactivity with lysine residues is covered neither by the kDPRA nor the testing schemes shown here. Such chemicals, if positive in both KS and h-CLAT may still be assessed with the regression models. Thus, the PoD for glutaraldehyde – a chemical not in the AD of the kDPRA and mispredicted for potency using kDPRA only – is predicted based on EQ6 with a PoD of 0.6%, which is still higher than the LLNA EC3 of 0.1% but in the correct GHS class. For another amine-reactive chemical, 3,4-dihydrocoumarin potency is underrated. While it is possible to measure amine reactivity of these chemicals, it may be a significant challenge to derive quantitative potency models based on the limited number of typical amine reactive chemicals as a training set (An exception are aldehydes, for which we provided such a model previously (Natsch et al., 2018)). A second limitation indicated for the kDPRA is “aromatic amines, catechols or hydroquinones” which may require further data to confirm their weak reactivity if their  $\log k_{\max}$  is  $< -2$ . Thus, there are two cases among the seven mispredicted chemicals rated as 1B instead of 1A (1,4-phenylenediamine and 2-amino-phenol). These are rated as 1A if EQ6 is applied.

Next to considering the applicability of the *in vitro* tests, it is also key to look at a WoE when assessing the wrong *in vitro* classifications vs. the LLNA outcome. Thus, the analysis of the LLNA data as performed by the OECD data review indicated a limitation of the LLNA for specificity vs. human data (Natsch et al., 2021; OECD, 2021a). This is partly because the review criteria required a higher maximal test concentration to conclude on a negative call in the LLNA as compared to the validation of the LLNA (Kolle et al., 2020). Also, the estimate of specificity vs. human data is based on a relatively low number of chemicals, but it indicates that the database does contain some false-positive chemicals in the LLNA. Indeed, among the 16 FN in 2o3 vs. LLNA data, there are seven chemicals for which the WoE indicates that they are not, or extremely weak, human sensitizers (ESM4<sup>5</sup>). On the other hand, among the over-predicted chemicals, as discussed above for the PoD, for several cases, the sensitization potency and correct GHS class could be underestimated by the LLNA and more correctly reflected by the *in vitro* PoD (ESM4<sup>5</sup>).

The integrated assessment discussed here is solely based on *in vitro* data from the three OECD TG and no *in silico* assessment is integrated into this approach, differently from almost all published approaches for an integrated evaluation of the sensitization potential (Del Bufalo et al., 2018; Hirota et al., 2018; Jaworska et al., 2015; Macmillan and Chilton, 2019; Strickland et al., 2017; Takenouchi et al., 2015). There are some benefits to the present approach conducting an assessment based solely on validated OECD test methods and not, from the start, integrating an *in silico* prediction: (i) Most *in silico* tools were developed and trained partly on the database with available *in vitro* and *in vivo* data and rule-based approaches based on structural alerts in principle have an unlimited number of degrees-of-freedom. Using *in silico* tools on the same database without separating test and training set may thus lead to an overfitted model. This problem is minimal for the PoD models used here as they are based on only 3 - 6 input variables and trained on  $> 180$  chemicals, (ii) when conducting an assessment solely based on *in vitro* data; an independent, parallel assessment can then be made applying the *in silico* tools to increase certainty and obtain a more holistic picture. If the *in silico* tool is already integrated into the initial prediction this is not possible without double-accounting, (iii) *in silico* tools as implemented, e.g., in TG497 (OECD, 2021a) have a relatively strict definition in their AD for known chemical features esp. to make conclusive negative predictions. Thus, using an *in silico* tool by default has implications to the overall AD for new chemicals to be assessed. Approaches to perform a WoE assessment on existing chemicals have been described using only human data (Basketter et al., 2014) or combining human, animal, *in vitro* and *in silico* data. For new chemicals, the human and animal part would be lacking, but the here proposed integrated *in vitro* approach can then be combined with parallel *in silico* predictions for a WoE.

