

Knowledge that will change your world

Choosing the metabolomics platform

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T argeted

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M etabolomics & P roteomics

L aboratory

1

Synopsis

- The analytical challenge
- · Early days of metabolomics
- Platforms
 - TLC
 - GC; GC x GC
 - GC-MS
 - LC-MS
- Mass analyzers
- What is mass?
- The LC
 - · Selected ion monitoring
 - Multiple Reaction ion Monitoring
- Ion mobility
- Imaging-MS

Challenges

- Unlike DNA, RNA and proteins, the metabolome is phenomenally chemically diverse
- Ranges from a gas (H₂) that prevades the universe and is the principal component of the Sun

to





- Earwax (long chain fatty acids, both saturated and unsaturated, alcohols, squalene, and cholesterol)
- No single method of analysis

3

Early beginnings of metabolomics in London

- Sir Ernst Chain (1945 Nobel Laureate - the biochemist who characterized penicillin)
 - · Also renown for his work on microanalysis
- Used 2D-paper chromatography to resolve glycolytic, Krebs cycle and amino acids derived from ¹⁴C-glucose
 - o Geiger counter mounted on a typewriter frame
 - o Digitized the collected data and prepared computer-generated figures



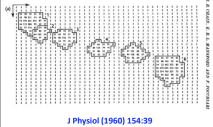




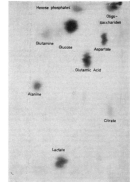


METABOLOMICS

Radiochromatography examples



J Physiol (1960) 154:39 E.B. Chain, K.R.L. Mansford and F. Pocchiari

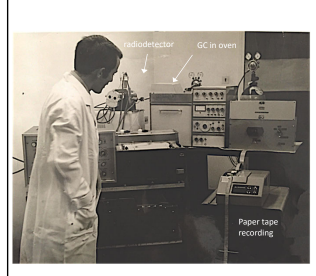


Autoradiogram of ¹⁴C-glucose metabolites from an isolated perfused Langendorff rat heart preparation. The metabolites were separated by 2D-paper chromatography.

The conditions were: 1st dimension: butan-l-ol-acetic acid-water (40:11:25, by vol.)

2nd dimension: (-) phenol-aq. NH₃ (sp.gr. 0.88)-water (80:1:20, by vol.) for 24hr.

Biochem. J. (1969) 115, 537 E.B. Chain, K.R.L. Mansford and L.H. Opie

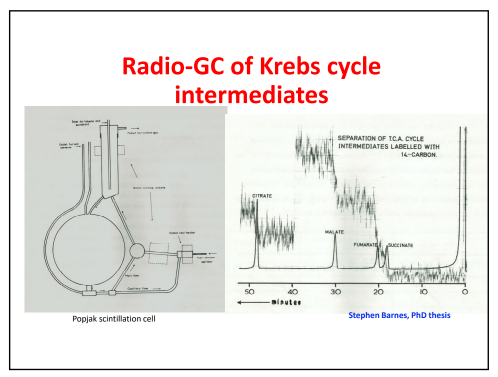


Radio-GC analysis

metabolomics in its infancy

Radio gas-liquid chromatography with digitization of collected data

Developed this for my PhD work (1967-1970) to study glucose metabolism in acellular slime mold, Physarum polycephalum



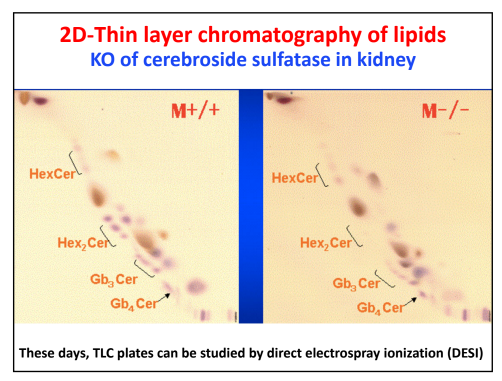
Software for data analysis on a PDP9 computer

