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Report on the European Partnership for Alternative Approaches to Animal Testing (EPAA) "New Approach Methodology (NAMs) User Forum", 30 – 31 October 2024, Helsinki, Finland

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#### **Article**

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Report on the European Partnership for Alternative Approaches to Animal Testing (EPAA) "New Approach Methodology (NAMs) User Forum", 30 – 31 October 2024, Helsinki, Finland

Mark T.D. Cronin, Sophie Cable, Christian Desaintes, Sylvia E. Escher, Ellen Hessel, Ellen Fritsche, Petra Kern, Gavin Maxwell, Gladys Ouedraogo, Tomasz Sobanski, Matthias Wehr, Andrew White

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### **Abstract**

The European Partnership for Alternative Approaches to Animal Testing (EPAA) held the "New Approach Methodology (NAMs) User Forum" at the European Chemicals Agency, Helsinki, Finland on 30 – 31 October 2024. The User Forum brought together stakeholders from regulatory agencies, industry, non-governmental organisations (NGOs) and academia, as well as European Union competent authorities. Lessons learned from applying NAMs for regulatory use were provided by the European Food Safety Authority (EFSA) and European Chemicals Agency (ECHA). Progress in the development of the developmental and neurotoxicity in vitro battery (DNT IVB) and Alternative Safety Profiling Algorithm (ASPA) were described, as well as five case studies describing uses of NAMs for chemical safety assessment. The presentations confirmed progress in NAMs and, in particular, the value of tiered testing strategies to bring together different lines of evidence. Specifically, tiered testing strategies for non-animal information are organised into three tiers, which may be relevant to hazard, exposure and toxicokinetic information. Progress into, and the needs for improvement of, the tiered strategies were discussed with a particular focus on the types of NAMs (in silico and in vitro) that may be required at each tier and the how confidence may be assigned to making a decision.

- **Keywords**: New Approach Methodology (NAM); Next Generation Risk Assessment (NGRA); tiered
- 46 testing strategy; chemical safety assessment; regulatory application

48	High	lights
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49	•	NAMs described for hazard identification and exposure assessment
50	•	Updates on application of NAMs from EFSA and ECHA
51	•	Tiered testing strategies can assist in the regulatory implementation of NAMs

• Case studies demonstrate the applicability of NAMs

Learnings and needs for NAMs' development identified

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# **Abbreviations**

56	ADME	Absorption, Distribution, Metabolism and Excretion
57	AOP	Adverse Outcome Pathway
58	APCRA	Accelerating the Pace of Chemical Risk Assessment
59	ASPA	Alternative Safety Profiling Algorithm
60	BER	Bioactivity-Exposure Ratio
61	CEP	Chemical Effect Predictor
62	Cmax	Maximum Concentration in Plasma
63	CRO	Clinical Research Organisation
64	CS	Case Study
65	DA	Defined Approach
66	DART	Developmental and Reproductive Toxicity
67	DEG	Diethylene Glycol
68	DNT	Developmental Neurotoxicity
69	DNT IVB	Developmental Neurotoxicity in vitro Battery
70	ECHA	European Chemicals Agency
71	EFSA	European Food Safety Authority
72	EPAA	European Partnership for Alternative Approaches to Animal Testing
73	EU	European Union
74	IATA	Integrated Approaches to Testing and Assessment
75	ISTNET	International STakeholder NETwork
76	IVIVE	In vitro-In vivo Extrapolation
77	KIC	Knowledge and Innovation Community
78	LOAEL	Lowest Observed Adverse Effect Level
79	MMP	Matched Molecular Pair
80	MOIE	Margin of Internal Exposure
81	MOS	Margin of Safety
82	NAM	New Approach Methodology
83	NGRA	Next Generation Risk Assessment
84	NGO	Non-Governmental Organisation
85	OECD	Organisation for Economic Cooperation and Development
86	PBK	Physiologically-Based Kinetic
87	PoD	Point of Departure
88	QAF	QSAR Assessment Framework
89	(Q)SAR	(Quantitative) Structure-Activity Relationship
90	RA	Retinoic Acid
91	qSIM	Quantifying Suitability of Analogues
92	SB	Sodium Benzoate
93	SCCS	Scientific Committee on Consumer Safety
94	STOT-RE	Specific Target Organ Toxicity - Repeated Exposure
95	TTC	Threshold of Toxicological Concern
96	VPA	Valproic Acid

#### 1. Introduction and Aims to the Workshop

This report summarises the presentations from, and the main findings of, the European Partnership for Alternative Approaches to Animal Testing's (EPAA's) "New Approach Methodology (NAMs) User Forum". The workshop was a hybrid event held at the European Chemicals Agency (ECHA) in Helsinki, Finland and on-line over two days (30 - 31 October 2024). It was attended by approximately 50 participants representing regulatory agencies, industry, non-governmental organisations (NGOs) and academia, as well as European Union (EU) competent authorities.

The aim of the User Forum was to explore further and share experiences with the use of New Approach Methodologies (NAMs) in chemical safety assessment. The particular focus of the meeting was the ability to make decisions with regard to chemical safety assessment from NAMs' data. This was mostly in the context of the use of NAMs as part of tiered testing strategies. This User Forum followed on the User Forum Kick-Off Workshop held 7-8 December 2023 (Cronin et al., 2025a). With regard to definitions of NAMs in the User Forum, a similar context can be applied as with the 2023 User Forum, where it was stated "NAMs were considered in a broad sense to include in silico, in chemico and in vitro approaches, -omics approaches or omic-enhanced in vivo studies combined as Defined Approaches (DAs) and/or Integrated Approaches to Testing and Assessment (IATA)" (Cronin et al., 2025a). With regard to the 2024 User Forum, tiered testing strategies were discussed more than IATA.

The purpose of this workshop report to summarise the presentations and case studies (Section 2) and key learnings from the presentations and discussion (Section 3). It is not intended to provide detailed minutes of the User Forum.

## 2. Summary of the Presentations and Case Studies at the User Forum

The User Forum heard a number of oral presentations (in person and hybrid). Section 2 summarises the content and main findings from the presentations, Section 2.1 is a summary of updates from the European Food Safety Authority (EFSA) and ECHA, Section 2.2 is a summary of two on-going initiatives, Section 2.3 summarises the case studies presented.

- 2.1 Updates from the European Food Safety Authority and European Chemicals Agency
- 2.1.1 European Food Safety Authority (EFSA): Lessons from Applying NAMs for Regulatory

**Use** 

127 An overview of the lessons learned from the application of NAMs from EFSA was given by Dr Sofia Batista Leite (EFSA). It was noted that EFSA works within many legal frameworks on EU Food Law, 128 129 which require different information requirements. The EFSA Strategy 2027 (EFSA, 2021b) highlights 130 EFSA's commitment to the development and integration of new scientific developments focusing on 131 NAM-based methods. To achieve their commitment, EFSA has published a road map for action on 132 NAMs and risk assessment (Escher et al., 2022) that helped with the prioritisation of EFSA's projects 133 on NAMs. In order to assist the harmonisation of approaches to the different legislations, Knowledge and 134 Innovation Communities (KICs) have been initiated. The KICs are intended to be dynamic knowledge 135 136 sharing and generating platforms, which aggregate information and discussion. One KIC focusses on 137 NAMs, the aim of which is to harmonise activities and identify stakeholders in NAMs. The KIC on NAMs 138 is also mapping the on-going activities in Europe to allow for aggregation of activities such as working 139 groups and the development of guidance and new tools. Currently, the EFSA funded projects on NAMs 140 can be grouped in four areas: cutting edge development and implementation; advancing 141 methodologies for toxicokinetics and toxicodynamics; protein safety assessment; and hazard 142 identification and characterisation. Two on-going EFSA-funded projects were described. The Developmental Neurotoxicity in vitro Battery (DNT IVB) is described in detail in Section 2.2.1 and its 143 use illustrated in Case Study 5. The EFSA NAMS4NANO Project aims to integrate NAMs chemical risk 144 145 assessments utilising information from case studies addressing nanoscale considerations. The work is 146 organised in 3 lots: i) the development of a qualification system for NAMs; ii) the development of 147 NAM-based case studies to fill data gaps in nanomaterial risk assessment; and iii) case studies to 148 improve methodology. 149 At the time of the meeting, an interim report had been published providing an initial proposal for a 150 "qualification system" for NAMs in food and the food sector, using nanomaterial risk assessment as example (Haase et al., 2024). EFSA recognise the implementation of approaches for nanoparticles risk 151 152 assessment is urgent. NAMs are seen to play a vital role in the risk assessment of nanoparticles and offer a unique opportunity to fill data gaps and address toxicity. Qualification is viewed as a promising 153 154 tool to assist the regulatory implementation of NAMs. 155 EFSA is contributing to the European Commission's roadmap for phasing out animal testing and 156 chemical safety assessments (Cronin et al., 2025b). It is acknowledged that its implementation into 157 the different legislations would be different as some follow data requirements that include animal testing (e.g., pesticides) whilst others do not (e.g., novel foods). Even if animal methods are still listed 158 159 in the respective guidance, EFSA's guidance is straightforward to update with new recommendations. 160 EFSA has identified a number of short-term actions that can support this work: phasing out of the use

