



Fundamentals of QSAR modeling: basic concepts and applications

Alexander Tropsha

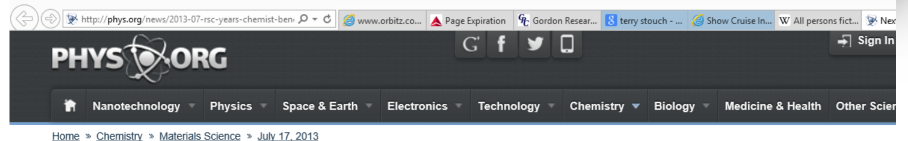
University of North Carolina, Chapel Hill, USA





- Basic concepts and best practices of QSAR modeling
- Data curation
- Case study and model interpretation: alerts about alerts
- Emerging approaches: Hybrid (chemicalbiological) QSAR modeling and Chemical Biological Read Across (CBRA)
- Summary of QSAR as (regulatory) decision support tool

The growing appreciation of molecular modeling and informatics



Next RSC president predicts that in 15 years no chemist will do bench experiments without computer-modelling them first

Jul 17, 2013

The newly-appointed President-Elect of the Royal Society of Chemistry today forecast the impact of advances in <u>modelling and computational informatics</u> on chemistry



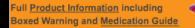
needed to guide people through the toughest times of their lives.

Will you answer the call?



president in 2014, said: "The speed and development of computers is now so rapid, and the advances in modelling and informatics are so dramatic that in 15 years' time, no chemist will be doing any experiments at the bench without trying to model them first."

Professor Tildesley is a world-leading expert in large-scale computational modelling and



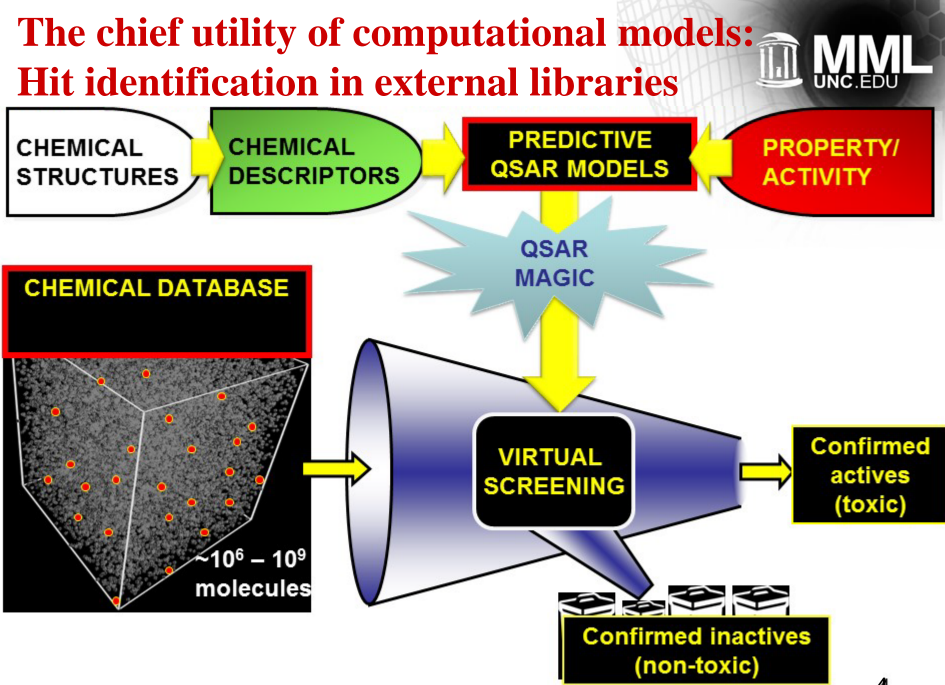
Download a Chronic Migra discussion guide and talk your doctor about BOTOX

Please scroll for Indication, Important Limitatio Important Safety Information including Boxed V

drooping eyelids, hoarseness or chang loss of voice (dysphonia), trouble sayin clearly (dysarthria), loss of bladder com trouble breathing, trouble swallowing. If happens, do not drive a car, operate

Popular

3



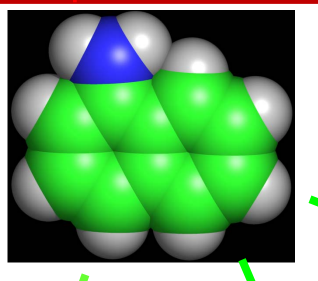
QSAR Modeling

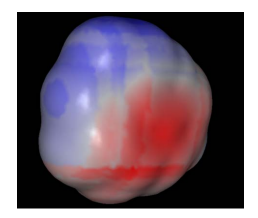


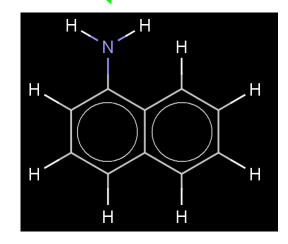
| $ f \\ f $ | 0.613 0.380 -0.22 1.450 0.708 1.146 0.491 - 0.30 0.141 0.956 0.256 0.799 | A C T I V I T Y |
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Slide credit: UNC MML

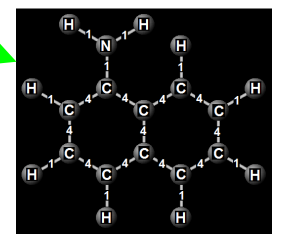
Structure representation



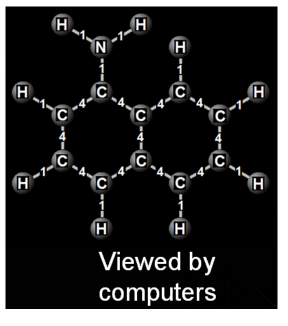




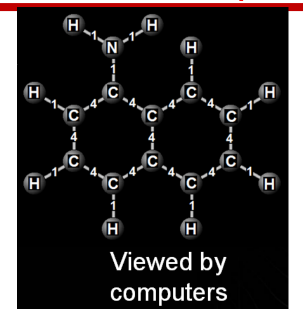








Structure representation



Graphs are widely used to represent and differentiate chemical structures, where atoms are vertices and bonds are expressed as edges connecting these vertices.

MOL File

Molecular graphs allow the computation of numerous indices to compare them quantitatively.

Molecular descriptors

| 1 | | | | -105 | | 020 | | | | | |
|---|-----------|------|---|------|-----|-----|-------|-----|------|--------|----------|
| 5 | 15 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0999 | V20 | 00 |
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| | 0.5 | 167 | | -2. | 550 | 0 | 0. | 000 | 00 C | 0 | 0 |
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| | | 667 | | -2. | 541 | 7 | | 000 | | 0 | 0 |
| | |)792 | | | 954 | | | 000 | | 0 | 0 |
| | | '917 | | -2. | | | | 000 | | 0 | 0 |
| | | 042 | | -2. | | | | 000 | | 0 | 0 |
| | | 250 | | -3. | | | | 000 | | 0 | 0 |
| | | 3083 | | -4. | 500 | 0 | | 000 | | 0 | 0 |
| | | 3917 | | | 083 | | | 000 | | 0 | 0 |
| | | 9750 | | | 795 | | | 000 | | 0 | 0 |
| | | 583 | | | 379 | | | 000 | | 0 | 0 |
| | | 9708 | | | 962 | | 0. | 000 | 00 C | 0 | 0 |
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| 4 | 11 | 1 | 0 | 0 | 0 | 0 | | _ ` | dg | | , |
| 5 | 6 | 1 | 0 | 0 | 0 | 0 | | | | | |
| 1 | 12 | 1 | 0 | 0 | 0 | 0 | - () | 20 | onr | ٦e | ct |
| 1 | 2 | 1 | 0 | 0 | 0 | 0 | - V | | | | |
| 2 | 13 | 1 | 0 | 0 | 0 | 0 | | | | 4. | |
| 2 | .7 | 1 | 0 | 0 | 0 | 0 | - Ié | 10 | oel- | -T\ | 0 |
| 3 | 14 | 1 | 0 | 0 | 0 | 0 | | | | | |
| 2 | 4 | 1 | 0 | 0 | 0 | 0 | | | | | |
| 4 | 15 | 1 | 0 | 0 | 0 | 0 | | | | | |
| 7 | 8 | 1 | 0 | 0 | 0 | 0 | | | | | |
| | 16 END | 1 | 0 | 0 | 0 | 0 | | | | | |
| | | | | | | | | | | | |

Vertices

(atomic type, coordinates etc.)

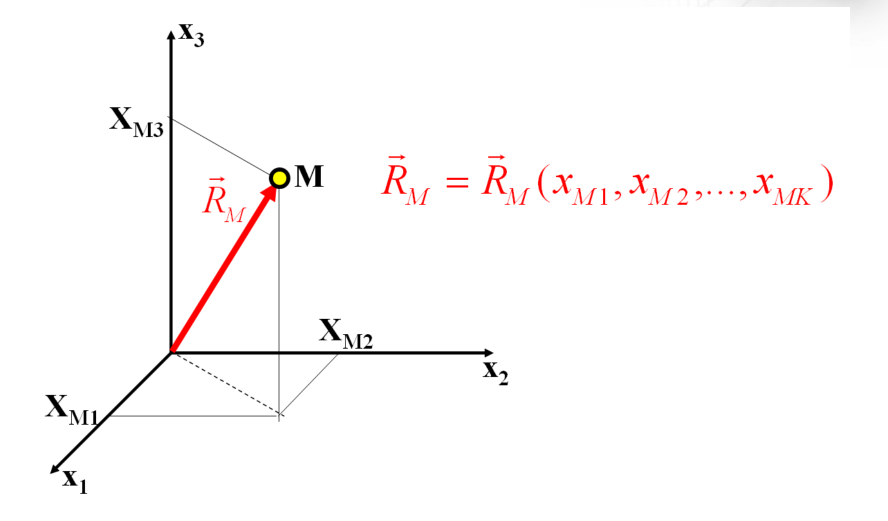
(connectivity table, label-types of bonds)

Datasets are represented by a matrix of molecular descriptors



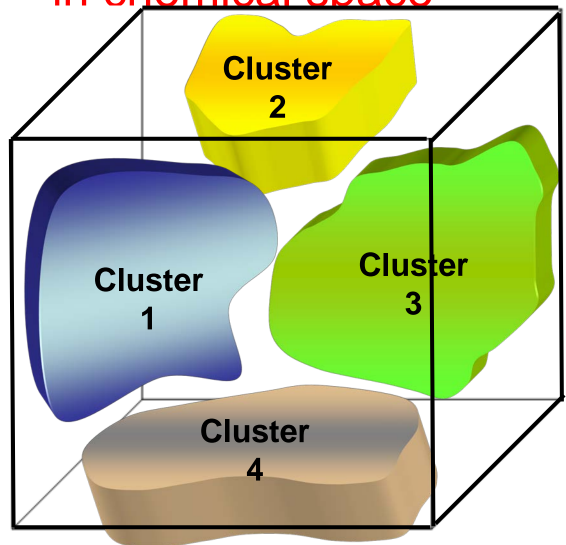
| Samples | Variables (descriptors) | | | | | | | |
|-------------|-------------------------|-----------------|-----|-----------------|--|--|--|--|
| (Compounds) | X ₁ | X ₂ | | X _m | | | | |
| 1 | X ₁₁ | X ₁₂ | ••• | X _{1m} | | | | |
| 2 | X ₂₁ | X ₂₂ | ••• | X _{2m} | | | | |
| | | | | | | | | |
| n | X _{n1} | X _{n2} | ••• | X _{nm} | | | | |

Compounds represented by vectors in a multidimensional descriptor space



Molecules may form clusters in chemical space





Molecules are considered as vectors in the space of descriptors (« chemical » space).

