Introduction

The progress achieved in developing new approach methodologies (NAM) for assessing the skin sensitization potential of chemicals over the last two decades has been remarkable. The Organisation for Economic Co-operation and Development (OECD) led the development of the adverse outcome pathway for skin sensitization divided into mechanistic key events (OECD, 2014). Three OECD guidelines have been published that cover these mechanistic events (covalent binding to protein, keratinocyte activation and dendritic cell activation). A total of eight non-animal test methods are approved in OECD TG 442C, 442D and 442E (OECD, 2018a,b, 2021d). However, none of the eight test methods are considered standalone replacements for complete hazard identification or potency determination.

The current focus is to find ways to combine in vitro, in chemico and in silico assessments with read-across predictions from similar chemicals to generate integrated approaches to testing and assessment (IATA) or defined approaches (DA). The OECD has published a new guideline (TG 497) that describes two simple DA for assessing skin sensitization (OECD, 2021a). This strategy of combining NAM data will enhance the vigor of skin sensitization hazard identification calls on chemicals and further address the critical need for predicting chemical potency. Potency prediction is needed for the UN Global Harmonized System (GHS) subclassification of sensitizers into 1A (strong sensitizers) and 1B (other sensitizers). For the specific needs of GHS subclassification into subcategory 1A, the kinetic direct peptide reactivity assay (kDPRA) is accepted as a standalone assay (Natsch et al., 2020; OECD, 2021d). However, assessing potency is also required for conducting next generation risk assessments (NGRA) on new chemical entities for which only non-animal information is available (Api et al., 2020; Bernauer et al., 2021; Dent et al., 2018; Gilmour et al., 2020). Potency prediction for risk assessment needs to be more granular than the GHS subclassification, and it is expected that this granularity can only be provided by combining multiple methods. Such
potency information is critical to ensure any new chemical is safe for exposed workers and consumers.

Multiple studies have addressed the application of quantitative in silico, in chemico and in vitro data from the validated assays for potency prediction (Gilmour et al., 2020; Kleinsteuber et al., 2018; Natsch et al., 2018; OECD, 2021a). A few NAMs have been designed to predict potency, including the SENS-IS® (Cottrez et al., 2015, 2016), the genomic allergen rapid detection (GARD) (Zeller et al., 2017), and the kDPRA (Waring et al., 2017). DA using data from NAMs have been shared for predicting potency, including a Bayesian network approach (Jaworska et al., 2015), regression models with KeratinoSens™ (KS) and peptide reactivity data (Natsch et al., 2015), artificial neural network model (Hirota et al., 2015, 2018), and integrated use of human cell line activation test (h-CLAT), DPRA and DEREK data (Takenouchi et al., 2015). It will be essential for risk assessors and regulators to evaluate these different approaches to identify which DA or individual methods provide an accurate point of departure (PoD) value for conducting sound risk assessments.

One promising approach based on generating linear regression models has been published using KS and kinetic peptide reactivity data to provide a predicted EC3 as a PoD (Natsch et al., 2015, 2018). Predicting an EC3 value offers the advantage of generating continuous potency values compared to predicting a chemical potency class (Cottrez et al., 2015; Jaworska et al., 2015; Zang et al., 2017; Zeller et al., 2017). It also provides the opportunity to manage uncertainty using statistical tools based on knowledge of the accuracy of the prediction. Such uncertainty could be factored in to refine the PoD value for conducting a skin sensitization risk assessment. The determination of potency has been primarily dependent on the use of the LLNA (Loveless et al., 2010; OECD, 2010), which has long been considered the “gold standard” for potency assessment because it yields quantitative data suitable for a dose-response evaluation. An alternative, non-animal approach is urgently needed.

The previous work on regression models (Natsch et al., 2015) used kinetic rate constants generated with the Cor1-C420 assay (Natsch and Gfeller, 2008). In this paper, updated linear regression models based on data from OECD validated methods (kDPRA, KS and/or h-CLAT) are described for predicting a PoD value for risk assessment purposes. A comprehensive database of 322 chemicals was assembled that contains data from previous papers (Natsch et al., 2015, 2020) merged with the database compiled within the framework of the OECD project on DA (OECD, 2021c). The paper addresses (1) contribution of the in vitro parameters for predicting LLNA potency, (2) comparison of prediction models based on an inclusive dataset versus the highly curated OECD dataset, (3) guidance in model selection with multiple input data, (4) comparison of kDPRA and Cor1-Cor420 reactivity data, and (5) case studies on chemicals with multiple LLNA EC3 values. This work further advances the 3Rs for skin sensitization testing as a standardized way to derive a PoD from validated methods is still a missing element in the application of NAM for sensitization assessment.

2 Materials and methods

Data sources of existing data
All in vivo and in vitro data are from our previous publication (Natsch et al., 2015), from data compilations by Urbisch et al. (2015) and Jaworska et al. (2015), and from the database compiled by the OECD working group on DA (OECD, 2021c). The database (ESM1-1) contains data from the DPRA and kDPRA (OECD TG 442C), KS assay (TG 442D), h-CLAT (TG 442E), LLNA (GL 429) and the Cor1-C420 reactivity assay (Natsch and Gfeller, 2008). All individual parameters are described in ESM1-11.

