

ChemTunes • ToxGPS

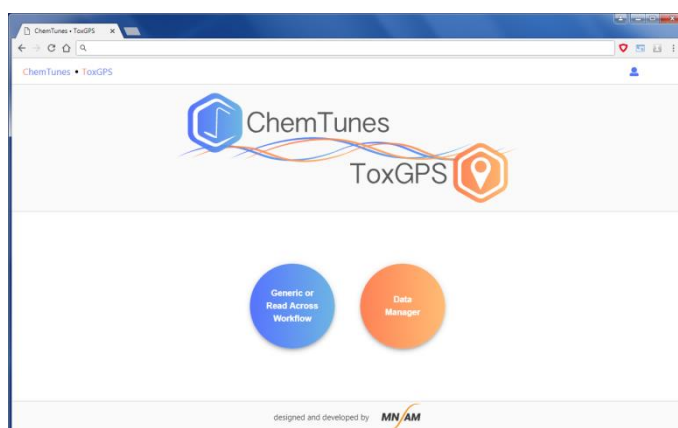
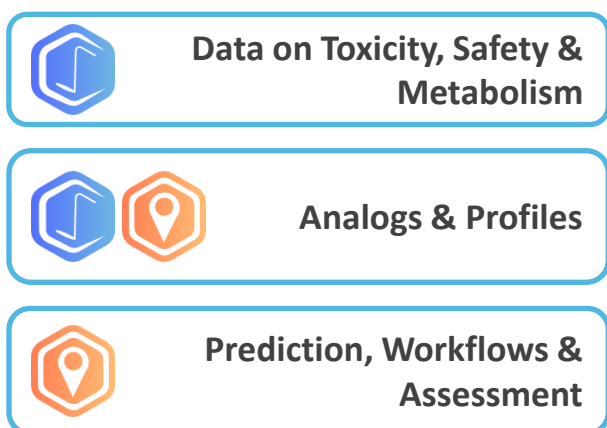
Database and Knowledgebase for
Safety Evaluation and Risk Assessment



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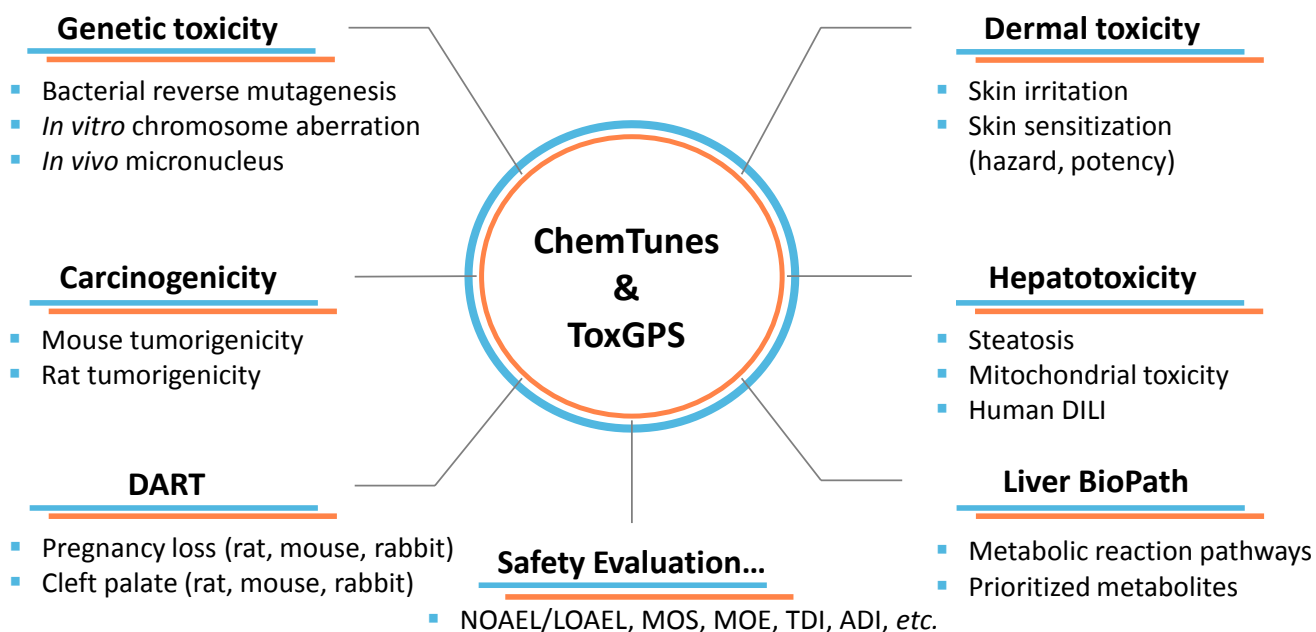
ChemTunes