Dimensions of this space correspond to the number of descriptors.

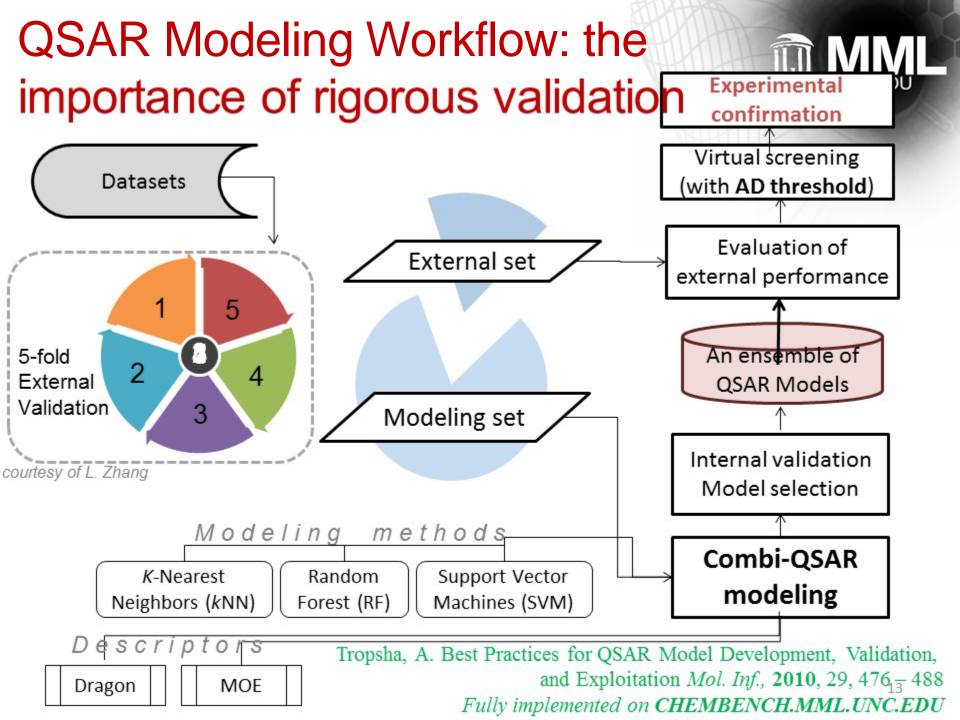
Clustering methods are employed to analyze distances between compounds and identify clusters.

QSAR Modeling

Establish <u>quantitative relationships</u> between descriptors and the target property capable of <u>predicting</u> activities of novel compounds.

| Chemistry | Bioactivity | Cheminformatics | | | | | |
|-----------------------|---------------------------|------------------|---------------------------|-----------------------|--------|-------------------------------|--|
| Chemistry | (IC50. Kd) | (M | Iolecular | Descri | ptors) | | |
| Comp.1 | (IC50, Kd) Value1 | D_1 | D_2 | D_3 | | D _n | |
| Comp.2 | Value2 | " | ** | " | | " | |
| Comp.3 | Value3 | " | " | •• | | ** | |
| Comp.N | ValueN | | | | | - | |
| | | | 2.5 - | | | | |
| BA = F(D) (linear | ~, (² | | - 2 - 1.5 - 1 - 1 - | and the second second | • | Training Linear (Training) | |
| e.g., -LogIC50 = k | $_1D_1+k_2D_2+\ldots+k_n$ | D _n) | Predicte | | | | |
| or non-linear, e.g. k | nearest neighbor | S | 0.5 - | | | | |
| | | | 0 | | | | |

Actual LogED50 (ED50 = mWkg)



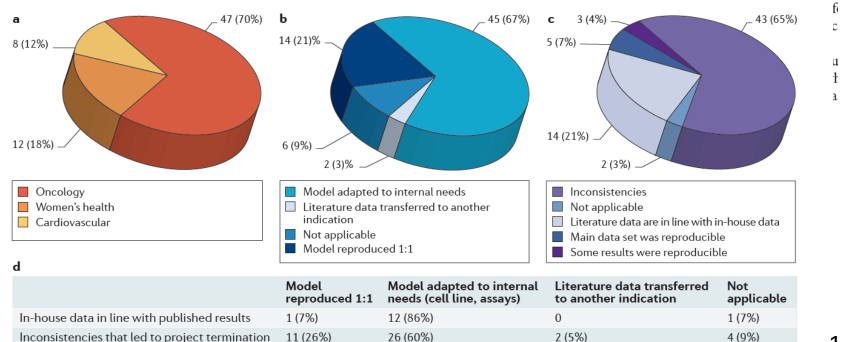
Data dependency and data quality are critical issues in QSAR modeling

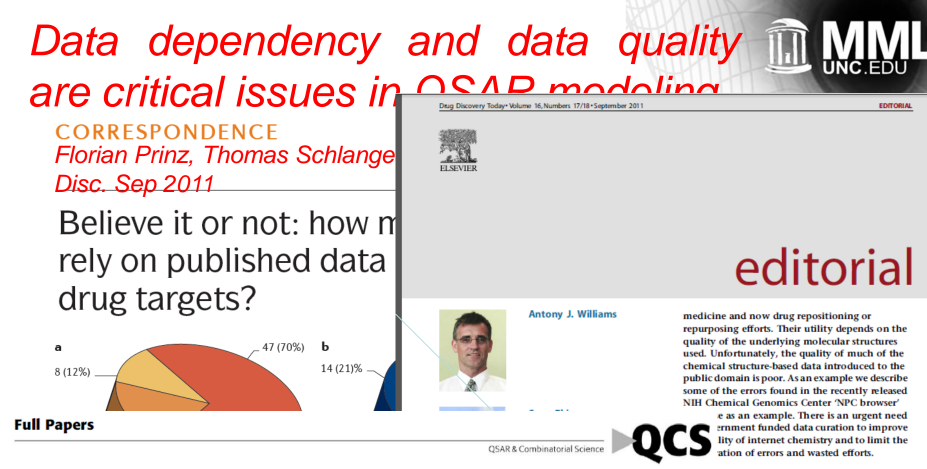
CORRESPONDENCE

LINK TO ORIGINAL ARTICL

Florian Prinz, Thomas Schlange and Khusru Asadullah. Nature Rev. Drug Disc. Sep 2011

Believe it or not: how much can we rely on published data on potential drug targets? results that are published are hard to reproduce. However, there is an imbalance betwee this apparently widespread impression and i public recognition (for example, see REFS 2,3 and the surprisingly few scientific public tions dealing with this topic. Indeed, to or knowledge, so far there has been no publishe in-depth, systematic analysis that compar





Are the Chemical Structures in Your QSAR Correct?

Douglas Young^a*, Todd Martin^a, Raghuraman Venkatapathy^b, and Paul Harten^a

- ^a US Environmental Protection Agency, 26 West Martin Luther King Drive, Cincinnati, OH 45268, USA; E-mail: young.douglas@epa.gov
- ^b Pegasus Technical Services, 26 West Martin Luther King Drive, Cincinnati, OH 45268, USA

Keywords: Databases, N-octanol/water partition coefficient, Quantitative structure-activity relationships, SMILES

Received: June 26, 2008; Revised: August 13, 2008; Accepted: August 21, 2008

DOI: 10.1002/qsar.200810084

In the last ten years, public online databases have

ling agencies have been investing in the development of main chemistry platforms with the primary attention in to the informatics platform itself rather the quality of

content. This is clearly exemplified by the recently PC browser from the NIH Chemical Genomics Center

1]. Public online databases such as PubChem, ChemIDid the EPA's ACTOR [3], to name just a few, have rapidly

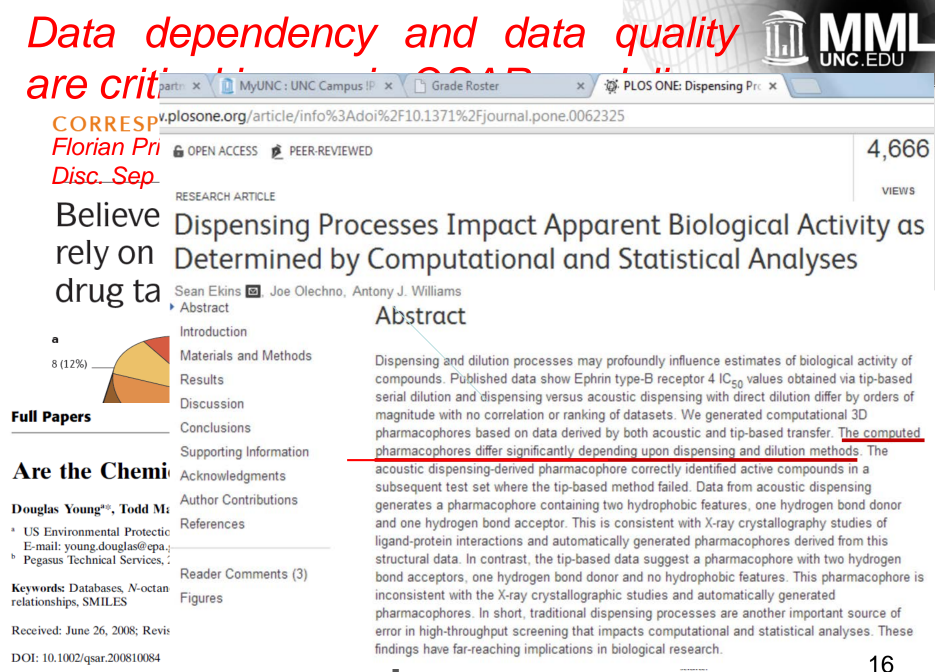
rusted valuable resources which researchers rely on for

able chemical structures and associated data. While emistry databases can certainly be of value, we feel the ould be immediately alerted to consider issues of data

hen using these resources and we call into question both

is and the trust we place in them. To our knowledge the raise, using the example of a recently released database, been described elsewhere and the user community, and

gencies, should not ignore them any longer. The develf cheminformatics platforms without due care given to