Data transformation and normalization for statistical analysis
All data were log-transformed and normalized as described (Natsch et al., 2015). In case multiple LLNA data were available, we took the geometric mean of the LLNA EC3 values. Data are expressed as pEC3, a logarithmic expression taking molecular weight into account since most in vitro data are expressed in molar concentrations and not on a per weight basis as in the LLNA.

\[ pEC3 = \log\left(\frac{MW}{EC3}\right) \]

For negative LLNA results, the pEC3 was set to zero, which is the special case for, e.g., molecules with a molecular weight of 100 and an EC3 of 100%. This approach treats all negatives equally, as not all negatives were tested up to 100%. Based on this normalization, the pEC3 spans a range from 0 for non-sensitizers to 4.86 for the potent sensitizer oxazolone.

In the KS assay, the IC50 value (concentration for 50% reduction in cellular viability) and the EC1.5 or EC3 values, i.e., concentration for 1.5- or 3-fold luciferase induction, are determined. Chemicals are tested up to a maximal concentration of 2000 μM. For chemicals with no cytotoxicity at this concentration, the numerical IC50 was set to the arbitrary value of 4000 μM. Similarly, if the luciferase gene was not induced above a given threshold, the EC1.5 or EC3 values were set to 4000 μM. In addition, in the KS prediction model only gene induction observed at non-cytotoxic levels (> 70% viability) is considered relevant. Thus, if 1.5-fold gene induction was only observed at cytotoxic levels,

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Abbreviations
DA, defined approach; DPRA, direct peptide reactivity assay; GHS, Global Harmonized System; h-CLAT, human cell line activation test; IATA, integrated approach to testing and assessment; kDPRA, kinetic direct peptide reactivity assay; KS, KeratinoSens™; LLNA, local lymph node assay; NAM, new approach methodologies; PoD, point of departure; OECD, Organisation for Economic Co-operation and Development; OECD DB, OECD reference database on defined approaches; TG, test guideline; VP, vapor pressure

1 doi:10.14579/altest.2201141s1
With these linear transformations, all in vitro parameters are set to 0 in case a molecule is non-reactive in kDPRA, non-toxic beyond the selected threshold in the cell-based assays, or it does not induce the cellular markers (luciferase or CD86/CD54 induction), while all positive in vitro evidence according to the prediction models of the individual assays has a positive value. These linear transformations of logarithmic data do not distort the data but make reading the regression equations easier.

Vapor pressure was calculated with TIMES SS software (Laboratory of Mathematical Chemistry). Vapor pressure is considered important as chemicals are tested in the LLNA with open application, while evaporation is limited in the in vitro assays. The relationship of calculated vapor pressure to evaporation from the LLNA vehicle acetone/olive oil 4:1 (AOO) at the typical skin surface temperature of 32°C was tested experimentally for 37 molecules, and a linear relationship between Log(VPcalc) and log half-life of the chemical in AOO was demonstrated as shown in Figure S2 in Natsch et al. (2015). Chemicals with a Log(VPcalc) < 10 Pa were found not to evaporate significantly from AOO within 60 min. Therefore, vapor pressure data were normalized as follows:

\[ \text{LogVP}_{\text{norm}} = \text{Log}(\text{VP}) - 1 \]

and values below zero were set to 0. With this approach, chemicals predicted to evaporate significantly from mouse ears within 60 min have a positive coefficient, whereas the coefficient for chemicals with low volatility (i.e., not significantly evaporating within 60 min) is set to 0.

Statistical analysis

Multiple linear regression against the pEC3 was performed with the MiniTab 15 software. The $R^2$-values and F- and $p$-values of the correlations are given throughout the manuscript as measures of correlation strengths. The regression equations were then used for predicting the behavior of new chemicals.
3 Results

3.1 A comprehensive database of in vitro and in vivo data

We compiled a comprehensive database on 322 chemicals (Tab. ESM1-1) with data from the OECD-validated LLNA, KS, h-CLAT, DPRA and kDPRAs. This database also contains physicochemical information and reaction rates measured using a method similar to the kDPRAs but with the more reactive peptide Cor1-C420, as reported earlier. This database is a merger of our previous database (Natsch et al., 2015) with the database used to validate the kDPRAs (Natsch et al., 2020) and the database put together within the frame of the OECD project on developing a DA for skin sensitization in the Series on Testing and Assessment No. 336, Annex 2 (OECD, 2021c). It also contains h-CLAT and DPRA data from Urbisch et al. (2015) and Jaworska et al. (2015). Table 1 indicates the different subsets of this database.

In the analysis below, we present regression analysis on these different datasets, and we provide a prediction spreadsheet (ESM2) integrating the key equations, which can be used to estimate a predicted LLNA pEC3 and an EC3 value as a PoD based on in vitro input parameters using the different regression equations.

