

Carcinogenicity and Mutagenicity Data: *New Initiatives to Improve Access & Utility for Modeling*

April 2-4, 2008
SCARLET Workshop, Milan, Italy

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY



<http://www.epa.gov/ncct/toxcast>

Part I

Data & Data Linkages



Distributed Structure-Searchable Toxicity (DSSTox) Public Database Network

<http://www.epa.gov/ncct/dsstox/>

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DSSTox

Distributed Structure-Searchable Toxicity (DSSTox) Database Network is a project of [EPA's National Center for Computational Toxicology](#), helping to build a public data foundation for improved structure-activity and predictive toxicology capabilities. The DSSTox website provides a public forum for publishing downloadable, structure-searchable, standardized chemical structure files associated with toxicity data. [More>](#)



[DSSTox Structure-Browser information Page](#)

25 February 2008

***File Updates and Enhancements:

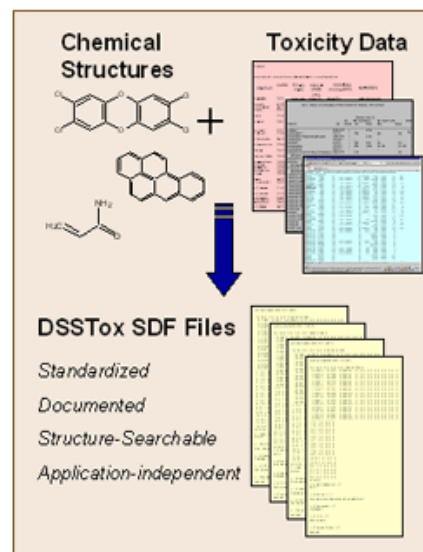
- Addition of new DSSTox Standard Chemical Field to all files: [STRUCTURE_InChIKey](#)
- Additional QA review, structure/CAS modifications, elimination of abbreviations in field entries, etc.

- Addition of categorical and ranked activity summary fields in 5 DSSTox Data Files (CPDBAS, DBPCAN, EPAFHM, FDAMDD, NCTRER), corresponding to standard [PubChem](#) bioassay activity fields:

PUBCHEM_ACTIVITY_OUTCOME (active/inactive/inconclusive):

[ActivityOutcome_CPDBAS_Rat](#)
[ActivityOutcome_CPDBAS_Mouse](#)
[ActivityOutcome_CPDBAS_Hamster](#)
[ActivityOutcome_CPDBAS_Dog_Primates](#)
[ActivityOutcome_CPDBAS_Mutagenicity](#)
[ActivityOutcome_CPDBAS_SingleCellCall](#)
[ActivityOutcome_CPDBAS_MultiCellCall](#)
[ActivityOutcome_DBPCAN](#)
[ActivityOutcome_EPAFHM](#)
[ActivityOutcome_NCTRER](#)


PUBCHEM_ACTIVITY_SCORE [log(1/ activity) mapped onto INTEGER[0-100] ran



- [DSSTox Graphic Flowchart](#)
- [DSSTox Project Goals](#)
- [DSSTox Publications](#)

DSSTox Data Files: [Details>](#) *All Updated

[CPDBAS v5b 1547 10Feb2008](#)
[DBPCAN v4b 209 15Feb2008](#)
[EPAFHM v4b 617 15Feb2008](#)
[FDAMDD v3b 1216 15Feb2008](#)
[HPVCSI v2c 3548 15Feb2008](#)
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[IRISTR v1b 544 15Feb2008](#)
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[NTPBSI v2b 2293 15Feb2008](#)
[NTPHTS v1b 1408 15Jul2008](#)
[TOXCST v2b 320 08Feb2008](#)

NAMEID	version #records date	Expanded DSSTox Data File Title & Description
CPDBAS	v5b 1547 10Feb2008	Carcinogenic Potency Database Summary Tables - All Species: Tumor target site incidence, TD50 potencies, summary activity calls for rat, mouse, hamster, dog, and/or non-human primate; data reviewed and compiled from literature and NTP studies.
DBPCAN	v4b 209 15Feb2008	EPA Water Disinfection By-Products with Carcinogenicity Estimates Database: Carcinogenicity estimates (high, moderate, low concern) by EPA experts using a mechanism-based analog SAR approach on a set of 209 water disinfection by-products, mostly small halogenated organics.
EPAFHM	v4b 617 15Feb2008	EPA Fathead Minnow Acute Toxicity Database: Acute toxicities of 617 chemicals tested in common assay, with mode-of-action assessments and confirmatory measures.
FDAMDD 	v3b 1216 15Feb2008	FDA Center for Drug Evaluation & Research - Maximum (Recommended) Daily Dose Database: Maximum (recommended) daily dose (MRDD) values for 1216 pharmaceuticals in mg/kg-body weight (bw)/day, converted to mmol and normalized to dataset; MRDD values extracted from public literature sources.
HPVCSI	v2c 3548 15Feb2008	EPA High Production Volume Challenge Program Structure-Index File : Compiled structures for three chemical lists provided on EPA HPV Challenge Program website; each record includes reference index to dated list.
HPVISD	v1b 1006 15Feb2008	EPA High Production Volume Information System (HPV-IS) Data Structure-Index Locator File : Compiled structures for the chemical inventory of the on-line EPA HPV-IS with chemical-specific URLs linking to HPV-IS data pages containing chemical properties, fate properties and toxicity data.
IRISTR	v1b 544 15Feb2008	EPA Integrated Risk Information System (IRIS) Toxicity Review Data File: Compiled structures for EPA IRIS website with chemical-specific URLs linking to risk assessment summary data pages for 544 chemical substances.
NCTRER	v4b 232 15Feb2008	FDA National Center for Toxicological Research (NCTR) - Estrogen Receptor Binding Database: Estrogen receptor relative binding affinities tested in a common in vitro assay for 232 chemicals, listed with chemical class-based structure-activity features.
NTPBSI 	v2b 2293 15Feb2008	National Toxicology Program (NTP) On-line Chemical Bioassay Database Structure-Index Locator File : Compiled structures for the NTP On-line Database with chemical-specific URLs linking to NTP study summary pages; file includes fields for each of 4 main bioassay study areas with indicator values specifying presence or absence of study data for the chemical substance record.
NTPHTS	v2b 1408 15Feb2008	National Toxicology Program (NTP) High-Throughput Screening Project Structure-Index File : Compiled structures for set of 1408 NTP chemical substances provided to the NIH Chemical Genomics Center for HTS bioassay testing and to PubChem (PubChem_CIDs and PubChem_SIDs included in NTPHTS_v2a file); NCGC HTS bioassay data are being deposited into PubChem and can be retrieved with these PubChem chemical CID and SID record listings.
TOXCST	v2b 320 08Feb2008	Research Chemical Inventory for EPA's ToxCast™ Program Structure-Index File : Compiled structures for 320 chemical substances that are candidates for Phase I High-Throughput screening (HTS) within the EPA ToxCast™ program. File will be updated with links to PubChem CIDs and SIDs for retrieving assay data, and with updates to chemical inventory as Program moves to Phase II and beyond.



Distributed Structure-Searchable Toxicity (DSSTox) Public Database Network

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SDF Download Page

CPDBAS: Carcinogenic Potency Database Summary Tables - All Species *Database File*

**** Version 5b, updated 10 February 2008:**

- Includes 7 new summary activity fields (TD50 Activity Score and Activity Outcome by species) to coordinate with PubChem CPDB Assay deposits.
 - Includes new InChIKey [Standard Chemical Field](#).
 - V5a includes 66 new chemical records and over 400 new or modified experimental results affecting nearly 100 existing records, corresponding to newly published CPDB website content.
 - Further chemical data
- [Procedures](#)

Quick & Easy File Download

- [Description](#)
- [Source Website](#)
- [Main Citations](#)
- [Guidance for Use](#)
- [Version 5 Update](#)
- [SDF Fields](#)
- [SDF Content Summary](#)
- [SDF Download](#)
- [Acknowledgements](#)

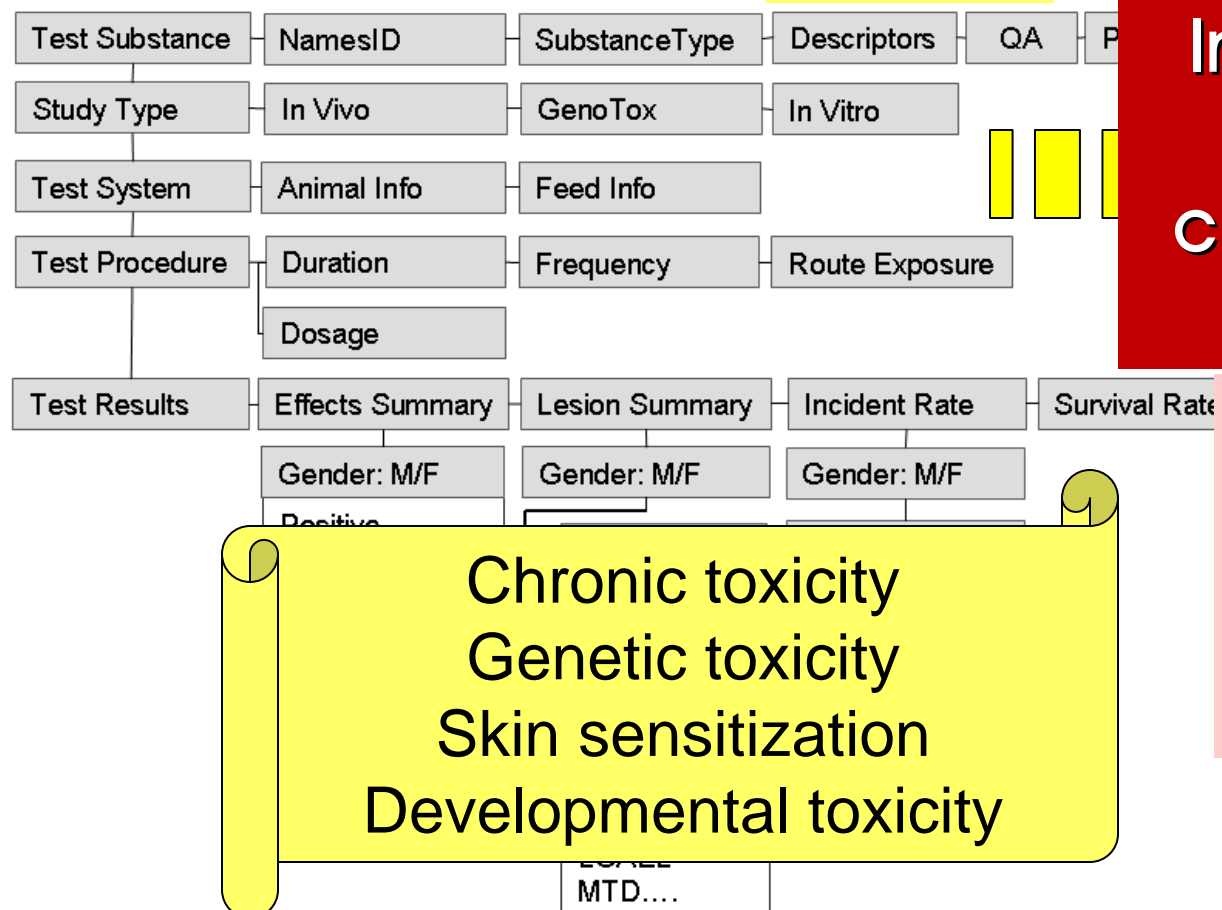
File Types	Description	File Size	Format
Documentation Files			
Log File	CPDBAS_LogFile_10Feb2008.pdf (PDF, 12 pp.)	92KB	
Field Definition File	CPDBAS_FieldDefFile_10Feb2008.pdf (PDF, 11 pp.)	62KB	
	CPDBAS_FieldDefFile_10Feb2008.doc	155KB	
Data Files: CPDBAS			
SDF Structure Data File	CPDBAS_v5b_1547_10Feb2008.sdf		
• Data Table (no structures)	CPDBAS_v5b_1547_10Feb2008_nostructures.xls	*.zip 3.7MB	
• Structures Table	CPDBAS_v5b_1547_10Feb2008_structures.pdf (PDF, 31 pp., 6.3MB)		

[File Error Report](#)

Toxicity Experimental Data → Summary Data:

ToxML / LIST Collaborations: FDA CDER/CFSAN

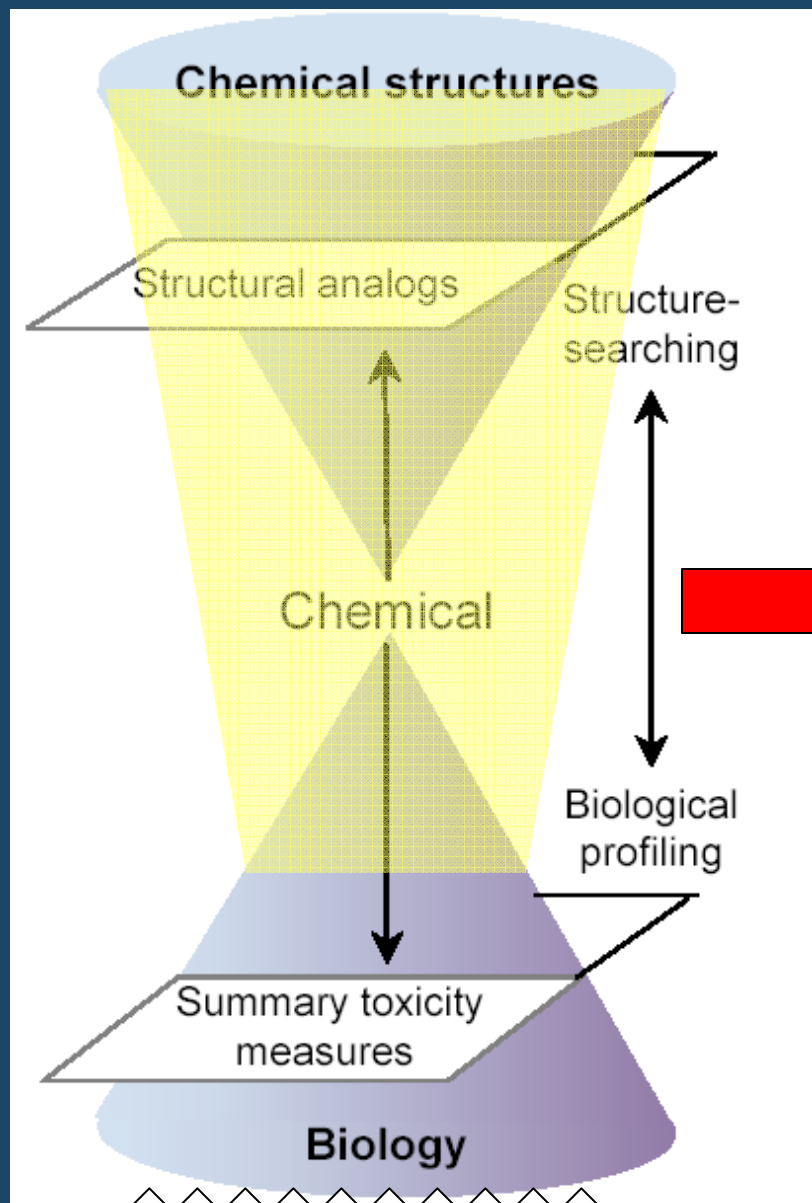
Toxicity Content Model ToxML



**Intermediate
toxicity
classifications
for SAR**

- ✦ Activity categories
- ✦ Potency categories
- ✦ Mode of action categories
- ✦ Summary calls

DSSTox Summary Toxicity Data Files



DSSTox Summary Toxicity Data Files

Compound	Tox1	Tox2	Tox3	Tox4	...
Chem1	rat	male	+	lung	...
Chem2					
Chem3					
Chem4					
Chem5					
Chem6					
Chem7					
Chem8					
...					

↑↑↑↑↑↑↑↑↑↑↑↑↑↑↑↑
Toxicity Data Models

CPDBAS_v5b_1547_10Feb2008

	ActivityCategory_		ActivityCategory_MultiCellCall_Details**			
Call	SingleCellCall	MultiCellCall**	multisite***	multisex	multispecies	Total Incidences* (CPDBAS_v5)
Active (1)	active					223
	active	active	✓			81
	active	active		✓		113
	active	active			✓	8
	active	active				
	active	active				
	active	active				
	active	active				
	active	active				
Inactive (0)	inactive					
	inactive	inactive				
	inactive	inactive				
	inactive	inactive				
	inactive	inactive				

If chemical is

ID:98

Aramite
140-57-8

ID:130

AZT
30516-87-1

ID:299

[4-Chloro-6-(2,3-xylylidino)-2-pyridyl]methyl sulfonate
50892-23-4

ID:316

Chloromethyl methyl ether
107-30-2

ID:715

alpha-1,2,3,4,5,6-Hexachlorocyclohexane
319-84-6

ID:912

Methylethylketoxime
96-29-7

ID:1000

3-Nitro-3-hexene
4812-22-0

ID:1076

p-Nitrosodiphenylamine
156-10-5

CPDBAS SDF Fields (61 total)*

[DSSTox Standard Chemical Fields](#) (19) * [STRUCTURE InChIKey](#) *field added in v5b*

[DSSTox Standard Toxicity Fields](#) (3)

[ActivityOutcome CPDBAS Mutagenicity](#) **modified in v5b* (formerly [Mutagenicity_SAL_CPDB](#))

[TD50 Rat mg](#)

[TD50 Rat mmol](#)

[ActivityScore CPDBAS Rat](#) **new to v5b*

[TD50 Rat Note](#)

[TargetSites Rat Male](#)

[TargetSites Rat Female](#)

[TargetSites Rat BothSexes](#)

[ActivityOutcome CPDBAS Rat](#) **new to v5b*

[TD50 Mouse mg](#)

[TD50 Mouse mmol](#)

[ActivityScore CPDBAS Mouse](#) **new to v5b*

[TD50 Mouse](#) [TD50 Dog mg](#)

[TargetSites M](#) [TargetSites Dog](#)

[TargetSites M](#) [TD50 Rhesus mg](#)

[TargetSites M](#) [TargetSites Rhesus](#)

[ActivityOutcome](#) [TD50 Cynomolgus mg](#)

[TD50 Hamste](#) [TargetSites Cynomolgus](#)

[TD50 Hamste](#) [TD50 Dog Primates Note](#) **modified in v5b*

[ActivityScore](#) [ActivityOutcome CPDBAS Dog Primates](#) **new to v5b*

[TD50 Hamste](#) [ActivityOutcome CPDBAS SingleCellCall](#) **modified in v5b*

[TargetSites H](#) [ActivityOutcome CPDBAS MultiCellCall](#) **modified in v5b*

[TargetSites H](#) [ActivityOutcome CPDBAS MultiCellCall Details](#) **modified in v5b*

[TargetSites H](#) [Note CPDBAS](#)contains controlled text entries for version content updates

[ActivityOutcome](#) [NTP TechnicalReport](#)

[TD50 Dog mg](#) [ChemicalPage URL](#)(formerly [Website_URL](#) in v4a), contains link to the record-specific [CPDB Chemical Index](#) data page, e.g. see [ACETALDEHYDE](#) [EXIT Disclaimer](#).

