

# Cheminformatics approaches to support chemical identification delivered via the EPA CompTox Chemicals Dashboard

#### Antony Williams<sup>1</sup>, Andrew D. McEachran<sup>2</sup>, Chris Grulke<sup>1</sup>, Elin Ulrich<sup>3</sup> and Jon R. Sobus<sup>3</sup>

1) National Center for Computational Toxicology, U.S. Environmental Protection Agency, RTP, NC

2) Oak Ridge Institute of Science and Education (ORISE) Research Participant, RTP, NC

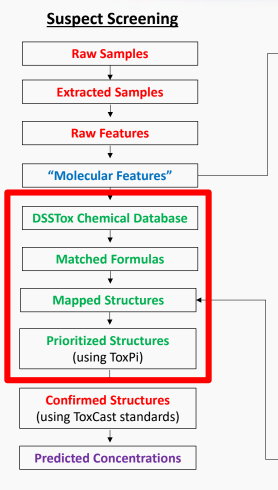
3) National Exposure Research Laboratory, U.S. Environmental Protection Agency, RTP, NC

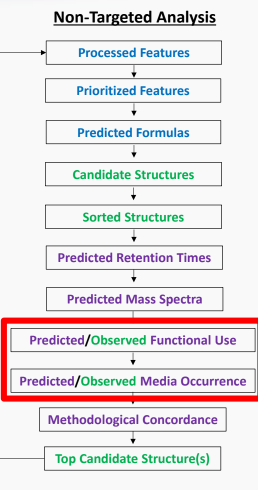
The views expressed in this presentation are those of the author and do not necessarily reflect the views or policies of the U.S. EPA

Spring 2019 ACS Spring Meeting, Orlando

#### Suspect Screening and Non-Targeted Analysis Workflows







#### <u>Color Key</u>

- Red = Analytical Chemistry
- **Blue** = Data Processing & Analysis

Purple = Mathematical & QSPR Modeling

Green = Informatics & Web Services



## **CompTox Chemicals Dashboard**

https://comptox.epa.gov/dashboard

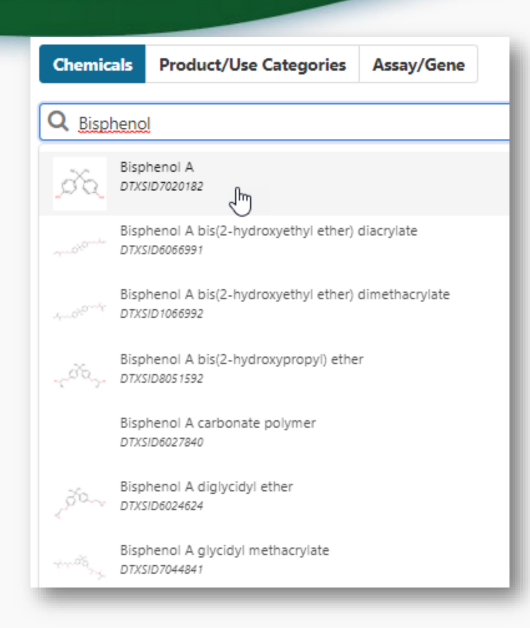


#### 875k Chemical Substances

Sepa United States Environmental Agency	Protection Home Advanced Search Batch Search Lists 🗸 Predictions Downloads	Share 🔻
UNITED STATES	875 Thousand Chemicals	
	Chemicals Product/Use Categories Assay/Gene	
Rommer	Q Search for chemical by systematic name, synonym, CAS number, DTXSID or InChIKey	
AL PROTECT	Identifier substring search See what people are saying, read the dashboard comments! Cite the Dashboard Publication click here	
	Latest News	
	Read more news	
	Journal of Cheminformatics article regarding "MS-Ready structures"	
	March 9th, 2019 at 1:09:45 PM	
	A recent article describes "MS-Ready structures", what they are, how they are generated and details regarding the benefits of these structures in navigating structure relationships across the dashboard. The article is published in the Journal of Cheminformatics here.	•
	• • • •	

#### **BASIC** Search





#### 80-05-7 | DTXSID7020182 Searched by DSSTox Substance Id. HAZARD Wikipedia -Bisphenol A (BPA) is an organic synthetic compound with the chemical formula (CH<sub>3</sub>)<sub>2</sub>C(C<sub>6</sub>H<sub>4</sub>OH)<sub>2</sub> belonging to the group of diphenylmethane derivatives and bisphenols, with two hydroxyphenyl groups. It is a colorless solid that is soluble in organic solvents, but poorly soluble in water (0.344 wt % at 83 °C). BPA is a starting material for the synthesis of plastics, primarily certain polycarbonates H<sub>3</sub>C CH3 Read more EXPOSURE Intrinsic Properties BIOACTIVITY Molecular Formula: C15H16O2 & Mol File Q. Find All Chemicals Average Mass: 228.291 g/mol Isotope Mass Distribution HO Monoisotopic Mass: 228.11503 g/mol SIMILAR COMPOUNDS ОH Structural Identifiers • GENRA (BETA) Linked Substances ۰. RELATED SUBSTANCES Presence in Lists 4 Record Information ۰. SYNONYMS **Quality Control Notes** ۰. LITERATURE COMMENTS

#### **Detailed Chemical Pages**

Advanced Search Batch Search Lists V Predictions Downloads

Bisphenol A

DETAILS

PROPERTIES

ADME

LINKS

EXECUTIVE SUMMARY

ENV. FATE/TRANSPORT



Copy 🕶 📕 Share 💌 📕 Submit Comment

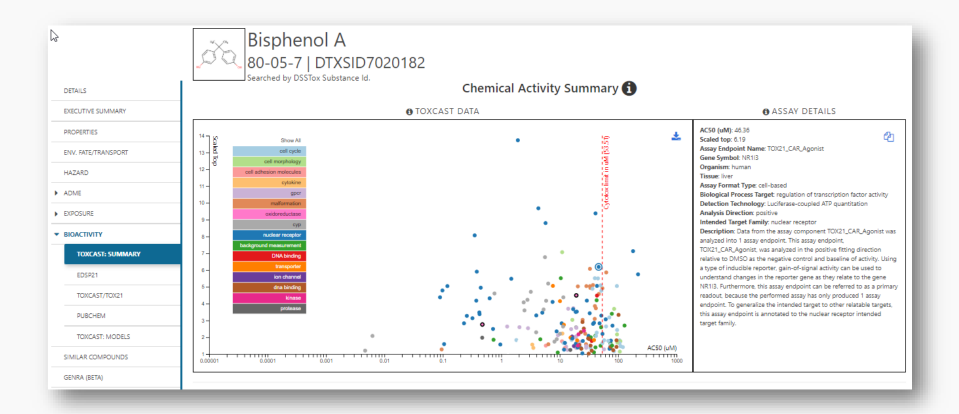
#### Access to Chemical Hazard Data



DETAILS							Ha	zard				
EXECUTIVE SUMMARY	DataType											
PROPERTIES	Toxic	tity Value	~									
ENV. FATE/TRANSPORT							🛉 Huma	an 💋 Eco				
HAZARD	a Down	nload 🔻	Columns	-								Search query
ADME		<b>-</b>	-	Subture 🗳			Units 🗘		-		Subsource \$	Source 🗘
EXPOSURE	More	Priority <sup>‡</sup>		Subtype	Risk assessment class 🕈	Value 🕈		Study type 🕈	Exposure route 🕈	Species *	Jubbuille	Source
BIOACTIVITY		7	MEG	Short-term Critical Air	short-term	500	mg/m3	-	inhalation	-	TG 230 Military Exposure Guidelines Table	DOD
SIMILAR COMPOUNDS		7	MEG	Short-term Marginal Air	short-term	100	mg/m3	-	inhalation	-	TG 230 Military Exposure Guidelines Table	DOD
		7	MEG	Short-term Negligible Air	short-term	15	mg/m3	-	inhalation	-	TG 230 Military Exposure Guidelines Table	DOD
GENRA (BETA)		7	MEG	Soil Negligible Soil	chronic	106000	mg/kg	-	soil	-	TG 230 Military Exposure Guidelines Table	DOD
RELATED SUBSTANCES		7	MEG	Long-Term, 5L/d Negligible Water	chronic	7	mg/L	-	oral	-	TG 230 Military Exposure Guidelines Table	DOD
SYNONYMS		6	RfD	-	chronic	0.05	mg/kg-day		oral	rat	Wignall	Wignall
LITERATURE		5	RfD		chronic	0.05	mg/kg-day			-	MSC Table 5	Pennsylvania DEP ToxValues
LINKS		-	100			0.05	ing/kgroby				inse labe s	Pennsylvania ber lokvalaes
COMMENTS		4	<u>RfD</u>	-	chronic	0.05	mg/kg-day	chronic	oral	rat	IRIS	Chiu
COMMENTS		3	<u>RfD</u>	-	chronic	0.6	mg/kg-day	-	oral	rat	EPA/ORNL/OLEM	HEAST
		1	<u>RfD</u>	-	chronic	0.05	mg/kg-day	-	oral	-	EPA NCEA	IRIS

#### In Vitro Bioassay Screening ToxCast and Tox21





#### Sources of Exposure to Chemicals



	Bisphenol A 80-05-7   DTXSID7020182 Searched by DSSTox Substance Id.								
DETAILS	Product and Use Categories (PUCs) 🚺								
EXECUTIVE SUMMARY	📩 Download 💌		_						
PROPERTIES	Columns V 10 V								
ENV. FATE/TRANSPORT	Product or Use Categorization	Categorization type	Number of Unique Products						
HAZARD	manufacturing, metals	CPCat Cassette	17						
ADME	adhesive	CPCat Cassette	17						
EXPOSURE		CPCat Cassette	16						
EXPOSURE		CPCat Cassette	12						
PRODUCT 9	USE CATEGORIES	CPCat Cassette	11						
FRODUCTA	SE CATEGORIES	CPCat Cassette	8						
		CPCat Cassette	8						
CHEMICAL V	VEIGHT FRACTION	CPCat Cassette	8						
		CPCat Cassette	7						
CHEMICAL F	UNCTIONAL USE	CPCat Cassette	6						
TOXICS REL	EASE INVENTORY	First << < 1 2 3 4 5	6 7 8 9 10 > >> Last						
MONITORING DATA									
EXDOSUDE	PREDICTIONS								

