



Applied Mass Spectrometry

TargetScreener HR

One workflow – all the answers

Innovation with Integrity

Answers today, tomorrow, every day

Always Accurate Results



Food Safety



Forensics



Toxicology



Environment

Environmental, food safety, toxicology and forensic laboratories need tools to rapidly screen for target compounds with accuracy and certainty. Bruker's workflow offers several benefits including providing data-independent acquisition and all ion fragmentation data, enabling the analysis of an unlimited number

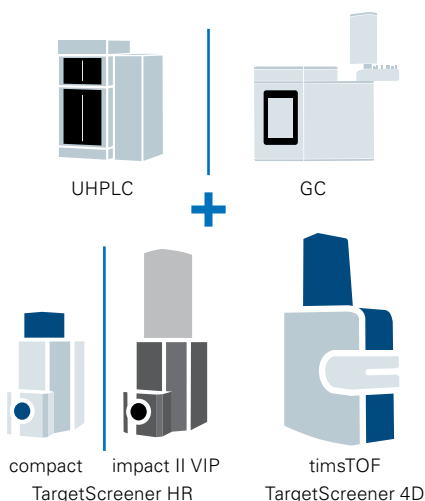
of compounds, and allowing retrospective analysis. TargetScreener HR encompasses all these features.

TargetScreener HR represents the modern way of monitoring: beyond obligatory testing, towards a complete picture to avoid surprises and headaches.

Comprehensive screening workflow

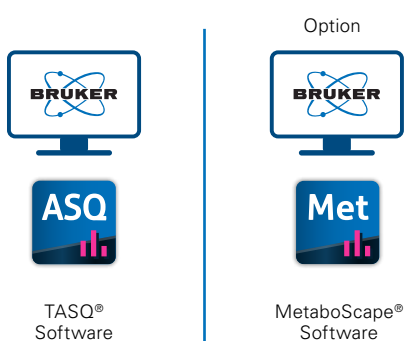
Combination of reliable data, powerful software, and robust database for sophisticated results

1 HARDWARE



Acquisition of full scan MS and bbCID accurate mass data

2 SOFTWARE



Target and untargeted analysis: Screening and quantitation

Discovery workflow: Statistical analysis Identification of unknown

3 REPORT



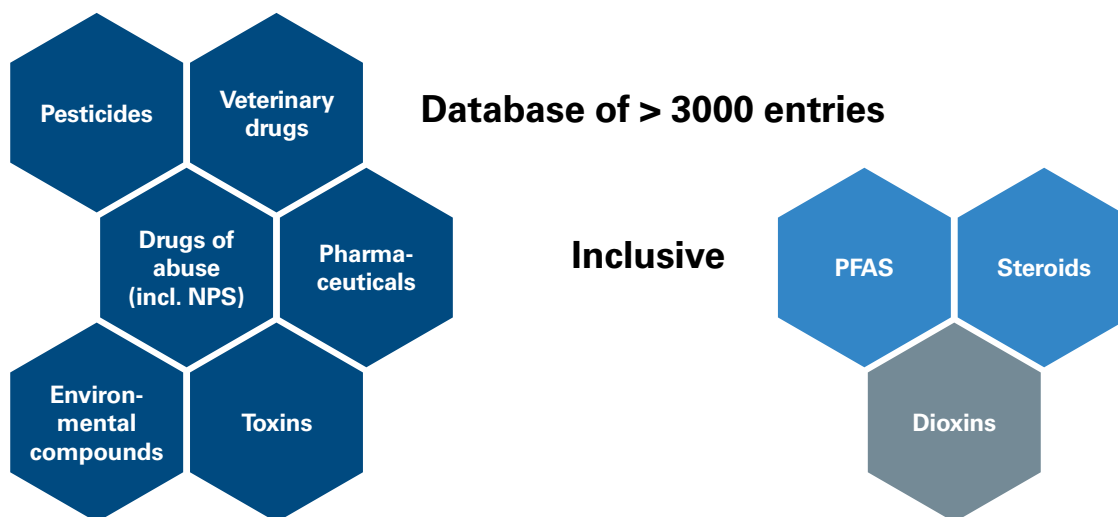
Report

Result reporting and/or LIMS export

- **TargetScreener HR**
 - Comprehensive screening – targeted and non-targeted
 - Minimizes false positive and false negative reporting
 - Accurate and precise quantitation meets stringent regulatory guidelines
- **TargetScreener 4D**
 - Plus Trapped Ion Mobility Spectrometry (TIMS) resulting in Collision Cross Section (CCS) values - a new 4th dimension of orthogonal selectivity for absolute confidence

