

Scientific Advances in Non-Targeted Chemistry



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Outline

What is Non-Targeted Analysis?

Benchmarking and Publications for Non-Targeted Analysis

Uses of NTA data

Future of Non-Targeted Analysis

What is Non-Targeted Analysis?

Targeted Analysis

"known knowns" Standards, calibration curves

Suspect Screening Analysis (SSA)

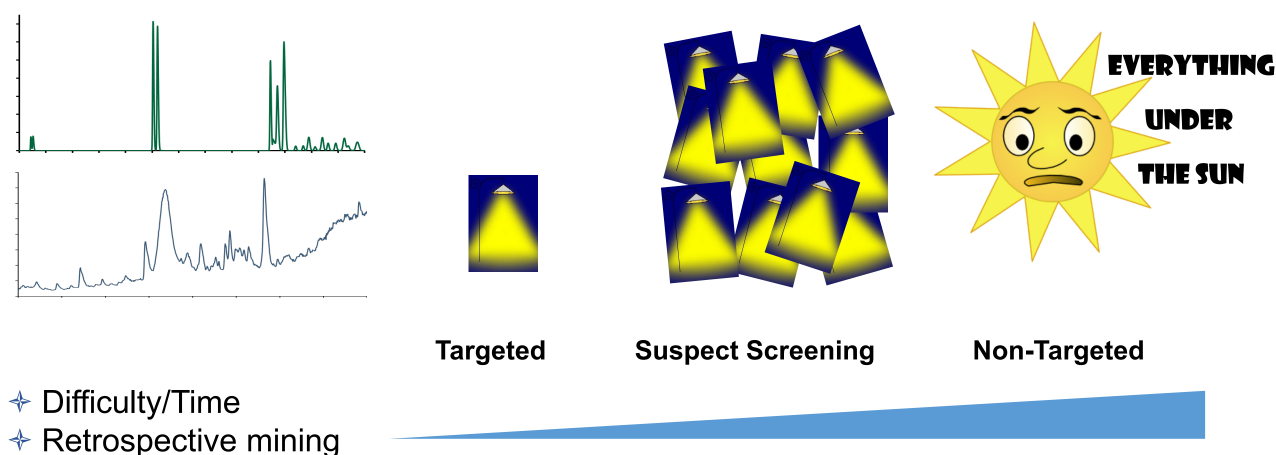
"known unknowns" Lists of compounds

Non-Targeted Analysis (NTA)

"unknown unknowns" MS first principles



Targeted vs. Non-Targeted Analysis



♦ Quantitative info

Structure confidence

How does High Resolution MS work?

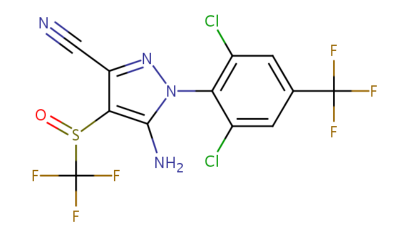
Atom	Natural Abundance	Exact Mass
¹ H	99.9885%	1.007825
² H	0.0115%	2.014102
¹² C	98.93%	12.000000
¹³ C	1.07%	13.003355
¹⁴ N	99.632%	14.003074
¹⁵ N	0.368%	15.000109
¹⁶ O	99.757%	15.994915
¹⁷ O	0.038%	16.999131
¹⁸ O	0.205%	17.999159
¹⁹ F	100%	18.998403
³² S	94.93%	31.972072
³³ S	0.76%	32.971459
³⁴ S	4.29%	33.967868
³⁶ S	0.02%	35.967079
³⁵ Cl	75.78%	34.968853
³⁷ Cl	24.22%	36.965903

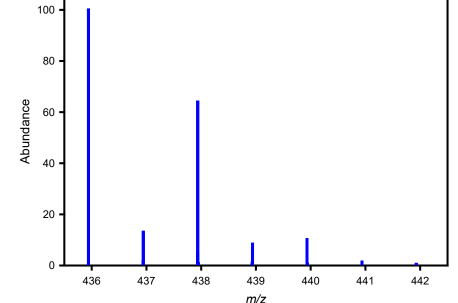
Example: Fipronil

Molecular Formula: $C_{12}H_4CI_2F_6N_4OS$

Monoisotopic Mass: 435.938706

= (12.0000*12 Carbon) + (1.007825*4 Hydrogen) + (34.968853*2 Chlorine) + (18.998403*6 Fluorine) + (14.003074*4 Nitrogen) + (15.994915*1 Oxygen) + (31.972072*1 Sulfur)





Benefits of Using Non-Targeted Analysis

Ability to detect many more compounds

- + Includes unknowns, things not in databases (like metabolites)
- + Broad range of chemical space covered (Define!)
- Rapidly screen for knowns
 - + Virtually unlimited in number
- Data is collected in a way to allow retrospective analysis

+ When did this compound start showing up?