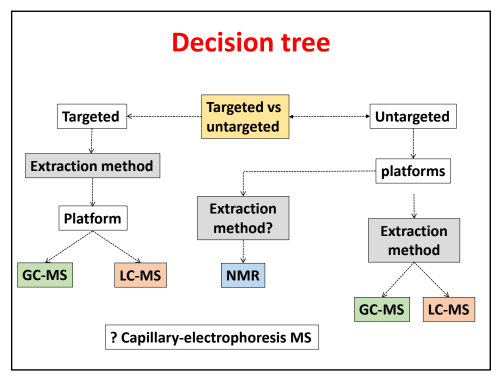




Punched tape data 1 data point/sec

Digital PDP computer Had a screen





Metabolomics and GC-MS

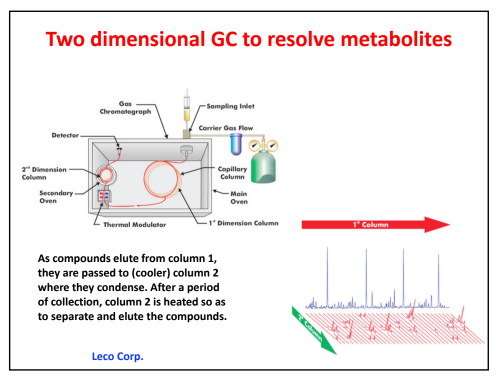
PROS

- Capillary columns can achieve very high chromatographic resolution
- Retention times are reproducible
- Mass spectral libraries are well developed

CONS

- Not all compounds can be analyzed by GC-MS
- Although amino acids, sugars, fatty acids, amines and organic acids can be derivatized, complex polyphenol glycosides and polar lipids are too unstable, even when derivatized, at the temperatures used to elute them
- Approximate mass limit of 400 Da

11



Nuclear Magnetic Resonance (NMR) Spectroscopy

- Detects NMR active nuclei
- Robust and highly reproducible
- Non-destructive
- Quantitative
- Used in
 - Structure elucidation
 - · Small molecules
 - Macromolecules (DNA, RNA, Proteins)
 - A number of techniques
 - 1D, 2D, 3D
- Molecular motion and dynamics



from Wimal Pathmasiri









NMR considerations

• Sample amount:

- Typical 600 MHz instrument requires 0.5 ml plasma/serum
- Higher field instrument and micro coil detector allows use of 0.1 ml

Quality control:

- In the UK Phenome Center, all samples are analyzed by NMR
 - · This allows for detection of outliers
 - Also found that there is a correlation between the NMR spectrum and whether problems occur in LC-MS analysis
 - NMR analysis used to filter out these samples

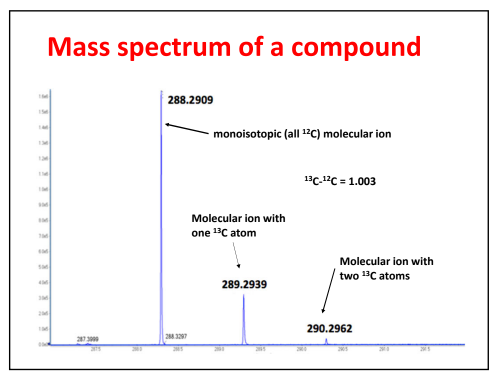
Hyperpolarization NMR

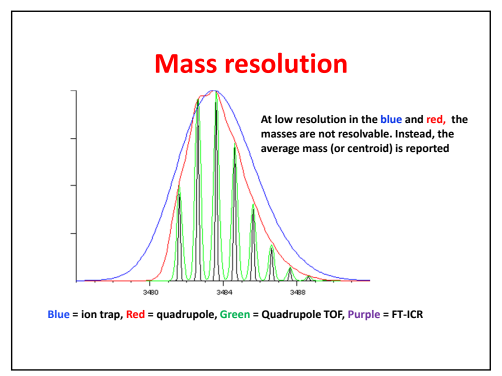
- The NMR signal comes from the not-quite, nonequilibrium between the two or more energy states a nuclei experience in a strong magnetic field
 - The natural excess population of the higher energy states is no more than 0.01%
 - This accounts for the low sensitivity of NMR
- By hyperpolarizing the compound, the excess population can be increased by 10⁴-10⁵.
 - · Much increased sensitivity
- Carbonaceous materials (metabolites) can be hyperpolarized by cooling to 1°K in a strong magnetic field (3 T or larger)
 - However, the lifetime of the hyperpolarized state is quite short (10-30 s) making metabolomics experiments quite difficult

15

Liquid chromatography-Mass Spectrometry

- PROS
 - Almost all compounds can be analyzed by LC-MS
 - · Exceptions hydrocarbons do not ionize
 - Several orders of magnitude increased sensitivity compared to NMR
 - Can collect MS, MSMS and ion mobility data
- CONS
 - Not uniformally quantitative
 - Mass spectral libraries are not well enough developed
 - Chromatographic separation not adequate
 - Retention time reproducibility not as good as GC-MS