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Send to

Tools:



Links: [Related Structures](#), [BioAssays](#), [Literature](#), [Other Links](#)

All: 12940

BioAssay: 3821

Protein3D: 0

Rule of 5: 7987

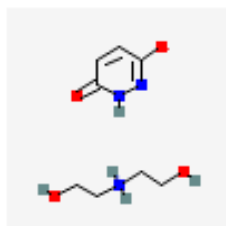


12940 DSSTox Substances

Items 1 - 20 of 12940

☐ 1: SID: 48423627

[Related Structures](#)



MALEIC HYDRAZIDE DIETHANOLAMINE; 2-hydroxy-N-(2-hydroxyethyl)ethanaminium 6-oxo-1,2,3,6-tetrahydropyridazin-3-olate; 5716-15-4

Compound ID: 24180705

Source: EPA DSSTox (31555)

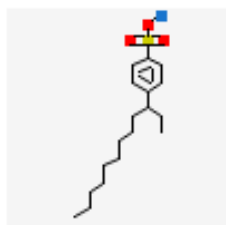
→ DSSTox NTPHTS Download Page

IUPAC: bis(2-hydroxyethyl)azanium; 6-oxo-1H-pyridazin-3-olate

MW: 217.222400 g/mol | MF: C8H15N3O4

☐ 2: SID: 48423362

[Related Structures](#), [Literature](#)



Dodecylbenzenesulfonic acid, sodium salt; 25155-30-0

Compound ID: 23707968

Source: EPA DSSTox (31261)

IUPAC: sodium 4-dodecan-3-ylbenzenesulfonate

MW: 348.475830 g/mol | MF: C18H29NaO3S

Search PubChem BioAssay for dsstox Go Clear Save Search

Limits Preview/Index History Clipboard Details

Display Summary Show 20 Sort by Send to

Tool: Links: Related BioAssays, Compounds, Literature, Other Links

All: 74 Confirmatory: 63 MLSCN: 63 Protein Target: 0 Screening: 1 Summary: 0

Items 1 - 20 of 74

1: AID: 1204 Summary | Data (Active) Related BioAssays, Compounds, Literature, Other Links

DSSTox (NCTRER) National Center for Toxicological Research Estrogen Receptor Binding Database [Screening Method]

Source: EPA DSSTox

Substances Tested: 232; Active: 131

2: AID: 1195 Summary | Data (Active) Related BioAssays, Compounds, Literature, Other Links

DSSTox (FDAMDD) FDA Maximum (Recommended) Daily Dose Database [Other Method]

Source: EPA DSSTox

Substances Tested: 1240; Active: 587

3: AID: 1205

DSSTox (CPDBAS)

Source: EPA DSSTox

Substances Tested: 403 /860 Active

4: AID: 1189

DSSTox (CPDBAS)

Source: EPA DSSTox

Substances Tested: 806 /1547 Active

5: AID: 1208

DSSTox (CPDBAS) Carcinogenic Potency Database Summary Rat Bioassay Results [Other Method]

Source: EPA DSSTox

Substances Tested: 582 /1152 Active

1. AID 1194: CPDBAS Salmonella Mutagenicity	403	/860 Active
2. AID 1189: CPDBAS SingleCellCall	806	/1547 Active
3. AID 1205: CPDBAS MultiCellCall	582	/1152 Active
4. AID 1208: CPDBAS Rat Bioassay (M/F/Both)	587	/1240 Active
5. AID 1199: CPDBAS Mouse Bioassay (M/F/Both)	445	/1007 Active
7. AID 1190: CPDBAS Dog & Primates Bioassay	15	/32 Active
8. AID 1195: FDAMDD – FDA Maximum Daily Dose	1216	/1216 Active
9. AID 1204: NCTRER – NCTR Estrogen Receptor Binding	131	/232 Active
10. AID 1188: EPA Fathead Minnow Acute Toxicity	580	/617 Active
11. AID 1201: EPA Disinfection By-Products Carcinogenicity Estimates	80	/209 Active

Related BioAssays by Activity Overlap

 **AID: 1205** 

Name: DSSTox (CPDBAS) Carcinogenic Potency Database Summary MultiCellCall Results

Data Source: [EPA DSSTox](#)



BioActivity Analysis: Structure-Activity

Activity Overlap for CPDBAS MultiCellCall Results

334 Related BioAssays by Activity Overlap of AID 1205

Total Pages: 17

Display:

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Page



#	<input type="checkbox"/>	Activity Similarity	Active in Both	BioAssay Name
1	<input type="checkbox"/>	72.6%	572	AID: 1189 , DSSTox (CPDBAS) Carcinogenic Potency Database Summary SingleCellCall Results
2	<input type="checkbox"/>	62.3%	441	AID: 1208 , DSSTox (CPDBAS) Carcinogenic Potency Database Summary Rat Bioassay Results
3	<input type="checkbox"/>	56%	362	AID: 1199 , DSSTox (CPDBAS) Carcinogenic Potency Database Summary Mouse Bioassay Results
4	<input type="checkbox"/>	32.8%	239	AID: 1194 , DSSTox (CPDBAS) Carcinogenic Potency Database Salmonella Mutagenicity
5	<input type="checkbox"/>	6.9%	40	AID: 1191 , DSSTox (CPDBAS) Carcinogenic Potency Database Summary Hamster Bioassay Results
6	<input type="checkbox"/>	4.3%	29	AID: 426 , Cell Viability - Jurkat
7	<input type="checkbox"/>	4.2%	29	AID: 544 , Cell Viability - SH-SY5Y
8	<input type="checkbox"/>	3.8%	42	AID: 1188 , DSSTox (EPAFHM) EPA Fathead Minnow Acute Toxicity
9	<input type="checkbox"/>	3.6%	24	AID: 540 , Cell Viability - N2a
10	<input type="checkbox"/>	3.3%	22	AID: 543 , Cell Viability - H-4-II-E
11	<input type="checkbox"/>	3.3%	22	AID: 981 , Cell Viability - LYMP2-010
12	<input type="checkbox"/>	3.2%	20	AID: 427 , Cell Viability - Hek293
13	<input type="checkbox"/>	3.2%	55	AID: 1195 , DSSTox (FDAMDD) FDA Maximum (Recommended) Daily Dose Database



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DSSTox

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25 February 2008

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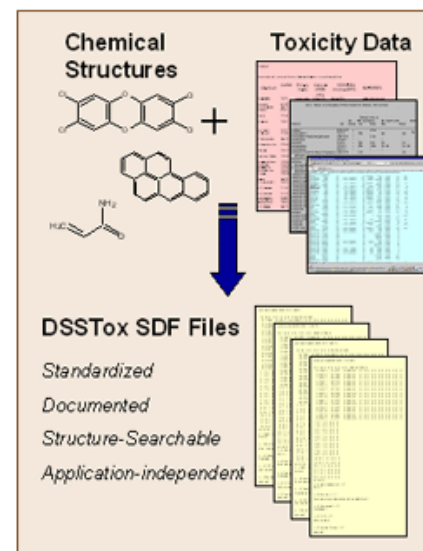
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allows users to search by [DSSTox Standard Chemical Fields](#) and includes options for:

- **Text Search:** Chemical Name, CAS RN, InChI, Formula
- **Structure Search (Exact, Substructure, Similarity):** SMILES or Structure Drawing Tool entry

***Revised Standard ID Fields for all DSSTox files:

- Modified [Record, File, Chemical, and Substance ID fields](#) to index all unique DSSTox structures and substances, also with respect to file record and version



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DSSTox Chemical Text Search

Choose search:

Enter search text:

Auto-detect

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Search

DSSTox Chemical Structure Search

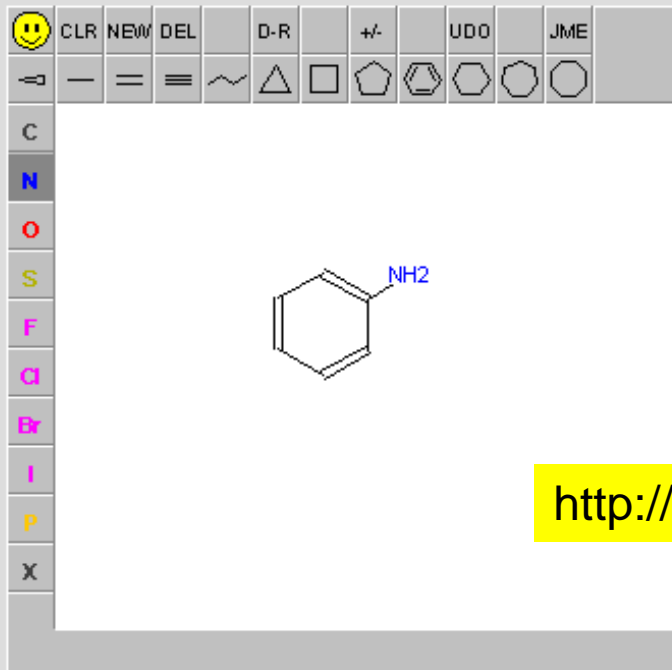
Enter SMILES string:

Preview below

Clear

Search

Or draw a molecule or substructure using the JME editor:



Search Options

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Search

EPA Integrated Risk Information System (IRIS)
Structure-Index Locator File (544 records)

Data Files to Search

☒ All DSSTox Files

☐ Selected DSSTox Files

☒ CPDBAS_v5b

☒ DBPCAN_v4b

☒ EPAFHM_v4b

☒ FDAMDD_v3b

☒ HPVCSI_v2c

☒ HPVISD_v1b

☒ IRISTR_v1b

☒ NCTRER_v4b

☒ SI_v2b

☒ TS_v2b

☒ TOXCST_v2b

http://www.epa.gov/dsstox_structurebrowser/

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Aniline Quickview (CASRN 62-53-3)

Quickview Navigation

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- [view Aniline Summary](#)

Health assessment information on a chemical substance is included in IRIS only after a comprehensive review of toxicity data by U.S. EPA health scientists from several Program Offices, Regional Offices, and the Office of Research and Development.

Disclaimer: This QuickView represents a snapshot of key information. We suggest that you read the [IRIS Summary](#) to put this information into complete context.

For definitions of terms in the IRIS Web site, refer to the [IRIS Glossary](#).

Status of Data for Aniline

File First On-Line: 09/07/1988

Last Significant Revision: 11/01/1990

Category (section)	Status	Last Revised
Oral RfD Assessment	No data	
Inhalation RfC Assessment	On-line	12/01/1993
Carcinogenicity Assessment	On-line	02/01/1994

Chronic Health Hazard Assessments for Noncarcinogenic Effects

[Reference Dose for Chronic Oral Exposure \(RfD\)](#)

Not Assessed under the IRIS Program.

[Reference Concentration for Chronic Inhalation Exposure \(RfC\)](#)

Critical Effect	Point of Departure	UF	MF	RfC
Lack of toxicity (other effect: Mild spleen toxicity.)	NOAEL (HEC): 3.4 mg/m3	3000	1	1x10⁻³ mg/m3



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IRIS (Integrated Risk Information System) is a compilation of electronic reports on specific substances found in the environment and their potential to cause human health effects. IRIS was initially developed for EPA staff in response to a growing demand for consistent information on substances for use in risk assessments, decision-making and regulatory activities. The information in IRIS is intended for those without extensive training in toxicology, but with some knowledge of health sciences.

Getting Started with IRIS



- [An overview of the web site](#)
- [What is IRIS?](#)
- [How does EPA decide which substances to add or update?](#)

[More frequent questions >>](#)

Search IRIS by Keyword

go

- ☒ IRIS Summaries/Toxicological Reviews
- ☐ Entire IRIS Website

[List of IRIS Substances >>](#)

Ask Peter



I'm Peter, the IRIS Virtual

Using the



Advanced Search

By Keyword,
Substance, or
CASRN

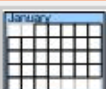
By Critical
Effect or
Tumor Type

By Toxicity
Value

By Human
Carcinogenicity

By
Uncertainty
Factor

Upcoming



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Search by Keyword, Substance, or CASRN

Go

Enter a Keyword, Substance, or CASRN:

☒ IRIS Summary/Toxicological Review

☐ Entire IRIS Website



Go

Search Scope

The search by **Keyword** allows you to enter any keyword or string of keywords to locate information on that topic. The search by [Chemical Name or CAS Registry Number \(CASRN\)](#) allows you to enter any chemical name or CASRN and retrieve a list of IRIS substances that match or are a synonym for that chemical name or CASRN.

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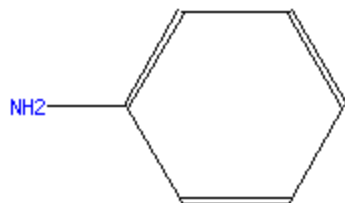
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EPA IRIS Website

DSSTox Structure-Browser



[IRISTR:](#)

EPA Integrated Risk Information System (IRIS) Structure-Index Locator File (544 records)

IRISTR_v1a_544_28Jul2007

[IRISTR Source Website](#)

Output Options

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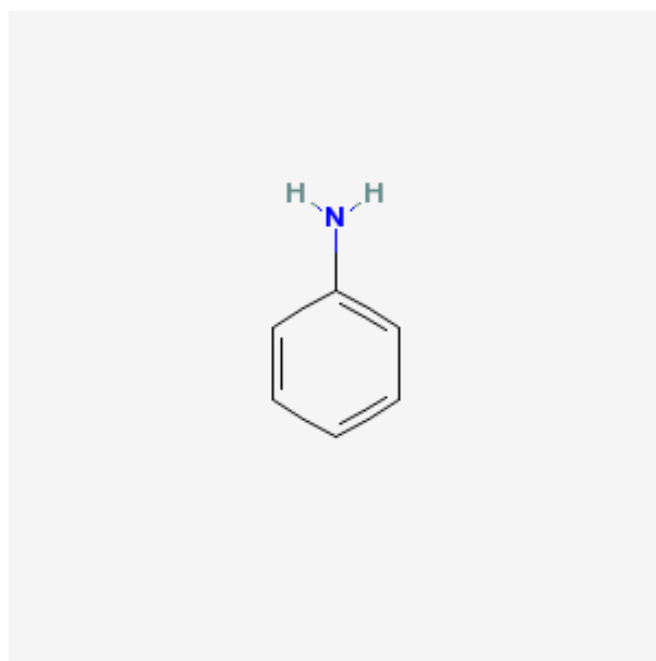
Print

[PubChem](#)

Links directly to chemical data page for Aniline in PubChem

DSSTox_RID	23877
DSSTox_Generic_SID	20090
StudyType	Human Health Exposure Toxicity Review for Risk Assessment
Endpoint	cancer; acute; short-term; sub-chronic; chronic; developmental
Species	rodent; human; dog; rabbit
STRUCTURE_Shown	tested chemical
TestSubstance_ChemicalName	Aniline
TestSubstance_CASRN	62-53-3
TestSubstance_Description	single chemical compound
Oral_RfD_Assessed	0
Oral_RfD_CriticalEffects	Not assessed under the IRIS program.
Inhalation_RfC_Assessed	1
Inhalation_RfC_CriticalEffects	mild spleen toxicity
Inhalation_RfC_mg_per_m3	0.001 mg/m3
Inhalation_RfC_mmol_per_m3	1.07380820711613E-05 mmol/m3
Inhalation_RfC_Notes	NOAEL (No observed adverse effect level) HEC (Human Equivalent Concentration): 3.4 mg/m3
Inhalation_RfC_Confidence	Low
WtOfEvidence_Cancer_Assessed	1
WtOfEvidence_Cancer_Concern	Medium
WtOfEvidence_1986GuidelineCategories	B2; Probable human carcinogen - based on sufficient evidence of carcinogenicity in animals
WtOfEvidence_Cancer_Narrative	Induction of tumors of the spleen and the body cavity in two strains of rat; and some supporting genetic toxicological evidence.

Compound Summary:



CID: [6115](#) [?](#) [+](#)



BioActivity: [Summary](#) [?](#)

All: [53 Links](#)

Active: [6 Links](#)

Inactive: [45 Links](#)

Inconclusive: [1 Link](#)



Protein Structures: [3 Links](#) [?](#)



Protein Sequences: [15 Links](#) [?](#)



NLM Toxicology: [Link](#) [?](#)



Substances: [?](#)

All: [270 Links](#)

Same: [28 Links](#)

Mixture: [242 Links](#)



Related Compounds: [?](#)

Same, Connectivity: [13 Links](#)



Similar Compounds: [164 Links](#) [?](#)



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DSSTox
Structure-
Browser



Part II

Toxicity Profiling

National Academy of Sciences Report (2007) *Toxicity Testing in the Twenty-first Century: A Vision and a Strategy*

NAS PANEL SEEKS MAJOR SHIFT IN HOW EPA ASSESSES CHEMICALS' TOXICITY

Date: June 22, 2007 -

Inside EPA

Online access provided by InsideEPA.com

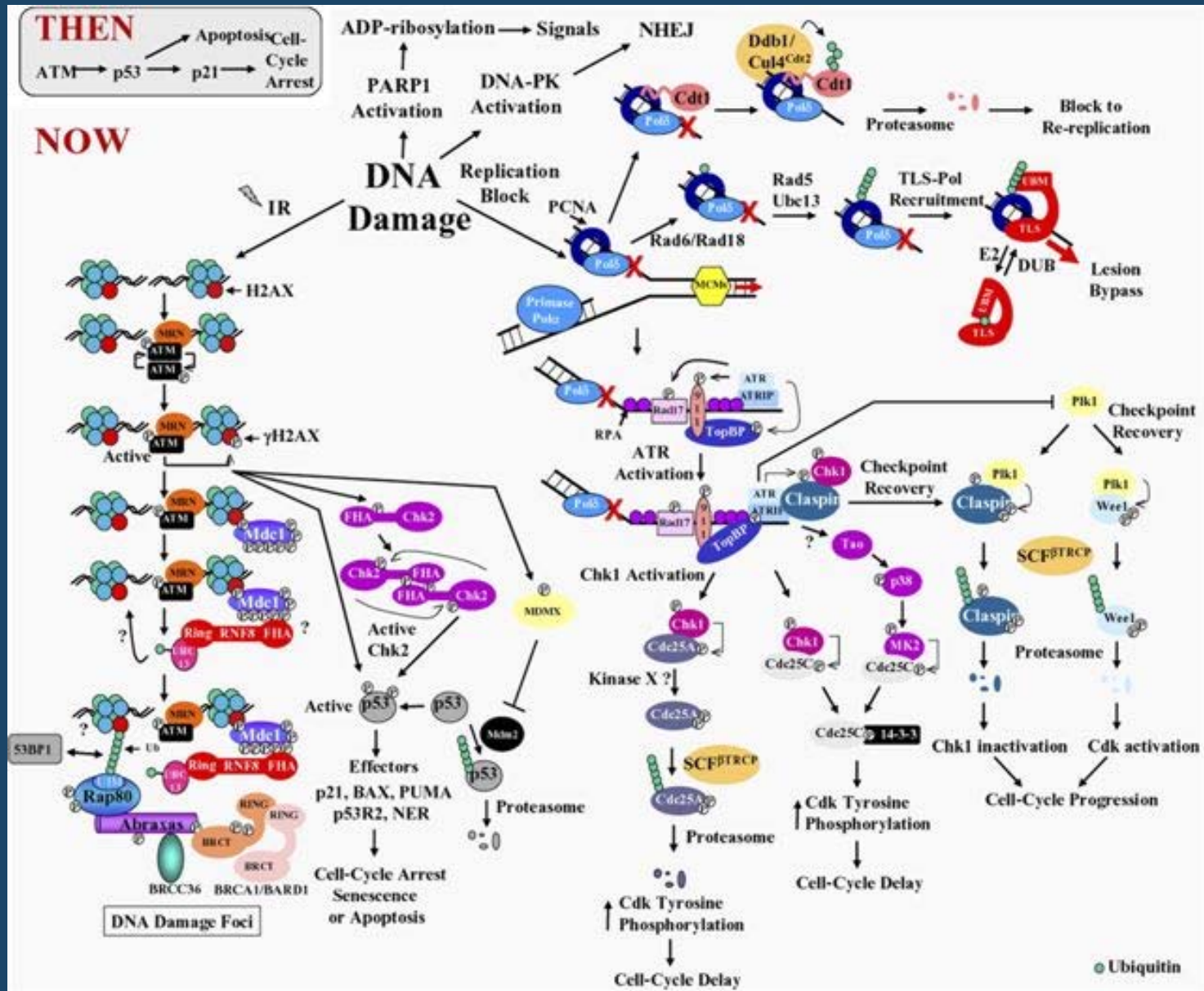
A National Academy of Sciences (NAS) panel is calling for a major shift in how EPA assesses chemicals' toxicity, recommending that the agency base its toxicological research and regulatory processes on how substances affect biological pathways -- which send information within and between cells -- rather than so-called health endpoints, such as cancer.