#### Link Access



#### Bisphenol A 80-05-7 | DTXSID7020182

	Searched by Approved	Name.			
DETAILS	General	Toxicology	Publications	Analytical	Prediction
EXECUTIVE SUMMARY	EPA Substance Registry Service	ACTOR	Toxline	FOR-IDENT	2D NMR HSQC/HMBC Prediction
PROPERTIES	Household Products Database	он, DrugPortal	Environmental Health Perspectives	NEMI: National Environmental Methods Index	Carbon-13 NMR Prediction
	Chemical Entities of Biological Interest	CCRIS	M NIEHS	RSC Analytical Abstracts	Proton NMR Prediction
ENV. FATE/TRANSPORT	(ChEBI)	ChemView	National Toxicology Program	🐴 Tox21 Analytical Data	ChemRTP Predictor
HAZARD	PubChem	<b>О</b> СТР	G Google Books	MONA: MassBank North America	LSERD
HAZARD	🛱 Chemspider	SechemPortal	G Google Scholar	and mzCloud	
► ADME	(CPCat	Gene-Tox	G Google Patents	NST NIST IR Spectrum	
	- 🧳 DrugBank	HSDB	PPRTVWEB	MST NIST MS Spectrum	
EXPOSURE	hmp HMDB	ToxCast Dashboard 2	PubMed		
BIOACTIVITY	W Wikipedia	LactMed	IRIS Assessments		
	Q MSDS Lookup	International Toxicity Estimates for Risk	🖲 EPA HERO		
SIMILAR COMPOUNDS	ChEMBL	ATSDR Toxic Substances Portal	🔍 NIOSH Skin Notation Profiles		
GENRA (BETA)	Q Chemical Vendors	🏽 Superfund Chemical Data matrix	🚾 NIOSH Pocket Guide		
	<ul> <li>CalEPA Office of Environmental Health Hazard Assessment</li> </ul>	🚾 NIOSH IDLH Values	C RSC Publications		
RELATED SUBSTANCES	NIOSH Chemical Safety Cards	ACToR PDF Report	🕌 BioCaddie DataMed		
SYNONYMS	ToxPlanet	Toxics Release Inventory	D Springer Materials		
	ACS Reagent Chemicals	CREST	Federal Register		
LITERATURE	W Wikidata	National Air Toxics Assessment	Regulations.gov		
LINKS	ChemHat: Hazards and Alternatives Toolbox		Bielefeld Academic Search Engine		
COMMENTS	🜞 Wolfram Alpha		CORE Literature Search		
COMMENTS	ScrubChem				
	ECHA Brief Profile				

#### NIST WebBook https://webbook.nist.gov/chemistry/



#### Analytical

☑ FOR-IDENT

B

NEMI: National Environmental Methods Index

C RSC Analytical Abstracts

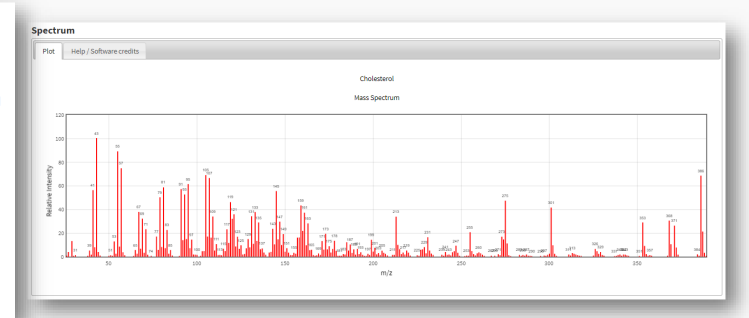
👌 Tox21 Analytical Data

MONA: MassBank North America

area mzCloud

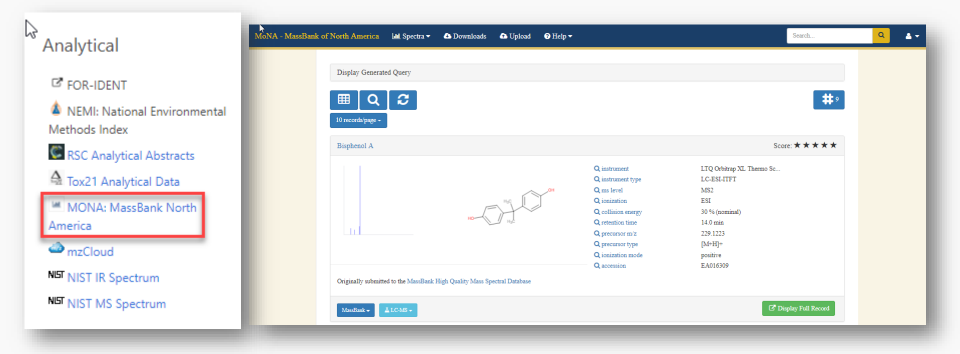
NIST IR Spectrum

NIST MS Spectrum



#### MassBank of North America https://mona.fiehnlab.ucdavis.edu





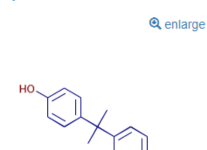
#### m/z CLOUD https://www.mzcloud.org/



# Analytical FOR-IDENT FOR-IDENT Kethods Index RSC Analytical Abstracts KSC Analytical Abstracts MONA: MassBank North America MONA: MassBank North America NET NIST IR Spectrum NET NIST MS Spectrum



#### **Bisphenol A**



Advanced Mass Spectral Database

Home

#### Systematic / IUPAC Name: 4,4'-(2,2-Propanediyl)diphenol

ID: Reference1477

Other Names: 2,2-Bis(4-hydroxyphenyl)propane; 2,2-Bis(*p*-hydroxyphenyl)propane; Phenol, (1-methylethylidene)bis-; Isopropylidene-bis(4-hydroxybenzene); 4-[1-(4-Hydroxyphenyl)-1-methylethyl]phenol; more

Formula: C<sub>15</sub>H<sub>16</sub>O<sub>2</sub>

Class: Industrial Chemicals



# DO WE REALLY NEED ANOTHER DATABASE?

# Is a bigger database better?



Journal of The American Society for Mass Spectrometry

January 2012, Volume 23, <u>Issue 1</u>, pp 179–185 | <u>Cite as</u>

#### Identification of "Known Unknowns" Utilizing Accurate Mass Data and ChemSpider

Authors

Authors and affiliations

James L. Little 🖂 , Antony J. Williams 🖂 , Alexey Pshenichnov, Valery Tkachenko

- ChemSpider was 26 million chemicals then
- Much BIGGER today
- Is bigger better??



# **Comparing Search Performance**



CrossMark

Ånal Bioanal Chem (2017) 409:1729–1735 DOI 10.1007/s00216-016-0139-z

RAPID COMMUNICATION

#### Identifying known unknowns using the US EPA's CompTox Chemistry Dashboard

Andrew D. McEachran<sup>1</sup> · Jon R. Sobus<sup>2</sup> · Antony J. Williams<sup>3</sup>

- Dashboard content was 720k chemicals
- Only 3% of ChemSpider size
- What was the comparison in performance?

## SAME dataset for comparison



Compound class	Number in class	Average rank		Number of compounds in each position rank-ordered				
			#1	#2	#3	#4	#5+	
Pharmaceutical drug	72	1.4	55	9	6	2		
Industrial chemicals	42	5.5	28	6	3		5	
Personal care products	8	6.1	3	1			4	
Steroid hormones	7	1.0	7					
Perfluorochemicals	6	1.2	5	1				
Pesticides	12	2.3	6	2	3		1	
Veterinary drugs	3	1.3	2	1				
Dyes	2	1.0	2					
Food product/natural compounds	4	3.8	2			1	1	
Illicit drugs	2	2.0	1		1			
Misc. molecules	3 <sup>a</sup>	1.3	2	1				



Summary statistics and rank-ordered position in the CompTox Chemistry Dashboard and ChemSpider of the 89 compound subset from the Little et al. [7] study

		Average rank	Num	ıber in ea	ach posi	tion ran	k-ordered
		(±SD)	#1	#2	#3	#4	#5+
Mass-based	Dashboard	$1.2 \pm 0.7$	77 <sup>a</sup>	5	3	3	
	ChemSpider	$2.2\pm6.1^b$	68	8	7	1	5
Formula-based	Dashboard	$1.1 \pm 0.4$	$78^{a}$	8	2		
	ChemSpider	$1.3 \pm 1.0$	77	8	2	1	2

<sup>a</sup>One chemical (tephrosin) not present in the Dashboard

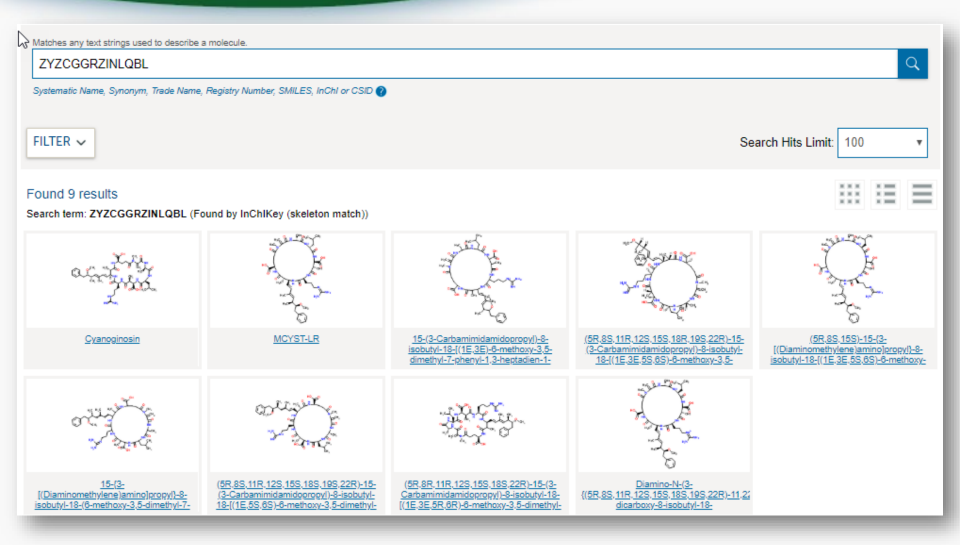
## Data Quality is important



#### Data quality in free web-based databases!



#### Will the correct Microcystin LR Stand Up? ChemSpider Skeleton Search

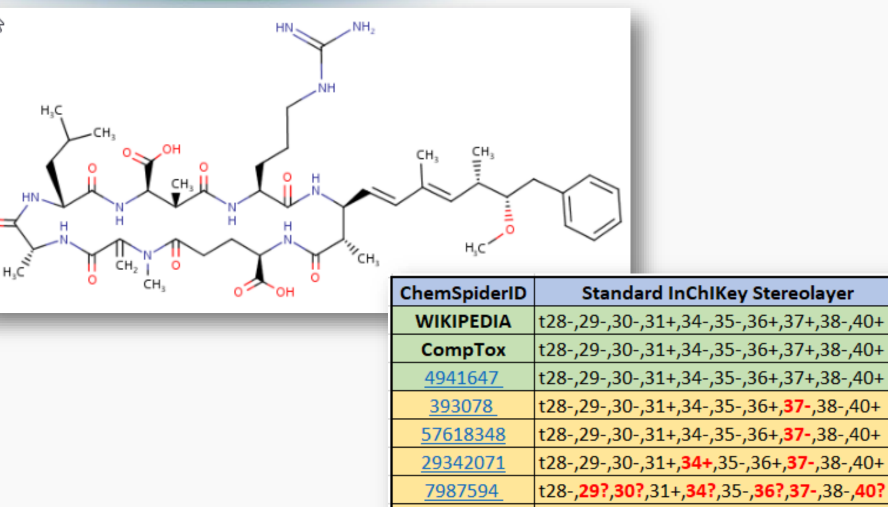


United States Environmental Protection

Agency

#### **Comparing ChemSpider Structures**

2



t28-,29-,30-,31+,34-,35-,36+, <b>37-</b> ,38-,40+
t28-,29-,30-,31+,34-,35-,36+, <b>37-</b> ,38-,40+
t28-,29-,30-,31+, <b>34+</b> ,35-,36+, <b>37-</b> ,38-,40+
t28-, <b>29?,30?</b> ,31+, <b>34?</b> ,35-, <b>36?,37-</b> ,38-, <b>40?</b>
t28-, <b>29?,30+,31-,34+,35+,36-,37-</b> ,38-, <b>40-</b>
NONE
NONE