Here we focused the analysis in relation to the LLNA outcome. When assessing hazard ID, looking at the human data is important (Natsch et al., 2021; OECD, 2021a) as the LLNA may also have limitations in specificity if used at too high concentrations or if not taking irritation into account as indicated above. However, only for a minority of chemicals quantitative human data on potency is available, and since we discuss here how to combine potency assessment into the 2o3 DA, the key analysis presented here was performed vs. the LLNA potency data. Nevertheless, we indicate in the discussion on individual chemicals (ESM3<sup>4</sup> and ESM4<sup>5</sup>) the semi-quantitative potency information from human data when available (Api et al., 2017; Basketter et al., 2014; OECD, 2021e) as these data further help to assess in which cases the *in vitro* data truly underestimate potency, but also highlight cases when the NAM assessment may lead to a more correct and more conservative human risk assessment.

The proposed testing sequences for (sub)-classifications and PoD determination are a proposal to make the best use of the data generated by testing according to TG 442c, 442d and 442e. The PoD could be used directly in risk assessment, and in an absence of other evidence, a default assessment factor may be introduced to account for uncertainty (Natsch et al., 2015). As risk assessors transition to using NAM data for potency assessment, a PoD derived from these regression models could be integrated into existing risk assessment schemes such as the Quantitative Risk Assessment QRA (Api et al., 2020). However, the assessment certainly does not stop there: Analysis of the prediction accuracy of close analogues with both *in vitro* and *in vivo* data will help for further refinement of the uncertainty for specific chemicals (Natsch et al., 2018). The large database provided in this and the parallel analysis (Natsch and Gerberick, 2022) and the increasing database from other initiatives will further help to investigate better in which chemical domain certainty is higher or lower and will provide read-across analogues to conduct such an uncertainty analysis in the specific chemical domain of the molecule to be assessed. Furthermore, depending on the chemical domain, further non-guideline methods can be applied to test specific parameters, such as metabolic activation by metabolic systems, reactivity with amine groups, or epidermal disposition. Such further evidence can then refine the PoD derived from the presented standard testing sequences.

### Electronic supplementary material

ESM1<sup>2</sup> gives the *in vitro* and *in vivo* data in Sheet 1, Sheet 2 gives all the individual predictions and fold-mispredictions for 188 chemicals with the three different testing sequences and Sheet 3 gives the graphical correlations between the different predictions and between predictions and *in vivo* results.

ESM2<sup>3</sup> gives the testing sequence starting with h-CLAT and comparison of predictions to OECD curated LLNA values.

ESM3<sup>4</sup> discusses  $> 5$ -fold misprediction vs. LLNA outcome also in light of other (e.g., human) evidence

ESM4<sup>5</sup> discusses GHS-misclassifications also in light of other (e.g., human) evidence

