OECD QSAR Toolbox and readacross Webinar

22 October 2014, 4:00pm BST





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Today's webinar

This webinar will cover:

- § Use of integrated approaches to testing and assessment and adverse outcome pathways to organize existing information and plan a non-animal testing strategy;
- § How QSARs and read-across can be used to meet REACH requirements;
- § Use of the OECD QSAR Toolbox;
- § Future research projects.







- **§ Dr. Amy Clippinger**, PETA International Science Consortium Ltd
- § Dr. Grace Patlewicz, DuPont
- § Dr. Mark Cronin, Liverpool John Moores University
- § Chair: Emma Chynoweth, Chemical Watch





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Questions

- § Please submit questions during the webinar using your chat box
- § Any unanswered questions can be raised on our Forum following the webinar: <u>http://forum.chemicalwatch.com/</u>





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Upcoming Webinars

Webinar 2: Skin Irritation and Corrosion

November 11, 2014,

11am ET, 4pm GMT

- Emilia Costin, Institute for In Vitro Sciences
- **Costanza Rovida**, CAAT Europe and REACH Mastery

Webinar 3: Serious Eye Damage and Eye Irritation

December 4, 2014

11am ET, 4pm GMT

- Kim Norman, Institute for In Vitro Sciences
- João Barroso, EURL ECVAM

Please contact the PETA International Science Consortium, Ltd., for assistance in avoiding animal testing

<u>pisc@piscltd.org.uk</u>





Non-testing approaches: How can (Q)SARs, read-across and the OECD QSAR Toolbox help in addressing REACH 2018?

Grace Patlewicz, DuPont, Newark, DE, USA Mark Cronin, Liverpool John Moores University, England





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Context – REACH Deadline

- 31st May 2018 marks the deadline for registration of phase-in substances manufactured or imported at 1-100 tonnes per year
- The information requirements for these tonnage bands are described in Annexes VII and VIII of the legal text
- This impacts 10,000s of substances





Context – REACH Legislation

- To address financial and animal welfare concerns, REACH explicitly expresses the need to use non-testing approaches to reduce the extent of experimental testing
- Article 25(1) states: "in order to avoid animal testing, testing on vertebrate animals for the purposes of this Regulation shall be undertaken only as a last resort."
- Article 13(1) states: "Information on intrinsic properties of substances may be generated by means other than tests, provided that the conditions set out in Annex XI are met. In particular for human toxicity, information shall be generated whenever possible by means other than vertebrate animal tests, through the use of alternative methods, for example, *in vitro* methods or qualitative or quantitative structure-activity relationship models or from information from structurally related substances (grouping or read-across)..."





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Aim(s) of this Webinar

- To provide an introduction of how non-testing approaches can be exploited as part of an Integrated Approach to Testing and Assessment (IATA) to address the information requirements within these Annexes
 - Focusing on *in silico* approaches
- To highlight advances in the Tox21 field that could in future impact the type of data that are generated to fulfil these information requirements





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Outline

- The IATA construct and related terms
 - Definitions
 - IATA under REACH
- Non-testing approaches
 - Definitions
 - (Q)SARs
- Chemical grouping, category and analogue approaches
 - Definitions
 - Considerations associated with read-across
 - Data gap filling within category/analogue approaches
- Future directions AOPs
 - Read-across enhancement
 - (Q)SAR and IATA development
- Take home messages
- Useful links





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Integrated Approaches to Testing and Assessment (IATA)

"IATA is a means of organising and analysing all the available relevant data on a given substance or group of substances coupled with mechanistic, exposure, and dosimetry information where possible, to focus testing when needed and facilitate an assessment conclusion" – OECD definition

"Integrated Testing Strategies (ITS) are approaches that integrate different types of data and information into the decision-making process. In addition to the information from individual assays, test batteries, and/or tiered test schemes, integrated testing strategies may incorporate approaches such as weight-of-evidence and exposure/population data into the final risk assessment for a substance" <u>http://www.alttox.org/ttrc/emerging-technologies/its/</u>

In practice:

A means of integrating existing data and non-testing data, determining what new information needs to be generated in order to make a decision





Integrated Testing Strategies (ITS)

- Under REACH, such IATA are termed ITS and one has been described for each of the endpoints of interest
- These ITS can be likened to workflows depicting the different steps of gathering (toxicity) information for a substance in order to evaluate its "fit for purposes" for classification & labelling and/or risk assessment
- Some ITS are more complex than others but the generic building blocks of considering existing data, *in vitro* alternatives, nontesting approaches BEFORE instigating new *in vivo* testing are the same
- Non-testing approaches fit within the context of these ITS schemes and should not be considered in vacuo





Typical Information within an ITS

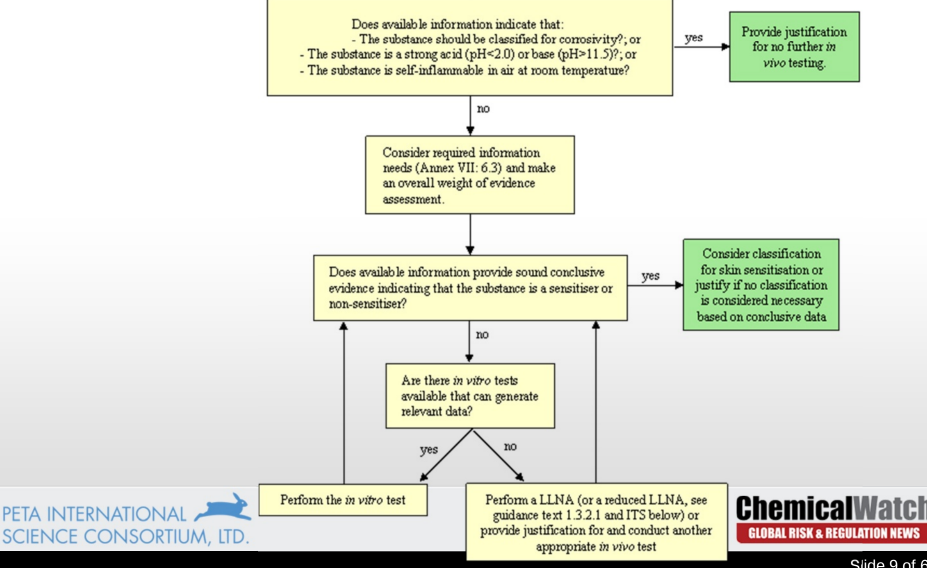
- Historical information on the chemical of interest
 - Non-standard in vivo tests
- Information from "similar" chemicals
- Predictions from other non-testing approaches such as (Q)SAR
- In chemico tests
- In vitro tests
- Molecular biology, -omics
- Exposure, (bio-)kinetics





REACH ITS for Skin Sensitisation

Gather and evaluate existing information (human-, animal-, in vitro-, (Q)SAR, read across and chemical category data) on skin sensitisation according to Annex VI, step 1



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Computational (In Silico) Toxicology

Databases of existing information Category formation (grouping) read-across

Structure-Activity Relationships (SAR)

Quantitative Structure-Activity Relationships (QSAR)

Expert Systems

Bioinformatics Chemoinformatics Biokinetics (PBPK)





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Computational (In Silico) Toxicology