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#### **Comparing ChemSpider Structures**



ChemSpiderID	InChlKey	# Stereocenters	# Different
WIKIPEDIA	ZYZCGGRZINLQBL-JCGNTXOTSA-N	10/10	0
CompTox	ZYZCGGRZINLQBL-JCGNTXOTSA-N	10/10	0
<u>4941647</u>	ZYZCGGRZINLQBL-JCGNTXOTSA-N	10/10	0
<u>393078</u>	ZYZCGGRZINLQBL-GWRQVWKTSA-N	10/10	1
<u>57618348</u>	ZYZCGGRZINLQBL-UPPCHHEJSA-N	10/10	1
<u>29342071</u>	ZYZCGGRZINLQBL-IIJTUTQBSA-N	10/10	2
<u>7987594</u>	ZYZCGGRZINLQBL-BESLYTPASA-N	5/10	6
<u>22900854</u>	ZYZCGGRZINLQBL-QAXSDTKVSA-N	9/10	8
<u>19692240</u>	ZYZCGGRZINLQBL-ORZJCNCZSA-N	0/10	10
<u>2831283</u>	ZYZCGGRZINLQBL-UHFFFAOYSA-N	0/10	10

## **Other Searches**



# UniChem

# Pub Chem About

#### ZYZCGGRZINLQBL

Treating this query as a text search.

Compounds (17)

Show	All  entries			
	CMR. Query InChl	src_id	Source	src_compound_id
	matches	1	ChEMBL	CHEMBL444092
	matches	4	Guide to Pharmacolog	y <u>4735</u>
	matches	6	KEGG Ligand	<u>C05371</u>
	matches	7	ChEBI	<u>6925</u>
	matches	9	ZINC	ZINC000169715525
	matches	9	ZINC	ZINC000255288110
	matches	9	ZINC	ZINC000255288111
	matches	9	ZINC	ZINC000255288112
	matches	9	ZINC	ZINC000255288113
	matches	9	ZINC	ZINC000255288114
	matches	9	ZINC	ZINC000255288115
	matches	9	ZINC	ZINC000583653042
	matches	9	ZINC	ZINC000669680403
	matches	10	eMolecules	<u>26754757</u>
	matches	10	eMolecules	<u>31239828</u>
	matches	11	IBM Patent System	DA3C2F25F29692734272194ED0E2C009
	matches	14	FDA SRS	EQ8332842Y

## **Delivering a Better Database**



- An ideal database would provide:
  - Curated CAS Number-Name mappings with "correct" chemical structures
- We have full time curators checking data

ecor	rd Information
ß	Citation: U.S. Environmental Protection Agency. Chemistry Dashboard.
	//comptox.epa.gov/dashboard/DTXSID3031654 (accessed Mar 17th, 2019), Microcystin LR
Data	Quality:
Leve	11: Expert curated, highest confidence in accuracy and consistency of unique chemical identifiers
Level	2: Expert curated, unique chemical identifiers using multiple sources
Level PubC	l 3: Programmatically curated from high quality EPA source, unique chemical identifiers have no conflicts in ChemID and Chem
Level	4: Programmatically curated from ChemID, unique chemical identifiers have no conflicts in PubChem
Level	5: Programmatically curated from ACToR or PubChem, unique chemical identifiers with low confidence, single public source



# **MASS AND** FORMULA SEARCHING (and metadata ranking)

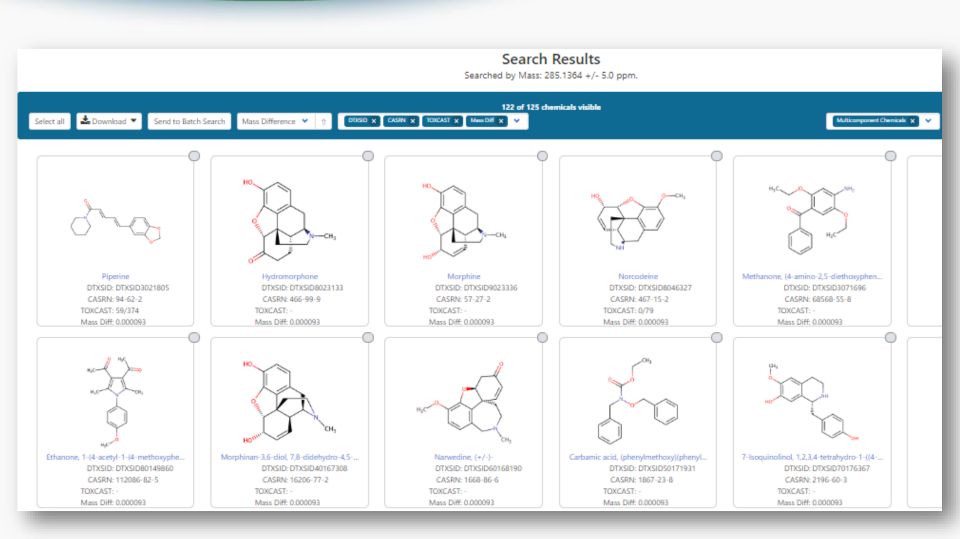
# Advanced Searches <u>Mass</u> and Formula Based Search



Mass Search 🕄
± Min/Max
Adduct All Adducts       Neutral     Choose adduct from dropdown
285.1364 Da ± 5 Da ppm
Search <b>Q</b>
Molecular Formula Search 🚯
💿 MS Ready Formula 🚯 🔘 Exact Formula 🚯
Formula
Please use the format of the following example: C6H8O2 or C6H(8-10)O(0-2)
Search <b>Q</b>

# Advanced Searches <u>Mass</u> and Formula Based Search





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# Using Metadata for Ranking



- Use available metadata to rank candidates
  - Associated data sources
    - Associated lists in DSSTox database
    - Associated sources in PubChem
    - Specific types (e.g. water, surfactants, pesticides etc.)
  - Number of associated PubMed articles
  - Number of products/categories containing the chemical

#### Metadata rank ordering





# SPECIFIC APPLICATIONS TO MASS SPEC.

#### Mass Spec Focused Applications

Journal of Exposure Science & Environmental Epidemiology (2018) 28:411–426 https://doi.org/10.1038/s41370-017-0012-y

#### **REVIEW ARTICLE**



# Integrating tools for non-targeted analysis research and chemical safety evaluations at the US EPA

Jon R. Sobus<sup>1</sup> · John F. Wambaugh<sup>2</sup> · Kristin K. Isaacs<sup>1</sup> · Antony J. Williams<sup>2</sup> · Andrew D. McEachran<sup>3</sup> · Ann M. Richard<sup>2</sup> · Christopher M. Grulke<sup>2</sup> · Elin M. Ulrich<sup>1</sup> · Julia E. Rager<sup>3,4</sup> · Mark J. Strynar<sup>1</sup> · Seth R. Newton<sup>1</sup>



Cite This: Environ. Sci. Technol. 2018, 52, 3125–3135

Article

pubs.acs.org/est

mental Protection

#### Suspect Screening Analysis of Chemicals in Consumer Products

Katherine A. Phillips,<sup>†</sup> Alice Yau,<sup>‡</sup> Kristin A. Favela,<sup>‡</sup> Kristin K. Isaacs,<sup>†</sup> Andrew McEachran,<sup>§,||</sup> Christopher Grulke,<sup>||</sup> Ann M. Richard,<sup>||</sup> Antony J. Williams,<sup>||</sup> Jon R. Sobus,<sup>†</sup> Russell S. Thomas,<sup>||</sup> and John F. Wambaugh<sup>\*,||</sup>

#### "MS-Ready Structures" https://doi.org/10.1186/s13321-018-0299-2



McEachran et al. J Cheminform (2018) 10:45 https://doi.org/10.1186/s13321-018-0299-2 Journal of Cheminformatics

#### METHODOLOGY



Open Access

# "MS-Ready" structures for non-targeted high-resolution mass spectrometry screening studies

Andrew D. McEachran<sup>1,2\*</sup>, Kamel Mansouri<sup>1,2,3</sup>, Chris Grulke<sup>2</sup>, Emma L. Schymanski<sup>4</sup>, Christoph Ruttkies<sup>5</sup> and Antony J. Williams<sup>2\*</sup>

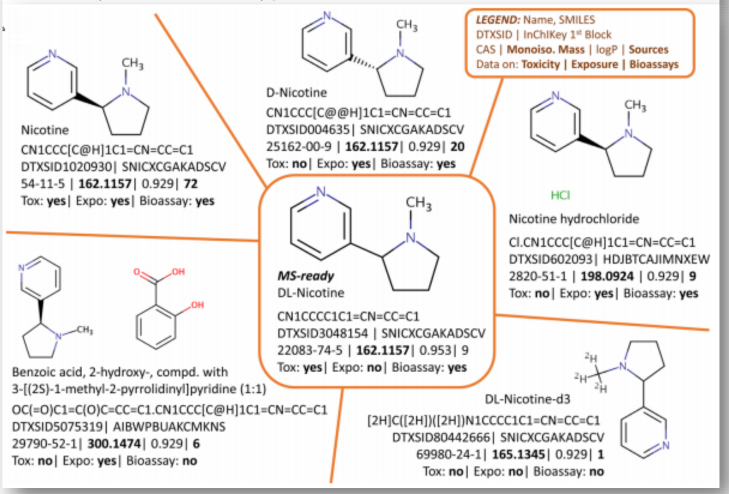




pubs.acs.org/est

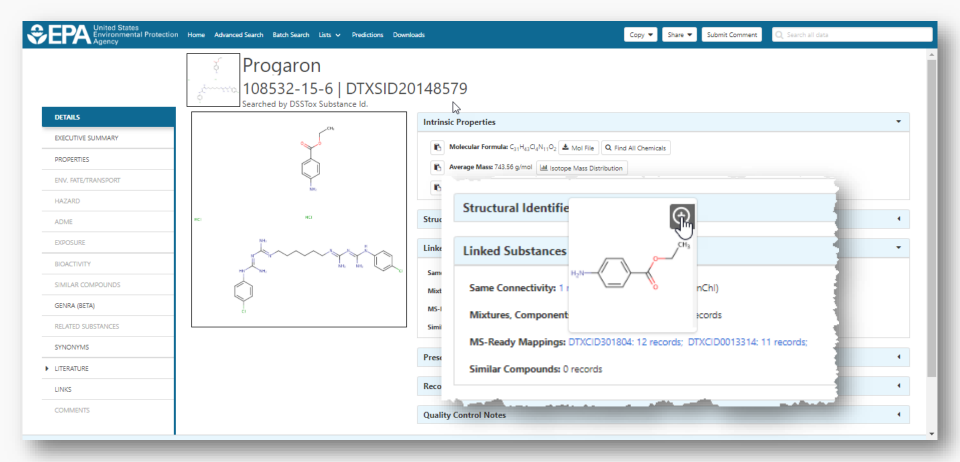
#### Open Science for Identifying "Known Unknown" Chemicals

