



Analysis of High-Throughput Profiling Assays in Chemical Safety Screening

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Outline

- Introduce Broad/High-Throughput Profiling Methods for Toxicology
- Overview of Transcriptomics
- Dose-Response Modeling of Transcriptomic Data
- Connectivity Mapping for Mechanistic Inference
- Cell Painting / High-Content Imaging



High-Throughput Profiling Methods

Broad-Coverage, Non-Targeted Assays



Tiered Chemical Safety Testing Strategy

Tier 1: Broad coverage, high content assays

- Must be cost-effective enough to rapidly screen 1000s of chemicals
 - e.g. Transcriptomics and/or cell imaging applied *in vitro*
 - Acute exposure: 6 - 24 hours
 - Multiple cell types with different metabolic profiles
- Goals: Prioritize chemicals by bioactivity & potency for further testing

Tier 2: Targeted *in vitro* assays

- Goals: confirm bioactivity & potency of chemicals flagged for potential safety issues

Tier 3: Organotypic assays, systems modeling, and more

- Goals: identify likely tissue, organ, or organism effect of chemical

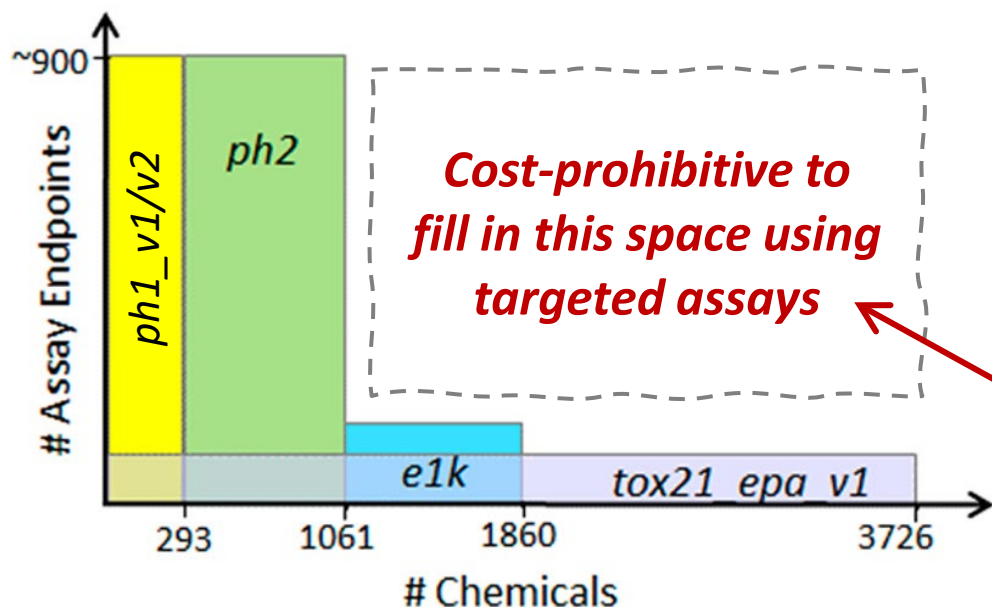
of Chemicals

Cost per Chemical



High-Throughput Screening (ToxCast)

Testing Phase	Chemical Set	Unique Chemicals	Assay Endpoints
ToxCast Phase I	ph1_v1	310	~700
ToxCast Phase II	ph1_v2	293	~200
	ph2	768	~900
	e1k	799	~50
Tox21	tox21_epa_v1	3726	~80



Richard et al. (2016)

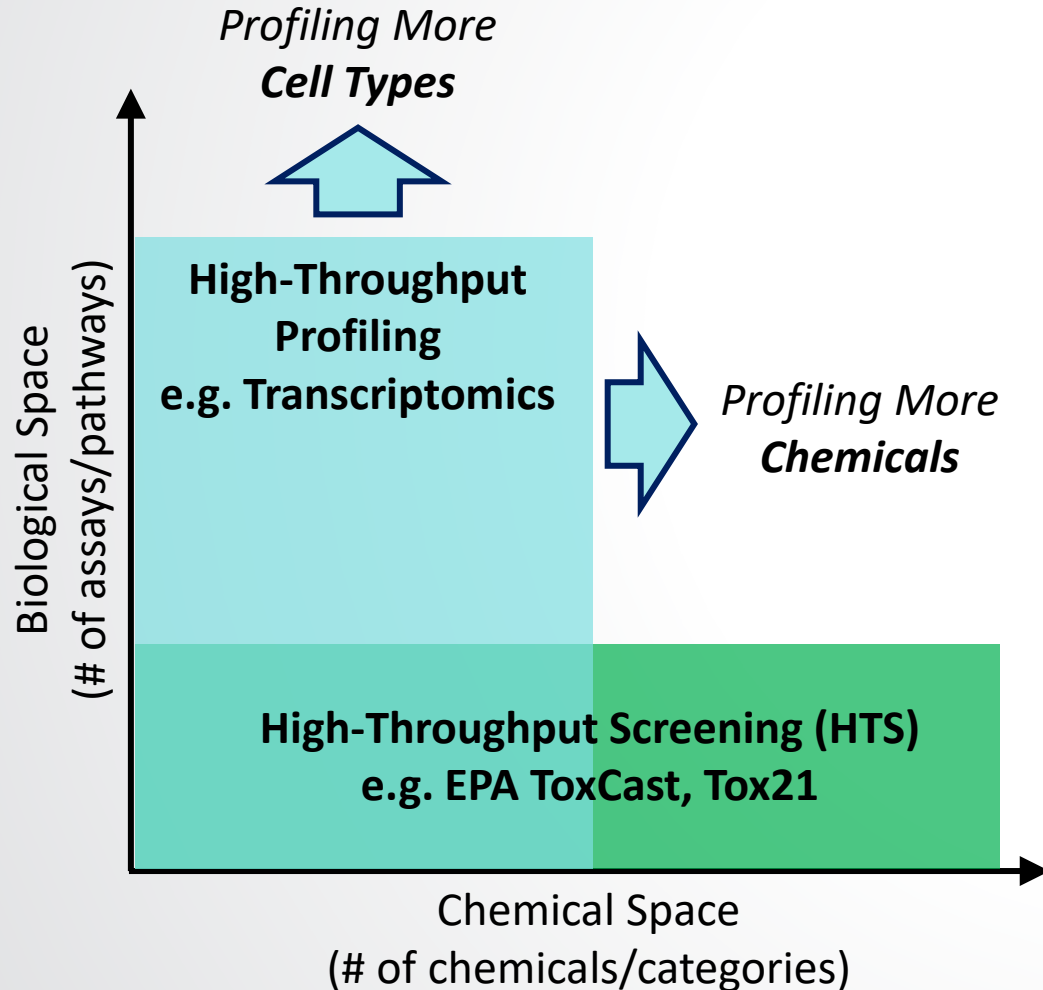
ToxCast: EPA-led effort using high-throughput screening (HTS) assays to assess bioactivity and potential toxic effects.

- Expose living cells or isolated proteins *in vitro* to pure chemicals in vehicle/culture media
- Maintain standard library of 1,000s of diverse chemicals
- Mostly targeted assays (*chemical X* → *target Y*) leading to **incomplete coverage of human biological space**
- See: Richard, et al. (2016)
DOI: [10.1021/acs.chemrestox6b00135](https://doi.org/10.1021/acs.chemrestox6b00135)

New Strategy for Hazard Evaluation: Improve efficiency and increase biological coverage by using broad-based (i.e. non-targeted) assays that capture many potential molecular and phenotypic responses of human cells to chemical exposure.



High-Throughput Profiling Methods



Definition: Any method that broadly profiles a range of biological pathways, functions, or features, as opposed to targeted testing of a single endpoint, including:

- **Transcriptomics** – profiling thousands of gene mRNA levels
- **Cell Painting** – profiling hundreds of cellular phenotypic features using microscopy
- *Broad batteries of targeted assays*
- *Other ‘Omics – e.g. metabolomics, proteomics*

Many Analysis Choices!



No single “best” method for analyzing high-throughput profiling data

- Are you interested in mechanism, or just want a threshold for general bioactivity?
- Is it more important to be **predictive** or **protective** of hazard level *in vivo*?
- What other data is available for the same/analogous chemicals?
- Different technologies require different statistical models, quality control, etc.
- Experimental design (*# of replicates, doses, etc.*) impacts analysis choices!





Overview of Transcriptomics

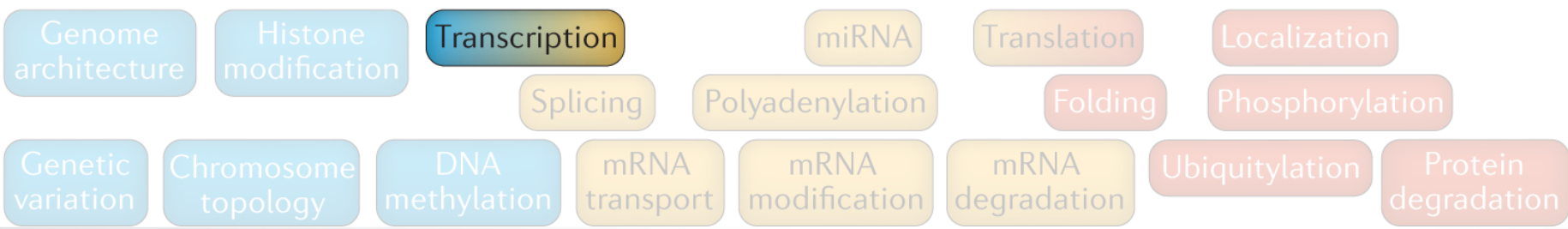
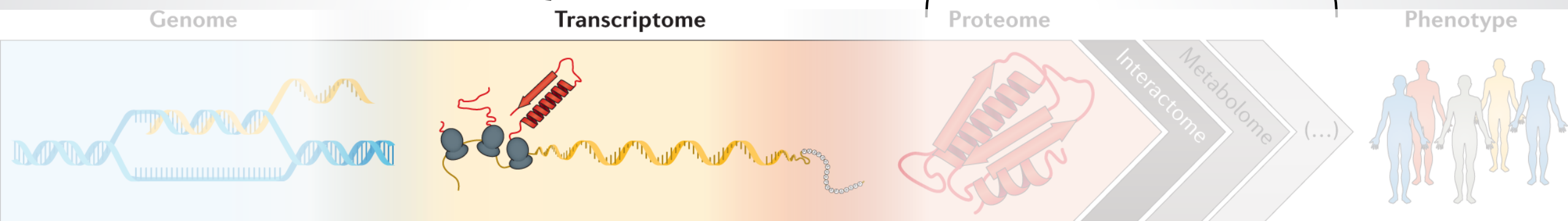
Profiling Genome-Wide Gene Expression



Why Transcriptomics?

