



European Partnership for Alternative Approaches to Animal Testing (EPAA)

Our vision

The replacement, reduction and refinement (3Rs) of animal use for meeting regulatory requirements through better & more predictive science (e.g. New Approach Methodologies (NAMs)). The partners are pooling knowledge and resources to accelerate the development, validation and acceptance of alternative approaches at national, European and global levels.

Our structure

EPAA Steering Committee takes strategic decisions on EPAA activities and is composed of representatives of the member companies and of services of the European Commission.

The EPAA covers **eight industry sectors** encompassing animal health, chemicals, cosmetics, fragrances, food and drinks, pharmaceuticals, soap and detergents and crop protection products. The co-chairs from industry and the European Commission convene the quarterly meetings.



The **EPAA** strategy is defined by a 5-year Action Programme; the current programme runs from 2021 to 2025. The EPAA reports progress against the strategy to its members and 3Rs-concerned stakeholders during the EPAA Annual Conference.

The **EPAA Mirror Group** consists of experts from the civil society including academia, animal welfare, laboratory animal science, 3Rs centres, and other third-party organizations. It acts as a consultation forum in an advisory capacity to the Steering Committee for the implementation of the Action Programme, providing comments from a broader societal perspective.



EPAA Annual Conference 2023

Our mission

Launched in 2005, EPAA is a unique partnership between industry and the European Commission with the aim of:

- Promoting the development & validation of animal alternatives for regulatory animal testing.
- Accelerating regulatory use of animal alternatives at member state, European, and global levels through fostering knowledge exchange among partners and stakeholders.
- Providing a platform for stakeholder dialogue on scientific, regulatory, and policy developments that impact regulatory animal testing.

In fulfilling its mission, EPAA also considers innovation, protection of intellectual property, and overall competitiveness of European industry.

In recent years EPAA has also sought to facilitate a successful implementation of the European Green Deal, Chemical Strategy for Sustainability, Pharmaceutical Strategy for Europe, and EU Commission roadmap towards ultimately phasing out animal testing for chemical safety assessments.





Our activities

EPAA projects are overseen by a Projects Platform & cover:

- Acute toxicity
- Harmonisation of 3Rs in Biologicals
- Carcinogenicity of Agrochemicals
- Skin sensitization NAM User Forum
- NAMs in regulatory decisions for chemical safety
 - -NAM Designathon (challenge for chemicals classification)
 - –NAM User Forum
- Environmental Safety Assessment

The EPAA also provides student travel grants and an annual prize for either lab technicians (Refinement Prize) or young scientists (3Rs Science Prize).





EPAA Annual report 2023



The Commission roadmap towards ultimately phasing out animal testing for chemical safety assessments

European Commission, DG GROW and DG ENV

Sofia Camorani, Marco Fabbri, Katrin Schutte, Georg Streck



Commission



COMMISSION ROADMAP

With Commission Communication C(2023) 5041 replying to the European Citizens' Initiative (ECI) 'Save Cruelty-free Cosmetics - Commit to a Europe without Animal Testing' the Commission announced the development of a roadmap towards ultimately phasing out animal testing for chemical safety assessments. The roadmap will

- Be finalised in the first quarter of the term of the next Commission (Q4 2025/Q1 2026).
- Outline milestones and specific actions, to be implemented in the short to longer term, to reduce animal testing and describe the transition towards an animal-free regulatory system.
- Analyse and describe the necessary steps to replace animal testing in all relevant pieces of chemical legislation (e.g. REACH, Biocidal Product Regulation, Plant Protection Products Regulation and human and veterinary medicines).
- Describe the path to expand and accelerate the development, validation and implementation of non-animal methods as well as means to facilitate their uptake across legislations.





What information do animal-based studies provide?

Options for short-term replacements?

Replacements after further method development?

Long- and short-term action points:

- Identify non-animal approaches
- List options to reduce animal needs
- List replacement options
- Define methods development needs
- Define protection goals
- Define animal-free regulatory system
- Identify required legislative changes
- ...

Transition to animal-free regulatory system

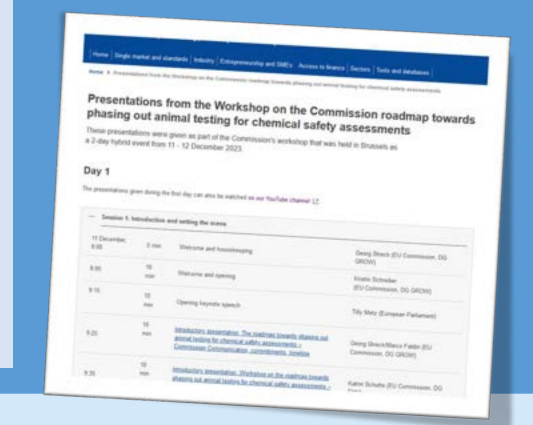
How to reach protection goals with non-animal methods?

Further elements of the roadmap

- 1. Analysis of strengths and weaknesses of the landscape of agencies, committees and working groups that provide advice on non-animal methods.
- 2. Analysis of the need and feasibility of an advisory expert scientific committee to provide advice on the development of non-animal approaches and their uptake and use in the regulatory context.
- 3. Analysis of possibilities to accelerate the validation and acceptance of new non-animal methods.
- 4. Analyse how to facilitate access to information such as upcoming events, calls, but also on guidance, e.g. through dedicated platforms and interactive communication tools.
- 5. A proposal on 'Streamlining EU scientific and technical work on chemicals through the EU agencies' that has the purpose to enhance the collaboration of the agencies and to increase their efficiency by making full use of synergies in the assessment of chemicals.
- 6. An analysis, in close collaboration with the agencies, of the possibilities to increase the agencies' visibility and impact in international forums.
- 7. Outline ways to improve outreach activities in the international dimension (e.g., GHS, OECD, ICH/VICH, ISO).
- 8. A proposal for a Regulation on chemicals data to improve accessibility to information on chemicals (proposal submitted).

OUTREACH AND INVOLVEMENT OF STAKEHOLDERS

Involving stakeholders is crucial for pooling the scientific knowledge that forms the basis of the roadmap and essential to receive support from Member States, agencies and all stakeholders, including Commission Partnerships such as the European Partnership for Alternative Approaches to Animal Testing (EPAA) or the Partnership for the Assessment of Risks from Chemicals (PARC), both during development of the roadmap and in the implementation phase. The roadmap development is therefore accompanied by multiple stakeholder activities and a stakeholder consultation plan according to the Better Regulation rules.



- Public consultation on the initiative on Have-your-say (https://ec.europa.eu/info/law/b etter-regulation/have-your-say en)
- Targeted stakeholder consultations in meetings/surveys
- Information of MS and stakeholders via Committees and Expert Groups
- Outreach activities with non-EU partners and international organisations

Organisation of workshops:

- A workshop end of 2023 to kick off the work on the roadmap
- A workshop in the second half of 2024 on the progress of the roadmap (25 Oct. 2024)
- Possible further workshops on scientific and regulatory aspects

Workshop on the Commission roadmap towards phasing out animal testing in chemical safety assessments (11/12 December 2023)

- Presentations and discussions with stakeholders (MS, EU agencies, industry, research community, NGOs) on ideas and approaches for the roadmap.
- Session in cooperation with the Partnership for the Assessment of Risks from Chemicals (PARC), WP2.2, on the Next Generation Risk Assessment.
- Presentations and recordings: https://single-market-economy.ec.europa.eu/presentations-workshop-commission-roadmap-towards-phasing-out-animal-testing-chemical-safety en





Transparency — a working tool to make progress together!

The European Commission (DG Environment) publishes data on the use of animals for scientific purposes based on information reported by the 27 Member States according to Directive 2010/63/EU. The Alures statistical and the Alures Non-Technical Summary databases provide not only reliable statistics but also insights into why and how animals are used in science. These transparency tools are crucial for knowing where our efforts to reduce or replace animal use in science would be most effective.

ALURES databases, world's most comprehensive depository on animal use taking transparency to another level!

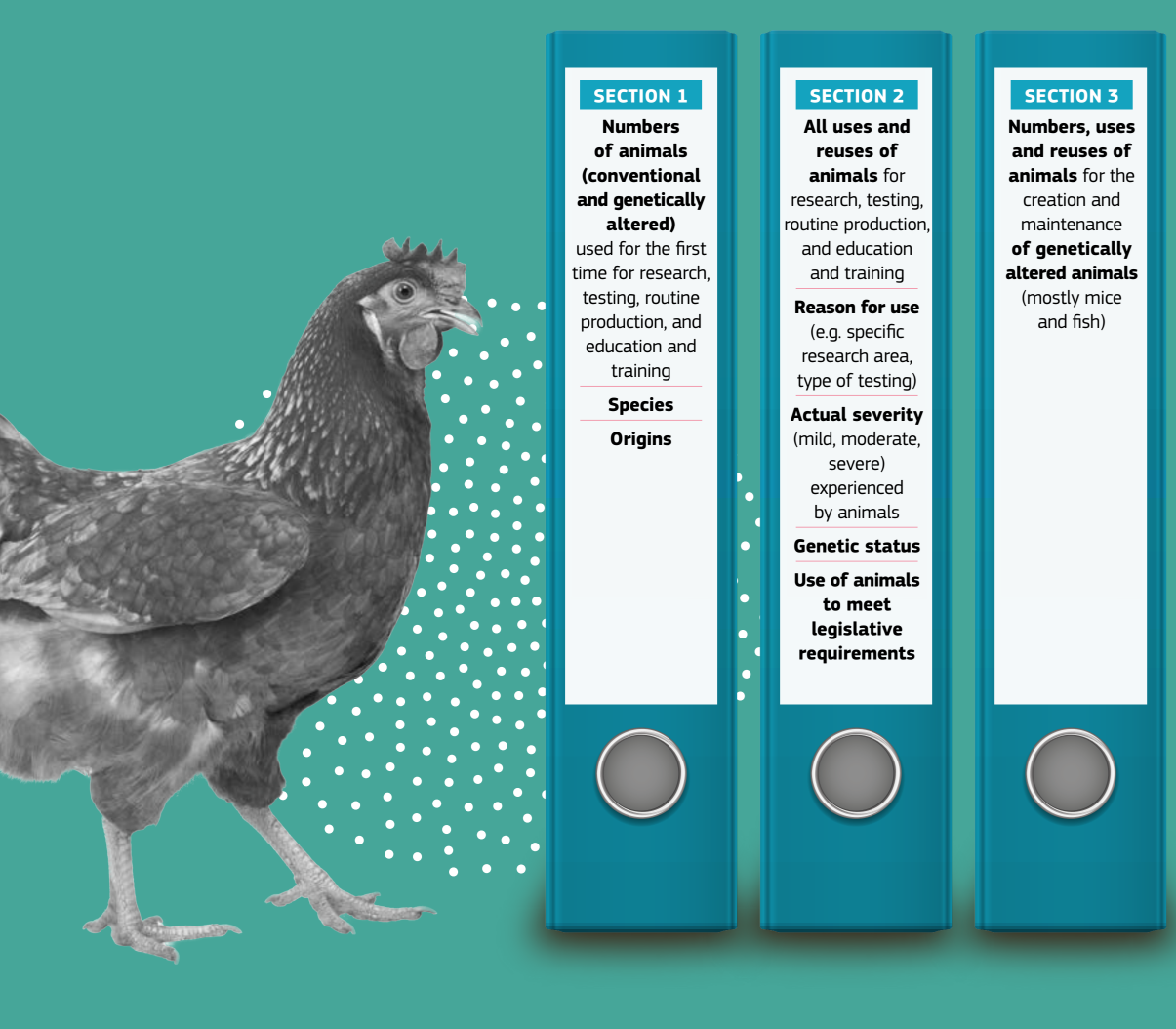
Reliable statistics with the understanding and openness in animal use allow

- The Public to have a complete overview of when and how animals in Europe are being used in science.
- **Researchers** and **funders** to determine urgent focus areas for the development of new alternative approaches
- **Policy makers** to make evidence-based, informed policy decisions

1. ALURES Statistical Database



What information is available?