The new studies envisioned by the panel would evaluate chemicals' effects on biological processes using cells or cell lines, preferably human, to examine how they react to exposure to different substances. Rather than focusing research and basing regulations on endpoints, such as a substance's apparent ability to create tumor cells or harm brain development in fetuses, EPA should center toxicity testing around "the perturbations in toxicity pathways that are expected to lead to adverse effects," the report says.

"In this framework, the goals of toxicity testing are to identify critical pathways that when perturbed can lead to adverse health outcomes and to . . . understand the effects of perturbations on human populations," says the report, *Toxicity Testing in the Twenty-first Century: A Vision and a Strategy*.

The DNA Damage Response: 10 Years After

[Harper & Elledge, Mol Cell 14:739, 2007]





National Center for Computational Toxicology

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ToxCast™ Program

Predicting Hazard, Characterizing Toxicity Pathways, and Prioritizing the Toxicity Testing of Environmental Chemicals

<http://www.epa.gov/ncct/toxcast/>

Introduction

In 2007, EPA launched ToxCast™ in order to develop a cost-effective approach for prioritizing the toxicity testing of large numbers of chemicals in a short period of time. Using data from state-of-the-art high throughput screening (HTS) bioassays developed in the pharmaceutical industry, ToxCast™ is building computational models to forecast the potential human toxicity of chemicals. These hazard predictions will provide EPA regulatory programs with science-based information helpful in prioritizing chemicals for more detailed toxicological evaluations, and lead to more efficient use of animal testing.

ToxCast™ Navigation

[Introduction](#)
[ToxCast™ Chemicals](#)
[ToxCast™ Assays](#)
[ToxCast™ Information Management](#)
[ToxCast™ Partnerships](#)
[ToxCast™ Contractors](#)
[ToxCast™ Presentations](#)
[ToxCast™ Publications](#)

POLICYFORUM

Science: Feb 15, 2008

TOXICOLOGY

Transforming Environmental Health Protection

Francis S. Collins,^{1†} George M. Gray,^{2*} John R. Bucher^{3*}

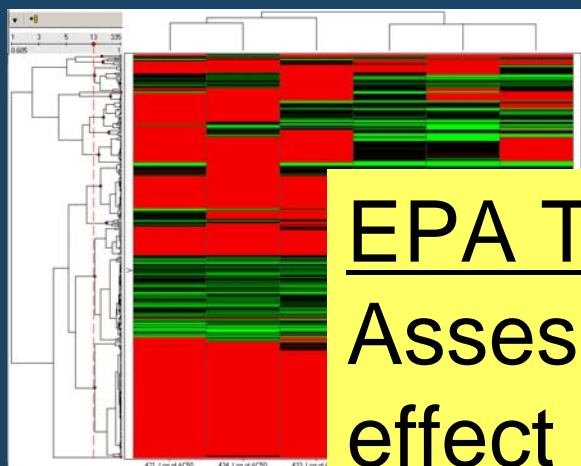
In 2005, the U.S. Environmental Protection Agency (EPA), with support from the U.S. throughput screening (HTS) and other automated screening assays into its testing tion, usually between 2 and 10 μ M, and tolerate high false-negative rates. In contrast, in

We propose a shift from primarily in vivo animal studies to in vitro assays, in vivo assays with lower organisms, and computational modeling for toxicity assessments.

Correlating Domain Outputs

EPA ToxCast Goal:

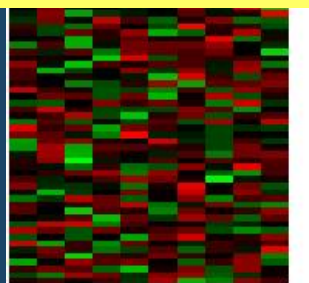
Assess broad biological effect patterns and correlate with the patterns of known toxicants for forecasting potential for hazard



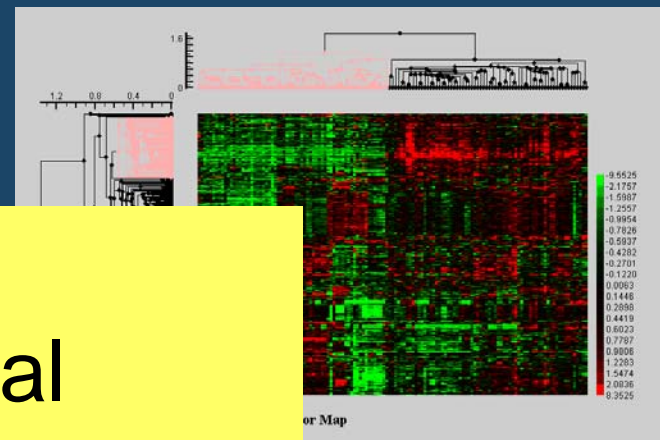
Cellular Assays



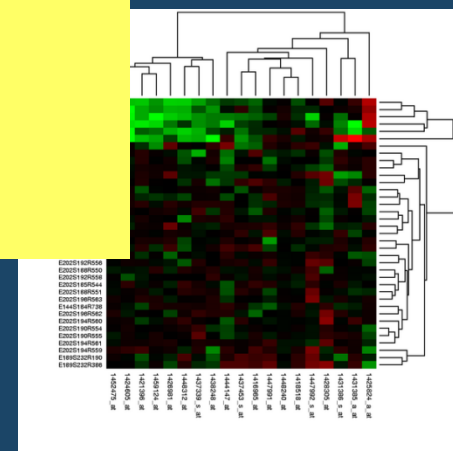
Biochemical Assays



Genomic Signatures



Chemical properties



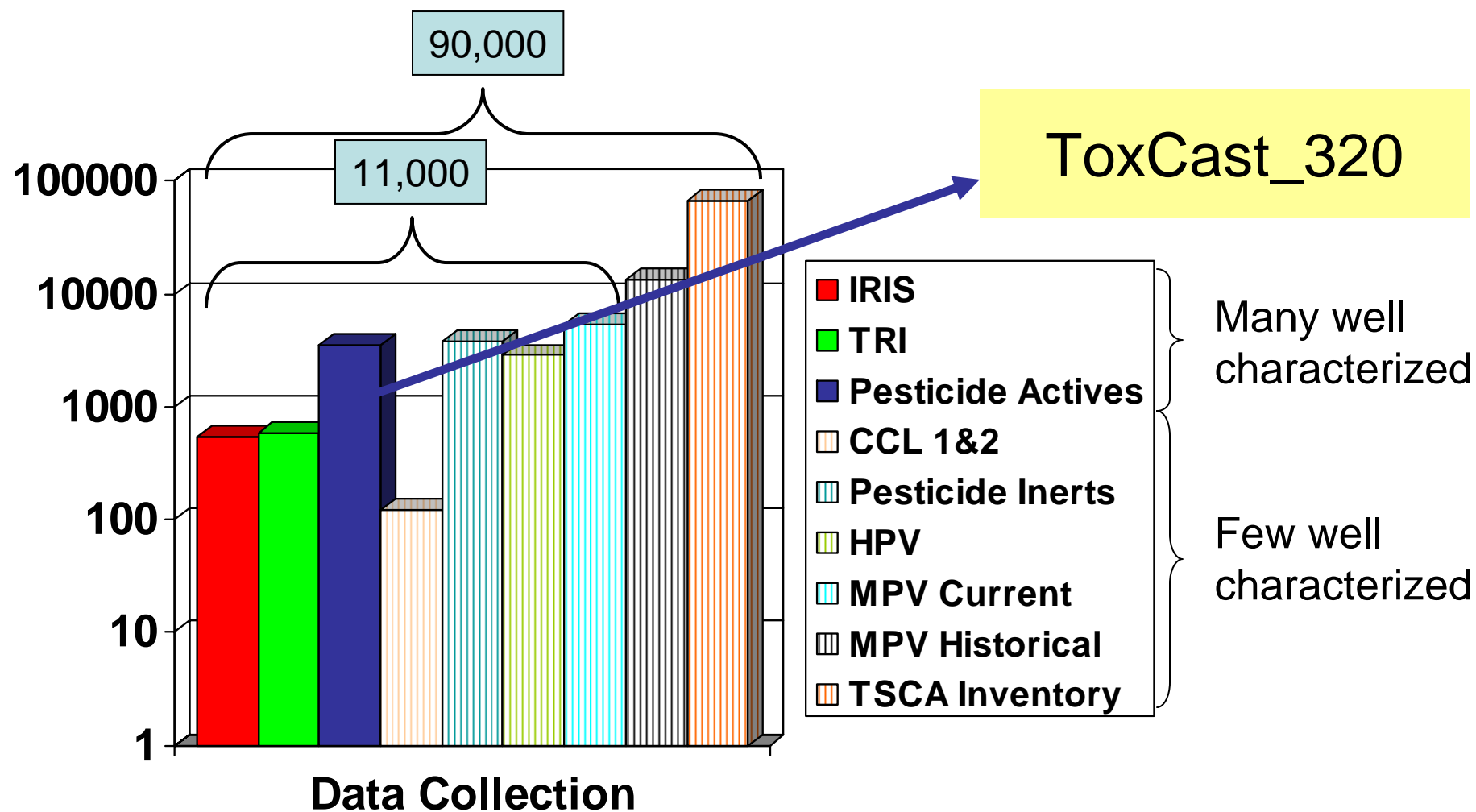
Toxicology Endpoints

ToxCast Phase I Objective: *Derive Predictive Signatures for Chemical Prioritization*

Requirements for Chemical Selection:

- Sufficient number and diversity of chemicals
- Availability of high quality reference *in vivo* toxicity data covering broad range of endpoints
- Expectation of broad spectrum biological activity
- Properties suitable for HTS
- Cost and availability

ToxCast Phase I Chemicals



Prioritize List of ~840 Pesticide Actives

- Data availability
- Toxicological interest
- Environmental relevance

CAS_NO	Parent Molecular Weight	ALogP	PCCODE	CHEMICALNAME	Exclude(X) Additional(+)	DER_TOTAL	Comment	Pesticides (3448)	Pesticides (Actives Supported) (1082)	Pesticide (Food Use Actives) (336)	AntiMicrobial (754)	AntiMicrobial (Food Use) (26)	Inerts (List 1) (8)	Inerts (List 2) (96)	HPV (2843)	HPV Challenge (1973)	EDSP (79)	CCL (41)	Number of lists	Activity/Chemical Classes	Activity/Pesticidal MOA
135158-54-2	210.3	2.19	061402	1,2,3-Benzothiadiazole-7-		6		1	1	1	0	0	0	0	0	0	0	0	3		unclass
123312-89-0	217.2	0.82	101103	1,2,4-Triazin-3(2H)-one, 4,5-		6		1	1	1	0	0	0	0	0	0	0	0	3	Triazine	
68049-83-2	338.2	3.74	119016	1,2,4-Triazolo(4,3-a)pyridin-3(2H)-		6		1	1	0	0	0	0	0	0	0	0	0	2		unclass
139-40-2	229.7	2.17	080808	1,3,5-Triazine-2,4-diamine, 6-		5		1	1	0	0	0	0	0	0	0	0	0	2	Triazine	chlorotri
122-34-9	201.7	1.39	080807	1,3,5-Triazine-2,4-diamine, 6-		6		1	1	1	1	1	0	0	0	0	1	0	6	Triazine	chlorotri
101-05-3	275.5	3.58	080811	1,3,5-Triazine-2-amine, 4,6-		6	Mouse	1	1	0	0	0	0	0	0	0	0	0	2	Triazine	triazine
29091-21-2	350.3	3.73	110201	1,3-Benzenediamine, 2,6-dinitro-		6		1	1	0	0	0	0	0	0	0	0	0	2	Amine,	phenyle
118134-30-8	297.5	4.07	120759	1,4-Dioxaspiro[4.5]undecane-2-		6		1	1	0	0	0	0	0	0	0	0	0	2		unclass
116255-48-2	377.1	3.66	120503	1H-1,2,4-Tria-															2	Bromin	conazol
119168-77-3	333.9	3.86	090102	1H-Pyrazole-															2	Amide,	pyrazol
119515-38-7	229.3	2.16	070705	1-Piperidinec															2	Alcohol	
101-10-0	200.6	2.19	021201	2-(m-Chlorop															2	Phenox	phenox
21564-17-0	238.3	3.56	035603	2-		6		1	1	0	1	0	1	0	0	0	0	0	4	Thiazol	benzoth
2971-36-0				2,2-Bis(4-hydroxyphenyl)-1,1,1-	+	0	metabol	0	0	0	0	0	0	0	0	0	0	0	0		organoc
94-75-7	221	2.46	030001	2,4-Dichlorophenoxyacetic acid		6	Repro	1	1	1	0	0	0	0	1	1	1	0	6	Carbox	phenox
16079-88-2	241.5	1.49	006315	2,4-Imidazolidinedione, 1-bromo-		6		1	1	0	1	0	0	0	1	0	0	0	4	Imidazo	
136-45-8	251.3	2.22	047201	2,5-Pyridinedicarboxylic acid,		6		1	1	1	0	0	0	0	0	0	0	0	3	Ester,	
120-32-1	218.7	3.64	062201	2-Benzyl-4-chlorophenol		5		1	1	0	1	0	1	0	0	0	0	0	4	Cresol,	
7786-34-7	224.1	0.41	015801	2-Butenoic acid, 3-		6		1	1	0	0	0	0	0	0	1	0	0	3	Organo	organop
1909-82-4	230.9	3.56	069203	2-Chloro-6-		6		1	1	1	0	0	0	0	1	1	0	0	5	Organo	

Pesticide Registration Records

Testsubstance_ChemicalName	Molecular Weight	ADMET SolubilityLeve	ALogP	PChem Sum (1-3)	EPAPTC 826	NTPHTS 1408	EPADNT 82	DNT GRADN	IMMTOX 87	HPVCSI 3548	ICCVAM 87	NCTRRER 232	NTPCSI 2415	NTPGTZ 1931	IRISSI 544	ECODEM 399	EPAFHM 617	FDAMDD 1217
1-Naphthaleneacetic acid	186.2	3	2.47	1	30	86										5		
2,4-D, dimethylamine salt	266.1	3	2.46	1	44	157							1373	509				
2,4-D	221	3	2.46	1	52	167				250		119	233	507	193	6		
Oethillone	213.3	3	3.35	1	584	280							382	1418		143		
Acetaminophen	151.2	4	1.35	1	104	436							725	1029			165	3
Acetochlor	269.8	3	3.12	1	114	440							731		5	17		
Alachlor	269.8	3	2.99	1	118	459						103	782		16	18	575	
Aldicarb	190.3	4	1.47	1	119	460	6	3	2				783	43	18	19	264	
Atrazine	215.7	3	1.78	1	149	501			52	1127	8	49	869	158	48	25		
Benomyl	290.3	3	2.57	1	156	511			2998				895	175	57	27		
1H-Purine-2,6-dione, 3,7-dihyd	194.2	4	0.06	1	26	584				20			168	1065	329		24	181
Capsaicin	305.4	3	3.79	1	222	587							1074					
Carbaryl	201.2	3										126	1620		102	38	33	
Carbendazim	191.2	3								2985			1079	342		293		
Chloropirrin	164.4	3											1124	419		48		
Citric acid	192.1	5								129								288
Dazomet	162.3	3	0.8	1	320	683				859			1224	515		379		
Diazinon	304.3	3	4.24	1	326	697	26						1256	546		302	334	
Dicloran	207	3	2.48	1	332	707				319			1273	575		67		
Dichlorvos	221	4	1.61	1	330	710	27			2786			1280	628	197	66		
Disulfoton	274.4	3	4.42	1	360	763	30						1364		233	365	328	
Camphor	152.2	3	2.4	1	221	767							1069					
Carbamothioic acid, dipropyl-	189.3	3	2.98	1	231	773	32						2135	865	248	84		
Fluometuron	232.2	3	2.8	1	428	808							1452	940	260	315		
Malathion	330.3	3	2.77	1	497	916	50	2					1664	1128	322	121	282	
Methyl parathion	263.2	3	3.14	1	539	961	54						1734	1259	348	153		

DSSTox File Incidence

Prioritize Chemicals for ToxCast Phase I: Chemical Characteristics

- Meet minimal HTS phys-chem requirements
 - ▶ Soluble in DMSO / water
 - ▶ Not highly volatile
 - ▶ Molecular Weight in approx range [100-1,000]
 - ▶ Available in ~pure or standard form with structure
- Span diverse structural space
- Include clusters of similar chemicals

Chemical Classes in ToxCast_320 (Phase I)