In the last ten years, public online databases have

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|--|---|--|-----|
| Di | « Hacks for Septa | Organometallics Responds t | 0 1 |
| B r€ | A Disturbing Note in a Recent SI File August 6th, 2013 | | |
| a 8 (12 | A recently published ASAP <u>article</u> in the journal Organome to raise some eyebrows in the chemical community. While itself is a straightforward study of palladium and platinum complexes, page 12 of the corresponding Supporting Info contains what appears to be an editorial note that was ina in the published document: | e the paper n bis-sulfoxide ormation <u>file</u> | |
| Full Pape | Emma, please insert NMR data here! where are they compound, just make up an elemental analysis | | _ |
| Are th Douglas Ye | This statement goes beyond a simple embarrassing failure edit the manuscript, as it appears the first author is being fabricate data. Elemental analyses would be very easy to long-time readers of this blog will recall how fake element <u>campaign of fraud</u> in the work she published from 2002 to | instructed to In fabricate, and tal analyses were pivotal to | Be |
| E-mail: yc ^b Pegasus T Keywords: I relationships Received: Ju | The compound labeled 14 (an acac complex) in the main compound 14 in the SI. In fact, the bridged-dichloride co intermediate in Scheme 5, which should raise more eyebr order to avoid having to provide robust characterization fo | mpound appears to be listed rows. Did the authors unlist | d a |
| DOI: 10.100 | ChemBark is contacting the <u>corresponding author</u> for com when we receive it. | • | b |

Data dependency and data quality

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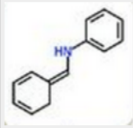
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In the Pipeline

http://pipeline.corante.com/archives/2014/04/11/biology_maybe_right_c hemistry_ridiculously_wrong.php

April 11, 2014

Biology Maybe Right, Chemistry Ridiculously Wrong



As my correspondent (a chemist himself) mentions, a close look at Figure 2 of the paper raises some real questions. Take a look at that cyclohexadiene enamine - can that really be drawn correctly, or isn't it just N-phenylbenzylamine? The problem is, that compound (drawn correctly) shows up elsewhere in Figure 2, *hitting a completely different pathway.* These two tautomers are not going to have different biological effects, partly because the first one would exist for about two molecular vibrations before it converted to the second. But how could both of them appear on the same figure?



And look at what they're calling "cyclohexa-2,4-dien-1-one". No such compound exists as such in the real world - we call it phenol, and we draw it as an aromatic ring with an OH coming from it. Thiazolidinedione is listed as "thiazolidine-2,4-quinone". Both of these would lead to red "X" marks on an undergraduate exam paper. It is clear that no chemist, not even someone who's been through second-year organic class, was involved in this work (or at the very least, involved in the preparation of Figure 2). Why not? Who reviewed this,

anyway?

DOI: 10.100

when we receive it.

In the last ten years, public online databases have In the

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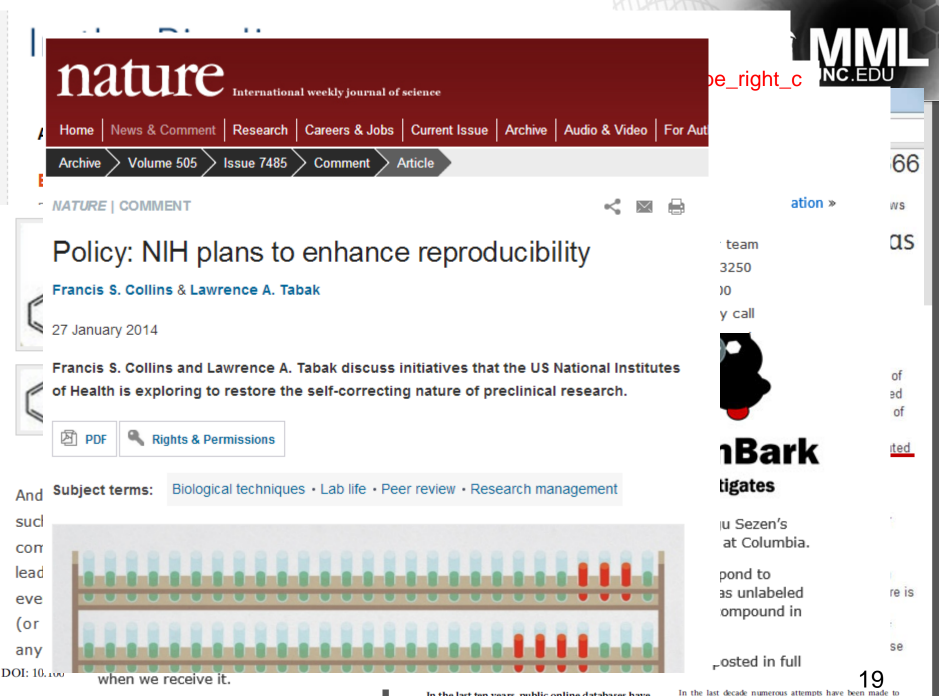
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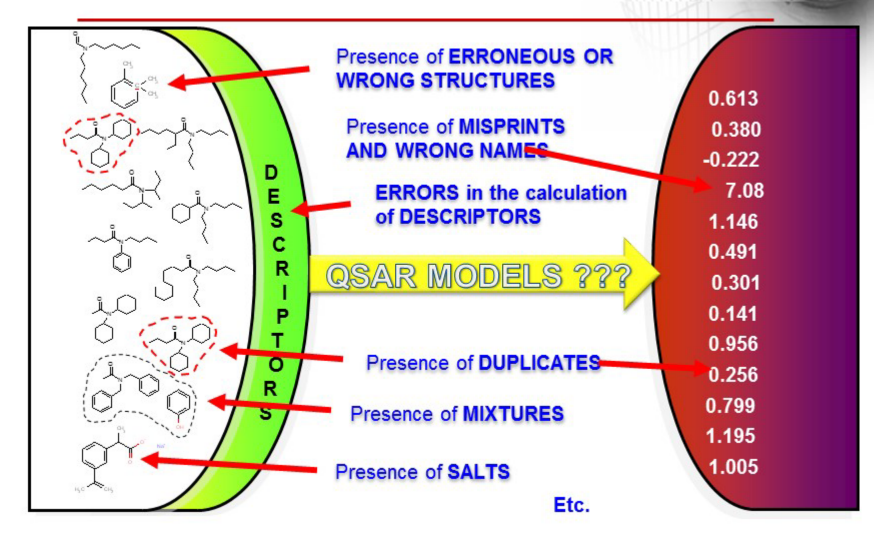
In the last ten years, public online databases have



In the last ten years, public online databases have

In the last decade numerous attempts have been made to

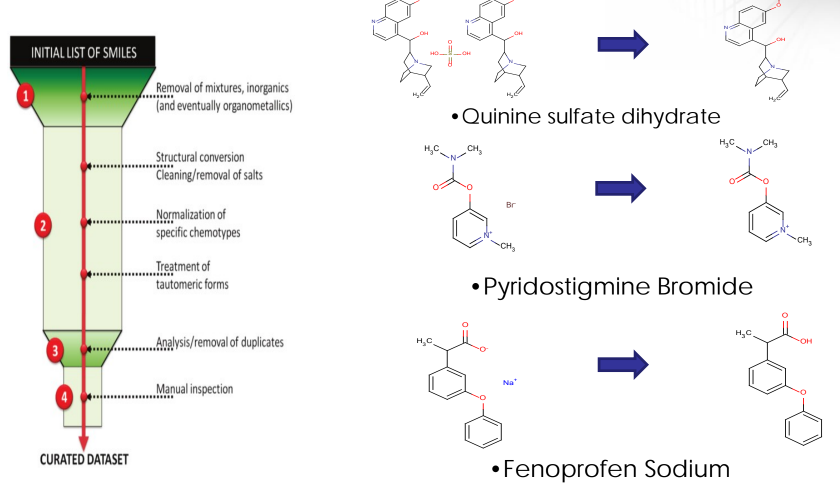
QSAR modeling with non-curated datasets



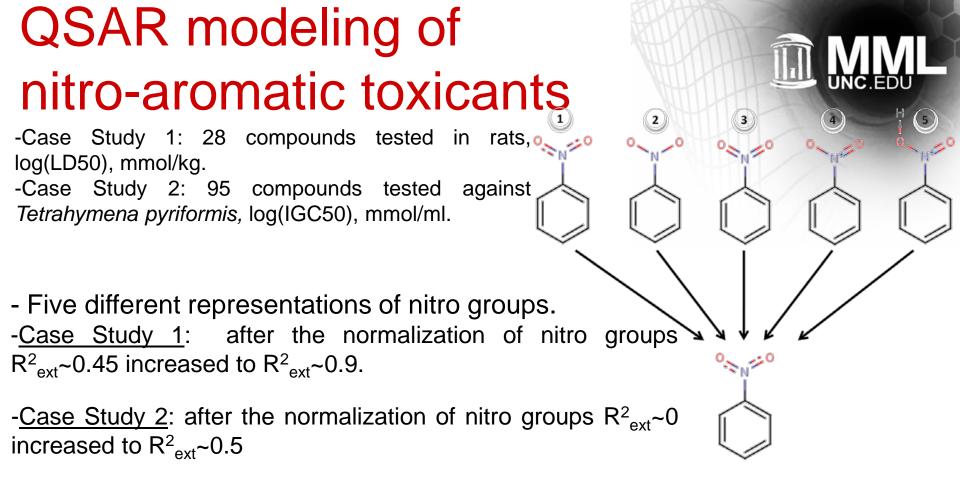
Chemical Structure Curation



Chemical structures should be cleaned and standardized (duplicates removed, salts stripped, neutral form, canonical tautomer, etc) to enable rigorous model development

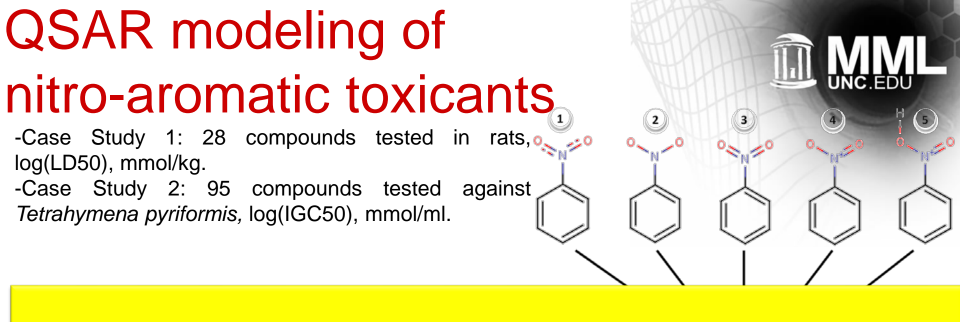


Muratov, Fourches, Tropsha. Trust but verify. *JC J. Chem. Inf. Model.* **2010**, 50, 1189-1204.



Even small differences in structure representation can lead to significant errors in prediction accuracy of models

Artemenko, Muratov et al. SAR QSAR 2011, 22 (5-6), 1-27.



Data curation affects the accuracy (up or down!) of QSAR models

Even small differences in structure representation can lead to significant errors in prediction accuracy of models

Artemenko, Muratov et al. SAR QSAR 2011, 22 (5-6), 1-27.