- Chemoinformatics platform providing toxicity, safety evaluation, and metabolism information
- Nearly 100,000 compounds over 30,000 studies covering > 70 endpoints



Human Health Endpoints

- Wide variety of endpoints important for safety assessment related to human health
- Data from toxicity and safety assessment databases, as well as *in silico* predictions



[1] Rathman JF, *et al.* Uncertainty Estimation and Quantitative Combination-of-Evidence using Dempster-Shafer Theory. *Comp. Tox.* **2018**, 6, 16-31

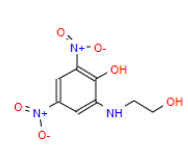
[2] Cherkasov A, *et al.* QSAR modeling: where have you been? Where are you going to? *J. Med. Chem.* **2014**, 57(12), 4977-5010

ToxGPS – Prediction






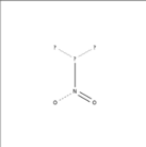
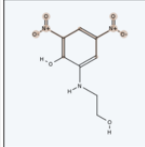
- Knowledgebase for compound location, MoA QSAR & rule-based toxicity prediction and metabolism
- In silico* safety assessment workflows including Read-Across and Threshold of Toxicological Concern

Prediction for Bacterial Reverse Mutagenesis

Prediction Results for compound #1



In database: Yes
 CHEMTUNES ID: CMS-62110
 Name: 2-HYDROXYETHYL PICRAMIC ACID
 Registry number(s): 99610-72-7
 # studies in CHEMTUNES: 9

Overall prediction	Positive probability	
	0.99	
QSAR Model	Positive probability	
Global	0.58-0.85	
MoA Models		
Aromatic amine	0.39-0.65	
Aromatic nitro	0.38-0.61	
Phenol	0.54-0.84	
Chemotype Alert	Chemotype Alert Match	Alert name Odds ratio
		Aromatic nitro 6.02

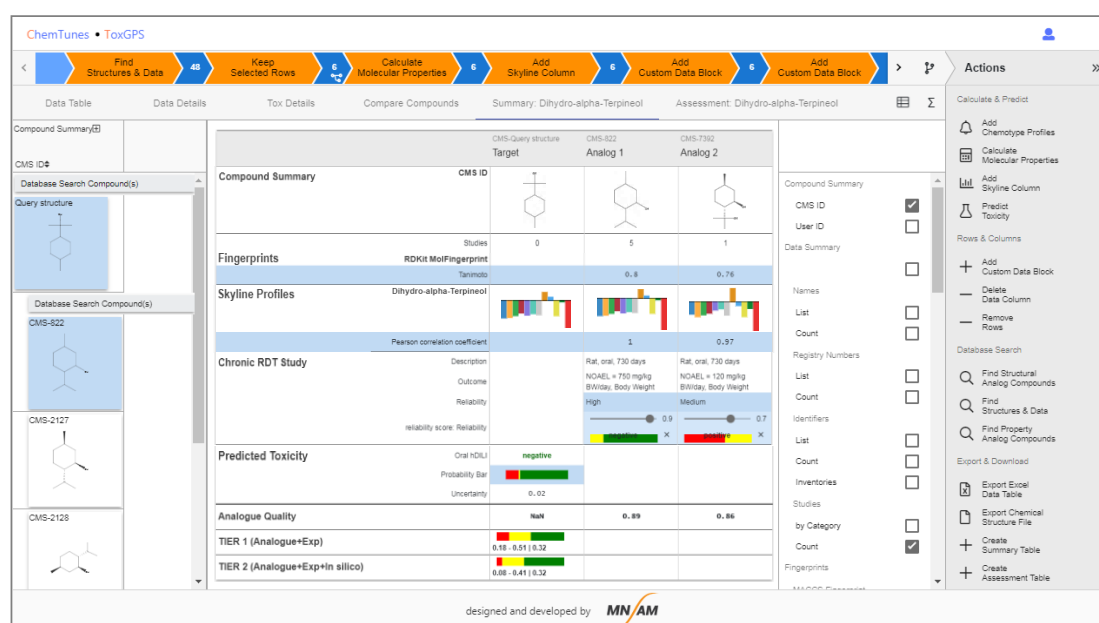
Combined outcome

QSAR (global and MoA) as well as rule-based results are combined into one final outcome by a rigorous decision theory approach¹

ToxGPS – Read-Across

- Chemical speciation – metabolites and tautomers
- Generate analogues (structure, properties, biological)
- Line up evidence
- Combine evidence for outcome and uncertainty estimation










