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## Retrospective and Prospective Case Studies to Accelerate the Pace of Chemical Risk Assessment Katie Paul Friedman<sup>1</sup>, Matthew Gagné<sup>2</sup>, Tara Barton-Maclaren<sup>2</sup>, John Bucher<sup>3</sup>, Russell Thomas<sup>3</sup>, Santé Canada EUROPEAN CUEV Mike Rasenberg<sup>4</sup>, Tomasz Sobanski<sup>4</sup>

<sup>1</sup>National Center for Computational Toxicology, US EPA, Research Triangle Park, NC USA; <sup>2</sup>Healthy Environments and Consumer Safety Branch, Health Canada, Ottawa ON; <sup>3</sup>National Toxicology Program, NIEHS, Research Triangle Park, NC USA; <sup>4</sup>Computational Assessment Unit, European Chemicals Agency, Helsinki, Finland

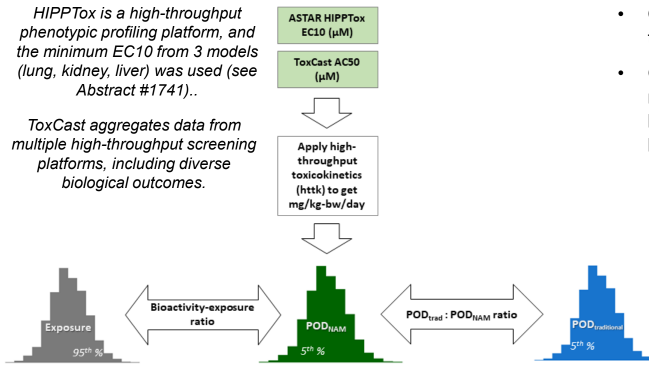
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## Abstract

Use of high-throughput, in vitro bioactivity data in setting a point-of-departure (POD) has the potential to accelerate the pace of human health risk assessments by chemical prioritization. Advancement toward this goal requires confidence that *in vitro* bioactivity data, in conjunction with high-throughput toxicokinetic information, can be used to estimate administered equivalent doses at or below the PODs from traditional animal studies. Further, hazard and exposure predictions, combined as a bioactivity:exposure ratio (BER) for use in risk-based prioritization, should be evaluated. In this work we describe two efforts of the Accelerating the Pace of Chemical Risk Assessment initiative, a consortium of international regulatory scientists, both with the same primary objective: to elucidate whether a POD derived from in vitro bioactivity would be a conservative estimate of traditional POD estimates, and if the BER is a useful prioritization metric. In the first project, we describe the outcome of a retrospective case study of 448 chemicals with high-throughput predictions of bioactivity, reverse dosimetry, and exposure, as well as traditional hazard information. For 92% of these chemicals, a POD derived from new approach methodologies (POD<sub>NAM</sub>) was a conservative prediction for the traditional POD (POD<sub>traditional</sub>) value. High-throughput exposure predictions were greater than the POD<sub>NAM</sub> for 26/448 chemicals, with BERs of less than zero, indicating higher priority for further investigation. The second, prospective study involves generation of NAM data for 200 chemicals to prioritize 20 chemicals for 90-day repeat dose testing in rats using a combination of the BER and bioactivity-based flags. Together these case studies enable regulatory scientists from different international contexts to develop efficient approaches for chemicals management, while possibly reducing the need for animal studies. This work demonstrates the feasibility, and continuing challenges, of using bioactivity and exposure NAMs in screening level safety assessment. This abstract does not necessarily reflect ECHA, Health Canada, NTP, or U.S. EPA policy.

