

Hazard Using New Approach Methodologies

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Precision Toxicology

- Aims at better protecting the health of people and the environment by establishing **new approach methodologies (NAMs)** for chemical safety testing
- Establishes a new regulatory paradigm of detecting toxicity using molecular biology for greater certainty at predicting which chemicals cause harm while avoiding traditional animal testing
- Accomplish this goal by identifying **molecular key** event (KE) biomarkers, predictive of chemically induced adverse health effects in humans, that feed directly into regulatory and industry practice

Participants of *PrecisionTox*

15 participating organizations across 8 countries



Distribution of participants



Molecular Key Event Biomarkers



- mKE biomarkers are quantifiable molecular indicator of a toxicological response predictive of the adverse outcome induced by exposure to a chemical substance
- Discovered via *in-silico* molecular biomarkers defined as a sparse network of interacting genes and their metabolic products





Linking Human Toxicology and Ecotoxicology

- **Challenge**: whole-organism testing is crucial, but no single model is a perfect human surrogate.
- **Solution**: evolutionarily diverse non-sentient organisms plus human cell line (PhyloToxicology)
 - Evolutionary origins of the interactome including toxicology-relevant networks
 - Induced by chemicals (chemical responsiveness)
 - Indicative of similar adversity (hazard relatedness)
 - Shared by multiple species (evolutionarily conserveness)
- Identification of molecular biomarkers of a chemical hazard

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Replacing Mammals by NSO Species

- Systematic use of distantly related non-sentient organism (NSO) species from across the tree of life
 - Daphnia magna (water flea)
 - Drosophila melanogaster (fruit fly)
 - Caenorhabditis elegans (nematode)
 - *Danio rerio* (zebra fish; embryo)
 - Xenopus laevis (frog; embryo)
 - Homo sapiens (human; cell-line)
- 250 21,00chemicals from various chemical classes, producing 0 samples for transcriptomics and metabolomics
- NAMs for chemical toxicity testing: combination of machine learning paradigms for multi-omics integration and mKE biomarkers identification





Step 1: Multi-Omics Molecular Data Production



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Step 2: Biomolecular Network Construction



Step 3: Association Quantification

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Step 4: Molecular Biomarker Identification



Mechanisms

Adversity

Predictability



Ortholog-based Annotation Explainability

Prediction Models

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Pathway Conservation

Probabilistic Learning





Pre-pilot Case Study



PrecisionTox & qAOP

- PrecisionTox uses a systems biology strategy
 - Collects large sets of untargeted multi-omics data
 - Data-driven modelling using combination of machine learning paradigms
 - Assesses the quality of findings with validation experiments
 - Discovers both known unknowns and unknown unknowns
- Efforts in qAOP
 - **Central Database** provides chemical information, known molecular biomarkers and pathways, and their related AOP information
 - **Computational Workflow** for quantifying the molecular KE biomarker associations
- Missing components
 - (Un)certainty measurement
 - From correlation to causality



Measurement of (Un)certainty

- Probabilistic learning
 - Classical machine learning lacks the measurement to account for model (un)certainty and prediction confidence
 - Probabilistic learning provides the opportunity to incorporate prior knowledge and hypotheses
- Explainable machine learning
 - In contrast to the conventional "black box" models, the behaviour and predictions made by the explainable models are understandable to humans
 - Explainability provides inspectable predictions and insights of the underlying mechanisms

13

Causal Inference

- Probabilistic graphical model combines probabilistic learning, explainable ML, and causal inference
- Sparse genetic regulatory network (GRN) inference using maximum-entropy probability model and multi-objective memetic algorithm (MMA-MEPM)



MEPM Method

find $\rho(x^t)$ that maximizing the Shannon entropy

$$S=-\sum_{t=1}^{T}
ho\left(x^{t}
ight){
m ln}\,
ho\left(x^{t}
ight)$$

subject to normalization,



 $ho(x^t) = rac{1}{Z} \mathrm{e}^{-H}$

 $H_t\!=rac{1}{2}\Sigma_{ij}x_i^tM_{ij}x_j^t$

energy function

Inferred information

 $M_{ij} \approx$ the pairwise gene interactions between gene *i* and gene *j*

$$M = C^{-1}$$

C = covariance matrix of gene expression matrix



pairwise gene interactions network



Fu, Y. (2022). Inferring sparse genetic regulatory networks based on maximum-entropy probability model and multi-objective memetic algorithm.



Causal Inference

• Modular meta-learning is particularly of interest due to the limited experimental data size and modular structure in biological networks



Lecca, P. (2021). Machine learning for causal inference in biological networks: Perspectives of this challenge. Frontiers in Bioinformatics, 1, 746712.





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https://precisiontox.org

Thank You