Curation of Bioactivity: Case study MMI

Predictive Models for Cytochrome P450 Isozymes Based on Quantitative High Throughput Screening Data

Hongmao Sun,*^{,†} Henrike Veith,[†] Menghang Xia,[†] Christopher P. Austin,[†] and Ruili Huang[†]

[†]National Institutes of Health (NIH) Chemical Genomics Center, NIH,

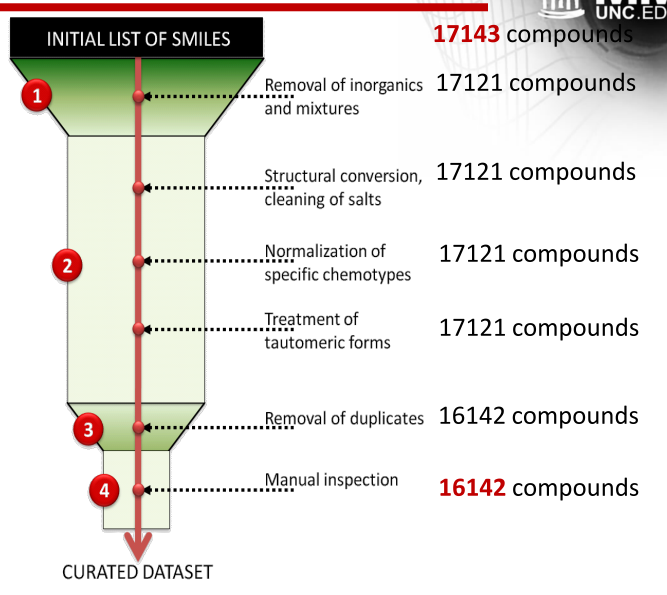
ABSTRACT: The human cytochrome P450 (CYP450) isozymes are the most important enzymes in the body to metabolize many endogenous and exogenous substances including environmental toxins and therapeutic drugs. Any unnecessary interactions between a small molecule and CYP450 isozymes may raise a potential to disarm the integrity of the protection. Accurately predicting the potential interactions between a small molecule and CYP450 isozymes is highly desirable for assessing the metabolic stability and toxicity of the molecule. The National Institutes of Health Chemical Genomics Center (NCGC) has screened a collection of over 17,000 compounds against the five major isozymes of CYP450 (1A2, 2C9, 2C19, 2D6, and 3A4) in a quantitative high throughput screening (qHTS) format. In this study, we developed support vector classification (SVC) models

CONCLUSION

SVM classification models have been built for the five most important isoforms of CYP450 (1A2, 2C9, 2C19, 2D6, and 3A4) based on a large gHTS data set with over 6000 compounds available for both model training and testing. The five CV optimized SVC models built by using the atom typing molecular descriptors exhibited consistently high predictive power when applied to the equally populated test sets with accuracies between 0.85 and 0.93, as measured by the AUC of ROC plots. The results indicated that the atom typing descriptors generated from a large, high quality data set were capable of feeding information rich learning materials to the SVM learner. Useful information of structural features was derived from feature importance analysis for each isozyme of CYP450. The privileged structural features that could result in inhibitory and stimulatory activity against different CYP450 isozymes can serve as valuable guidelines in the drug discovery process.

for these five isozymes using a set of customized generic atom types. The CYP450 data sets were randomly split into equal-sized training and test sets. The optimized SVC models exhibited high predictive power against the test sets for all five CYP450 isozymes with accuracies of 0.93, 0.89, 0.89, 0.85, and 0.87 for 1A2, 2C9, 2C19, 2D6, and 3A4, respectively, as measured by the area under the receiver operating characteristic (ROC) curves. The important atom types and features extracted from the five models are consistent with the structural preferences for different CYP450 substrates reported in the literature. We also identified novel features with significant discerning power to separate CYP450 actives from inactives. These models can be useful in prioritizing compounds in a drug discovery pipeline or recognizing the toxic potential of environmental chemicals.

Dataset Curation summary



Fourches D, et al. J Chem Inf Model. 2010 50(7):1189-204.

NCGC dataset: analysis of duplicates



- Out of 1280 duplicate couples :
 - 406 had no discrepancies-no values or no values for comparison
 - 874 had biological profile differences
- A total of 1535 discrepancies were found in the

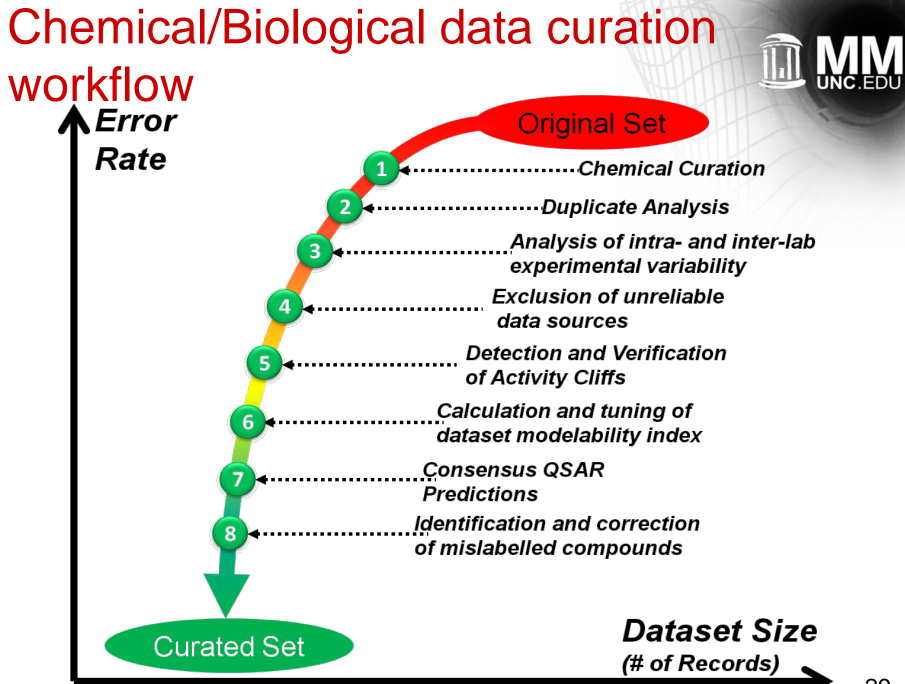
| | CYP2C9 | CYP1A2 | CYP3A4 | CYP2D6 | CYP2C19 |
|-----------------------|--------|--------|--------|--------|---------|
| # of discrepancies | 154 | 363 | 426 | 422 | 170 |

Neighborhood Analysis for Duplicates MM

17,000 compounds screened against five major CYP450 isozymes. 1,280 pairs of duplicates couples were found (874 had different bioprofiles)

| Tocris-0740 | SID | Supplier | 2C9 | 1A2 | 3A4 | 2D6 | 2C19 |
|-------------|----------|---------------|------|------|------|------|------|
| CID_6603937 | 11113673 | Tocris | -4.6 | -4.4 | -4.6 | -6.2 | -4.5 |
| CID_6603937 | 11111504 | Sigma Aldrich | -4.4 | | R | -5.6 | -5 |

| 5 Nearest neighbors | Tanimoto Similarity | SID | Supplier | 2C9 | 1A2 | 3A4 | 2D6 | 2C19 | |
|------------------------|------------------------|----------|---------------|-----|------|------|------|------|---------|
| 6604862 | 0.98 | 11114071 | Tocris | | | -4.5 | | -5.5 | |
| 6604106 | 0.98 | 11112029 | Sigma Aldrich | | | -5.1 | | | |
| 6604846 | 0.98 | 11114012 | Tocris | | | | | | |
| 6604136 | 0.95 | 11112054 | Sigma Aldrich | | | -4.8 | -5.9 | | HO HO N |
| 6604137 | 0.95 | 11113764 | Tocris | | -4.4 | -4.7 | -4.5 | | 28 |



Fourches, Muratov, Tropsha. Nat Chem Biol. 2015, 11(8):535.

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Published guidance on model development and validation: The OECD Principles



To facilitate the consideration of a QSAR model for regulatory purposes, it should be associated with the following information:

- a defined endpoint
- > an unambiguous algorithm;
- a defined domain of applicability

> appropriate measures of goodness-of-fit, robustness and predictivity

>a mechanistic interpretation if possible

Should be added: data used for modeling should be carefully curated

21 "how not to do QSAR" principles MI

Table 1. Types of error in QSAR/QSPR development and use.

| No. | Type of error | Relevant OECD principle(s) |
|-----|--|-------------------------------|
| 1 | Failure to take account of data heterogeneity | 1 |
| 2 | Use of inappropriate endpoint data | 1 |
| 3 | Use of collinear descriptors | 2, 4, 5 |
| 4 | Use of incomprehensible descriptors | 2, 5 |
| 5 | Error in descriptor values | 2 2 |
| 6 | Poor transferability of QSAR/QSPR | 2 |
| 7 | Inadequate/undefined applicability domain | 3 |
| 8 | Unacknowledged omission of data points | 3 3 |
| 9 | Use of inadequate data | |
| 10 | Replication of compounds in dataset | 3 |
| 11 | Too narrow a range of endpoint values | 3 |
| 12 | Over-fitting of data | 4 |
| 13 | Use of excessive numbers of descriptors in a QSAR/QSPR | 4 |
| 14 | Lack of/inadequate statistics | 4 |
| 15 | Incorrect calculation | 4 |
| 16 | Lack of descriptor auto-scaling | 4 |
| 17 | Misuse/misinterpretation of statistics | 4 |
| 18 | No consideration of distribution of residuals | 4 |
| 19 | Inadequate training/test set selection | 4 |
| 20 | Failure to validate a QSAR/QSPR correctly | 4 |
| 21 | Lack of mechanistic interpretation | 5 |

Dearden JC et al., 2009, SAR and QSAR in Environmental Research, Vol. 20, Nos. 3–4, April–June 2009, 241

Model accuracy and interpretation: ³² Case studies (modeling of skin sensitization ³² and Ames genotoxicity)

- The Local Lymph Node Assay (LLNA) is generally regarded as the preferred test for evaluating skin sensitization.¹
- Although LLNA has a good correlation with human skin sensitization, it has been shown that LLNA fails in several cases to predict human skin sensitization.²
- Ca. 3.89% (39,090) of the 1,004,873 animals used for safety testing in Europe are used in skin sensitization/irritation tests²; this creates a strong need to evaluate skin sensitization potential for a chemical without expensive and time-consuming animal testing.

In silico methods are highly recommended for time and cost saving of skin-related

research.⁴

¹OECD. Test No. 429: Skin Sensitisation http://iccvam.niehs.nih.gov/SuppDocs/FedDocs/OECD/OECD-TG429-2010.pdf (accessed Jan 23, 2013).

²Api, A. M.; Basketter, D.; Lalko, J.; Basketter, D.; Lalko, J. Cutan. Ocul. Toxicol. 2014, 9527, 1–5.

²European Commission. Seventh teport on the statistics on the number of animals used for experimental and other scientific purposes in the member states of the **2013** ⁴European Commission. On the animal testing and marketing ban and on the state of play in relation to alternative methods in the field of cosmetics **2013**.