3.2 A quantitative global regression model integrating kDPRAs data with KS data

The database contains kDPRAs and KS data for a set of 203 chemicals. For this dataset, a regression model based on the normalized EC1.5 and IC50 from KS, Log kmax from the kDPRAs and normalized vapor pressure was derived, similar to the global model in the 2015 study (EQ1), by replacing reaction rates with the Cor1-C420 peptide with data from the validated kDPRAs.

EQ1 pEC3 = 0.42 + 0.40 × Log kmax norm + 0.15 × Log EC1.5 norm + 0.36 × Log IC50 norm − 0.21 × Log VP norm

| Term                      | Coefficient | p-value |
|---------------------------|-------------|---------|
| Constant                  | T = 4.61, p < 0.0005 |
| Log kmax norm             | T = 9.1, p < 0.0005 |
| Log EC1.5 norm            | T = 2.5, p = 0.013 |
| Log IC50 norm             | T = 4.47, p < 0.0005 |
| Log VP norm               | T = -3.44, p = 0.001 |
| N                          | 203; S = 0.73; R² = 62.0%; R² (adj) = 61.3% |

In this equation, kmax from kDPRA and IC50 from KS have the highest statistical weight. This equation is derived from the most comprehensive dataset with available kDPRAs data. EQ1 is therefore proposed as the key model in the prediction spreadsheet (ESM2) for chemicals with available kDPRAs and KS data only.

For comparison, the same regression analysis was performed on the subset of data from the core set used in the 2015 study with kDPRAs data (n = 158; EQ2). Similarly, the same model was also calculated on the subset of data with available h-CLAT data. Each can then be the only (i.e., based on the same chemicals) compared to the models below integrating h-CLAT data (n = 188; EQ3).

EQ2 pEC3 = 0.34 + 0.38 × Log kmax norm + 0.16 × Log EC1.5 norm + 0.42 × Log IC50 norm − 0.18 × Log VP norm

n = 158; S = 0.72; R² = 63.0%; R² (adj) = 61.2%

EQ3 pEC3 = 0.40 + 0.40 × Log kmax norm + 0.14 × Log EC1.5 norm + 0.38 × Log IC50 norm − 0.21 × Log VP norm

n = 188; S = 0.75; R² = 58.4%; R² (adj) = 57.5%

The resulting equations are very similar to EQ1, indicating that the chemicals added to the core set used previously have no major impact on the overall predictive model and that the model is relatively stable when used with different subsets of the input data.

3.3 A quantitative global regression model integrating kDPRAs data with h-CLAT data

Next, a similar model was calculated using only data from the kDPRAs and h-CLAT to predict the potency of chemicals with these two sources of data available, thus normalized EC1.5 and IC50 were replaced with the normalized MIT and the CV75 from h-CLAT, both calculated in µM (n = 188; EQ4).

EQ4 pEC3 = 0.18 + 0.36 × Log kmax norm + 0.21 × Log MIT norm + 0.35 × Log CV75 norm − 0.19 × Log VP norm

| Term                      | Coefficient | p-value |
|---------------------------|-------------|---------|
| Constant                  | T = 1.65, p = 0.102 |
| Log kmax norm             | T = 7.59, p < 0.0005 |
| Log MIT norm              | T = 3.06, p = 0.004 |
| Log CV75 norm             | T = 3.69, p < 0.0005 |
| Log VP norm               | T = -3.0, p = 0.003 |
| N                          | 188; S = 0.72; R² = 61.1%; R² (adj) = 60.3% |

Compared to EQ3, on the same dataset, kmax has a similar weight in this equation, and MIT and CV75 can replace the KS data with a very similar overall statistical weight. EQ4 is thus added to the prediction spreadsheet (ESM2) to be used for those chemicals for which only kDPRAs and h-CLAT data are available.

3.4 A quantitative global regression model integrating kDPRAs data with KS and h-CLAT data

For some chemicals, comprehensive data from all three OECD TGs will be available. Thus, the most comprehensive model integrates kDPRAs, KS and h-CLAT data (n = 188; EQ5).
result in the kDPRA and the DPRA. For these and complex extracts or multi-constituent substances with data gaps in the kDPRA due to incompatibility with the assay, a model based on KS and h-CLAT data only may be of interest. This model is presented in Equation 6.

\[
\text{EQ6 } p\text{EC}_3 = 0.09 + 0.276 \times \log \text{MIT}_{\text{norm}} + 0.22 \times \log \text{EC}_{1.5}\n_{\text{norm}} + 0.34 \times \log \text{CV}_{75}\n_{\text{norm}} - 0.12 \times \log \text{VP}_{\text{norm}}
\]

\[
\text{Constant } T = 0.67, \ p = 0.50
\]

\[
\log \text{MIT}_{\text{norm}} T = 3.75, \ p < 0.0005
\]

\[
\log \text{EC}_{1.5}\n_{\text{norm}} T = 1.41, \ p = 0.159
\]

\[
\log \text{IC}_{50}\n_{\text{norm}} T = 0.94, \ p = 0.347
\]

\[
\log \text{CV}_{75}\n_{\text{norm}} T = 2.74, \ p = 0.004
\]

\[
\log \text{VP}_{\text{norm}} T = -2.84, \ p = 0.005
\]

\[n = 188; S = 0.72; R^2 = 61.6\%; R^2 (adj) = 60.6\%\]