Extended compound database: Drugs, pesticides, and environmental pollutants

One solution for GC and LC workflows



The core of the TargetScreener HR workflow lies in its precise database. This comprehensive database includes high-resolution monoisotopic precursor ion masses, retention time data, and up to ten detailed qualifier ions. The qualifier ion data encompasses accurate mass b/cID fragments, 'in source' fragment ions,

adduct information, and isomer details. Additionally, CCS values are included to enhance the confidence in identification.

A perfect match to the TargetScreener HR database is indicated by the green color of the MRSQC or MRSQC score.

Color-coded screening results

Straightforward reviewing of either a sample or an analyte as desired by the analyst. Once reviewed, qualitative and quantitative sample reports are then easily generated.

M R S Q C



Mass accuracy
Retention time
mSigma value (Isotopic pattern)
Qualifier ions (Diagnostic ions)
CCS value



Parameter within narrow window

Value between narrow and wide limits

Parameter outside the wide tolerance window

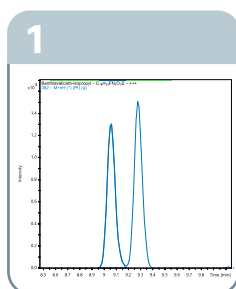
No score available

5 Steps from sample to report

Powerful for in-depth investigative screening and quantitation
as well as retrospective data mining

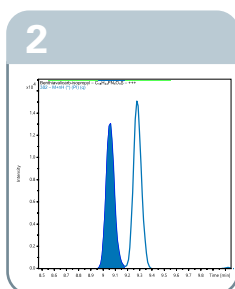


Ready-to-use LC/GC-, MS- and software methods reduce lab time.
Reaching confident results in just 5 steps with the TASQ software:



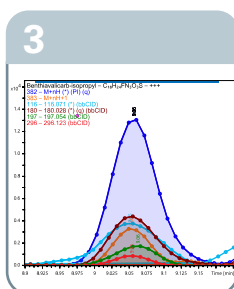
Screening

Generation of
EIC traces for
Principal Ions



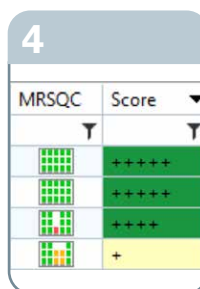
Peak detection

Automatic Retention
Time check



Verification

Check diagnostic
ions: in-source and
bbCID fragments,
isotopes and adducts



Scoring

Scoring by mass
accuracy (M),
Retention Time (R),
Isotope pattern (S),
diagnostic Qualifier
ions (Q) and CCS
values (C)



Quantitation

Instant calculation of
concentration of
confirmed compounds

TargetScreener HR searches all ions all of the time by full scan MS and MS/MS without compromising sensitivity, resolution or mass accuracy. Datasets are complete and unrestricted. The diagnostic qualifier ions

are used as an enhanced confirmation criterion to eliminate false positives as well as deliver unambiguous identification for co-eluting isobaric drugs.



Compliant with international guidelines



Start with ready-to-use methods

With Bruker's innovative technologies and default software methods, your lab will pass the method validation and analytical quality

control requirements that are necessary to support the validity of data.