Emma L. Schymanski\*<sup>,†</sup><sup>©</sup> and Antony J. Williams<sup>\*,‡</sup><sup>©</sup>



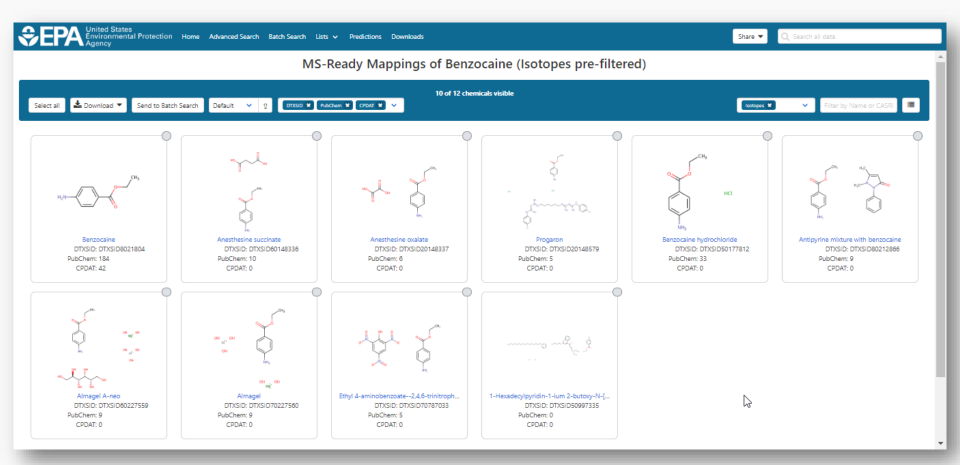
## **MS-Ready Mappings**





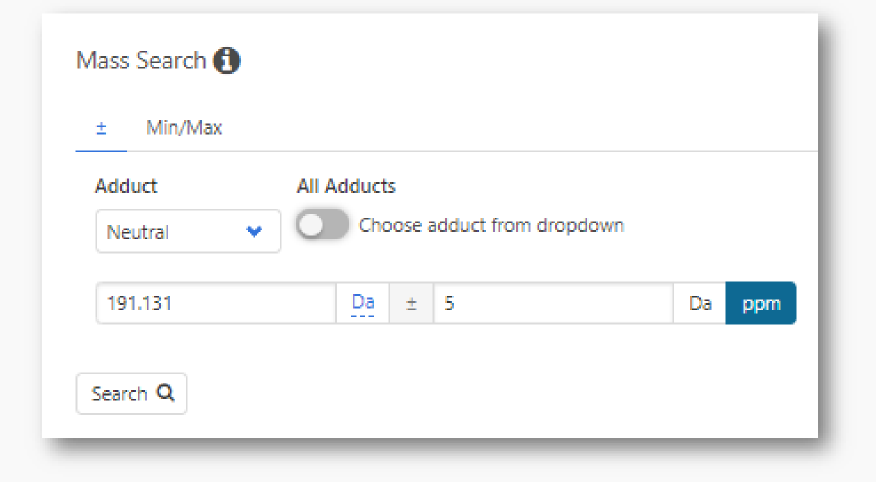
## MS-Ready Mappings Set





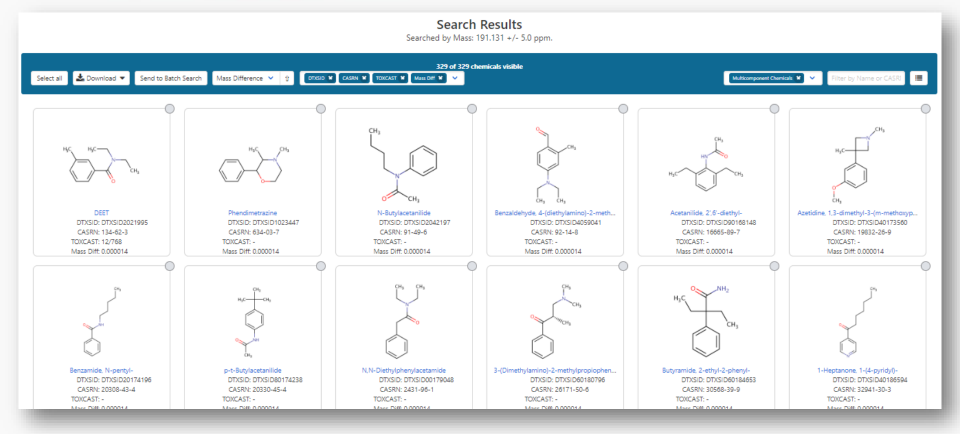
#### Advanced Searches Mass Search





#### Advanced Searches Mass Search





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# MS-Ready Structures for **Formula Search**



### Molecular Formula Search 🚺

💿 MS Ready Formula 🚯 🔿 Exact Formula 🚯

### Formula

Please use the format of the following example: C6H8O2 or C6H(8-10)O(0-2)

Search 🔍

## **MS-Ready Mappings**



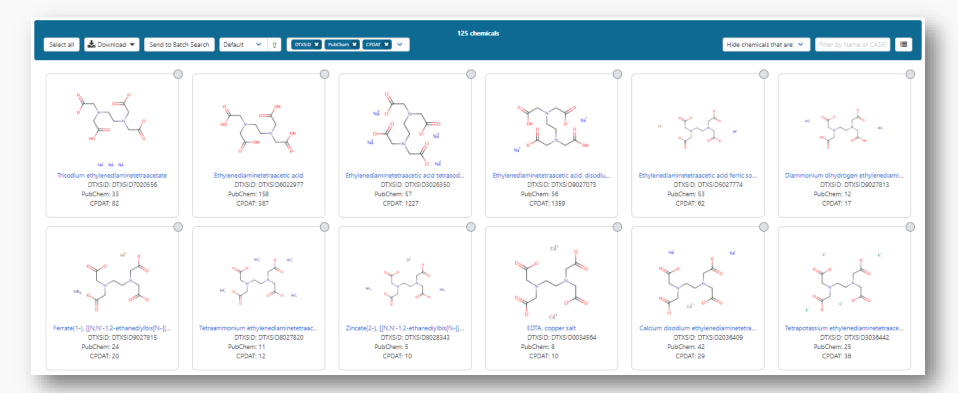
### • EXACT Formula: C10H16N2O8: 3 Hits

Ň	O MS Formula C10H1		ıla 🚯
S	Select all 🛃 Download 🔻 Send to B	Batch Search Default 👻 🕆 DTXSID 🗙	3 of 3 chemi
	$\begin{aligned} & \stackrel{\bullet}{ } \stackrel{\bullet}{          $	$ \begin{array}{c} & \stackrel{\circ}{\leftarrow} \stackrel{\circ}$	$ \begin{array}{c} & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ & $

## **MS-Ready Mappings**



- Same Input Formula: C10H16N2O8
- MS Ready Formula Search: 125 Chemicals







- 125 chemicals returned in total
  - 8 of the 125 are single component chemicals
  - 3 of the 8 are isotope-labeled
  - 3 are neutral compounds and 2 are charged

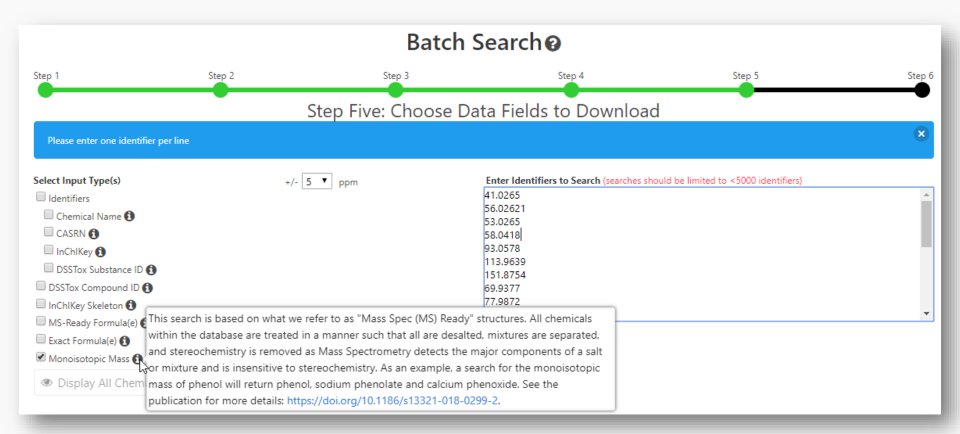
### **Batch Searching**



• Singleton searches are useful but we work with **thousands** of masses and formulae!

- Typical questions
  - What is the list of chemicals for the formula  $C_xH_yO_z$
  - What is the list of chemicals for a mass +/- error
  - Can I get chemical lists in Excel files? In SDF files?
  - Can I include properties in the download file?

### **Batch Searching Formula/Mass**



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### Searching batches using MS-Ready Formula (or mass) searching