# Part I: Retrospective case study

### Figure 1. Overall retrospective workflow



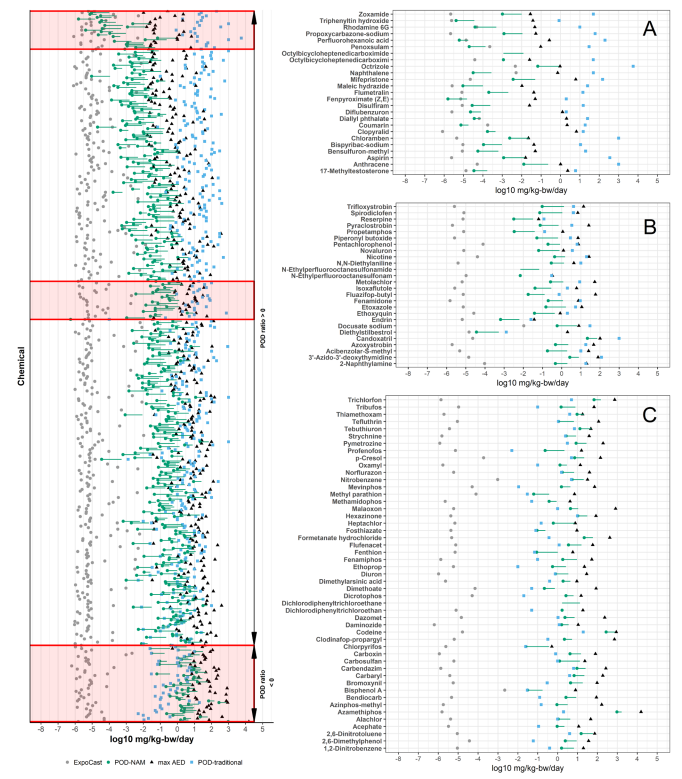
- 448 substances with exposure predictions (ExpoCast SEEM2 95<sup>th</sup> percentile for total US population), *in vitro* assay data, HTTK information, and *in vivo* hazard information.
- 50<sup>th</sup> and 95<sup>th</sup> percentile from the Monte Carlo simulation of inter-individual toxicokinetic variability were used to estimate administered equivalent doses (AEDs) for the minimum HIPPTox EC10 and the 5<sup>th</sup> percentile of credible ToxCast AC50 values for each substance.
- The minimum of either the ToxCast or HIPPTox-based AEDs were selected as the  $POD_{NAM, 50}$  or  $POD_{NAM, 95}$ . The  $POD_{NAM}$  estimates were compared to the 5<sup>th</sup> percentile from the distribution of the  $POD_{traditional}$  values obtained from multiple sources to obtain the  $log_{10}POD$  ratio.
- The log<sub>10</sub>BER was obtained by comparing the POD<sub>NAM</sub> estimates to exposure predictions. All values used for computation were in log<sub>10</sub>-mg/kg-bw/day units.

#### **POD**<sub>NAM.95</sub> would have been conservative for screening and prioritization purposes when compared to POD<sub>traditional</sub> for 89% (400/448) of the substances.

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### Figure 2. Comparison of Exposure, POD<sub>NAM</sub>, and POD<sub>traditiona</sub>

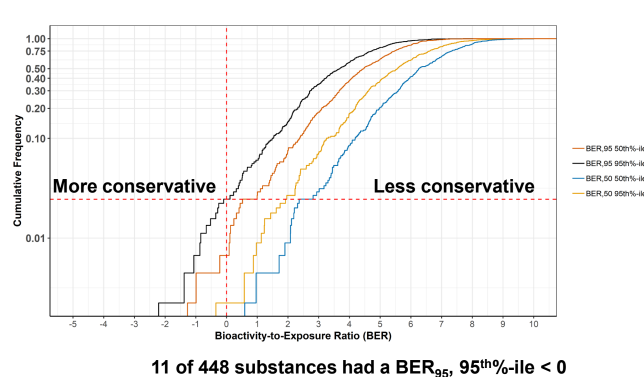
- Comparison of ExpoCast (SEEM2; gray circles), POD<sub>NAM</sub> (green circles), maximum AED (black triangles), and POD<sub>traditional</sub> values (blue boxes) for 448 substances.
- Green line segment indicates the POD<sub>NAM,95</sub> to POD<sub>NAM,50</sub>. Inset images A, B, and C correspond to the red boxes overlaid on the main plot. Image 3A provides a magnification on the substances with the largest log<sub>10</sub>POD ratio values. Image 3B displays a sample of substances that approach the median log<sub>10</sub>POD ratio. Image 3C includes all 48 substances for which the POD<sub>NAM, 95</sub> > POD<sub>traditional</sub>



