

EU scenario on alternatives in cosmetic safety evaluation

State-of-play, impact & recommendations

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- I. EU measures affecting animal testing
- **II. State-of-the-art of science**
- **III. Impact of EU regulatory measures**
- **IV. Recommendations**



I. EU measures affecting animal testing

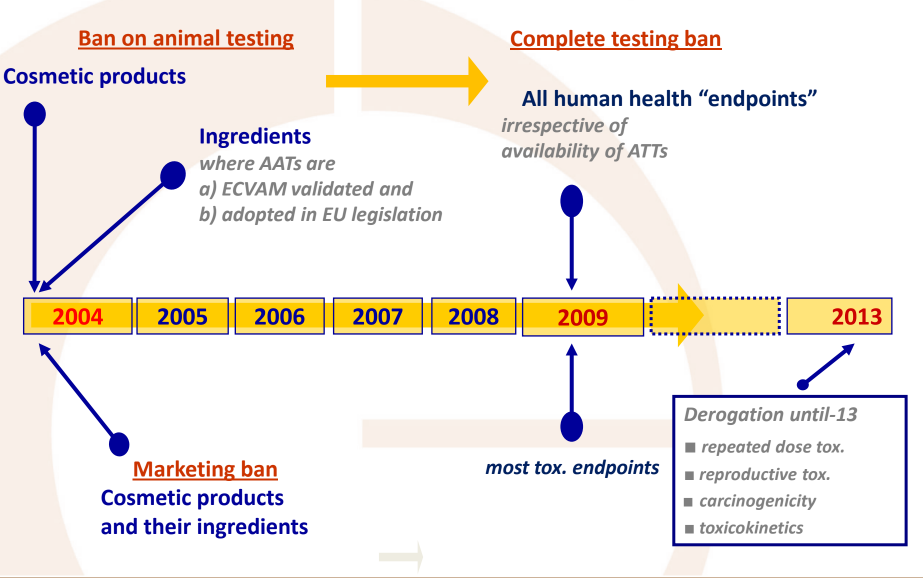
EU measures affecting animal testing



Testing and marketing bans "for the purpose of the Cosmetics Regulation" successively entering into force between 2004 and 2013

EU measures affecting animal testing





European Commission Interpretation



In 2013, the European Commission clarified its own interpretation of the scope of the ban explaining that it is not a blanket ban:

«The Commission considers that animal testing that has clearly been motivated by compliance with non-cosmetics related legislative frameworks should not be considered to have been carried out 'in order to meet the requirements of this Directive/Regulation' »

«The Commission considers that the marketing ban is triggered by the reliance on the animal data for the safety assessment under the Cosmetics Directive/Regulation, not by the testing as such . In case animal testing was carried out for compliance with cosmetics requirements in third countries, this data cannot be relied on in the Union for the safety assessment of cosmetics. »

European Commission Interpretation

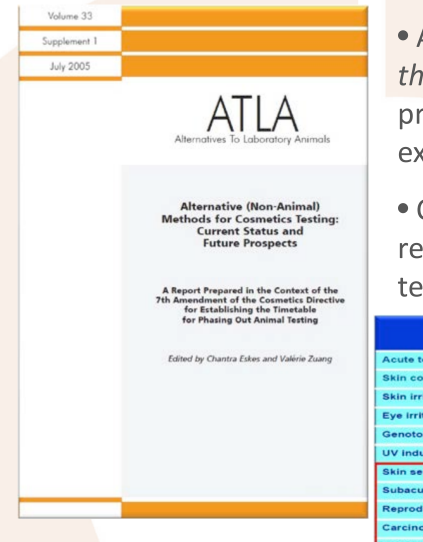


- Products are banned only if ingredients or products tested for cosmetics purposes (in or outside of the EU)
- Possibility to market products if ingredients tested for multiple purposes (in and outside of the EU)
- The marketing ban is only triggered by the use of data
- In exceptional circumstances: derogation for existing, nonreplacable ingredient of which the use raises a specific human health problem (substantiation required)