Highly dynamic in response to environmental stimuli,
Well-established, cost-effective, high-throughput methods

Broad profiling methods are newer
or less cost-effective



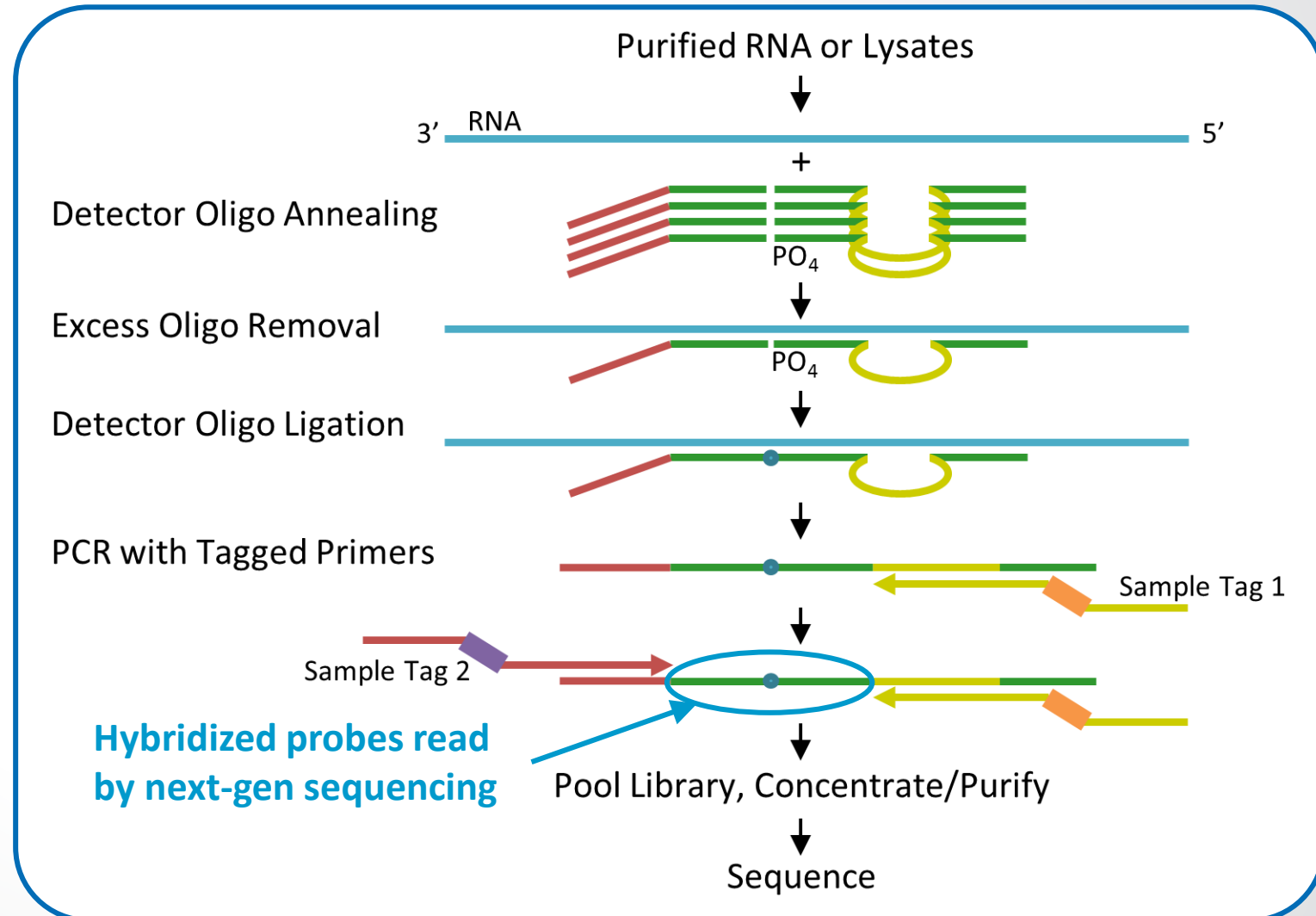
Hazardous exposures can cause perturbations
at multiple levels of gene regulation





Targeted RNA-seq Assay (TempO-seq)

- Next-Gen sequencing of targeted probes hybridized to expressed transcripts
- Scalable: can measure a few thousand genes (S1500+) or up to whole transcriptome
- Captures gene expression at lower cost than RNA-seq or microarrays
- Compatible with raw cell lysates – *ideal for large-scale screening*





Transcriptomics Knowledgebases

Signature Databases – any collection that links sets of genes to specific biological categories or patterns (e.g. functions, pathways, responses):

- Gene Ontology (geneontology.org) - *Nucleic Acids Res* (2021) DOI:[10.1093/nar/gkaa1113](https://doi.org/10.1093/nar/gkaa1113)
- MSigDB (gsea-msigdb.org) - *Bioinformatics* (2011) DOI:[10.1093/bioinformatics/btr260](https://doi.org/10.1093/bioinformatics/btr260)
- Reactome (reactome.org) - *Nucleic Acids Res* (2022) DOI:[10.1093/nar/gkab1028](https://doi.org/10.1093/nar/gkab1028)

Databases of Toxicogenomic/Transcriptomic Profiles:

- TG-GATES (biosciencedb.jp) - *Nucleic Acids Res* (2015) DOI:[10.1093/nar/gku955](https://doi.org/10.1093/nar/gku955)
- Connectivity Map (clue.io) – *Cell* (2018) DOI:[10.1016/j.cell.2017.10.049](https://doi.org/10.1016/j.cell.2017.10.049)

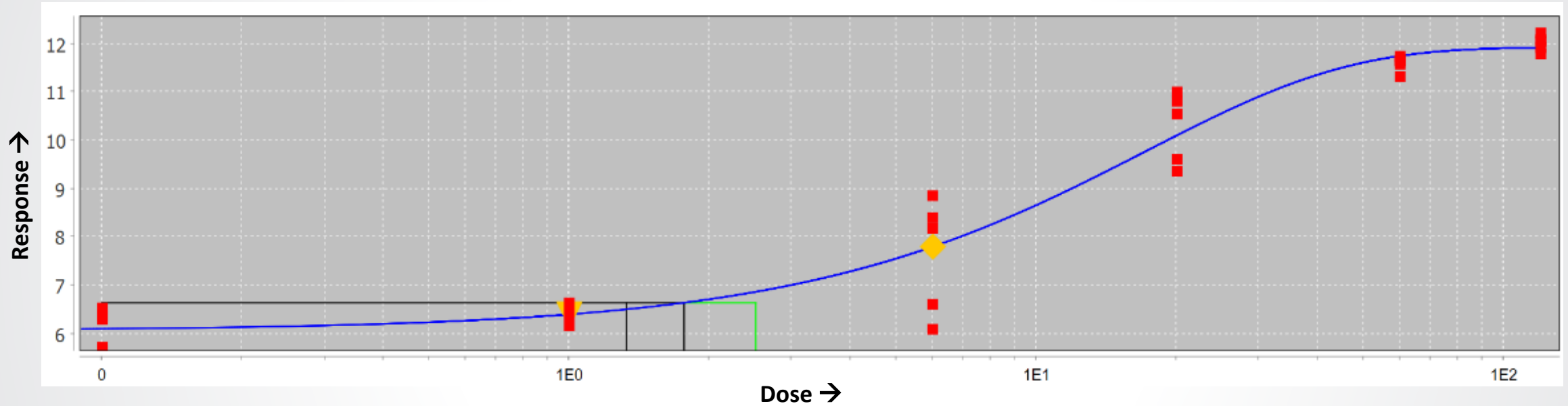
General Transcriptomic DBs: [Gene Expression Omnibus](#) (NCBI) [ArrayExpress](#) (EMBL)