161	of animal studies that have shown redundancy or lack of relevant information (ongoing work regarding
162	the use of dog for agrochemical risk assessment); better use of NAMs for absorption, distribution,
163	metabolism and excretion (ADME) assessment; and to encourage advocacy and guidance of NAMs in
164	EFSA panels period.
165	In summary, EFSA has a number of commitments to NAMs including the avoidance of redundant
166	animal studies; increasing the acceptance and confidence in the use of NAMs; development of
167	strategies to speed up the acceptance of NAMs; and to collaborate with key partners. It also provides
168	a number of resources including its journal ( <a href="https://www.efsa.europa.eu/en/publications/corporate">https://www.efsa.europa.eu/en/publications/corporate</a> ),
169	and databases ( <a href="https://www.efsa.europa.eu/en/applications/pesticides">https://www.efsa.europa.eu/en/applications/pesticides</a> ) and
170	$(\underline{https://www.efsa.europa.eu/en/data-report/chemical-hazardsdatabase-openfoodtox}).$
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172	2.1.2 European Chemicals Agency (ECHA): Experience in Developing and Applying NAMs for
173	Regulatory Use
174	An update on the experience of developing and applying NAMs for regulatory purposes was provided
175	by Dr Tomasz Sobanski (ECHA). A number of challenges to the regulatory acceptance of NAMs were
176	outlined. This includes the limitations of the current regulatory frameworks which may not yet
177	incorporate the new methods. This means there is still a heavy reliance on in vivo testing, whilst there
178	is policy and societal pressure for animal-free testing. There is also a need to build capacity in a number

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A three-step process for the use of NAMs for animal-free hazard assessment was described. Step 1 is to identify and address critical needs to enable the use of NAMs. It is essential to demonstrate that NAMs have applicability for a particular purpose. Firstly, NAM batteries must be demonstrated to be efficient for hazard identification for a given regulatory endpoint. Secondly, NAMs' ability to characterise hazard based on molecular and or cellular changes as opposed to the currently used observed adversity at a higher level is required. Thirdly, there is a need for reliable extrapolation to convert doses tested in the NAMs to the external equivalent dose or exposure. Existing *in vitro-in vivo* extrapolation (IVIVE) approaches are currently an area of high uncertainty and more reliable approaches are required.

of areas of the use of NAMs as well as developing them further in emerging topics, e.g., polymers,

nanomaterials, endocrine disruption, immunotoxicity and neurotoxicity, amongst others.

Step 2 is to demonstrate and apply NAMs under the current regulatory systems to build experience and gain confidence. ECHA is currently focusing efforts in a number of areas where there is a significant potential for reduction of animal use. i) Wider application of *in silico* approaches such as (quantitative)

structure-activity relationships ((Q)SARs) for less complex endpoints. The QSAR Assessment 193 194 Framework (QAF), recently released by the Organisation for Economic Cooperation and Development 195 (OECD), enables the evaluation of individual predictions for regulatory acceptance and will lead to 196 broader acceptance of QSARs. ii) Improving the use of read-across and the better integration of NAMs 197 such as -omics as bridging evidence. iii) Establishing robust protocols for Physiologically-Based Kinetic 198 (PBK) and toxicokinetic in vitro measurements and modelling, with a better understanding how to 199 optimise them to cover broad chemical space. iv) The better integration of -omics data in regulatory 200 methods and gaining confidence in their use. 201 Step 3 is to consider the requirements for a new regulatory framework that incorporates NAMs. This 202 includes the fact that a new framework may not rely on the same endpoints as currently used; gaining 203 knowledge in how to derive Points of Departure (PoDs) from molecular data; calibration of NAM 204 assays and data with well-defined protection goals; revision of Classification, Labelling and Packaging 205 (CLP) criteria to comply with NAM data; performance, throughput and cost from a business perspective; and improving the validation system for in vitro tests. It is appreciated that 206 207 communication is a key aspect to the implementation of NAMs, ECHA publishes an annual report on 208 key areas of regulatory challenge (ECHA, 2024). 209 ECHA is supporting a number of projects relating to the use of NAMs for regulatory purposes. In 210 addition to those noted above, there are efforts to encourage the sharing of data and knowledge 211 including the evolution of IUCLID. ECHA supports several of these initiatives to develop NAM-based 212 tools for hazard identification and characterisation through external contracts. ECHA is also an active 213 partner in the initiatives associated with the Accelerating the Pace of Chemical Risk Assessment 214 (APCRA). The APCRA case studies have demonstrated that NAMs can be used for conservative priority 215 setting (Paul Friedman et al., 2020) as well as investigating the integration of NAMs assays for the 216 assessment of data poor chemicals (Paul Friedman et al., 2025). The APCRA case study has demonstrated that PoDs from NAMs are not predictive of in vivo endpoints but may provide an 217 218 empirical PoD indication for data poor substances which could be used alongside other techniques 219 such as the threshold of toxicological concern (TTC) and QSAR. Other research by APCRA partners has 220 demonstrated that there may be considerable uncertainty in exposure estimates which are required 221 for Bioactivity-Exposure Ratio (BER) calculation. 222 ECHA concluded by summarising the lessons that have been learned in their investigation of the use 223 of NAMs. These are discussed in more detail in the context of the whole User Forum in Section 3, but include appreciation that one-to-one replacement of in vivo tests will not be possible, solutions will 224 225 be based around a combination of data-driven and knowledge driven approaches, the new

approaches must demonstrate performance within the remit of realistic expectations, and for

systemic toxicity it is essential to include toxicokinetics and metabolic activation, with the understanding that for industrial chemicals the current uncertainties associated with toxicokinetics are high.

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#### 2.2 On-Going Initiatives in NAMs

Invited presentations were made regarding two approaches to developing and implementing NAMs and tiered testing strategies.

#### 2.2.1 The Developmental Neurotoxicity *In Vitro* Battery (DNT IVB))

An in vitro battery for developmental neurotoxicity (DNT) was described by Prof Ellen Fritsche (SCAHT - Swiss Centre for Applied Human Toxicology and DNTOX GmbH). The growth in neurodevelopmental disorders is recognised and prioritised internationally, however, around 200 chemicals, mostly pesticides, have been tested for DNT. Current in vivo testing (OECD TG426 and TG443) is resource intensive with known uncertainties (Paparella et al., 2020). There is therefore an incentive for further DNT testing of chemicals and specifically the regulatory uptake of NAMs focusing on fit-for-purpose methods with high throughput and human relevance. Since 2005 there has been much effort in preparing acceptable NAMs for DNT (Smirnova et al., 2024). A particular turning point was a workshop which formulated an International STakeholder NETwork (ISTNET) to create a DNT in vitro testing road map (Bal-Price et al., 2015). The ISTNET brought together relevant stakeholders to agree how to move the tests forward as well as formulating the biology that control the development of the human brain. The overarching processes of human brain development that, if perturbed, may result in an adverse outcome were identified allowing for a battery of eight endpoints covered by 17 assays to be defined - the so-called DNT in vitro battery (DNT IVB) (Aschner et al., 2017; Fritsche et al., 2018 Masjosthusmann et al., 2020). On-going case studies are assisting in the understanding of the confidence and applicability of the DNT IVB. The initial findings of the case studies and recommendations for guidance and interpretation of the information from them have been published by the OECD (OECD, 2023). The development of the assays within the DNT IVB requires demonstration of scientific validity to gain confidence in their biological relevance and predictivity, with an example being Koch et al. (2022). Performance was assessed against reference chemicals to determine sensitivity and specificity (Carstens et al., 2022; Blum et al., 2023). An important aspect to make the DNT IVB usable has been to ensure lab-to-lab transfer. To enable this, the DNT IVB is currently being transferred to a contract research organisation (CRO) "DNTOX" (www.dntox.de). Transferring assays to a CRO is an important process to demonstrate transferability and make the assays available at the same time. NAM availability through CROs is an important step on the path to their regulatory acceptance (Blum et al. 2025). An example of an IATA case study utilising the DNT IVB was performed by EFSA for the re-evaluation of the pesticide deltamethrin applying Adverse Outcome Pathway (AOP)-based knowledge to demonstrate altered oligodendrocyte differentiation and neuronal network function (EFSA, 2021a). There are a number of on-going activities to gain more confidence in the DNT IVB, namely further compound testing to optimise the battery, assay refinement and development, and further AOP/ IATA development. The DNT IVB has also been considered in the context of endocrine disruption with the possibility to extend it to include other nuclear receptor-guided pathways beyond thyroid hormone disruption (Koch et al., 2025).