2. ALURES NTS Database



Gaining insight into the use of animals in science.

Non-technical project summaries (NTS)

 provide clear and concise descriptions of authorised animal projects in the EU

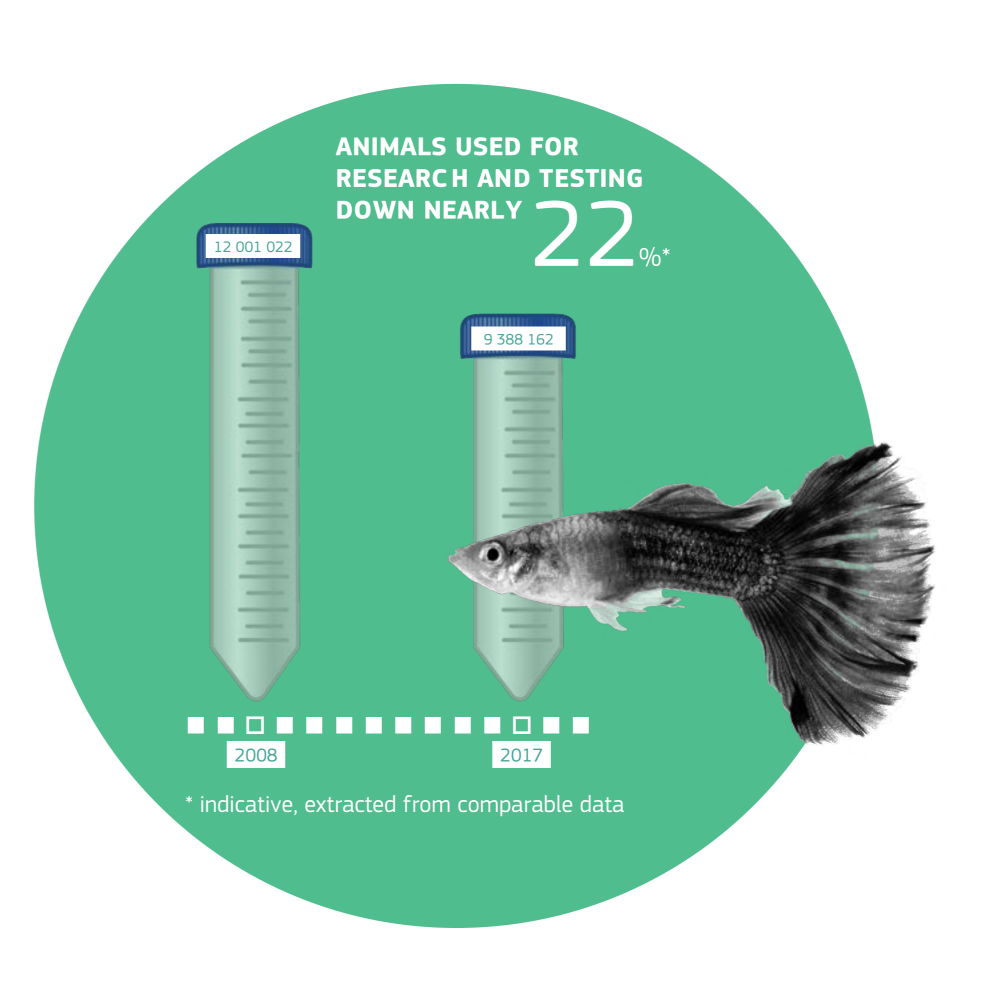
 facilitate understanding of why and how animals are used and how the work is done

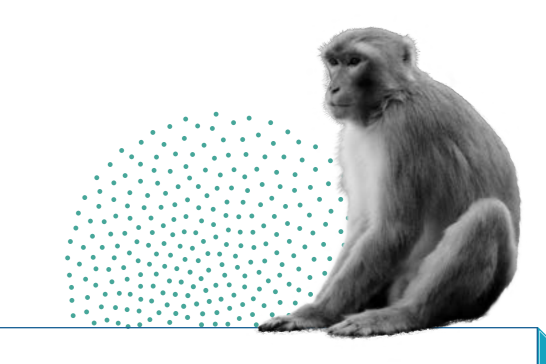
 explain how the Three Rs are applied and what limitations they have



Making the EU a global leader in transparency

The EU is committed to phasing out the use of animals in science when it is scientifically possible to do so. It is working towards the ultimate goal of replacing all animals used for scientific purposes, but more time is needed to develop alternative approaches that do not involve animals.





WHAT ARE **SCIENTIFIC PURPOSES?**

All uses of animals for basic, translational and applied research, regulatory testing and production, education and taining, as well as the creation and maintenance of genetically altered animal lines

Protecting and improving the welfare of animals in scientific research

Why is it still necessary to use animals in research?

Animals have played key roles in medical advancements of the last century. We would not enjoy better health, improved quality of life and longer life expectancy without the knowledge gained from animal research.

Technological advances, computer simulations and test tube methods already greatly reduce the number of animals used, but are not yet able to fully replicate living organisms' complexities and reactions.

Hence, alternative solutions are not yet always available.

It is the EU's ultimate goal to completely replace animals in science. Until this becomes reality, it is committed to reducing the number of animals and respecting the welfare of the animals used for scientific purposes.

How are animals protected?

In Europe, all living animals used in science are protected by very strict legislation, Directive 2010/63/EU. All animal studies must comply with this legislation. Animals cannot be used for scientific purposes without prior authorisation. Authorities can only allow the use of animals when there are no alternative, non-animal methods available. In addition, the use of the animals must be justified by the expected benefits, also taking into account ethical considerations.

The Three Rs, to Replace, Reduce and Refine the use of animals, at the heart of the EU legislation

- There is a legal obligation to eliminate or minimise pain, suffering, distress and lasting harm on animals to a minimum level possible.
- All efforts to minimise pain, suffering and distress have to be made from the planning stage.
- Every establishment must have a named person responsible for the welfare and care of animals, as well as a designated veterinarian.



poster

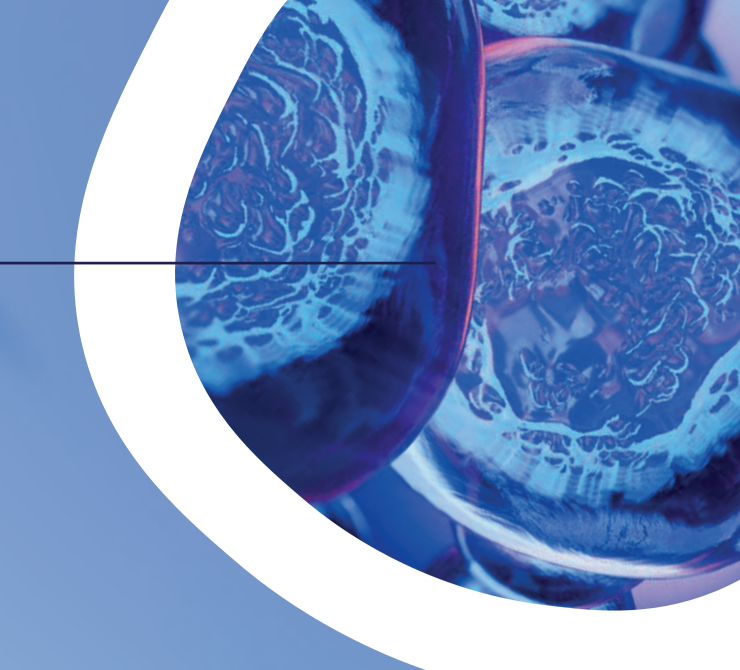
science

Animals in EC website

Luxembourg: Publications Office of the European Union, 2024

© European Union, 2024 Reuse of this document is allowed, provided appropriate credit is given and any changes are indicated (Creative Commons Attribution 4.0 International license). For any use or reproduction of elements that are not owned by the EU, permission may need to be sought directly from the respective right holders. All images $\mathbb C$ stock.adobe.com – all rights reserved

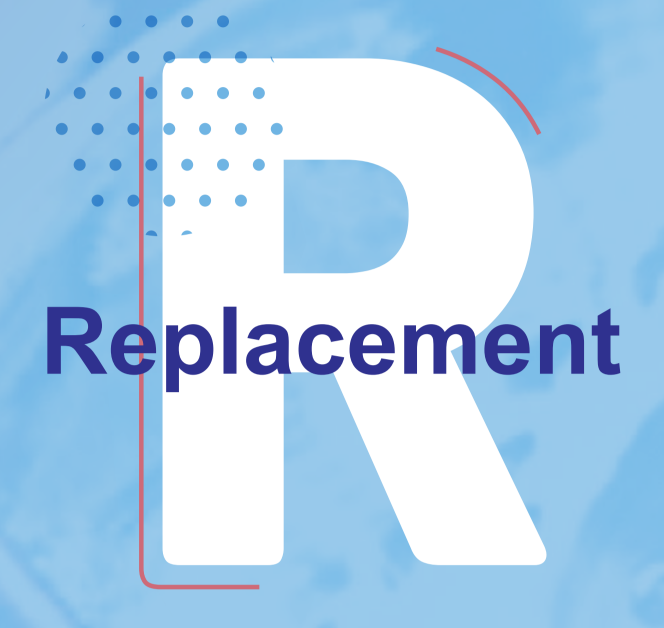




EURL ECVAM

Non-animal methods in science and regulation

FOR OVER 30 YEARS THE JRC HAS BEEN WORKING ON THE THREE RS



use alternatives instead of animals



use fewer animals



minimise pain and distress

The mandate of EURL ECVAM is set out in EU legislation to protect animals used for scientific purposes and includes the following duties



research and development



method validation



knowledge sharing



promotion of the three Rs



EU Reference Laboratory for alternatives to animal testing (EURL ECVAM)

joint-research-centre.ec.europa.eu/eu-reference-laboratory-alternatives-animal-testing-eurl-ecvam_en