Bioinformatics Chemoinformatics

Biokinetics (PBPK)





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Structure Activity Relationships and Structural Alerts

- A SAR is a (qualitative) association between a chemical substructure and the potential of a chemical containing the substructure to exhibit a certain biological effect
- E.g. Carcinogenicity alerts reflected in the "**Supramolecule**" Ashby and Tennant (1988) Mut. Res. 204:17-115

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(Quantitative) Structure-Activity Relationships ([Q]SARs)

- A (Q)SAR attempts to relate (statistically or otherwise) the activity of one or more molecules to their physico-chemical properties or structural descriptors
- QSAR can be used to predict:
 - Quantitative endpoints e.g. potency
 - Qualitative endpoints e.g. active / inactive





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Collections of (Q)SARs

- An Expert System is a formalised system, usually computerised that enables an end-user to make rational predictions of toxicity based on structure alone
- Expert systems are typically categorised by whether they are underpinned by:
 - empirically based algorithms such as QSARs e.g. TOPKAT
 - knowledge bases such as SARs e.g. Derek Nexus
 - or a hybrid of the two e.g. TIMES





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Regulatory Applications of (Q)SARs

"Packaged mature knowledge for systematic reuse"

- For data gap filling to provide an estimate for a given (eco)toxicity/e-fate/phys chem endpoint in lieu of testing (replacement or supporting information)
- To substantiate waivers or as part of ITS by providing another line of reasoning
- To rationalise spurious results in experimental data since the (Q)SAR is based on a larger body of data, provides a more compelling WoE to rationalise the validity of a potential outlier
- Essential for category development and associated read-across justification - to provide a context of endpoint mechanistic similarity





Using (Q)SARs to Fill Data Gaps

- Under REACH, some of the information requirements within Annexes VII and VIII readily lend themselves to QSAR use
- Examples could include: providing LC₅₀ or EC₅₀ estimates for fish, daphnid, algae toxicity especially for difficult to test compounds such as gases, providing Log Koc and Log Kow estimates, supporting data for mutagenicity endpoints, skin/eye irritation, skin sensitisation...
- However under REACH certain conditions have to be met and specific documentation has to be provided

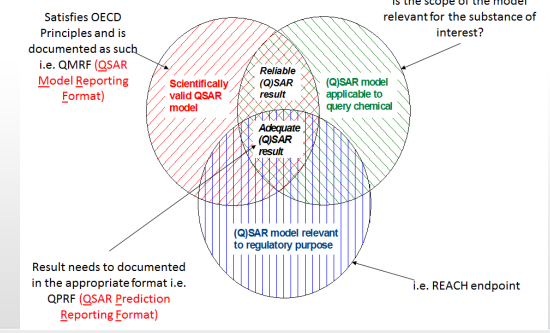




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Annex XI – Use of (Q)SARs

- Results obtained from valid qualitative or quantitative structure-activity relationship models ((Q)SARs) may indicate the presence or absence of a certain dangerous property.
- Results of (Q)SARs may be used instead of testing when the following conditions are met:
 Applicability domain Is the scope of the model







Assessing Scientific Validity: OECD Principles for (Q)SAR Validation

A (Q)SAR should be associated with the following information:

- a defined endpoint
- an unambiguous algorithm
- a defined applicability domain
- appropriate measures of goodness-of-fit, robustness and predictivity
- a mechanistic interpretation, if possible

• Published as OECD guidance





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Other Practical Considerations for (Q)SAR Use

- Is it possible to re-create the (Q)SAR model? what is the availability of the underlying training set, what descriptors were used in the (Q)SAR development?
- To what extent & how can the domain be extracted? What threshold should be set for a substance to be considered within domain? Does that depend on how the prediction is intended to be used?
- What other information exists that might be relevant for the endpoint under consideration (i.e. the ITS) to help determine whether the QSAR estimate should or can be used as a 'true' replacement value or as part of a WoE?





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Assessing Applicability Domain to Determine if the Model is Valid for Use for a Specific Molecule

- Applicability domain may be characterised using:
 - Descriptors
 - Structural features e.g. fragments, fingerprints
 - Metabolic transformations
 - Mechanistic information
- Tools exist to assess applicability domains
 - e.g. LMC Domain Manager, AMBIT Discovery etc.



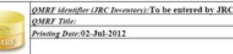


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Documenting the Model: **QSAR Model Reporting Format (QMRF)**

QSAR Model Reporting Format (QMRF) is a harmonised template for summarising and reporting key information on (Q)SAR models, including the results of any validation studies

- The information is structured according to the OECD (Q)SAR validation principles.
- A freely available editor is available
- http://ihcp.jrc.ec.europa.eu/our labs/pr edictive toxicology/gsar tools/QRF
- http://echa.europa.eu/documents/1016 2/13632/information requirements r6 en.pdf



1.QSAR identifier

1.1.QSAR identifier (title): 1.2.Other related models:

1.3.Software coding the model:

2.General information

2.1.Date of QMRF:

- 2.2.QMRF author(s) and contact details:
- 2.3.Date of QMRF update(s):
- 2.4.QMRF update(s):
- 2.5.Model developer(s) and contact details:
- 2.6.Date of model development and/or publication:
- 2.7.Reference(s) to main scientific papers and/or software package:
- 2.8. Availability of information about the model:
- 2.9. Availability of another QMRF for exactly the same model:

3.Defining the endpoint - OECD Principle 1

- 3.2.Endpoint:
- 3.3.Comment on endpoint:
- 3.4.Endpoint units:
- 3.5.Dependent variable:
- 3.6.Experimental protocol:





^{3.1.}Species:

QSAR Prediction Reporting Format (QPRF)

- The QSAR Prediction Reporting Format (QPRF) is a harmonised template for summarising and reporting substance-specific predictions generated by (Q)SAR models
- QPRF requires information on:
 - The substance
 - General information (e.g. date and author)
 - Description of QSAR according to OECD Principles and how it relates to target substance
 - Adequacy (optional)

http://ihcp.jrc.ec.europa.eu/our_labs/predictive_toxicology/qsar_tools/QRF

http://echa.europa.eu/documents/10162/13632/information_requirements_r6_en.pd f





Current Experiences of (Q)SAR Approaches

- As replacements most promising for physicochemical, ecotoxicity and environmental fate properties e.g. Log Kow, acute fish toxicity, ready biodegradability. Much progress has also been made in the area of genetox specifically - Ames mutagenicity and to a large extent on skin sensitisation
- As supporting information in category approaches or as additional information as part of an WoE – most progress has been made with (Q)SARs for endpoints such as skin/eye irritation, or other genotoxicity endpoints
- (Q)SARs for repeated dose toxicity endpoints are not sufficiently evolved to be used as replacements but can play an useful role in supporting read-across within category/analogue approaches





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Category Formation (Grouping) for Read-across