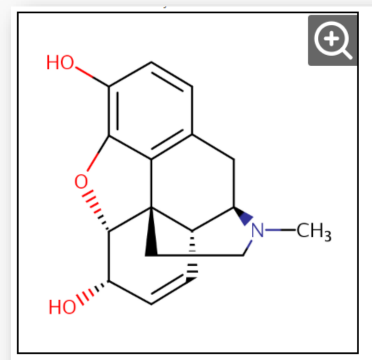
4	А	В	С	D	E	F	G
1	INPUT	DTXSID	CASRN	PREFERRED NAME	MOL FORMULA	MONOISOTOPIC MASS	DATA SOURCES
2	C14H22N2O3	DTXSID2022628	29122-68-7	Atenolol	C14H22N2O3	266.163042576	46
3	C14H22N2O3	DTXSID0021179	6673-35-4	Practolol	C14H22N2O3	266.163042576	32
4	C14H22N2O3	DTXSID4048854	841-73-6	Bucolome	C14H22N2O3	266.163042576	20
5	C14H22N2O3	DTXSID1045407	13171-25-0	Trimetazidine dihydrochloride	C14H24Cl2N2O3	338.116398	19
6	C14H22N2O3	DTXSID0045753	56715-13-0	R-(+)-Atenolol	C14H22N2O3	266.163042576	19
7	C14H22N2O3	DTXSID2048531	5011-34-7	Trimetazidine	C14H22N2O3	266.163042576	14
8	C14H22N2O3	DTXSID10239405	93379-54-5	Esatenolol	C14H22N2O3	266.163042576	12
9	C14H22N2O3	DTXSID50200634	52662-27-8	N-(2-Diethylaminoethyl)-2-(4-hydroxyphenoxy)acetamide	C14H22N2O3	266.163042576	7
10	C14H22N2O3	DTXSID4020111	51706-40-2	dl-Atenolol hydrochloride	C14H23CIN2O3	302.1397203	6
11	C14H22N2O3	DTXSID1068693	51963-82-7	Benzenamine, 2,5-diethoxy-4-(4-morpholinyl)-	C14H22N2O3	266.163042576	5
12	C18H34N2O6S	DTXSID3023215	154-21-2	Lincomycin	C18H34N2O6S	406.213757997	35
13	C18H34N2O6S	DTXSID7047803	859-18-7	Lincomycin hydrochloride	C18H35CIN2O6S	442.1904357	22
		DTXSID20849438			C18H35CIN2O6S	442.1904357	1
	C10H12N2O		486-56-6	Cotinine	C10H12N2O	176.094963014	40
16	C10H12N2O	DTXSID8075330	50-67-9	Serotonin	C10H12N2O	176.094963014	22
17	C10H12N2O	DTXSID8044412	2654-57-1	4-Methyl-1-phenylpyrazolidin-3-one	C10H12N2O	176.094963014	18
18	C10H12N2O	DTXSID80165186	153-98-0	Serotonin hydrochloride	C10H13CIN2O	212.0716407	11
19	C10H12N2O	DTXSID2048870	29493-77-4	(4R,5S)-4-methyl-5-phenyl-4,5-dihydro-1,3-oxazol-2-amine	C10H12N2O	176.094963014	10
20	C10H12N2O	DTXSID10196105	443-31-2	6-Hydroxytryptamine	C10H12N2O	176.094963014	9
21	C10H12N2O	DTXSID90185693	31822-84-1	1,4,5,6-Tetrahydro-5-phenoxypyrimidine	C10H12N2O	176.094963014	7
22	C10H12N2O	DTXSID40178777	2403-66-9	2-Benzimidazolepropanol	C10H12N2O	176.094963014	7
23	C10H12N2O	DTXSID80157026	13140-86-8	N-Cyclopropyl-N'-phenylurea	C10H12N2O	176.094963014	6
24	C10H12N2O	DTXSID30205607	570-14-9	4-Hydroxytryptamine	C10H12N2O	176.094963014	6
25	C14H18N4O3		17804-35-2	Benomyl	C14H18N4O3	290.137890456	68
26	C14H18N4O3		738-70-5		C14H18N4O3		51
27	C14H18N4O3	DTXSID40209671			C14H19CIN4O3	326.1145682	8
28	C14H18N4O3	DTXSID70204210	55687-49-5	Benzenemethanol, 4-((2,4-diamino-5-pyrimidinyl)methyl)-2,	C14H18N4O3	290.137890456	5
29	C14H18N4O3	DTXSID20152671		6-Methoxy-4-(3-(N,N-dimethylamino)propylamino)-5,8-quina		290.137890456	4
30	C14H18N4O3	DTXSID30213742		1H-1,2,4-Benzotriazepine-3-carboxylic acid, 4,5-dihydro-4-		290.137890456	3
	C14H18N4O3	DTXSID30219608	69449-07-6	2,4-Pyrimidinediamine, 5-((3,4,5-trimethoxyphenyl)methyl)·		308.14845514	3
	C14H18N4O3	DTXSID20241155		L-Aspartic acid, compound with 5-((3,4,5-trimethoxyphenyl		423.175398165	3
33	C14H18N4O3	DTXSID80241156		L-Glutamic acid, compound with 5-((3,4,5-trimethoxypheny		437.191048229	3
	C14H18N4O3	DTXSID20143781	101204-93-7	1H-Pyrido(2,3-e)-1,4-diazepine-2,3,5-trione, 4-(2-(diethylam	C14H18N4O3	290.137890456	3
35	C12H11N7	DTXSID6021373	396-01-0		C12H11N7	253.107593382	52
36	C12H11N7	DTXSID00204465			C12H11N7	253.107593382	7
37	C12H11N7		7300-26-7		C12H9N7	251.091943318	4
38	C12H11N7	DTXSID00848025			C12H13N7O4S	351.074973101	1
39	C12H11N7	DTXSID50575293	92310-83-3	(1E)-N-Phenyl-1,2-bis(1H-1,2,4-triazol-1-yl)ethan-1-imine	C12H11N7	253.107593382	1
40	C8H9NO2	DTXSID2020006	103-90-2	Acetaminophen	C8H9NO2		75
11	CSHONO2	DTVSID6026667	13/ 20 3	Mothyl 2 aminohonzoato	C8H0NO2	161.063328534	50



# RELATED **APPLICATIONS OF INTEREST TO** MASS SPEC.

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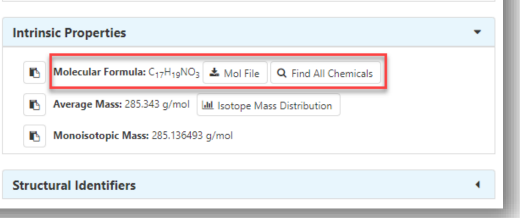




#### Wikipedia

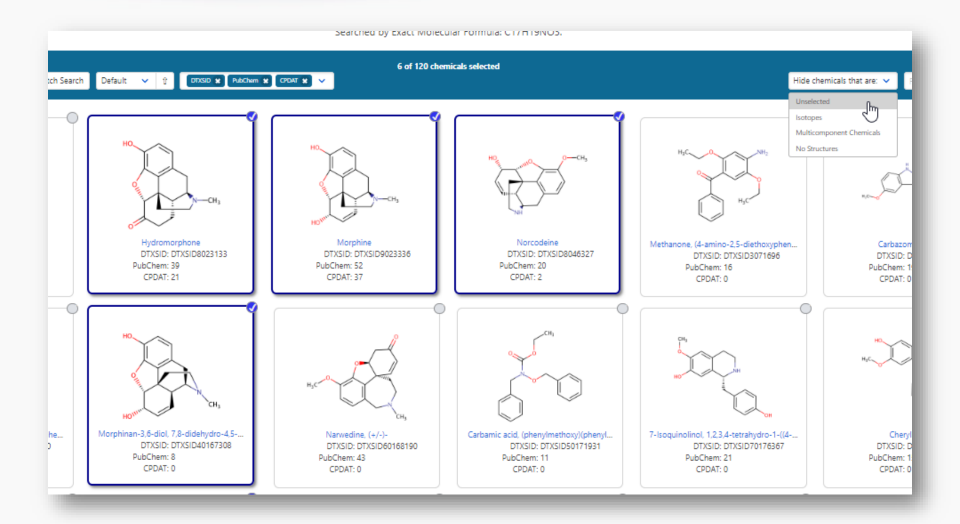
**Morphine** is a pain medication of the opiate family which is found naturally in a number of plants and animals. It acts directly on the central nervous system (CNS) to decrease the feeling of pain. It can be taken for both acute pain and chronic pain. It is frequently used for pain from myocardial infarction and during labor. It can be given by mouth, by injection into a muscle, by injection under the skin, intravenously, injection into the space around the

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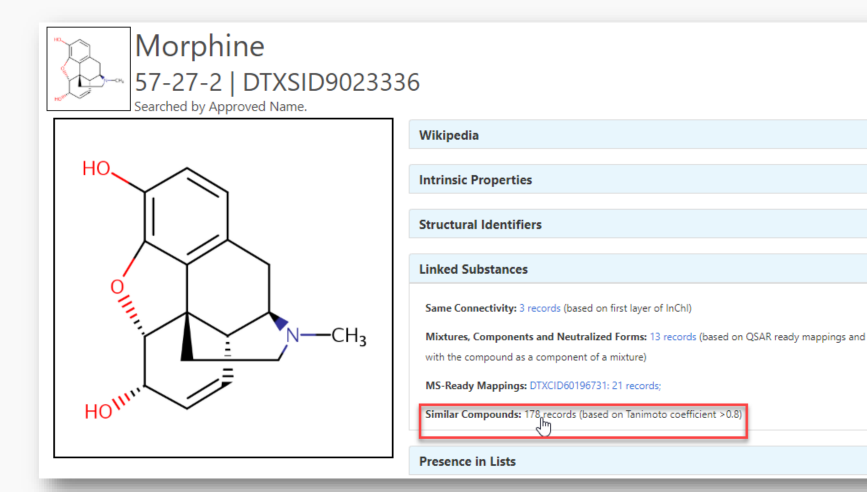


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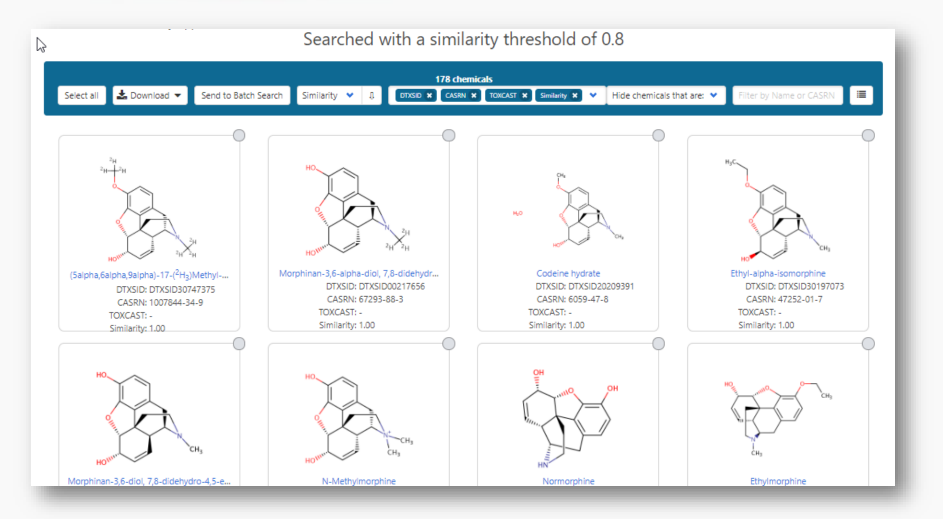
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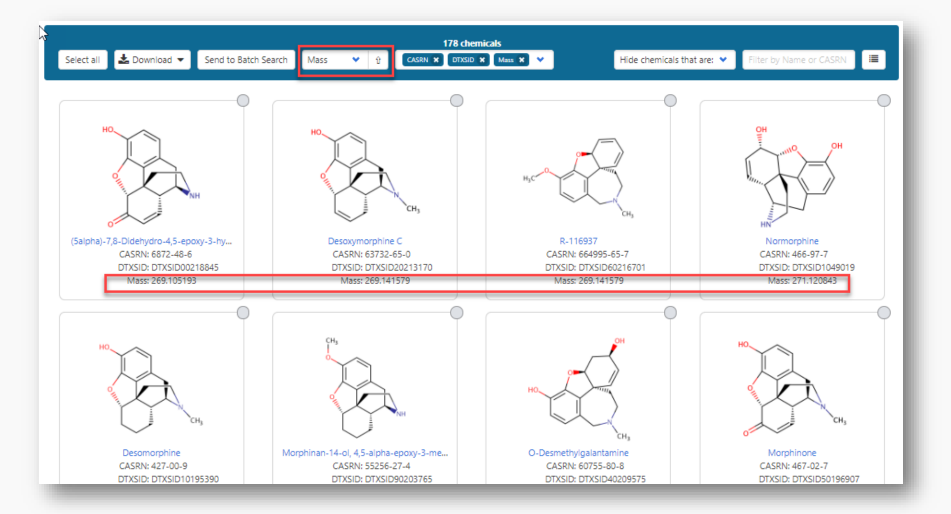
### Find me "related structures" Based on Structure Similarity





### Find me "related structures" Structure Similarity – sort on mass





### Literature Searching





### **Abstract Sifter**

#### 1) Select PubMed starting point query then 2) click on Retrieve.

Select a Query Term	Retrieve Articles
Select a Query Term	
Hazard	
Fate and Transport	
Metabolism/PK/PD	
Chemical Properties	
Exposure	
Mixtures	
Male Reproduction	
Androgen Disruption	
Female Reproduction	
GeneTox	
Cancer	
Clinical Trials	
Embryo and embryonic development	
Child (infant through adolescent)	
Dust and Exposure	
Food and Exposure	
Water and Exposure	
Algae	
Disaster / Emergency	

#### Optionally, edit the query before retrieving.

"57-27-2" OR "Morphine"

### Literature Searching



Child (Intant through adolescent)		
Dust and Exposure		
Food and Exposure		
Water and Exposure	N	
Algae	W	
Disaster / Emergency		

voptionally, edit the query before retrieving.