Summary of SANTE 11312/2021 guideline criteria

- 1 RT tolerance (standard sample): ± 0.1 min

Analytes settings		
RT tol. [min] $\pm$	RT narrow [min] $\pm$	RT wide [min] $\pm$
0.50	0.25	0.40

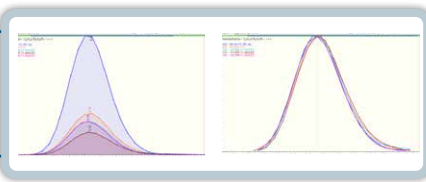
- 3 2 Ions with mass accuracy ≤ 5 ppm (or ≤ 1 mDa below 200 m/z)

Ion	Spectrum type	Mandatory	mDa narrow $\pm$	ppm narrow $\pm$	m/z
M=nH	FullScan	☒	0.9	3.00	284.0538
M=nH+1	FullScan	☒	0.9	3.00	285.0566
104.016	bbCID	☒	1.0	9.61	104.0165
199.960	bbCID	☒	1.0	5.00	199.9599

- 2 $S/N \geq 3$ or signal should be present in at least in 5 subsequent scans

Method Editor	
Minimal datapoints over a peak	5
Signal/noise filter	0.500

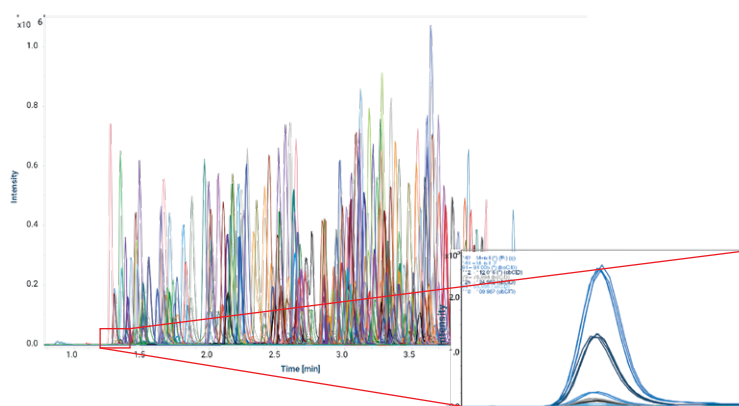
- 4 Full overlap of precursor and product ion EICs



Accelerate – Pesticides in wheat matrix in just 5 min

Analysis of 294 pesticides in wheat matrix with a 5 min method.

The pesticides were spiked at concentrations of 5, 10 and 20 $\mu\text{g/kg}$ resulting in an analyzed concentration of 1.25, 2.5 and 5 ng/mL . The data was evaluated with the TASQ software.

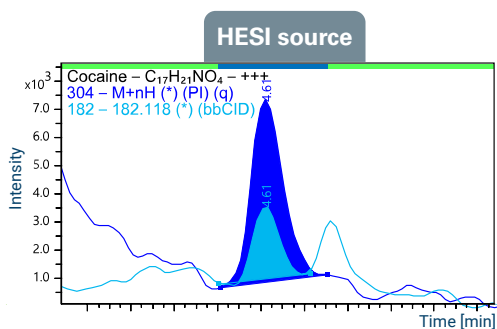
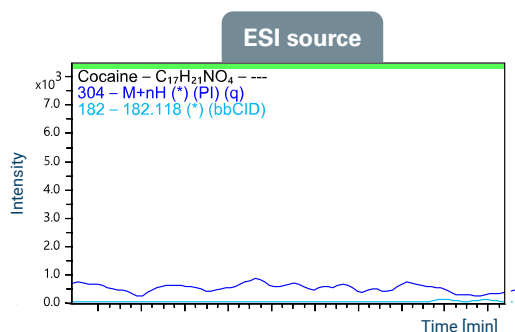