The predictive power of EQ5 is only very marginally improved vs EQ3 and EQ4, indicating that with both KS and h-CLAT data, the overall predictivity improves only slightly as compared to the situation where only one cellular assay is available. This has already been observed in our previous comprehensive analysis (Natsch et al., 2020), indicating significant data redundancy between these two cell-based assays. This is especially the case for the cytotoxicity readout. When both IC50 and CV75 are used in EQ5, they contribute additively, but the coefficient and statistical parameters indicate a weak contribution of each parameter. Alternative models using either CV75 or IC50 alone have the same R² value (61%), but in both cases, the contribution of the cytotoxicity parameter has more statistical weight (p = 0.014 for IC50 and p = 0.005 for CV75; regression equations not shown), confirming data redundancy especially for the two cytotoxicity indicators, which is not a surprising finding.

3.5 A quantitative global regression model integrating KS and h-CLAT data in the absence of kDPRA data

Finally, in some cases, chemicals are rated as positive in a DA based on a positive result in the cell-based assays only despite a negative result in the kDPRA and the DPRA. For these and complex extracts or multi-constituent substances with data gaps in the kDPRA due to incompatibility with the assay, a model based on KS and h-CLAT data only may be of interest. This model is presented in Equation 6.

\[
\text{EQ6 } p\text{EC}_3 = 0.09 + 0.276 \times \log \text{MIT}_{\text{norm}} + 0.22 \times \log \text{EC}_{1.5}\n_{\text{norm}} + 0.34 \times \log \text{CV}_{75}\n_{\text{norm}} + 0.06 \times \log \text{IC}_{50}\n_{\text{norm}} - 0.12 \times \log \text{VP}_{\text{norm}}
\]

\[
\text{Constant } T = 0.67, \ p = 0.50
\]

\[
\log \text{EC}_{1.5}\n_{\text{norm}} T = 3.22, \ p = 0.002
\]

\[
\log \text{IC}_{50}\n_{\text{norm}} T = 0.43, \ p = 0.671
\]

\[
\log \text{MIT}_{\text{norm}} T = 3.75, \ p < 0.0005
\]

\[
\log \text{CV}_{75}\n_{\text{norm}} T = 2.29, \ p = 0.025
\]

\[
\log \text{VP}_{\text{norm}} T = -1.81, \ p = 0.072
\]

\[n = 188; S = 0.81; R^2 = 51.8\%; R^2 (adj) = 50.5\%

All the above models (EQ1, 4 and 5) were also calculated with KS EC3 derived from KS, i.e., the concentration for threefold stimulation of the luciferase signal in KS, instead of EC 1.5, and models with very similar R² were obtained (data not shown). However, in the situation without reactivity data, EQ7 using KS EC3 has improved statistical power, with KS EC3 having the highest statistical weight.

\[
\text{EQ7 } p\text{EC}_3 = 0.202 + 0.222 \times \log \text{MIT}_{\text{norm}} + 0.40 \times \log \text{EC}_{3}\n_{\text{norm}} + 0.313 \times \log \text{CV}_{75}\n_{\text{norm}} + 0.023 \times \log \text{IC}_{50}\n_{\text{norm}} - 0.151 \times \log \text{VP}_{\text{norm}}
\]

\[
\text{Constant } T = 1.67, \ p = 0.098
\]

\[
\log \text{EC}_{3}\n_{\text{norm}} T = 5.29, \ p < 0.0005
\]

\[
\log \text{IC}_{50}\n_{\text{norm}} T = 0.18, \ p = 0.855
\]

\[
\log \text{MIT}_{\text{norm}} T = 3.02, \ p = 0.003
\]

\[
\log \text{CV}_{75}\n_{\text{norm}} T = 2.28, \ p = 0.024
\]

\[
\log \text{VP}_{\text{norm}} T = -2.28, \ p = 0.023
\]

\[S = 0.77; R^2 = 55.9\%; R^2 (adj) = 54.7\%\]
a) Does the smaller data subset collected by the OECD group lead to significantly different equations compared to those presented above on a more comprehensive set? 
b) Do the LLNA EC3 values after the data curation lead to significantly different predictive equations compared to the LLNA data in the previously published databases? 
Table 3 provides comparisons of the model parameters as calculated for the chemicals in the OECD database only (n = 149) vs (i) published LLNA data and (ii) curated LLNA data, and these models are compared to the models established on the more comprehensive sets of chemicals (EQ1, EQ4 and EQ5). 
Overall, the additional models (EQ8-EQ13) derived from the subset in the OECD database are similar to EQ1, EQ4 and EQ5, integrating the larger datasets of all available data for a given data combination. Especially, only minor differences are observed between the models on the historical and the curated LLNA data (EQ8 vs 9 / EQ10 vs 11 / EQ12 vs 13).