• 309 Unique Structures

• Replicates for QC

• 291 Pesticide Actives

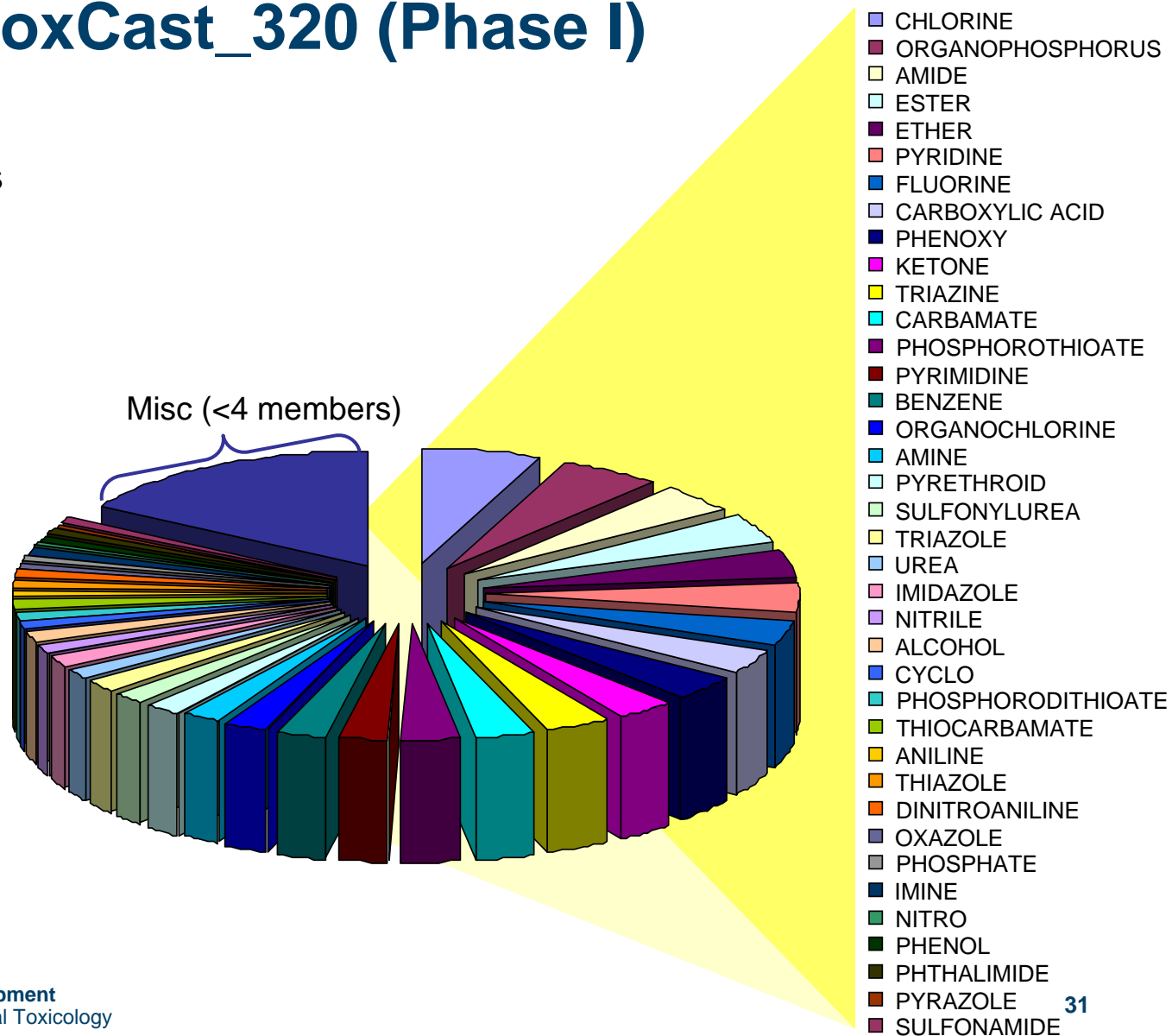
• 9 Industrial Chemicals

• 8 Metabolites

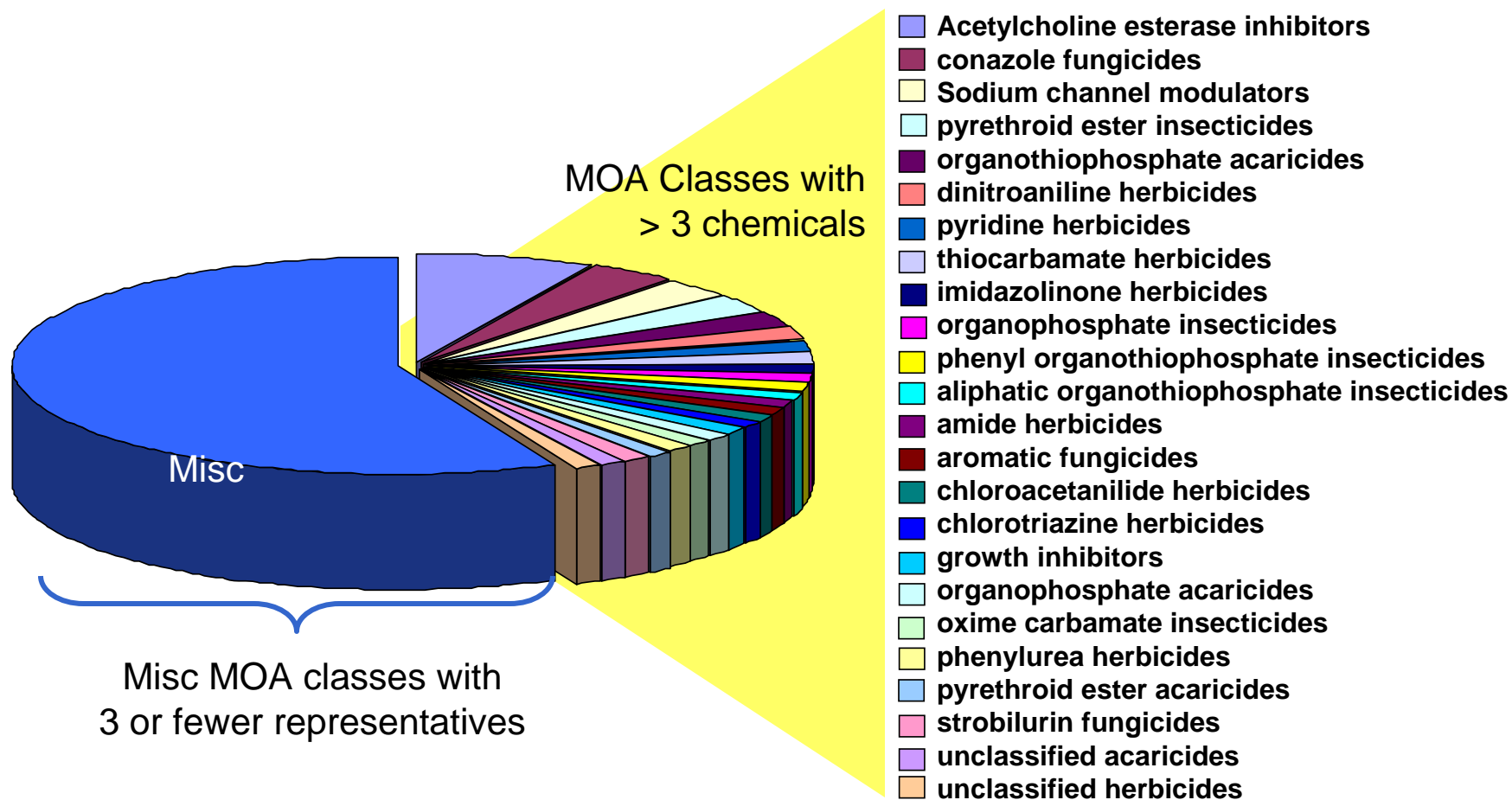
• 56/73 Proposed Tier 1
EDSP

• 14 HPV

• 11 HPV Challenge



Mode-of-Action Classes in ToxCast_320 (Phase I)





Distributed Structure-Searchable Toxicity (DSSTox) Public Database Network

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TOXCST: Research Chemical Inventory for EPA's Structure-Index File

**** Version 2b DSSTox Structure-Index File 08 February 2008**

➤ Incorporates new InChIKey field and a few structure modifications after additional quality of Analysis (COAs) of purchased chemicals.

Quick & Easy File Downloads: [FTP Download Instructions](#)

- [Description](#)
- [Source Website & Contact](#)
- [Main Citation](#)
- [SDF Fields](#)
- [SDF Content Summary](#)

File Types	Description	File Size
Documentation Files: TOXCST		
Log File	TOXCST LogFile 08Feb2008.pdf	37KB
Data Files: TOXCST		
SDF Structure Data File	TOXCST v2b 320 08Feb2008.sdf	922KB
• Data Table (no structures)	TOXCST v2b 320 08Feb2008_nostructures.xls	207KB
• Structures Table	TOXCST v2b 320 08Feb2008_structures.pdf (7 pp.)	1.4 MB

TOXCST SDF Content Summary

TOXCST SDF Content	Totals_v2b*
# Records	320
DSSTox Standard Chemical Fields	19
TOXCST Source Fields	2
Total # Fields	21
Chemical Content	Counts_v2b
STRUCTURE_ChemicalType:	
defined organic	313
inorganic	1
organometallic	6
no structure	0
STRUCTURE_TestForm_DefinedOrganic:	
parent	299
complex	8
salt	6
salt complex	0
TestSubstance_Description:	
single chemical compound	308
macromolecule	0
unspecified or multiple forms	0
mixture or formulation	12

EPA Pesticide Programs: Data Evaluation Records (DERs)

- Used for hazard identification and characterization
- Study Types
 - Chronic
 - Cancer
 - Subchronic
 - Multigeneration
 - Developmental
 - Others: DNT, Neurotox, Immu
- Derive Endpoints (NOAEL)
 - Systemic
 - Parental
 - Offspring
 - Reproductive
 - Maternal
 - Developmental
- Critical Effects for Endpoints

\$10,000,000

DER Format

- Study Identifiers
 - Tested Chemical Information
 - IDs
 - Name
 - Purity
 - Study Type IDs
 - Reviewer Information
- Citation(s)
- Executive Summary
 - Summary Study Design
 - Summary Effects
 - Endpoints (NOAEL/LOAEL)
- Test Materials
 - Chemical Properties
 - Animal Information
 - Species
 - Strain
 - Husbandry
- Results (full dose-response)
 - Clinical signs
 - Body weight
 - Clinical Chemistry/ Hematology
 - Gross Pathology
 - Non-neoplastic Pathology
 - Neoplastic Pathology
 - Parental vs. Offspring
 - Maternal vs. Fetal

Extraction of DER information

STUDY TYPE: Combined chronic toxicity/oncogenicity feeding – Rat
OPPTS 870.4300 [§83-5]

DP BARCODE: D257223
P.C. CODE: 111901

SUBMISSION CODE: S564270
TOX. CHEM. NO.: 497AB

TEST MATERIAL (PURITY): Imazalil (purity ≥97.4%)
SYNONYMS: R023979

CITATION: Van Deun, K. 1999. Combined oral chronic toxicity/carcinogenicity study with Imazalil in the SPF Wistar rat. Dept. Toxicology, Janssen Research Foundation, 2340 Beerse, Belgium. Laboratory report number, 3817, June 8, 1999. MRID 44858001. Unpublished.

SPONSOR: Janssen Pharmaceutica N.V., 2340 Beerse, Belgium

EXECUTIVE SUMMARY:

In a chronic toxicity/oncogenicity study (MRID 44858001), Imazalil (≥97.4% a.i.) was administered in the diet to groups of 50 male and 50 female Hannover substrain (SPF) Wistar-derived rats at concentrations of 0, 50, 200, 1200, or 2400 ppm (equivalent to 0.0, 2.7, 10.8, 65.8, and 134.8 mg/kg/day for males and 0.0, 3.6, 14.6, 85.2, and 168.3 mg/kg/day for females) for two years. All rats were observed daily for clinical signs of toxicity and morbidity, weighed weekly, and food consumption monitored biweekly. Blood and urine samples were collected after 6, 12, and 18 months of treatment and at study end. Surviving rats were sacrificed after 104 weeks of treatment. All rats were necropsied and the tissues and organs inspected grossly and microscopically for toxicity-related effects and the carcinogenic potential of Imazalil.

The absolute weights of most organs were decreased while their weights relative to body weight increased for male and female rats in the 1200 and 2400 ppm treatment groups. These effects are considered related to inanition and inappetence and not a direct result of Imazalil treatment. However, effects found in the liver and thyroid were considered directly related to treatment. The absolute liver weight of male rats in the 2400 ppm group was increased while it was decreased in female rats. The associated relative liver weights of male and female rats in the 1200 and 2400 ppm groups were significantly increased 9-26%. In addition, the absolute and relative thyroid weights of male but not female rats in the 1200 and 2400 ppm groups were increased.

The effect of treatment on the liver (males and females) and thyroid (males only) were confirmed microscopically, but had distinct sex-related etiologies. The incidence of clear cell and basophilic foci was equivocal while eosinophilic foci were significantly increased for male rats in the 2400 ppm group. In female rats of the 2400 ppm group, the incidences of clear cell and basophilic foci were significantly decreased but the incidence of eosinophilic foci was unaffected. Also, the incidence of hepatocyte fatty vacuolation was increased only in male rats of the 1200 ppm and 2400 ppm groups while the incidence of pigmentation was increased only in females of the 200, 1200, and 2400 ppm groups. In addition, the location of hepatocellular hypertrophy was distinctly different. Female rats in the 1200 and 2400 ppm groups had significant increases in centrilobular and pericentral hypertrophy while male rats only had centrilobular hypertrophy. Finally, the incidence of thyroid follicular cell hyperplasia was increased only in male rats of the 1200 and 2400 ppm groups.

The lowest observed adverse effect level (LOAEL) for male and female rats was 1200 ppm (65.8 and 85.2 mg/kg/day, respectively) with a corresponding no observed adverse effect level (NOAEL) of 200 ppm (10.8 mg/kg/day males, 14.6 mg/kg/day females). These are based on the effects found on body weight, weight gain, and the macro- and microscopic effects noted in the liver of all rats and the thyroid of male rats.

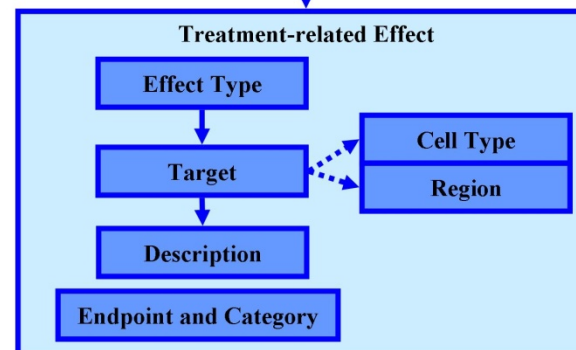
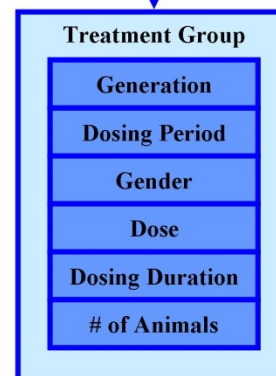
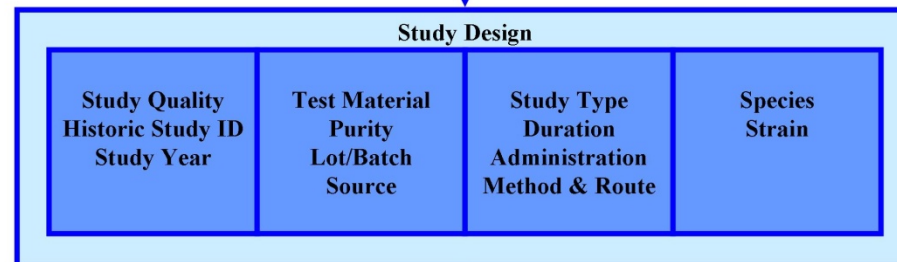
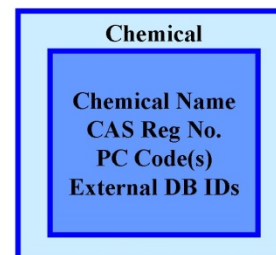
Male rats had a significant increase in the incidence of hepatocellular adenomas and thyroid follicular neoplasia while no increase was found for female rats. These results indicate a difference in the disposition of Imazalil between the sexes increases hepatic and thyroid neoplasia in male rats, likely through differences in metabolic activation of the test material.

This chronic toxicity/oncogenicity study in the rat is **Acceptable/guideline** and satisfies the guideline requirement for a combined chronic toxicity/oncogenicity study in rats [83-5]. No deficiencies were noted for this study.

ToxRefDB: *Toxicity Reference Database*

Fields/Data Elements Utilizing Standardized Vocabulary

- Chemical
- Study Quality
- Study Type
- Method & Route of Admin
- Species
- Strain
- Generation
- Dosing Period
- Gender
- Dosing Duration
- Effect Type
- Effect Target
- Effect Description
- Target Cell Type
- Target Region



Data Entry Completeness Score

Partially Complete (Effect Data)

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

ToxRefDB
Input Form

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY



Historic Study Identifiers

MRID# 44858001

Primary Study Year 1999

Supplemental MRID/Historic ID(s)

Study/Data Quality

Data Usability Acceptable Guideline (post-1998)

Study-Level Comments

Note: Thyroid weights inc in male and dec in female.
Thyroid neoplasia increase in male and decrease in female (both statistically significant)

Test Material Information

Search Chemical List

Search PC Code

Chemical Imazalil

Purity (%) 97.4

Lot/Batch# ZR023979G3F661

Source

Test Material (Chemical) Comments

ZR023979G3F661 / >97.4% a.i. /// ZR023979G3G641 / >98.6% a.i.

Study Type

Study Type Combined chronic toxicity/carcinogenicity

Study Duration Start 0 day

Additional Study Duration Information

Finish 104 week

Animal and Dose Information

Species rat

Method/Route of Administration

Strain [Other]

Feed

Animal and Dose Administration Comments (Including Not In List)

Strain: Hannover substrain (SPF) Wistar-derived

Study Effect List

Upload Form Info
Use Excel upload
form to add
treatment groups.
Click "Bulk
Upload"; Copy and
paste into form
and upload groups.

[Excel Treatment
Group Form](#)

Bulk Upload

Update List

EFFECT DATA

Click on "View or
Add Critical Effect
Data by Type" to
input effect data
for any treatment
group by effect
type.

Treatment Group List

Treatment Group Category	Gender Category	Dose Period Type	Dose	Duration	# / Group	View or Add Effect Data by Type
Adult (P1)	M	Initial-to-Terminal	2.7 mg/kg/day	104 week	50	
Adult (P1)	F	Initial-to-Terminal	3.6 mg/kg/day	104 week	50	
Adult (P1)	M	Initial-to-Terminal	10.8 mg/kg/day	104 week	50	
Adult (P1)	F	Initial-to-Terminal	14.6 mg/kg/day	104 week	50	
Adult (P1)	M	Initial-to-Terminal	65.8 mg/kg/day	104 week	50	
Adult (P1)	F	Initial-to-Terminal	85.2 mg/kg/day	104 week	50	
Adult (P1)	M	Initial-to-Terminal	134.8 mg/kg/day	104 week	50	
Adult (P1)	F	Initial-to-Terminal	168.8 mg/kg/day	104 week	50	

Delete Selected Treatment Group

Search Effect Vocabulary

Fisher's Exact Test

Toggle to Critical Effects Form

Edit Uploaded
Treatment GroupTreatment Group
Category

Adult (P1)

Gender #/group

M 50

Dose Period Type

Initial-to-Terminal

Dose Units

2.7 mg/kg/day

Duration Units

104 week

Save Delete New

Navigation buttons

Show all
Effects
[Assign
LOAELs]