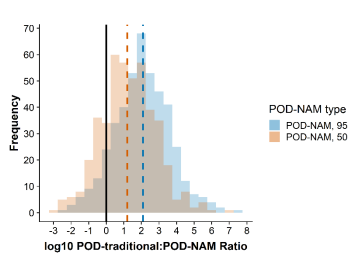
## NAM-based approach informs reasonable, conservative screening and prioritization

#### Figure 3. Cumulative frequency of bioactivityexposure ratio (BER)

- BER<sub>95</sub> used 95<sup>th</sup> percentile from the credible interval to predict median total US population exposure (ExpoCast SEEM2);BER<sub>50</sub> the 50<sup>th</sup> percentile.
- BER<sub>95</sub> and BER<sub>50</sub> values were calculated as the "95<sup>th</sup>%-ile" and "50<sup>th</sup>%-ile," using the POD<sub>NAM 95</sub> and POD<sub>NAM 50</sub>, respectively.



# 0 using the POD<sub>NAM 50</sub>.



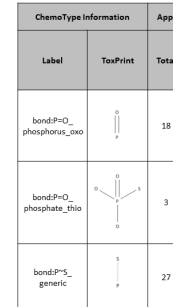
### Figure 7. When the $log_{10}$ POD ratio < 0, was it driven by a specific study type (as a surrogate for phenotypes)?

Condi log10-POD r log10-POD ra

#### Condi log10-POD ra log10-POD r

### Figure 8. When the $log_{10}POD$ ratio < 0, was it driven by a specific chemical features?

The enriched chemical structural features represented by ToxPrints for the log<sub>10</sub>POD  $ratio_{95} < 0$  set.

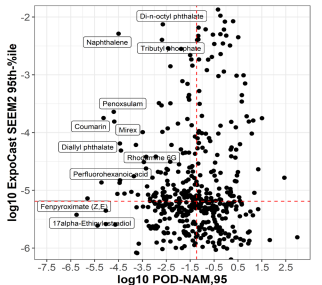


Based on a Fisher's exact test, chemical features associated with organophosphate pesticides and carbamates are more likely to drive a  $log_{10}POD$  ratio < 0.

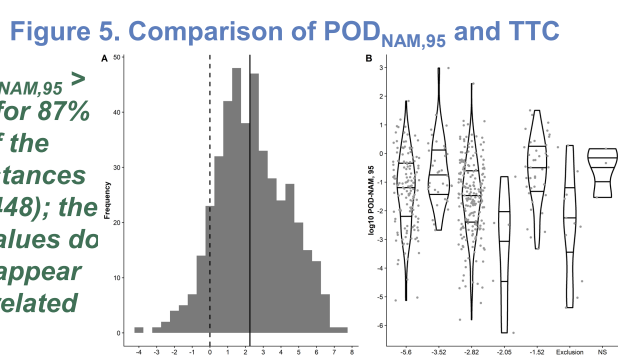
### BER<sub>95</sub>, 95<sup>th</sup> percentile did not prioritize an unreasonable number of substances; the BER selected reflects the level of conservatism and uncertainty considered within a screening assessment

#### Figure 4. Did exposure or bioactivity appear to drive the BER-based priority?

- Compared 95<sup>th</sup> percentile from the credible interval to predict total US population exposure (ExpoCast SEEM2) to the POD<sub>NAM 95</sub>.
- Dashed lines indicate the median exposure and POD<sub>NAM.95</sub> estimates for the 448 substances in the case study.



#### In general for $log_{10}BER < 0$ , the POD was relatively low. For certain substances the exposure estimates were relatively low.



log10 POD-NAM 95:TTC Ratio

*POD<sub>NAM,95</sub>* > **TTC for 87%** of the substances (389/448); the two values do not appear correlated

### **Figure 6.** log<sub>10</sub>**POD** ratio distribution

•  $\log_{10}$  POD ratio is illustrated for the POD<sub>NAM 95</sub> and the POD<sub>NAM 50</sub>. Using the more conservative (i.e., lower) POD<sub>NAM.95</sub>, 48 of the 448 substances (10.7%) demonstrated a  $\log_{10}$ POD ratio < 0 (to the left of the solid vertical line), whereas 92 of the 448 substances (20.5%) demonstrated a log10-POD ratio <

• The medians of the log10-POD ratio distributions are indicated by dashed lines for POD<sub>NAM. 95</sub> and  $POD_{NAM, 50}$  as 2 and 1.2, respectively.