3.8 Practical application: An open spreadsheet for calculating a PoD and case studies
To further facilitate application, here we provide a prediction spreadsheet as ESM2. Users can simply enter the experimental data from the different in vitro assays along with the molecular weight and a calculated vapor pressure to calculate a predicted EC3, which can be used as a PoD and, if applying adequate safety factors and taking into account all available information and applicability domain limitations, for a quantitative risk assessment. To illustrate its use, the prediction spreadsheet also contains the input data for two examples, DNCB and cinnamic aldehyde.

These models without kDPRA data have less statistical power, but they would primarily be used either in the case of data gaps for the kDPRA, for chemicals that are negative in the kDPRA, or for chemicals outside of the applicability domain of the kDPRA (see ESM3.4). These latter chemicals, in most cases, are weak or moderate sensitizers, as the majority of strong sensizers have a direct peptide reactivity in the kDPRA and DPRA (Natsch et al., 2020).
Here we show how different combinations of source data from OECD TG 442C, 442D and 442E can be used in different regression equations. Depending on the available data, a user is faced with two critical questions:

a) Once data from two positive tests are available, is it worth collecting data from the third test for a more accurate potency assessment?

b) If the available data allows calculating a predicted PoD with different equations, which result should be taken forward?

The detailed data analysis to answer these questions is presented in ESM34, and a summary is given here. If two positive results are available, including kDPRA data, studying the third key event, in general, is not needed because (i) if the third outcome were negative, the output of the two positive tests could be used directly with the corresponding equation EQ1 or EQ4, and if (ii) the

For all chemicals, additional spreadsheets in ESM1 (Tab. ESM1-21 – ESM2-4) give the individual predictions by the different equations on the different data subsets shown in Table 1. Here we show individual data on key case studies (Tab. 4). We selected all chemicals for which the OECD LLNA database provides n ≥ 5 individual EC3 values. Selecting these chemicals as case studies has two advantages: (i) The overall weight of evidence for the LLNA EC3 is strong as it comes from multiple independent tests and (ii) for these chemicals, we have a good understanding of the variability of the target, i.e., the range of LLNA EC3 measured, and thus the predictions can also be compared to this intrinsic variability of the prediction target.

For most of the 16 case studies, the predicted EC3 values fall within the experimental variability of the LLNA or are close to this range. In general, the predictions based on kDPRA and either cell-based assay are quite close for these chemicals.

### Tab. 4: Case studies: Predictions by the different models for chemicals with multiple LLNA EC3 (n ≥ 5) values in the OECD curated database

| OECD MLLP LLNA EC3a | LLNA EC3 measured rangeb | LLNA EC3 statistical rangec | Pred. EC3 EQ1d | Pred. EC3 EQ4e | Pred. EC3 EQ5f | Pred. EC3 EQ13g | Pred. global model 2015i |
|---------------------|------------------------|---------------------------|--------------|---------------|---------------|----------------|------------------------|
| Aniline             | NC                     | 13.25 - (> 100)           | NC           | 60            | 52            | 57             | 52 > 100              |
| Penicillin G        | 31.3                    | 11.2 - 46.5               | 15.2 - 47.2  | > 100         | > 100         | > 100 > 100     |
| Hydroxyceptralinall | 21.1                    | 18.8 - 33                 | 18.5 - 27.2  | 18.7          | 11.3          | 10.9 8.0 3.5    |
| Geraniol            | 16.1                    | 5.6 - 57                  | 7.6 - 37.5   | 18.3          | 14.3          | 14.2 10.3 5.1    |
| Eugenol             | 11.6                    | 3.8 - 16.6                | 5.8 - 15.4   | 19.9          | 6.8           | 10.4 10.2 14.1   |
| Alpha-hexyl cinnamatic aldehyde | 10.8    | 1.2 - 33.8                | 4.9 - 17.4   | 5.9           | (25)h         | 17.4 15.0 1.8    |
| Lilial              | 8.6                     | 3 - 18.6                  | 3.6 - 16.8   | 20.5          | 9.3           | 12.5 13.9 6.2    |
| Citral              | 5.8                     | 1.5 - 26.8                | 3 - 12.7     | 9.4           | 5.0           | 4.8 3.5 12.6     |
| Formaldehyde        | 3.8                     | 0.35 - 14.5               | 0.6 - 9.6    | 1.5           | 0.8           | 1.0 2.0 4        |
| 3-dimethylaminopropylamine | 3.5   | 1.8 - (> 10)              | 1.9 - 5.7    | 40            | 37            | 32 27 53        |
| Isoeugenol          | 1.3                     | 0.5 - 6.4                 | 0.8 - 3      | 1.8           | (4.6)h         | 4.2 4.1 1.6      |
| Cinnamic aldehyde   | 1                       | 0.5 - 3.1                 | 0.7 - 1.8    | 1.0           | 0.8           | 0.8 0.9 1.1     |
| Hydroquinone        | 0.19                    | 0.07 - 1.6                | 0.1 - 0.5    | 0.9           | 0.4           | 0.4 0.4 0.8     |
| PPD                 | 0.11                    | 0.06 - 0.2                | 0.07 - 0.17  | 3.5           | 1.9           | 1.7 1.2 0.72    |
| DNCB                | 0.054                   | 0.02 - 0.096              | 0.02 - 0.08  | 0.18          | 0.19          | 0.17 0.24 0.21  |
| Kathon CG           | 0.008                   | 0.0049 - 0.063            | 0.004 - 0.035| 0.05          | 0.05          | 0.05 0.08 n.d.   |
| Oxazolone           | 0.002                   | 0.0011 - 0.0026           | 0.001 - 0.003| 1.5           | 0.5           | 0.7 0.9 0.9     |