II. State-of-the-art of science

Timetables for phasing out Animal tests

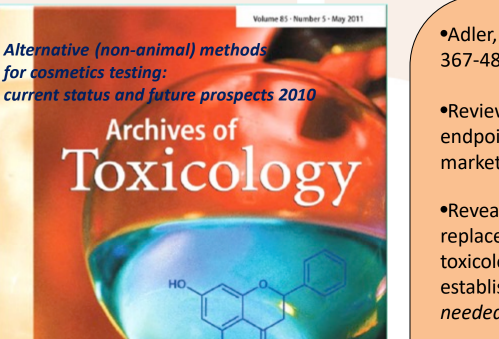


• A first technical report on timetables for the phasing out of animal testing was prepared by nominated independent experts in 2003 and published in 2005

 Covered all toxicological endpoints relevant for cosmetics (ingredients) testing

Toxicological Endpoint	Cut-off-dates according article 4a "Testing Ban"	Cut-off-dates according article 4a "Marketing Ban"	2004 EU estimate Availability of Alternatives
Acute toxicity	11 March 2009	11 March 2009	> 2014
Skin corrosion	11 March 2009	11 March 2009	-
Skin irritation	11 March 2009	11 March 2009	> 2009
Eye irritation	11 March 2009	11 March 2009	2010
Genotoxicity / Mutagenicity	11 March 2009	11 March 2009	> 2016
UV induced effects (Allergy)	11 March 2009	11 Mach 2013	> 2016
Skin sensitization	11 March 2009	11 March 2013	2016 - 2018
Subacute / Subchronic toxicity	11 March 2009	11 March 2013	not estimated
Reproductive toxicity	11 March 2009	11 March 2013	not estimated
Carcinogenicity	11 March 2009	11 March 2013	not estimated
Toxicokinetics and Metabolism	11 March 2009	11 March 2013	> 2016

Current status and Outlook beyond the Cosmetics Europe ban



•Adler, S. *et al.* (2011) Arch. Toxicol., 85: 367-485.

•Review considered the toxicological endpoints important for the 2013 marketing ban deadline

•Revealed that the scientific basis to fully replace animal testing for the five toxicological key areas is still not established (*additional time beyond 2013 needed*)

•Confirmed that it could take at least another 7 – 9 years for the replacement of some of the current *in vivo animal* tests necessary for the safety assessment

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OECD Guidelines for Toxicology studies Cosmetics Europe

N° Type

- 401 Acute Oral Toxicity
- 402 Acute Dermal Toxicity
- 403 Acute Inhalation Toxicity
- 404 Acute Dermal Irritation/Corrosion
- 405 Acute Eye Irritation/Corrosion
- 406 Skin Sensitisation
- 407 Repeated Dose 28-Day Oral Toxicity Study in Rodents
- 408 Repeated Dose 90-Day Oral Toxicity Study in Rodents
- 409 Repeated Dose 90-Day Oral Toxicity Study in Non-Rodents
- 410 Repeated Dose Dermal Toxicity:90-Day
- 411 Subchronic Inhalation Toxicity: 90-Day
- 412 Subacute Inhalation Toxicity: 28-Day Study
- 413 Subchronic Inhalation Toxicity: 90-Day Study
- 414 Prenatal Developmental Toxicity Study
- 415 One-Generation Reproduction Toxicity
- 416 Two-generation Reproduction Toxicity Study
- 417 Toxicokinetics
- 418 Delayed Neurotoxicity of Organophosphorus Substances Following Acute Exposure
- 419 Delayed Neurotoxicity of Organophosphorus Substances: 29-Day Repeated Dose Study
- 420 Acute Oral toxicity Fixe Dose Procedure
- 421 Reproduction/Developmental Toxicity Screening Test
- 422 Combined Repeated Dose Toxicity Study with the Reproduction/Developmental Toxicity Screening Test
- 423 Acute Oral Toxicity Acute Toxic Class Method
- 424 Neurotoxicity Study in Rodents
- 425 Acute Oral Toxicity: Up-and-Down Procedure
- 426 Developmental Neurotoxicity Study
- 427 Skin Absorption: In Vivo Method
- 428 Skin Absorption: In Vitro Method
- 429 Skin Sensitisation: Local Lymph Node Assay