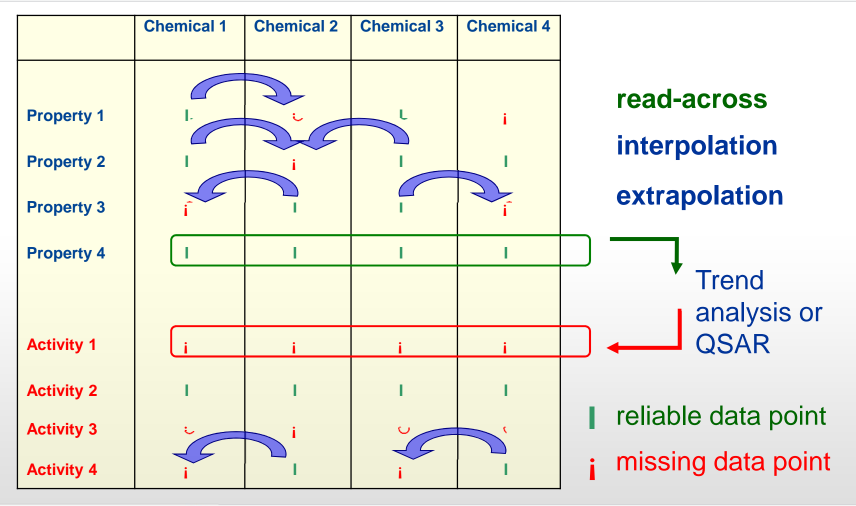
- "Analogue approach" refers to grouping based on a very limited number of chemicals (e.g. target substance) + source substance)
- "Category approach" is used when grouping is based on a more extensive range of analogues (e.g. 3 or more members) and there may be an apparent trend in property
- Read-across describes one of the methods for filling data gaps in either the analogue or category approaches i.e. <u>not to be</u> confused with the "analogue approach"





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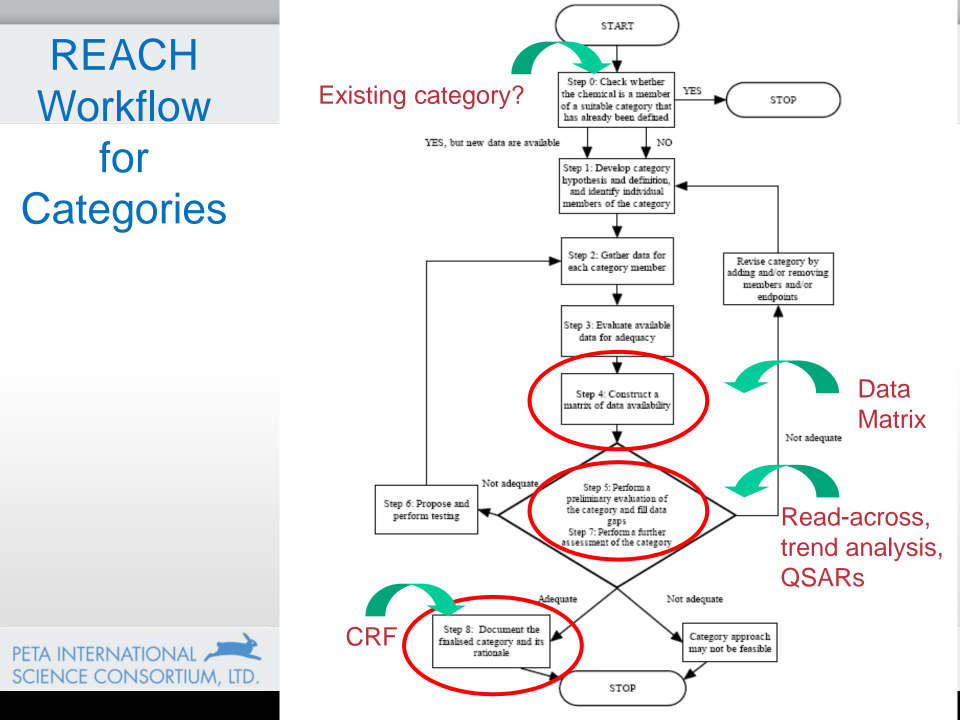
Read-across





ChemicalWatch Global Risk & Regulation News

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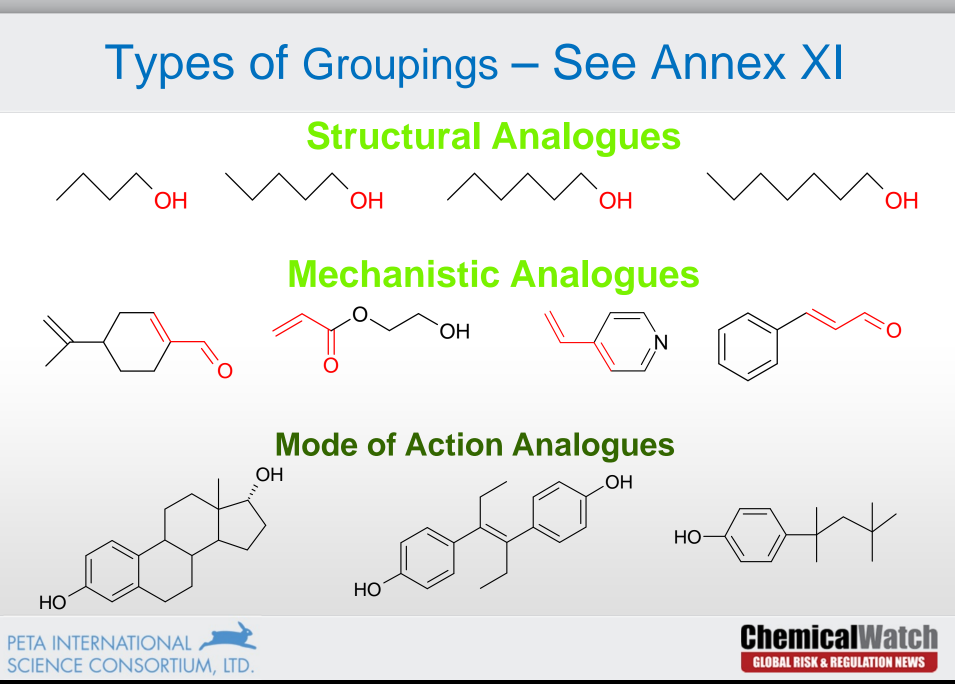
Considerations Before Embarking on a "Read-across"

- How many data gaps? And for which endpoints?
- Legitimate access to sufficient, reliable data?
- Plausible hypothesis for grouping substances and ease and cost of substantiating that hypothesis?
- Accurate and credible assessment of the hazards for the substance in question? Is the scientific confidence sufficient for the purpose required?
- Consequence and cost of the read-across approach not being accepted?





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Types of Groupings

Substances that are **metabolised** to a common molecule Substances that are **degraded** rapidly to common products

• The rationale underpinning the category/analogue approach might be based on 1 or more of these rationales





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Identifying Source Analogues...With Data

Based on own internal company inventory

Using computational tools to help identify potential analogues and in some cases to help evaluate those analogues for their suitability

OECD QSAR Toolbox

ToxMatch

Toxtree

ChemProp

Leadscope

Analogue Identification Method

AMBIT

VITIC

ECHA dissemination databasehas the substance been registered already?