a) LLNA EC3, median-like location parameter derived by OECD group from multiple EC3 values; b) range between minimal and maximal measured LLNA EC3 in OECD DB; c) geometric mean and geometric standard deviation were calculated, range around the geometric mean defined by the geometric standard deviation is given; d) based on KS and kDPRA; e) based on h-CLAT and kDPRA; f) based on KS, h-CLAT and kDPRA; trained on comprehensive database (n = 188) and historical LLNA database; g) based on KS, h-CLAT and kDPRA; trained on curated OECD LLNA data only (n = 149); EQ13: pEC3 = 0.26 + 0.25 × Log kmax norm + 0.19 × Log MITnorm + 0.17 × EC1.5norm + 0.27 × Log CV75norm - 0.02 × Log IC50norm - 0.19 × Log VPnorm; h) values in brackets: chemicals negative in h-CLAT, EQ4 is not recommended for chemicals negative in h-CLAT; i) predictive model using the Cor1-C420 assay published previously.

### Model selection with multiple input data

Here we show how different combinations of source data from OECD TG 442C, 442D and 442E can be used in different regression equations. Depending on the available data, a user is faced with two critical questions:

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The detailed data analysis to answer these questions is presented in ESM34, and a summary is given here. If two positive results are available, including kDPRA data, studying the third key event, in general, is not needed because (i) if the third outcome were negative, the output of the two positive tests could be used directly with the corresponding equation EQ1 or EQ4, and if (ii) the

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b) If the available data allows calculating a predicted PoD with different equations, which result should be taken forward?

The detailed data analysis to answer these questions is presented in ESM34, and a summary is given here. If two positive results are available, including kDPRA data, studying the third key event, in general, is not needed because (i) if the third outcome were negative, the output of the two positive tests could be used directly with the corresponding equation EQ1 or EQ4, and if (ii) the
third outcome were positive, either equation EQ1, EQ4 or EQ5 could then be used, but these would lead to very similar predictions. Thus, for the 73 chemicals with three positive tests, the correlation between the predicted pEC3 from EQ1 (KS + kDPRA) and EQ5 (KS, h-CLAT and kDPRA) has an R² value of 0.95 with a slope close to 1 and y-intercept close to zero. Similarly, the correlation between EQ4 (h-CLAT and kDPRA) and EQ5 has an R² value of 0.99. Thus, adding the third test and moving from EQ1 or EQ4 to EQ5 has little effect on the final prediction. There would also not be a significant difference, whether the testing started with either 442D or 442E, as there is also a high correlation (R² value of 0.91 with a slope close to 1 and y-intercept close to zero) between the predicted pEC3 from EQ1 (KS and kDPRA) and EQ4 (h-CLAT and kDPRA) for chemicals positive in all assays. This relatively close alignment of EQ1, EQ4 and EQ5 for chemicals positive in all three tests is also illustrated by the case studies in Table 4. If all three tests are positive and data available, then using EQ5 appears most appropriate. A conservative alternative would be using the lowest EC3 value from EQ1, EQ4 and EQ5 to perform a more stringent risk assessment (see ESM3⁴).

A different question is what to do when all three tests are available and one of them is negative. Then the choice is to (i) use the model with the data from the two positive tests only or (ii) use EQ5 (KS, h-CLAT and kDPRA). Theoretically, the former choice is the more conservative one, as negative evidence is not factored in, while the latter uses all available data. Interestingly, there is also a high correlation between the predicted pEC3 for both choices (R² value of 0.92 with a slope close to 1) (n = 43 chemicals with 2 positive tests) and the correlation with the in vivo data is better for the latter option (see ESM3⁴). Thus, based on this analysis, it appears that using EQ5 is optimal in all cases when all data are available (2 or 3 positive tests). However, further expert-guided analysis can lead to different choices in selected cases, e.g., in cases of putative pro- or pre-haptens, which may be better predicted by the cellular assays and best predicted by EQ6.