("57-27-2" OR "Morphine") AND ((water OR groundwater OR drinking water) AND Environmental Exposure)

### Literature Searching



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wastewater	Spectrom	netry ↓	EPA	Total	PMID	Year	Title		Authors	Journal	Rev	*
4	2		0	6	29274731	2017	Simultaneous analysis of opioid analgesics and thei		Krizman-Matasic; Kostanjevecki; Ahel; Terzic	Journal of chromatography. A		
0	1		0	1	25768972	2015	Evaluating external contamination of polybrominate		Poon; Aleksa; Carnevale; Kapur; Goodyer; Koren	Therapeutic drug monitoring		
0	1		0	1	22544551	2012	Spatial distribution of illicit drugs in surface waters o		Vazquez-Roig; Andreu; Blasco; Morillas; Picó	Environmental science and pollution research inter		
1	1		0	2	20801487	2010	Analysis	of llicit and illicit drugs in waste, surface an	Berset; Brenneisen; Mathieu	Chemosphere		
1	1		0	2	17935751	2007	Illicit dru	gs, a novel group of environmental contami	Zuccato; Castiglioni; Bagnati; Chiabrando; Grassi;	Water research		
2	1		1	4	17607391	2007	Using environmental analytical data to estimate lev		Bones; Thomas; Paull	Journal of environmental monitoring : JEM		
3	1		2	6	17180984	2006	Simultaneous determination of psychoactive drugs		Hummel; Löffler; Fink; Ternes	Environmental science & technology		
6	0		0	6	30583189	2018	Assessm	nent of drugs of abuse in a wastewater trea	Kumar; Tscharke; O'Brien; Mueller; Wilkins; Padhye	The Science of the total environment		
0	0		3	3	30488421	2018	Effect of	enriched environment during adolescence	Mohammadian; Najafi; Miladi-Gorji	Developmental psychobiology		
3	0		0	3	29574368	2018	Estimatio	on of the consumption of illicit drugs during $\dots$	Foppe; Hammond-Weinberger; Subedi	The Science of the total environment		
1	0		0	1	28787791	2017	Evaluatio	on of in-sewer transformation of selected illi	Gao; Banks; Li; Jiang; Lai; Mueller; Thai	The Science of the total environment		
9	0		0	9	28472697	2017	Occurren	nce and fate of illicit drugs and pharmaceuti	Causanilles; Ruepert; Ibáñez; Emke; Hernández; d	The Science of the total environment		
0	0		0	0	28010888	2016	2016 Dose-dependent effects of morphine on lipopolysac Mottaz; Schönenberger; Fischer; Eggen; Sc		Mottaz; Schönenberger; Fischer; Eggen; Schirmer;	Environmental pollution (Barking, Essex : 1987)		
0	0		0	0	27746311	2016	Effects of voluntary exercise on the viability, prolifer		Haydari; Safari; Zarbakhsh; Bandegi; Miladi-Gorji	Neuroscience letters		
0	0		0	0	27261879	2016	Genotox	ic effects induced by the exposure to an en	Parolini; Magni; Castiglioni; Binelli	Ecotoxicology and environmental safety		
3	0		0	3	27179320	2016	Tempora	al trends in drug use in Adelaide, South Aus	Tscharke; Chen; Gerber; White	The Science of the total environment		-
	wastewater           4           0           1           2           3           6           0           3           1           9           0           0           0	wastewater         Spectron           4         2           0         1           0         1           1         1           1         1           2         1           3         1           6         0           0         0           3         0           1         0           9         0           0         0           0         0           0         0           0         0	stewater         Spectrom           wastewater         Spectrometry           4         2           0         1           0         1           1         1           1         1           2         1           3         1           6         0           0         0           3         0           1         0           9         0           0         0           0         0           0         0           0         0           0         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PMID       Year       Title       Authors       Journal       Journal       PMID       Year       Year       Title       Authors       Journal       Journal       PMID       Year       Year



# FOCUSED CHEMICAL LISTS OF INTEREST

### **Chemical Lists**



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List Acronym 🗢	List Name 🗘	Last Updated 🗘	Number of Chemicals 🕈	List Description \$
HDXEXCH	MASSPECDB: Hydrogen Deuterium Exchange Standard Set - Under HDX Conditions	2018-11-07	592	Observed species (deuterated and undeuterated) from the HDXNOEX list under hydrogen deuterium exchange conditions (Ruttkies, Schymanski et al. in prep.)
HDXNOEX	MASSPECDB: Hydrogen Deuterium Exchange Standard Set - No Exchange	2018-11-07	765	Environmental standard set used to investigate hydrogen deuterium exchange in small molecule high resolution mass spectrometry (Ruttkies, Schymanski et al. in prep.)
MASSBANKEUSP	MASSPECDB: MassBank.EU Collection: Special Cases	2017-07-16	263	The MassBank.EU list contains curated chemicals (Schymanski/Williams) associated with the literature/tentative/unknown/SI spectra available on MassBank.EU that are not available as part of the full MassBank collection of reference standard spectra.
MASSBANKREF	MASSPECDB: MassBank Reference Spectra Collection	2017-07-13	1267	This MassBank list contains chemicals associated with the full MassBank collection of reference standard spectra available on MassBank.EU, MassBank.JP and MassBank of North America as well as the Open Data collection, curated by Williams/Schymanski.
MYCOTOXINS	MASSPECDB: Mycotoxins from MassBank.EU	2017-08-02	88	This is a set of mycotoxins, initiated by the contribution of spectra of 90 mycotoxins to MassBank.EU by Justin Renaud and colleagues from Agriculture and Agri-Food Canada, Government of Canada

### **EPAHFR: Hydraulic Fracturing**



54

2

### WATER|EPA; Chemicals associated with hydraulic fracturing

🔍 Search EPAHFR Chemicals

Identifier substring search

#### **List Details**

**Description:** Chemicals used in hydraulic fracturing fluids and/or identified in produced water from 2005-2013, corresponding to chemicals listed in Appendix H of EPA's Hydraulic Fracking Drinking Water Assessment Final Report (Dec 2016). Citation: U.S. EPA, Hydraulic Fracturing for Oil and Gas: Impacts from the Hydraulic Fracturing Water Cycle on Drinking Water Resources in the United States (Final Report). U.S. Environmental Protection Agency, Washington, D.C. EPA/600/R-16/236F, 2016. <u>https://www.epa.gov/hfstudy</u>

\*Note that Appendix H chemical listings in Tables H-2 and H-4 were mapped to current DSSTox content, which has undergone additional curation since the publication of the original EPA HF Report (Dec 2016). In the few cases where a Chemical Name and CASRN from the original report map to distinct substances (as of Jan 2018), both were included in the current EPAHFR chemical listing for completeness; additionally, 34 previously unmapped chemicals in Table H-5 are now registered in DSSTox (all but 2 assigned CASRN) and, thus, have been added to the current EPAHFR listing. **Number of Chemicals:** 1640

2 Cſ NĤa NH. OH Alkylbenzenesulfonate linear Ammonium chloride Ammonium hydroxide Diammonium citrate DTXSID: DTXSID3020041 DTXSID: DTXSID0020078 DTXSID: DTXSID5020079 DTXSID: DTXSID4020080 PubChem: 82 PubChem: 0 PubChem: 19 PubChem: 83 CPDAT: 83 CPDAT: 260 CPDAT: 18 CPDAT: 857

### **PFAS** lists of Chemicals



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PFAS

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List Acronym \$	List Name 🗘	Last Updated 🗘	Number of Chemicals 🕈	List Description \$
EPAPFAS75S1	PFAS EPA: List of 75 Test Samples (Set 1)	2018-06-29	74	PFAS list corresponds to 75 samples (Set 1) submitted for initial testing screens conducted by EPA researchers in collaboration with researchers at the National Toxicology Program.
EPAPFAS75S2	PFAS EPA: List of 75 Test Samples (Set 2)	2019-02-21	75	PFAS list corresponds to a second set of 75 samples (Set 2) submitted for testing screens conducted by EPA researchers in collaboration with researchers at the National Toxicology Program.
EPAPFASCAT	PFAS EPA Structure- based Categories	2018-06-29	64	List of registered DSSTox "category substances" representing PFAS categories created using ChemAxon's Markush structure-based query representations.
EPAPFASINSOL	PFAS EPA: Chemical Inventory Insoluble in DMSO	2018-06-29	43	PFAS chemicals included in EPA's expanded ToxCast chemical inventory found to be insoluble in DMSO above 5mM.
EPAPFASINV	PFAS EPA: ToxCast Chemical Inventory	2018-06-29	430	PFAS chemicals included in EPA's expanded ToxCast chemical inventory and available for testing.
EPAPFASRL	PFAS EPA: Cross-Agency Research List	2017-11-16	199	EPAPFASRL is a manually curated listing of mainly straight-chain and branched PFAS (Per- & Poly-fluorinated alkyl substances) compiled from various internal, literature and public sources by EPA researchers and program office representatives.
PFASKEMI	PFAS: List from the Swedish Chemicals Agency (KEMI) Report	2017-02-09	2416	Perfluorinated substances from a Swedish Chemicals Agency (KEMI) Report on the occurrence and use of highly fluorinated substances.
PFASMASTER	PFAS Master List of PFAS Substances	2018-07-26	5061	PFASMASTER is a consolidated list of PFAS substances spanning and bounded by the below lists of current interest to researchers and regulators worldwide.
PFASOECD	PFAS: Listed in OECD Global Database	2018-05-16	4729	OECD released a New Comprehensive Global Database of Per- and Polyfluoroalkyl Substances, (PFASs) listing more than 4700 new PFAS
PFASTRIER	PFAS Community- Compiled List (Trier et al., 2015)	2017-07-16	597	PFASTRIER community-compiled public listing of PFAS (Trier et al, 2015)