Benchmarking and Publications for Non-targeted Analysis (BP4NTA)

✤ ~90 international members

Membership based on interest in NTA

- + Experience with NTA varies from beginners to experts
- Wide range of applications: metabolomics, exposure, food, biological, medical devices, environmental

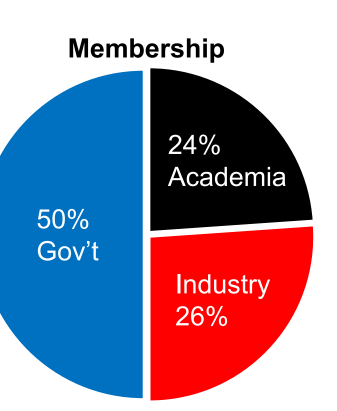
Leads Ben Place (NIST) and Elin Ulrich (EPA)



Interested? Contact us! benjamin.place@nist.gov

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Working Group Objectives

- Create a list of commonly-used NTA terms, concepts, and performance calculation and provide definitions/equations
 - + Publish guidance document with terms, use community consensus and feedback
 - + Audience includes new researchers, journal reviewers/editors, and experts
- Reporting recommendations and scientific best practices to promote transparency and reproducibility
- Build and maintain coalitions/communications with other groups that have similar interests including metabolomics, NORMAN, mQACC
- Move toward proficiency testing levels for SSA and NTA (ASTM/ISO)
 Define proficiency expert, competent, etc. (10 years out)

Examples from Working Group

Suspect screening is a methodology that aims to identify chemicals detected via mass spectrometry by <u>comparing to a predefined user</u> <u>list or library</u> containing known chemicals of interest using information such as accurate mass and isotope ratios.

Non-Targeted Analysis (NTA) aims to identify chemicals of interest detected by mass spectrometry <u>without a predefined list of</u> <u>chemicals</u>. Note NTA may include suspect screening, for example, by using a defined list and looking for those features in the data then attempting the identification of any remaining features without a defined list. Due to this complimentary usage of NTA and suspect screening, NTA often encompasses both suspect screening and true NTA.

Results Reporting: Data Processing & Analysis Performance								
Rec	ommended Data to Assess and Report							
Perf. Metric	Performance Data	Proposed Questions for Evaluation						
Quality	QC checks along data analysis workflow (e.g., check for QC compounds, check for outlier samples)	Were QC benchmarks established? Did results meet those benchmarks?						
Boundary	Quantification or semi-quantification of identified compounds and associated limits of detection/identification Chemical space (e.g., K _{ow} , ionizability, etc.) of the data processing/analysis method (e.g., of the library/database used; constraints introduced by approaches such as mass defect analysis or molecular networking, etc.)	Did sensitivity impact compound detection? Was the chemical space covered by the data analysis method assessed? Did the data processing/analysis methods impact the chemical space of detected/identified compounds?						
Accuracy	Performance calculations, such as the True Positive Rate (TPR), False Positive Rate (FPR), True Negative Rate (TNR), and False Negative Rate (FNR) either 1. <u>at the annotation (compound) level</u> , for known compounds in QC spikes/controls, or 2. <u>at the sample level</u> , for samples with known classification/grouping	<u>Annotation level:</u> Do these metrics provide insight about the accuracy/selectivity of the detection and identification workflow with respect to certain compound classes or chemical characteristics? <u>Sample level:</u> Do these metrics provide insight about the accuracy/selectivity of the classification method with respect to certain sample types?						
Precision	Reproducibility of detection and identification for QC spikes across sample types or in QC controls (e.g., re-injected pooled samples). Performance calculations, such as the False Discovery Rate (FDR) or Precision.	Was a threshold for repeatable detection and identification listed? Was detection and identification sufficiently reproducible? What factors impacted reproducibility?						

	Section	Category	Sub-Category	Example Information to Report	
		Experimental Design	Objectives & Scope	•Study goals, hypotheses, and use/definitions of NTA/SS in the study •Expected chemical coverage of approach (e.g., volatile, nonpolar compounds by GC-EI-MS)	
r			Sample Information & Preparation		
			QC Spikes & QC Controls	 Use of isotopically labeled standards and/or RT reference material Use of Development and use of positive (spiked) controls, pooled matrix controls, blanks for QC 	
		Data Acquisition	Run Order Preparation	•Sample replication and randomization •Inclusion of blanks and QC samples in the acquisition sequence	
			Chromatography	 Instrument specifications Method settings (e.g., column, mobiles phase, gradient, injection techniques) 	
	Methods		Mass Spectrometry	 Instrument specifications and Method settings (e.g., resolution, acquisition parameters, DDA vs. DIA vs. AIF) Instrument calibration and/or tuning procedures 	
		Data Processing & Analysis	Data Processing	 Software program(s) used, including file conversion Workflow steps (e.g., centroiding, peak picking, alignment, gap filling), methods, and settings Peak detection thresholds (e.g., replicate detection criteria; minimum height, area, or S/N levels; comparison to blanks) Data correction or normalization methods (e.g., RT calibration/indices, peak area/height normalization with IS, blank subtraction) 	
			Statistical Analyses	 Software program(s) used Method goals (e.g., feature prioritization, compound class identification, sample classification) Method type (e.g., clustering, classification, hypothesis testing) and settings 	
Kathy Peter (NIST) Lead			Annotation & Identification	 Software program(s), libraries, and databases used (including information about in-house databases) Workflow steps (e.g., formula assignment, suspect screening, MS/MS spectra interpretation, library MS/MS matching) Workflow methods (e.g., formula prediction method, scoring algorithms) and settings Thresholds for annotation/identification (e.g., mass error and RT; scores for formula assignment, MS/MS spectral matching) 	
	Results	Data Outputs	Identification & Confidence Levels	 Reported identifications and associated confidence levels (e.g., Schymanski et al. levels) Supporting annotated data (e.g., formula, RT, MS/MS match scores, fine isotope pattern, source of MS/MS spectra) For unidentified features (i.e., not standard-confirmed), proposed tentative structures and other annotated data Exported MS/MS spectra (e.g., as a library, database, or deposition into online repository) 	
			Statistical Outputs	 Visuals/plots (e.g., heatmaps, PCA and loading plots) and statistical output (e.g., adj. p-values) Reported classifications or groupings of features, identifications, or samples New statistical packages or code 	
		QA/QC and Other Performance Metrics	Data Acquisition	 Quality: Deviations from QA practices and results from QC checks for sample preparation and data acquisition Boundary: Description of the capabilities/chemical space of sample prep, chromatographic, and MS methods Accuracy: Reported chromatographic and mass accuracy Precision: Reported variability of retention time, precursor mass error, and abundance 	
			Data Processing & Analysis	 Quality: Outcomes of QC checks along data analysis workflow Boundary: Quantification or semi-quantification of identified compounds, limits of detection/identification Accuracy: Calculations such as TPR, FPR, etc. at annotation level for QC spikes/control samples or at sample level Precision: Reproducibility of identification for QC spikes across sample types 	