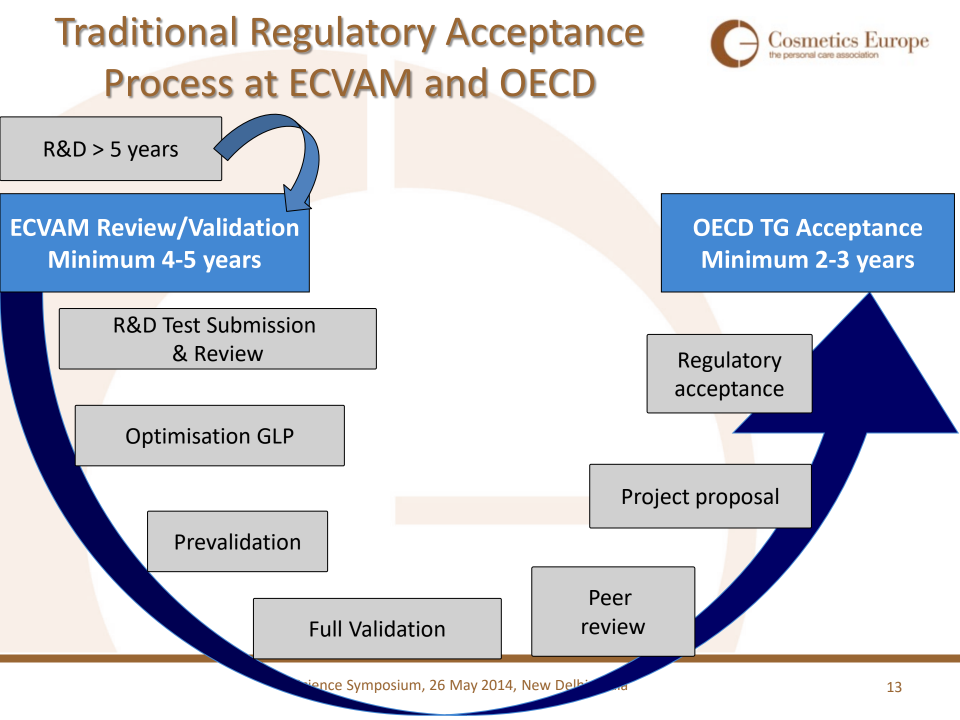
N° Type

- 430 In Vitro Skin Corrosion: Transcutaneous Electrical Resistance Test (TER)
- 431 In Vitro Skin Corrosion: Human Skin Model Test
- 432 In Vitro 3T3 NRU Phototoxicity Test
- 435 In Vitro Membrane Barrier Test Method for Skin Corrosion
- 436 Acute Inhalation Toxicity Acute Toxic Class Method
- 437 Bovine Corneal Opacity and Permeability Test Method for Identifying Ocular Corrosives and Severe Irritants
- 438 Isolated Chicken Eye Test Method for Identifying Ocular Corrosives and Severe Irritants
- 440 Uterotrophic Bioassay in Rodents: A short-term screening test for oestrogenic properties
- 441 Hershberger Bioassay in Rats: A short-term Screening Assay for (Anti)Androgenic Properties
- 451 Carcinogenicity Studies
- 452 Chronic Toxicity Studies
- 453 Combined Chronic Toxicity/Carcinogenicity Studies
- 455 Stably Transfected Human Estrogen Receptor-α Transcriptional Activation Assay for the Detection of Estrogenic Agonist-Activity of Chemicals
- 471 Bacterial Reverse Mutation Test
- 472 Genetic Toxicology: Escherichia coli, Reverse Assay
- 473 In Vitro Mammalian Chromosome Aberration Test
- 474 Mammalian Erythrocyte Micronucleus Test
- 475 Mammalian Bone Marrow Chromosome Aberration Test
- 476 In Vitro Mammalian Cell Gene Mutation Test
- 477 Genetic Toxicology: Sex-Linked Recessive Lethal Test in Drosophilia melanogaster
- 478 Genetic Toxicology: Rodent dominant Lethal Test
- 479 Genetic Toxicology: In Vitro Sister Chromatid Exchange assay in Mammalian Cells
- 480 Genetic Toxicology: Saccharomyces cerevisiae, Gene Mutation Assay
- 481 Genetic Toxicology: Saccharomyces cerevisiae, Mitotic Recombination Assay
- 482 Genetic Toxicology: DNA Damage and Repair, Unscheduled DNA Synthesis in Mammalian Cells In Vitro
- 483 Mammalian Spermatagonial Chromosome Aberration Test
- 484 Genetic Toxicology: Mouse Spot Test
- 485 Genetic Toxicology: Mouse Heritable Translocation Assay
- 486 Unscheduled DNA Synthesis (UDS) Test with Mammalian Liver Cells In Vivo