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OECD (Q)SAR Toolbox

- A software tool which facilitates the development, evaluation, justification and documentation of chemical categories for read-across
- Software workflow mimics that described in the OECD and REACH guidance on categories
- Contains regulatory inventories and data plus "profilers" which encode SAR type information which represent molecular initiating events (MIEs) within Adverse Outcome Pathways (AOPs)
- Profilers include those for "DNA Binding", "Protein Binding", "Aquatic toxicity MOAs" etc – hence works best for skin sensitisation, mutagenicity and aquatic toxicity endpoints
- Ongoing development is focusing on how to implement new MIEs and AOPs into the Toolbox to facilitate read-across for repeated dose toxicity endpoints
- First AOP implemented into the OECD Toolbox skin sensitisation





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Is Substance Already a Member of an Existing Category?

 Is there an existing HPV category already available e.g. HPVIS, OECD, OECD Toolbox

http://webnet.oecd.org/hpv/ui/Default.aspx

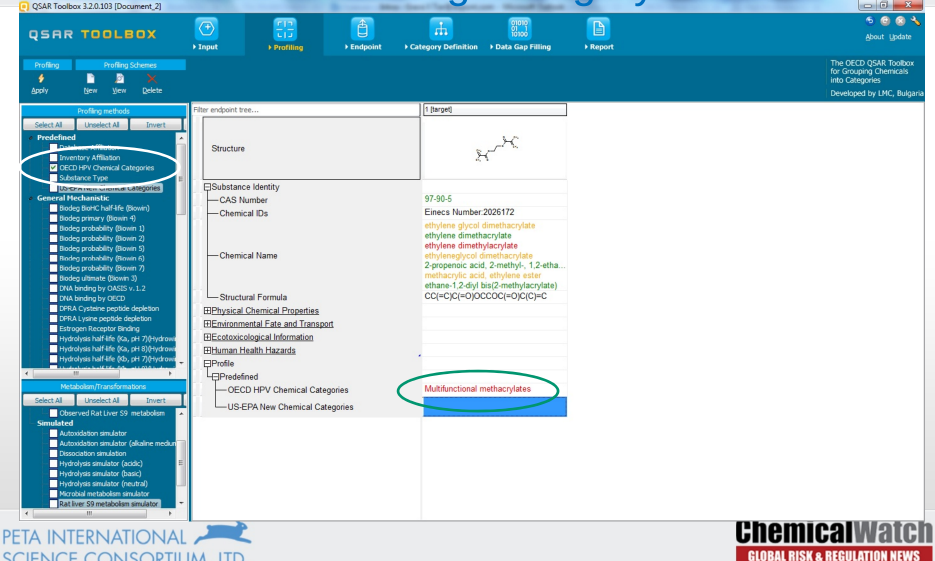
http://www.epa.gov/hpvis/

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	The High Production Volume Information System (HPVIS) is a database that provides access to health and environmental effects information obtained	Highlights	Click on an item 🗸		Open all / Toggle all
HPV Challenge Home	through the High Production Volume (HPU) Challenge. This program "challenges" companies to make this data publicly available on chemicals produced or	On November 6, 2008, EPA	>Home		open an / Toggle an
	imported into the United States in quantities of 1 million pounds or more per year.	posted a Risk Based Prioritization document for	> Search		
HPVIS Home	On this Web site, HPVIS enables users to search for summary information, test plans, and new data on HPV chemicals as they are received by the Agency.	elemental mercury in certain	SIDS contacts	Acid Chloride Category (4)	
About HPV Chemical	Currently, the HPVIS database contains over 340 submissions, representing almost 900 chemical substances, either as a single chemical submission or as a member of a chemical category.	products and substitutes (PDF) (7 pp., 67K8, about PDF), and	Sponsored chemicals		
Hazard Characterizations		determined that mercury in these products poses a "high	Category chemicals	Sponsor: United States	
	EPA is carefully reviewing HPV chemical data to characterize the hazards and risks associated with HPV chemicals. <u>HPVS contains HPV Chemical Hazard</u> Characterizations prepared during EPA's oncoing review of the health and environmental effects data contained with each HPV chalence Program	priority, special concern."	>Login	Current Status; Publication available Hexanovi chloride, 2-ethyl- (CAS 760-67-8)	
Hazard Characterizations	submission. HPVIS also contains Risk-Based Prioritization documents prepared from EPA's examination of HPV Challenge hazard data along with chemical	Start here to look up data on	>Help	Neodecanovi chloride (CAS 40292-82-8)	
	use and exposure information collected from the 2006 Inwentory Update Reporting (2008). These recommendation documents prioritize HPV chemicals for follow-up data collection or management actions based on their potential risks.	a high production volume chemical.		Nonanovi chloride (CAS 764-85-2)	
About HPV Chemical Risk-Based		Enter partial chemical name or	Reports 🗸	Propanovi chloride, 2,2-dimethyl- (CAS 3282-30-2)	
Prioritizations	On November 6, 2009, EPA posted a <u>Bisk Based Prioritization document for elemental mercury in certain products and substitutes (PDF)</u> (7 pp., 67KB, about PDF), and determined that mercury in these products poses a "high priority, special concern."	Chemical Abstract Service (CAS) Number to search the almost 900	>Overall Status		
		chemical substances in the HPVIS	All Sponsored Substances	Aliphatic acids (78)	
Risk-Based Prioritization	Data Collection and HPVIS Content	database.	> Publications		
Documents	In the HPV Challenge Program, companies have sponsored more than 2,200 HPV chemicals, with approximately 1,400 chemicals sponsored directly through			Sponsor: Italy	
	the HPV Challenge Program and over 860 chemicals sponsored indirectly through international efforts.	Search		Current Status: Cheminal assessment in discussion 9-Hexadecenoic acid. (Z)- (CAS 2091-29-4)	
	Under the program, when companies, such as chemical manufacturers and trade associations, voluntarily sponsor a set of HPV chemicals, they provide			Carboxylic acids, di-, C4-11 (CAS 68937-72-4)	
	existing data or perform tests on the chemicals, and submit their test data to this database. To ensure consistency, sponsors follow the <u>Screening</u> Information Data Set (SIDS), developed by the Organization for Economic Cooperation and Development (OECD). SIDS provides internationally agreed upon	Other options:		Fatty acids, C12-20 and C12-20 unsaturated (CAS 68334-03-2)	
	tests for screening chemicals for human and environmental hazards.	 Standard Query Report - Access data from lists of 		Fatty acids, C16 and C18-unsaturated (CAS 67701-07-9)	
	HPVIS consists of basic hazard (toxicity) and environmental fate information on HPV chemicals that can be used by environmental managers, public decision-	MPV ohernical submissions. Use this query to access		Fatty acids, coco, heavy fractions (CAS 68937-85-9)	
	makers, and others in their own health and environmental protection activities.	data, including available		13-Docosenoic acid, (13Z)- (CAS 112-86-7)	
	HPVIS submissions contain data on up to 50 endpoints organized into the following four disciplines described at the end of this paragraph. Click on any of	EPA Mazard Characterizations and Risk-		Ammonium dodecanoate (CAS 2437-23-2)	
	the disciplines to view a list of the MPVIS endpoints included in that discipline. Click on any of the individual endpoints to view the specific data fields defined	Based Prioritization Documents, by searching		Azanium octadecanoate; Octadecanoic acid, ammonium salt (CAS 1002-89-7)	
	for each endpoint. Note: this information is metadata - information describing the data that is included in HPVIS. To view the actual data, use the Search box on the right side of this page.	lists of submissions		Calcium dipalmitate (CAS 542-42-7)	
		organized by either chemical, category,		Carboxylic acids, C5-9 (CAS 68603-84-9)	
	Physical/chemical properties (e.g., metring point, vapor pressure) Environmental fate, and pathways (e.g., biologgradatism, stability in soil)	submission name, sponsor, or submitter.		Carboxylic acids, C6-18 and C8-15 di- (CAS 68937-70-2) Decanoic acid (CAS 334-48-5)	
	Fortrolity (a.e., (sh molecum, midry to accurate data) Fortrolity (a.e., (sh molecum, midry to accurate data)	* Ad Hoc Overy- Create a		Fatty acids, C10-16 (CAS 68002-90-4)	
£		🙂 Internet		Eatty acids, C10-10 (CAS 00002-90-7)	





