PDA Training Course Extractables & Leachables
19-20 October 2023

Analytical techniques used in E&L studies

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Overview

- Analysis of extractables & leachables is a challenge!
- Analytical techniques for organic compounds
- Analytical techniques for inorganic compounds
- Screening methods vs validated methods
- Structural elucidation





The diverse world of extractables

Why the analysis of E&L is a challenge...





Diversity in CCS

Broad spectrum of:

- Types of Containers
- Types of Materials used in the Manufacture of Containers
- Number of Suppliers per Material
- Number of Grades (per supplier) for each type of Material
- Type of Sterilization (impact on material impurity profile)





Pharmaceutical CCS

INHALATION

- Metered Dose Inhaler Components e.g.:
 - Gaskets
 - Stem
 - Body
 - Metering Chamber
 - Protection Ring
 - Actuator
 - Canister
- o Dry Powder Inhaler Components
- Nasal Spray Systems
- o Nasal Dropper Systems
- o Blow-Fill Seal containers
- Nebulizers

0...

OPHTHALMIC

- Eye Dropper Systems
- o Tubes
- o Blow-Fill-Seal containers

0 ...

PARENTERAL

- o Bottles
- o Vials
- o (Pre-Filled) Syringes
- Cartridges
- o (Rubber) Stoppers
- o Rubber Plungers
- Sealing Discs
- o Needle Shields
- Tip Caps
- o I.V. Bags
- o Administration Sets
- 0 ...

DERMAL/TOPICAL

- Spray Systems
- Tube systems
- 0 ...

SINGLE USE SYSTEMS

- o (Multilayer) Bags
- Tubings
- Connectors
- o Ports
- Filters (+ Housing)
- Chromatographic Columns
- Lyo trays
- 0 ...

SECONDARY PACKAGING

- Labels
- Adhesive/Glue (e.g. on labels)
- o Ink
- Overwrap foils
- o Blisters
- Cardboard packaging
- 0 ...





Materials of construction for CCS

- Low Density Polyethylene
- High Density Polyethylene
- Polypropylene
- o Rubbers
- Butyl Rubbers
- Chlorobutyl Rubbers w/o Coating
- o Bromobutyl Rubbers w/o Coating
- EPDM Rubbers
- o Isoprene Rubbers
- Nitrile Rubbers
- Latex Rubbers
- Other Rubbers
- Multi-layer Films and Foils
- Polyurethane (PU)
- Ethylvinyl Acetate (EVA)
- Ethylvinyl Alcohol (EVOH)

- Polyamide (Nylon-6, Nylon-66)
- Cyclic Olefin Copolymers (COC)
- Cyclic Olefin Polymers (COP)
- Polyethylene Terephthalate (PET, PETG)
- Polybutylene Terephthalate (PBT)
- Polyacetal (POM)
- Polymethylmethacrylate (PMMA)
- Acrylonitrile Butadiene Styrene (ABS)
- Silicone
- Thermo Plastic Elastomers (TPE's)
- Polycarbonate
- o PTFE
- o PEEK
- Glass w/o Coating
- Metals
- 0...





Suppliers for a given material

Polyethylene - produced by:

- o Borealis
- LyondellBasell
- o SABIC
- Dupont
- o Enichem
- o INEOS
- o TOTAL
- 0 ...

Pharmaceutical Rubbers – main Global Suppliers:

- o Datwyler
- West Pharmaceutical
- o Stelmi

Each supplier has different (health care) grades!





Each supplier: different grades

Polyethylene (PE) - produced by:

- o Borealis: over 30 different Medical Grades
- o LyondellBasell: over 30 different Medical Grades
- o SABIC: over 30 different Medical Grades
- Dupont: different grades
- o Enichem: different grades
- INEOS: different grades
- o TOTAL: different grades

0 ...

Pharmaceutical Rubbers - main global suppliers:

- o Datwyler: over 100 different commercial rubber formulations
- West Pharmaceutical: over 100 different commercial rubber formulations
- o Stelmi: also, a broad range of commercial rubber formulations





Impurity profile of 1 grade

INTENTIONALLY ADDED

- o Pigments / colorants
- Clarifying agents
- Catalysts and Curing Agents
- Fillers
- Anti-oxidants
- Plasticizers
- Photostabilizers
- Slip agents
- Acid scavengers
- o ...