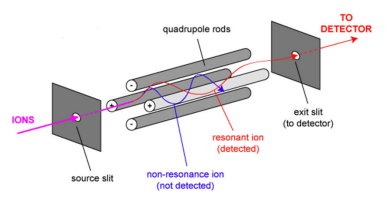


Selecting the mass spectrometer

- It is necessary to use an instrument to measure:
 - The mass of the metabolites accurately
 - To provide sufficient mass resolution to distinguish the isotopes associated with each metabolite
- There are several types of MS detectors
 - Quadrupole
 - ion trap
 - time-of-flight (TOF)
 - Orbitrap
 - Fourier Transform-Ion Cyclotron Resonance (FT-ICR)

19

Quadrupole mass filter

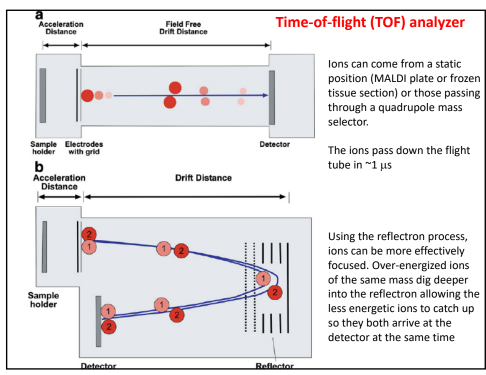


Consists of four parallel rods. Each opposing rod pair is connected together electrically, and a radio frequency (RF) voltage with a DC offset voltage is applied between one pair of rods and the other. This causes the ions to rotate in spirals as they go through the quadrupole. For a given voltage, only ions of a specific m/z can pass through. The voltage can be scanned to generate a mass spectrum or held constant to allow one ion to pass through.

The mass spectrometer

- For untargeted analysis it is important to have high mass resolution, accuracy and speed
 - Initial data analysis is performed on the molecular ions
 - Each metabolite has a unique mass (m/z)
 - Nonetheless, a particular mass, however exact, is not necessarily a unique metabolite
- Fourier transform-ion cyclotron resonance and Orbitrap instruments have the greatest mass accuracy
 - However, their performance is time-dependent and is degraded significantly using short acquisition times (<100 ms)
 - They are best used for follow up experiments

21



TOF is the mass analyzer of choice for untargeted metabolomics

Quadrupole-orthogonal time-of-flight (Q-tof)



Current models have 30-80,000 mass resolution and 1 ppm or better mass accuracy

23

Links to the different Q-TOFs

- Agilent 6560 Ion Mobility Q-TOF LC/MS
 - https://www.agilent.com/cs/library/brochures/5991-3640EN 6560 Ion Mobility QTOF LCMS Brochure Final singlepgs.pdf
- Bruker TimsTof
 - https://www.bruker.com/products/mass-spectrometry-and-separations/lc-ms/o-tof/timstof/overview.html
- SCIEX 6600+ TripleTOF/SelexION
 - https://sciex.com/products/mass-spectrometers/qtof-systems/tripletof-systems/tripletof-6600-system
- Waters Synapt G2Si
 - http://www.waters.com/waters/en_US/SYNAPT-G2-Si-High-Definition-Mass-Spectrometry/nav.htm?cid=134740622&locale=en_US

Masses of elements and their isotopes

- Mass is defined using the mass of carbon-12 being 12.0000 (exactly) – the others have non-integer mass defects
- · On this scale,
 - ¹H is 1.007825 and ²H is 2.014102 (extra neutron)
 - ¹⁴N is 14.003074 and ¹⁵N is 15.000108 (extra neutron)
 - 16O is 15.994915, 17O is 16.999132 and 18O is 17.999161
 - ³¹P is 30.973761
 - 32S is 31.972071 and 34S is 33.967867 (4%)
- You can find the mass of every element and its isotopes and their natural abundances at