NEW Status Report



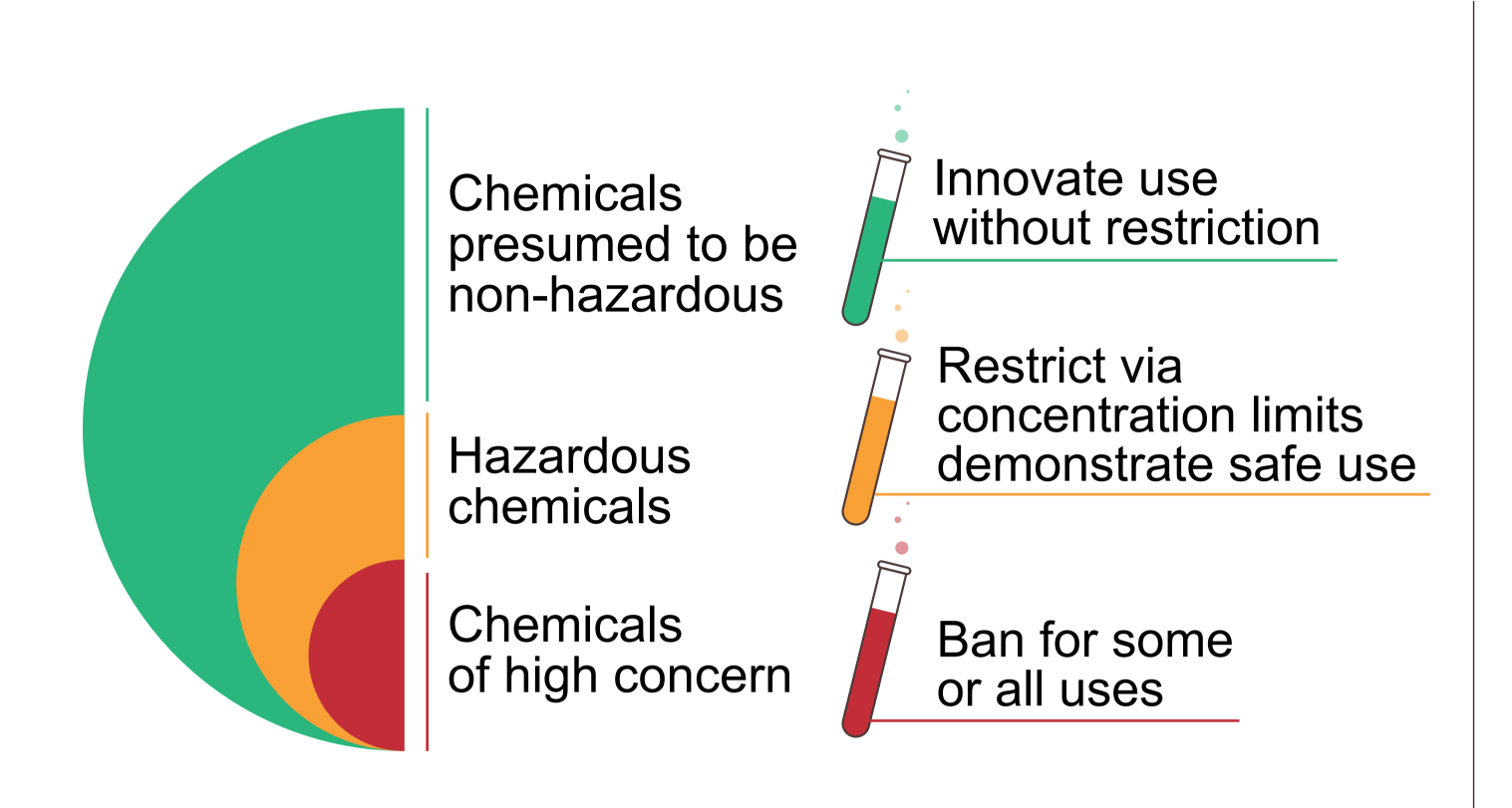


NAM DESIGNATHON



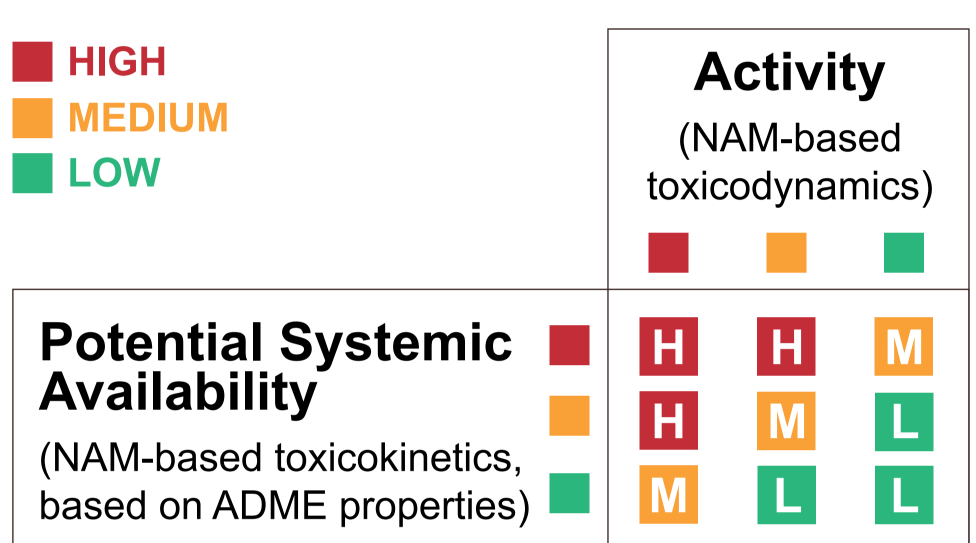
In 2023, the European Partnership for Alternative Approaches to Animal Testing (EPAA) invited the research community developing New Approach Methodologies (NAMs) to submit NAM-based solutions to inform the development of a future classification system for systemic toxicity in humans.

PRINCIPLE OF EQUIVALENT PROTECTION: MAKE THE SAME DECISIONS, NOT NECESSARILY THE SAME PREDICTIONS



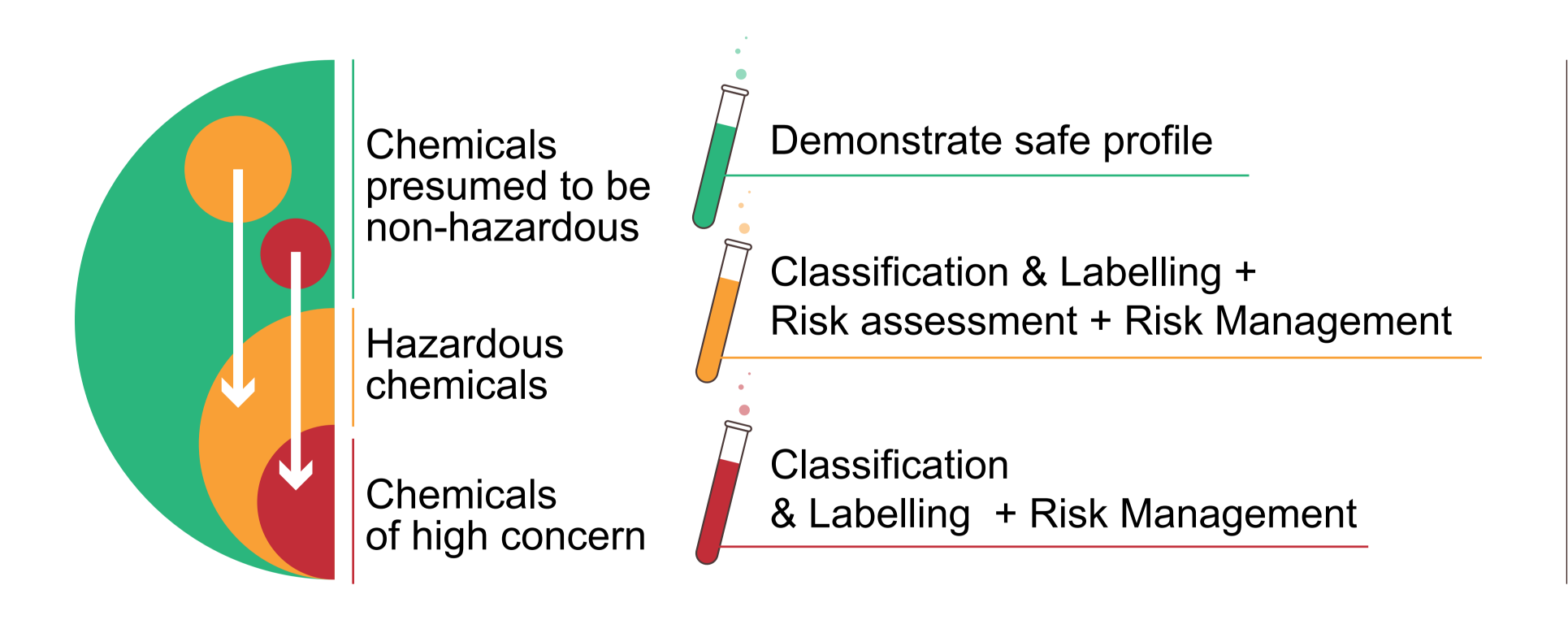
Assign chemicals to groups 1-3 (low, medium & high concern)

Existing data for already classified chemicals (high & medium concern) are used to calibrate the classification scheme resulting in equivalent protection





NAMS CAPTURE CHEMICALS CURRENTLY TREATED AS GREEN, BUT BASED ON NO OR LIMITED INFORMATION



Besides identifying unknown chemicals of concern, we hope to identify low concern chemicals supporting replacement and sustainable by design.

We received 23 NAM-based solutions to inform the development of a future classification system for human systemic toxicity.

In March we met with all contributors to compare and contrast the different solutions and co-create!

Now we are starting Phase 2 of the EPAA NAM Designathon.

If you are curious about our progress or want to contribute, CONNECT TO US HERE: 自多熟度











Towards an animal testing-free system for industrial chemicals

Identifying the needs and working to replace animal testing within the current chemicals management system

EU chemicals management system

To protect Human health and the environment, we rely on a horizontal generic approach with some fundamental elements:



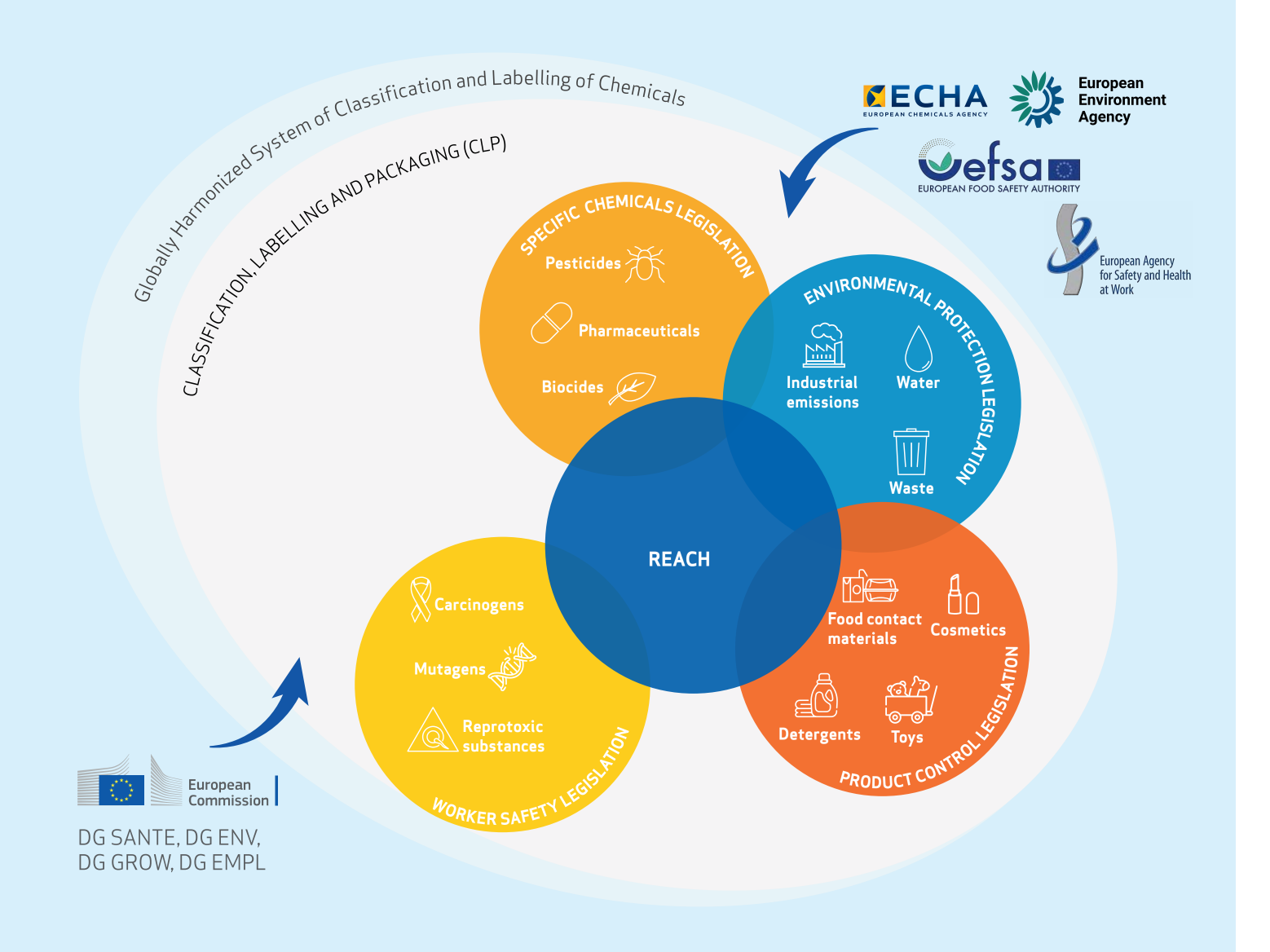
defined hazard classes

 $\Rightarrow \square$ clear criteria for consistent classification

standard information requirements for hazard assessment

quality data for decision making

consistent regulatory actions within chemicals legislation



Critical needs

To achieve an animal testing-free regulatory system, we need to consider elements of chemical assessment:



Hazard identification NAMs should allow a conclusive outcome on the (lack of) hazard



Hazard characterization NAMs should identify hazard based on new information (e.g. at molecular/cellular level)



Extrapolation NAMs data should lead to set safety levels, to communicate the hazard and assess the risks

1. to close major gaps

2. to gain experience and confidence

3. to propose a NAMs compatible system



Identify critical needs for an to animal free

Apply already available NAMs efficient transition more effectively in the current system



Re-think the system to enable NAMs fully compatibility

Now

system

Short-term

Mid-term

Long-term



Areas for development



Application of **toxicokinetics** in chemical assessments



Development of improved in silico methods (e.g. QSARs) for lower tier endpoints (e.g. Acute oral toxicity)



Utilisation of NAMs to support read-across and grouping



Inclusion of 'omics in current test guidelines for higher tier endpoints (e.g. Repeated dose toxicity)

Learn more



Read ECHA's report on the use of alternatives to testing on animals



Read ECHA's report on the key areas of regulatory challenges



Alternatives to animal testing.

https://echa.europa.eu/animal-testing-under-reach



We protect health and the environment through our work for chemical safety https://echa.europa.eu





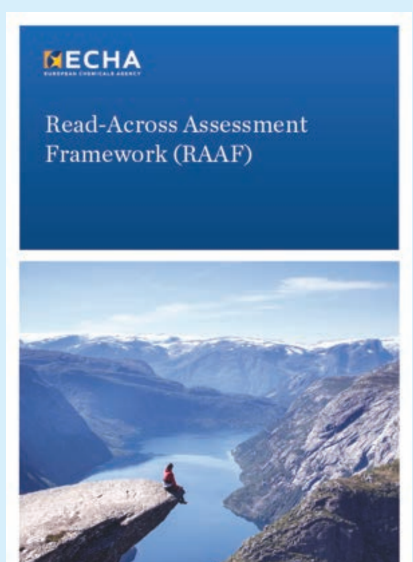
ECHLA Activities to promote New Approach Methodologies - NAMs

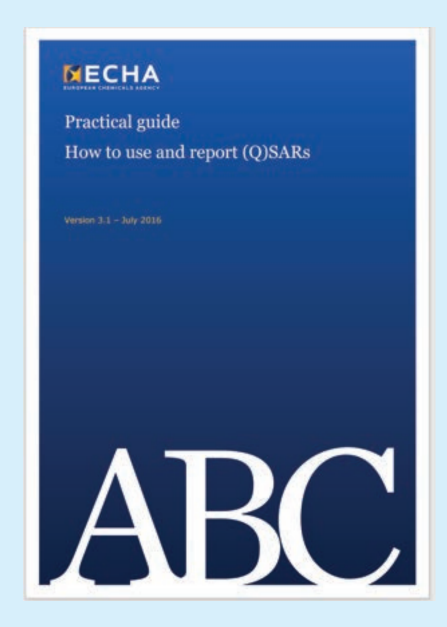
ECHA is committed and active in promoting alternatives to animal testing

Guidance and standardisation

- We provide practical guidance and training to registrants
- We support the development of harmonised guidance and assessment frameworks at OECD







Collaboration

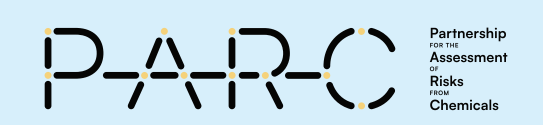
We collaborate in research projects and case studies

Globally

Within EU Research consortia

With academia and industry

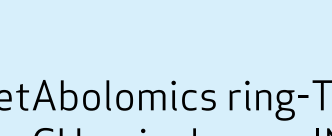
















MetAbolomics ring-Trial for CHemical groupING (MATCHING)

Making data available

We facilitate access to non-confidential data in IUCLID format



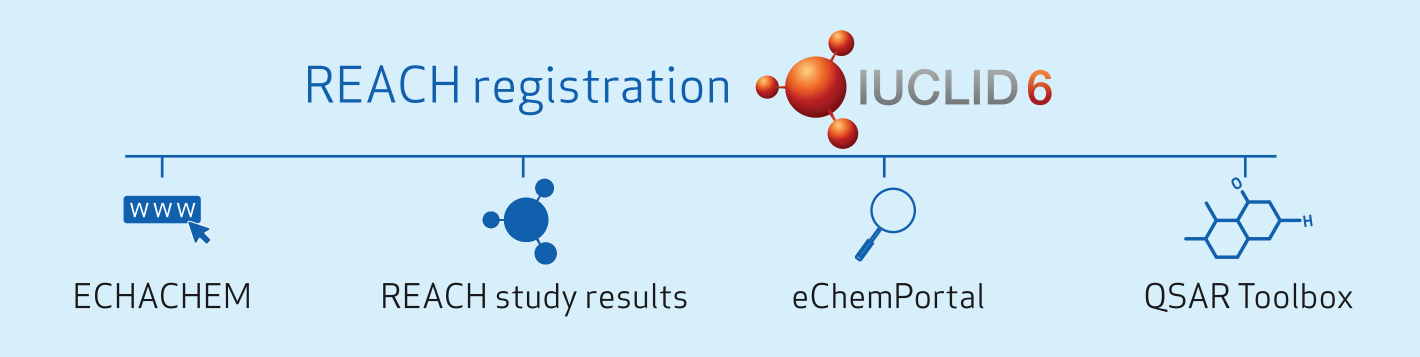
Development of predictive computational models.



Correlation and concordance analyses



Safety data sheets and classification and labelling of chemicals



ECHLA projects

We invest in services related to methodological developments of NAMs related to addressing critical needs in regulatory hazard assessment



Developments of QSARs and their assessment framework



Guidance on sample cryopreservation for omics measurement



Better utilisation of 'omics to support read-across and grouping



Toxicokinetics for industrial chemicals

Learn more

Read ECHA's workshop report on New Approach Methodologies



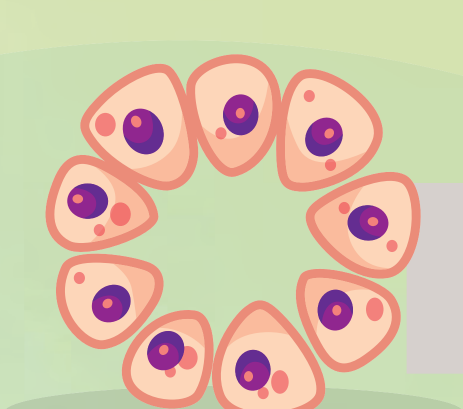


We protect health and the environment through our work for chemical safety https://echa.europa.eu





ALTERNATIVES TO ANIMAL TESTING



THE COMMISSION INVESTED

>€1
billion

> 300 projects

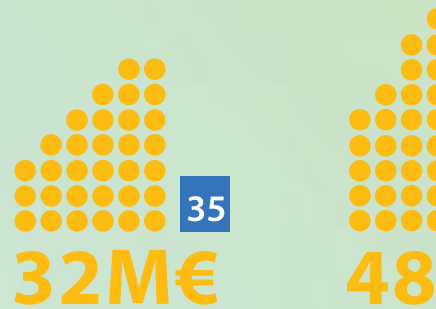


Over the last two decades, the European Commission has substantially supported alternatives to animal testing

AVERAGE ANNUAL BUDGET



FP5
5th Framework
Programme



FP66th Framework Programme

84 48 M€

FP77th Framework
Programme

76M€

H2020 Horizon 2020

Projects supported by the European Commission



WHAT DO WE WANT TO DO WITH IT?

Changing the paradigm in safety testing of chemicals and drugs and developing efficient complementary tools for biomedical research

WHY THEY ARE IMPORTANT?

HUMAN-RELEVANT

Improve predictability and robustness of studies for scientists and regulators

VERSATILE
Answer questions that current methods cannot address

Animal protection: replace/reduce/refine animals (3Rs)

SUSTAINABLE/AUTONOMOUS
Independent of shortage/quality with animals

WHY IS UPTAKE OF ALTERNATIVES TO ANIMAL TESTING SLOW?



Limited awareness of and confidence in Alternatives to animal testing from regulators

Insufficient availability of Alternatives to animal testing to cover complex regulatory endpoints or all aspects of biomedical sciences

Lack of investment in Member States to fund additional Alternatives to animal testing and dedicated infrastructures

Lack of resources in Member States to qualify/standardize/validate
Alternatives to animal testing

Limited coverage in university teaching

IT TAKES TIME ...