NON-INTENTIONALLY ADDED

- Related to the Polymer
 - Polymer Degradation Compounds
- Related to the Polymerization Process
 - Solvent residues
 - Monomers
 - Catalysts
 - Oligomers
- Related to the additives
 - Additive degradation compounds
- Related to secondary packaging
 - Glue, Labels, Carton/Paper
- Processing Impurities
 - Lubricants, surfactants, solvents

0 ...





Conclusion: diverse chemistry!

PHYSICO-CHEMICAL PROPERTIES OF EXTRACTABLES

UNIVERSE OF EXTRACTABLES: 10.000 – 100.000 compounds

Analytical method: identification and quantification

COMBINATION OF ANALYTICAL TECHNIQUES REQUIRED

- For routine screening: labs need to be cost-effective
- Only possible with extensive material knowledge & databases





Sample Preparation

The most important & most underestimated activity in the lab





Trace analysis is a challenge

- o Have very experienced people in sample preparation team
- o Very intensive training for new staff in sample prep team
- o QC on solvents used select batches of clean solvents with suppliers
- QC on extraction equipment
- Separate glassware
- Precleaning of glassware validation of cleaning procedures
- Sampling of test articles how to handle test articles?
- o **UPW sample prep** should be **separated** from solvent sample prep
- Orrection for absorbed solvents?
- How to concentrate extracts while avoiding cross contaminations
- Storage of extracts under controlled conditions
- Holding times of extracts
- Selection of type of containers for storage of extracts
- O How to keep **DEHP** out of the Lab!









Organic compounds

Chromatography – Mass Spectrometry





Chromatography - Mass Spectrometry

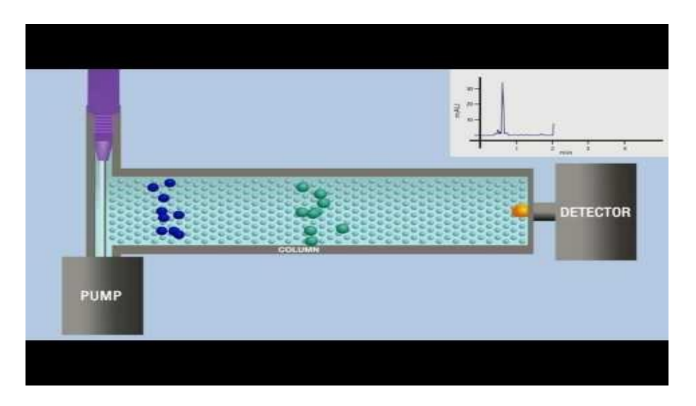
- Complex mixture of compounds!
- Analysis is 2-step process:
 - Separation
 - Detection (+ structural information of detected compound)
- Chromatography:
 - Separation technique
 - Involves 2 'phases': stationary phase + mobile phase
- Mass Spectrometry:
 - Detection technique hyphenated to the chromatography system
 - Mass information of detected compounds





Chromatography - Mass Spectrometry

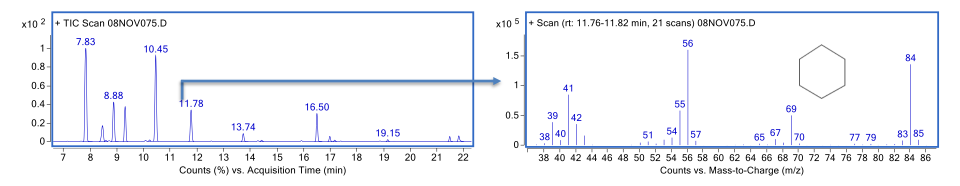
Video animation on chromatography separation principle







Chromatography – MS output



Chromatogram

- Analytical output from chromatography system
- Detector signal intensity in function of analysis time
- Compound separation
- Retention time → depends on compound properties
- Peak area → measure of quantity

Mass spectrum

- Analytical output from mass spectrometer
- Compound detection, but does more!
- Mass (fragment) information for each peak in chromatogram
- Very powerful tool for identification