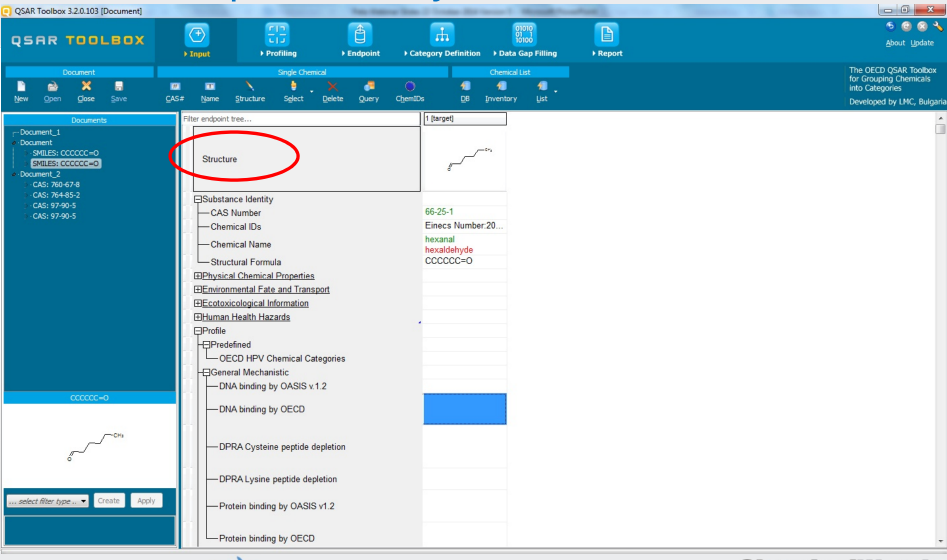
Is Substance Already a Member of an Existing Category?



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Compound Entry and Data Retrieval

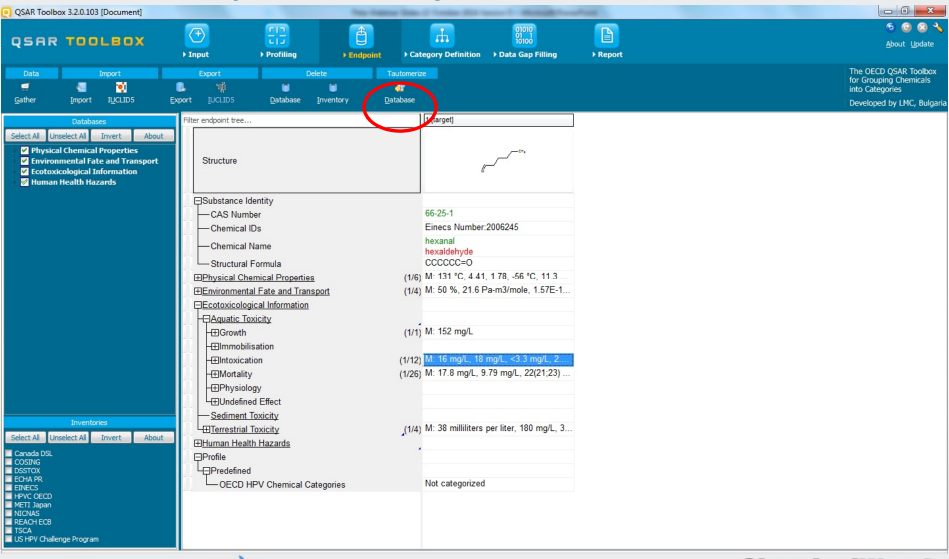






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Compound Entry and Data Retrieval







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Profiling Outcomes

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 Endpoint Specific Acute aquatic toxicity classification by 	CAS Number Chemical IDs	66-25-1 Einecs Number:20	Profiling Profiling Schemes		
Acute aquatic toxicity MOA by OASIS Aquatic toxicity classification by ECOS	- Chemical Name	hexanal	🔸 📄 🖻 🗙		
Bioaccumulation – metabolism alerts Bioaccumulation – metabolism half-lives	Structural Formula	hexaldehyde CCCCCC=0	<u>A</u> pply <u>N</u> ew <u>V</u> iew <u>D</u> elete		
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DNA alerts for AMES, MN and CA by O Eye irritation/corrosion Exclusion rules			Select All Unselect All Invert		
Eye irritation/corrosion Inclusion rules	EHuman Health Hazards		Protein binding by OECD		CY3
 in vitro mutagenicity (Ames test) alerts in vivo mutagenicity (Micronudeus) ale 	Profile Predefined		Protein binding potency Superfragments	Structure	
Keratinocyte gene expression Oncologic Primary Classification	OECD HPV Chemical Categories		Toxic hazard classification by Cramer (
Protein binding alerts for skin sensitization rtER Expert System ver. 1 - USEPA	General Mechanistic		Toxic hazard classification by Cramer (
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Metabolism/Transformations Select All Unselect All Invert	DNA binding by OECD		Acute aquatic toxicity classification by Acute aquatic toxicity MOA by OASIS	- Predefined	
Observed Rat Liver S9 metabolism			Aquatic toxicity dassification by ECOS/	OECD HPV Chemical Categories	Not categorized
Simulated Autoxidation simulator	DPRA Cysteine peptide depletion		Bioaccumulation – metabolism alerts	–⊟General Mechanistic	
Autoxidation simulator (alkaline mediun Dissociation simulation	DPRA Lysine peptide depletion		Bioaccumulation – metabolism half-lives Biodegradation fragments (BioWIN MIT	DNA binding by OASIS v.1.2	No alert found
Hydrolysis simulator (acidic) Hydrolysis simulator (basic)			Carcinogenicity (genotox and nongenc	DNA binding by OECD	Schiff base formers Schiff base formers >> Direct Acting
Hydrolysis sinulator (basic) Hydrolysis simulator (neutral) Microbial metabolism simulator	Protein binding by OASIS v1.2		DNA alerts for AMES, MN and CA by O Eye irritation/corrosion Exclusion rules	brive	Schiff base formers >> Direct Acting
Rat liver S9 metabolism simulator	Dubie biefer hu 050D		Eye irritation/corrosion Inclusion rules l		Low reactive
4	Protein binding by OECD		✓ in vitro mutagenicity (Ames test) alerts in vivo mutagenicity (Micronucleus) ale	DPRA Cysteine peptide depletion	Low reactive >> Long-chain aliphatic Moderate reactive
			Keratinocyte gene expression		Moderate reactive >> Saturated alde
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Creating an Endpoint Specific Category