## References

- Api, A. M., Basketter, D., Bridges, J., Cadby, P. et al. (2020). Updating exposure assessment for skin sensitization quantitative risk assessment for fragrance materials. *Regul Toxicol Pharmacol* 118, 104805. doi:10.1016/j.yrtph.2020.104805
- Api, A. M., Parakhia, R., O'Brien, D. and Basketter, D. A. (2017). Fragrances Categorized According to Relative Human Skin Sensitization Potency. *Dermatitis* 28, 299-307. doi:10.1097/DER.0000000000000304
- Basketter, D. A., Alepee, N., Ashikaga, T., Barroso, J. et al. (2014). Categorization of chemicals according to their relative human skin sensitizing potency. *Dermatitis* 25, 11-21. doi:10.1097/DER.0000000000000003
- Bauch, C., Kolle, S. N., Ramirez, T., Eltze, T. et al. (2012). Putting the parts together: Combining in vitro methods to test for skin sensitizing potentials. *Regul Toxicol Pharmacol* 63, 489-504. doi:10.1016/j.yrtph.2012.05.013
- Bernauer, U., Bodin, L., Chaudhry, Q., Coenraads, P. J. et al. (2021). The SCCS Notes of Guidance for the testing of cosmetic ingredients and their safety evaluation, 11th revision, 30–31 March 2021, SCCS/1628/21. *Regulatory Toxicology and Pharmacology* 127, 105052. doi:10.1016/j.yrtph.2021.105052
- Del Bufalo, A., Pauloin, T., Alepee, N., Clouzeau, J. et al. (2018). Alternative Integrated Testing for Skin Sensitization: Assuring Consumer Safety. *Applied In Vitro Toxicology* 4, 30-43. doi:10.1089/aivt.2017.0023
- Dent, M., Amaral, R. T., Da Silva, P. A., Ansell, J. et al. (2018). Principles underpinning the use of new methodologies in the risk assessment of cosmetic ingredients. *Computational Toxicology* 7, 20-26. doi:10.1016/j.comtox.2018.06.001
- Dumont, C., Barroso, J., Matys, I., Worth, A. et al. (2016). Analysis of the Local Lymph Node Assay (LLNA) variability for assessing the prediction of skin sensitisation potential and potency of chemicals with non-animal approaches. *Toxicol In Vitro* 34, 220-228. doi:10.1016/j.tiv.2016.04.008
- Ezendam, J., Braakhuis, H. M. and Vandebriel, R. J. (2016). State of the art in non-animal approaches for skin sensitization testing: from individual test methods towards testing strategies. *Arch Toxicol* 90, 2861-2883. doi:10.1007/s00204-016-1842-4
- Gabbert, S., Mathea, M., Kolle, S. N. and Landsiedel, R. (2020). Accounting for Precision Uncertainty of Toxicity Testing: Methods to Define Borderline Ranges and Implications for Hazard Assessment of Chemicals. *Risk Anal* Article in press. doi:10.1111/risa.13648
- Gilmour, N., Kern, P. S., Alepee, N., Boislevé, F. et al. (2020). Development of a next generation risk assessment framework for the evaluation of skin sensitisation of cosmetic ingredients. *Regul Toxicol Pharmacol* 116, 104721. doi:10.1016/j.yrtph.2020.104721
- Hirota, M., Ashikaga, T. and Kouzuki, H. (2018). Development of an artificial neural network model for risk assessment of skin sensitization using human cell line activation test, direct peptide reactivity assay, KeratinoSens and in silico structure alert parameter. *J. Appl. Toxicol.* 38, 514-526. doi:10.1002/jat.3558
- Hoffmann, S. (2015). LLNA variability: An essential ingredient for a comprehensive assessment of non-animal skin sensitization test methods and strategies. *ALTEX* 32, 379-83. doi:10.14573/altex.1505051
- Hoffmann, S., Kleinstreuer, N., Alépée, N., Allen, D. et al. (2018). Non-animal methods to predict skin sensitization (I): the Cosmetics Europe database<sup/>. *Crit Rev Toxicol* 48, 344-358. doi:10.1080/10408444.2018.1429385
- Jaworska, J. S., Natsch, A., Ryan, C., Strickland, J. et al. (2015). Bayesian integrated testing strategy (ITS) for skin sensitization potency assessment: a decision support system for quantitative weight of evidence and adaptive testing strategy. *Arch. Toxicol.* 89, 2355-83. doi:10.1007/s00204-015-1634-2
- Kleinstreuer, N. C., Hoffmann, S., Alépée, N., Allen, D. et al. (2018). Non-animal methods to predict skin sensitization (II): an assessment of defined approaches (\*). *Crit Rev Toxicol* 48, 359-374. doi:10.1080/10408444.2018.1429386
- Kolle, S. N., Landsiedel, R. and Natsch, A. (2020). Replacing the refinement for skin sensitization testing: Considerations to the implementation of adverse outcome pathway (AOP)-based defined approaches (DA) in OECD guidelines. *Regul Toxicol Pharmacol* 115, 104713. doi:10.1016/j.yrtph.2020.104713
- Leontaridou, M., Urbisch, D., Kolle, S. N., Ott, K. et al. (2017). The borderline range of toxicological methods: Quantification and implications for evaluating precision. *Altex* 34, 525-538. doi:10.14573/altex.1606271
- Macmillan, D. S. and Chilton, M. L. (2019). A defined approach for predicting skin sensitisation hazard and potency based on the guided integration of in silico, in chemico and in vitro data using exclusion criteria. *Regul Toxicol Pharmacol* 101, 35-47. doi:10.1016/j.yrtph.2018.11.001
- Natsch, A., Emter, R., Gfeller, H., Haupt, T. et al. (2015). Predicting Skin Sensitizer Potency Based on In Vitro Data from KeratinoSens and Kinetic Peptide Binding: Global Versus Domain-Based Assessment. *Toxicol. Sci.* 143, 319-32. doi:10.1093/toxsci/kfu229
- Natsch, A., Emter, R., Haupt, T. and Ellis, G. (2018). Deriving a No Expected Sensitization Induction Level for Fragrance Ingredients Without Animal Testing: An Integrated Approach Applied to Specific Case Studies. *Toxicol Sci* 165, 170-185. doi:10.1093/toxsci/kfy135
- Natsch, A., Gerberick, G.F. (2022). Integrated skin sensitization assessment based on OECD methods (I): Deriving a Point of Departure for risk assessment. *ALTEX* Online ahead of print. doi:10.14573/altex.2201141
- Natsch, A., Haupt, T., Wareing, B., Landsiedel, R. et al. (2020). Predictivity of the kinetic direct peptide reactivity assay (kDPRA) for sensitizer potency assessment and GHS subclassification. *ALTEX* 37, 652-664. doi:10.14573/altex.2004292
- Natsch, A., Landsiedel, R. and Kolle, S. N. (2021). A triangular approach for the validation of new approach methods for skin sensitization. *ALTEX* 38, 669-677. doi:10.14573/altex.2105111
- OECD (2014). The Adverse Outcome Pathway for Skin Sensitisation Initiated by Covalent Binding to Proteins.
- OECD (2018a). In vitro skin sensitisation assays addressing the AOP key event on keratinocyte activation. *OECD testing guidelines* 442d. doi:10.1787/9789264229822-en
- OECD (2018b). In vitro skin sensitisation assays addressing the key event on activation of dendritic cells on the adverse outcome pathway for skin sensitisation. *OECD testing guidelines* 442e. doi:10.1787/9789264264359-en
- OECD (2021a). Guideline No. 497: Defined Approaches on Skin Sensitisation. *OECD Guidelines for the Testing of Chemicals, Section 4, OECD Publishing, Paris.* doi:10.1177/026119290703500311
- OECD (2021b). In Chemico Skin Sensitisation Assays addressing the Adverse Outcome Pathway, key event on covalent binding to proteins. *OECD testing guidelines* 442c. doi:10.1787/9789264229709-en