> POD<sub>NAM.95</sub> includes interindividual variability in the in vitro to in vivo extrapolation process to a greater extent, and is more often a conservative estimate of POD<sub>traditional</sub>

ratio, $95 < 0$ 3 45	ition	Dev/Repro is min POD	Dev/Repro is not min POD
ratio $95 > 0$ 41 359	ratio,95 < 0	3	45
	ratio,95 > 0	41	359

ition	Chronic is min POD	Chronic is not min POD
ratio,95 < 0	28	20
ratio,95 > 0	244	156

#### Based on a Fisher's exact test, when log<sub>10</sub>POD ratio <0. it was not driven by a specific study type.

opearance of the ToxPrint			Metrics		ChemoType Information App		Appea	Appearance of the ToxPrint			Metrics		
tal	POD ratio ≤ 0	POD ratio > 0	ВА	OR	p-value	Label	ToxPrint	Total	POD ratio ≤ 0	POD ratio > 0	ва	OR	p-value
8	12	6	0.62	22	7.4E-09	bond:P~N_ generic	P	5	4	1	0.54	36	0.00055
3	3	0	0.53	NA	0.0012	bond:C(=O)N_ carbamate	N 0 0 0	20	6	14	0.54	3.9	0.014
7	13	14	0.62	10	3.5E-7	bond:CS_sulfide	c   s	53	15	38	0.61	4.3	0.00011

# Part II: Prospective case study

- How well does a NAM-based approach perform in the prospective case? This prospective case study builds upon learnings from the retrospective case study, addressing questions including Can NAM-based POD estimates be improved using additional technologies or assumptions? • Are reasonable NAM-based POD estimates attainable for substances with limited in vitro bioactivity?
- o Can BER, and additional hazard flags, be used to select substances for in vivo screening? dentification of substances with: · Limited hazard information and exposure potential Compatibility for currently available in vitro screening methodology Completion of a NAM battery for 200 substances within the substances identified • Multiple in vitro platforms: ToxCast, high-throughput transcriptomics, high-throughput phenotypic profiling, Immunotoxicity assays, acute neurotoxicity assays, developmental toxicity assays, endocrine-relevant assays and models • High-throughput toxicokinetic information for in vitro to in vivo extrapolation Figure 8. Output from Steps 1 and 2 Confirmatory 5-day *in vivo* testing based on BER and hazard flags
- Transcriptomics in liver Classical in vivo observations and toxicokinetics Further confirmation of a small subset from Step 3 in a 90-day subchronic study Evaluation • Comparison of Step 2-4 data (if available), and any other traditional hazard information

## Conclusions

- A major premise of this work is that the minimal concentration corresponding to *in vitro* bioactivity is likely to be a conservative threshold for any specific effects or toxicities that might be observed *in vivo*.
- IVIVE that is included in development of the POD<sub>NAM</sub> and (2) the exposure predictions, highlighting that for different screening applications differing amounts of uncertainty can be included in this workflow.
- The prospective case study furthers confidence, and identifies possible limitations, in NAM-based screening assessments.
- The collaborative, international consideration of these issues in screening level assessments demonstrates the current stateof-the-science and presents a transparent and adaptable basis for utilization of HTS information.



-		In 2019, any gaps in this heatmap will be filled.    Scenario 1   Substance present on the EU, Canada, and/or US market, with a potential for consumer use and
		significant data gaps for systemic toxicity (105).
		Scenario 2
		Substance present on the EU, Canada, and/or US market, with known toxicity and potential interspecies differences (8).
		Scenario 3
		Substance selected from the retrospective case study, by sampling substances with varying log10POD ratios.
EPA Bioavailability (CaCo2)	EPA HTTK	The BER (<10 <sup>4</sup> ) from Step 2, and hazard flags based on potential endocrine, developmental, neuro, and/or immuno-toxicity, will be used to advance ~20 substances to Step 3.

• BER may be a reasonable data-driven metric for prioritization that is tunable based on the amount of uncertainty in (1) the