The development of the DNT IVB demonstrates the lifecycle for sustainable regulatory application of NAMs. This starts with the available test systems, a roadmap that has consensus from different stakeholders on how to move forward, the requirement for test methods that are ready for use, reliable and relevant as well as OECD input for guidance, and lastly a CRO that makes the test method(s) available for use and ultimately into regulation (Blum et al. 2025). Using the DNT IVB as a role model, the approach has been extended to developmental and reproductive toxicity (DART), with an ISTNET – DART Meeting setting out a road map for this highly complex endpoint (Fritsche et al., 2024).

#### 2.2.2 The Alternative Safety Profiling Algorithm (ASPA)

The Alternative Safety Profiling Algorithm (ASPA) was presented by Dr Andrew White (Unilever). ASPA has been developed within the European Union ASPIS Cluster of three projects (ONTOX, PrecisionTox and RISK-HUNT3R). ASPA intends to act as a workflow to implement and operationalise Next Generation Risk Assessment (NGRA) to support chemical safety assessment. It builds on existing tiered strategies for chemical safety assessment, including, but not limited to, workflows from SEURAT-1, US EPA, RISK21, ICCS, OECD guidance and those summarised by Browne et al. (2024). It is being developed and supported by case studies within the APSIS cluster, e.g. see the summaries of Case Studies 4 and 5 (Sections 2.3.4 and 2.3.5) in this report as well as Leist et al. (2025).

ASPA intends to support the assessment of systemic chronic health effects ensuring the protection of human health. Further, it is designed to be applicable to different regulations, being feasible, flexible and extendable to apply mechanistically-based NAMs. The aim is to provide an understandable and interpretable output demonstrating a degree of confidence for the user. As such, the ASPA workflow serves as a guide for data generation and interpretation for the assessment of systemic toxicity. The ASPA workflow intends to define, through a tiered approach, which tools and methods to use and how to evaluate data including an assessment of uncertainty. The workflow also provides context for the

data in terms of a hazard or risk assessment scenario with multiple exit points at which a decision can be made. The case studies within the RISK-HUNT3R project (see Case Studies 4 (Section 2.3.4) and 5 (Section 2.3.5)) are using existing data to evaluate the workflow and thus to demonstrate its applicability, and to build confidence in determining human relevant protective doses.

The ASPA is modular and based around a series of options, questions and provides guidance on how to make a decision. It has three distinct elements (or columns) to determine hazard, exposure, and ADME properties for particular safety assessment scenarios. The outputs from these three elements feed into the risk assessment. The structure of the three elements is intended to be efficient in terms of resources, starting where possible with *in silico* approaches, going forward to experimental NAM data to increase confidence for a particular purpose. The APSA can be visualised as a decision tree using building blocks and decision points as the main elements. Each of the building blocks and decision points has a unique identifier and will be provided a link to dedicated guidance. The tiers within each of the three elements of the ASPA are described in more detail in Section 3.2.1 in the light of other similar strategies and discussion within the User Forum.

ASPA and its implementation is ongoing and is considered to be a "living document". Whilst its implementation will be demonstrated through various case studies, a number of clear needs are already apparent. Amongst these are the requirement for the use of standard reporting formats, a greater and better appreciation of the role of uncertainty and how this informs the decision-making process and demonstration of how and where the APSA workflow could be applied within different regulatory contexts and for different industrial sectors. The workflow is currently being developed as a web-based tool and dashboard termed NAMASTOX.

### 2.3 Summary of the Case Studies

Five case studies (CS1-CS5), representing different endpoints and uses for NAMs were presented to the User Forum. The case studies were predominantly based on published material and are summarised, along with the relevant publication(s) below. The case studies were requested to provide specific comments, learnings and perspectives on topics such as the status of regulatory use of the described NAM, along with technical and performance aspects, as well as opportunities for future use and development. The learnings and insights from the case studies are compiled in Section 3.

# 2.3.1 Case Study 1 - Using Next Generation Risk Assessment to Make Safety Decisions for Cosmetic Ingredients Under Regulatory Scrutiny

The objective of Case Study 1 (CS1), presented by Dr Sophie Cable (Unilever), was to demonstrate
human safety assessment could be undertaken using NGRA. Specifically, NGRA for four case study
chemicals was described, these were selected from the Scientific Committee on Consumer Safety
(SCCS) priority list. NGRA was described as being exposure-led, hypothesis driven and designed to
$ensure\ the\ prevention\ of\ harm.\ \textit{Ab\ initio}\ assessments\ were\ performed\ to\ benchmark\ the\ outputs\ from$
a NAM-based safety assessment. Previous case studies have illustrated the use of NGRA for coumarin
(Baltazar et al., 2020) and benzophenone-4 (Baltazar et al., 2025). NGRA was based on a tiered
framework incorporating in vitro data for hazard and exposure. Three tiers are applied, Tier 0 being
problem formulation, in silico approaches and the application of TTC; Tier 1 being hazard and exposure
(in vitro) data generation; and Tier 2 is the refinement of the assessment to increase decision certainty.
Exit points exist within the three tiers if a safety decision can be made.
CS1 described in detail NGRA for climbazole in a use scenario of a preservative at 0.2% in a face cream.
The NGRA described in CS1 applies a systemic toolbox for early tier-testing. The toolbox is based on
the determination of the PoD using transcriptomics and assays for cellular stress pathways for non-
specific effects and in vitro pharmacological profiling assays for specific effects. In silico approaches
such as QSARs and structural alerts provide leads to direct the specific testing. Exposure in the 0.2%
formulation was above TTC thresholds, and further information to inform risk assessment was
required. Internal exposure was estimated through PBK modelling to provide a maximum
concentration in plasma (Cmax). A BER distribution is calculated from the PoD and exposure estimate.
The case study on climbazole was performed ab initio, on the assumption that there were no historic
data on which to base a safety decision. In silico analysis indicated alerts for reproductive toxicity and
carcinogenicity which informed the <i>in vitro</i> tests. NGRA demonstrated that it is possible to use NAM
data from the systemic toolbox to make safety decisions protective of human health. <i>In silico</i> models
such as PBK assessment could be over predictive and required refinement, this could be achieved with

#### 2.3.2 Case Study 2 - Improving Efficiency and Accuracy of NGRA for Low Toxicity Substances

the inclusion of in vitro biokinetic data. With regard to determining hazard, the transcriptomics and

cell stress assays covered most adverse effects, although there were concerns over the reliability of

cellular effects and the metabolic competence in the minimal set of cell lines. More knowledge is

required on the use of BER and associated variability and uncertainty in BER, with benchmarking of

#### A Case Study with Benzoic Acid

BER being a vital process to demonstrate its applicability.

The objective of Case Study 2 (CS2), presented by Dr Petra Kern (Procter and Gamble), was to demonstrate that a category could be created for substances with low toxicity to enable read-across to be performed to fill missing data gaps. Specifically, CS2 considered the quantitative assessment of the similarity of benzoic acid analogues using a variety of approaches. For the purposes of CS2, benzoic acid was the source substance with reliable toxicity data and a PoD of 500 mg/kg/d. Analogues were initially sought from the OECD QSAR Toolbox, however the profilers and similarity measures were not able to identify suitable analogues. Analogue identification was improved using a Matched Molecular Pair (MMP) approach that identifies molecules that differ only by a structural change at a single site or small portion of the molecule (Lester and Yan, 2021; Yan et al., 2023). It is well established that rating of analogues for read-across requires expert judgment (Lester et al., 2018). In order to optimise the process of analogue identification and reduce reliance on expert judgment, the "Quantifying Suitability of Analogues" (qSIM) approach has been developed (Lester et al., 2023). This incorporates information from metabolism, physico-chemical properties and structural alerts (coded as fingerprints) relating to reactivity and other toxicologically important properties.

In order to improve confidence in the use of read-across analogues, NAM data for *in vitro* metabolism were obtained. Confidence was also increased through the application of mechanistic NAM data, for instance existing ToxCast data as well as the generation of transcriptional and adapted pharmacological profiling (Burbank et al., 2024). A key aspect of the *in vitro* NAM testing was to set the highest concentration at a values consistent with exposure and the PoD in *in vivo* testing. Overall, CS2 demonstrated the need for better means to select analogues and that confidence could be achieved within a group of compounds associated with low toxicity when further, metabolically and mechanistically relevant, information was included.

# 2.3.3 Case Study 3 - Read-Across and New Approach Methodologies Applied in a 10-Step Framework for Cosmetic Safety Assessment – A Case Study with Parabens

The objective of Case Study (CS3), presented by Dr Gladys Ouédraogo (L'Oréal), was to describe and illustrate the 10-step process for read-across in the context of NGRA. Full details of the 10-step process for read-across supported by NAMs have been published by Alexander-White et al. (2022).