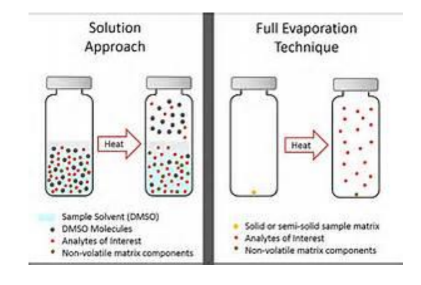


Volatile Organic Compounds (VOC)

Headspace – Gas chromatography – Mass Spectrometry (HS-GC/MS)



- Monomer residues
- Solvent residues from production steps
- Residues from polymer treatments
- Small polymer degradation products



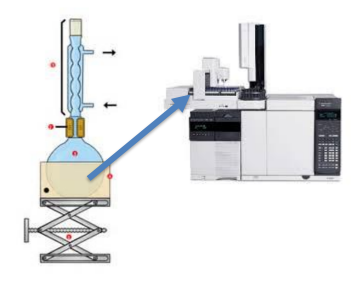




Semi-Volatile Organic Compounds (SVOC)

Gas chromatography – Mass Spectrometry (GC/MS)

- Lubricants
- Plasticizers
- Antioxidants
- Polymer degradation products
- Solvents with an elevated boiling point







GC

SEPARATION of (semi-)volatile organic compounds (Mw < 650 Da)

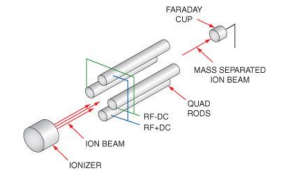
- Gas phase separation technique using narrow open tubular columns coated with a film of stationary phase, mounted in temperatureprogrammable oven
- Separation of compounds based on boiling point and polarity owing to variations in affinity with the stationary phase
- A higher film thickness of stationary phase increases retention:
 - VOCs: high film thickness (≥ 1 µm)
 - SVOCs: low film thickness (≤ 0.25 µm)
- Length of capillary column increases resolution (but increases analysis time as well)
- Not well suited for polar compounds like acids, amines, diols... Where specific conditions may need to be applied





MS (coupled to GC)

DETECTION & MASS-BASED SEPARATION



- 3 events: ionization / mass separation / detection all happening under high vacuum
- o Ionization: electrion ionization (70 eV) → convert molecule into ion and induce further fragmentation
- Quadrupole mass analyzer:
 - Scanning mass filter → only 1 mass can pass through a given electric field
 → other masses are removed
 - By rapidly sweeping the electric field → scanning of a mass range
 - Scanning goes extremely fast: milliseconds
 - Ions that reach the detector induce a signal that is measured
 - Mass spectrum: bar-graph plot of signal intensity vs. mass (unit)
 - Multiple mass spectra are recorded each second of the analysis (~ 3 scans/second)

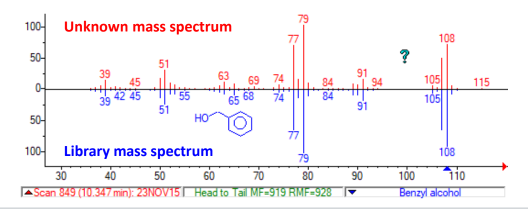




GC/MS spectrum

GC/MS spectra are "standardized"

- Most GC/MS instruments for routine use make use of electron ionization single quad technology
- Electron ionization (and associated molecule fragmentation) is a very reproducible event
 - → Reproducible mass spectra are obtained across different instruments across the world
- Obtained mass spectra can be compared to commercial databases or in-house databases
 - → In case of a good match may lead to identification of the compound







Non-Volatile Organic Compounds (NVOC)

Ultra Performance Liquid chromatography – Mass Spectrometry (UPLC/MS)

- Fillers
- Plasticizers
- Antioxidants
- Anti-slip agents
- Oligomers







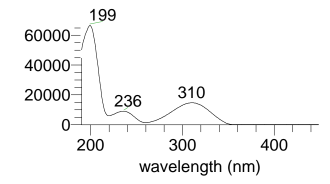
UPLC

- Separation technique suited for non-volatile organic compounds (NVOCs)
- Liquid-based separation technique using columns packed with stationary phase
- Using high pressure to pump sample dissolved in mobile phase through packed column
- Separation of compounds based on affinity for the stationary phase
 - Polar stationary phases: straight phase chromatography
 - Apolar stationary phases: reversed phase chromatography (most used)
- Optimizing separation by
 - Selection of chromatographic column (length, polarity of stationary phase)
 - Selection of mobile phase (water, methanol, acetonitrile)
- o **Detection**:
 - Diode Array Detection (DAD using UV spectrum)
 - (high resolution accurate mass) Mass Spectrometry (primary choice)