OECD Guidelines for Toxicology studies O Cosmetics Europe

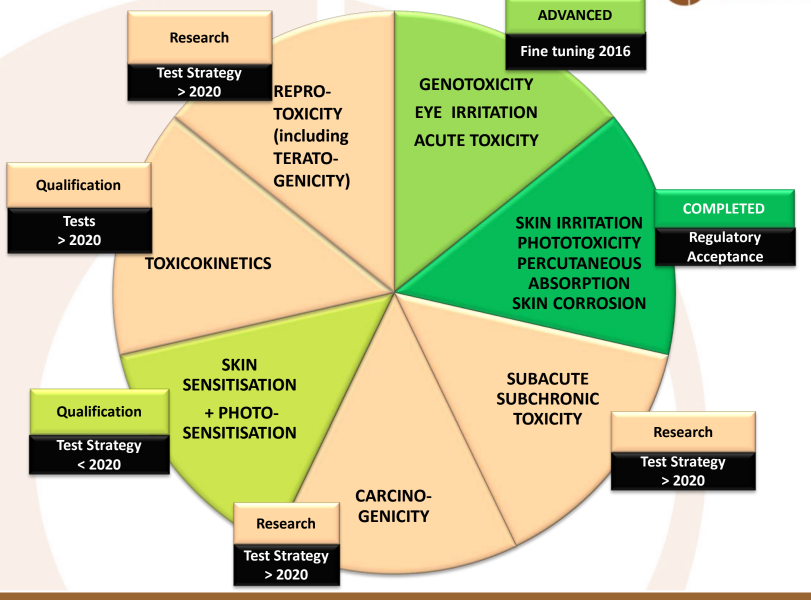


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402	Acute Dermal Toxicity	431	In Vitro Skin Corrosion: Human Skin Model Test
403	Acute Inhalation Toxicity	432	In Vitro 3T3 NRU Phototoxicity Test
404	Acute Dermal Irritation/Corrosion	435	In Vitro Membrane Barrier Test Method for Skin Corrosion
405	Acute Eye Irritation/Corrosion	436	Acute Inhalation Toxicity - Acute Toxic Class Method
406	Skin Sensitisation	437	Bovine Corneal Opacity and Permeability Test Method for Identifying Ocular Corrosives and Severe Irritants
407	Repeated Dose 28-Day Oral Toxicity Study in Rodents	438	Isolated Chicken Eye Test Method for Identifying Ocular Corrosives and Severe Irritants
408	Repeated Dose 90-Day Oral Toxicity Study in Rodents	440	Uterotrophic Bioassay in Rodents: A short-term screening test for oestrogenic
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409	Repeated Dose 90-Day Or <mark>al Toxicity Study in Non-Rodents</mark>	441	Hershberger Bioassay in Rats: A short-term Screening Assay for (Anti)Androgenic
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410	Repeated Dose Dermal Toxicity Repeat	45	nalniy contains <i>in</i>
411	Subchronic Inhalation Toxicity: 90-Day	452	Chronic Toxicity Studies
412	Subacute Inhalation Toxicity 28-Day Study Subchronic Inhalation Toxicity VIVO studies. These co	453	Combined Chronic Toxicity/Carcinogenicity Studies
413	Subchronic Inhalation Toxic ty VIVO SLUUIES. IIIESE CO		Cally TS GUILESstUp Receptor-α Transcriptional Activation Assay for