Study Design

Treatment Groups

Study Design Level Controls



Search



Enter New Study

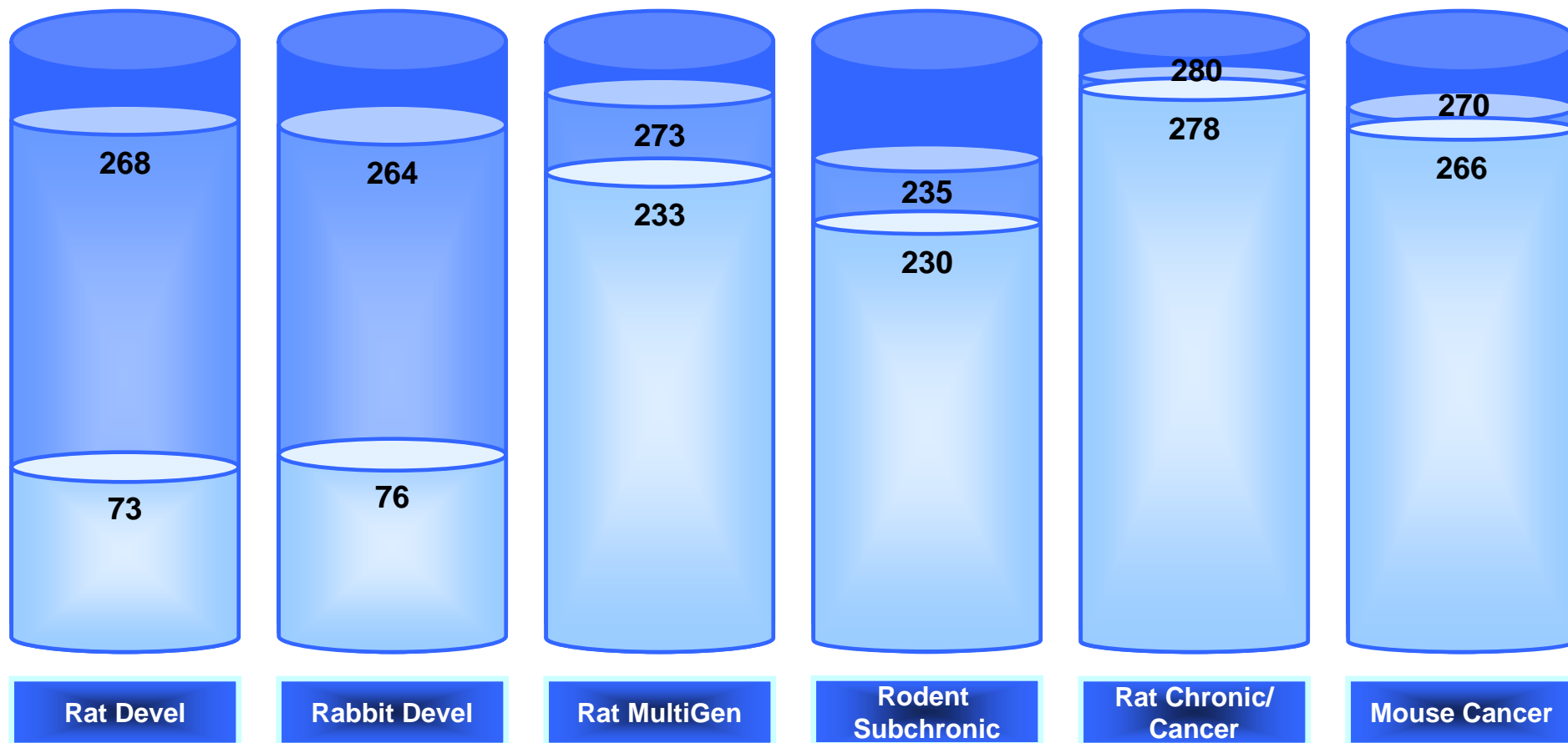
Filename/
Citation

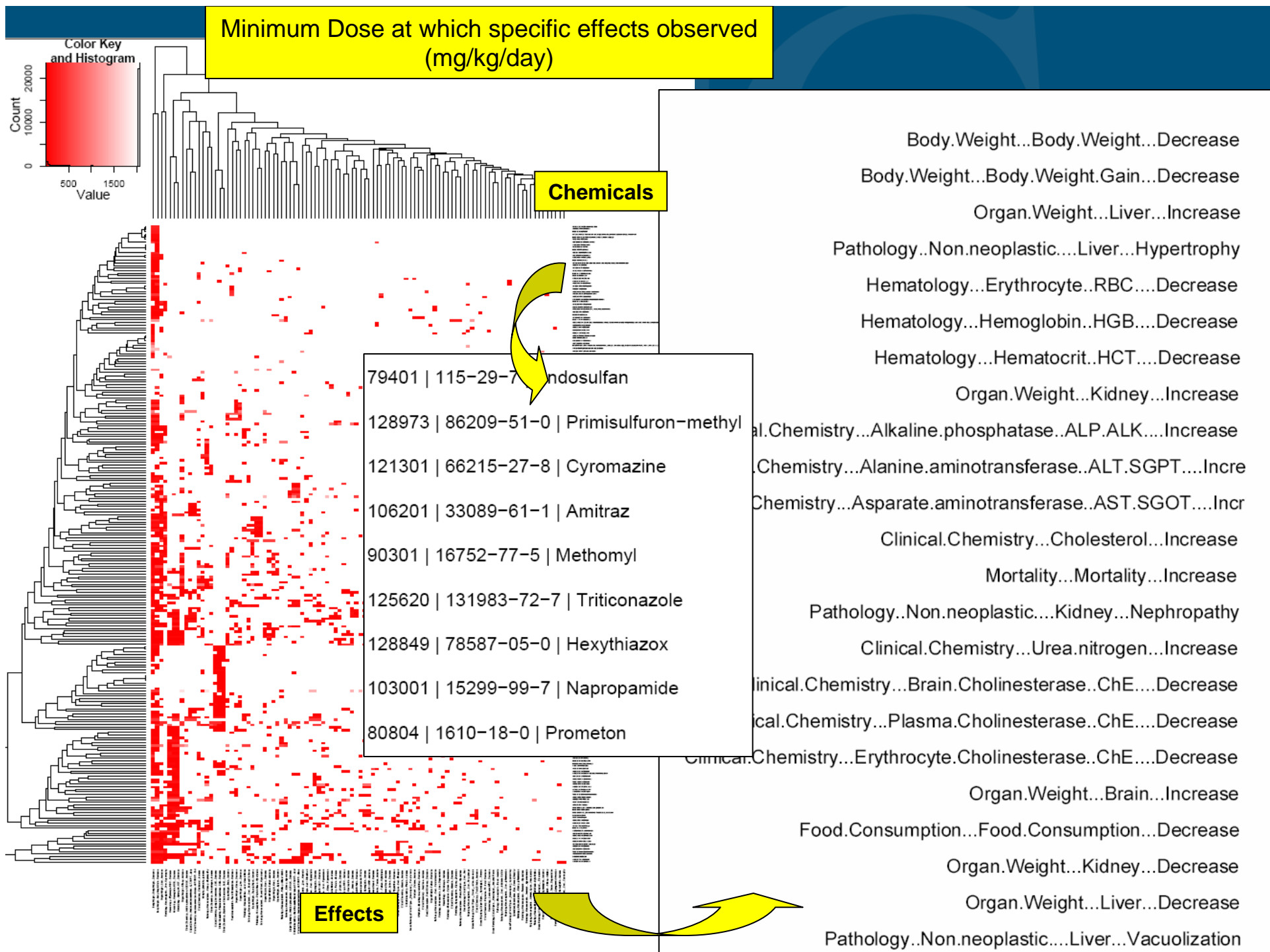
Toggle back to ToxRefDB Switchboard

Data Entry Status

ToxCast Phase I Chemicals Only

Total: 291 Pesticides







ToxCast HTS Assay Contractors and Collaborators

Compound Focus,

a subsidiary of

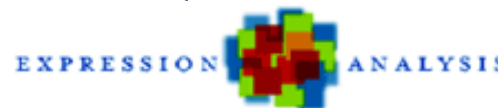
BioFocus DPI
A Galápagos Company



BioSeek



Nine contracts, one IAG and an MOU provide chemical procurement; hundreds of biochemical, cellular, tissue and genomic assays; model organisms; and the capacity to screen up to 10,000 chemicals

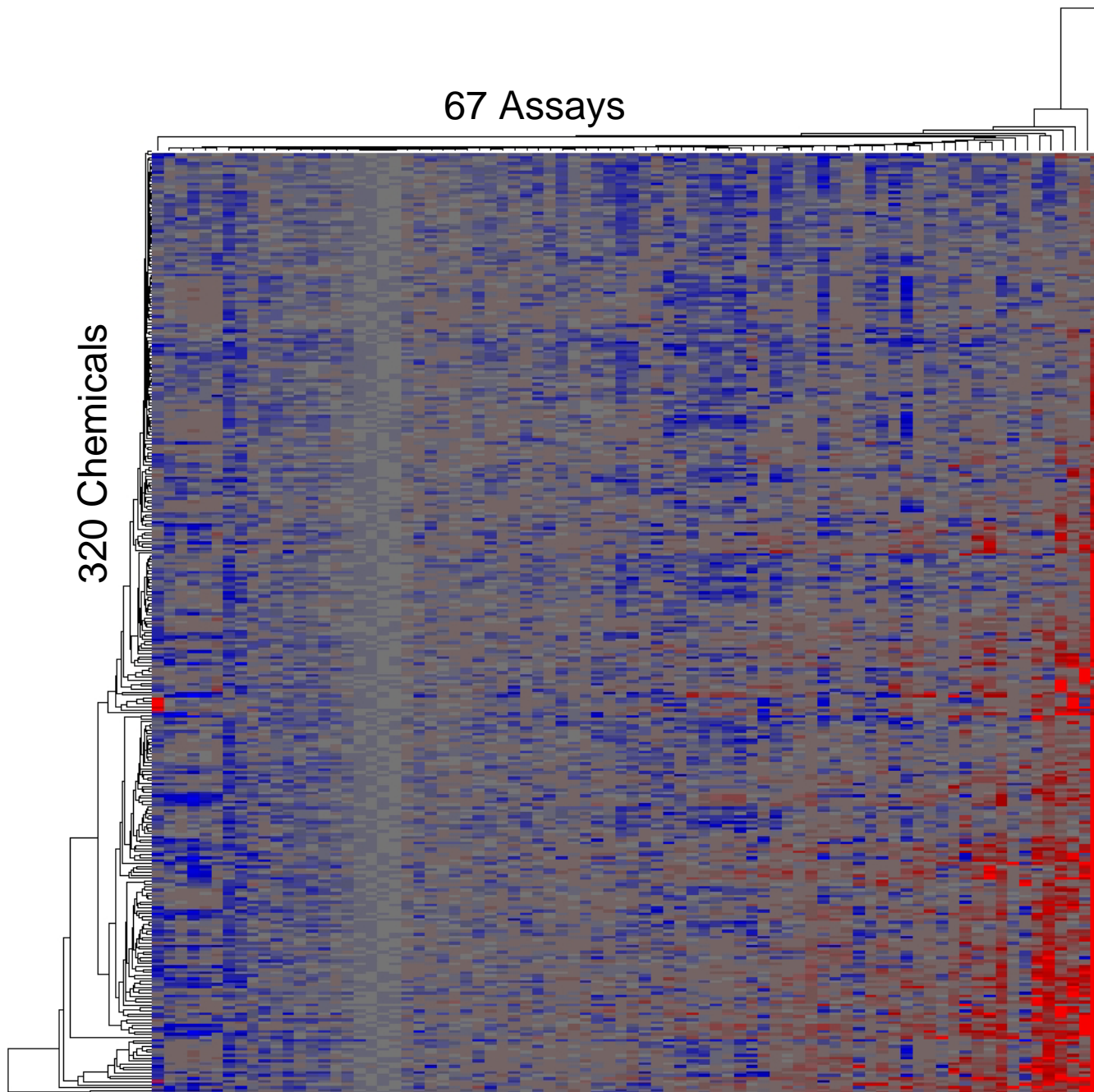


Assay Type	# Assays	# Endpoints	Assay Source	Comment	Source	ToxCast_320 Status 04dec2007
HTS	240	240	Human, rat, other	Enzyme inhibition, receptor binding	NovaScreen	Single concentration data delivered; multiple concentration to follow
uHTS	10+	10+	Human and rodent	Nuclear receptor reporter gene assays	NIH Chemical Genomics Center	Multiple concentration (11) data from 2 of 10 assays delivered
Reporter Gene Assays	2	67	HepG2 cells (human liver)	Nuclear receptor, transcription factor	Attagene	Single concentration data delivered; multiple concentration to follow
Genomics	1	22,000	Hepatocyte-Kupffer co-culture	PCR, microarrays	IVAL and Expression Analysis	Multiple concentration (5) in rat system underway
Kinetic Cell Growth	1	Kinetic	A549 cells (human lung)	Real time electrical impedance	ACEA Biosciences	Multiple concentration (8) data delivered
Cell Co-Culture	1	6	Human liver, lung, kidney cells	Cytotoxicity, shared metabolism	IVAL	Multiple concentration (8) data underway
Complex Cell Culture	8	87	Primary human cells	Cell signaling pathways	Bioseek	Multiple concentration (4) data delivered
HCS	1	11	HepG2 cells (human liver)	Imaging cytotoxicity	Cellumen	Multiple concentration (10) data delivered
Tissue Slice Culture	1	1	Rat liver, lung, kidney	Precision-cut Tissue slices	Hamner Institutes	Multiple concentration (5) data underway
Zebrafish	1	11	Danio rerio	Teratogenesis	Phylonix	Multiple concentration (3) data underway for 20 chemicals
TOTAL	265	22,433				