http://www.nist.gov/pml/data/comp.cfm

• The mass of a proton is 1.0072766 and that of an electron is 0.000548597

25

Predicted mass defects for C_xH_nO_m

H atoms	O=0	Oxygen atoms									
		O=1	O=2	O=3	O=4	O=5	O=6	O=7	O=8	O=9	O=10
1	0.0078	0.0027	-0.0023	-0.0074	-0.0125	-0.0176	-0.0227	-0.0278	-0.0329	-0.0379	-0.0430
2	0.0157	0.0106	0.0055	0.0004	-0.0047	-0.0098	-0.0149	-0.0199	-0.0250	-0.0301	-0.0352
3	0.0235	0.0184	0.0133	0.0082	0.0031	-0.0020	-0.0070	-0.0121	-0.0172	-0.0223	-0.0274
4	0.0313	0.0262	0.0211	0.0160	0.0110	0.0059	0.0008	-0.0043	-0.0094	-0.0145	-0.0196
5	0.0391	0.0340	0.0290	0.0239	0.0188	0.0137	0.0086	0.0035	-0.0016	-0.0066	-0.0117
6	0.0470	0.0419	0.0368	0.0317	0.0266	0.0215	0.0164	0.0114	0.0063	0.0012	-0.0039
7	0.0548	0.0497	0.0446	0.0395	0.0344	0.0294	0.0243	0.0192	0.0141	0.0090	0.0039
8	0.0626	0.0575	0.0524	0.0473	0.0423	0.0372	0.0321	0.0270	0.0219	0.0168	0.0117
9	0.0704	0.0653	0.0603	0.0552	0.0501	0.0450	0.0399	0.0348	0.0297	0.0247	0.0196
10	0.0783	0.0732	0.0681	0.0630	0.0579	0.0528	0.0477	0.0427	0.0376	0.0325	0.0274
11	0.0861	0.0810	0.0759	0.0708	0.0657	0.0607	0.0556	0.0505	0.0454	0.0403	0.0352
12	0.0939	0.0888	0.0837	0.0786	0.0736	0.0685	0.0634	0.0583	0.0532	0.0481	0.0430
13	0.1017	0.0966	0.0916	0.0865	0.0814	0.0763	0.0712	0.0661	0.0610	0.0560	0.0509
14	0.1096	0.1045	0.0994	0.0943	0.0892	0.0841	0.0790	0.0740	0.0689	0.0638	0.0587
15	0.1174	0.1123	0.1072	0.1021	0.0970	0.0920	0.0869	0.0818	0.0767	0.0716	0.0665
16	0.1252	0.1201	0.1150	0.1099	0.1049	0.0998	0.0947	0.0896	0.0845	0.0794	0.0743
17	0.1330	0.1279	0.1229	0.1178	0.1127	0.1076	0.1025	0.0974	0.0923	0.0873	0.0822
18	0.1409	0.1358	0.1307	0.1256	0.1205	0.1154	0.1103	0.1053	0.1002	0.0951	0.0900
19	0.1487	0.1436	0.1385	0.1334	0.1283	0.1233	0.1182	0.1131	0.1080	0.1029	0.0978
20	0.1565	0.1514	0.1463	0.1412	0.1362	0.1311	0.1260	0.1209	0.1158	0.1107	0.1057

For positively charged ions, add **1.007276** to the overall m/z value For negatively charged ions, subtract **1.007276** from the overall m/z value

Empirical formula

If the mass of an ion is known accurately enough, then it is possible to write down its empirical formula

27

What is the mass of a metabolite?

Hexanol

 $C_6H_{14}O$ = 6*12.0 + 14*1.007825 + 15.994915 = 102.1044651

Glucose

 $C_6H_{12}O_6$ = 6*12.0 + 12*1.007825 + 6*15.994915 = 180.063388

Masses of genistein's ions

• Genistein, C₁₅H₁₀O₅

Mass = 15*12.0 + 10*1.007825 + 5*15.994915

 $[M+H]^+ = M + 1.00727638 = 271.060073$ $[M-H]^- = M - 1.00727638 = 269.045547$

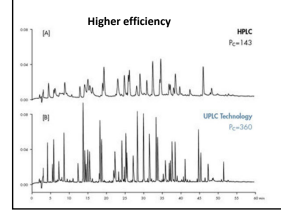
- If glucose is joined to genistein and water (H₂O) is eliminated, what are the values of the [M+H]⁺ ion and the [M-H]⁻ ion?
- Please send me the answer before class on Wednesday, Jan 29th.