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			-					Developed by LMC, Bulgaria	
Grouping methods	Filter endpoint tree		1 [target]	2	3	4	5	6 ^	
Predefined Database Affiliation Inventory Affiliation OECD HPV Chemical Categories Substance Type US-EPA New Chemical Categories	Structure		gcrs	H₂C≿O	8 CH4	^۲ ۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰ (۲۰۰۰	//СН3 О	/снз	
General Mechanistic	Substance Identity								
Biodeg BioHC half-life (Biowin) Biodeg primary (Biowin 4)	CAS Number		66-25-1	50-00-0	110-62-3	3268-49-3	75-07-0	123-72-8	
Biodeg probability (Biowin 1)	Chemical IDs	- Chemical IDs		Einecs Number:20	Einecs Number:20	Einecs Number:22	Einecs Number:20	Einecs Number:20	
Biodeg probability (Biowin 2) Biodeg probability (Biowin 5) Biodeg probability (Biowin 6) Biodeg probability (Biowin 7) Biodeg mobability (Biowin 7) Jiodeg probability (Biowin 3) Jind binding by OASIS V.1.2	Chemical Name				valeraldehyde pentanal 1-pentanal	3-(methylthio)propi methional propanal, 3-(methy 3-(methylthio) prop propionaldehyde, 3	acetaldehyde hydrol	butyraldehyde butyraldehyde, n- butanal	
DNA binding by OECD	Structural Formula	cal Chemical Properties (173/917) nmental Fate and Transport (53/232) xicological Information (56/1562)	0=00000	C=0	0=0000	3-(methylsulfanyl)p CSCCC=0	CC=0	0=2222	
OPRA Cysteine peptide deviction DPRA Cysteine peptide depletion				M: -19.1 °C, 0.35,			M: 16.8, 16.7, 13.6		
DPko Crime pentic upletion Estrogen Receptor Binding				M: 71 %, 0.0341 P				M: 100 %, 11.7 Pa	
Hydrolysis half-life (Ka, pH 7)(Hydrowin)				M: 23.8 mg/L, 27.4					
 Hydrolysis half-life (Ka, pH 8)(Hydrowin) Hydrolysis half-life (Kb, pH 7)(Hydrowin) 	⊞Human Health Hazards		-			M: Negative, Negat			
Hydrolysis half-life (Kb, pH 8)(Hydrowin)					5 . 5	5 , 5	5 . 5		
Hydrolysis half-life (pH 6.5-7.4)									
 Ionization at pH = 1 Ionization at pH = 4 	OECD HPV Chemical Categori	es							
Ionization at pH = 7.4	General Mechanistic								
Ionization at pH = 9	DNA binding by OASIS v.1.2	DNA binding by OASIS v.1.2							
Protein binding by OASIS v1.2 Protein binding by OECD	DNA binding by OECD		Schiff base formers Schiff base former Schiff base former						
Documer.c [2(3) Schiff base formers <and3 base<="" schiff="" td=""><td>e f</td><td>on</td><td>Low reactive Low reactive >> Lo Moderate reactive Moderate reactive</td><td></td><td></td><td></td><td></td><td></td></and3>	e f	on	Low reactive Low reactive >> Lo Moderate reactive Moderate reactive						
	DPRA Lysine peptide depletion		Moderate reactive						
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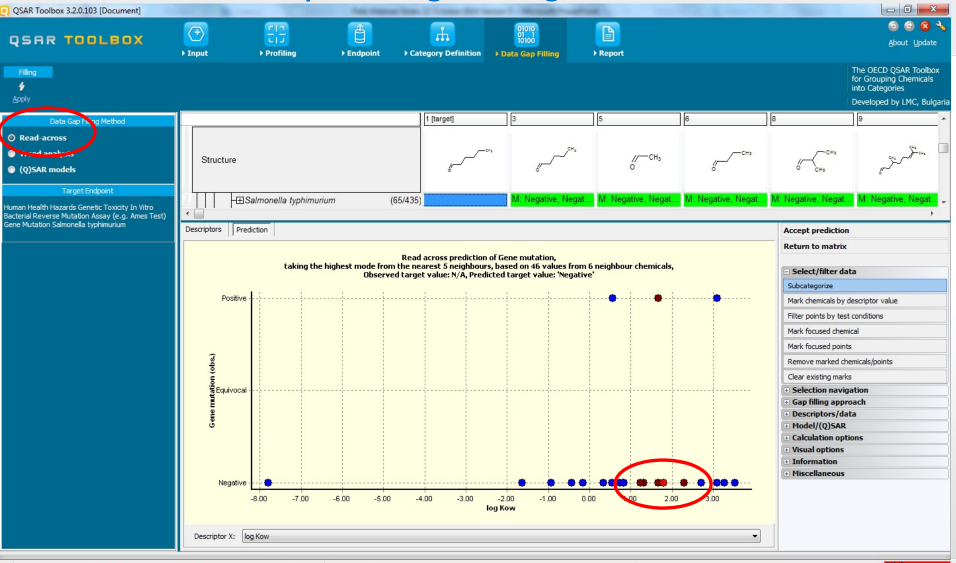
263 Schiff base formers<AND>Schiff base formers >> Direct Acting Schiff Base Formers<AN





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Data Gap Filling Using Read-across



263 Schiff base formers<AND>Schiff base formers >> Direct Acting Schiff Base Formers<AN Create prediction by gap filling





Annex XI of REACH: Grouping and Read-across

- If the group concept is applied, substances shall be classified and labelled on this basis.
- In all cases results should:
 - be adequate for the purpose of classification and labelling and/or risk assessment
 - have adequate and reliable coverage of the key parameters addressed in the corresponding test method referred to in Article 13(3)
 - cover an exposure duration comparable to or longer than the corresponding test method referred to in Article 13(3) if exposure duration is a relevant parameter, and
 - adequate and reliable documentation of the applied method shall be provided





Category Reporting Format (CRF) Should Provide a Detailed Account of the Rationale for Performing a Category or Analogue Approach

- A) information about the category members
- B) what the rationale/hypothesis for formulating the grouping
- C) whether the purities/impurities will affect toxicity
- D) the scope (domain) of the grouping
- E) the endpoints covered and the extent to which the group formulated aims to address all endpoints or a subset of these
- F) Rationale for the validity of the grouping_
- G) Data matrix providing a summary of experimental data for the grouping members
- H) Classification & Labelling information

This document is not trivial to prepare

R.6.2.6.2 Reporting Format for a chemical category

1.	Category definition and its members
1.1.	Category Definition
1.1.a.	Category Hypothesis
	Describe the molecular structure a chemical must have to be included in the category. Provide a brief hypothesis for why the category was formed: the hypothetical relational features of the category i.e. the chemical similarities (analogues), purported mechanisms and trends in properties and/or activities that are thought to collectively generate an association between the members. All functional groups of the category members need to be identified. If there is a mechanistic reasoning to the category, describe the foreseen mode of action for each category member and if relevant describe the influence of the mode of administration (oral dermal, inhalation).
1.1.b.	Applicability domain (AD) of the category
	Describe the set of inclusion and/or exclusion rules that identify the ranges of values within which reliable estimations can be made for category members. Clearly indicate the borders of the category and for which chemicals the category does not hold. For example, the range of log K _{ow} values or carbon chain lengths over which the category is applicable. The justification for the inclusion and/or exclusion rules should be reported under Section 2) <i>Category justification</i> below.
1.2.	Category Members
	Describe all category members as comprehensively as possible. Provide CAS numbers, names and chemical structures of all category members.
1.3.	Purity / Impurities
	Provide purity/impurity profiles for each member of the category, including their likely impact on the category endpoints. It should be discussed which influence these impurities are thought to have on physico-chemical parameters, fate and (eco)toxicology, and hence on the read-across.