Attagene Heatmap

320 Chemicals

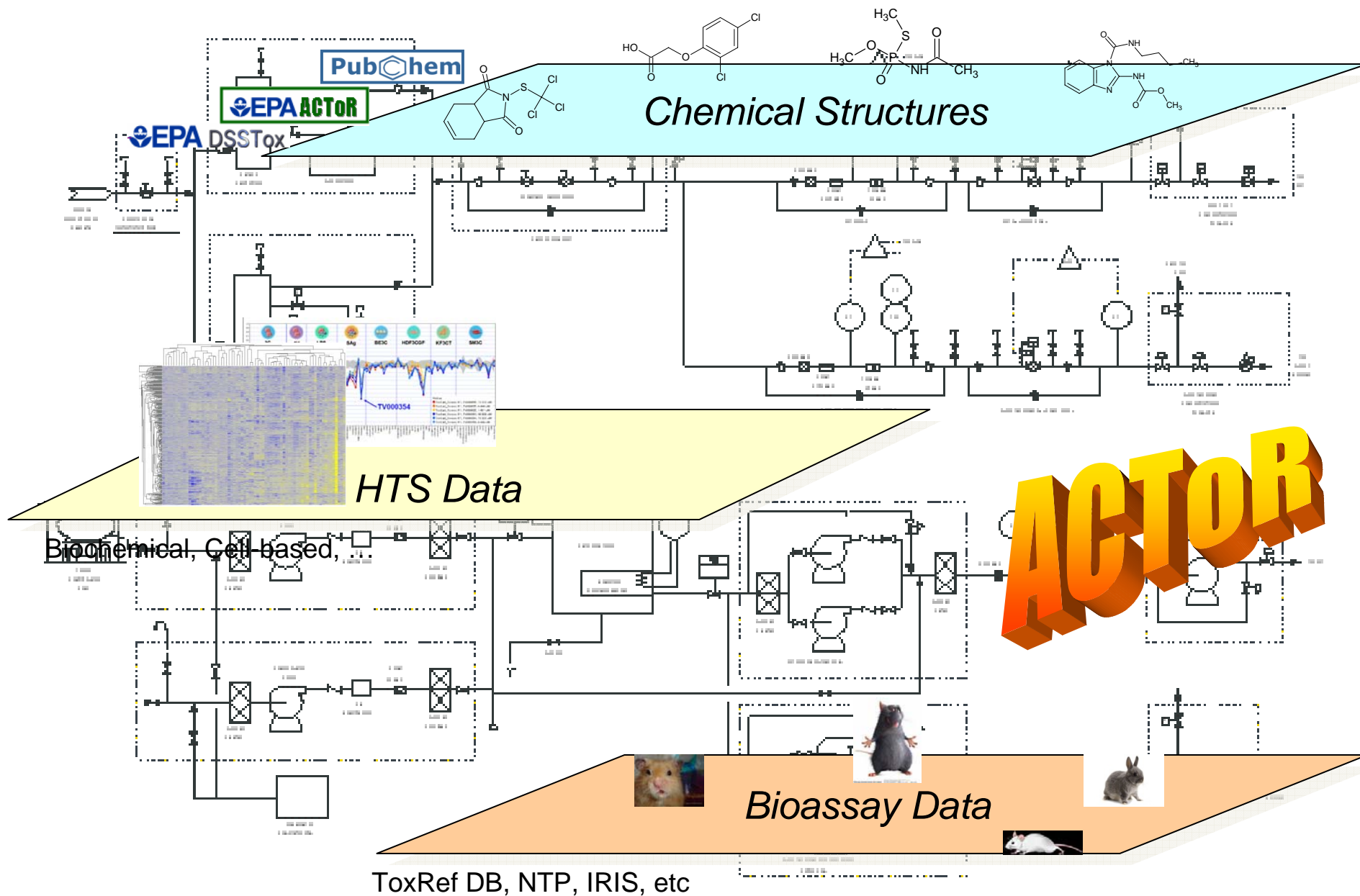
67 Assays



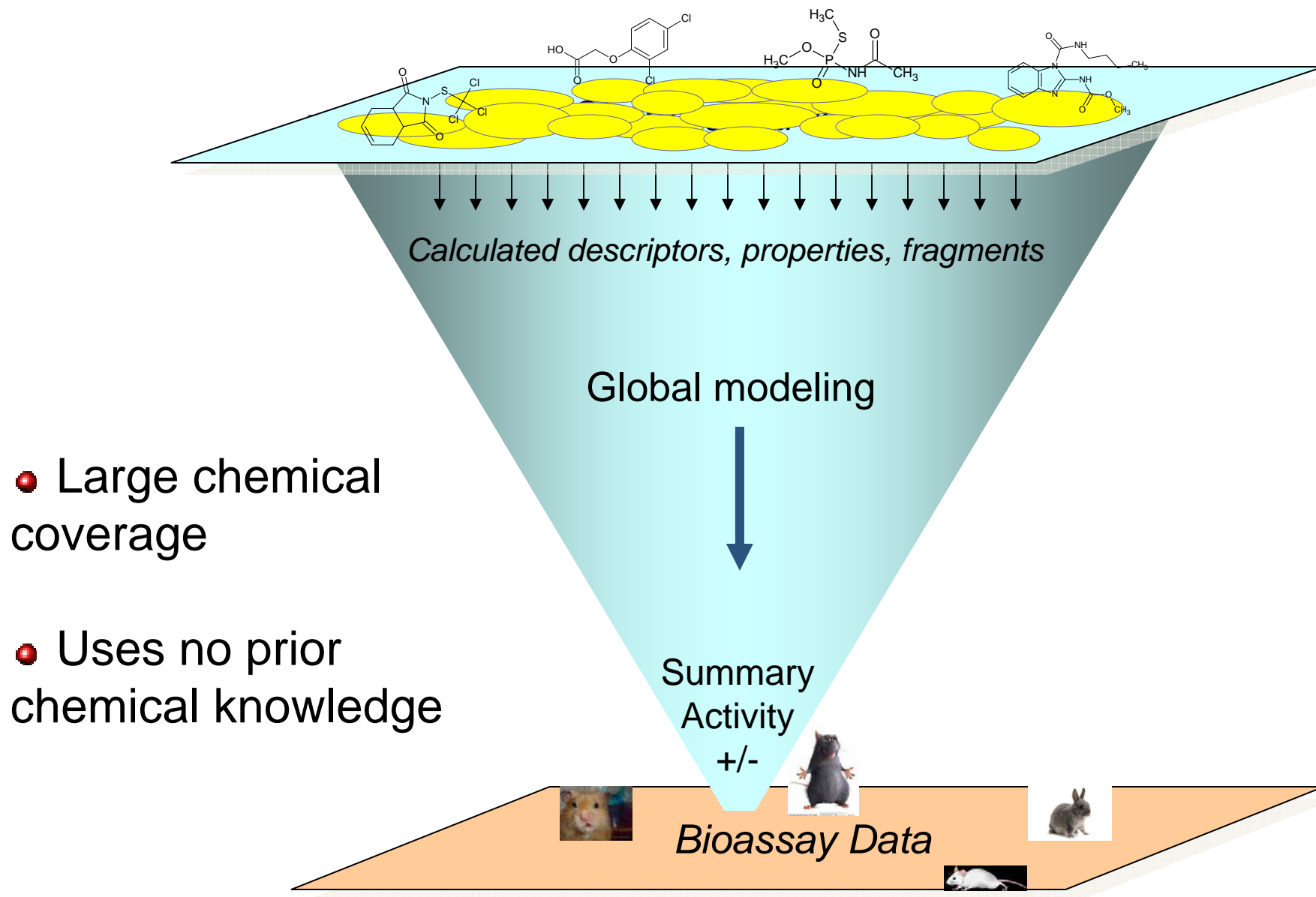
Part III

Incorporating SAR Concepts into ToxCast

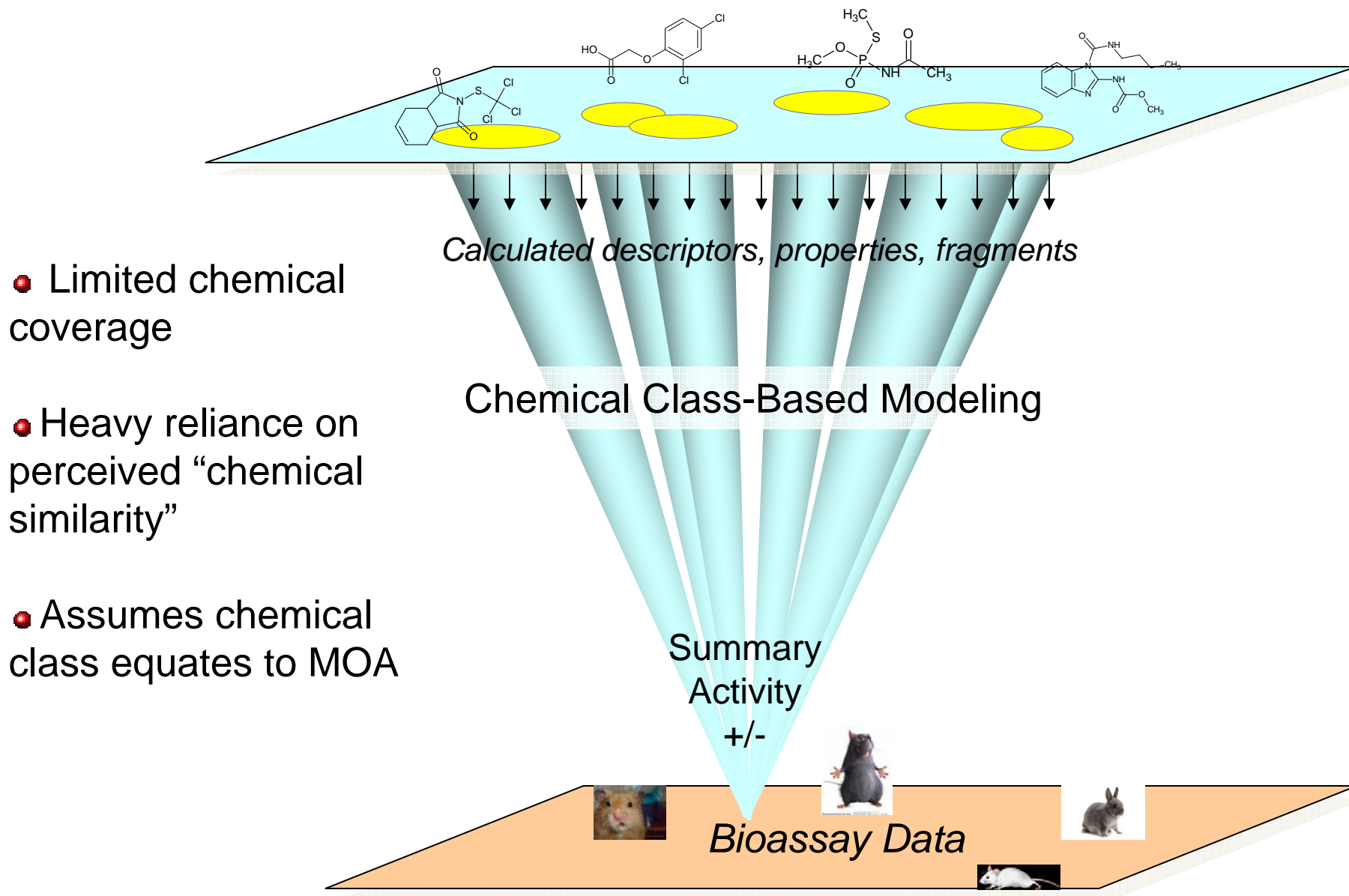
ToxCast: Multidimensional Data



Structure-Activity Approaches to Toxicity Prediction



Structure-Activity Approaches to Toxicity Prediction



Oncologic Carcinogenicity Estimation Expert System:

Chemical Class – Based Prediction Modules

Acyl and Benzoyl Halide Type Compounds
Acrylate Reactive Functional Groups
Acrylamide Reactive Functional Groups
Aflatoxin Type Compounds
Aldehyde Type Compounds
Aliphatic Azo and Azoxy Type Compounds
Alkanesulfonyl Ester Type Compounds
Alkyl Sulfate and Alkyl Alkanesulfonate Type Compounds
Aromatic Amine Type Compounds
Anhydride Type Compounds
Aryldiazonium Salts
C-Nitroso and Oxime Type Compounds
Carbamate Type Compounds
Carbamyl Halide Type Compounds
Coumarin and Furocoumarin Type Compounds
Dicarbonyl Type Compounds
Epoxide Reactive Functional Groups
Ethyleneimine Reactive Functional Groups
Haloalkylamine Reactive Functional Groups
Haloether Reactive Functional Groups
Halogenated Aromatic Hydrocarbon Type Compounds
Halogenated Cycloalkane Type Compounds

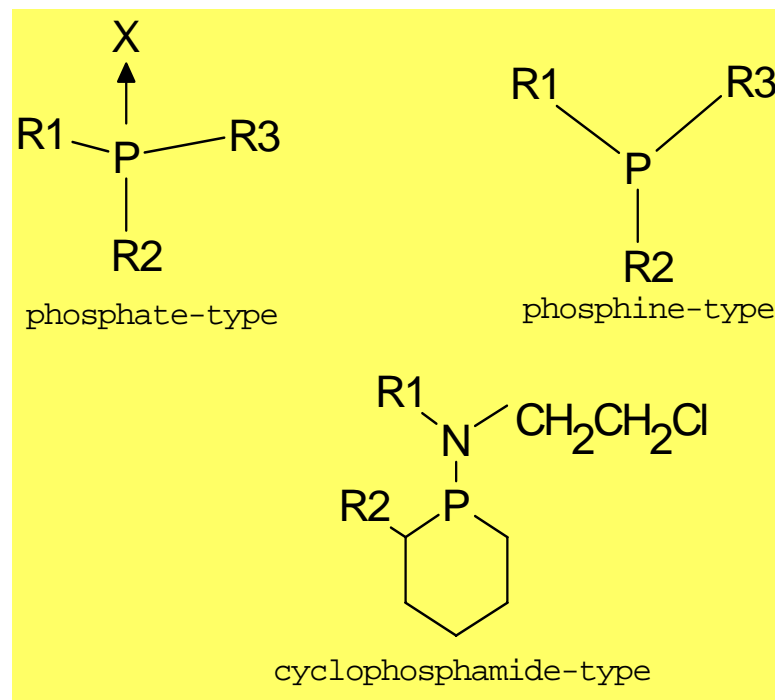
ortho-Halogenated Heterocyclic Type Compounds
Halogenated Nitroaromatic Type Compound
Halogenated Linear Aliphatic Type
Compounds
Halothioether Reactive Functional Groups
Hydrazo Type Compounds
Reactive Ketone Reactive Functional Groups
Lactone Type Reactive Functional Groups
Nitrosamide Type Compounds
Nitrosamine Type Compounds
Nitroalkane and Nitroalkene Type Compounds
Nitrogen Mustard Reactive Functional Groups
Organophosphorus Type Compound
Peroxide Type Compounds
Phenol Type Compounds
Phosgene Type Compounds
Polycyclic Aromatic Hydrocarbons -
Heterocyclic Polycyclic Aromatic Hydrocarbons -
Homocyclic Siloxane Type Compounds
Reactive Sulfone Reactive Functional Groups
Sulfur Mustard Reactive Functional Groups
Sultone Reactive Functional Groups
Thiocarbamate Type Compounds
Thiocarbonyl Type Compounds
Triazene Type Compounds
Urea Type Compounds

Oncologic Carcinogenicity Estimation Expert System:

Organophosphate

R1/R2/R3 = alkyl groups or aryl groups

P X-alkyl or X-aryl where X is oxygen, sulfur or in some cases nitrogen, linking the phosphorus to the alkyl/aryl group



phosphine-type → phosphine and phosphine-oxide

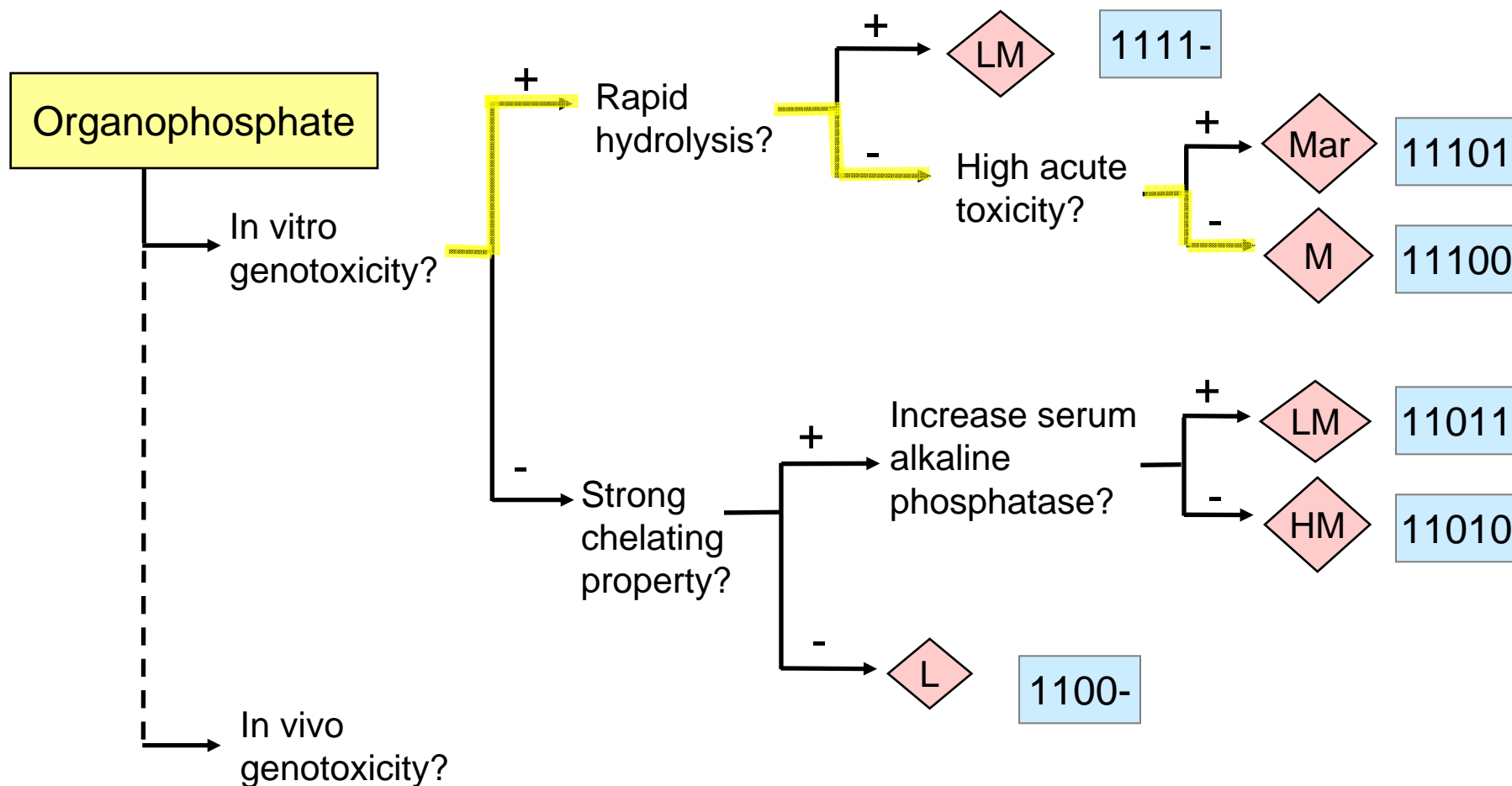
cyclophosphamide-type → cyclophosphamide, isophosphamide, trophosphamide

R1/R2/R3: alkyl (Cn), hydrogen (H), benzyl ($\text{CH}_2\text{C}_6\text{H}_5$), phenyl (C_6H_5), Morpholino, $\text{NR}'\text{R}''$ (where $\text{R}'\text{R}''$ can be one of the above).

X1/X2/X3/X4: Oxygen (O), Sulfur (S)

Substituents: Halogens (Cl, Br, I, F), hydroxyl (OH), carboxylic acid (COOH), sulfonic acid (SO_3H) and additionally alkyl (Cn) on the aryl ring..

Combining SAR and Biofunctional Information in Oncologic: *Predicting Carcinogenicity of Organophosphates*

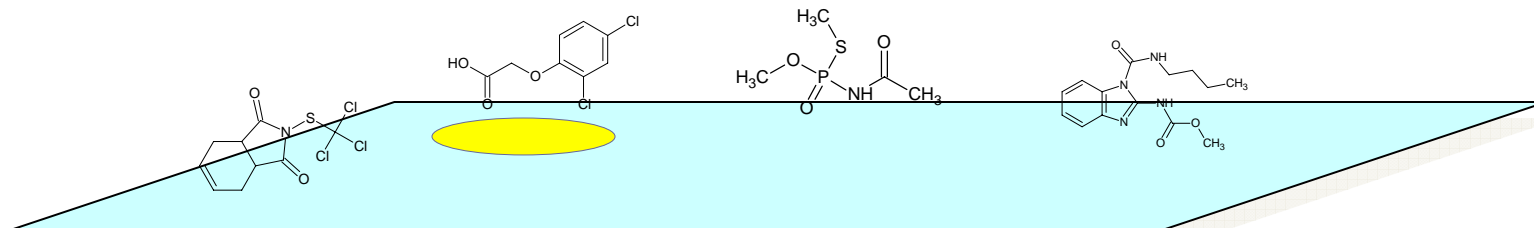


Oncologic Carcinogenicity Estimation Expert System: *Chemical Class – Based Prediction Modules*

Acyl and Benzoyl Halide Type Compounds
 Acrylate Reactive Functional Groups
 Acrylamide Reactive Functional Groups
 Aflatoxin Type Compounds
 Aldehyde Type Compounds
 Aliphatic Azo and Azoxy Type Compounds
 Alkanesulfonyl Ester Type Compounds
 Alkyl Sulfate and Alkyl Alkanesulfonate Type Compounds
 Aromatic Amine Type Compounds
 Anhydride Type Compounds
 Arylazo Type Compounds
 Aryldiazonium Salts
 C-Nitroso and Oxime Type Compounds
 Carbamate Type Compounds
 Carbamyl Halide Type Compounds
 Coumarin and Furocoumarin Type Compounds
 Dicarbonyl Type Compounds
 Epoxide Reactive Functional Groups
 Ethyleneimine Reactive Functional Groups
 Haloalkyl 1100101010000... 101001111100000
 Haloether Reactive Functional Groups
 Halogenated Aromatic Hydrocarbon Type Compounds
 Halogenated Cycloalkane Type Compounds

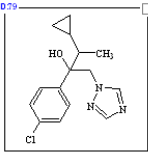
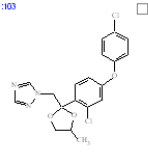
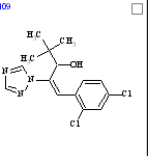
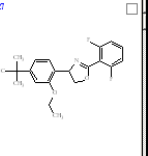
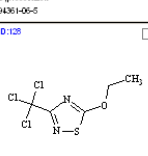
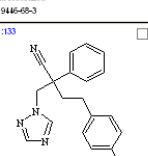
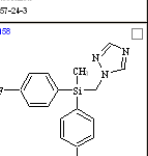
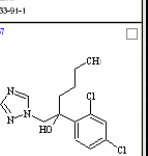
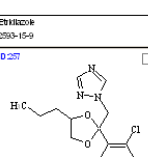
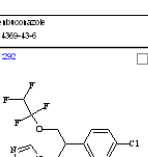
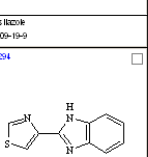
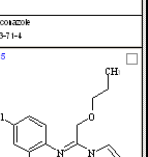

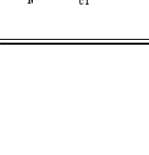
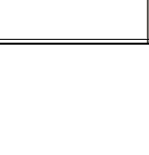
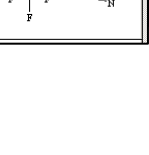
ortho-Halogenated Heterocyclic Type Compounds
Halogenated Nitroaromatic Type Compound
Halogenated Linear 1111- Type
CompoundsHalothioether Reactive Functional Groups
Hydrazo Type Compounds
Reactive Ketone Reactive Functional Gro 11101
Lactone Type Reactive Functional Groups
Nitrosamide Type Compounds
Nitrosamine Type Compounds 11100
Nitroalkane and Nitroalkene Type Compounds
Nitrogen Mustard Reactive Functional Groups
Organophosphorus Type Compound
Peroxide Type Compounds 11011
Phenol Type Compounds
Phosgene Type Compounds
Polycyclic Aromatic Hydrocarbons - 11010
HeterocyclicPolycyclic Aromatic Hydrocarbons -
Homocyclic Siloxane Type Compounds
Reactive Sulfone Reactive Functional Groups
00000010100000000101001... 1100-0
Sulfone Reactive Functional Groups
Thiocarbamate Type Compounds
Thiocarbonyl Type Compounds
Triazene Type Compounds
Urea Type Compounds

Bioactivity Profile of Structure Class

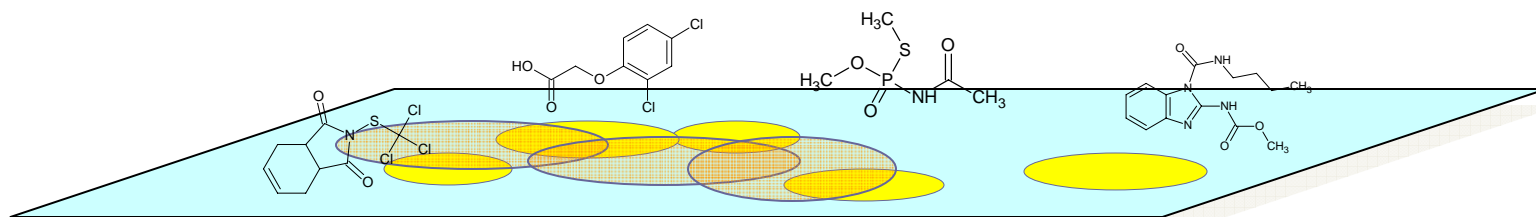


Sample HTS Results for Conazoles

- Patterns can inform SAR
- Chemical structures can suggest basis for activity differences

	NAME	CYP2C19	CYP2C9	CYP3A1	Dopamine Transporter (Human)	CYP2D2	Androgen Receptor	Dopamine Transporter (Rat)	CYP2B6	CYP2D1	CYP3A4	Progesterone Receptor	Benzodiazepine Receptor
	Cyproconazole	1	1	1	1	1	0	1	0	0	1	0	0
	Difenconazole	1	1	1	1	1	0	0	1	1	0	0	0
	Diniconazole	1	1	1	0	1	0	0	0	1	1	1	0
	Fenbuconazole	1	1	0	0	0	0	0	0	0	1	0	0
	Flusilazole	1	1	1	0	1	1	0	1	1	NA	1	1
	Hexaconazole	1	1	1	1	1	0	1	1	1	NA	1	0
	Imazalil	1	1	1	1	1	1	1	1	1	1	1	1
	Myclobutanil	1	1	1	1	0	0	0	0	0	NA	0	0
	Paclobutrazol	1	0	1	1	0	1	1	0	1	1	0	0
	Prochloraz	1	1	1	1	1	1	1	1	1	NA	1	1
	Propiconazole	1	1	1	0	0	0	0	1	0	NA	0	1
	Tetraconazole	1	1	1	0	1	1	0	1	0	1	1	0
	Triadimefon	1	1	0	1	1	1	1	0	0	1	0	1
	Triadimenol	1	0	0	1	0	1	1	0	0	0	0	0
	Triflumizole	1	1	1	1	1	1	0	1	1	1	1	1
	Triticonazole	1	1	1	1	0	1	1	0	0	NA	0	0
	Totals	16	14	13	11	10	9	8	8	8	8	7	6