Example of in vitro method for skin senzitisation From FP6 SENS-IT-IV project to GARD Method (Genomic Allergen Rapid Detection)

START 2005

SENS-IT-IV

Development

END 2011 VALIDATION (EURL ECVAM)

2017-2021

EU LEGISLATION

2023

Regulatory Approval

GARD Method

Market

2014
SPIN-OFF
Lund University

2022 OECD Test Guideline



for the development of animal-free safety assessment, and expand it beyond

INCREASED DIALOGUE WITH STAKEHOLDERS (INDUSTRY, REGULATORS, CROs)

H2020

Horizon Europe



Animal-free Safety
Project cluster for Implementation
of novel Strategies



Partnership for the Assessment of Risk from Chemicals

Innovative Health Initiative

DISCOVER FUNDING OPPORTUNITIES:





PARTNERSHIP FOR THE ASSESSMENT

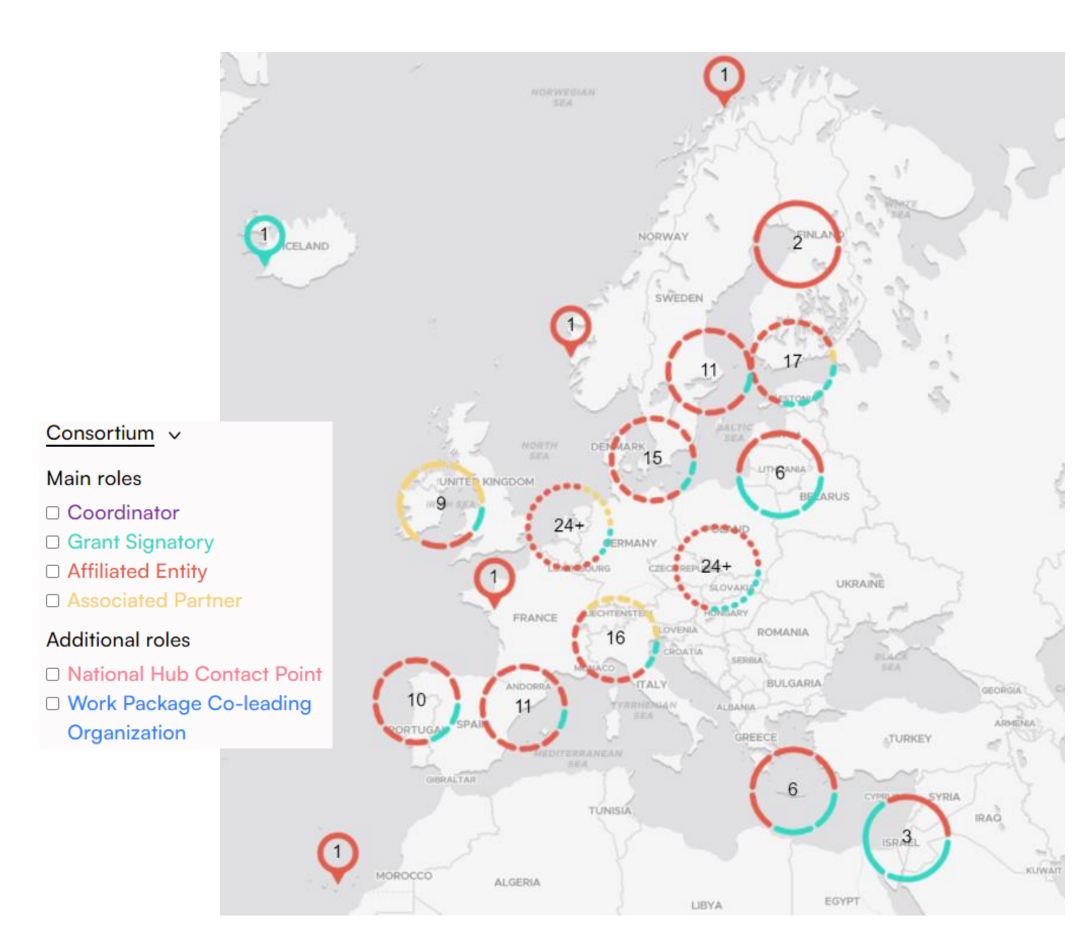
OF RISKS FROM CHEMICALS

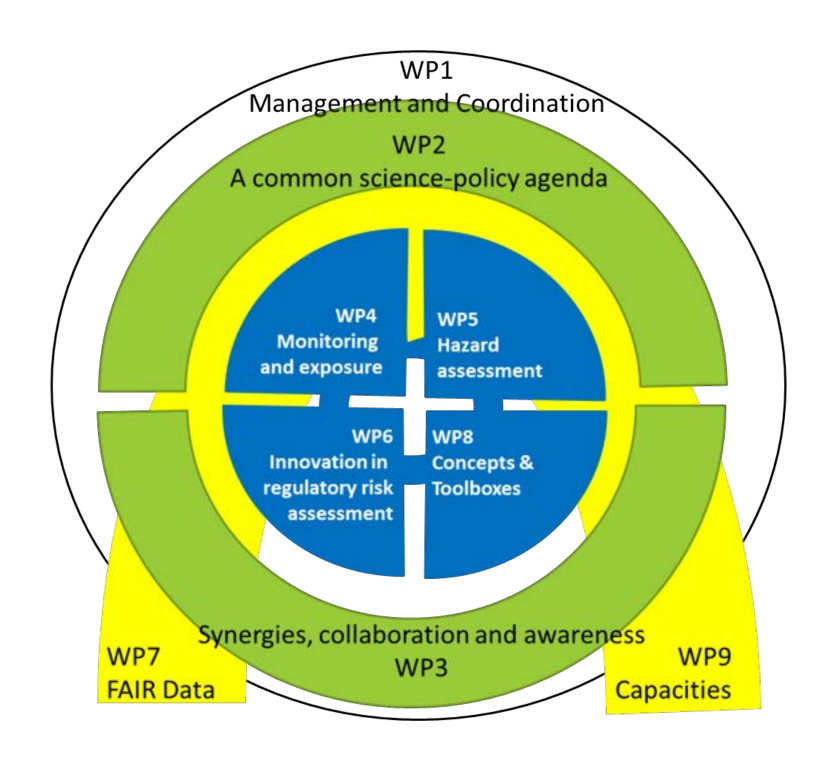


PARC Coordination Team, T2.2 co-leaders ANSES, BfR

PARC in a nutshell

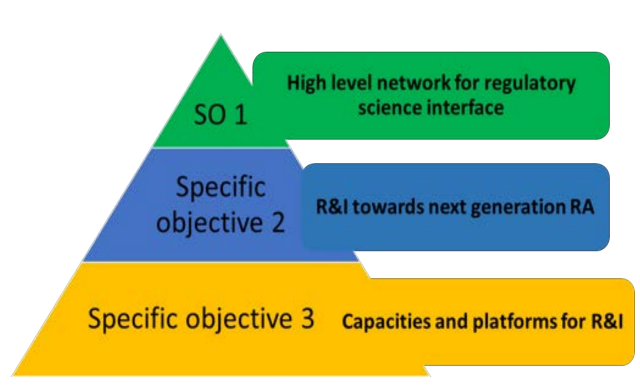
- Public-public European partnership under Horizon Europe
- 7 years length launched in May 2022
- Cofunded 50/50 by the European Commission and PARC partners
- 400M€ budget, max 200M€ reimbursed by the European Commission
- 198 partners from 28 countries 23 member states, 3 assiciated countries, 2 non associated countries (Switzerland, UK), 3 European Agencies (EFSA, ECHA et EEA)

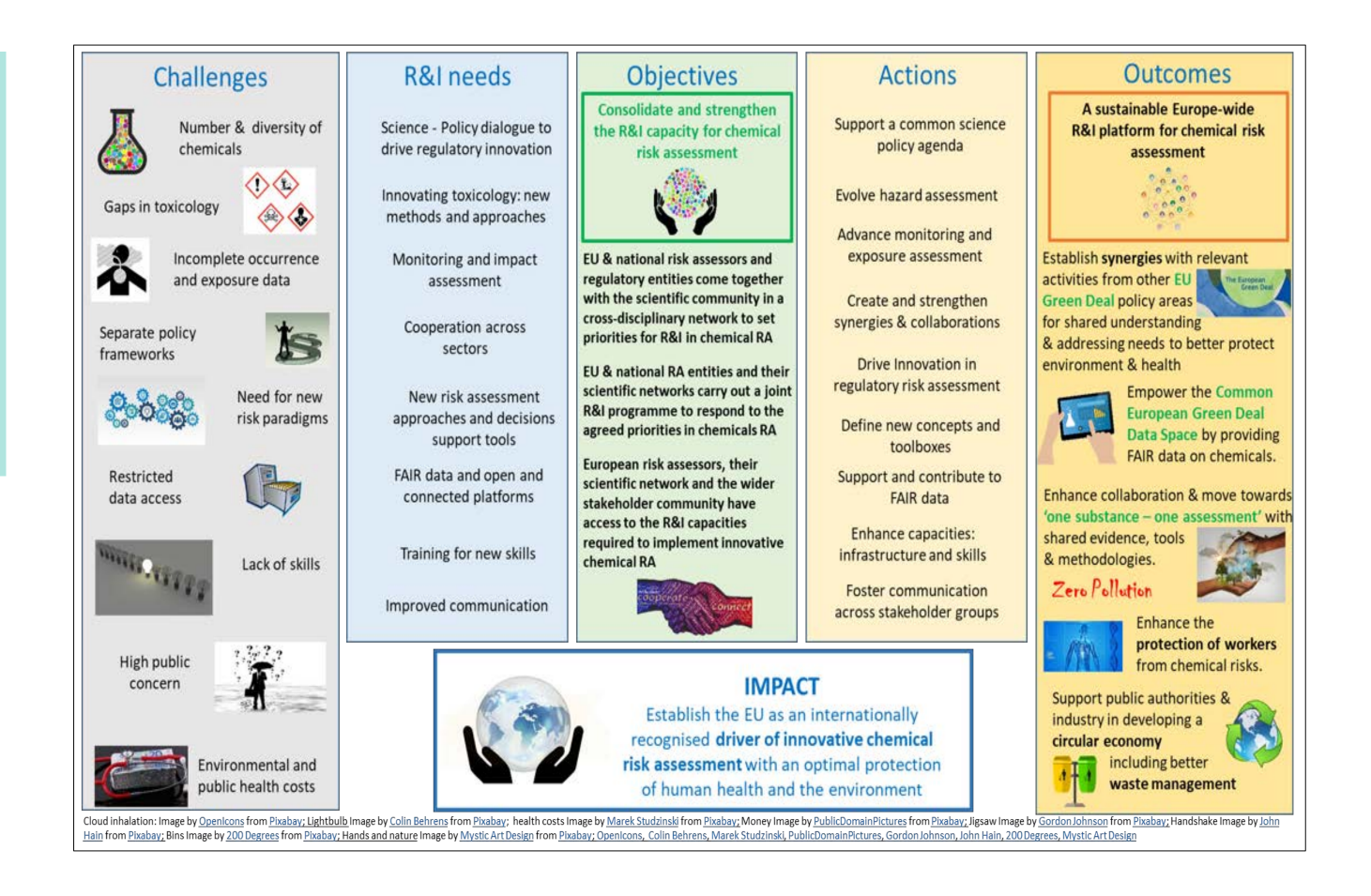




PARC actively contributes to the Chemicals Strategy for Sustainability:

- Dialogue at the EU level,
- Development of a strong network
- Innovative methods and tools





From scientific innovation to policy: NGRAroute

- To provide a concrete and applicable roadmap proposal for implementing Next-Generation Risk Assessment (NGRA) as the default approach to chemical risk assessment in EU chemicals legislation
- In scope: all European chemicals legislation with a risk assessment component of its own. Human health and environmental risk assessment.