Example Uses and Requirements

Decision Context		≠xt	/	
Sample	Chemical	Semi-	Example Uses of NTA Data	Example
Classification	Annotation	Quantitation		Stakeholders
	, I		- Classify locations impacted by point-source emitters	- EPA, USGS
	1	1	- Classify locations impacted by inadvertent environmental releases	- FEMA, EPA
Req	Opt	Opt	- Classify exposure status for active or former military personnel	- DoD, VA
	<u> </u>		- Classify food items not meeting criteria for product certification	- FDA, NIST
	, I		- Identify natural or synthetic chemical nerve agents	- DHS, CDC
	1	1	- Identify chemicals associated with product-related illness	- CPSC, FDA
Req	Req	Opt	- Identify chemicals released in emergency response scenarios	- FEMA, EPA
			- Identify designer drugs used for athletic performance enhancement	- DEA, FDA
	1	1	- Assess occupational health risks from exposure to fire-fighting foams	- NIOSH, DoD
	1	1	- Assess consumer health risks from exposure to household products	- CPSC, EPA
Req	Req	Req	- Assess ecological health risks from exposure to urban wastewater	- USGS, EPA
	1	1	- Assess maternal and infant health risk from exposure during pregnancy	- NIEHS, EPA
	·'	<u> </u>		

Uses of NTA Data

Exposure surveillance

- + What chemicals are in food, water, products, dust, blood, etc.?
- Starting point for generation of targeted methods to allow addition to lists like Unregulated Contaminant Monitoring Rule (UCMR)

Chemical prioritization

- + What are relevant chemicals & mixtures?
- Lautenberg Act- Risk-based process to determine which chemicals to prioritize for assessment, identifying them as high/low priority substances

+ Exposure forensics

- + What are chemical signatures of exposure sources?
- + Can assist with enforcement/cleanup efforts

Biomarker discovery

- + What chemicals are associated with health impairment?
- + Provides data for AOPs, new tests for HT toxicity screening efforts

The Future of NTA

- Standardized QA/QC, terminology, review, reporting
 As possible, standardize methods
- Benchmarking, performance metrics
 True/False Positives/Negative, chemical space coverage
- Learning from related fields (e.g., metabolomics)
- Semi-quantitative analysis
- Regulatory uses
- * "Make non-targeted the new targeted" Thomas Burke

Acknowledgements and References

Ben Place (NIST) Jon Sobus (US EPA) Kathy Peter (NIST) Sara Nason (CT Agr Exp Stn) Seth Newton (US EPA) BP4NTA working group members EPA NTA researchers

BP4NTA website (Sara and Seth)- <u>https://nontargetedanalysis.org/</u> CompTox Chemicals Dashboard- <u>https://comptox.epa.gov/dashboard/</u> SETAC Focused Topic Meeting Nontarget Analysis for Environmental Risk Assessment- <u>https://nta.setac.org/</u>

Integrating tools for non-targeted analysis research and chemical safety evaluations at the US EPA https://www.nature.com/articles/s41370-017-0012-y

EPA's non-targeted analysis collaborative trial (ENTACT): Genesis, design, and initial findings https://link.springer.com/article/10.1007/s00216-018-1435-6

Using prepared mixtures of ToxCast chemicals to evaluate non-targeted analysis (NTA) method performance https://link.springer.com/article/10.1007%2Fs00216-018-1526-4

Examining NTA performance and potential using fortified and reference house dust as part of ENTACT https://link.springer.com/article/10.1007%2Fs00216-020-02658-w