# COMPLEX CHEMICAL SUBSTANCES

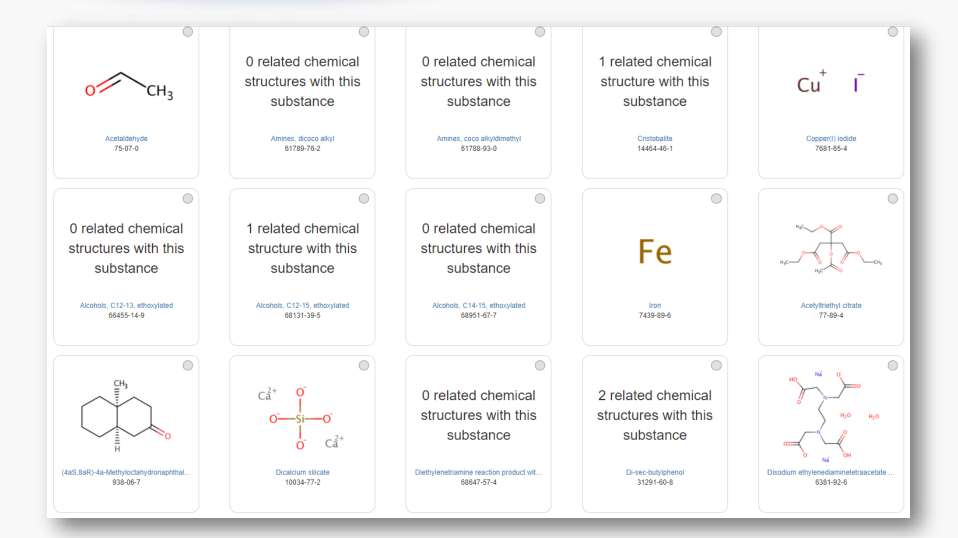


## Chemical Substances of Unknown or Variable Composition, Complex Reaction Products and Biological Materials (UVCB Substance) on the TSCA Inventory

This paper is a compendium of information related to the broad class of chemical substances referred to as UVCBs for the Toxic Substances Control Act (TSCA) Chemical Substance Inventory. These chemical substances cannot be represented by unique structures and molecular formulas.

### Many Hydraulic Fracturing Chemicals are "Complex"

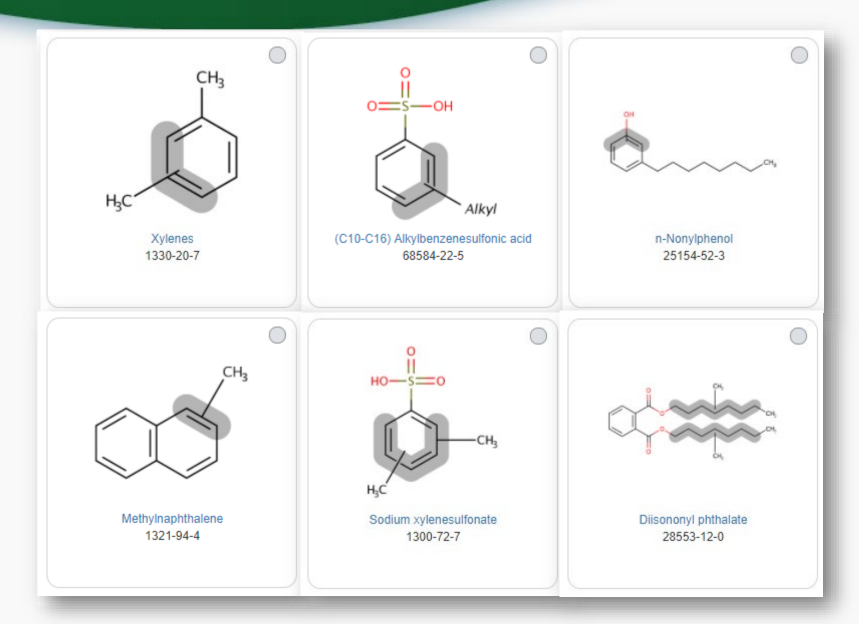




### "Markush Structures"

https://en.wikipedia.org/wiki/Markush\_structure



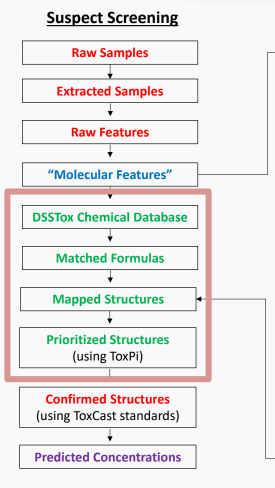


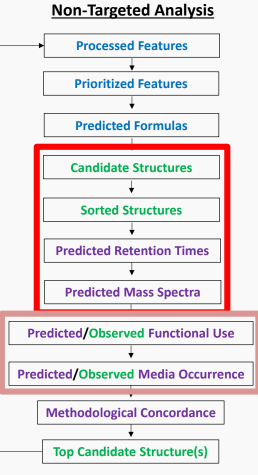


# WORK IN PROGRESS

### Suspect Screening and Non-Targeted Analysis Workflow







#### <u>Color Key</u>

- Red = Analytical Chemistry
- **Blue** = Data Processing & Analysis

Purple = Mathematical & QSPR Modeling

Green = Informatics & Web Services



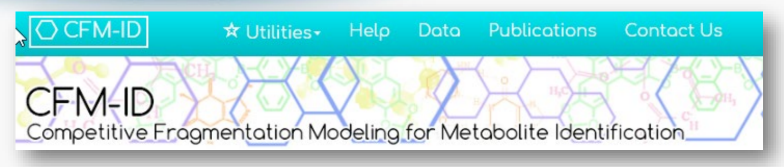


- Predicted Spectra for candidate ranking
  - Viewing and Downloading pre-predicted spectra
  - Search spectra against the database

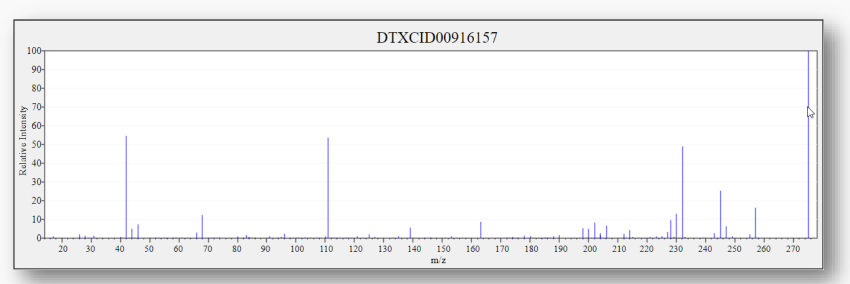
### **Predicted Mass Spectra**

http://cfmid.wishartlab.com/





- MS/MS spectra prediction for ESI+, ESI-, and EI
- Predictions generated and stored for >800,000 structures, to be accessible via Dashboard



### Search Expt. vs. Predicted Spectra



SEPA United States Environmental Protection Agency	Home Advanced Search Batch Search Lists 🛩 Predictions Downloads	Share - Q. Search all data
	Non Target Analysis Prototype	Â
	Mass Search <u>± Min/Max</u> 321.138493476 Da <u>± 0.0000002 Da ppm</u>	
	Molecular Formula Search	
	Mass or Formula must be entered before searching spectrum Ionization Type          ESI+           ESI+           ESI-           EI           Spectra Input	
	Single Energy         Multiple           304.1332052 11.6199475         •           198.0913404 7.306439699         •           123.0440559 6.538348292         •           196.0756904 6.269463115         •           216.1019051 4.700461978         •	
	Peak Match Window: 0.02 Da ppm	

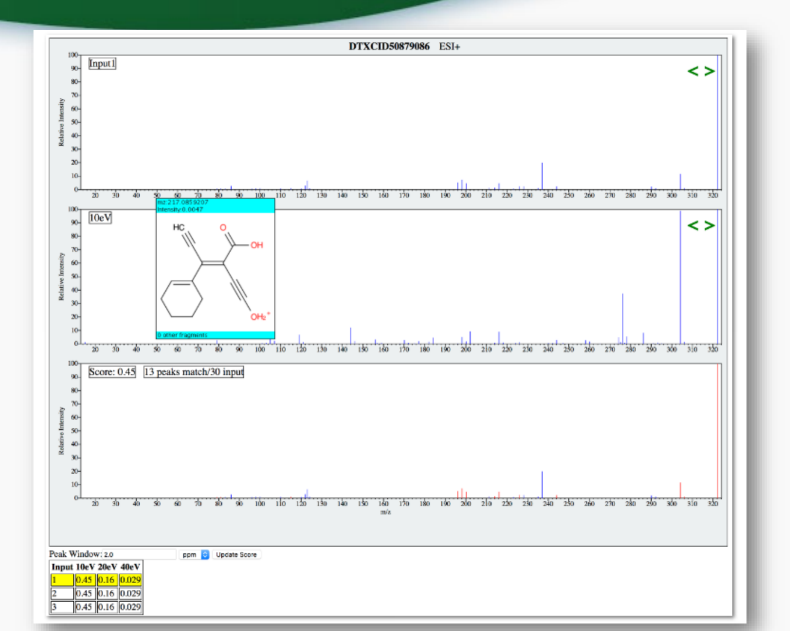
### Search Expt. vs. Predicted Spectra



United States Environmental Protection Home Adv Agency	ranced Search Batch Search Lists 🛩 Predictions Downloads	Share 🔻 🔍 Search all data
Spectra Inpu Single Ener	Chemical Structure ID	Score (10eV)
304.1332052 11.61 198.0913404 7.30 123.0440559 8.53 196.0756904 5.26	DTXCID101048191	0.22
218.1019051 4.70	DTXCID101181567	0.19
Peak Match Search	DTXCID50879086	0.17
TSV CSV Excel	DTXCID60686349	0.14
Chemical Structure ID	DTXCID00830900	0.13 m of Scores
DTXCID101181567	DTXCID10971176	0.12
DTXCID60688349 DTXCID00830900	DTXCID60301242	0.12
DTXCID10971178 DTXCID60301242	DTXCID40703048	0.11
DTXCID40703048 DTXCID60349982	DTXCID60349982	0.11
DTXCID10316649 .	DTXCID10316649	0.09

### **Spectral Viewer Comparison**





## Work in Progress



- Predicted Spectra for candidate ranking
  - Viewing and Downloading pre-predicted spectra
  - Search spectra against the database
- Retention Time Index Prediction

### **Retention Time Prediction for Ranking**





Talanta

Volume 182, 15 May 2018, Pages 371-379



## A comparison of three liquid chromatography (LC) retention time prediction models

Andrew D. McEachran <sup>a, b</sup> 은 岵, Kamel Mansouri <sup>a, b, 1</sup>岵, Seth R. Newton °岵, Brandiese E.J. Beverly <sup>a, c, 2</sup>岵, Jon R. Sobus °岵, Antony J. Williams <sup>b</sup> 은 岵

Show more

https://doi.org/10.1016/j.talanta.2018.01.022

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### Moving to Relative Retention Times





### Journal of Hazardous Materials

Volume 363, 5 February 2019, Pages 277-285



Development and application of retention time prediction models in the suspect and non-target screening of emerging contaminants