3.10 Comparison of the reaction rate measured with the kDPRA and the Cor1-C420 assay used previously

Our previous study (Natsch et al., 2015) was entirely based on Cor1-C420 peptide reactivity data. In a final analysis, we compared data from the two peptide reactivity assays and regression models developed with the two reactivity assays. This helps to understand assay redundancy and similarity between the reactivity assays and between the current and the previous model, which provides information on the robustness of the approach proposed here and an indication of how the two peptide assays can be used interchangeably. This analysis is detailed in ESM4⁵, and here we give a brief summary of the results.

A linear correlation is observed when comparing the reaction constants from kDPRA and the Cor1-C420 assay. Due to the higher reactivity of the Cor1-C420 peptide as compared to the Cys-peptide (Natsch et al., 2007), the correlation has a negative y-intercept:

\[ \text{EQ14:} \quad \log k_{\text{max}} (\text{kDPRA}) = 0.9 \times \log k_{\text{max}} (\text{Cor1-C420 assay}) - 0.59 \]

Thus, based on the measured Cor1-C420 data, we calculated predicted kDPRA values based on EQ14 and used these data to train the same regression model on the same chemicals as in EQ1 (EQ15).

\[ \text{EQ15:} \quad \text{pEC3} = 0.40 + 0.49 \times \log k_{\text{max, norm}} (\text{calculated from Cor1-420 result}) + 0.12 \times \log \text{EC1.5}_{\text{norm}} + 0.28 \times \log \text{IC50}_{\text{norm}} - 0.22 \times \log V_P_{\text{norm}} \]

\[ S = \text{0.73; R}^2 = 61.5\%; \text{R}^2 (\text{adj}) = 60.7\% \]

This equation has a similar predictive capacity as EQ1 (calculated on the same dataset) and overall a similar contribution of the individual parameters, although contribution of the reactivity parameter is slightly higher and the contribution of KS slightly weaker in EQ15 vs EQ1. The reactivity rates for the Cor1-C420 assay presented in the full database are all presented as measured values and also transformed to predicted kDPRA k_max values according to EQ14. In another data column, data for measured kDPRA k_max are shown and data gaps were filled with the predicted kDPRA k_max values according to EQ14. Due to data redundancy between the two tests, these data can be used in the absence of a true kDPRA experimental value for any further modeling on the larger database, e.g., to perform domain-based assessments or quantitative read-across assessment based on data from the target and the read-across substance using all the chemicals in the full database (For details see ESM4²).

4 Discussion

This paper presents global regression models generated from different NAM datasets for producing a PoD value for risk assessment purposes. The models are based on the use of OECD-validated NAMs that include the recently accepted kDPRA (OECD, 2021d) as well as the KS and h-CLAT (OECD, 2018a,b). A comprehensive database of 322 chemicals was assembled for this analysis (Natsch et al., 2015, 2020; OECD, 2021c) that may also be useful for further refinements or the development of alternative models.

4.1 Robustness of the models when compared to previous models

Previous work describing the development of regression models using NAM data used a more reactive peptide Cor1-C420 (Natsch and Gfeller, 2008) for assessing reactivity (Natsch et al., 2015, 2018). With the update of OECD TG 442C (OECD, 2021d) that now includes the kDPRA, it made sense to evaluate this equation generated (EQ15) has the same predictive capacity as EQ1 and, overall, a similar contribution of the individual parameters, indicating
similar information content of the two reactivity assays and a robustness of the overall approach.

The regression model using a comprehensive dataset with available kDPRA and KS data (n = 203) was based on the normalized EC1.5 and IC50 from KS, Log kmax from the kDPRA, and normalized vapor pressure. This model (EQ1) is based on the highest number of chemicals compared to other models and is very similar to the global model published previously (Natsch et al., 2015), but replaces Cor1-C420 reactivity data with kDPRA data.

4.2 Complementary and redundancy of the cell-based assays
A vital feature described in the paper is the flexibility in which input NAM data is used to make a PoD prediction. A model integrating kDPRA data with h-CLAT instead of KS showed a similar statistical weight using normalized MIT and CV75 data (n = 188; EQ4). This shows that either KS or h-CLAT data can be used to obtain comparable predictions. It is well known that having data from all three assays is not always possible due to assay compatibility factors (Kolle et al., 2019). However, in cases where data from all three OECD TG data is available, a comprehensive model can be obtained that integrates kDPRA, KS and h-CLAT data (n = 188; EQ5).

Interestingly, having data from both KS and h-CLAT along with kDPRA data only slightly improves the predictive power compared to one cellular assay. This redundancy has been observed previously (Natsch et al., 2020) and indicates that there is significant redundancy between the two cell-based assays. For situations where kDPRA data is absent, possibly due to an incompatibility issue using complex mixtures or technical issues (e.g., thiols), a regression model integrating KS and h-CLAT data is available (n = 188; EQ6), too. In the absence of reactivity data, the parameters for luciferase induction in KS or surface marker induction in h-CLAT receive more statistical weight. The best statistical power is obtained using KS EC3 instead of EC1.5, suggesting that this parameter for strong Nrf2-dependent luciferase induction can partially compensate for the lack of reactivity data (EQ7).