DAD/UV detector



Advantages:

- Standard equipment in analytical lab
- Low cost
- UV detection simultaneous with MS detection: can be used as add-on detector

Disadvantages:

- Not universal / generic (chromophore needed for detection)
- Limited sensitivity, depending on chromophore(s)
- Poor specificity, even for Diode Array Detectors (scanning UV)
 - → Information about detected molecule is limited (e.g. link with API?)





MS (coupled to LC)

Advantages:

- Increased specificity: (exact) mass
- Increased sensitivity
- Mass spectra may reveal more information about the identity of the compound
- Allows for building (in-house) mass spectral databases

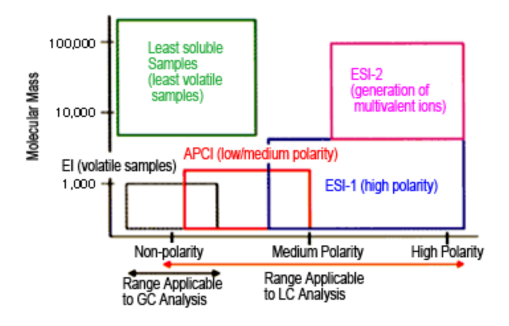
Disadvantages:

- Higher cost
- Contrary to GC/MS: no universal spectra (depends on ion source design, mobile phase, MS settings, ...) → no universal libraries!
- Need for multiple ionization methods to allow a broader range of target compounds for UPLC/MS





Ionization vs Compound Range



- Electron Ionization: only works in gas phase under vacuum → not LC compatible
- Atmospheric Pressure Chemical Ionization: LC up to medium polarity
- ElectroSpray Ionization: LC medium polarity high polarity
 Nowadays: more and more both APCI & ESI in E&L study design





Modern LC/MS instrumentation

Older systems:

- Quadrupole or ion trap (cf. GC/MS)
- Low resolution: unit mass e.g. m/z 220 can be distinguished from m/z 221

Nowadays:

- Q-TOF or Orbitrap technology
- High resolution & mass accuracy (HRAM) e.g. m/z 220.000 can be distinguished from m/z 220.002
- High accuracy may allow determination of elemental formula when molecular ion is detected
- Extremely powerful technique in combination with UPLC when developing inhouse high resolution MS databases in combination with retention time of reference compounds
- Contrary to GC/MS, UPLC/HRAM-MS is used in "first-pass" screening to compensate for the lack of mass spectral fingerprinting and availability of commercial databases like in GC/MS

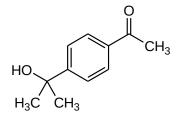




Modern LC/MS instrumentation

LC-QUADRUPOLE (LOW RESOLUTION)

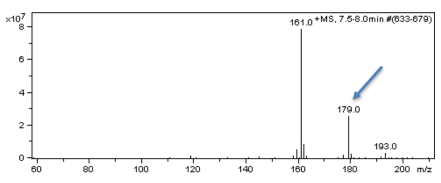


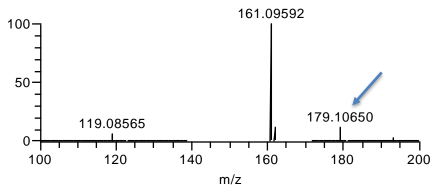


Peroxide curative related compound from EPDM rubber **Exact mass: 179.10666**

LC-ORBITRAP (HRAM)







Mass error: 0.16 mDa or 1 ppm





Inorganic Compounds

Analytical Techniques





Elements

Inductively Coupled Plasma / Optical Emission Spectroscopy or MS



Origin of elements

- Metals from glass
- Metals from rubbers
- Catalysts, used during polymerization process
- Fillers, added to polymer materials
- Acid scavengers
- Activators for rubber polymerization