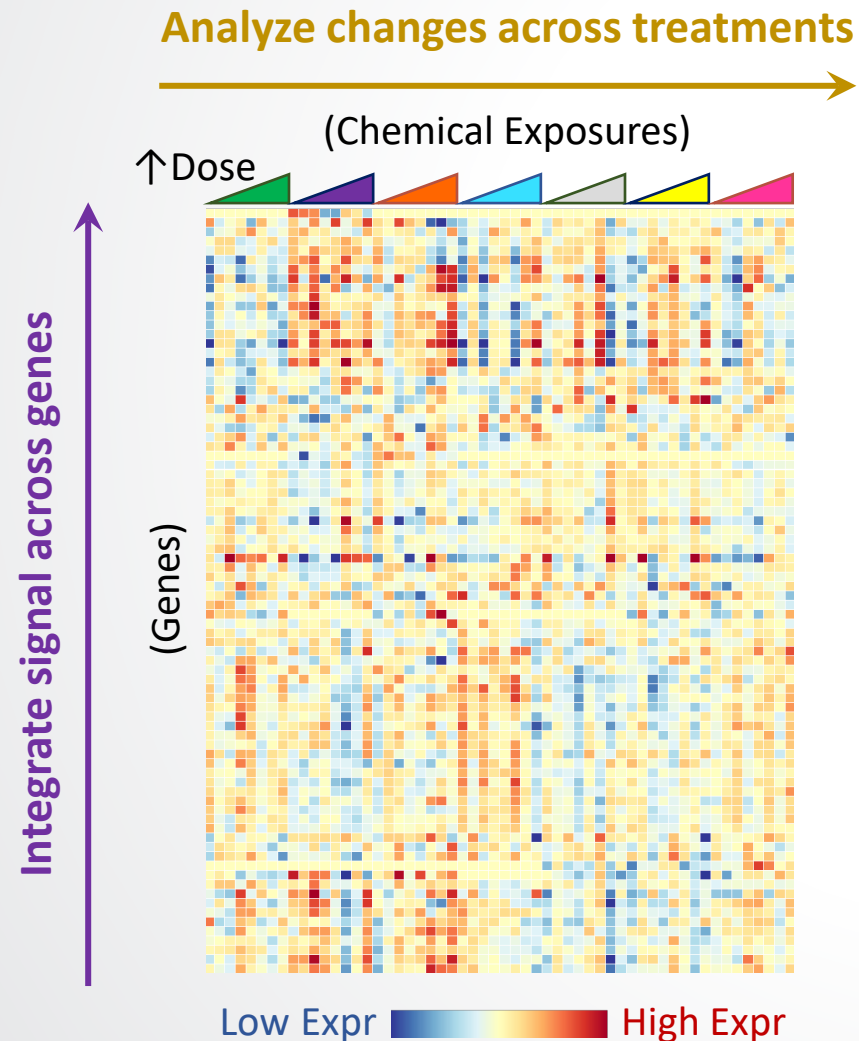
Dose-Response Modeling of Transcriptomic Data



Dose-Response Models



- Commonly used for apical endpoint and targeted assay data
- Benchmark Dose Software (BMDS): www.epa.gov/bmds
- ToxCast Pipeline (tcpl): cran.r-project.org/package=tcpl

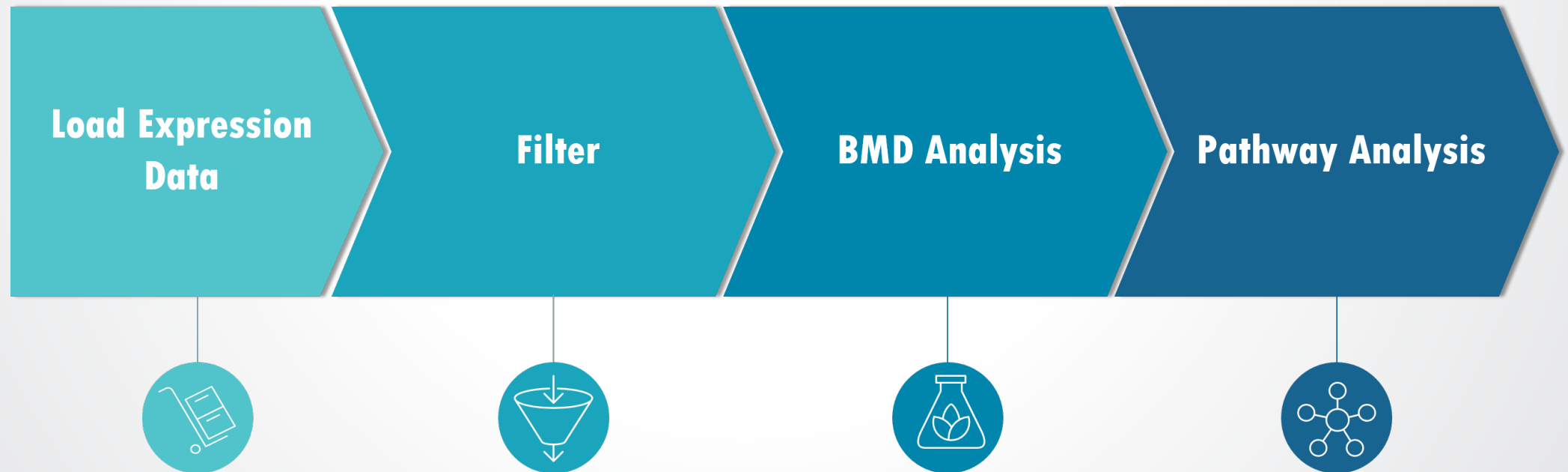


- Different genes may respond at different doses of a given exposure!
- Need to analyze both:
 - Dose-responsive trends
 - Coordinated changes in gene expression
- Gene-level data noisier in transcriptomics than targeted measurements (e.g. RT-qPCR)
- Dose-response modeling thousands of features (e.g. mRNA levels) leads to computational & statistical challenges



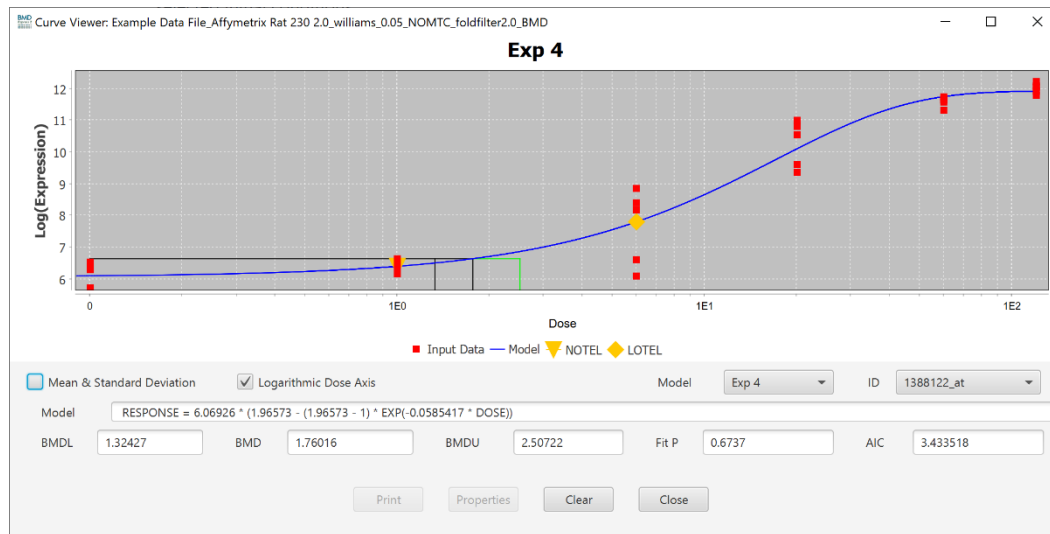
BMDExpress Software

- **Benchmark Dose (BMD):** Lowest dose/conc when an effect exceeds the background response rate
- BMDExpress automates fitting & summarizing multiple models (BMDS software) on many genes
- More information: Phillips, et al. *Bioinformatics* 2019 DOI: [10.1093/bioinformatics/bty878](https://doi.org/10.1093/bioinformatics/bty878)



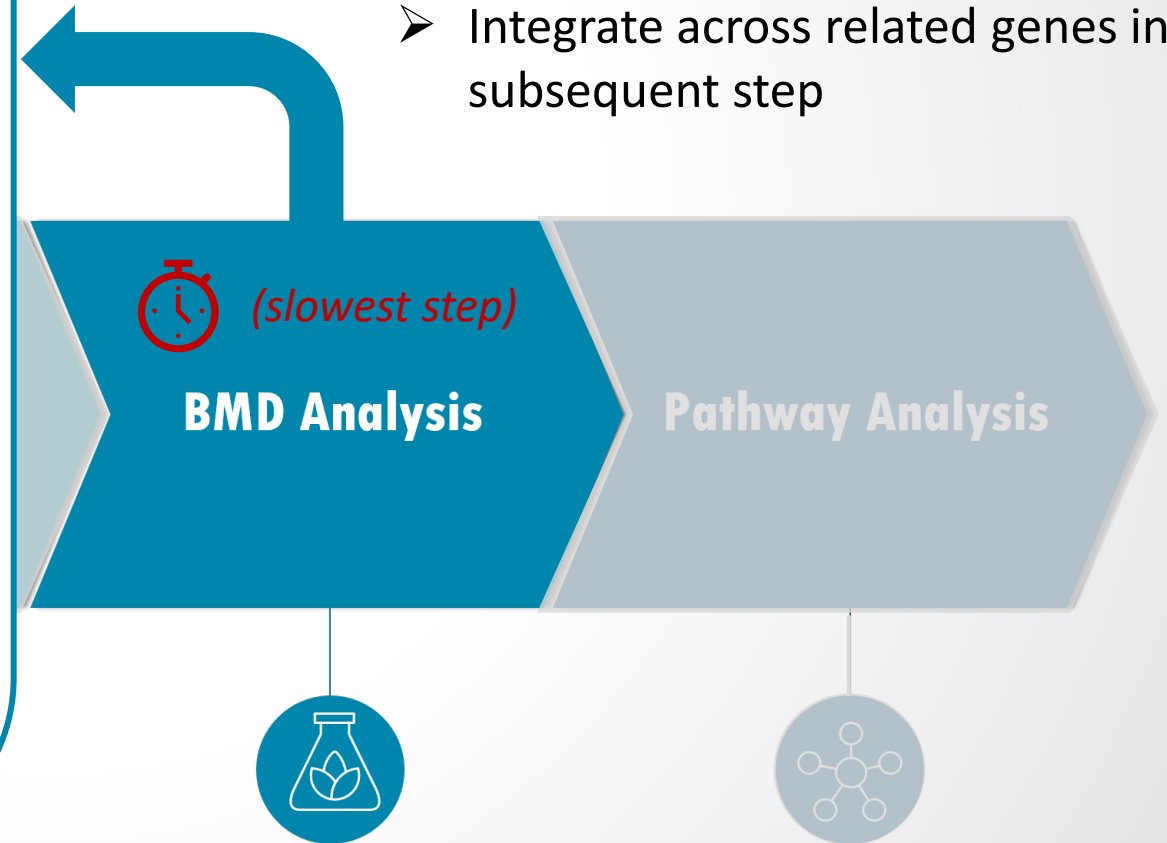
Run independently for each probe/gene:

- Fit multiple curve shapes to data
- Select best-fit model



Gene-level BMD typically computed using **BMR_f = 1** or **1.349**, many tunable parameters

- Perform dose-response analysis on individual probes/genes
- Integrate across related genes in subsequent step





BMDExpress Software

Summarize dose-response models for **biologically related** sets of gene

- Identify pathways/gene sets with multiple dose-responsive genes
- Category-level POD = median of active gene-level PODs

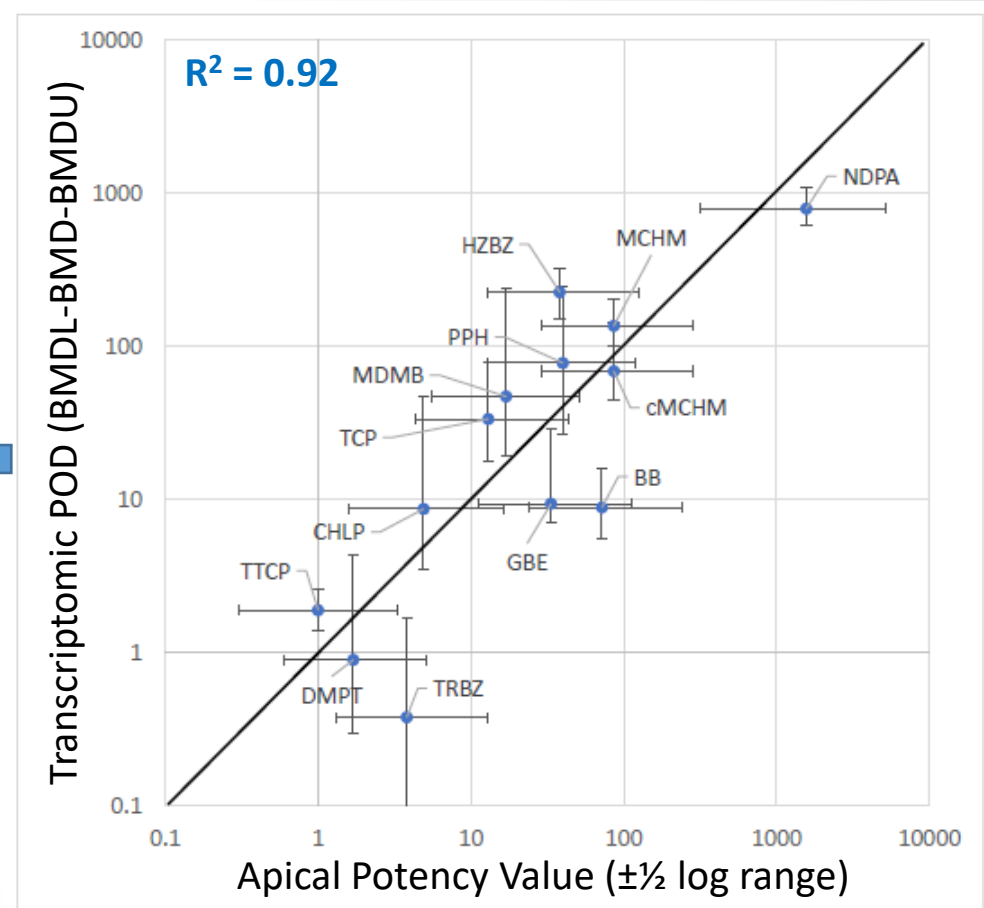




Overall Transcriptomic POD (tPOD)

Common methods for deriving overall tPOD:

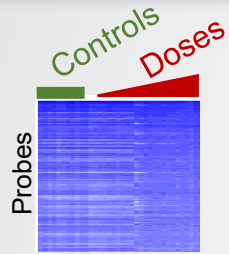
Gene-based Methods	
Nth Percentile (e.g. 5 th %ile) BMD	Reardon, et al. Tox Sci 2021
Nth Lowest (e.g. 25 th) BMD	
Pathway/Category-based Methods	
Lowest Active Pathway BMD	Gwinn, et al. Tox Sci 2020
5 th Percentile Pathway BMD	Harrill, et al. Tox Sci 2021
Global Methods:	
Distance-based POD (e.g. Mahalanobis Distance)	Nyffeler, et al. SLAS Discov 2021



Important to benchmark methods for the intended purpose, e.g. prediction of PODs from animal studies



Dose-Response Modeling of Gene Sets/Signatures



Concentration Series of Whole Transcriptome Profiles + Vehicle Controls

Love et al. (2014)

DESeq2 Moderated Fold-Changes

Barbie et al. (2009)

Gene Set Enrichment Analysis

Sheffield et al. (2022)

Benchmark Dose Modeling

Curated Signature Collection

Catalog of gene set signatures with toxicological relevance, annotated for known molecular targets

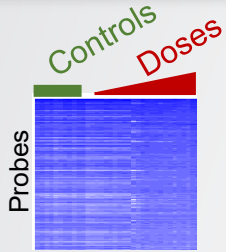
- **Bioplanet** (*Huang, et al. Front Pharmacol 2019*)
- **CMap** (*Subramanian, et al. Cell 2017*)
- **DisGeNET** (*Piñero, et al. Database 2015*)
- **MSigDB** (*Liberzon, et al. Cell Syst 2015*)

Open Source: github.com/USEPA/CompTox-httrpathway

- EPA/CCTE method for summarizing large-scale transcriptomic screening studies
- Integrates signal across known gene set (a.k.a. signature) **before** dose-response modeling



Dose-Response Modeling of Gene Signature Scores



Concentration Series of Whole Transcriptome Profiles + Vehicle Controls

Love et al. (2014)

DESeq2 Moderated Fold-Changes

Curated Signature Collection

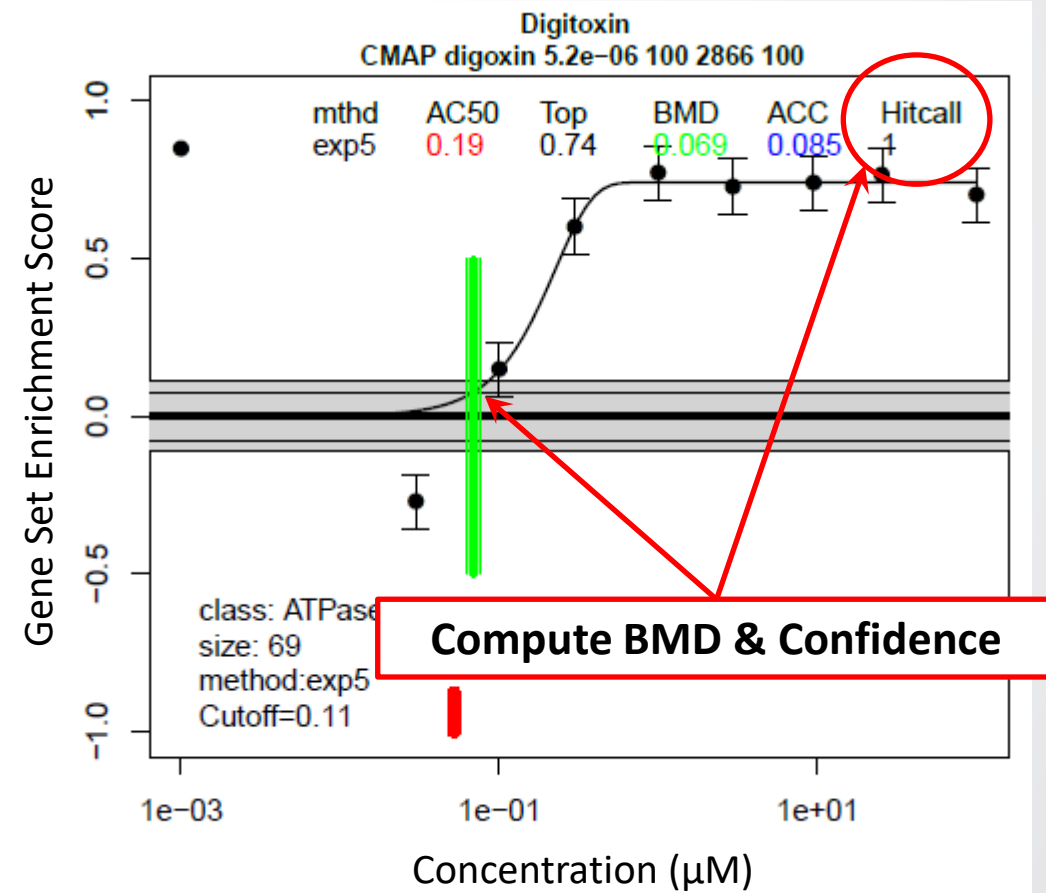
Barbie et al. (2009)

Gene Set Enrichment Analysis

Sheffield et al. (2022)