Workshop on Commission roadmap for phasing out animal testing in chemical safety assessments (Dec 11/12, 2023)

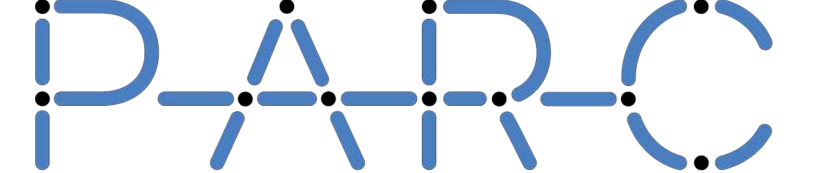
 Ongoing collaboration with EU Commission and EPAA, roadmap proposal expected for 2025

A new co-operative space for the risk assessment community

PARCopedia

Knowledge management and community platform for chemical risk assessment professionals in and beyond PARC













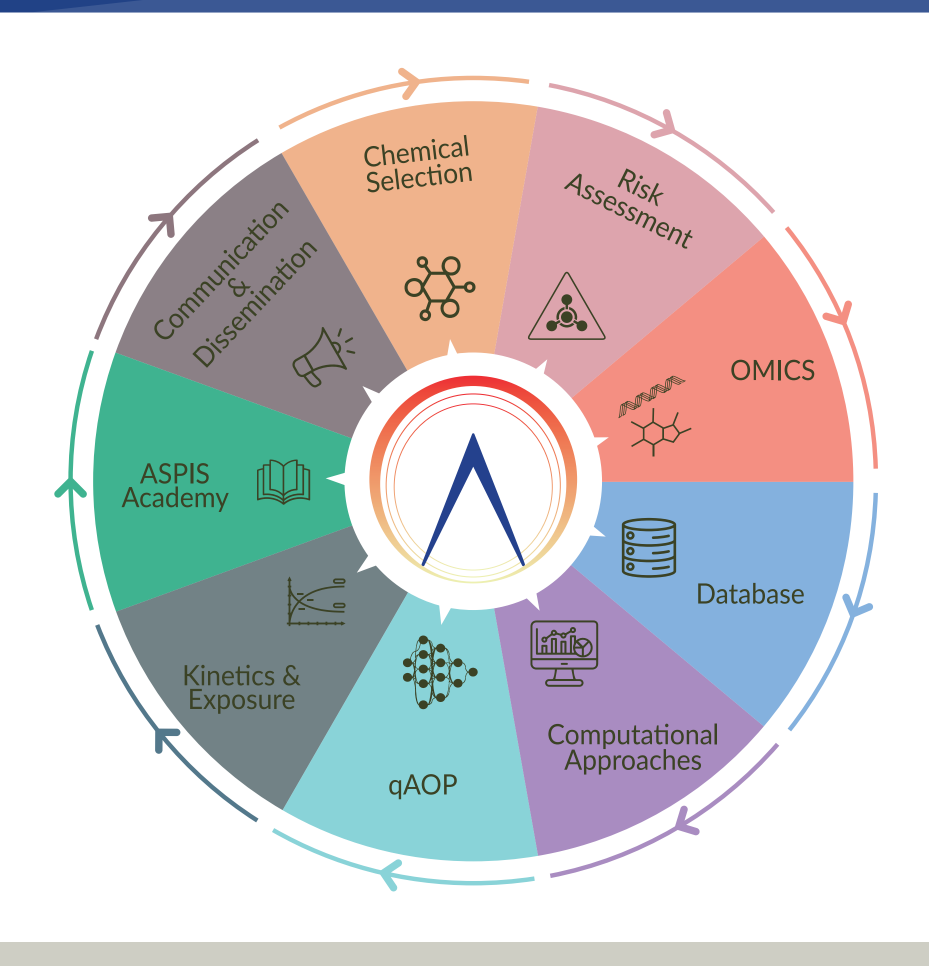




ASPIS - Animal-free Safety Assessment of Chemicals: Project Cluster for Implementation of Novel Strategies

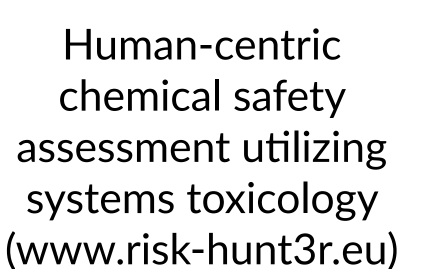
J. Freedman¹, E. Benfenati, F. Busquet, F. Caiment, G. Callegaro, M. Cronin, G. Ecker, S. Escher, B. Hardy, G. Hayot, B. Islam, N. Keith, N. Kramer, E. Kuchovska, L. Ladeira, T. Luechtefeld, M. Luijten, R. Martínez, G. Palucca, E. Roggen, S. Scholz, J. Shaw, B. van de Water, K. Veltman, M. Vinken, P. Xia, H. Yang and J. Colbourne.

Working groups



Consortium







Synthesizing toxicology knowledge to support next-generation risk assessment (ontox-project.eu)



Leveraging evolutionary diversity to reveal the molecular basis of toxicity (precisiontox.org)

Mission

To establish a next generation risk assessment (NGRA) framework based on new approach methodologies (NAMs), encompassing in vivo to in silico technologies. Its goal is to unite three distinct ideas to better understand chemical toxicity and provide, together, innovative methods of assessing and regulating hazardous chemicals without traditional toxicity testing using laboratory animals.

Background

ASPIS is a confluence of three Horizon 2020-funded projects: **PrecisionTox**, **ONTOX** and **RISK-HUNT3R**. It represents Europe's effort towards the sustainable, animal-free, and reliable chemical risk assessment of tomorrow. It includes more than 70 institutions across the European Union, United Kingdom and United States.

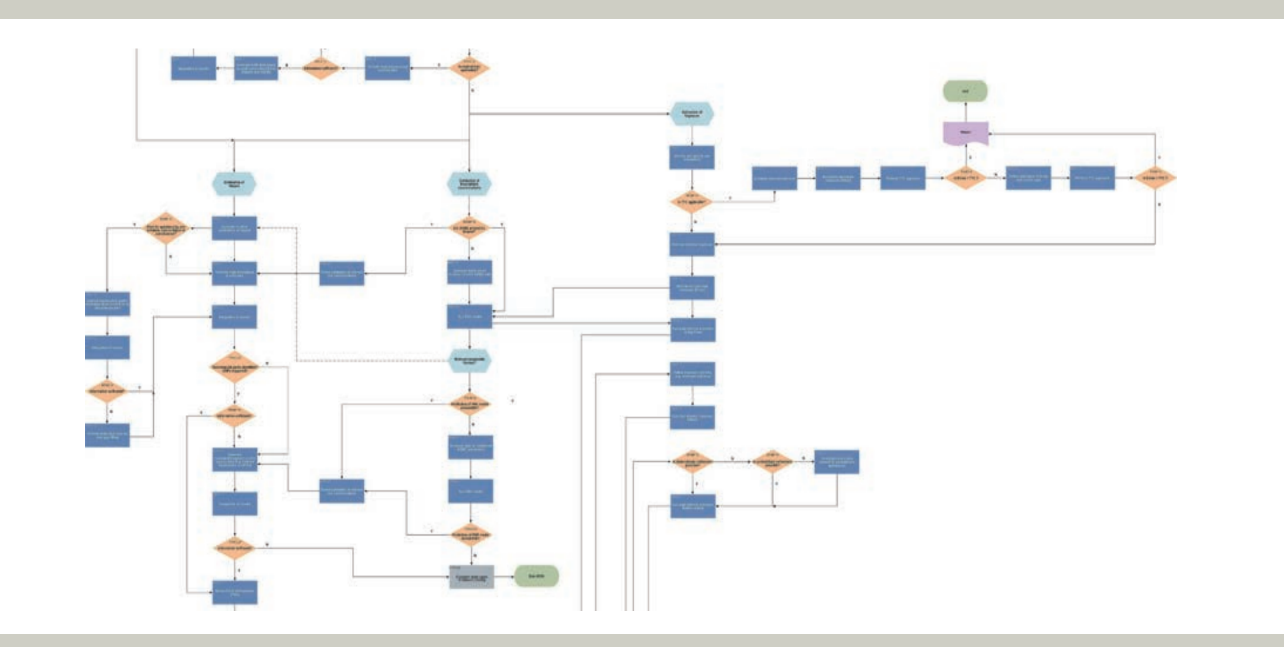
Activities and Perspectives

- ASPIS is a unique partnership consolidating the regulatory science research community by producing innovative and pragmatic solutions for industry, policy makers (EU Commission) and regulators (PARC)
- ASPIS takes an active role in the societal aspects by generating tools to replace and reduce the use of laboratory animals in regulatory science
- ASPIS contributes to the discussion on the EU Commission roadmap to phase out animal testing, as well as participating at EPAA designathon 20-22 March 2024

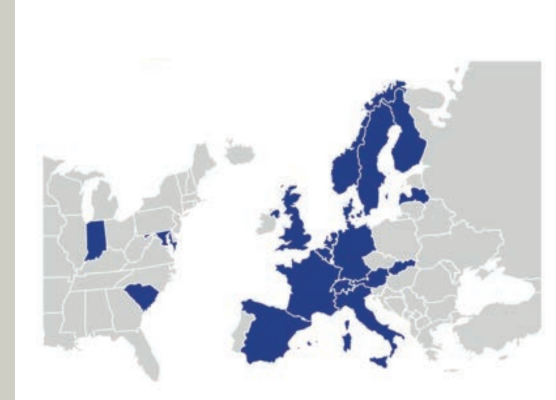
Inter-ASPIS Activities

ASPIS Next Generation Risk Assessment framework (ASPA)

- ASPIS is developing case studies to operationalize NGRA, by developing a well-guided workflow for safety assessment of chemicals
- ASPA defines a tiered approach on what tools/methods to use; at which steps to obtain and evaluate data; and how to put data into a context of a hazard or risk assessment scenario.
- ASPA defines a decision logic with one entry but multiple exit points, activating/deactivating specific modules, and prioritizing and filtering of information.
- Currently being applied to steatotic chemicals and DNTs



ASPIS Academy



Established in March 2023, the ASPIS Academy is a network of >120 early-stage researchers (ESRs) from 65 partners in the EU, UK and USA interested in developing and using New Approach Methodologies (NAMs) in toxicology. The goal of the Academy is build a viable ESR network focused on the use of NAMs for chemical risk assessment, to improve the careers of ESRs through training, championing their needs while setting up a space where ideas and dreams of a new generation of young scientists can shine. Additionally, creating an ASPIS legacy by training a new generation of young scientists.

Programs



In person trainings at ASPIS Open Symposia, project meetings, and online webinars and workshops. This includes science communication, effective presentations, AI in Risk Assessment and Summer School.



Involvement of ESRs in ASPIS working groups. Goals include gaining specialized insights, expanding network, exposure to diverse perspectives, and co-authorship opportunities.