The 10-step framework for read-across is organised into three tiers which are broadly associated with the ICCS principles for NGRA (Dent et al., 2018). Tier 0 includes steps 1-4 to identify the structure, supporting data and search for analogues. Assessment or estimate of exposure is key for Tier 0 from e.g., use scenarios, and can be defined in different ways which may be refined as further information is made available. If sufficient information is available at Tier 0 (or after Tier 1 or 2), a decision can be

made. If insufficient information is available, the data collection proceeds to the next Tier. Tier 1 includes steps 5 and 6, which relate to ADME properties controlling bioavailability as well as data to inform on mode of action to better characterise the compounds. Tier 2 includes steps 7-10 and adds further refinement to the read-across through the collection of further information, e.g. through targeted use of NAMs testing or biokinetics, deriving a PoD, performing a Margin of Safety (MOS) evaluation and determining whether the level of confidence is acceptable. If sufficient information is not available, then the read-across will be ended.

The 10-step read-across framework was applied to the safety assessment for the use of propyl paraben as a preservative at 0.18% in cosmetics. A full description of the propyl paraben case study is available in Ouedraogo et al. (2022). Calculation of systemic exposure, which also included aggregate exposure, was above TTC, therefore the read-across assessment was initiated. The MMP approach (see CS2 (Section 2.3.2)) was applied and three significant analogues were identified on the basis of structural, reactivity and metabolic similarity. A variety of physico-chemical and in silico information was obtained for the target and source compounds including those associated with reproductive toxicity and endocrine disruption. Comparator molecules, with known activity, were also included to improve understanding of the in silico assessments. On the first attempt at read-across the MOS was too low, thus the systemic bioavailability was refined by the inclusion of further information, for instance for metabolism from studies in primary human hepatocytes, as well as a comparison of skin vs liver metabolism. Existing ToxCast and newly generated transcriptomics data (Naciff et al., 2022) were utilised to support mode of action. The NAM data confirmed the relationship between activity and alkyl chain, also that propyl paraben had lower activity than source compounds such as butyl paraben. This allowed for a refinement of internal exposure and bioactivity in Tier 2. Further tier 2 testing allowed further refinement and the use of Margin of Internal Exposure (MoiE).

The 10-step read-across framework provided a number of learnings with regard to the use of NAM data to support analogue selection and justification as well as making risk assessment decisions. Read-across based on chemical similarity alone has limitations. However, the similarities and differences in toxicokinetics and toxicodynamics were informed by appropriate NAM assays which strengthened potency assessment and internal exposure estimates. The safety assessment decision was assisted by the use of the MoIE. Overall, NAM data were shown to make read-across more robust and assessment of the confidence was valuable.

#### 2.3.4 Case Study 4 - Prioritisation and Screening: Which Testing Scope is Sufficient?

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The objective of Case Study 4 (CS4), presented by Matthias Wehr (Fraunhofer Institute for Toxicology and Experimental Medicine), was to evaluate an in vitro NAM assay battery for specific target organ toxicity - repeated exposure (STOT-RE) classification, within the context of the APSA workflow (see Section 2.2.2). The basis for the case study was an appreciation of the large number of chemicals which are used commercially but for which there are few, or no, toxicity data (EEA, 2019). CS4 focused specifically on the hazard element (column) within ASPA. Hazard identification in ASPA focuses on two steps, the first being the use of high throughput NAM assays for, e.g., prioritisation and screening, the second being to follow up on possible toxicological alerts or to reduce uncertainties with further mechanistic evidence based around the testing of AOPs. For prioritisation and screening, a key focus of CS4 was to determine the minimum in vitro testing approach to provide sufficient information to make a decision. Previous work has demonstrated for in vivo data that the Lowest Observed (Adverse) Effect Level LO(A)EL is driven by a relatively small number of main targets (Batke et al., 2013), therefore NAMs would not necessarily be required to cover every aspect of physiology and toxicology. The hypothesis is that assays for general signs of toxicity and effects on the main target organs could be sufficient for prioritisation and screening. A training set of about 30 toxic (STOT-RE1) and 30 low toxicity (no effect up to 1,000 mg/kg bw/d) compounds was established. Compounds were assessed in two tiers, the first using existing in silico tools and in vitro data, the second tier with an enriched test battery covering a broad biological space. Approximately three quarters of the compounds had ToxCast data – these showed good specificity but poorer sensitivity, and there were difficulties with when there were fewer data. Other information was obtained for about two thirds of the compounds from Chemical Effect Predictor (CEP) from DISGENET. CEP showed good sensitivity but poorer specificity (excluding data poor compounds). Further information was obtained from in silico predictions and alerts for liver metabolism and clearance. As a second part to CS4, the data were included into a scheme to assign compounds to levels of concern based on that activity and potential systemic availability (as defined by Berggren and Worth, 2023). In summary, the Tier 1 information applied through an ASPA workflow was able to distinguish toxic from low toxicity compounds. As STOT-RE does not take account of mechanism of action, broad testing methods may be suitable to obtain a protective PoD. However, it is difficult to compare existing in vitro data with each other and between compounds, therefore Tier 2 testing was applied to enrich the biological coverage and information. This involved broad mechanistic testing using seven unique human liver reporter cell lines covering 31 reporter gene (Calux) assays as well as seven stress pathways, phenomics cell painting from HepG2 cells and whole transcriptome analysis in three different cell systems. The concordance of the different assays was analysed and for more than 50% of the compounds the assays agreed and were consistent with the in vivo data. CS4 is on-going and

intends to demonstrate how to best combine the information from the assays and use machine learning to identify the most discriminative approaches. This will assist in the identification of the minimal *in vitro* testing required in ASPA.

# 2.3.5 Case Study 5 - Developmental Neurotoxicity Classification Labelling and Packaging Case Study

The objective of Case Study 5 (CS5), presented by Dr Ellen Hessel (RIVM), was to evaluate the potential of the use of NAMs for CLP purposes. The particular focus of CS5 was to identify the barriers, gaps and challenges of using of NAMs for CLP of DNT within the APSA workflow (refer to Section 2.2.2). In this context, it was confirmed that CS5 related to providing information regarding the intrinsic properties of a substance that are associated with its potential to cause harm, as stipulated by the criteria for classification. Thus, the exposure element (column) of the ASPA workflow was not considered in CS5, however, the ADME element will be considered to investigate if the compounds will enter the brain and cross barriers during pregnancy. Under CLP, DNT is currently considered under reproductive toxicity, mainly related to functional deficiency. The precedent in using *in vitro* NAMs for CLP, in the context of local toxicity (skin and eye irritation and skin sensitisation), through the use of defined approaches, was noted.

CS5 utilised five compounds and collected information from different *in silico* and *in vitro* NAMs. Valproic acid (VPA) and retinoic acid (RA) were chosen as positive control compounds due to their strong association with human DNT effects, consistent with findings reported by Aschner et al. (2017). 2-Ethylhexanoic acid showed DNT effects in mice and was therefore also included as positive control. Diethylene glycol (DEG) and sodium benzoate (SB) were selected as negative control reference compounds (Blum et al., 2023). Key questions were addressed regarding the sufficiency of existing AOPs and AOP networks relating to the complexity of the human brain, which is the basis for many of the currently used and proposed NAMs. Other challenges identified included understanding the information required from NAMs assays and when there is sufficient information, as well as whether adversity can be measured *in vitro* and considerations of assay performance. Knowledge of the processes of brain development is available and is the basis of the DNT IVB (see Section 2.2.1 and Fritsche et al., 2018), in addition there is an AOP network for DNT (Spînu et al., 2019). However, whether these summaries of the main process in brain development are sufficient to describe it is essential and not yet known.

The hazard identification of DNT was performed using *in silico* alerts and QSAR predictions (at Tier 0 of the ASPA workflow) for DNT itself and MIE predictions. Tier 1 assessment used a variety of high-

throughput *in vitro* assays including CALUX, cell painting, etc. The information from Tiers 0 and 1 will be combined to identify potential alerts to direct testing at Tier 2 – this process remains a clear challenge and will be informed by CS5. Tier 2 allows for hazard characterisation and is utilising the DNT *in vitro* battery (Crofton and Mundy, 2021; OECD, 2023) as well as complex assays within the RISK-HUNT3R project. Other key challenges include whether the complex Tier 2 assays cover all DNT effects, e.g., those associated with neurobehaviour and covering the complexity of the developing brain, and how this will relate to CLP for DNT. Other NAMs are investigating the use of systems biology networks and an *in silico* model for the closure of the neural tube.

The ADME element of the ASPA workflow was also considered. Whilst a PoD is not required for CLP, information was sought on whether compounds cross barriers as well as their bioavailability and metabolism. At Tier 1, toxicokinetics information from the literature will be utilised, in addition to knowledge of bioavailability and PBK modelling to the foetus. Tier 2 testing will include *in vitro* measurement of placental and blood-brain barrier passage. CS5 is on-going and will investigate further the use of the data, which are the most significant assays and how decisions can be made of CLP of DNT.