For each model based on a dataset of 188 chemicals with data from KS, h-CLAT or kDPRA, their predictive capacity is similar. For all models, the median of the fold-misprediction is around 2.5-fold, while the geometric mean is around 3.3-fold (Tab. 2). This finding demonstrates the flexibility in the utility of using these models based on what test data is available. A description of how to select an appropriate model is provided in ESM3^2, and a parallel paper describes how the models can be combined with the “2 out of 3” DA (Natsch and Gerberick, 2022). The analysis shows that for chemicals with three positive tests, adding the third test has little effect on the final prediction. Thus, a third key event study is not needed if two positive tests, including kDPRA data, are available. There is also no significant difference whether one starts with the kDPRA or either of the two cell-based assays. When multiple PoD values are available, the most predictive choice would be to use EQ5, integrating all three assays, although a different choice might be made when the chemical is a putative pro- or pre-hapten, in which case EQ6 might be optimal.

4.3 Assessment of using the comprehensive database vs the curated OECD DA database
Very similar models to EQ1 were obtained using the core set of data from Natsch et al. (2015) for which kDPRA data are available (n = 158; EQ2) and the set of chemicals with available h-CLAT data (n = 188; EQ3), indicating the model is robust and not highly dependent on the training set used. A curated database developed by an OECD expert group has been made publicly available (OECD, 2021c), which contains 196 chemicals, of which 154 have LLNA data and 149 are available with NAM data in our database. The regression coefficients for the models described (EQ1, EQ4, EQ5) are very similar to equivalent models derived from the OECD curated subset (Tab. 3). However, the contribution of reactivity from kDPRA tends to be lower. This may be attributed to the fact that the OECD group had stringent criteria to allow for extrapolation of LLNA results (OECD, 2021b). This led to the exclusion of LLNA data for several strong and extreme sensitizers, which are highly reactive chemicals and for which the reaction rate has a high weight in the assessment. On the other hand, the KS EC1.5 has a higher weight on this selected subset. When, for the same chemical set, the LLNA EC3 from previously published databases is used as the target as compared to the curated EC3 values (e.g., EQ8 vs EQ9), there is only a very small difference in the resulting regression equation, indicating that the data curation had no major impact on the overall picture. In addition, models established on a large number of chemicals are not inferior to the models built upon the OECD curated data (see results for EQ13 and EQ5 in Tab. 4), although the data curation certainly makes a significant difference for some individual chemicals. Going forward, it therefore appears appropriate to use models incorporating the maximal weight of evidence from the largest databases publicly available rather than to base models only on the curated OECD data.

4.4 Transparency of the regression models
As compared to complex tools like neural networks or Bayesian nets, linear regression models may appear to be relatively simple statistical tools. However, they have the advantage of high transparency. The prediction spreadsheet (ESM2^3) is designed to allow one to enter experimental data from the different NAMs along with molecular weight and a calculated vapor pressure to calculate a PoD value. This should facilitate practical application.

4.5 Key misprediction
For most of the 16 case studies, it is evident that the PoD values, based on using kDPRA and one or both cellular assays, are quite similar to the LLNA values based on ≥ 5 individual EC3 values. Clear outliers under-predicted by the models are 3-dimethylaminopropylamine and p-phenylenediamine (see discussion on applicability domain in ESM3^2). The extreme sensitizer oxazolone is predicted as a strong sensitizer by the models, but the predicted PoD value is clearly underestimated. The unique reactivity of oxazolone with lysine residues, explaining its unsurpassed sensitization potential, has been investigated before and is not captured by the kDPRA (Natsch et al., 2010). A more detailed discussion
on mispredicted chemicals is provided in a parallel paper (Natsch and Gerberick, 2022).

4.6 Outlook
For risk assessors responsible for assessing the skin sensitization risk of new chemical entities or chemicals lacking sufficient data, it is critical to have tools available that are predominantly dependent on using NAM data. The PoD value obtained from these regression models can thus be used to assist in the conduct of skin sensitization risk assessments. The predictive regression models (EQ1, 4, 5, 6 and 7 as an alternate) have been built from a comprehensive database of 188-203 chemicals along with a comparative analysis using the curated OECD database (OECD, 2021c). They show similar statistical strength and prediction accuracy. Performance is similar when using different data input parameters or when comparing models generated from different datasets. The PoD derived from these models may, using appropriate assessment factors to account for uncertainty and taking all information into account, be used as a starting point to determine safe use levels in products. The application and guidance on how to use these regression models when using the “2 out 3” DA is covered in a separate paper (Natsch and Gerberick, 2022).

Electronic supplementary material
ESM1 contains the full database in Sheet 1 and the predictions by the different models for the individual chemicals for different data subsets and vs different LLNA datasets in Sheets 2-4.

ESM2 is the prediction spreadsheet to calculate the PoD from in vitro data.

ESM3 discusses choice of the different regression models based on test availability.

ESM4 compares predictivity of the different reactivity assays.

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Conflict of interest
The authors declare no competing interests.

Data availability
All data of this publication are made publicly available and all models and tests used are freely available.