Analytical tools in forensic science

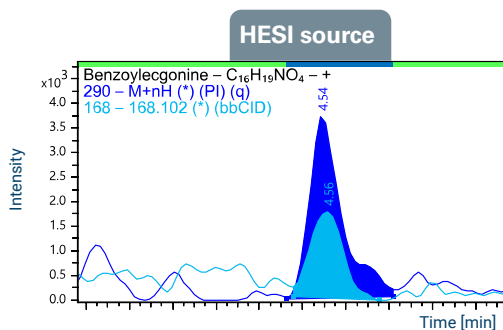
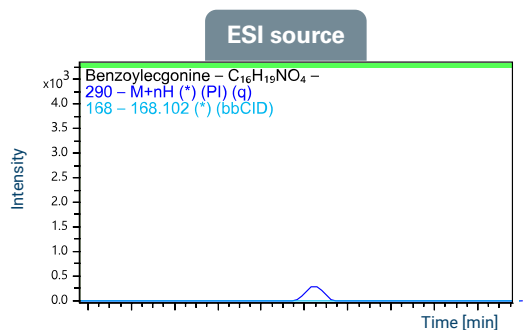
Increased sensitivity for forensic drug investigations

The VIP-HESI source, a novel dual source, is equipped with Heated Electrospray and Atmospheric Pressure Chemical Ionization (APCI) probes. Less thermal degradation and high-efficient ionization leads to a highly improved limit of detection (LOD). The reporting of false positives is minimized due to accurate mass data and the highly curated database.

Cocaine



Benzoylecgonine



Hair samples analyzed with ESI or VIP-HESI. Hair samples were cut into small pieces. After sample preparation a solution was injected into the UHPLC and analyzed with the impact II VIP HR-MS. Shown are the EICs from Cocaine (concentration 0.3 ng/mL in hair) and its metabolite Benzoylecgonine.

TargetScreener HR workflow allows for an **easy addition of new compounds** to the HR-MS database, which is of extreme importance for the rapid emergence of new drugs. TargetScreener HR collects all data, all of the time, allowing for retrospective analysis of previously screened samples with the addition of new compounds to the database.

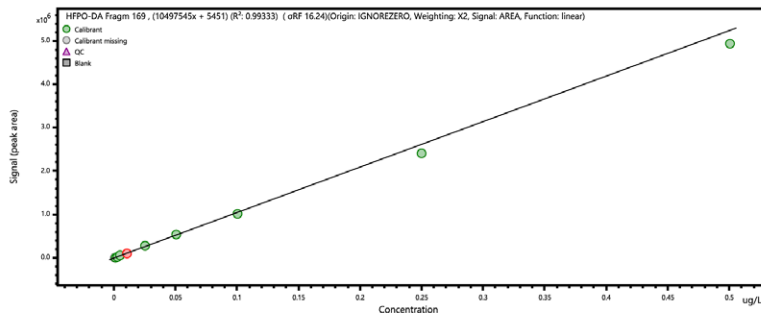
Powerful for screening and quantitation

Accurate quantitation of analytes

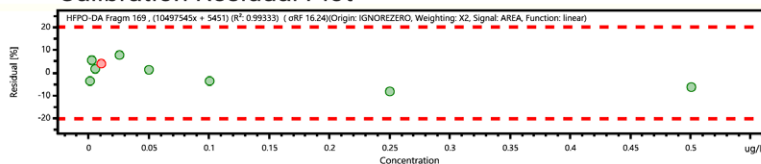
PFAS, known as per- and polyfluorinated alkyl substances, represent a broad category of industrial chemicals that have been widely utilized. As a result of their potential toxicity, PFAS pose an escalating danger to both humans and wildlife. Consequently, the analysis of these substances has become an increasingly crucial undertaking for analytical chemists. With the TargetScreener HR workflow it is possible to screen and quantitate PFAS.

GenX (HFPO-DA)