## 3. Summary of the Learnings and Insights from the NAMs User Forum

The User Forum illustrated the ongoing development and application of NAMs for chemical safety assessment and discussed in detail some of the practical aspects required for acceptance and decision making. There was a clear commitment to implementation of NAMs in chemical safety assessment from the participants in the User Forum, specifically from ECHA and EFSA.

This section summarises not only the main findings from the presentations and case studies, but also the discussion and comments submitted online and elsewhere. Where appropriate, reference is made to specific presentations or case studies. This section is organised around the needs to implement NAMs as well as their practical implementation.

### 3.1 Learnings from the Development of the DNT *In vitro* Battery (IVB)

The development of the DNT IVB (Section 2.2.1) represents significant progress in the development of NAMs for complex endpoints. It acknowledges that there will be no one-to-one replacement for complex *in vivo* endpoints. A number of significant aspects of the development of the DNT IVB could form a blueprint and be applied for further endpoints. These are summarised briefly according to Blum et al. (2025):

521 There is a benefit to gain international agreement of biology, e.g. by one, or more, expert 0 workshops that bring together relevant stakeholders to map the biological and 522 523 physiological processes involved. The purpose here is to identify the key biological 524 processes that result in adversity such that NAMs can be identified for them. 525 Once the key biological processes have been identified, there is a need to evaluate 0 526 currently available assays that cover these processes and which are adequate for use, as well as identifying gaps where further developed assays are required. For the DNT IVB this 527 528 was again achieved gaining agreement from experts and stakeholders. 529 There is a need to demonstrate reliability and relevance of NAM assays selected and 0 benchmark against known activities. An assessment of performance of the test battery -530 531 determining false negatives and positives – is required. To demonstrate performance, there is a need for a reference set of chemicals and test 532 0 533 results that cover recognised modes / mechanisms of action, as well as acknowledging which pathways are missing. 534 Case studies are highly beneficial to investigate the performance of a test battery and build 535 0 confidence. These will allow for the demonstration of the application of the test battery. 536 CS5 is an example of such a case study that is ongoing that applies a tiered approach 537 538 including the DNT IVB data and additional more specific assays to follow up on mechanistic 539 leads that measure the functionality of the nervous system to investigate whether NAMs can be used for hazard identification in CLP. 540 Once developed, any NAM or battery of NAMs needs to demonstrate transferability, for 541 542 instance from laboratory-to-laboratory. Such transferability goes beyond the development 543 of the NAM assay itself and will require funding. In the case of the DNT IVB EFSA funds this 544 transfer. The transferability was enabled by the foundation of a bespoke CRO, although more than a single organisation may be necessary. 545 A tiered approach, including one or more Defined Approaches, is useful to make the IVB 546 0 547 even more applicable. CS5 is demonstrating the use of NAMs, including data from the DNT-548 IVB, for CLP purposes within the ASPA framework. 549 It is essential to identify and characterise uncertainties in a test battery. This has been 0 550 achieved for the DNT IVB where uncertainties are known and can be addressed. It is

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#### 3.2 Application of NAMs in Tiered Strategies

hamper their application.

important that uncertainties of NAMs such as those identified for the DNT IVB do not

555	A variety of tiered testing strategies to implement NAMs for chemical safety assessment were
556	described at the User Forum. These attempt to combine information to allow decisions to be made
557	with regard to, e.g., hazard identification or risk assessment. The tiered testing strategies frequently
558	described three, or more, tiers as described in Section 3.2.1. The organisation of the tiered testing
559	strategies is designed to have decision points when sufficient confidence can be placed to make a
560	specific decision. Fundamental questions, which are expanded upon below, were:
561	Is the coverage provided by the cell lines protective?
562	Do NAMs provide the same level of protection?
563 564	<ul> <li>What is the extent of the biological coverage of the NAMs applied in the Tiers of the testing strategies?</li> </ul>
565	<ul> <li>What are the protection goals of a particular tiered testing strategy?</li> </ul>
566 567	The User Forum heard specific examples and learnings with the use of tiered testing strategies, which are summarised below.
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569	3.2.1 Tiered Strategies, Frameworks and Approaches will be Utilised for Safety Assessment:
570	An Increase in Understanding of Their Use is Required.
571	Various examples of tiered frameworks were presented at the User Forum (e.g. CS1, ASPA). Whilst
572	there are differences between the tiered frameworks, they have the same structure (Tiers $0-2$ and
573	decision points). The ASPA framework was described in detail with illustrative case studies (see Section
574	2.2.2)
575	There was broad consensus in how the Tiers in a framework are organised, as illustrated by the ASPA,
576	DNT IVB and case studies:
577	• Tier 0 involves the problem formulation, collection of existing information and data, for
578	instance on hazard and exposure. Techniques such as TTC may be applicable. In silico methods
579	such as QSARs, structural alerts, read-across can provide pointers for effects to follow up at
580	higher tiers (these can also be applied at Tier 1).
581	• Tier 1 generally comprises a broad set of general <i>in vitro</i> or molecular biological NAM assays.
582	• Tier 2 generally comprises more specific assays to follow up on mechanistic leads. This should
583	increase confidence in the decision being made.

There was agreement that the application of NAMs in tiered strategies can be used to make safety

decisions. Associated with this is a need to combine data-driven and knowledge driven (NAM)

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approaches with performance demonstrated, or benchmarked, against a reference test set. An example of the need for, and utility of, reference test sets was provided by the DNT IVB.

Decision points within and between Tiers are critical. Should sufficient confidence in the data be apparent, the decision can be made and testing stopped. If there is insufficient confidence, then further information is required, for instance by passing to the subsequent Tier. The User Forum agreed that there is a need for more information on when to go to a higher tier or exit the tiered strategy. An example could be the types of *in silico* or *in vitro* alerts that would trigger moving to a higher Tier.

Currently the definition of the scope of a protective NAM battery of tests (at Tier 1) is limited. In addition, how can tiered strategies, such as ASPA, be applied to different industrial sectors should be investigated. To achieve such goals, case studies were seen as being useful to demonstrate the utility of tiered testing strategies, as well as address the on-going questions such as decision points, sufficiency of information etc.

#### 3.2.2 Consensus on Which NAMs and Tools to Use in a Tiered Strategy

In the descriptions of tiered testing strategies (e.g. in the case studies) the User Forum was presented with a variety of types of NAMs for different endpoints and purposes. There was no attempt to reach agreement or consensus in the User Forum as to which are appropriate. There is a recognised challenge to make NAMs applicable across all legislations.

There was agreement that regardless of which NAMs are used, there should be consideration of whether they are relevant for the context of use and the issue(s) being addressed, protective, sensitive etc. To ensure NAMs within tiered strategies are protective, benchmarks for NAMs and the tiered strategies should be considered (analogous to the benchmarking of the NAMs themselves). As part of the benchmarking process, the conservatism in NAMs to enable a decision to be made should be considered. The implementation of NAMs should find a balance such that they are not overprotective.

There was also agreement for the need to identify commonalities, confidence and limitations (uncertainties) of NAMs for use in tiered strategies. It is likely that a number of NAMs will be applied, machine learning may be able to identify the optimum combination in terms of efficiency, i.e., minimum data required to make a decision (see CS4). There is a need to demonstrate a baseline set of NAM assays that if nothing was observed, then no adverse effects would be expected *in vivo*.

The biological coverage of NAMs is largely unknown and needs to be defined and described. It was acknowledged that NAMs cannot have universal coverage and for successful and appropriate application their applicability domain should be defined. Specifically, further knowledge is required on

618	whether NAMs (e.g. transcriptomics and cell stress assays, e.g. CS1), cover most / all adverse effects.
619	One suggestion to assess the utility of NAMs and tiered testing strategies was to consider repeated-
620	dose toxicity where there are data for many chemicals with a broad coverage of chemical space.
621	Some other specific recommendations and needs were identified:
622	• There is a need for compound selection in tiered strategy that will cover relevant mechanisms.
623	• There is a requirement for better understanding of NAM data, with regard to their capability
624	to identify adversity as opposed to (bio)activity or adaptation.
625	• There is a requirement, for instance at Tier 2, that the NAMs cover the complexity of the
626	endpoint being modelled. As example is the DNT IVB which needs to cover the complexity of
627	the brain to a sufficient level to identify adverse effects.
628	• The lack of consistent NAM data for the existing assays is perceived to be a problem. There is
629	a need for consistent data and to be able to identify where more are required to fill data gaps
630	(CS4).
631	• The maximum in vitro concentration to be tested (that may be used to demonstrate no
632	activity) is not consistent and will require more consideration (CS4).
633	•
634	The metabolic competence of NAMs is not known. Many are performed without a metabolic
635	component and the significance of this should be considered, also whether this should be part
636	of Tiers 1 or 2 of a tiered strategy.
637	More information may come from the APCRA studies and EPAA Designathon in terms of how
638	to refine the information that is available from NAMs.
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640	3.2.3 ASPA – An Alternative Safety Profiling Algorithm
641	The APSA is an example of a tiered testing strategy to implement a NGRA workflow (Leist et al., 2025).
642	Progress on the APSA is on-going with the purpose to enable various decisions for chemical safety
643	assessment. ASPA builds up evidence as defined in Tier 0-2 (Section 3.2.1). There are three main
644	elements (columns) to ASPA: hazard, ADME and exposure leading to risk assessment. These can be
645	adapted to specific purposes, e.g. for CLP purposes, hazard identification is key and does not require
646	exposure (CS5). It is designed to have a standard reporting approach.
647	The ASPA is designed with a number of decision / exit points. When there is sufficient confidence in
648	the information, a decision may be made. The identification and characterisation of uncertainty is
649	essential and vital aspect to make a decision – this should be documented adequately. ASPA is

designed to reduce uncertainty within the tiered approach, allowing for a conservative assessment of