Reza Aalizadeh, Maria-Christina Nika, Nikolaos S. Thomaidis Ӓ 🖾

Show more

https://doi.org/10.1016/j.jhazmat.2018.09.047

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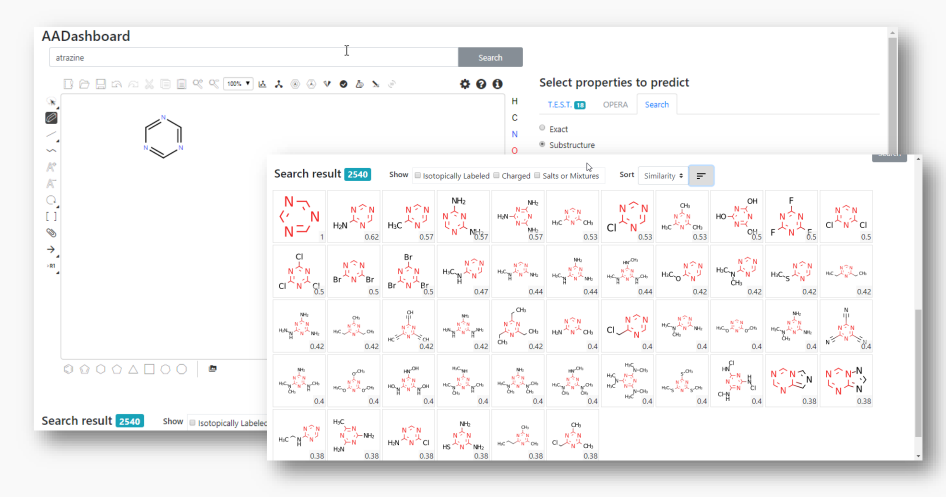
## Work in Progress



- Predicted Spectra for candidate ranking
  - Viewing and Downloading pre-predicted spectra
  - Search spectra against the database
- Retention Time Index Prediction
- Structure/substructure/similarity search

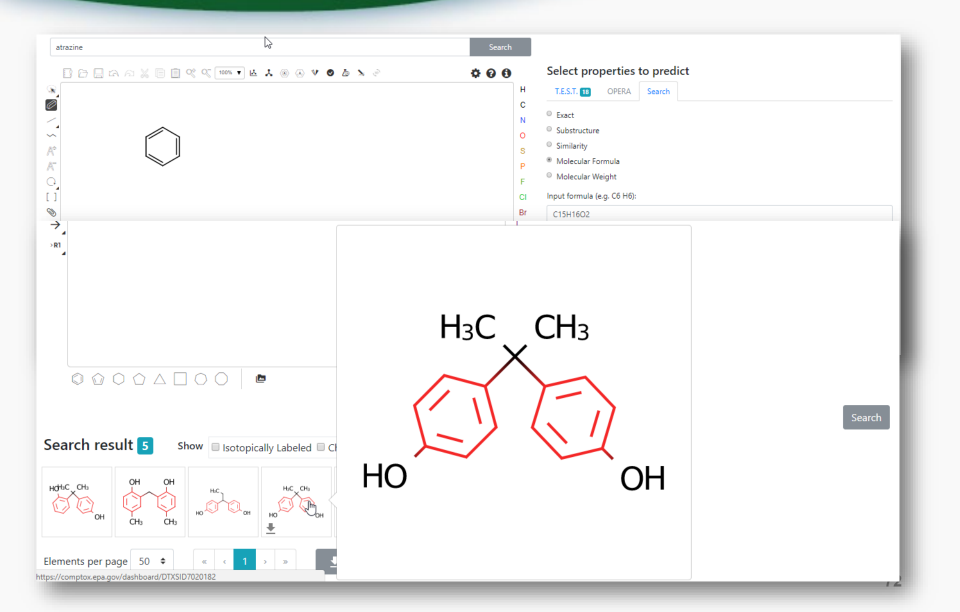
### Prototype Development





### Prototype Development





## Work in Progress



- Predicted Spectra for candidate ranking
  - Viewing and Downloading pre-predicted spectra
  - Search spectra against the database
- Retention Time Index Prediction
- Structure/substructure/similarity search
- Access to API and web services for programmatic access

### API services and Open Data



- Groups waiting on our API and web services
- Mass Spec companies instrument integration
- Release will be in iterations but for now our data are available

1 Ca 2 26 3 10 4 60	<b>asrn</b> 6148-68-5 07-29-9	dsstox_substance_id DTXSID7020001	:ludes the CAS Number, DSSTox substance identifier (D preferred_name A-alpha-C			
2 26 3 10 4 60	6148-68-5 07-29-9	DTXSID7020001				
4 60						
		DTXSID2020004	Acetaldehyde oxime			
5 10	0-35-5	DTXSID7020005	Acetamide			
	03-90-2	DTXSID2020006	Acetaminophen			
6 96	68-81-0	DTXSID7020007	Acetohexamide			
7 18	8523-69-8	DTXSID2020008	Acetone[4-(5-nitro-2-furyl)-2-thiazolyl] hydrazone			
8 75	5-05-8	DTXSID7020009	Acetonitrile			
9 12	27-06-0	DTXSID6020010	Acetoxime			
10 65	5734-38-5	DTXSID6020012	N'-Acetyl-4-(hydroxymethyl) phenylhydrazine			
The Cor associat formula	ited with any c ae searches sh	stry Dashboard can be used by hemical, whether it include so ould be based on desalted, ar	y mass spectrometrists for the purpose of structure iden lvents of hydration, salts or multiple components. How Id desolvated structures with stereochemistry removed. ferred Name, CAS-RN. DTXSID, Formula, Formula of the	ever, mass spectrometry de . We refer to these as "MS r	tects ionized chemical structures and molecu eady structures" and the MS-ready mapping:	ılar s are



# SIDE EFFECTS OF SHARING OPEN DATA

### **NORMAN Suspect List Exchange**

https://www.norman-network.com/?q=node/236



### NORMAN

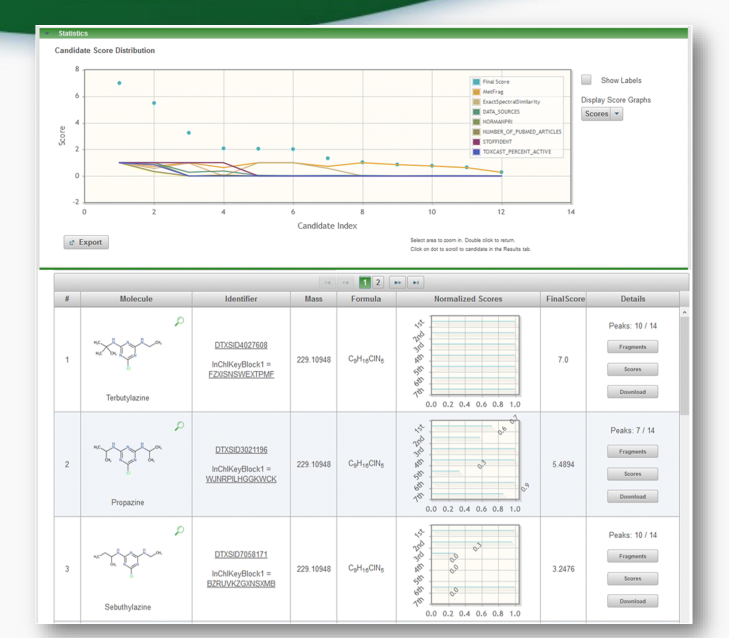
Network of reference laboratories, research centres and related organisations for monitoring of emerging environmental substances

Wastewater Suspect List based on Swedish Product Data	Wastewater Suspect List Original File with Mapped DTXSIDs (12/02/2019)	KEMIWWSUS InChlKeys (12/02/2019)	A prioritized list of 1,123 substances relevant for wastewater based on Swedish product registry data, including scores. Provided by Stellan Fischer, KEMI.
Algal toxins list from CompTox	ALGALTOX XLSX, CSV (14/02/2019) CompTox ALGALTOX List	ALGALTOX InChlKeys (14/02/2019)	List of algal toxins (generated during blooms) from the CompTox Chemicals Dashboard.
CCL 4 Chemical Candidate List	CCL4 XLSX, CSV (14/02/2019) CompTox CCL4 List	CCL4 InChIKeys (14/02/2019)	Contaminants that are not (yet) regulated in the USA but are known or anticipated to occur in public water systems; from CompTox.
Hydrogen Deuterium Exchange (HDX) Standard Set	HDXNOEX XLSX, CSV (14/02/2019) CompTox HDXNOEX List CompTox HDXEXCH List	HDXNOEX InChlKeys (14/02/2019)	Environmental standard set used to investigate hydrogen deuterium exchange in small molecule HRMS (Ruttkies et al. submitted). HDXEXCH list also contains observed deuterated species.
Neurotoxicants Collection from Public Resources	NEUROTOXINS XLSX, CSV (14/02/2019) CompTox NEUROTOXINS List	NEUROTOXINS InChlKeys (14/02/2019)	A list of neurotoxicants compiled from public resources, details on CompTox and Schymanski <i>et al.</i> (submitted).
Statins Collection from Public Resources	STATINS XLSX, CSV (14/02/2019) CompTox STATINS List	STATINS InChlKeys (14/02/2019)	A list of statins (lipid-lowering medications) compiled from public resources, details on CompTox.
Synthetic Cannabinoids and Psychoactive Compounds	SYNTHCANNAB XLSX, CSV (14/02/2019) CompTox SYNTHCANNAB List	SYNTHCANNAB InChlKeys (14/02/2019)	A list of synthetic cannabinoids and psychoactive compounds assembled from public resources, from CompTox.

### Integration to MetFrag in place

https://jcheminf.biomedcentral.com/articles/10.1186/s13321-018-0299-2

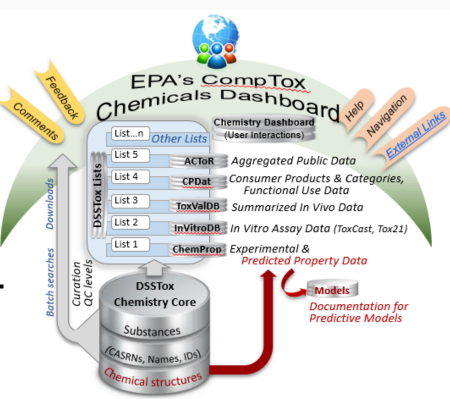




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## Conclusion

- Dashboard access to data for ~875,000 chemicals
- MS-Ready data facilitates structure identification
- Related metadata facilitates candidate ranking
- Relationship mappings and chemical lists of great utility
- Dashboard and contents are one part of the solution
- We are committed to open API development with time..



nvironmental Protection

Agency



- THANK YOU for the invitation!
- IT Development team especially Jeff
   Edwards and Jeremy Dunne
- Chris Grulke for the ChemReg system
- NERL colleagues Jon Sobus, Elin Ulrich, Mark Strynar, Seth Newton
- Emma Schymanski, LCSB, Luxembourg
- The NORMAN Network and all contributors





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