Structure Class vs Bioactivity Class

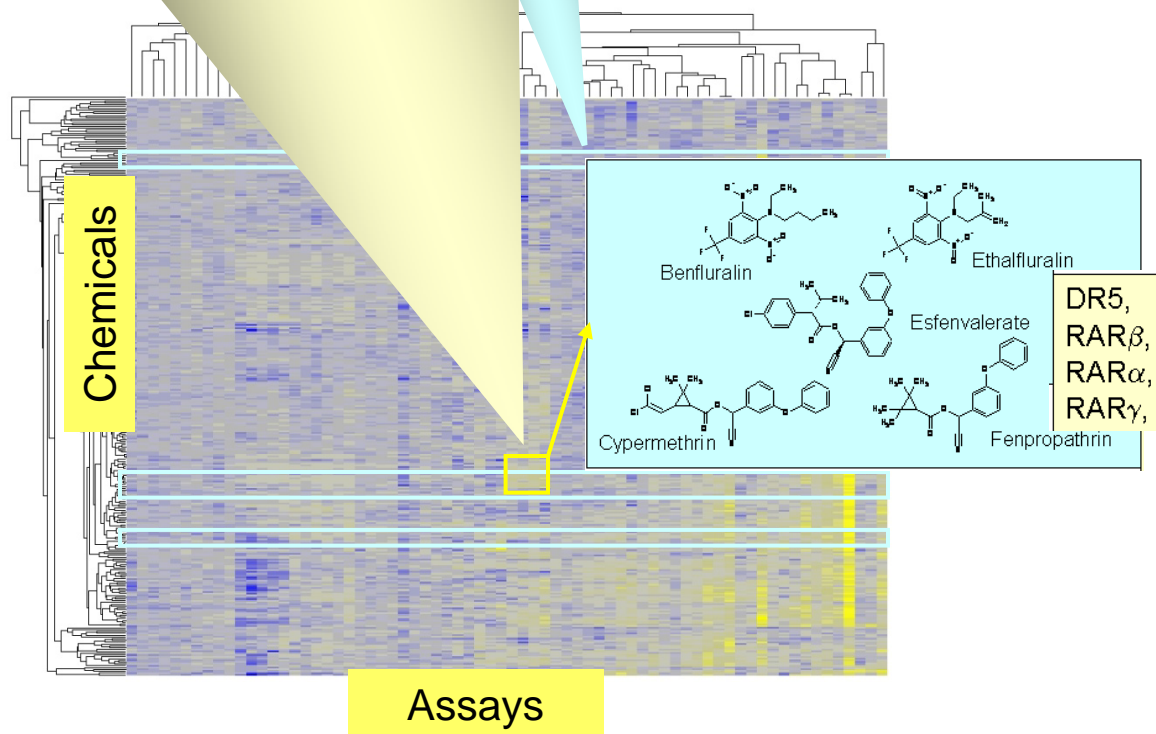


Chemical structure class:

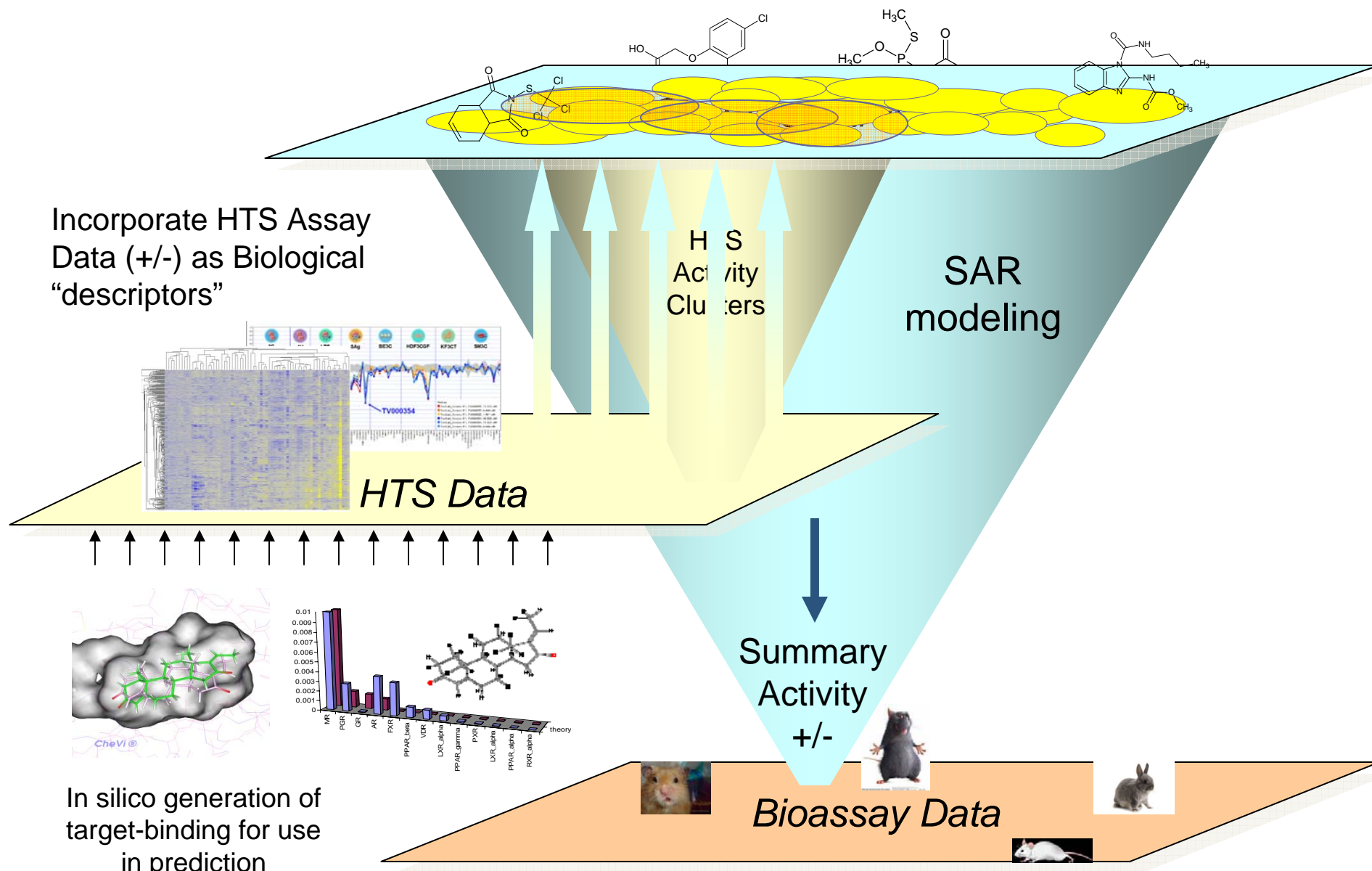
- Cluster according to activity and mechanism
- Differences in activity profiles can discriminate within structure class

Bioactivity profile class:

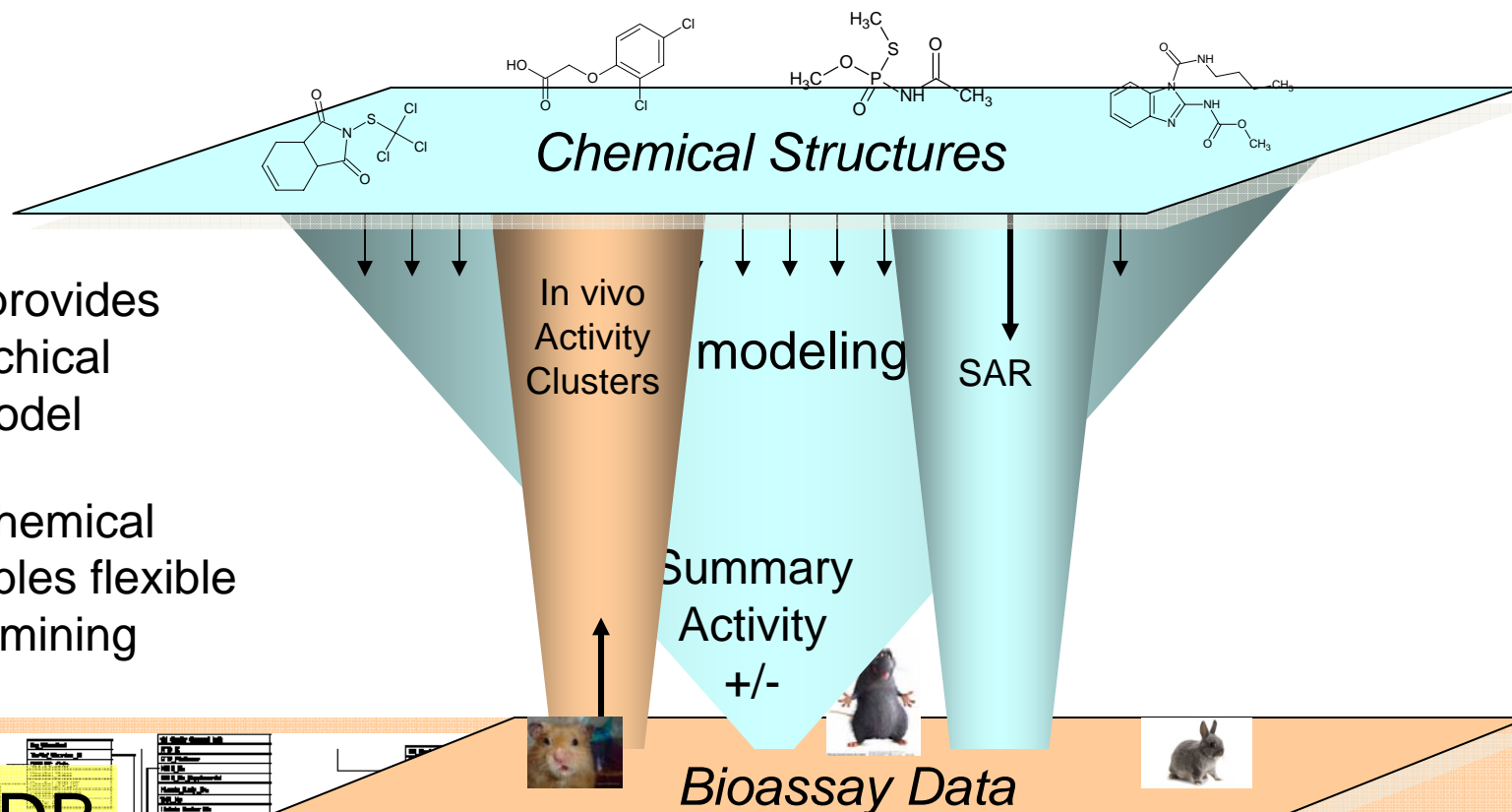
- Can project onto multiple chemical classes
- Potentially broader coverage of chemical space
- Implies mechanistic similarity



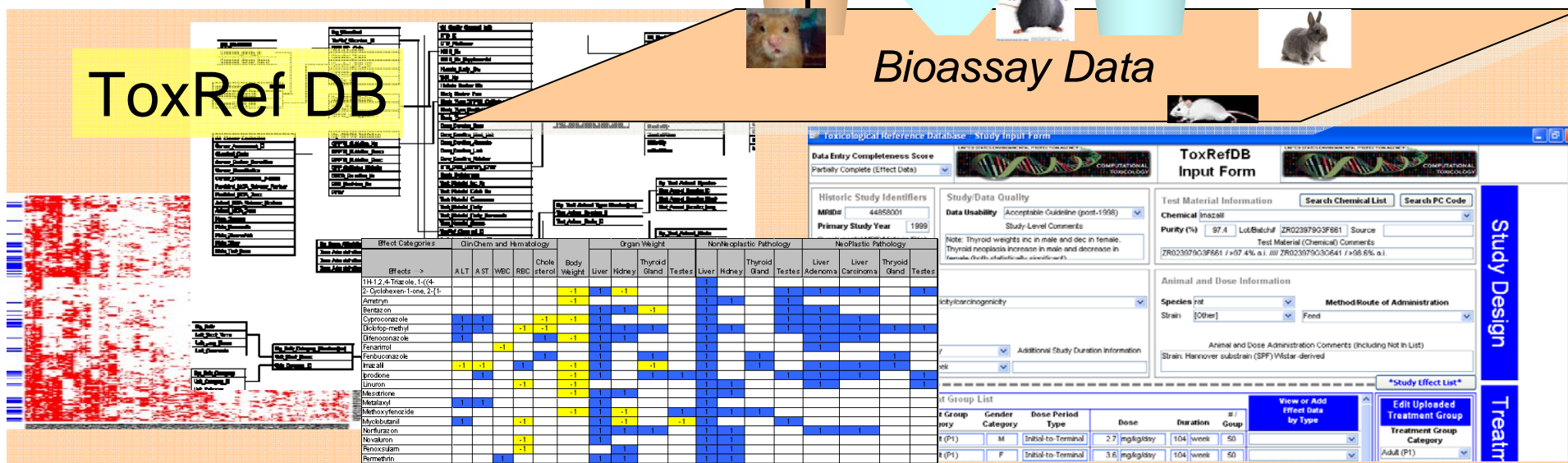
Structure-Activity Approaches to Toxicity Prediction



Use of Bioassay Activity Categories in SAR



- ToxRef DB provides detailed hierarchical toxicity data model
- Linkage to chemical structures enables flexible SAR and data mining



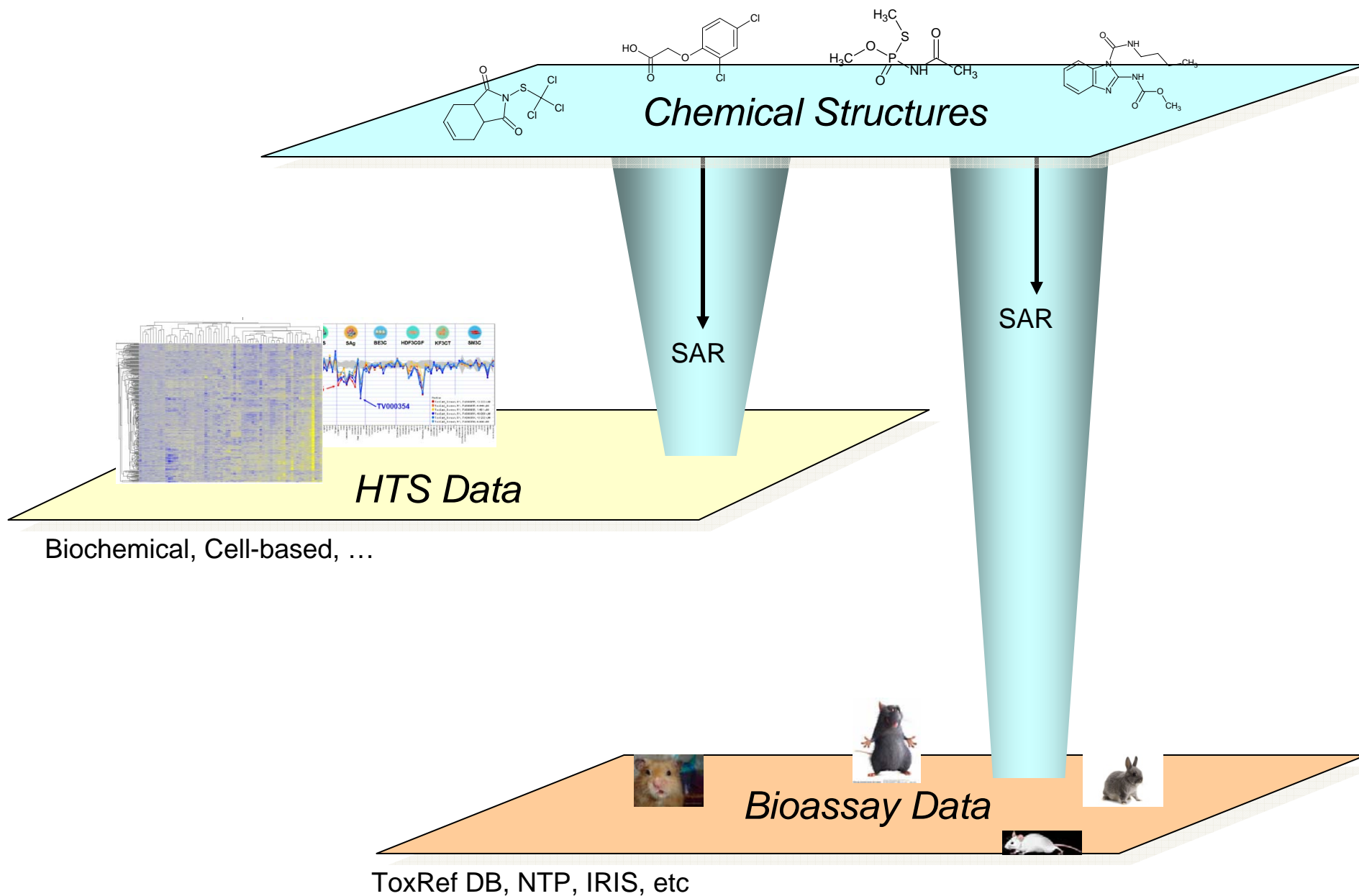
ToxRefDB Profiling of Liver Effects for Pesticides

Liver non-neoplastic histopathology and increased organ weight are often associated with tumors and cancer