651	hazard and exposure. An essential challenge is how to make a decision and when there is sufficient
652	information – to answer this question needs the input of regulators and PARC. In addition, the ASPA
653	is designed to be flexible, adaptable and updateable.
654	A number of ASPA case studies are being conducted in the RISK-HUNT3R Project. Case studies are
655	valuable to demonstrate the ASPA, how it can be applied to make decisions and develop it further. It
656	is intended that the ASPA will be provided with guidance and a digital version (NAMASTOX) to
657	implement it. Other recommendations included evaluating the ASPA to determine which parts could
658	be applicable for regulatory use and how to promote consensus building within the ASPA (see also
659	Section 3.4).
660	
661	3.3 On-Going Needs Identified for the Implementation of NAMs
662	The User Forum recognised that there is still considerable development needed in some areas of
663	NAMs. Various needs for the development of NAMs that have been previously stated through EPAA
664	workshops (Westmoreland et al., 2022) and User Forums (Cronin et al., 2025a) are not repeated here.
665	However, some clear additional needs were identified in the User Forum, particularly with regard to
666	regulatory implementation. These are summarised below.
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668	3.3.1 Appreciation of Uncertainty in Data and Decision Making
669	The appreciation of uncertainty in all aspects of the use of NAMs and their application in tiered testing
670	strategies and NGRA is crucial. This is often a neglected and underdeveloped topic that requires
671	further understanding. Specifically, there is a need to determine the acceptable level of uncertainty in
672	NGRA, for instance with the use of NAMs in a tiered strategy. Assessment of uncertainty is recognised
673	as being a vital component in the decision making process within strategies such as the ASPA
674	framework.
675	Assessment and understanding of uncertainty is crucial for all the data inputs into chemical safety
676	assessment. There was discussion in the User Forum regarding uncertainty in in vivo data. This is
677	important because in vivo data are currently required under many legislations, as well as being the
678	benchmark for the performance of many NAMs. It was acknowledged that uncertainty in in vivo data
679	may be large and is often undefined.
680	The uncertainty associated with in vitro NAM data should be characterised. Given the possible high

uncertainty in in vivo data, in vitro NAMs should not be expected to have lower uncertainty than the

in vivo data. Currently there may also be high uncertainty in toxicokinetic data. Uncertainty in

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toxicokinetic data is not acceptable as it will propagate through the safety decision making process
e.g. as part of the BER. Various strategies to reduce uncertainty were presented, including the
inclusion of NAMs data into ADME and exposure estimates (CS1) (see also Section 3.4).

#### 3.3.2 There is a Need to Set Goals and Performance Standards for NAMs

The importance of the validation of NAMs as part of their regulatory acceptance, and the challenges associated with that, are well acknowledged, for instance the discussion from a previous EPAA User Forum (Cronin et al., 2025a). However, the current User Forum acknowledged that a clear definition of success with regard to the use of NAMs is required. For instance, there could be an agreement of realistic goals and performance metrics for individual NAMs or groups of NAMs, such as specificity and balanced accuracy.

"Success criteria" for NAMs could be defined *a priori*. Once verified against these criteria, NAMs could be applied. This would support the easier development of tiered strategies and frameworks for chemical safety assessment. Clearly defined success criteria will allow the research community to understand what is required and expected when NAMs are being developed. There is also a need to benchmark the performance of NAMs / tiered strategies against the previous information requirements and decisions made. This should define and take account of the limitations of the new systems.

#### 3.3.3 Further Development of the Bioactivity-Exposure Ratio (BER)

The calculation of BER, or Margin of Exposure, is vital to apply NAMs in NGRA and with tiered testing strategies. This has been discussed previously in an EPAA User Forum (Cronin et al., 2025a). A variety of approaches to the application of BER were presented (e.g. CS1). However, no consensus was sought or reached in the User Forum as to how BER should be applied, the uncertainty in it and how it can be used to make a decision or be utilised in tiered testing strategy. The needs to benchmark BER to ensure it is protective, and better understand its uncertainty, were acknowledged.

#### 3.4 NAMs to Improve Exposure Assessment

The User Forum acknowledged that understanding exposure to chemicals is fundamental to the implementation of NAMs and application of tiered testing frameworks (e.g., CS1, CS3). There are a

713	number of aspects to this relating to estimates of internal exposure, relevance of doses in NAMs assays
714	through to aggregate exposure.

Exposure is fundamental for the application of NGRA, with several examples given in the User Forum (e.g., CS1, CS3). It is also one of the key elements within the ASPA. Further, knowledge of (internal and external) exposure is crucial to support the application of NAMs and tiered strategies for safety assessment. However, concerns were raised regarding the quality of the information relating to exposure and the possible high levels of uncertainties, e.g., in TK data (Section 3.3.1). As a fundamental part of NGRA, uncertainty in exposure assessment should be low, where possible.

It was observed that *in vitro* NAM data and information help improve exposure estimates and improve confidence. There is a definite approach to reduce uncertainty in exposure assessment in NGRA. This may assist in refining the exposure estimates as there is progression from Tier 0 to 1 to 2. Key NAM data for improving confidence in PBK models included hepatic clearance, fraction unbound and blood-plasma data. PBK models were calibrated against human clinical data (CS1).

Overall, there is a need to determine the best use of exposure information in NGRA and gain greater certainty in exposure estimates. For systemic toxicity, all cases should incorporate toxicokinetic and /or ADME information. Various approaches using PBK modelling to determine exposure were presented (e.g. CS1) although there is no consensus in their use. There is also a need to map exposure scenarios across industrial sectors and uses of chemicals.

# 3.4 Progress in *In Silico* and Other NAMs: Read-Across, -Omics Data and Category Formation

A number of other NAMs were described in the User Forum. A key *in silico* NAM is read-across, however read-across based on chemical structure and / or similarity alone was found to be limited. Structural similarity-based read-across may have too much uncertainty to be able to make a decision. There is value in combining a variety of metabolic, physico-chemical and reactivity data to improve confidence in analogue selection whereby similarity can be quantified by considering multiple streams of data. The use of profilers with the OECD QSAR Toolbox was not sufficient to identify meaningful analogues, approaches such as MMP were found to be more sophisticated (CS2). A variety of NAM data (e.g. -omics and ToxCast data) to support read-across were presented, based on both toxicodynamics and toxicokinetics. Transcriptional profiling assisted in identifying analogues with similar mechanisms of action. It is recognised that ToxCast data are incomplete and their use is challenging, it is preferable (where possible) to consider only data from shared assays, although this reduces the number of data to be considered.

Read-across / category formation can be used to group low toxicity substances, this is well supported by NAM data and can assist in addressing low toxicity substances. There is still debate on how to provide confidence in confirming an assessment of "low toxicity". Extending the application of read-across, the Cosmetics Europe 10-step read-across strategy is a tiered approach which incorporates elements of NGRA. It covers parts of Tiers 0-2 as described in Section 3.2.1. It has different decision / exit points. This read-across strategy also allows for the inclusion of NAM data to support read-across and increase confidence (CS3).

#### 4. Conclusions

The NAMs User Forum provided an opportunity to share learnings and experiences from a variety of stakeholders applying NAM data in NGRA. A variety of presentations were made which described the development and application of NAMs, typically within tiered testing strategies. A focus of the User Forum was determining the ability to make decisions from NAMs. Whilst some areas have made significant progress, e.g. DNT, for many areas of hazard identification and risk assessment further effort is required. The User Forum has provided an opportunity to identify areas where progress in implementing NAMs, through the use of tiered testing strategies, is required and essential to demonstrate the implementation of NGRA into practice.

### Disclaimer

The views and opinions expressed in this manuscript are those of the authors and contributors to the workshop, they do not represent those of the European Commission, the European Chemicals Agency and the European Food Safety Authority.