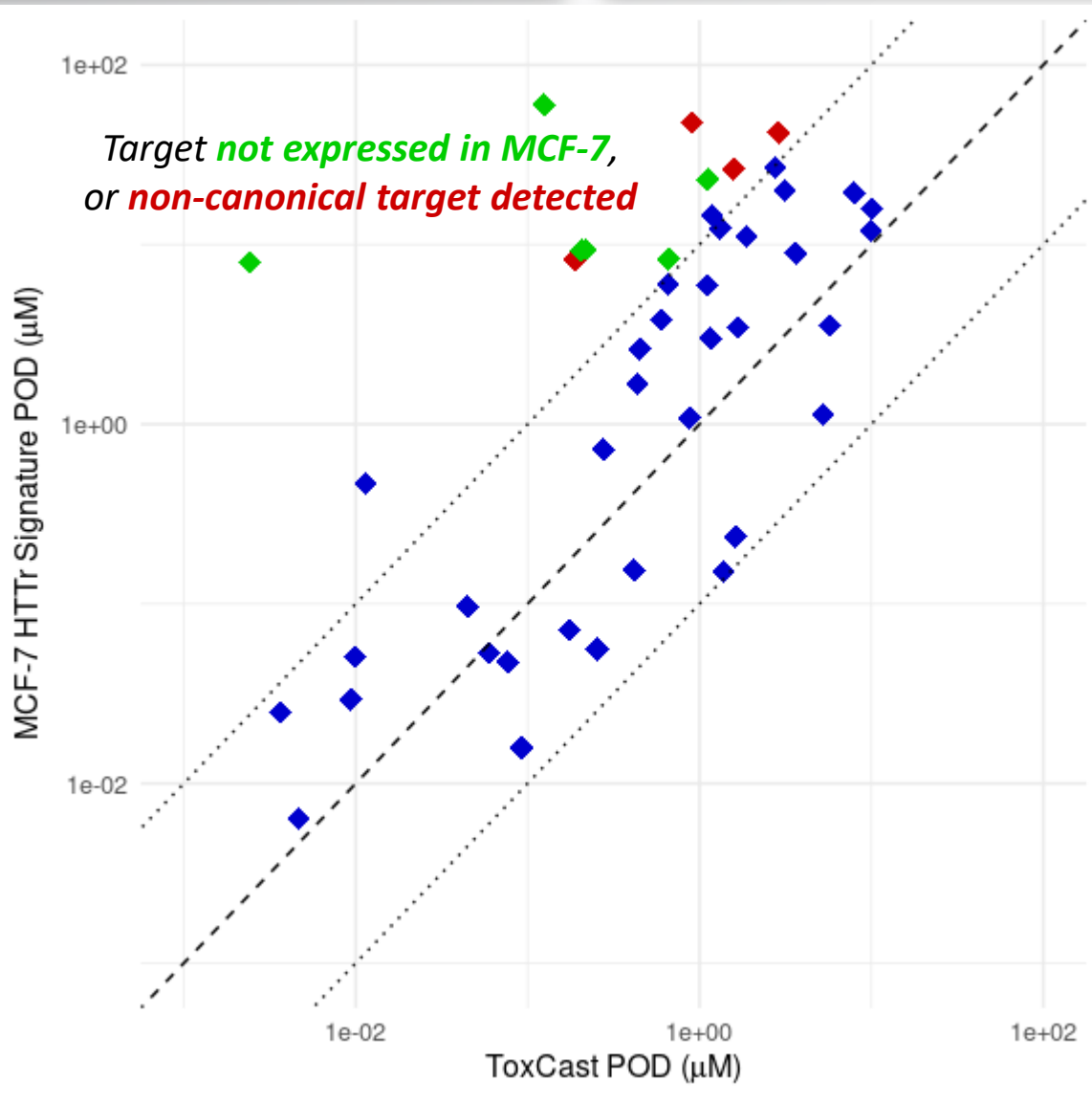
Benchmark Dose Modeling

Overall tPOD = 5th percentile of active signature BMDs
(Hitcall captures confidence that signature is truly dose-responsive)





tPODs Are Concordant With ToxCast



- Pilot study of 44 well-characterized chemicals in MCF-7 cells, 6h exposure
Harrill, et al. *Toxicol Sci* (2021)
DOI: [10.1093/toxsci/kfab009](https://doi.org/10.1093/toxsci/kfab009)
- Compared transcriptomic PODs to previous ToxCast targeted assay results
(multiple cell types, assays, and exposure lengths)
Paul-Friedman, et al. *Toxicol Sci* (2020)
DOI: [10.1093/toxsci/kfz201](https://doi.org/10.1093/toxsci/kfz201)
- Signature-based PODs are highly concordant with ToxCast results for the majority of test chemicals in pilot study



Alternate Dose-Response Modeling Methods

Many other analysis methods proposed, this is an active area of research!

Bayesian Methods:

- BIFROST – Reynolds, et al. *Comp Tox* (2020)
DOI: [10.1016/j.comtox.2020.100138](https://doi.org/10.1016/j.comtox.2020.100138)
- BBMD – Shao & Shapiro, *EHP* (2018) URL: benchmarkdose.com
- ToxicR – Wheeler, et al. *Environmetrics* (2022) CRAN: [ToxicR](https://cran.r-project.org/web/packages/ToxicR/index.html)

Integration across genes using latent variables:

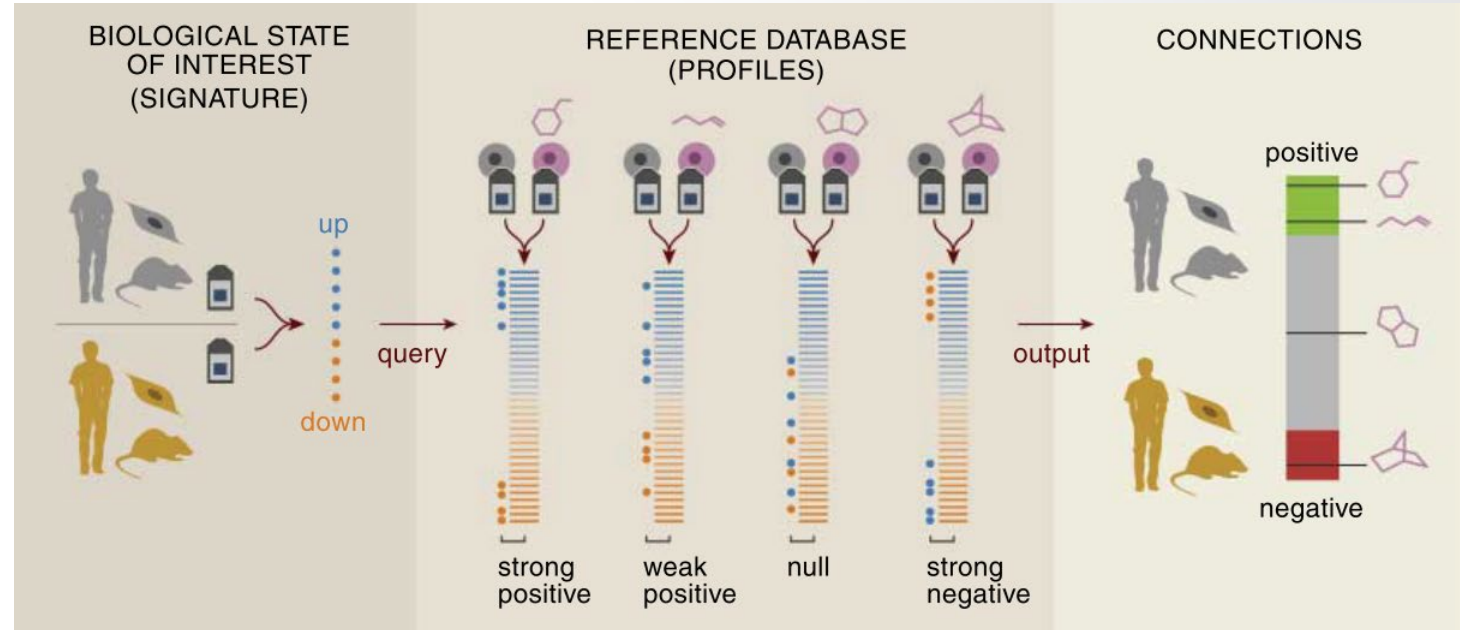
- Basili, et al. *Chem Res Tox* (2022) DOI: [10.1021/acs.chemrestox.1c00444](https://doi.org/10.1021/acs.chemrestox.1c00444)