Model accuracy and interpretation: Case studies

- QSAR models of skin sensitization and their application to identify potentially hazardous compounds (Alves VM, Muratov E, Fourches D, Strickland J, Kleinstreuer N, Andrade CH, Tropsha A. <u>Toxicol Appl Pharmacol. 2015</u> 284(2):262-72)
- QSAR models of skin permeability and the relationships between skin permeability and skin sensitization (Alves VM, Muratov E, Fourches D, Strickland J, Kleinstreuer N, Andrade CH, Tropsha A. <u>Toxicol Appl Pharmacol. 2015</u> 284(2):273-80)
- QSAR models of human data could replace mLLNA test for predicting human skin sensitization potential of chemicals (Alves VM, Muratov E, Fourches D, Strickland J, Kleinstreuer N, Andrade CH, Tropsha A. In preparation).

Skin Sensitization Dataset (mLLNA)



Source

ICCVAM (Interagency Coordinating Committee on the Validation of Alternative Methods) report 2009

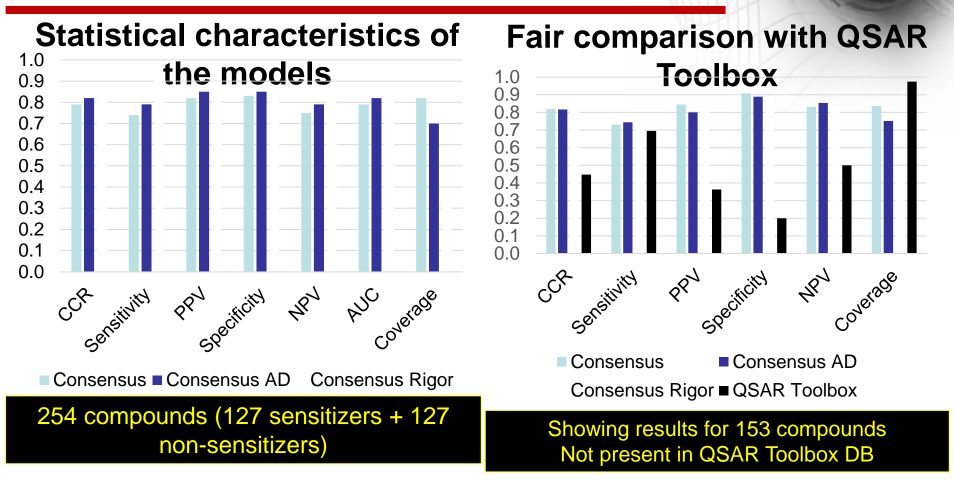
| Vehicle type | Non- sensitizer | Sensitizer | Total | 471 records |
|----------------------|----------------------|---------------------|---------------|---------------------------------|
| ACE | 14 | 31 | 45 | |
| AOO | 51 | 178 | 229 | 408 found by |
| dH ₂ O | 2 | 2 | 4 | name |
| DMF | 40 | 27 | 67 | Removal of |
| DMSO | 16 | 15 | 31 | Organometallic compound |
| PG | 6 | 8 | 14 | Inorganic salts |
| Pluronic L92 (1%) | 2 | 5 | 7 | Duplicates Dataset balancing |
| Others | 4 | 7 | 11 | Dataset balancing |
| Total | 135 | 273 | 408 | 387 |
| bbreviations: AOO. | acetone&olive oil (4 | :1 by volume); ACE, | acetone: DMF. | e e nom e u mele |

acelone&olive oli (4.1 by volume), ACE, a dimethyl formamide; DMSO, dimethyl sulfoxide; PG, propylene glycol. compounds

254 compounds were retained for QSAR modeling: 127 non-sensitizers + 127 sensitizers

133 remaining sensitizers were used for additional external validation³⁴

QSAR models of skin sensitization (mLLNA)



Models were built using **Random Forest** approach – 5-fold External CV results

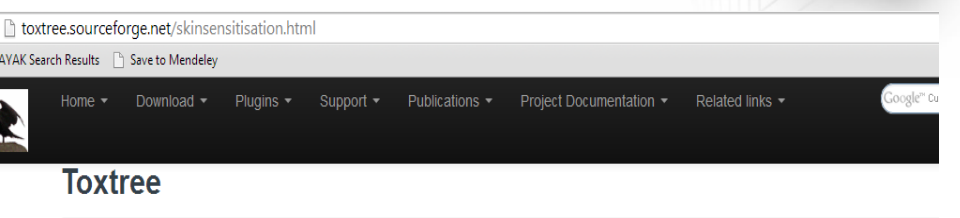
ALERTS vs. QSAR: ACTIVATED PYRIDINE/PYRIMIDINE

| 0 0 | QSAR Toolbox | QSAR | Experiment |
|--|---|-------------------|-------------------|
| H ₃ C CI Ethyl 2,6-dichloro-5-fluoro-b-oxo-3- pyridinepropanoate | Contains Activated Pyridine Sensitizer | Non Sensitizer | Non Sensitizer |
| H ₃ C N CH ₃ H ₃ C N CH ₃ H ₃ C N CH ₃ C CH | Contains Activated Pyridine Sensitizer | Non Sensitizer | Non Sensitizer |
| H ₃ C – N – V – V – V – V – V – V – V – V – V | Contains Activated Pyridine Sensitizer | Non Sensitizer | Non Sensitizer |

ALERTS vs. QSAR: NO PROTEIN BINDING ALERTS

| CH ₃ | QSAR Toolbox | QSAR | Experiment |
|---|----------------------------|-------------------|-------------------|
| H ₃ C F F F F F F F F F F F F F F F F F F F | No alert Sensitizer | Non Sensitizer | Non Sensitizer |
| 1-[3-(Cyclopentyloxy)-4-methoxy-phenyl]-4- oxocyclohexane carbonitrile | No alert Sensitizer | Non Sensitizer | Non Sensitizer |
| H ₃ C CH ₃ H ₂ N H ₃ C NH ₂ 3-Aminomethyl- 3,5,5- trimethylcyclohexyl amine | No alert Non sensitizer | Sensitizer | Sensitizer |

Chemical Alerts (rules) of Toxicity: are they truly reliable?



Last Published: 2014-06-15 | Version: 2.6.6

Skin sensitisation reactivity domains

Identification of mechanisms of toxic action for skin sensitisation using a SMARTS pattern based approach.

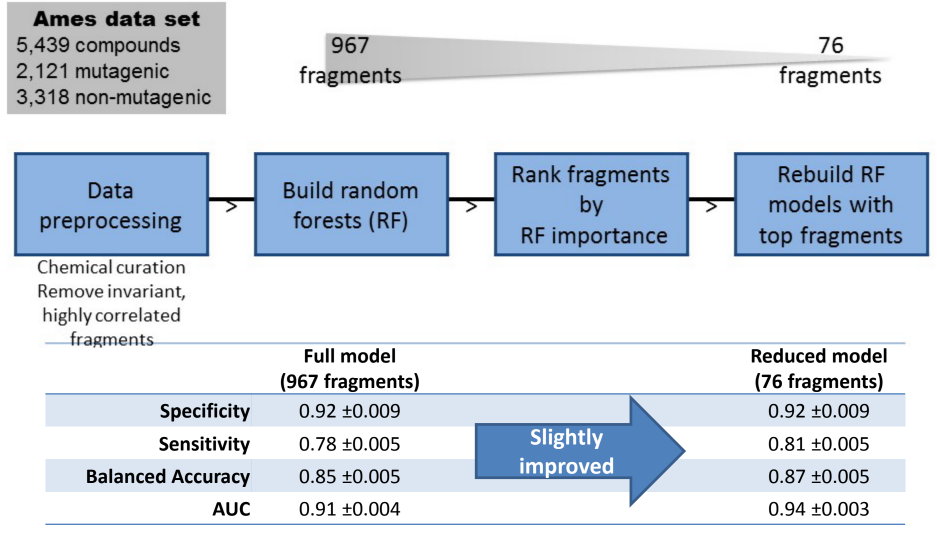
Available since ToxTree 2.1.0 (under name "Skin sensitisation alerts" and "Skin sensitisation alerts (M.Cronin)"). The name is changed to "Skin sensitisation reactivity domain" by P&G team suggestion in order to reflect the fact the alerts provide grouping into reactivity mode of action and do not predict skin sensitisation potential.

Developed by IdeaConsult Ltd. (2), (Sofia, Bulgaria), with collaboration with and support from Procter and Gamble (2) 2010

Chemical Alerts (rules) of Toxicity: are they truly reliable?



Model interpretation: identifying statistically important fragments as complex alerts

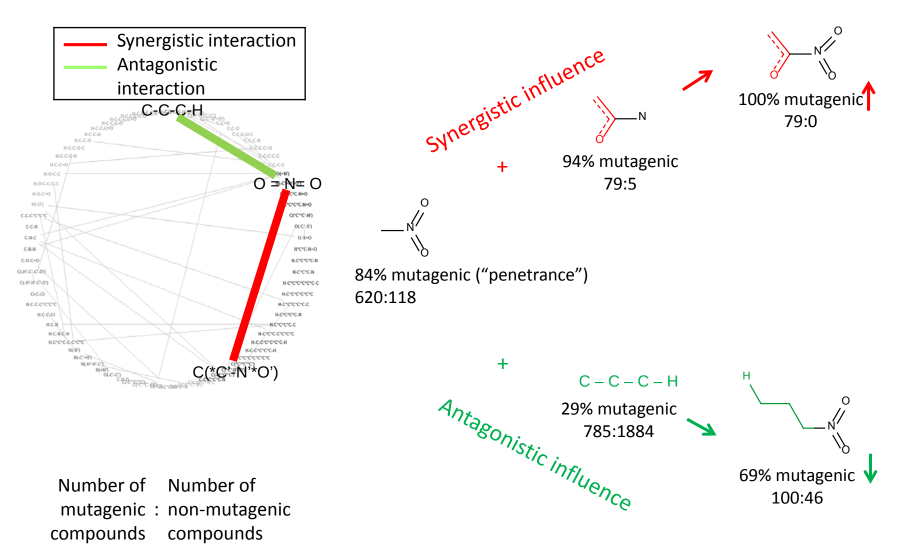


Results from 5-fold external cross validation

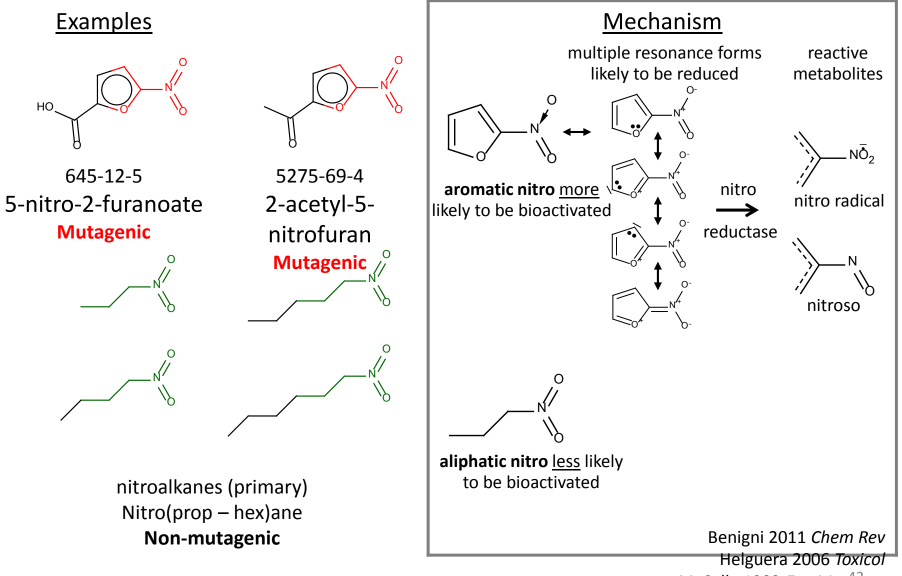
Example of fragment (alert) interaction

Nitro's mutagenic effect is:

increased by furan (synergism) decreased by primary alkanes(antagonism)