### **Declaration of Competing Interest**

770 Prof Ellen Fritsche is co-founder, shareholder and scientific managing director of DNTOX GmbH.

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785

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#### 5. References

- 787 Alexander-White C, Bury D, Cronin M, Dent M, Hack E, Hewitt NJ, Kenna G, Naciff J, Ouedraogo G,
- 788 Schepky A, Mahony C, Europe C (2022) A 10-step framework for use of read-across (RAX) in next
- 789 generation risk assessment (NGRA) for cosmetics safety assessment. Regul. Toxicol. Pharmacol. 129:
- 790 105094. doi: 10.1016/j.yrtph.2021
- 791 Aschner M, Ceccatelli S, Daneshian M, Fritsche E, Hasiwa N, Hartung T, Hogberg HT, Leist M, Li A,
- 792 Mundi WR, Padilla S, Piersma AH, Bal-Price A, Seiler A, Westerink RH, Zimmer B, Lein PJ (2017)
- 793 Reference compounds for alternative test methods to indicate developmental neurotoxicity (DNT)
- 794 potential of chemicals: example lists and criteria for their selection and use. ALTEX 34: 49-74. doi:
- 795 10.14573/altex.1604201Bal-Price A, Crofton KM, Leist M, Allen S, Arand M, Buetler T, Delrue N,
- 796 FitzGerald RE, Hartung T, Heinonen T, Hogberg H, Bennekou SH, Lichtensteiger W, Oggier D, Paparella
- 797 M, Axelstad M, Piersma A, Rached E, Schilter B, Schmuck G, Stoppini L, Tongiorgi E, Tiramani M,
- 798 Monnet-Tschudi F, Wilks MF, Ylikomi T, Fritsche E (2015) International STakeholder NETwork (ISTNET):
- 799 creating a developmental neurotoxicity (DNT) testing road map for regulatory purposes. Arch. Toxicol.
- 800 89: 269-287. doi: 10.1007/s00204-015-1464-2
- 801 Baltazar MT, Cable S, Cubberley R, Hewitt NJ, Houghton J, Kukic P, Li H, Malcomber S, Nicol B,
- Pendlington R, Punt A, Reynolds J, Scott S, Spriggs S, Dent MP (2025) Making safety decisions for a
- sunscreen active ingredient using next-generation risk assessment: Benzophenone-4 case study.
- 804 *ALTEX Altern. Anim. Exper.* doi 10.14573/altex.2501201
- 805 Baltazar MT, Cable S, Carmichael PL, Cubberley R, Cull T, Delagrange M, Dent MP, Hatherell S,
- Houghton J, Kukic P, Li H, Lee MY, Malcomber S, Middleton AM, Moxon TE, Nathanail AV, Nicol B,
- 807 Pendlington R, Reynolds G, Reynolds J, White A, Westmoreland C (2020) A Next-Generation Risk

808	Assessment case study for coumarin in cosmetic products. <i>Toxicol. Sci.</i> 176: 236-252. doi:
809	10.1093/toxsci/kfaa048
810	Batke M, Aldenberg T, Escher S, Mangelsdorf I (2013) Relevance of non-guideline studies for risk
811	assessment: the coverage model based on most frequent targets in repeated dose toxicity studies.
812	Toxicol. Lett. 218: 293-298. doi: 10.1016/j.toxlet.2012.09.002
813	Berggren E, Worth AP (2023) Towards a future regulatory framework for chemicals in the European
814	Union - Chemicals 2.0. Regul. Toxicol. Pharmacol. 142: 105431. doi: 10.1016/j.yrtph.2023.105431
815	Blum J, Bartmann K, de Paula Souza J, Fritsche E (2025) Developmental neurotoxicity as a case example
816	for a six-step framework for the sustainable regulatory implementation of NAMs. Curr. Opin. Toxicol.
817	42: 100528. doi: /10.1016/j.cotox.2025.100528
818	Blum J, Masjosthusmann S, Bartmann K, Bendt F, Dolde X, Dönmez A, Förster N, Holzer AK, Hübenthal
819	U, Keßel HE, Kilic S, Klose J, Pahl M, Stürzl LC, Mangas I, Terron A, Crofton KM, Scholze M, Mosig A,
820	Leist M, Fritsche E (2023) Establishment of a human cell-based <i>in vitro</i> battery to assess developmental
821	neurotoxicity hazard of chemicals. <i>Chemosphere</i> 311(Pt 2): 137035. doi:
822	10.1016/j.chemosphere.2022.137035
823	Browne P, Paul Friedman K, Boekelheide K, Thomas RS (2024) Adverse effects in traditional and
824	alternative toxicity tests. Regul. Toxicol. Pharmacol. 148: 105579. doi: 10.1016/j.yrtph.2024.105579
825	Burbank M, Kukic P, Ouedraogo G, Kenna JG, Hewitt NJ, Armstrong D, Otto-Bruc A, Ebmeyer J,
826	Boettcher M, Willox I, Mahony C (2024) In vitro pharmacologic profiling aids systemic toxicity
827	assessment of chemicals. Toxicol. Appl. Pharmacol. 492: 117131. doi: 10.1016/j.taap.2024.117131
828	Carstens KE, Carpenter AF, Martin MM, Harrill JA, Shafer TJ, Paul Friedman K (2022) Integrating data
829	from in vitro New Approach Methodologies for developmental neurotoxicity. Toxicol. Sci. 187: 62-79.
830	doi: 10.1093/toxsci/kfac018
831	Crofton KM, Mundy WR (2021) External scientific report on the interpretation of data from the

- 832 developmental neurotoxicity in vitro testing assays for use in Integrated Approaches for Testing and
- 833 Assessment. EFSA Supporting Publication 18: EN-6924. 42 pp. doi: 10.2903/sp.efsa.2021.EN-6924
- Cronin MTD, Baltazar MT, Barton-Maclaren TS, Bercaru O, De Abrew KN, Desaintes C, Escher SE, Kern 834
- 835 P, Maxwell G, Rogiers V, Schutte K, Sobanski T (2025a) Report on the European Partnership for
- 836 Alternative Approaches to Animal Testing (EPAA) "New Approach Methodologies (NAMs) User Forum
- 837 Kick-off Workshop". Regul. Toxicol. Pharmacol. 159: 105796. doi: 10.1016/j.yrtph.2025.105796.
- 838 Cronin MTD, Berggren E, Camorani S, Desaintes C, Fabbri M, Fabrega J, Herzler M, Ingram JDE, Lacasse
- 839 K, Louhimies S, Maxwell G, Schutte K, Sobanski T, Streck G, Terron A, Worth AP (2025b) Report of the

- 840 European Commission workshop on "The roadmap towards phasing out animal testing for chemical
- safety assessments", Brussels, 11–12 December 2023. Regul. Toxicol. Pharmacol. 161: 105818. doi:
- 842 10.1016/j.yrtph.2025.105818
- Dent M, Teixeira Amaral R, Amores Da Silva P, Ansell J, Boisleve F, Hatao M, Hirose A, Kasai Y, Kern P,
- Kreiling R, Milstein S, Montemayor B, Oliveira J, Richarz A, Taalman R, Vaillancourt E, Verma R, Vieira
- 845 O'Reilly Cabral Posada N, Weiss C, Kojima H (2018) Principles underpinning the use of new
- methodologies in the risk assessment of cosmetic ingredients. *Comput. Toxicol.* 7: 20-26.
- 847 ECHA (2024) Key Areas of Regulatory Challenge. ECHA-23-R-08-EN, ECHA, Helsinki, Finland. doi:
- 848 10.2823/858284.
- 849 EFSA (2021a) EFSA PPR Panel (EFSA Panel on Plant Protection Products and their Residues),
- Hernández-Jerez A, Adriaanse P, Aldrich A, Berny P, Coja T, Duquesne S, Focks A, Marinovich M, Millet
- 851 M, Pelkonen O, Pieper S, Tiktak A, Topping C, Widenfalk A, Wilks M, Wolterink G, Crofton K, Hougaard
- 852 Bennekou S, Paparella M and Tzoulaki I, 2021. Scientific Opinion on the development of Integrated
- Approaches to Testing and Assessment (IATA) case studies on developmental neurotoxicity (DNT) risk
- assessment. EFSA Journal 2021;19(6):6599, 63 pp. doi: 10.2903/j.efsa.2021.6599
- 855 EFSA (2021b) EFSA Strategy 2027. Science, safe food and sustainability. EFSA, Palma, Italy. ISBN 978-
- 856 92-9499-263-5. doi:10.2805/274627. Available from:
- 857 https://www.efsa.europa.eu/sites/default/files/2021-07/efsa-strategy-2027.pdf (accessed 12
- 858 January 2025).
- 859 Escher SE, Partosch F, Konzok S, Jennings P, Luijten M, Kienhuis A, de Leeuw V, Reuss R, Lindemann K-
- 860 M, Hougaard Bennekou S (2022) Development of a Roadmap for Action on New Approach
- Methodologies in Risk Assessment. 153 pp. doi:10.2903/sp.efsa.2022
- 862 European Environment Agency (2019) The European Environment State and Outlook 2020 -
- 863 Knowledge for Transition to a Sustainable Europe, Publications Office,
- 864 https://data.europa.eu/doi/10.2800/96749
- 865 Fritsche E, Aspiroz LS, Arand M, Faustman E, Müller I (2024) International STakeholder NETwork
- 866 (ISTNET) Workshop for creating a developmental and reproductive toxicity (DART) testing roadmap
- 867 for regulatory purposes. ALTEX 41: 671-673. doi: 10.14573/altex.2410081
- 868 Fritsche E, Barenys M, Klose J, Masjosthusmann S, Nimtz L, Schmuck M, Wuttke S, Tigges J (2018)
- Development of the concept for stem cell-based developmental neurotoxicity evaluation. *Toxicol. Sci.*
- 870 165: 14-20. doi: 10.1093/toxsci/kfy175