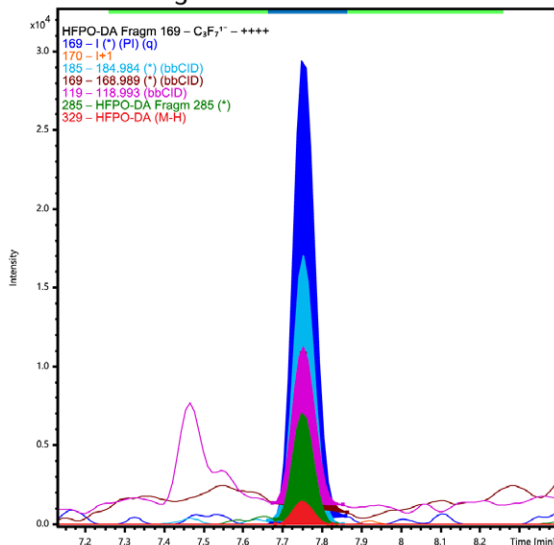
Calibration Curve



Calibration Residual Plot



Chromatogram



Quantitation of GenX – a PFAS compound

Extract from TASQ software. Shown are the calibration curve (concentration range: 0.001-0.5 $\mu\text{g/L}$), the calibration residual plot and the chromatogram of GenX/HFPO-DA (concentration 0.01 $\mu\text{g/L}$).

Addressing the crucial challenge of screening and quantitation of analytes in diverse matrices

Bruker's TASQ software, included in the TargetScreener HR workflow package, is specifically designed to the comprehensive requirements of customers. It offers a user-friendly method setup with a menu-driven, quick start wizard, expertly guiding users through qualitative and quantitative workflows. TASQ provides the convenience of preconfigured reports or the option to export data to a Laboratory Information Management System (LIMS) for further customization in processing and reporting.

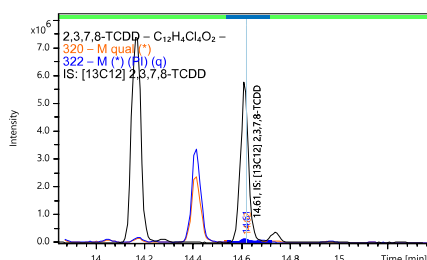
A new standard in GC-APCI



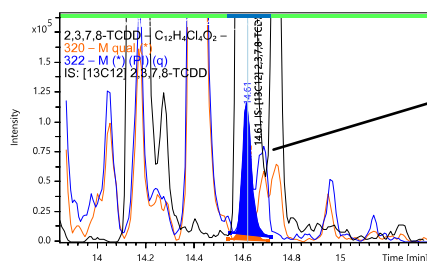
High-speed TIMS coupled to GC-APCI-HR-MS: The ultimate screening power

TargetScreener 4D provides a ready-to-use method for the analysis of Dioxins, Furans, and PCBs in the same run and matching Regulation (EU) 2017/644. Bruker's solution enables monitoring of the levels of dioxins,

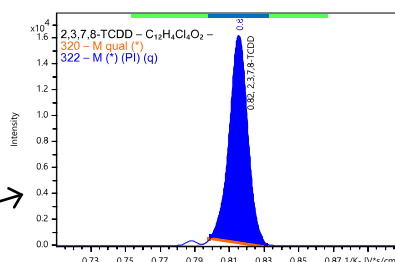
dioxin-like PCBs and non-dioxin-like PCBs in certain foods. Save lab time and increase productivity with Bruker's ready-to-use method for 350 contaminants.



↓ in zoom



CCS values
as additional
ID criterion



Quantitation of 2,3,7,8-TCDD in certified reference material. Sludge extract spiked with a mixture containing several congeners of dioxins and furans (specially several congeners of TCDF and TCDD) and diluted 10 times before injecting.

§ REGULATION (EU) 2017/644 (METHODS OF ANALYSIS FOR THE CONTROL OF LEVELS OF DIOXINS)

- RT tolerance (relative RT in relation to IS or ref. standards): ± 0.25 %
- Repeatability (RSDr) < 20 %
- Intermediate precision (RSDR) < 25 %
- Monitoring of at least two specific ions

TargetScreener 4D

One package – Unlimited solutions!

Add a new dimension for more confidence



1 RT

2 m/z

3 MS/MS

4 Mobility

Additional ID criterion

Separation and quantitation of isomers

Clean-up of chromatograms and spectra

Trapped ion mobility spectrometry (TIMS) provides an additional dimension of separation, unlocking and eliminating all uncertainties of the sample analysis and generates ultimate confidence in compound identification.