Relevant
Reliable
Consistent
Reproducible



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Workflow: Find Structures & Data (48) → Keep Selected Rows (6) → Calculate Molecular Properties (6) → Add Skyline Column (6) → Add Custom Data Block (6) → Add Custom Data Block (6)

Summary: Dihydro-alpha-Terpineol Assessment: Dihydro-alpha-Terpineol

Compound Summary	CMS ID	CMS-Query structure	CMS-822	CMS-730
Target				
Studies	0	5	1	
Fingerprints	RDKit MolFingerprint		0.8	0.76
	Tanimoto			
Skyline Profiles	Dihydro-alpha-Terpineol			
Pearson correlation coefficient		1	0.97	
Chronic RDT Study	Description	Rat, oral, 730 days	Rat, oral, 730 days	
	Outcome	NOAEL = 750 mg/kg B/Wday, Body Weight	NOAEL = 120 mg/kg B/Wday, Body Weight	
	Reliability	High	Medium	
	reliability score: Reliability	0.9	0.7	
Predicted Toxicity	Oral NDILI	negative		
	Probability Bar			
	Uncertainty	0.62		
Analogue Quality		NaN	0.89	0.86
TIER 1 (Analogue+Exp)				
		0.18 - 0.51 0.32		
TIER 2 (Analogue+Exp+in silico)				
		0.08 - 0.41 0.32		

designed and developed by **MN/AM**

[3] Leist M, Yang C, *et al.* Novel Technologies and an Overall Strategy to Allow Hazard Assessment and Risk Prediction of Chemicals, Cosmetics, and Drugs with Animal-Free Methods. *Altex* **2012**, *29*, 4/12, 373-388

[4] Yang C, *et al.* Computational Toxicology Approaches at the US Food and Drug Administration. *ATLA* **2009**, *37*, 523-531

Marquee Features



- Mechanism-based (MoA) QSAR models
- Rule-based and endpoint specific alerts^{2,3}



- Proven approach used at US FDA for risk assessment^{3,4}



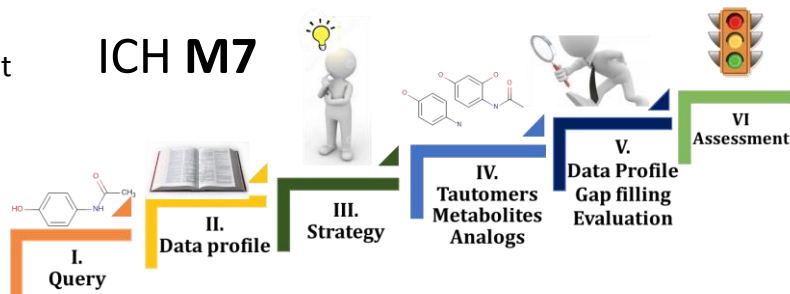
- Data- and knowledge-based system
- Established chemoinformatics methods



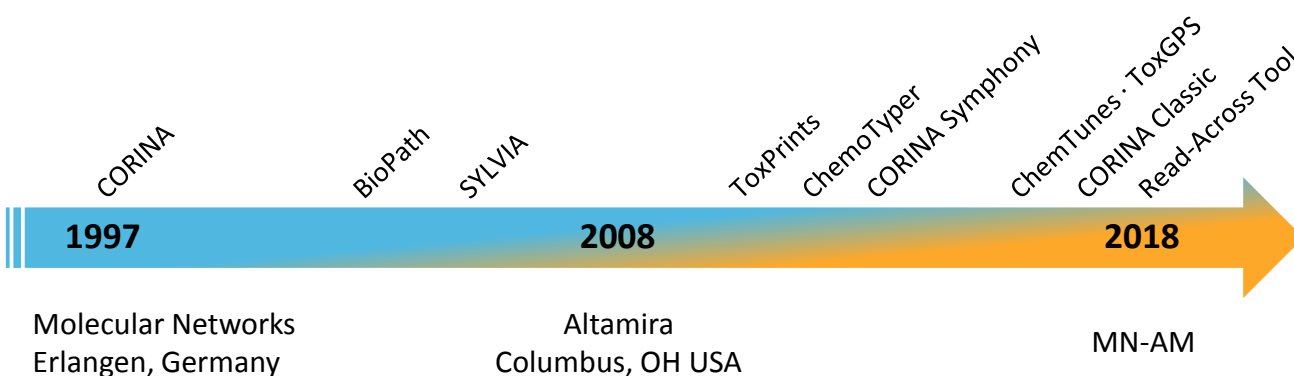
- Decision theory approach¹
- Weight-of-evidence (WoE) outcome and uncertainty¹

Applications

- Compound location service
- Platform for *in silico* safety assessment
- Read-Across workflow
- ICH M7 genotoxic impurities in drug products



MN-AM at a Glance



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