- OECD (2021c). Series on Testing and Assessment No. 336: Supporting document to the Guideline (GL) on Defined Approaches (DAs) for Skin Sensitisation- Annex 2. Organisation for Economic Cooperation and Development, Paris. <https://www.oecd.org/chemicalsafety/testing/series-testing-assessment-publications-number.htm>
- OECD (2021d). Series on Testing and Assessment No. 336: Supporting document to the Guideline (GL) on Defined Approaches (DAs) for Skin Sensitisation- Annex 3. Organisation for Economic Cooperation and Development, Paris. <https://www.oecd.org/chemicalsafety/testing/series-testing-assessment-publications-number.htm>
- OECD (2021e). Series on Testing and Assessment No. 336: Supporting document to the Guideline (GL) on Defined Approaches (DAs) for Skin Sensitisation- Annex 4. Organisation for Economic Cooperation and Development, Paris. <https://www.oecd.org/chemicalsafety/testing/series-testing-assessment-publications-number.htm>
- Strickland, J., Zang, Q., Paris, M., Lehmann, D. M. et al. (2017). Multivariate models for prediction of human skin sensitization hazard. *J Appl Toxicol* 37, 347-360. doi:10.1002/jat.3366
- Takenouchi, O., Fukui, S., Okamoto, K., Kurotani, S. et al. (2015). Test battery with the human cell line activation test, direct peptide reactivity assay and DEREK based on a 139 chemical data set for predicting skin sensitizing potential and potency of chemicals. *Journal of Applied Toxicology* 35, 1318-1332. doi:10.1002/jat.3127
- Urbisch, D., Mehling, A., Guth, K., Ramirez, T. et al. (2015). Assessing skin sensitization hazard in mice and men using non-animal test methods. *Regul Toxicol Pharmacol* 71, 337-51. doi:10.1016/j.yrtph.2014.12.008
- Wareing, B., Kolle, S. N., Birk, B., Alepee, N. et al. (2020). The kinetic direct peptide reactivity assay (kDPRA): Intra- and inter-laboratory reproducibility in a seven-laboratory ring trial. *ALTEX* 37, 639-651. doi:10.14573/altex.2004291

### **Conflict of interest statement**

The authors declare no competing interests. All data of this publication are made publicly available and all models and tests used are freely available.

### **Data availability statement**

All input data are available in ESM1<sup>2</sup> to this manuscript on the first datasheet. All individual predictions are included in ESM1<sup>2</sup> on the second spreadsheet.