Additional ID criterion

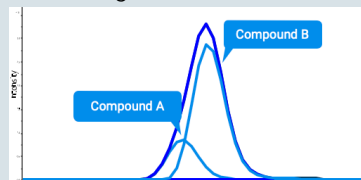
M R S Q C



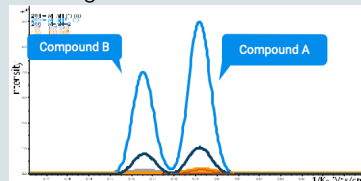
Mass accuracy
Retention time
mSigma value (Isotopic pattern)
Qualifier ions (Diagnostic ions)
CCS value

Separation and quantitation of isomers

Chromatogram

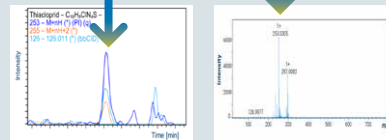
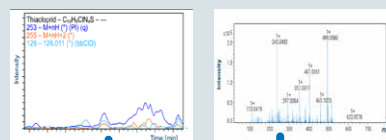


Mobilogram



Clean-up of chromatograms and spectra

Mobility filter OFF

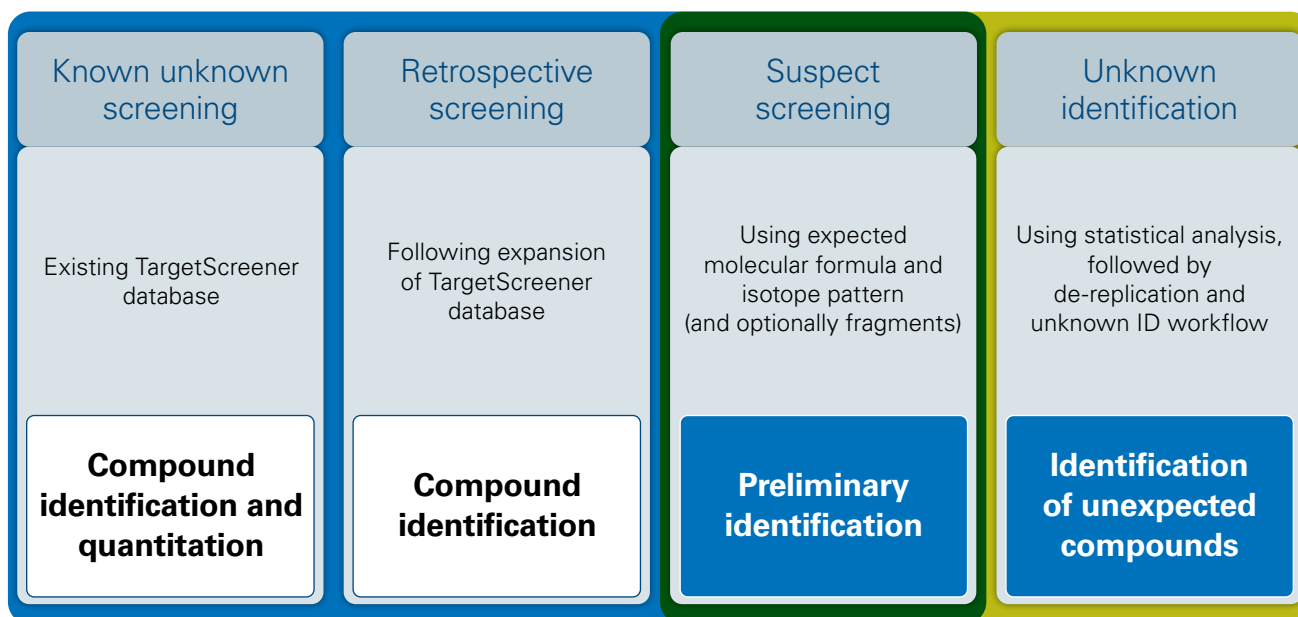


Mobility filter ON

Full array of identification workflows

Bruker's software suites **TASQ®** and **MetaboScape® software** process target and non-targeted workflows. TASQ supports various screening methodologies, plus accurate quantification based on an available calibration curve. MetaboScape® uses statistical analysis to detect unexpected compounds and identifies 'known unknowns' using public databases and spectral library searches (e.g. MetaboBase® library, NIST 2023).