Mentors and mentees work together to build a path to success, innovation, and excellence in toxicology

Career development sessions and mentorship pairs across the three consortia.



ESR mobility facilitation among the cluster partners. Goals include sharing of knowledge and expertise and accelerating collaborations.



at EUROTOX 2024

1 For additional information contact the ASPIS Working

Group Coordinator at Jon.Freedman@wormtox.org

Join us at the ASPIS Open Symposium





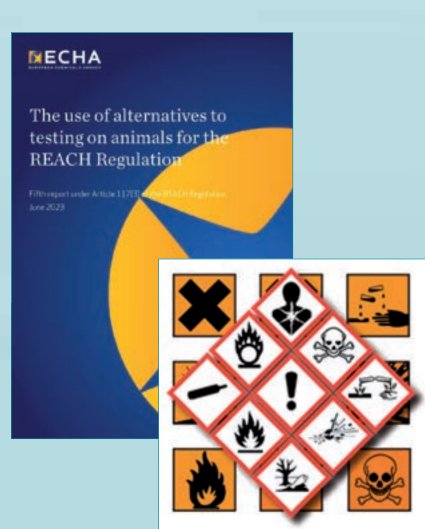
The European detergents industry is committed to closing the gap between science and regulation to prevent unnecessary animal testing

- No animal tests are carried out on finished detergent products.
- A.I.S.E. has a long history of pro-active product stewardship projects, e.g. on safe handling and use of enzymes.
- The safety of detergent ingredients is regulated under REACH. A.I.S.E. advocates for the proper use of New Approach Methodologies (NAMs) in the revision of REACH, which offer opportunities to prevent unnecessary animal testing.
- A.I.S.E. and its members contribute to activities on NAMs through refinement of use and exposure and other alternative approaches to animal testing.

The role of enzymes in determinant to influence the control of the

REGULATORY CHALLENGES FOR DETERGENT INGREDIENTS INCLUDE:

- Current & upcoming changes to REACH (hazard data PBT/vPvB / PMT/vPvM and safety assessment)
- REACH surfactant category registrations (endpoint data requests vs. use of Weight of Evidence (WoE) and read-across)
- Classification schemes used:
- Globally Harmonised System of Classification and Labelling of Chemicals (GHS)
- EU Classification, Labelling and Packaging of Substances and Mixtures (CLP) Regulation (EC No 1272/2008) new hazard classes introduced by Commission Delegated Regulation (EU) 2023/707



A.I.S.E.'s commitment to finding alternatives to animal testing is implemented through our scientific engagement with partners in a variety of research initiatives.



Non-animal science in regulatory decisions for chemical safety

UN Sub-Committee of Experts on the GHS Informal Working Group on Non-Animal Testing Methods



• Classification of health hazards, in particular skin corrosion/irritation, serious eye damage/irritation and respiratory or skin sensitisation



Collaboration with other formulating sectors to contribute to the successful implementation of the requirements of the REACH and CLP Regulations



with



- Eye Damage & Irritation
- Physiologically-Based Kinetic (PBK) Model validation for surfactants
- Assessing Membrane-Water Partitioning of Surfactants
- Suitability of the RTgill-W1 cell line assay (OECD 249) for surfactants



A.I.S.E. has new & ongoing projects on alternative tests for eye and skin irritation as well as several scientific publications.









ABOUT A.I.S.E.

A.I.S.E. has been the voice of the detergents and maintenance products industry to EU regulators since 1952, representing over 900 companies supplying household and professional cleaning products and services across Europe. A.I.S.E. has pursued scientific research to finding alternatives to animal testing for over three decades. www.aise.eu/alternativestoanimaltesting



www.aise.eu





Modern science to protect people and environment

A commitment to progressively transition to New Approach Methods (NAMs) and phase out animal testing

The chemicals industry is developing non-animal testing methods and innovating new and sustainable chemicals.



Computational models



In vitro assays & predictive screening





Exposure-driven and regulatoryrelevant concepts in (eco)toxicology





Cefic's Long-range Research Initiative (LRI) advances scientific assessment of chemicals safety through rigorous research.



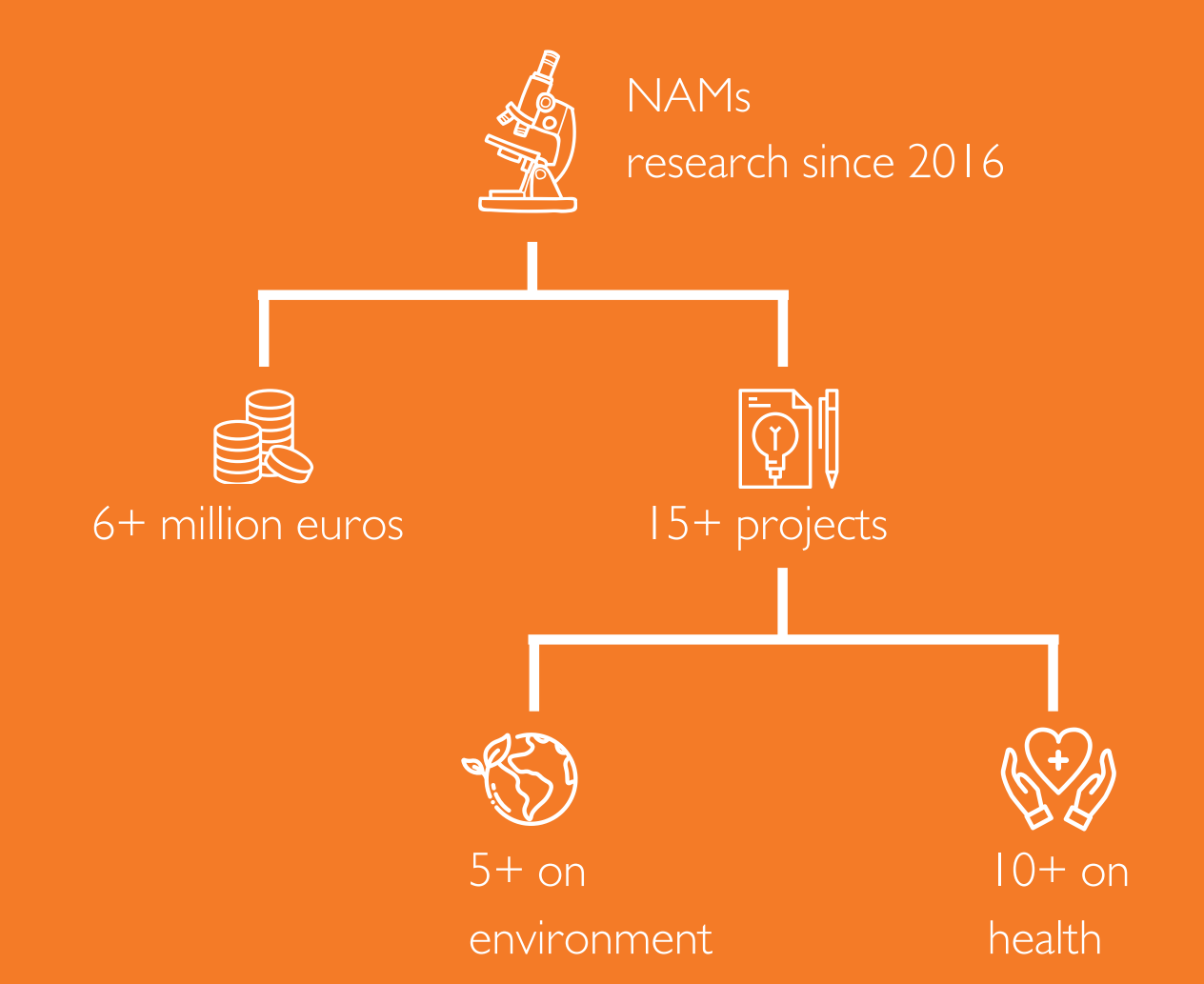
25 years of funding



200+ projects



100 million euros for research, studies & awards



Find out
more about
Cefic-LRI
projects:



What is needed to build confidence and increase deployment of NAMs in REACH?

NOW

Regulatory acceptance & policy uptake of available & reliable NAMs.

VISION

Paradigm shift in REACH: Decisions on testing is based on exposure and the extent to which a chemical becomes available to a biological system.



- A safe space for knowledge exchange & to create trust
- International alignment & global standards



Cefic, the European Chemical Industry
Council is the voice chemical companies
across Europe, which provide 1.2 million
jobs and account for 14% of world
chemicals production.

www.cefic.be - info@cefic.be

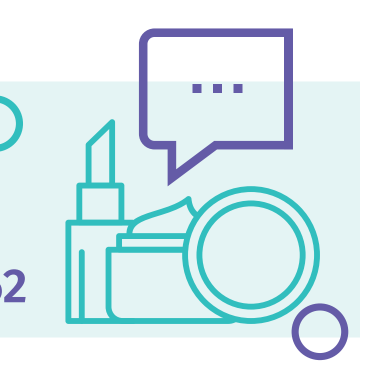


Cosmetics Europe's key achievements in advancing Non - Animal Methods (NAMs)



*** 1**

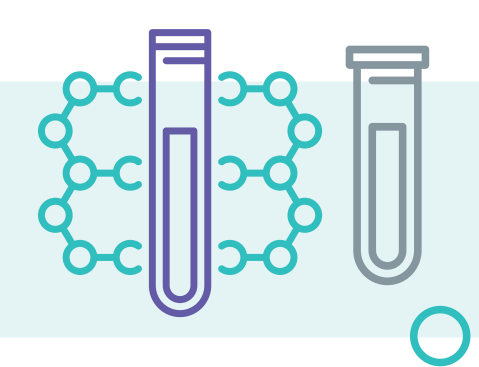
COSMETICS EUROPE, THE VOICE OF THE COSMETICS INDUSTRY SINCE 1962



Cosmetics Europe represents cosmetics and personal care manufacturers, as well as associations advocating for our industry at national level, right across Europe.



FOR OVER 30 YEARS, THE COSMETICS SECTOR HAS LED THE ADVANCEMENT OF NAMS



Ensuring the safety of cosmetic products and promoting the development and acceptance of alternatives to animal testing are key priorities for our sector. The industry's dedication to upholding the highest safety standards is reflected in the continuous refinement of testing and assessment capabilities for all our cosmetic product ingredients.



THE LONG-RANGE SCIENCE STRATEGY (LRSS): RESEARCH AND SCIENCE PROGRAMME ON NAMs (2016 -2022)

The LRSS programme drove our specific research programme on alternatives to animal testing. It was founded on multidisciplinary partnerships between cosmetics companies and other groups that have a deep interest in NAMs and Next Generation Risk Assessment (NGRA), including the international regulatory community, validating agencies, academia, research institutes and industry partners.



THE GOAL OF THE LRSS PROGRAMME

TO ENABLE ANIMAL-FREE SAFETY ASSESSMENTS OF COSMETIC INGREDIENTS

3 PILLARS

Filling critical science gaps related to specific endpoints of toxicity and the use of NAMs to understand biological mechanisms of adverse effects caused by a substance to which the body or environment is exposed to.