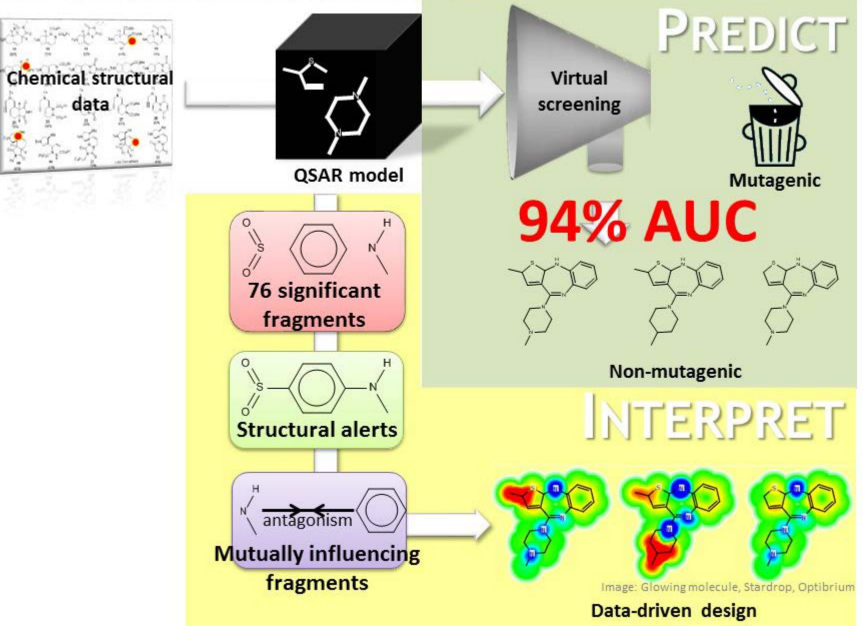


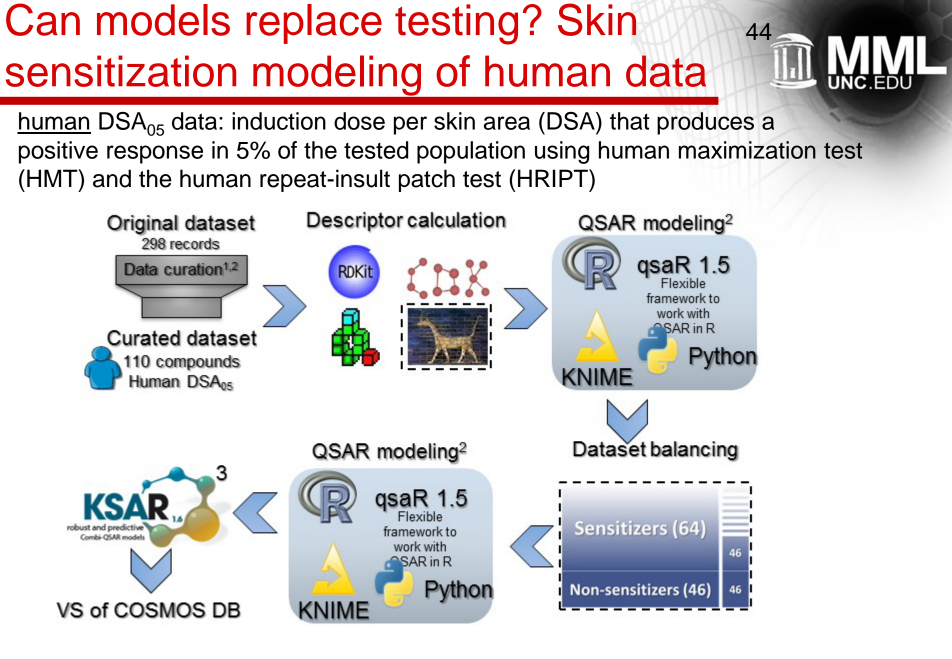
Nitro compounds are <u>active</u> when paired with aromatic rings inactive when paired with primary alkanes



McCalla 1983 Env Mutagen

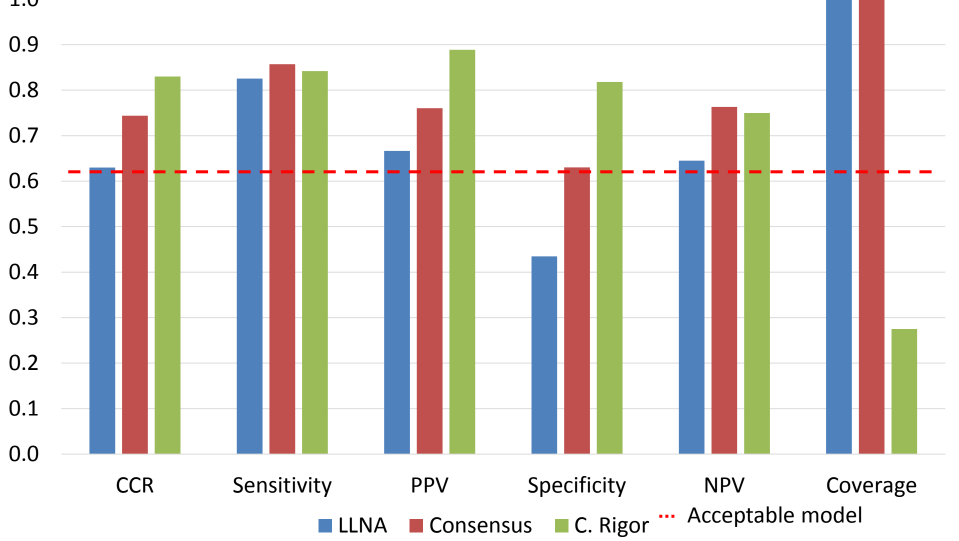
Marrying SAR and QSAR in CWAS: Deriving alerts from validated QSAR models





¹Fourches, D.; Muratov, E.; Tropsha, A. *J. Chem. Inf. Model.* **2010**, *50*, 1189–1204. ²Tropsha, A. *Mol. Inform.* **2010**, *29*, 476–488. ³Braga, R. C.; Alves, V. M. et al. *Curr. Top. Med. Chem.* **2014**, *14*, 1399–1415.

Comparison of external predictive accuracy for human data: QSAR gives more reliable predictions than mLLNA



Accessed by 5-fold external cross validation; SVM: Support Vector Machine; AD: Applicability Domain. No. of compounds = 63 sensitizers + 46 non sensitizers

QSAR and toxicity testing in the 21st century

REPORT

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BRIE

July 2007

Toxicity Testing in the 21st Century: A Vision and a Strategy

Advances in molecular biology, biotechnology, and other fields are paving the way for major improvements in how scientists evaluate the health risks posed by potentially toxic chemicals found at low levels in the environment. These advances would make toxicity testing quicker, less expensive, and more directly relevant to human exposures. They could also reduce the need for animal testing by substituting more laboratory tests based on human cells. This National Research Council report creates a far-reaching vision for the future of toxicity testing.

oxicity tests on laboratory animals are conducted to evaluate chemicals-including medicines, food additives, and industrial, consumer, and agricultural chemicals-for their potential to cause cancer, birth defects, and other adverse health effects. Information from toxicity testing serves as an important part of the basis for public health and regulatory decisions concerning toxic chemicals. Current test

methods were developed incrementally over the past 50 to 60 years and are conducted using laboratory animals, such as rats and mice. Using the results of animal tests to predict human health effects involves a number of assumptions and extrapolations that remain controversial. Test animals are often exposed to higher doses than would be expected for typical human exposures, requiring assumptions about



effects at lower doses or exposures. Test animals are typically observed for overt signs of adverse health effects, which provide little information about biological changes leading to such health effects. Often controversial uncertainty factors must be applied to account for differences between test animals and humans. Finally, use of animals in testing is expensive and time consuming, and it sometimes raises ethical issues.

Today, toxicological evaluation of chemicals is poised to take advantage of the on-going revolution in biology and biotechnology. This revolution is making it increasingly possible to study the effects of chemicals using cells, cellular components, and tissues-preferably of human origin—rather than whole animals.

POLICYFORUM

TOXICOLOGY

Transforming Environmental **Health Protection**

Francis S. Collins,1*1 George M. Gray,2* John R. Bucher3*

n 2005, the U.S. Environmental Protection Agency (EPA), with support from the U.S. National Toxicology Program (NTP), funded a project at the National Research Council (NRC) to develop a long-range vision for toxicity testing and a strategic plan for implementing that vision. Both agencies wanted future toxicity testing and assessment paradigms to meet evolving regulatory needs. Challenges include the large numbers of substances that need to be tested and how to incorporate recent advances in molecular toxicology, computational sciences, and information technology; to rely increasingly on human as opposed to animal data; and to offer increased

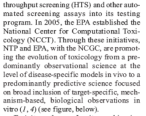
d costs (1-5). In mittee on Toxicity of Environmental ports that reviewed lentified key issues, nd implementation shift in the assessrd and risk (6, 7). have laid out a solid prehensive and rigd comparisons with determine whether ments will be realourpose, NTP, EPA, of Health Chemical GC) (organizations

with expertise in experimental toxicology, computational toxicology, and high-throughput technologies, respectively) have established a collaborative research program.