Technique

- ICP to produce excited atoms
- Excited atoms recombine, giving off electromagnetic radiation at wavelengths characteristic for each element
- o Emission wavelengths detected by the spectrophotometer
- Or ions detected by mass spectrometry
- o Intensity correlates to concentration → quantitative technique



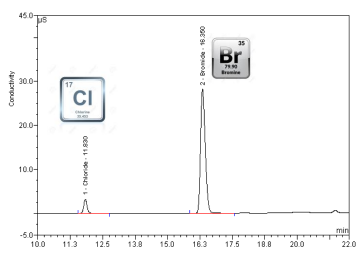


Anions

Ion Chromatography (IC)

Origin of anions

- Polyolefins: formate / acetate as oxidation products
- Halobutyl rubbers: bromide, chloride, fluoride
- o Fluoropolymers: fluoride
- Trace impurities: nitrite, nitrate, phosphate, sulfate



Example: UPW extract of a halobutyl rubber

Technique

- Special liquid chromatography technique
- Designed for separation and detection of ions
- <u>Detection</u>: conducitivy or amperometry





Other specific analytical methods

- GF-AAS for silicone oil detection and quantification
- HPLC-UV for TMPTMA (glue residue)
- HPLC-UV for S₈ (cross-linker)
- pH (release of acidic/alkalinic agents in UPW)
- Conductivity (release of salts in UPW)
- Non-Volatile Residue (gravimetric residue after evaporation of extract)
- FTIR characterization of NVR
- Total Organic Carbon: reconcilliation with concentration of organic compounds from chromatographic techniques
- 0 ...





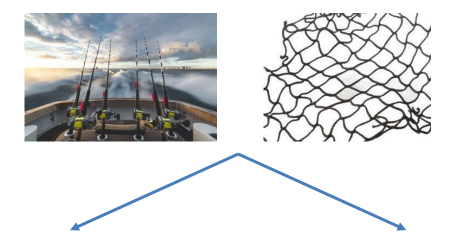
Screening | Discovery

Identification & Quantification





Different fishing techniques











Different analytical techniques



IDENTIFICATION

CAS No 2 to 7 digits 2 digits 1 check digit



QUANTIFICATION



Concentration

μg/L

μg/unit μg/g





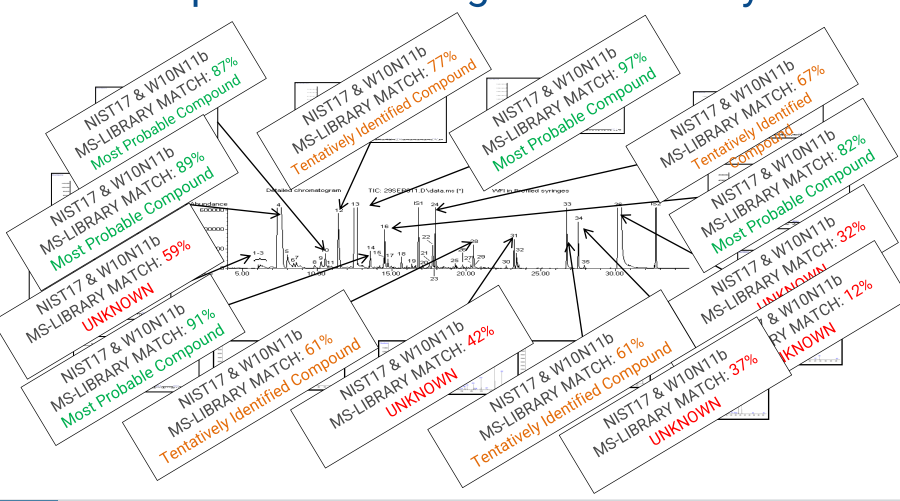
Concept of "screening" or "discovery"

- Untargeted analysis mode used in extractables studies (organic comp)
- Trying to IDENTIFY every peak in a chromatogram...
- o ... above a certain threshold:
 - Either based on analytical feasibility (reporting threshold)
 - Or based on toxicological threshold (e.g. AET)
- Generate a list of extractables from the tested material with focus on identification
- Screening is estimated or semi-quantitative: estimation of concentration
- Useful for follow-up in a leachables study