Areas in which The LRSS programme scientists evaluated the safety of cosmetic ingredients using NAMs:

- Skin Sensitization
- Eye Irritation and Severe Eye Damage
- Genotoxicity/ Mutagenicity
- Toxicokinetics (absorption, distribution, metabolism and elimination (ADME))
- Toxicodynamics
- Dermal and Inhalation Exposure
- Environment

Implementation of NAMs in NGRAs, to show, through case studies, that safety assessments are possible on multiple toxicity endpoints, especially systemic toxicity.



- Case studies have been instrumental in the LRSS program, guiding the practical implementation of scientific workflows
- These case studies demonstrated the feasibility of safety assessment solely relying on non-animal data, integrating findings from in silico and in vitro NAMs
- LRSS led ~20 case studies to highlight the practical application of NAMs in tiered NGRAs

Development and validation of NAMs as well as confidence building in the NGRA framework to advocate uptake by industry and gain regulatory acceptance.



- Several scientific contributions from Cosmetics Europe were included in the Scientific Committee on Consumer Safety (SCCS) Notes of Guidance (NoG) for testing the safety of ingredients
- Only within the LRSS program, Cosmetics Europe delivered >15 OECD documents

Outside LRSS, Cosmetics Europe submitted the 1st regulatory NGRA dossier to evaluate the systemic safety of Benzophenone- 4 (Under evaluation by the SCCS)

COLLABORATION, PARTNERSHIPS AND THE FUTURE



- Cosmetics Europe continuously engages in partnerships with organizations (EPAA, ICCR) and collaborations (RiskHunter, ONTOX, PARC, VHP4Sfatey) with similar initiatives and goals.
- Cosmetics Europe played a key role in the establishment of the International Collaboration on Cosmetics Safety (ICCS), a global initiative focused on advancing the adoption of animal-free assessments of cosmetics, and their ingredients, for human health and environmental safety. Since its establishment, Cosmetics Europe has been actively involved in various projects of the organisation in its capacity as an ICCS member.





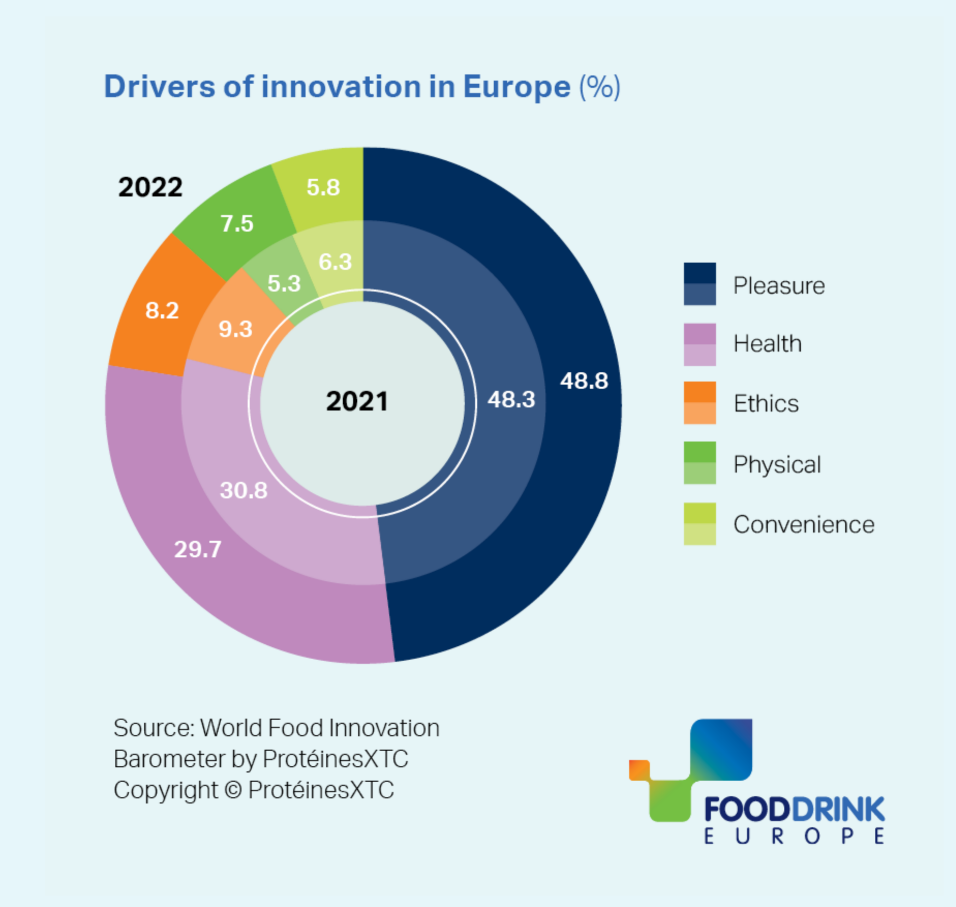






Towards the integration of new approach methodologies (NAMs) in food safety risk assessments

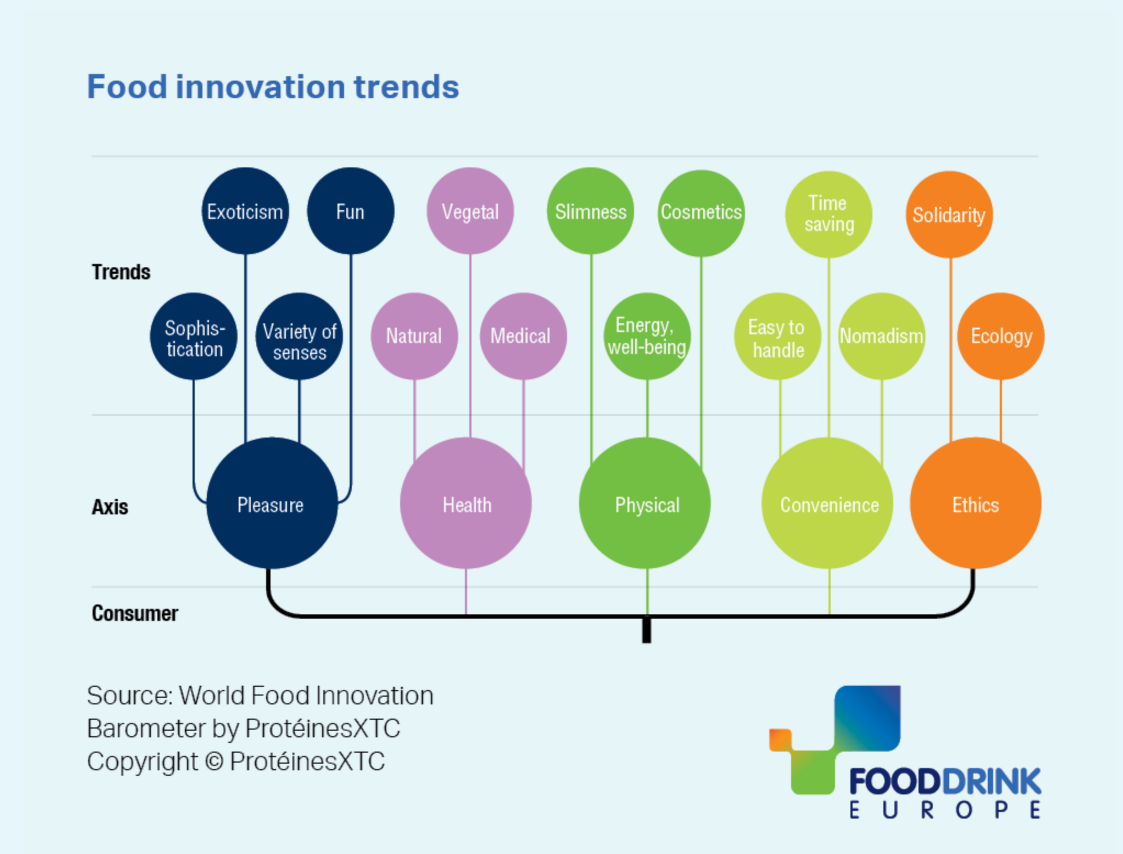




Novel foods and food ingredients can help meet consumer needs and expectations, improve health, and contribute to sustainable food systems.

All 'food and food ingredients' must be demonstrated to be safe following their consumption and prior to their placement on the market.

Application of NAMs can improve the relevance of data available for food safety assessment and avoid unnecessary animal use.



WHY?

- Next generation or non-animal approaches have advantages over animal approaches e.g. mechanistic investigations
- Risk assessment questions can be better tailored to the human situation
- Political and societal calls to phase out animal testing
- Increasing popularity of vegan foods
- Current EU Regulatory framework allows for the use of NAMs
- Need for more consistency in use of NAMs for food safety risk assessments
- Time associated with traditional studies impacts competitiveness of European businesses and speed to market
- EFSA guidance documents in food safety risk assessments must reflect the new science and provide flexibility for use of NAMs



OUR POSITION

FoodDrinkEurope¹ calls for the routine application of NAMs in scientific and regulatory food safety assessment in Europe²:

Need to close the gap between modern safety science and regulatory requirements

- More flexibility to use fit-for-purpose NAMs within regulatory framework
- Focus on specific human centric safety questions and avoid check list of animal studies

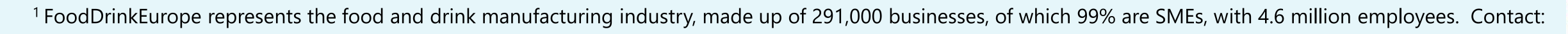
Innovation requires the right regulatory setting and a NAMs approach is important for innovation in the food industry

- NAMs provide more informative high-quality data (e.g. mechanistic understanding), and can be quicker to perform
- Risk of innovation in the EU food sector slowing or stopping without regulatory acceptance of NAMs

The food industry welcomes partnership with the broad base of stakeholders

- Partnership with academia, industry and regulatory authorities to progress adoption of NAMs
- Opportunity to feedback on latest EFSA guidance documents and pilots launched for integrating NAMs in risk assessment







² Position paper co-signed by FoodDrinkEurope, EU Specialty Food Ingredients, the Association of Manufacturers and Formulators of Enzyme Products (AMFEP), the European Flavour Association (EFFA), and the International Organization of the Flavor Industry (IOFI)

Accelerating the transition

Towards animal-free, sustainable innovation



Close cooperation

Enhancing regulatory application of NAMs through close cooperation with stakeholders and collaboration in various scientific coordination platforms.



for Alternative Approaches to Animal Testing







Fragrance initiatives

Actively supporting the development and regulatory acceptance of animal-free approaches for the advancement of fragrance safety.



Research Institute Fragrance
Materials (RIFM)- An independent
nonprofit scientific organization
that uses historical data, NAMs and
advanced exposure tools to the
greatest extent possible to
evaluate the safe use of fragrance
ingredients. Realistic exposure data
in consumer products are derived
via the Creme-RIFM Aggregate
Exposure Model. RIFM's safety
assessment conclusions inform the
IFRA Standards.



IFRA Standards – IFRA's safe use policy restricting the use of certain fragrance ingredients based on the RIFM safety assessments.



International Dialogue for the Evaluation of Allergens (IDEA) - An industry-driven multi-stakeholder forum developing a NAM-based framework for Quantitative Risk Assessment for skin sensitisation. Find out more: ideaproject.info



Safe and Sustainable by Design (SSbD) - IFRA supports the SSbD framework development as active partner within IRISS, to use NAMs in fragrance innovation and the evaluation of alternatives.