EPA, NCGC, and NTP Joint Activities

In 2004, the NTP released its vision and roadmap for the 21st century (1), which established initiatives to integrate high-

¹Director, National Human Genome Research Institute (NHGRI), National Institutes of Health, Bethesda, MD 20892; ²Assistant Administrator for the Office of Research and Development, U.S. Environmental Protection Agency, Washington, DC 20460; ³Associate Director, U.S. National Toxicology Program, National Institute of Environmental



Toxicity pathways. In vitro and in vivo tools are being used to identify cellular responses after chemical exposure expected to result in adverse health effects (7). HTS methods are a primary means of discovery for drug development, and screening of >100,000 compounds per day is routine (8). However, drug-discovery HTS methods traditionally test compounds at one concentra-

15 FEBRUARY 2008 VOL 319 SCIENCE www.sciencemag.org

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.

tion, usually between 2 and 10 µM, and tolerate high false-negative rates. In contrast, in the EPA, NCGC, and NTP combined effort, all compounds are tested at as many as 15 concentrations, generally ranging from ~5 nM to ~100 µM, to generate a concentrationresponse curve (9). This approach is highly reproducible, produces significantly lower false-positive and false-negative rates than the traditional HTS methods (9), and facilitates multiassay comparisons. Finally, an informatics platform has been built to compare results among HTS screens; this is being expanded to allow comparisons with historical toxicologic NTP and EPA data (http://ncgc.nih.gov/pub/openhts). HTS data collected by EPA and NTP, as well as by the NCGC and other Molecular Libraries Initiative centers (http://mli.nih.gov/), are being made publicly available through Webbased databases [e.g., PubChem (http:// pubchem.ncbi.nlm.nih.gov)]. In addition,

Biochemical- and cell-based Human experience in vitro assays ical tests 1-3 studies/vear 10-100/vear 100-10.000/v Computational toxic ology Critical toxicity pathways



National Academy of Scie



Slide courtesy of Dr. Ann Richard, EPA (modified)

EPAs Contribution: The ToxCast Research Program

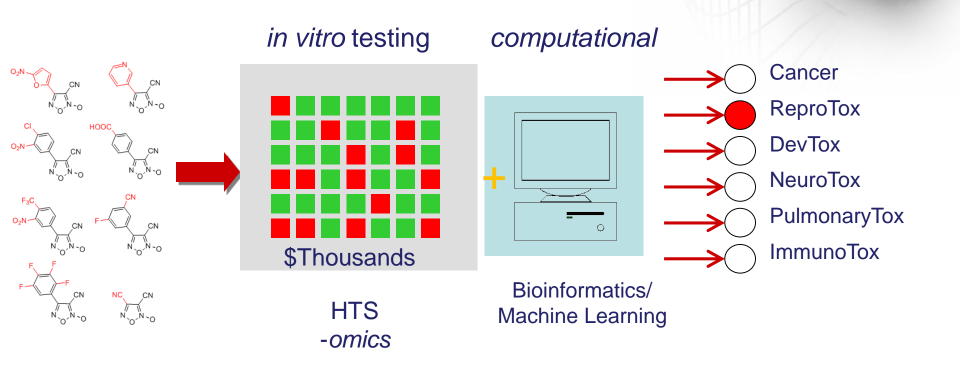
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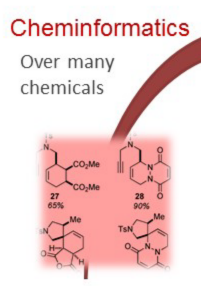
QSAR and Chemical Toxicity Testing in the 21 Century





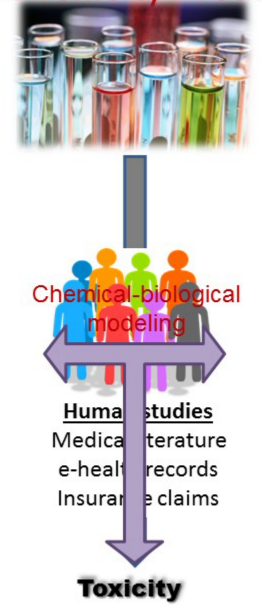
Slide courtesy of Dr. Ann Richard, EPA (modified)

Integration of Diverse Data Streams into QSAR Modeling to Improve Toxicity Prediction



Chemical descriptors (in silico):

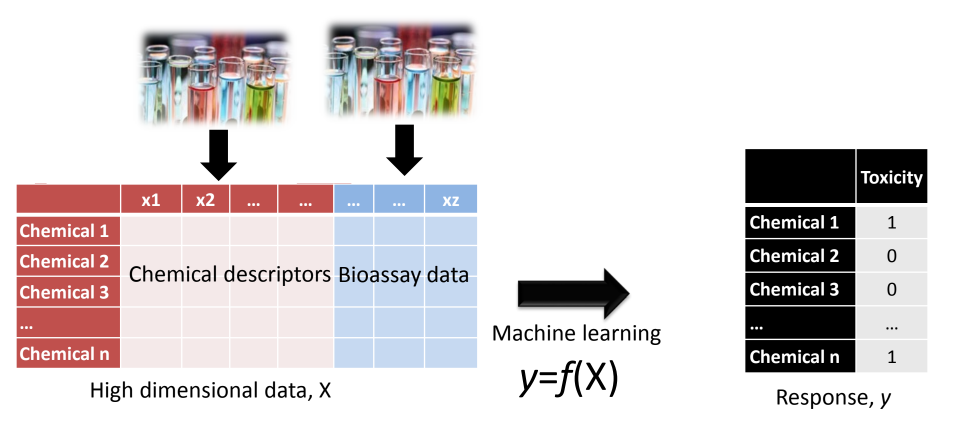
Molecular weight, Connectivity indices Presence/absence of fragment, Hydrophobicity, etc.



Bioinformatics Over many biological assays

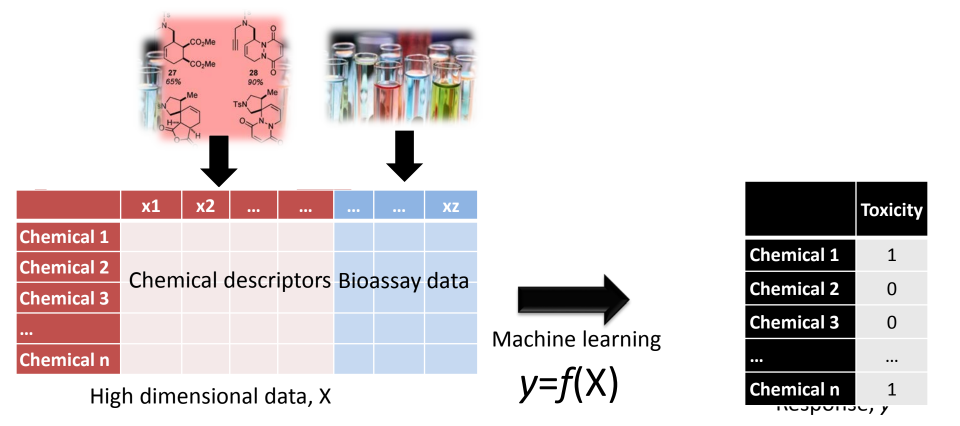
<u>Short-term biological assays</u> Transcriptomics, Metabolomics, Cytotoxicity, Genotype, etc

QSAR modeling: chemical descriptors



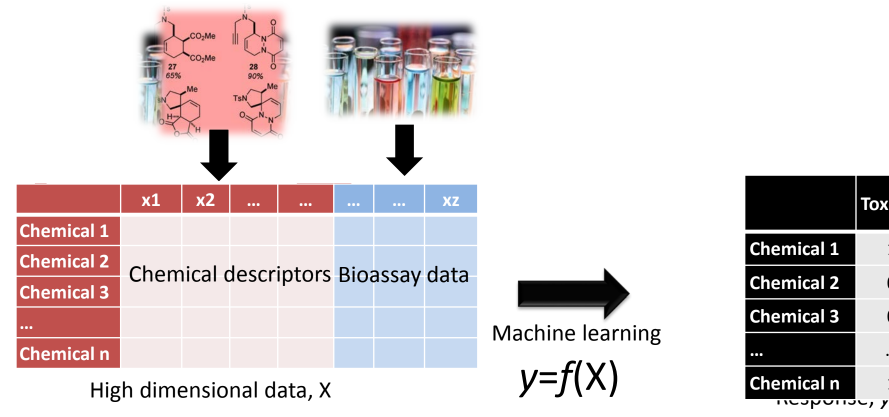
Zhu H et al. (2008) *Environ. Health Perspect.* 116, 506-513; Low Y et al. (2011) *Chem. Res. Toxicol.* 24,1251-1262; Sedykh A *et al.* (2011) *Environ. Health Perspect.* (119): 364-370

QSAR modeling: in vitro assay descriptors



Zhu H et al. (2008) *Environ. Health Perspect.* 116, 506-513; Low Y et al. (2011) *Chem. Res. Toxicol.* 24,1251-1262; Sedykh A *et al.* (2011) *Environ. Health Perspect.* (119): 364-370

QSAR modeling: hybrid descriptors



Toxicity

1

0

0

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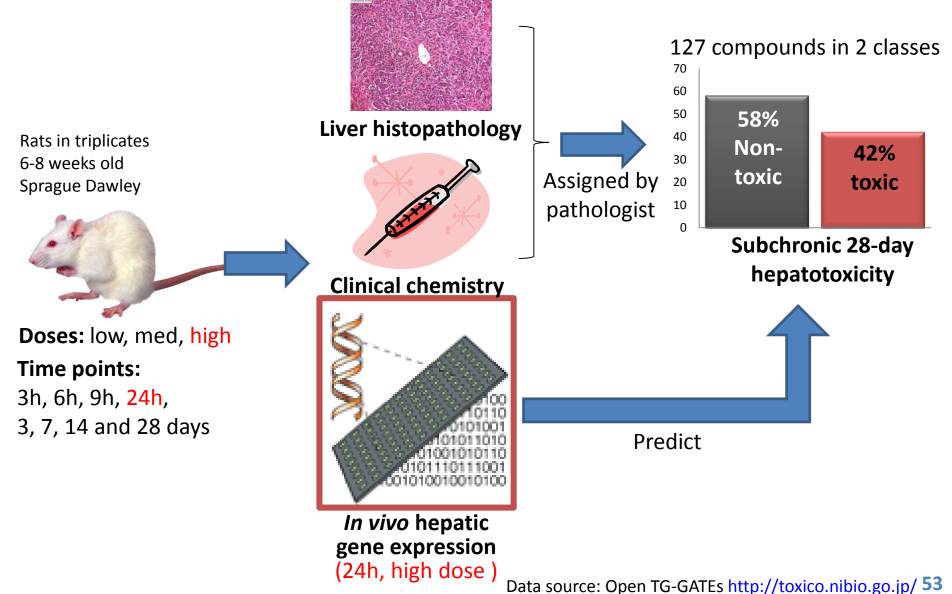
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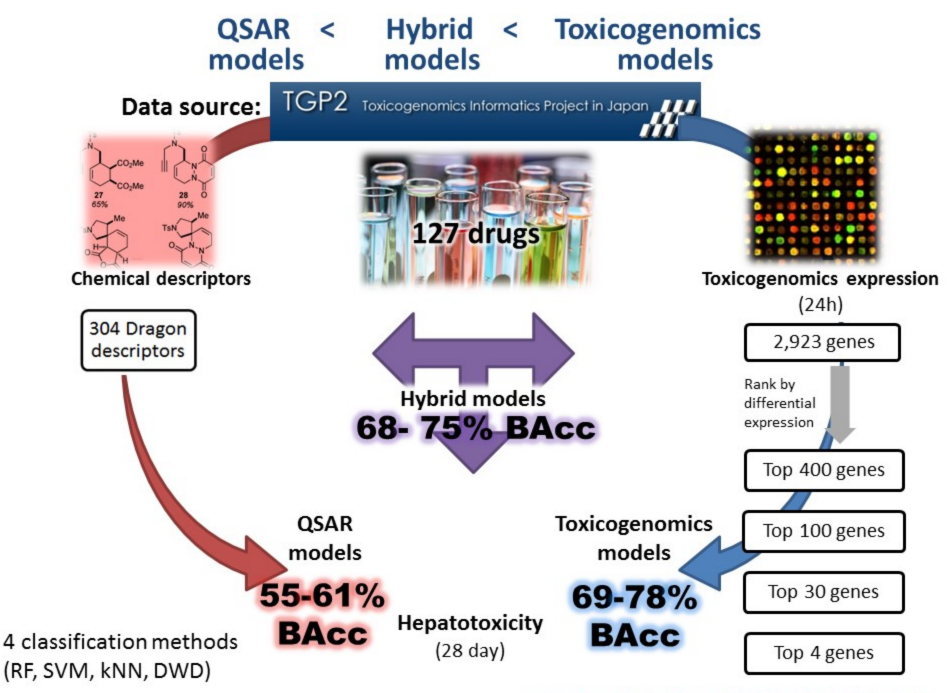
Zhu H et al. (2008) *Environ. Health Perspect.* 116, 506-513; Low Y et al. (2011) *Chem. Res. Toxicol.* 24,1251-1262; Sedykh A *et al.* (2011) *Environ. Health Perspect.* (119): 364-370