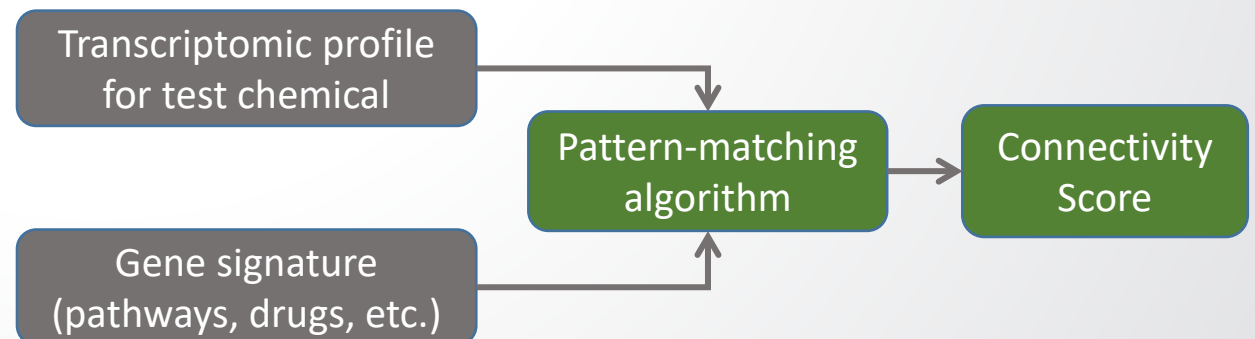
Connectivity Mapping

Inference of common mechanism/effects by transcriptomic similarity

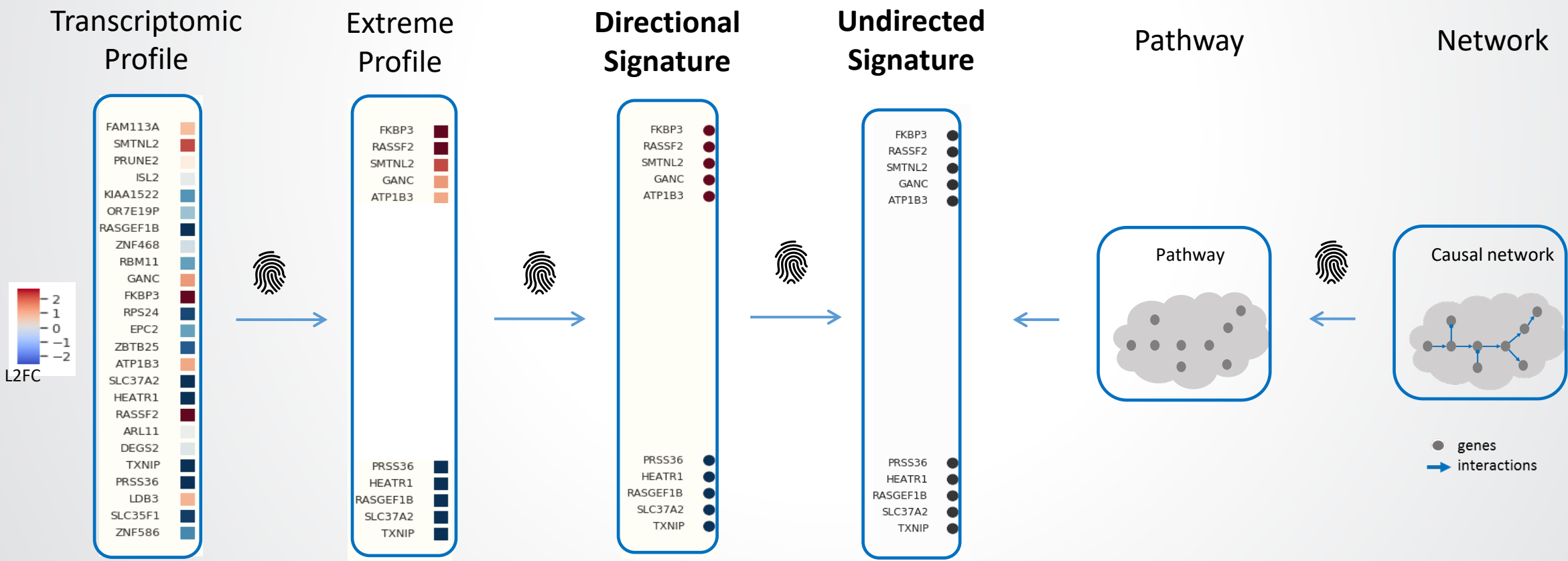
- Postulates that similarity between transcriptomic profiles reflects common biological state or mechanism
- “Fingerprint” transcriptomic profiles for similarity analysis
- Increasing utility in toxicology to infer mechanism by similarity
[DeAbrew et al., 2016](#)
[Wang et al., 2016](#)
- Web-based tool: <https://clue.io/>
 (Broad Institute)



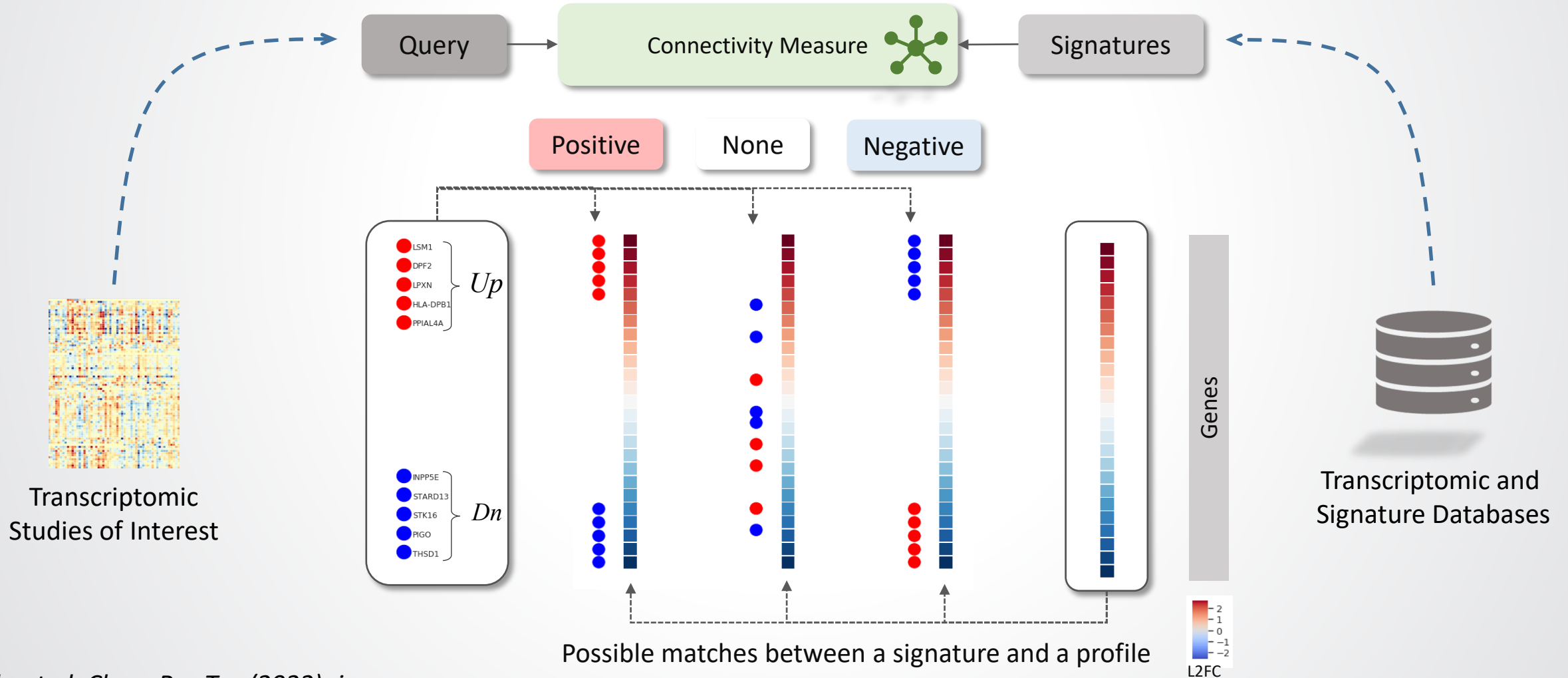
Lamb et al., Science 2006 DOI: [10.1126/science.1132939](https://doi.org/10.1126/science.1132939)



Signatures can be derived from specific gene expression profiles as well as conceptual models of biological pathways & networks, different representations are also possible.



Multiple connectivity "score" functions proposed





Applications of Connectivity Mapping

- **Group chemicals by Mode of Action (MOA)**, e.g.:
De Abrew, et al. *Tox Sci* (2016) DOI: [10.1093/toxsci/kfw058](https://doi.org/10.1093/toxsci/kfw058)
- **Predict mechanism** based on similarity to reference chemicals, e.g.:
Wang, et al. *BMD Genomics* (2016) DOI: [10.1186/s12864-016-2406-y](https://doi.org/10.1186/s12864-016-2406-y)
- Select chemicals for **Read Across analysis**, e.g.:
De Abrew, et al. *Toxicology* (2019) DOI: [10.1016/j.tox.2019.05.008](https://doi.org/10.1016/j.tox.2019.05.008)
GenRA: Helman, et al. *ALTEX* (2019) DOI: [10.14573/altex.1811292](https://doi.org/10.14573/altex.1811292)



Cell Painting

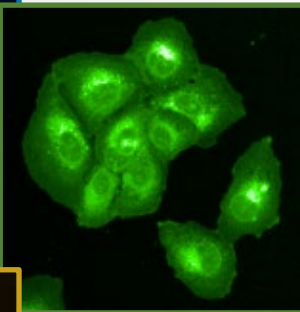
High-Throughput Phenotypic Profiling (HTPP) of cells using High-Content Imaging

Cell Painting with Multiple Markers

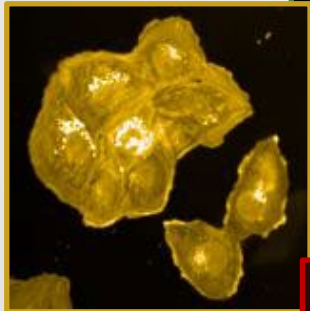
DNA



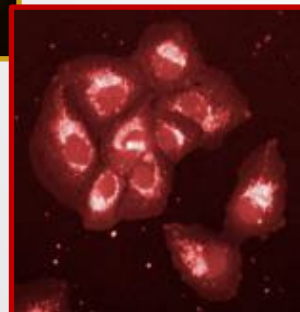
RNA + ER



Golgi + membrane
+ actin skeleton



Mitochondria



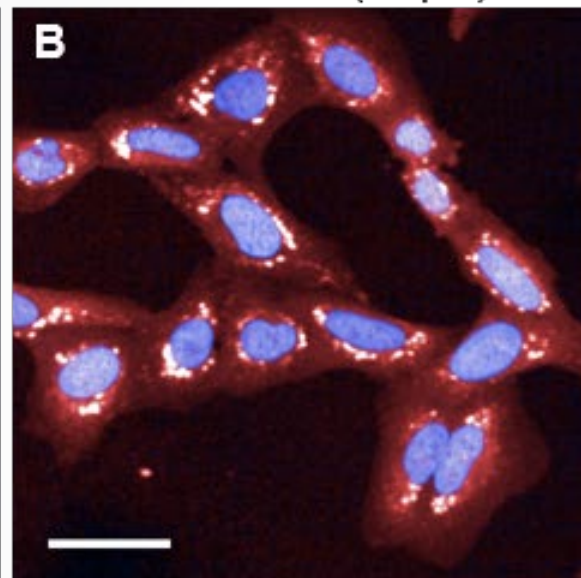
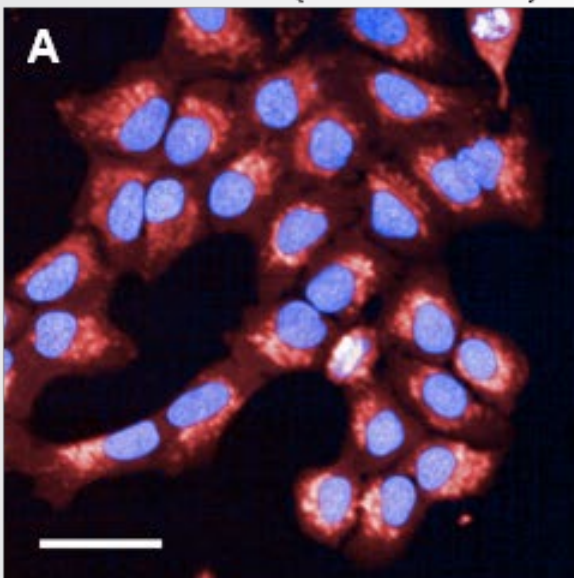
- Measures a large variety of phenotypic features in fluoroprobe labeled cells *in vitro*.
- Originally introduced in Bray, et al. *Nat Protoc* (2016) DOI: [10.1038/nprot.2016.105](https://doi.org/10.1038/nprot.2016.105)