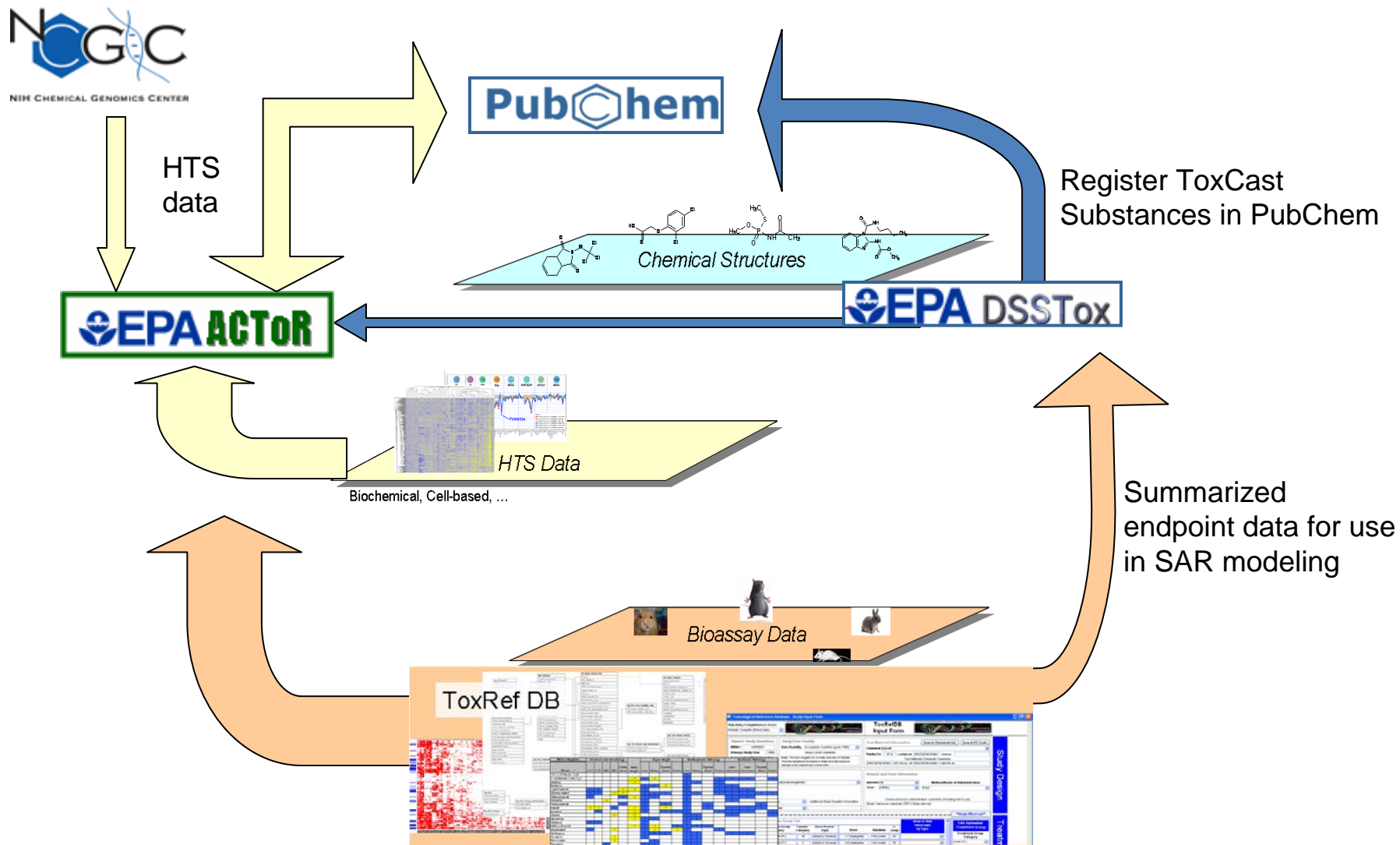
Effect Categories	ClinChem and Hematology						Organ Weight				NonNeoplastic Pathology				NeoPlastic Pathology			
	ALT	AST	WBC	RBC	Cholesterol	Body Weight	Liver	Kidney	Thyroid Gland	Testes	Liver	Kidney	Thyroid Gland	Testes	Liver Adenoma	Liver Carcinoma	Thyroid Gland	Testes
1H-1,2,4-Triazole, 1-((4-2-Cyclohexen-1-yl)-2-((1-1-1,1,1-trifluoro-2-methyl-2-phenylethoxy)ethyl)-1H-1,2,4-triazol-5-yl)-1H-1,2,4-triazole						-1	1	-1			1			1	1			1
Ametryn						-1	1				1	1		1				
Bentazone							1	1	-1		1			1	1			
Cyproconazole	1	1			-1	-1	1				1			1	1	1		
Diclofop-methyl	1	1		-1	-1		1	1	1		1	1		1	1	1	1	1
Difenoconazole	1				1	-1	1	1			1				1	1		
Fenarimol			-1				1				1				1			
Fenbuconazole					1		1		1		1		1				1	
Imazalil	-1	-1		1		-1	1		-1		1		1		1	1	1	
Iprodione		1					1		1	1				1	1	1		1
Linuron				-1		-1	1				1	1			1			1
Mesotrione						-1	1	1			1	1						
Metazachlor	1	1					1				1							
Methoxyfenozide						-1	1	-1		1	1	1	1					
Myclobutanil	1						1	-1		-1	1			1				
Norfenoxuron							1	1	1		1	1			1	1		
Novaluron							1				1	1						
Penoxsulam							1	1			1	1						
Permethrin							1	1			1	1						
Picloram	-1						1				1	1						
Primisulfuron-methyl						-1	1	-1			1			1	1	1		
Propiconazole					-1	-1	1				1				1			
						-1	1	-1			1							
						-1	1	1	-1		1							1
						-1	1			1	1		1		1		1	
							1				1	1			1			
						-1	1			1	1	1		1				
Acetochlor					1	-1		1				1						
Amdro						-1				-1				1				
Atrazine						-1						1						
Azinphosmethyl																		
Bifenazate			-1	-1	-1	-1		-1							1			
Cyfluthrin						-1												
Deltamethrin						-1												
Diethyl 1-(2,4-dichlorophenyl)-3-methyl-5-pyrazolone-4-carboxylate																		
Fenitrothion				-1		-1												
Imazapyr													1					
lambda-Cyhalothrin						-1												
Mesosulfuron-methyl			1	1														
Methyl parathion				-1	1		1	1										
Metolachlor						-1												
Molinate										-1				1				1
Oryzalin				-1		-1	1	1	1	1		1	1				1	
Oxamyl						-1												
Parathion		-1																

Activity Profile is refined "Endpoint" for SAR modeling

Structure-Analog Approaches



ToxCast: Data Publication & Exploration



ToxCast_320

Bioactivity Analysis:

Retrieve all
bioassay data
in PubChem for
ToxCast_320

482 Bioassays
45 Compounds

NCBI PubChem

PubChem BioAssay [GO]
PubMed | Entrez | Structure | PubChem | Help

PubChem » BioAssay Services » BioActivity Analysis: Summary

BioActivity Analysis: 482 Bioassays (473 Tested) and 45 Compounds

Summary [DataTable] [Structure-Activity]

Compounds: 45 (9 shown) [?]
Revise Compound Selection:
[Select Active](#)
[Add Active](#)
[Add Tested](#)
[Add Similar Compounds](#)

BioAssays: 482 [?]
Revise BioAssay Selection:
[Select Active](#)
[Select Tested](#)
[Add Active](#)
[Add Tested](#)
[Add Related BioAssays](#)
[Selected BioAssays](#)

9 chemical structures are displayed in a 3x3 grid.

Total Pages: 24 Display: 20 Go To Page 1 [Navigation Buttons]

#	<input type="checkbox"/>	AID	Active	Inactive	Discrepant	Tested	Outcome Method	Name
1	<input checked="" type="checkbox"/>	884	16	15	3	39	Confirmatory	qHTS Assay for Inhibitors and Substrates of Cytochrome P450 3A4
2	<input checked="" type="checkbox"/>	544	9	15	1	30	Confirmatory	Cell Viability - SH-SY5Y
3	<input checked="" type="checkbox"/>	541	7	20		30	Confirmatory	Cell Viability - NIH 3T3
4	<input checked="" type="checkbox"/>	426	7	20	1	30	Confirmatory	Cell Viability - Jurkat
5	<input checked="" type="checkbox"/>	964	6	20	1	30	Confirmatory	Cell Viability - LYMP1-003 - Assay at 40 hr
6	<input checked="" type="checkbox"/>	981	6	23	1	30	Confirmatory	Cell Viability - LYMP2-010
7	<input checked="" type="checkbox"/>	893	6	20	1	30	Confirmatory	qHTS Assay for Inhibitors of HSD17B4, hydroxysteroid (17-beta) dehydrogenase 4
8	<input type="checkbox"/>	167	6	20	1	30	Confirmatory	NCI Yeast Anticancer Drug Screen. Data for the bub3 strain
9	<input type="checkbox"/>	165	5	6	1	10	Confirmatory	NCI Yeast Anticancer Drug Screen. Data for the cln2 rad14 strain

Selected bioassays

BioActivity Analysis: 7 BioAssays and 45 Compounds (8 Tested)

[Summary](#) [Data Table](#) [Structure-Activity](#)

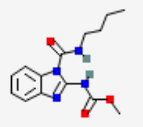
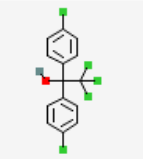
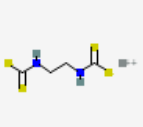
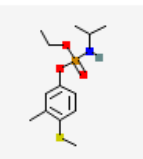
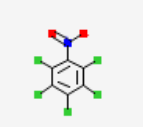
AIDs: 426 541 544 884 893 964 981

Total BioAssay Result Count: 8

[Data Table, Concise](#) [Data Table, Complete](#) [Plot](#) [Select](#)

Show details for
7 Bioassays

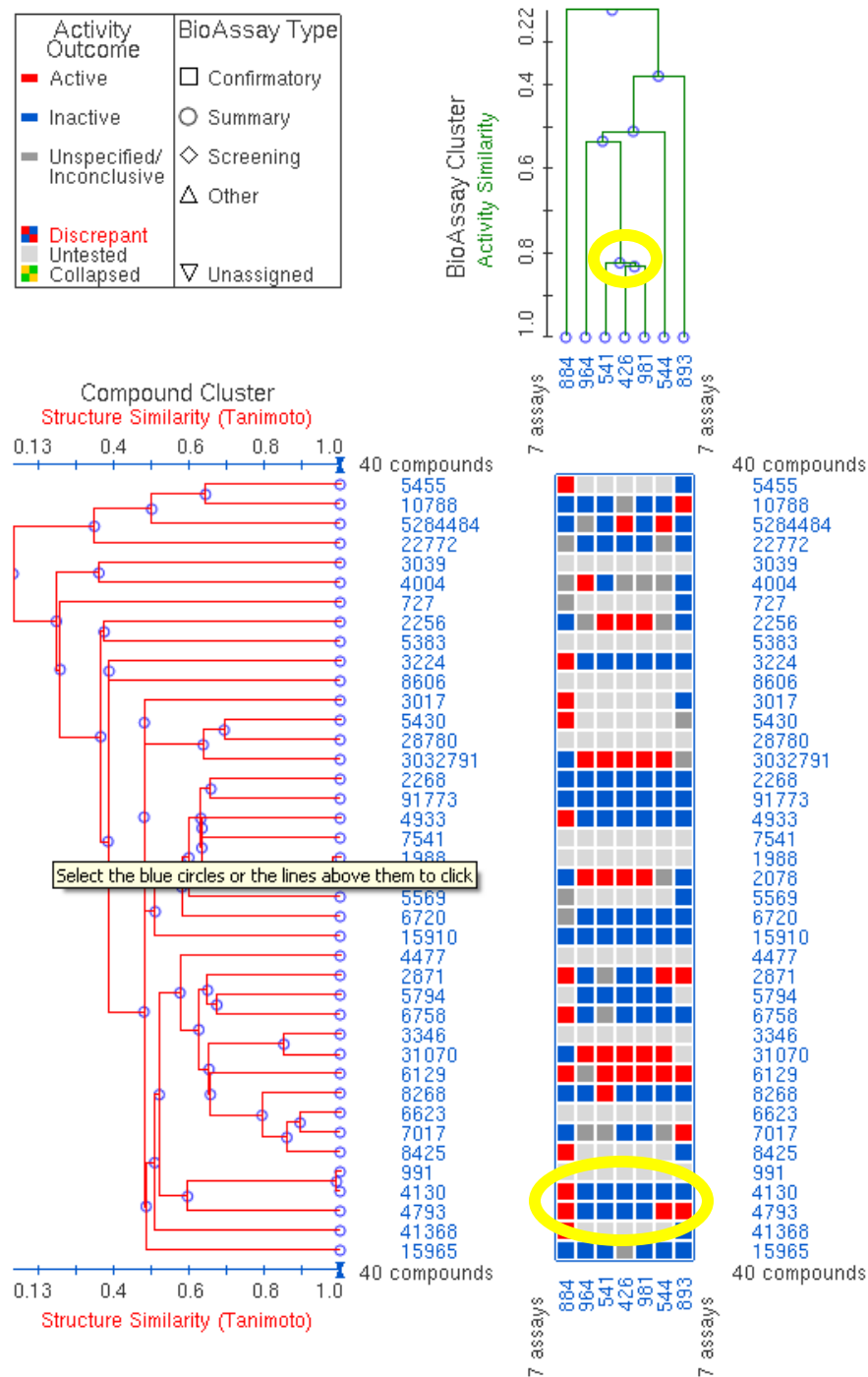
Page: 1 of 1 Sort: ☐ ☒ ☐ ☐ (Click the result table header to sort.)

#	Structure	SID	CID	Active Count	AID_426 Score	AID_426 Outcome	AID_541 Score	AID_541 Outcome	AID_544 Score	AID_544 Outcome	AID_884 Score	AID_884 Outcome	AID_884 Potency [uM]*	AID_893 Score	AID_893 Outcome	AID_893 Potency [uM]*	AID_964 Score	AID_964 Outcome	AID_981 Score	AID_981 Outcome
1		17389169	28780	2	54	Active	0	Inactive	56	Active	0	Inactive		0	Inactive		21	Inconclusive	12	Inconclusive
2		17389924	8268	0	39	Inconclusive	0	Inactive	0	Inactive	0	Inactive		0	Inactive		0	Inactive		Inactive
3		17389681	5284484	1	0	Inactive	0	Inactive	0	Inactive	20	Inconclusive	25.1189	64	Active	5.0119	0	Inactive		Inactive
4		17388788	31070	1	0	Inactive	0	Inactive	70	Active	21	Inconclusive	10	0	Inactive		0	Inactive		Inactive
5		17389772	6720	1	0	Inactive	73	Active	0	Inactive	0	Inactive		0	Inactive		0	Inactive		Inactive

Structure-Activity Bioactivity Analysis:

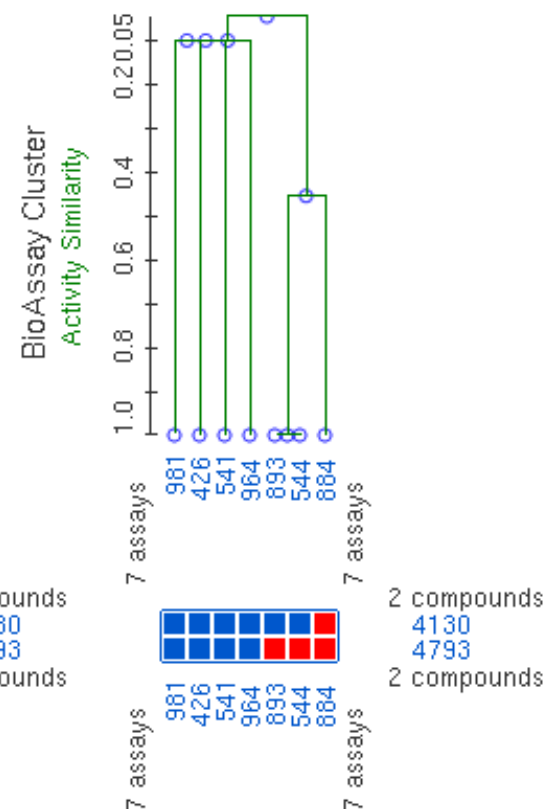
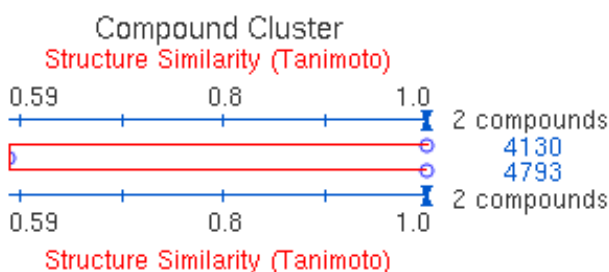
7 bioassays, 45 Actives

View Bioassay Profile
by Structure Similarity
Cluster

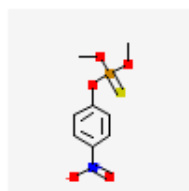


View bioassay profile of structure similarity cluster

Activity Outcome	BioAssay Type
Active	Confirmatory
Inactive	Summary
Unspecified/Inconclusive	Screening
Discrepant	Other
Untested	Unassigned
Collapsed	



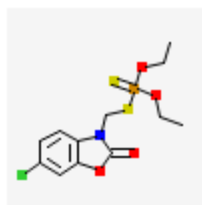
1: CID: [4130](#)



Parathion-methyl; METHYL PARATHION; Methylparathion ...
IUPAC: dimethoxy-(4-nitrophenoxy)-sulfanylidene phosphorane
MW: 263.207461 | MF: C8H10NO5PS

[Related Structures, Assays, Literature](#)

1: CID: [4793](#)



phosalone; Benzophosphate; Benzophos ...
IUPAC: 6-chloro-3-(diethoxyphosphinothioylsulfanylmethyl)-1,3-benzoxazol-2-one
MW: 367.808561 | MF: C12H15ClNO4PS2

[Related Structures, Assays, Literature](#)

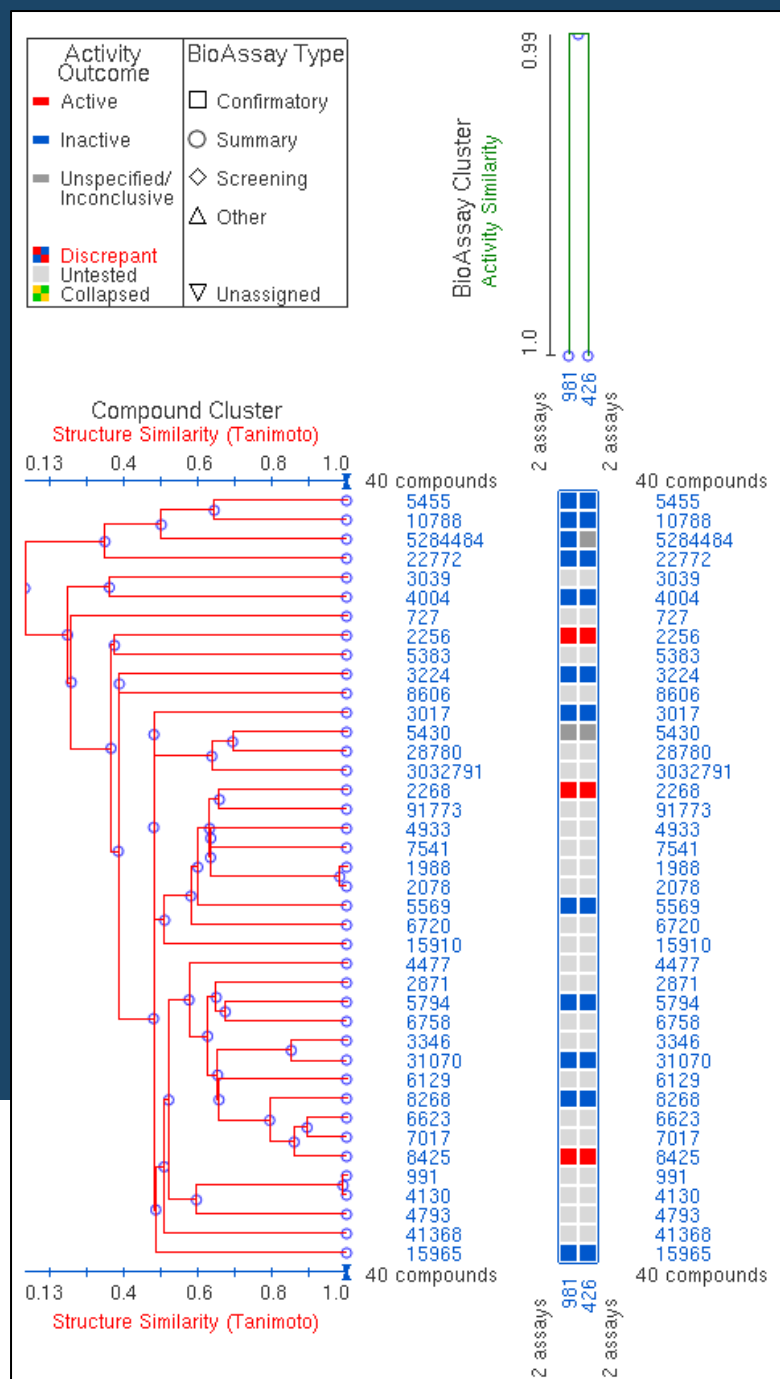
Similar Bioassay Profile across structure space

1: AID: [426](#)

Cell Viability - Jurkat
Source: NCGC

1: AID: [981](#)

Cell Viability - LYMP2-010
Source: NCGC



Incorporating SAR Concepts into ToxCast:

Conclusions

- HTS data offers:
 - ▶ activity-based clustering of chemicals
 - ▶ biofunctional information for refining class-based SAR
 - ▶ new “biological” descriptors for global SAR
- In vivo bioassay profiles expanding “endpoints” for SAR
- Structure-analog approaches coupled with HTS and ToxRef in vivo data offer powerful data mining tools
- Public tools for structure-based exploration of data becoming available

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