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Endpoint Justification

- Overarching rationale (type of groupings) provides a basis to grouping the chemicals together but is essentially a starting hypothesis
- Next step is to justify the grouping on the basis of considerations such as bioavailability, reactivity, metabolism
- And factor how these impact individual endpoints in turn this is where QSARs and other information from the Toolbox can help





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(Q)SAR Endpoint Justifications

- Acute oral considerations include bioavailability, chemical reactivity and metabolism, similarity in structure, Cramer structural classifications.
- Acute dermal concordance with oral results? skin penetration?
- Acute inhalation volatile substances neutral organics appear to be well correlated with Vapour pressure.
- Skin/Eye irritation some overlap with the alerting groups for electrophilicity, pKa?
- Sensitisation alerting groups encoding electrophilic features, Log Kow may be a consideration for some reaction types.





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(Q)SAR Endpoint Justifications

- Mutagenicity lots of focus on Ames but little on other endpoints let alone *in vivo* endpoints
- Carcinogenicity empirical binary QSAR models exist i.e. yes/no prediction but are of limited utility in terms of providing mechanistic justification
- Reproductive/Development handful of empirical models, some (Q)SARs on estrogen binding
- Repeated dose toxicity handful of empirical models which aim to predict LOAEL but not sophisticated to estimate likely target organs.

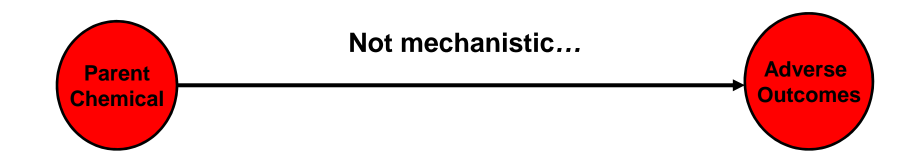
Read-across prone to uncertainty – how can one relate structure to such a downstream endpoint with any reliability?





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Current Approach for Non-testing Development and Application



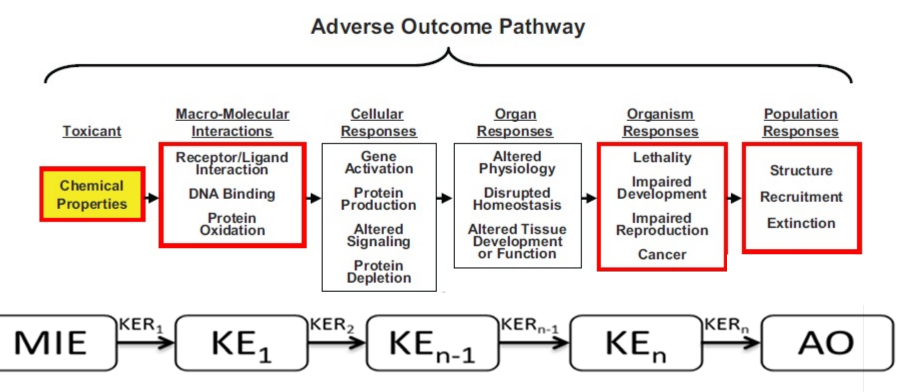
Can relating structure to such downstream adverse outcomes be performed with sufficient scientific confidence?





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AOP: Offers a Framework for Developing Nontesting Approaches Differently



An AOP represents existing knowledge concerning the sequence of events and causal linkages between initial molecular events, ensuing key events and an adverse outcome at the individual or population level.





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Why are AOPs Important?

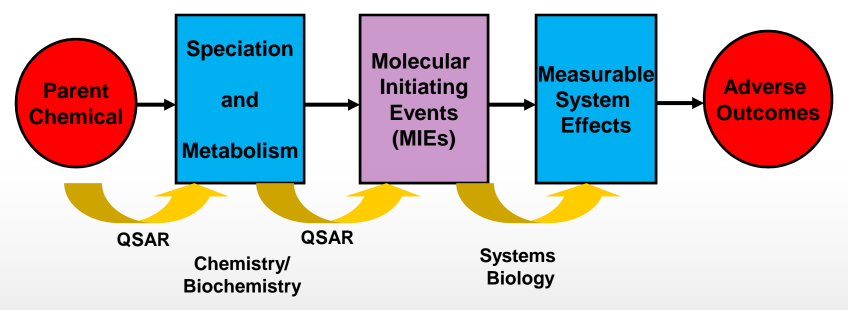
- A framework to organise information
- AOPs provide the linkage from chemistry, through the MIE to Adverse Effect
- Data from key events provides support to, and will enhance, readacross especially for regulatory acceptance
- Data from key events will support definition of domains for MIEs
- Will inform ITS or IATA for risk assessment and provide a roadmap for future QSAR development





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Refining how non-testing approaches are developed in the context of an AOP



1. Identify Plausible MIEs

2. Explore Linkages in Pathways to Downstream Effects 3. Develop QSARs to predict MIEs from Structure or characterise other KEs as SARs





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Implementation of AOP for Skin Sensitisation in the OECD QSAR Toolbox

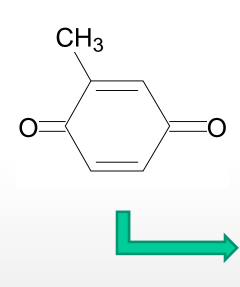
Input target chemical by CAS number

QS

P New

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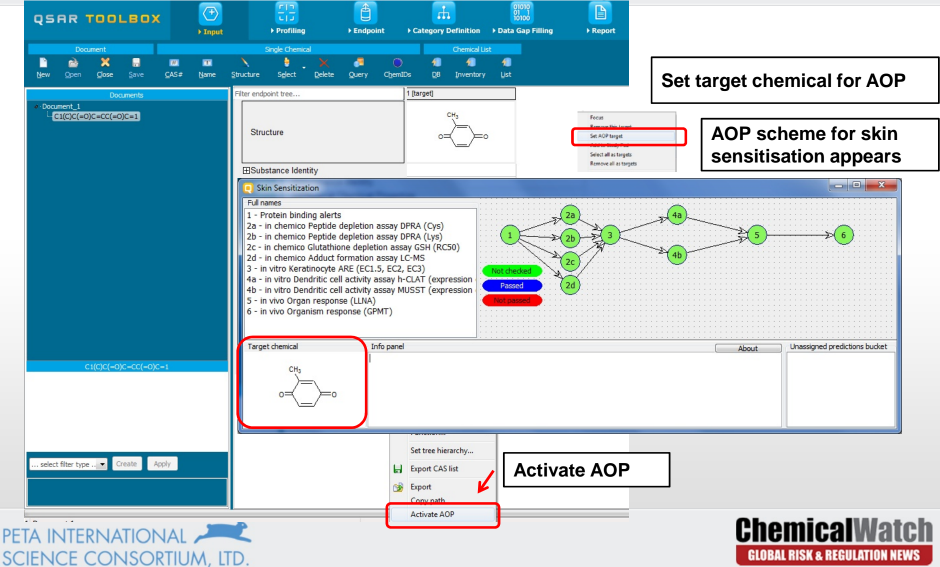
autoxidised to methyl quinone