Concept of "screening" or "discovery"







Quantification in screening

Screening is untargeted → no prior knowledge about extractables / leachables profile

In case many extractables reported → accurate quantification for all is not practically feasible

Estimated quantification

- o Internal standard (I.S.) compound spiked to each (final) extract
- Assumption that response of analyte = response of I.S. (response factor = 1)
- Accounts for instrument variation
- Does not account for different response vs I.S. or liquid/liquid recovery

Semi-quantitative quantification

- o Internal standard (I.S.) compound spiked to each (final) extract
- \circ Record analytical response of standard vs response of I.S. \rightarrow relative response factor (RRF)
- Correct concentrations of confirmed ID's with RRF
- Accounts for instrument variation + response variation of analyte vs I.S.





Validated Methods

For accurate quantification



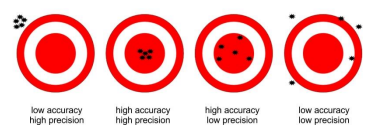


Validated methods

- Chromatography Mass Spectrometry instrumentation more or less the same
- Except: triple quadrupole (QqQ) instead of single quadrupole (selectivity + sensitivity)
- Validated methods are targeted → leachables to be quantified are a priori known
- Methods are specifically developed and optimized for the target leachables

Validated quantification

- Specific internal standard for each target leachable
- Quantitative performance of method is validated:
 - Selectivity / Specificity → no interference from blank signal, drug matrix, other leachables...
 - Limit of detection / Limit of quantification → lowest concentration level for accurate quant
 - Linear range → concentration range validated for accurate quantification
 - Precision → variability of analytical method
 - Accuracy → closeness to true value







Structural Elucidation

High-end Mass Spectrometry





Structural elucidation - Introduction

- Unknown / Partially identified compounds > AET in 1st pass screening
 - Unknowns are treated as carcinogenic/mutagenic
 - To allow de-risking by tox assessment, a structure is needed!
- Request to further increase ID level (e.g. low margin of safety)
 - Tentative to Confident
 - Confident to Confirmed (standard should be available or synthesized)
- Goal of identification studies: generate / collect comprehensive set of supporting data to increase the identification level of a target compound





Structural elucidation - Instrumentation



Liquid Chromatography

- Orbitrap
- FT-Ion Cyclotron Resonance

Requirements

- High-end mass spectrometers
- o (Very) high resolution
- High mass accuracy
- Multiple ionization methods
- Tandem mass spectrometry

Gas Chromatography

- Q-TOF
- Orbitrap







Structural elucidation - HRAM

Element	Nominal Mass	Exact Mass
Hydrogen (H)	1	1.0078
Carbon (C)	12	12.0000
Nitrogen (N)	14	14.0031
Oxygen (O)	16	15.9949



Nitrogen gas: N₂

Nominal mass: 28 Da Exact mass: 28.0062 Da



Carbon monoxide: CO

Nominal mass: 28 Da Exact mass: 27.9949 Da



Difference: 0.0113 Da

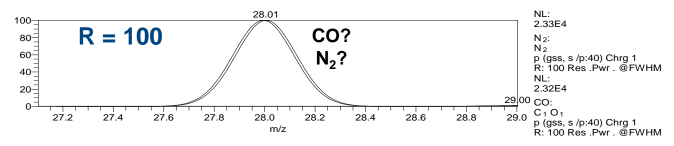




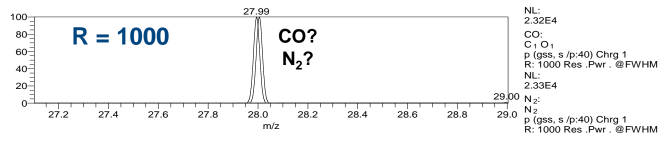
Structural elucidation - HRAM

Misidentification of a compound with a mass of 28 Da can be fatal... how to be sure?