Marker	Cellular Component	Labeling Chemistry
Hoechst 33342	Nucleus	Bisbenzamide probe that binds to dsDNA
Concanavalin A – AlexaFluor 488	Endoplasmic reticulum	Lectin that selectively binds to α -mannopyranosyl and α -glucopyranosyl residues enriched in rough endoplasmic reticulum
SYTO 14 nucleic acid stain	Nucleoli	Cyanine probe that binds to ssRNA
Wheat germ agglutinin (WGA) – AlexaFluor 555	Golgi Apparatus and Plasma Membrane	Lectin that selectively binds to sialic acid and N-acetylglucosaminyl residues enriched in the trans-Golgi network and plasma membrane
Phalloidin – AlexaFluor 568	F-actin (cytoskeleton)	Phallotoxin (bicyclic heptapeptide) that binds filamentous actin
MitoTracker Deep Red	Mitochondria	Accumulates in active mitochondria

Example Chemicals

Solvent control (0.5% DMSO)

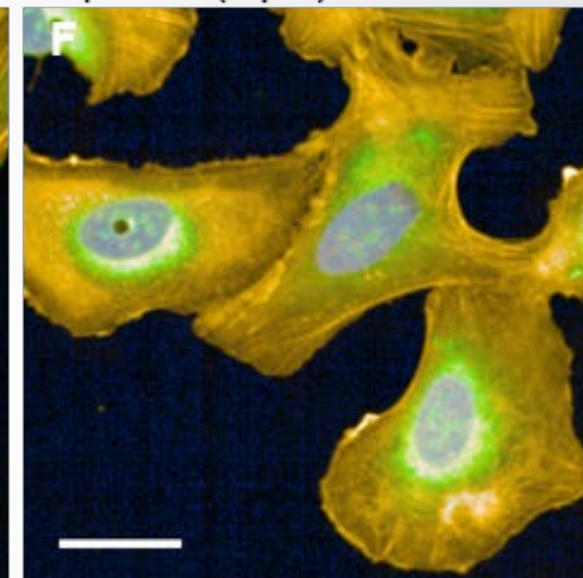
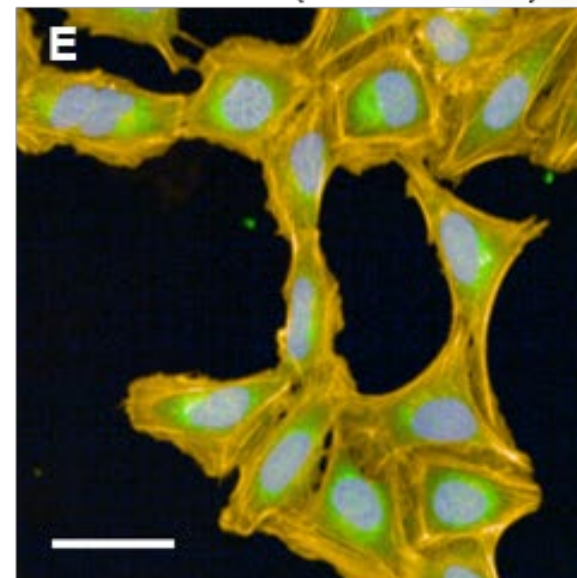
Berberine chloride (10 μ M)



→ Mitochondrial compactness/texture

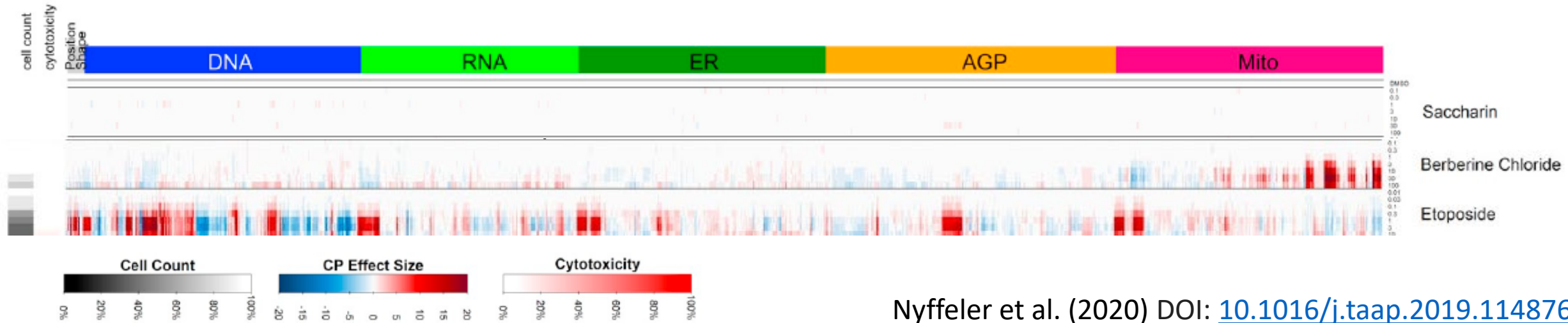
Solvent control (0.5% DMSO)

Etoposide (3 μ M)



→ Cells are larger

Strong phenotypes are observable qualitatively and can be measured quantitatively using imaging processing software



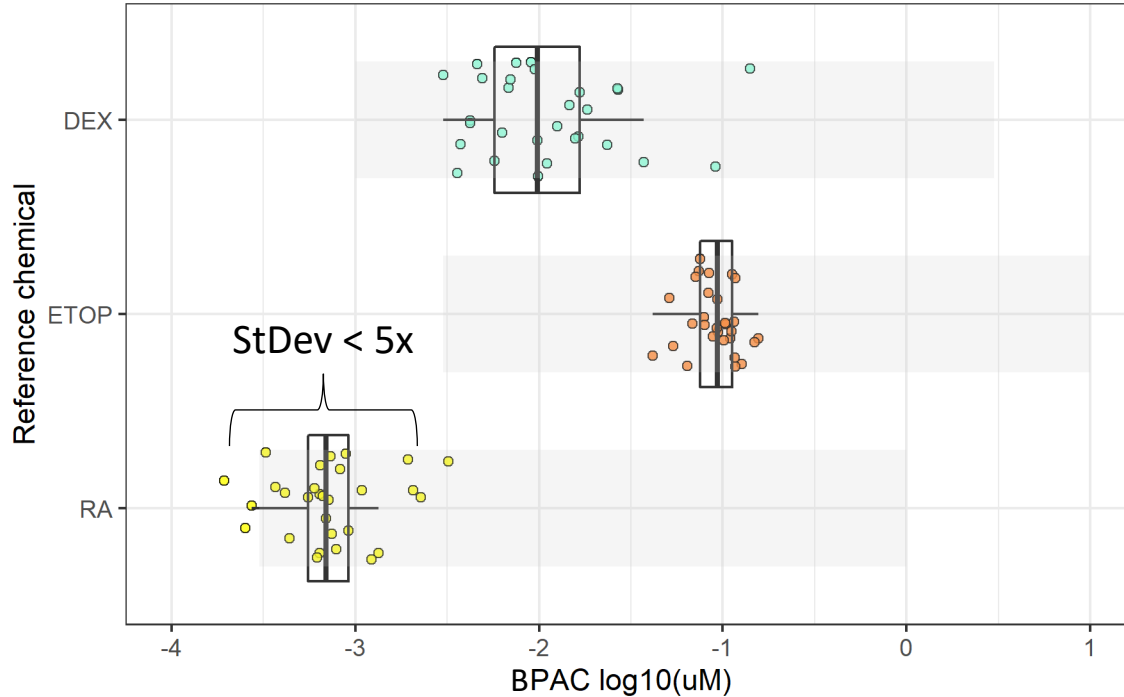
Nyffeler et al. (2020) DOI: [10.1016/j.taap.2019.114876](https://doi.org/10.1016/j.taap.2019.114876)

- Image analysis software quantifies multiple features per cell & fluorescence channel (intensity, size, texture, etc.): [CellProfiler](#) (*open source*) [Harmony](#) (*commercial*)
- Cell-level features are summarized per well & normalized to controls
- Different chemicals induce distinct, dose-responsive profiles

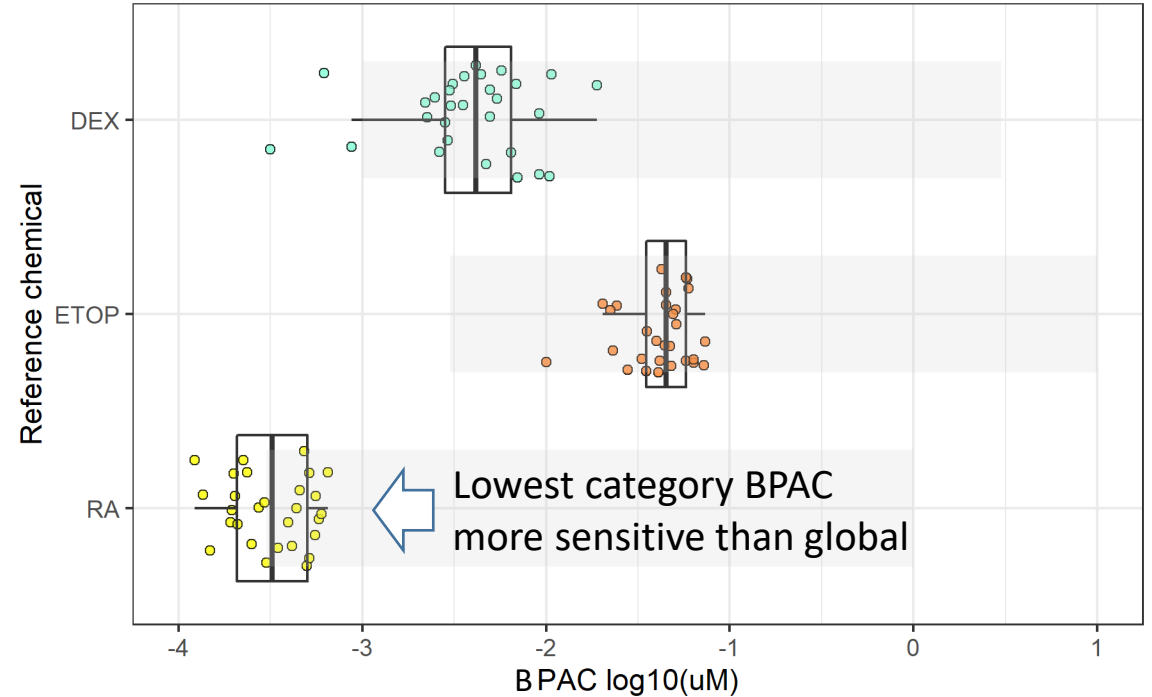


Assay Performance / Reproducibility

Global Mahalanobis:



Category-level Mahalanobis:



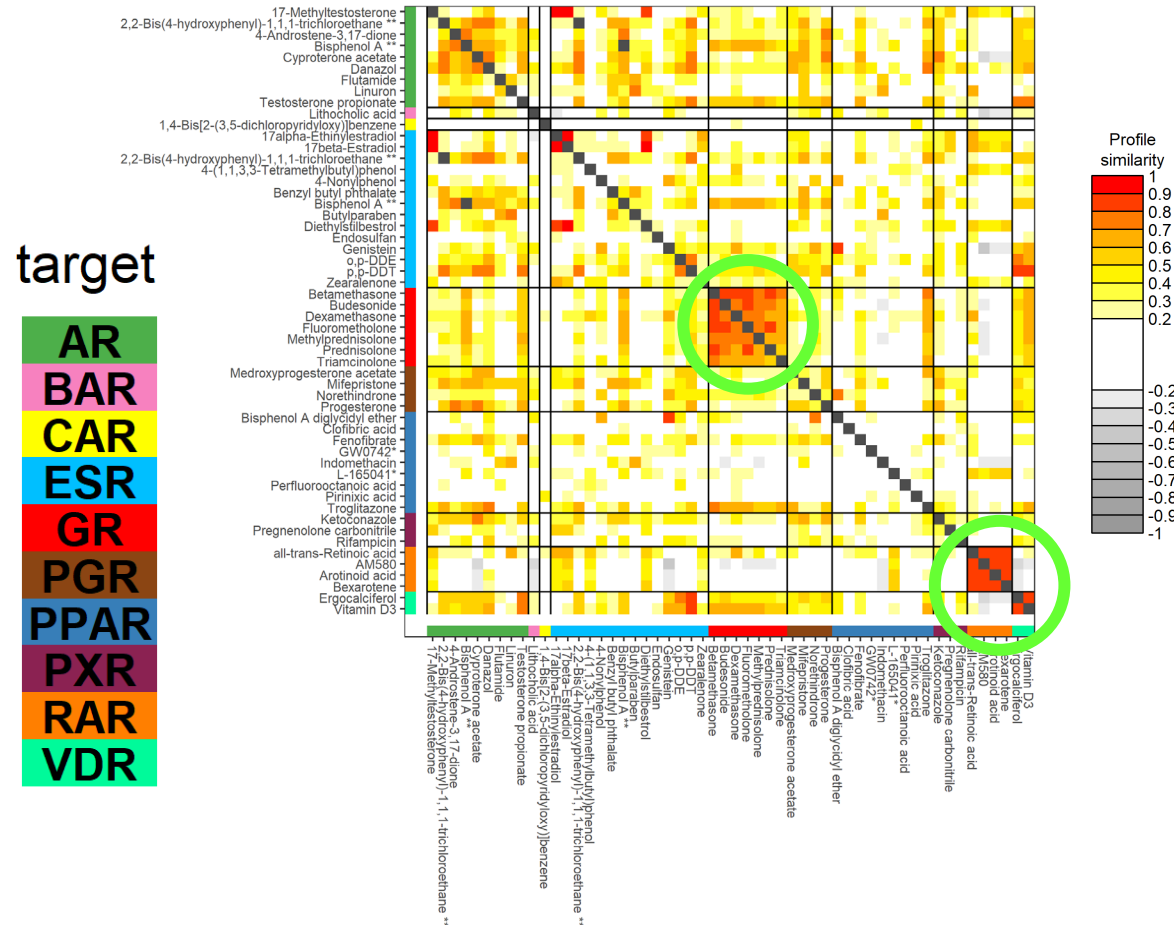
Used distance-based metrics (Mahalanobis) to derive the **Biological Phenotype Altering Concentration (BPAC)**

BPAC	
Retinoic Acid:	~ 0.3 nM
Dexamethasone:	~ 3 nM
Etoposide:	~ 30 nM



Profile Similarity Reflects Common Targets

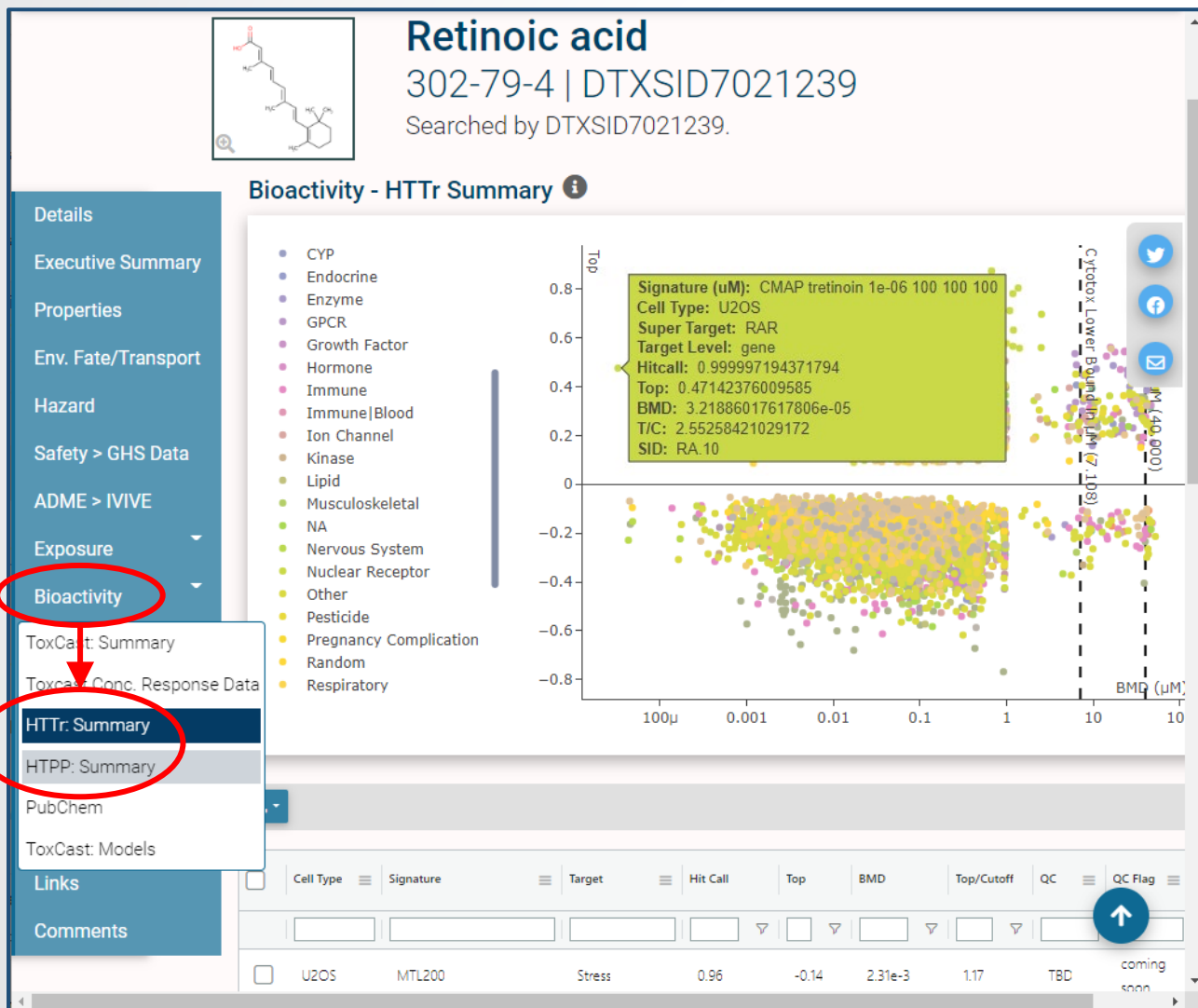
Biological Similarity in Cell Painting



- Cell Painting results for multiple chemicals in U-2 OS Cells
- Glucocorticoid & Retinoic Acid agonists display characteristic profiles
- Potential for similar applications to Connectivity Mapping in transcriptomics!



EPA High-Throughput Profiling Results



- Now available on EPA's CompTox Chemicals Dashboard! (comptox.epa.gov)
- Signature-level results viewable and downloadable (HTTr)
- Cell Painting results are also available (HTPP)



Summary

- High-throughput/broad profiling technologies can assess many types of bioactivity at once
- No single best way to analyze the data – depends on technology, experimental design, and use-case!
- Transcriptomics & Cell Painting can be used for:
 - Mechanism-agnostic POD determination, e.g. transcriptional POD (tPOD)
 - Mechanistic/Mode of Action (MOA) inference
 - Guiding/prioritizing further targeted testing
 - Generalized Read Across



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Questions?

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Contributors

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