The Use of Biological Screening Data as Additional Biological Descriptors Improves the Prediction Accuracy of Conventional QSAR Models of Chemical Toxicity

- Zhu, H., *et al.* Use of cell viability assay data improves the prediction accuracy of conventional quantitative structure-activity relationship models of animal carcinogenicity. *EHP*, **2008**, (116): 506-513
- Sedykh A, *et al.* Use of in vitro HTS-derived concentration-response data as biological descriptors improves the accuracy of QSAR models of in vivo toxicity. *EHP*, **2011**, 119(3):364-70.
- Low *et al.*, Predicting drug-induced hepatotoxicity using QSAR and toxicogenomics approaches. *Chem Res Toxicol*. **2011** Aug 15;24(8):1251-62
- Rusyn *et al*, Predictive modeling of chemical hazard by integrating numerical descriptors of chemical structures and short-term toxicity assay data. *Tox. Sci.*, **2012**, 127(1):1-9
- Low Y, *et al.* Integrative chemical-biological read-across approach for chemical hazard classification. *Chem Res Toxicol.* **2013**, 26(8):1199-208
- Low, Y, *et al.* Integrative Approaches for Predicting In Vivo Effects of Chemicals from their Structural Descriptors and the Results of Short-Term Biological Assays. *Curr. Top. Med. Chem.*, 2014, 14(11):1356-64
- Low et al, Cheminformatics-Aided Pharmacovigilance: Application to Stevens Johnson Syndrome. JAMIA, 2015 (in press).

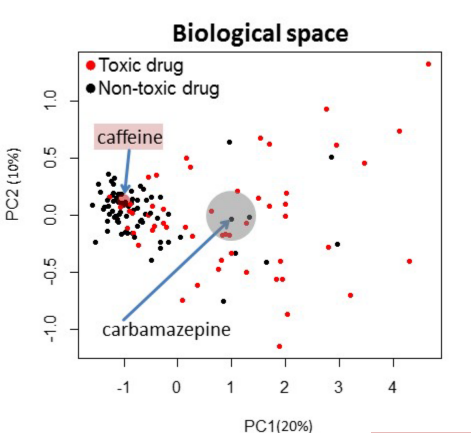
Predicting Subchronic Hepatotoxicity from 24h Toxicogenomics Profiles





Low et al. (2011) Chem. Res. Toxicol. 24,1251-1262

Conflicting Predictions by QSAR and Toxicogenomics Models



Carbamazepine

Distant biological neighbors
Close chemical neighbors
Chemical similarity works
better

Caffeine

Close biological neighbors
Distant chemical neighbors
TGx similarity works
better

PC1 (70%)

0

Chemical space

Toxic drug

caffeine

3

2

0

7

2

-4

PC2 (4%)

Non-toxic drug

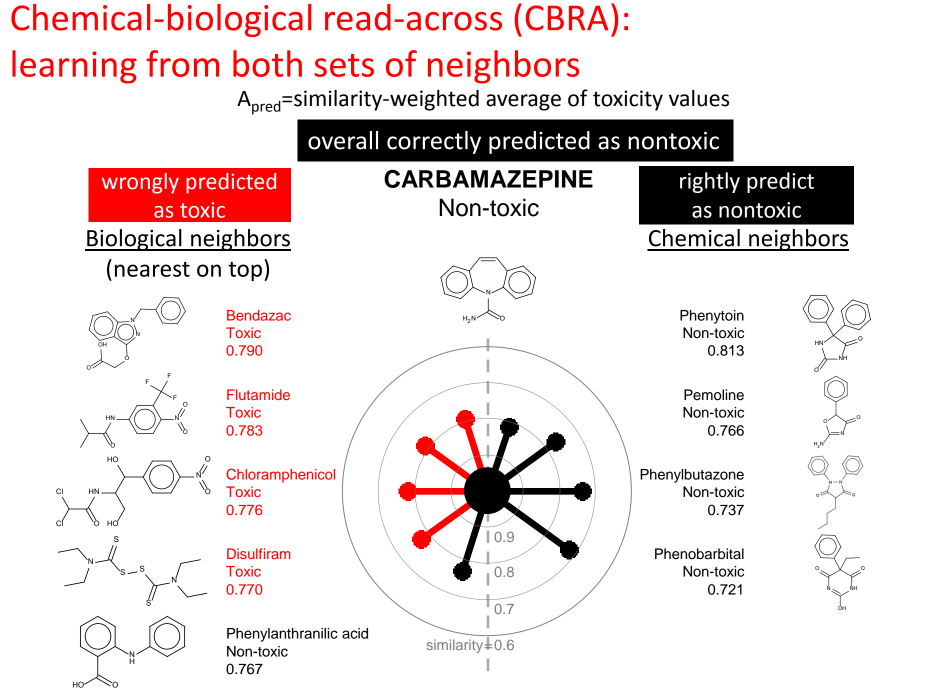
carbamazepir

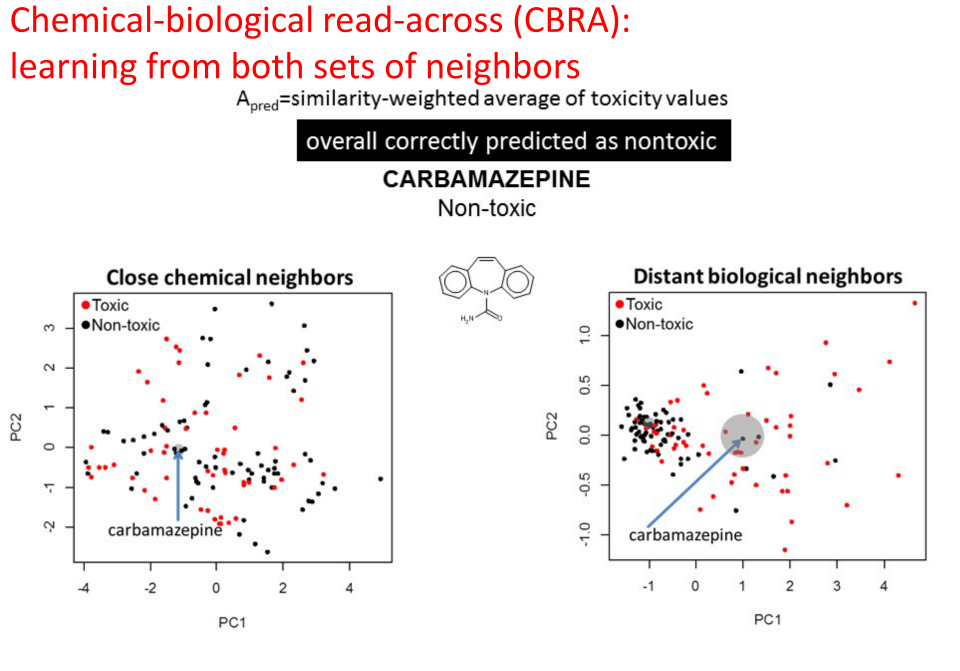
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<u>Improved</u>

2

- prediction:
- Learn from both sets of neighbors



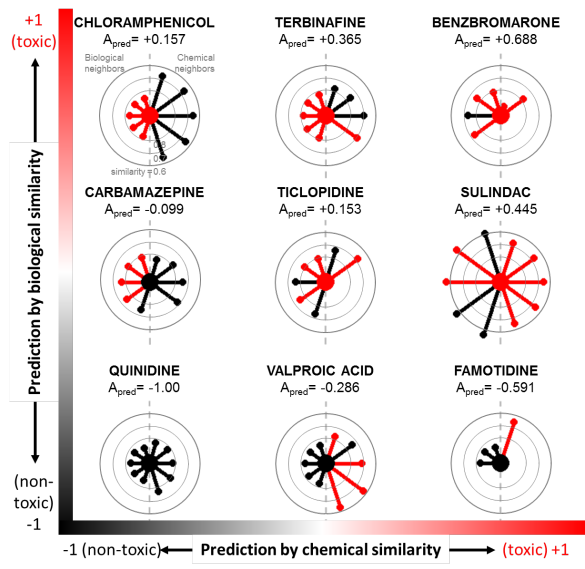


CBRA outperforms other models

| Model | Specificity | Sensitivity | Balanced accuracy (CCR) |
|---------------------------------|----------------------------|-----------------------------------|--|
| Chemical read-across | 0.73 ± 0.07 | 0.34 ± 0.05 | 0.53 ± 0.04 |
| Biological read-across | 0.85 ± 0.07 | 0.66 ± 0.04 | 0.76 ± 0.04 |
| Hybrid read-across | 0.85 ± 0.07 | 0.58 ± 0.04 | 0.72 ± 0.04 |
| Multi- space read- across | 0.89 ± 0.07 Results | 0.66 ± 0.04 of 5-fold externa | 0.78 ± 0.04 al cross-validation |

- Single space approaches replicated previous results: TGx > hybrid > QSAR
- Multi-space kNN read-across, using both chemical and toxicogenomic neighbors, had the highest predictive power

Radial Plots Visualize both Chemical and Biological Similarity to Help Forming the Read-across Argument



Conclusions and Outlook

- Rapid accumulation of large biomolecular datasets (especially, in public domain):
 - Strong need for both chemical and biological data curation
 - Cheminformatics approaches support <u>biological</u> data curation
- Novel approaches towards Integration of inherent chemical properties with <u>short term</u> biological profiles (biological descriptors)
 - improve the outcome of structure in vitro in vivo extrapolation
- Interpretation of significant chemical and biological descriptors emerging from externally validated models
 - inform the selection or <u>design</u> of effective and safe chemicals and focus the selection of assays/interpretation in terms of MoA
- Tool and data sharing
 - Pubic web portals (e.g., Chembench, OCHEM)

Alexander Tropsha

Principal Investigator Postdoctoral Fellows

Olexander Isayev, **Regina Politi**

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Research Professors Alexander Golbraikh, Denis Fourches (now at NCSU),

Eugene Muratov

Adjunct Members

Weifan Zheng, Shubin Liu

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- R01-GM068665
- NSF

- ABI 9179-1165

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- RD834999

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