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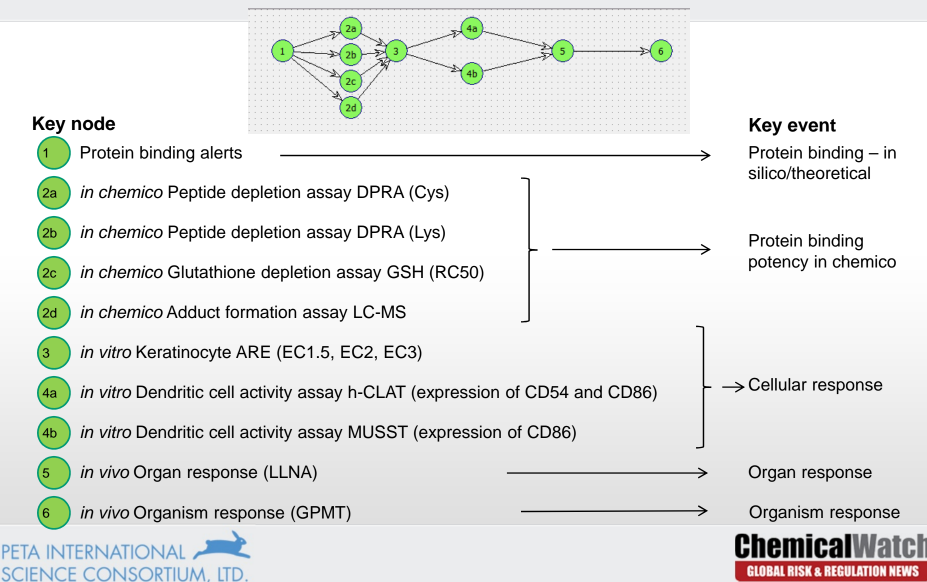


Overview of the AOP implementation in the OECD QSAR Toolbox: Activating the AOP



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Overview of implemented AOP scheme



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Enhancing Read-across

- AOP for skin sensitisation is the first AOP that has been implemented into the Toolbox
- Enables a read-across to be enhanced with information from other downstream key events thereby increasing the confidence in the prediction made and thus its regulatory applicability





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AOP-informed IATA

a) What existing data and data types are available?

> Additional Data, Method Needs

Insufficient confidence What AOP-IATA tools/assays can be applied or need to be developed to generate data to make the decision?



IATA e.g. QSARs, Read-across, ITS Is data input adequate to make regulatory decision? Regulatory decisions

b) Is there an AOP that is applicable to the regulatory application of interest?

- c) Regulatory Applications
- Screening
- Prioritization
- Classification & Labeling
- Hazard Assessment
- Risk Assessment



Take Home Messages - 1

- REACH 2018 represents a significant task of compiling the information requirements for Annexes VII and VIII for a large number of substances
- Annex IX provides opportunities for using adaptations prior to any experimental testing
- Considerations include:
- Has the substance already been registered by another party?
- Are there promising analogues to explore read-across within an analogue/category approach?
- How many datagaps and for which endpoints? This will drive the practical strategy of whether QSARs or grouping approaches are more feasible





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Take Home Messages - 2

- QSARs are most effectively used for ecotox, efate and physchem endpoints as replacement values and as supporting information for "simpler" mammalian endpoints within an IATA
- The OECD principles need to be evaluated for the QSAR(s) and documented in an QMRF together with an QPRF for the prediction itself
- For "more complex" endpoints such as repeated dose 28 day or developmental toxicity screening tests – an analogue/category approach is likely to be more effective – an overarching hypothesis and evidence to support the read-across is essential – (Q)SARs can be helpful in providing some of this evidence





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Take Home Messages - 3

- In future, Tox21 approaches using an AOP construct offer prospects for providing different type of information that is structured in an mechanistic IATA
- This also has implications for how read-across could be justified in future or how QSARs might be developed and applied
- To date an AOP for skin sensitisation has been successfully implemented into the OECD Toolbox to facilitate such a step change in read-across enhancement
- A number of software tools, technical guidance and literature references are available that could be helpful – see useful links pages for a non exhaustive selection





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Useful Links - 1

Domain tools

http://ambit.sourceforge.net/download_ambitdiscovery.html

http://oasis-Imc.org/

Technical regulatory guidance

https://eurl-ecvam.jrc.ec.europa.eu/laboratories-research/predictive_toxicology

http://echa.europa.eu/documents/10162/13632/information_requirements_r6_en.pdf

http://echa.europa.eu/support/grouping-of-substances-and-read-across

http://echa.europa.eu/practical-guides

OECD Toolbox

http://www.qsartoolbox.org/

Industry guidance and experiences

ECETOC TR116 Category approaches, read-across, (Q)SAR

Blackburn, K., and Stuard, S. B. A framework to facilitate consistent characterization of read across uncertainty. *RegToxicol Pharmacol* **2014**, 68, 353-362.





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Useful Links - 2

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- Patlewicz G, Chen MW, Bellin CA. Non-Testing approaches under REACH help or hindrance? Perspectives from a practitioner within Industry. SAR QSAR Environ. Res. 2011, 22(1-2): 67-88.
- Cronin MTD et al (2013) Chemical Toxicity Prediction: Category Formation and Read-Across. Royal Society of Chemistry.
- Cronin MTD and Madden JC (2010) In Silico Toxicology. Principles and Applications. Royal Society of Chemistry.
- Tollefsen, K. E, Scholz, S., **Cronin, M. T.,** et al. (2014). Applying Adverse Outcome Pathways (AOPs) to support Integrated Approaches to Testing and Assessment (IATA). *Reg Toxicol Pharmacol*, in press.





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Useful Links - 3

The next training course for the OECD Toolbox is in Barcelona from Nov. 17-to Nov. 21, organized by ReachMonitor (<u>http://www.reachmonitor.com/index.php?lang=2&aptd=0</u>) and delivered by LMC – developers of the OECD Toolbox (<u>http://www.oasis-Imc.org/</u>).





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... any questions?





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Thank you for attending



What did you think about the webinar? Please take part in our email survey (in your inbox now)

A downloadable recording of this presentation (with slides) will be available shortly.

If you have any questions, please contact Lorna <u>(lorna@chemicalwatch.com)</u>



Webinar 2: Skin irritation and corrosion, 11 Nov, 4pm GMT Click here to register

Webinar 3: Serious Eye Damage and Eye Irritation 4 Dec, 4pm GMT Click here to register