- Haase A, Barroso J, Bogni A, Bremer-Hoffmann S, Fessard V, Gutleb AC, Mast J, McVey E, Mertens B,
- 872 Oomen AG, Ritz V, Serchi T, Siewert K, Stanco D, Usmani SM, Verleysen E, Vincentini O, van der Zande
- 873 M, Cubadda F (2024) Proposal for a fit for purpose qualification system for New Approach
- Methodologies (NAMs) in the food and feed sector. EFSA supporting publication 2024:EN-9008. 96 pp.
- 875 doi: 10.2903/sp.efsa.2024
- Koch K, Bartmann K, Hartmann J, Kapr J, Klose J, Kuchovská E, Pahl M, Schlüppmann K, Zühr E, Fritsche
- 877 E (2022) Scientific validation of human neurosphere assays for developmental neurotoxicity
- 878 evaluation. *Front. Toxicol.* 4: 816370. doi: 10.3389/ftox.2022.816370
- Koch K, Schlüppmann K, Hüsken S, Merit Stark L, Förster N, Masjosthusmann S, Klose J, Dönmez A,
- 880 Fritsche E (2025) Nuclear hormone receptors control fundamental processes of human fetal
- 881 neurodevelopment: Basis for endocrine disruption assessment. Environ. Int. 198: 109400. doi:
- 882 10.1016/j.envint.2025.109400
- 883 Leist M, Tangianu S, Affourtit F, Braakhuis H, Colbourne J, Cöllen E, Dreser N, Escher SE, Gardner I,
- Hahn S, Hardy B, Herzler M, Islam B, Kamp H, Magel V, Marx-Stoelting P, Moné MJ, Lundquist P,
- Ottenbros I, Ouedraogo G, Pallocca G, van de Water B, Vinken M, White A, Pastor M, Luijten M (2025)
- 886 An Alternative Safety Profiling Algorithm (ASPA) to transform next generation risk assessment into a
- structured and transparent process. *ALTEX* doi: 10.14573/altex.2509081.
- Lester C, Byrd E, Shobair M, Yan G (2023) Quantifying analogue suitability for SAR-based read-across
- toxicological assessment. Chem. Res. Toxicol. 36: 230-242. doi: 10.1021/acs.chemrestox.2c00311
- 890 Lester C, Reis A, Laufersweiler M, Wu S, Blackburn K (2018) Structure activity relationship (SAR)
- 891 toxicological assessments: The role of expert judgment. *Regul. Toxicol. Pharmacol.* 92: 390-406. doi:
- 892 10.1016/j.yrtph.2017.12.026
- Lester CC, Yan G (2021) A matched molecular pair (MMP) approach for selecting analogs suitable for
- 894 structure activity relationship (SAR)-based read across. Regul. Toxicol. Pharmacol. 124: 104966. doi:
- 895 10.1016/j.yrtph.2021.104966
- Masjosthusmann S, Blum J, Bartmann K, Dolde X, Holzer AK, Stürzl LC, Keßel EH, Förster N, Dönmez A,
- 897 Klose J (2020) Establishment of an a priori protocol for the implementation and interpretation of an
- 898 in-vitro testing battery for the assessment of developmental neurotoxicity. EFSA Supporting
- 899 Publications. 17(10):1938E.
- 900 Naciff JM, Shan YK, Wang X, Daston GP (2022) Transcriptional profiling efficacy to define biological
- activity similarity for cosmetic ingredients' safety assessment based on next-generation read-across.
- 902 Front. Toxicol. 4: 1082222. doi: 10.3389/ftox.2022.1082222

- 903 OECD (2023) Initial Recommendations on Evaluation of Data from the Developmental Neurotoxicity
- 904 (DNT) In-Vitro Testing Battery, OECD Series on Testing and Assessment, No. 377, OECD Publishing,
- 905 Paris, <a href="https://doi.org/10.1787/91964ef3-en">https://doi.org/10.1787/91964ef3-en</a>.
- 906 Ouedraogo G, Alexander-White C, Bury D, Clewell HJ 3rd, Cronin M, Cull T, Dent M, Desprez B,
- 907 Detroyer A, Ellison C, Giammanco S, Hack E, Hewitt NJ, Kenna G, Klaric M, Kreiling R, Lester C, Mahony
- 908 C, Mombelli E, Naciff J, O'Brien J, Schepky A, Tozer S, van der Burg B, van Vugt-Lussenburg B, Stuard
- 909 S, Cosmetics Europe (2022) Read-across and new approach methodologies applied in a 10-step
- 910 framework for cosmetics safety assessment A case study with parabens. Regul. Toxicol. Pharmacol.
- 911 132: 105161. doi: 10.1016/j.yrtph.2022.105161
- Paparella M, Bennekou SH, Bal-Price A (2020) An analysis of the limitations and uncertainties of in vivo
- 913 developmental neurotoxicity testing and assessment to identify the potential for alternative
- 914 approaches. *Reprod. Toxicol.* 96: 327-336. doi: 10.1016/j.reprotox.2020.08.002
- 915 Paul Friedman K, Gagne M, Loo LH, Karamertzanis P, Netzeva T, Sobanski T, Franzosa JA, Richard AM,
- 916 Lougee RR, Gissi A, Lee JJ, Angrish M, Dorne JL, Foster S, Raffaele K, Bahadori T, Gwinn MR, Lambert J,
- 917 Whelan M, Rasenberg M, Barton-Maclaren T, Thomas RS (2020) Utility of in vitro bioactivity as a lower
- bound estimate of *in vivo* adverse effect levels and in risk-based prioritization. *Toxicol. Sci.* 173: 202-
- 919 225. doi: 10.1093/toxsci/kfz201
- 920 Paul Friedman K, Thomas RS, Wambaugh JF, Harrill JA, Judson RS, Shafer TJ, Williams AJ, Joey Lee J-Y,
- Loo L-H, Gagné M, Long AS, Barton-Maclaren TS, Whelan M, Bouhifd M, Rasenberg M, Simanainen U,
- 922 Sobanski T (2025) Integration of new approach methods for the assessment of data-poor
- 923 chemicals, *Toxicol. Sci.* 205: 74–105. doi: 10.1093/toxsci/kfaf019
- 924 Smirnova L, Hogberg HT, Leist M, Hartung T (2024) Revolutionizing developmental neurotoxicity
- 925 testing a journey from animal models to advanced in vitro systems. ALTEX 41: 152-178. doi:
- 926 10.14573/altex.2403281
- 927 Spînu N, Bal-Price A, Cronin MTD, Enoch SJ, Madden JC, Worth AP (2019) Development and analysis
- 928 of an adverse outcome pathway network for human neurotoxicity. *Arch. Toxicol.* 93: 2759-2772. doi:
- 929 10.1007/s00204-019-02551-1
- 930 Westmoreland C, Bender HJ, Doe JE, Jacobs MN, Kass GEN, Madia F, Mahony C, Manou I, Maxwell G,
- 931 Prieto P, Roggeband R, Sobanski T, Schütte K, Worth AP, Zvonar Z, Cronin MTD (2022) Use of New
- 932 Approach Methodologies (NAMs) in regulatory decisions for chemical safety: Report from an EPAA
- 933 Deep Dive Workshop. *Regul. Toxicol. Pharmaco.* 135: 105261. doi: 10.1016/j.yrtph.2022.105261

934	Yan G, Rose J, Ellison C, Mudd AM, Zhang X, Wu S (2023) Refine and strengthen SAR-based read-across
935	by considering bioactivation and modes of action. Chem. Res. Toxicol. 36: 1532-1548. doi:
936	10.1021/acs.chemrestox.3c00156
937	

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# **Highlights**

- NAMs described for hazard identification and exposure assessment
- Updates on application of NAMs from EFSA and ECHA
- Tiered testing strategies can assist in the regulatory implementation of NAMs
- Case studies demonstrate the applicability of NAMs
- Learnings and needs for NAMs' development identified

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Declaration of interests
$\Box$ The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.
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Prof Ellen Fritsche is co-founder, shareholder and scientific managing director of DNTOX GmbH.