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	Toxicity Screening Test		
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425	Acute Oral Toxicity: Up-and-Down Procedure	482	Genetic Toxicology: DNA Damage and Repair, Unscheduled DNA Synthesis in
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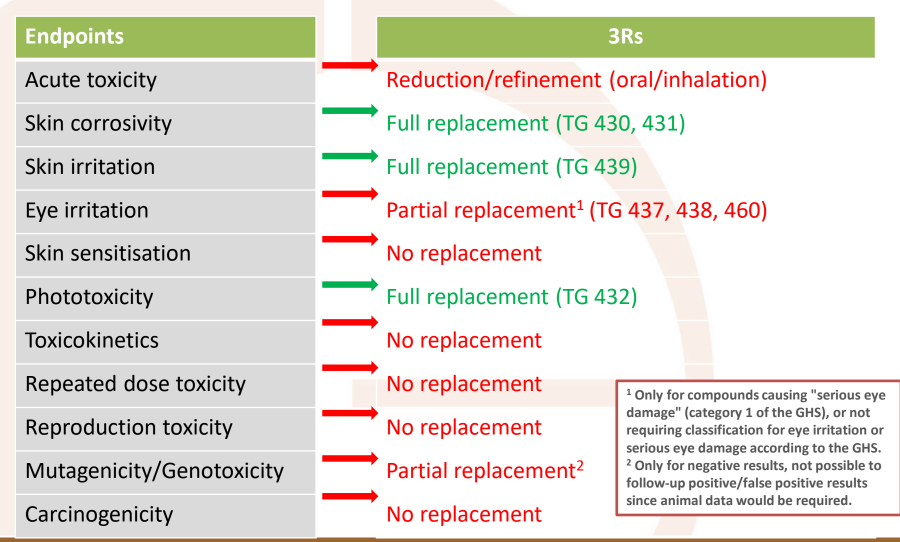
Status of Science





Available validated Alternative Methods for Human Health Safety Assessments in the SCCS Notes of Guidance





Science - Concluding remarks



- Only for a limited number of toxicological endpoints, replacement methods are available and validated (Report by the European Commission Joint Research Centre (JRC)¹)
- Replacement methods for skin allergy testing potentially available within 2-4 years but for the more complex endpoints a timeline is difficult to anticipate
- State-of-the-art of regulatory accepted test methods (OECD Test Guidelines²)
- ¹<u>http://ihcp.jrc.ec.europa.eu/our_labs/eurl-ecvam/eurl-ecvam-releases-2013-progress-report-development-validation-regulatory-acceptance-alternative-methods</u>
- ²<u>http://www.oecd-ilibrary.org/environment/oecd-guidelines-for-the-testing-of-</u> <u>chemicals-section-4-health-effects</u> 20745788

Science - Concluding remarks



More time and efforts are needed to develop the complete set of alternative methods:

- New internationally agreed tools and testing approaches required for successful development of replacements
- New streamlined validation criteria required for mechanistic tests; not realistic that a single screening test should take 6-8 years to reach regulatory acceptance