Identification workflows



TASQ

Target screening & quantitation

MetaboScape

Discovery metabolomics & phenomics

High-resolution mass spectrometry (HR-MS) is made with data-independent acquisition, with unbiased ion fragment generation and detection, enabling analysis of an unlimited number of compounds.

Spectral libraries with high mass accuracy are then utilized for the identification of unknowns. For added flexibility, users can add new compounds to the HR-MS database as they become available. This is future proof for any investment for years to come, enabling retrospective data mining of previously screened samples.

- High sensitivity with the novel VIP-HESI source down to ppt levels.
- Generation of HR accurate mass parent ions, true isotope pattern intensities and fragment qualifier ions in combination with Retention Time.
- Multiple level MRSQ(C) scoring of results provides efficient reduction of false positive and negative results by simple default methods.
- Reliable quantitation and comprehensive screening possible using the same data sets.
- Unique addition of CCS values in the 4D mode.

Sample Screening and Quantification

Analysis: Bruker CEC 8_50 ng/mL_timsON b6CID POS_45_1_8541

Creation Date: 2023-04-03 15:31 Sample Type: Calibrant
 Method: Carved Demo_240 Pesticides Mass Calib. Date: 2023-03-02 14:39
 +CEC_timsTOP POS (4)

Station Name: Default Operator: Demo User
 Instrument: timsTOP Pro Instrument SN: 1854399.122
 Data Path: E:\Data\Demo Data\CEC Dilution Series_POS_timsON (TASQ)\Bruker CEC 8_50 ng/mL_timsON b6CID POS_45_1_8541.d

Analysis	m/z exp.	Δ m/z (ppm)	Δ RT (min)	Mand. ions	Quantity	Chromatogram
Azinphos-methyl Fragment 143	143.0627	2.62	-0.03	2/2		
RT theo [min]	m/z Score	RT Score	ions Score	RT		
6.30	***	***	***			
Azinphos-methyl Fragment 160	132.0444	2.19	-0.11	3/3		
RT theo [min]	m/z Score	RT Score	ions Score	RT		
9.81	***	***	***			
Boscalid 4	362.1154	0.36	0.16	3/3	55.30 ng/mL	
RT theo [min]	m/z Score	RT Score	ions Score			
7.83	***	***	***	0.9965		
Chloridazon 4	222.0429	-0.98	-0.02	4/4	38.32 ng/mL	
RT theo [min]	m/z Score	RT Score	ions Score			
5.11	***	***	***	0.9992		
Chlorthal-methyl 4	213.0799	0.75	-0.06	2/3		
RT theo [min]	m/z Score	RT Score	ions Score	RT		
7.41	***	***	***			
Chlorpyrifos 4	291.0895	-1.57	-0.10	2/3		
RT theo [min]	m/z Score	RT Score	ions Score	RT		
9.40	***	***	***			

Bruker April 25, 2023 1 / 7

Sample to report just in a few steps



Kristoffer Kilpinen

Development chemist, Eurofins Environment, Denmark

"After using TargetScreener for more than two years, we have found it to be a valuable tool for conducting wide-range suspect screenings of environmental samples."

TargetScreeners HR – Meeting the demands



Food safety



Dioxins



Pesticides



Doping



Veterinary drugs



Mycotoxins



PFAS



Drugs of abuse



Homeland security

Bruker Applied Mass Spectrometry is continually improving its products and reserves the right to change specifications without notice. © BDAL 06-2023, 1900369

For Research Use Only. Not for use in clinical diagnostic procedures.

Bruker Switzerland AG

Fällanden · Switzerland
Phone +41 44 825 91 11

marketing.bams.us@bruker.com - www.bruker.com

Bruker Scientific LLC

Billerica, MA · USA
Phone +1 (978) 663-3660

Online information

bruker.com