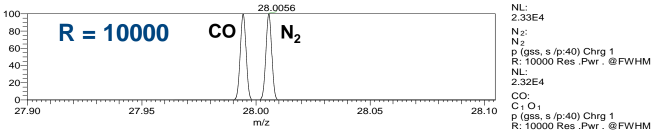
Not separated



Not separated



Separated





NL:

2.22E4

2.21E4 C₅ H₁₀ O₂:

C₅ H₁₀ O₂

p (gss, s /p:40) Chrg 1

R: 100000 Res .Pwr . @FWHM

C4 H₁₀ N₂ O:

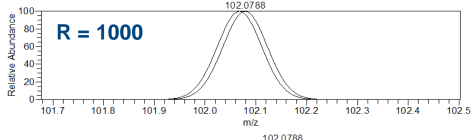
C₄ H₁₀ N₂ O₁ p (gss, s /p:40) Chrg 1 R: 1000 Res .Pwr . @FWHM



Structural elucidation - HRAM

E&L example: 2 compounds where both have nominal mass 102...

Not separated

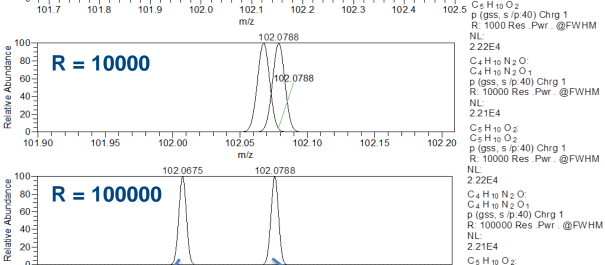


102.07

Close...



Separated



102.08

m/z

 $C_5H_{10}O_2$ - isopropyl acetate

102.06

102.05

 $C_4H_{10}N_2O$ - N-nitrosodiethylamine

102.09





HRAM – Important take-aways

accurate mass alone does not deliver a structure...

... but delivers the elemental formula of the molecule and fragments of the molecule

high resolution does not deliver a structure...

... but enables to **separate molecules** with the same nominal mass but different elemental formulas

...but assists in confirming the elemental formula using isotope matching

Mass spectral interpretation skills and expertise are required

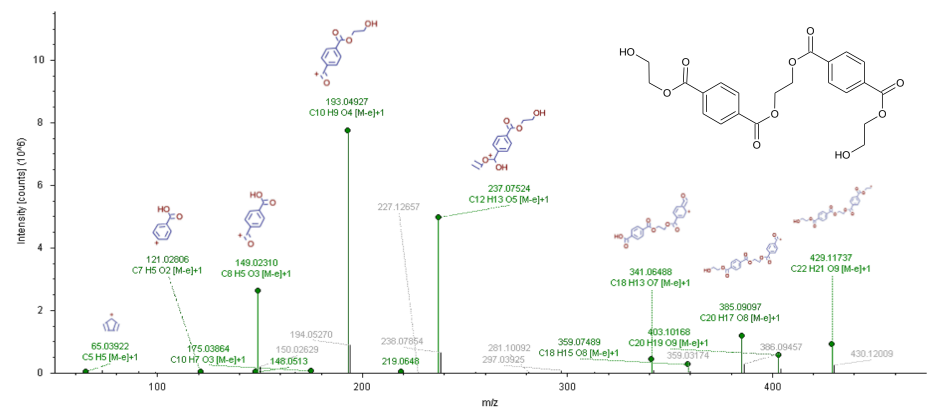






Tandem Mass Spectrometry (MSⁿ)

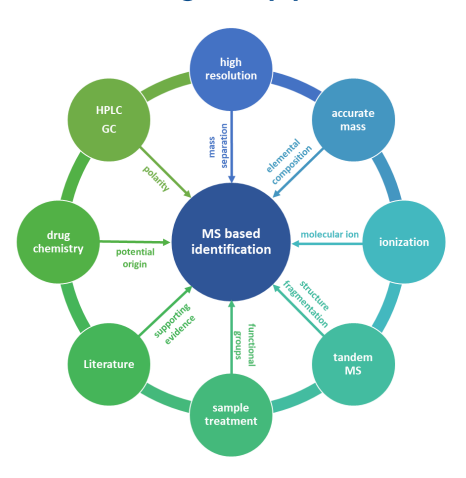
- 1. Select molecular ion & induce fragmentation
- 2. Measure all molecule fragments with HRAM







Multi-angle approach required



TEAM effort!

Mass spec expert(s)

Drug chemistry expert

Material engineer