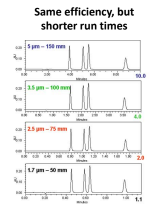
29

The LC

• 1D-approach

- Use of reverse-phase, normal phase and HILIC phase
- particle size smaller is more efficient, but back pressure is a problem





LC flow rate

- MS Sensitivity is inversely related to flow rate
 - Slower flow gives more sensitivity







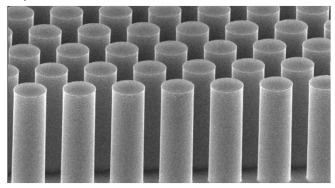
normal flow (0.2-0.4 ml/min)

nanoflow (0.3-5 µl/min)

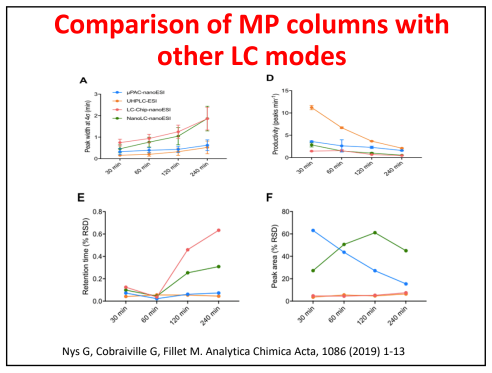
31

Improving the chromatography

• Micropillar columns



Non-turbulent fluid flow and reduced back pressure results in long-g-g columns



Optimizing nanoLC for metabolomics

- Objective is to develop metabolomics for small animal model systems
 - D. melangaster
 - C. elegans
 - D. rerio







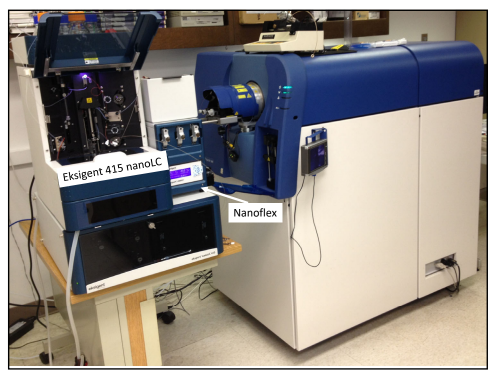
- A single zebrafish yields about 1 μ l of plasma
- Need to move down to the nanoscale
- Important to maintain consistency and quantitation
 - Reproducible columns and temperature

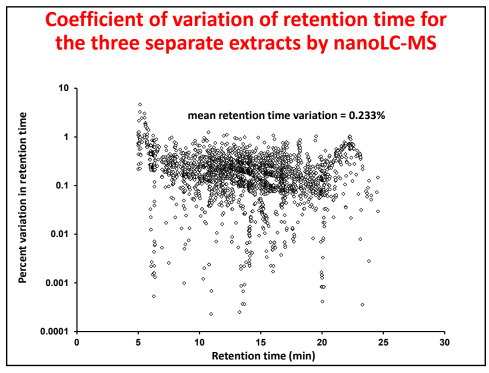


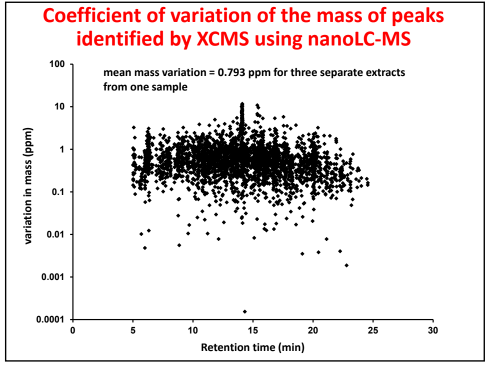
Close up of a nanochipLC cartridge (15 cm x 0.2 mm ID).

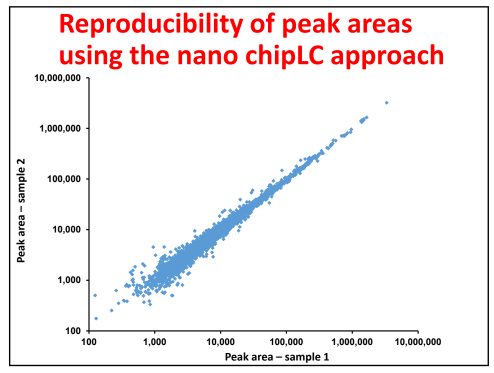
- Each long section of the column is ~2.5 cm (1 inch).
- Can be machined to a better tolerance.
- Simpler connections to the liquid stream.
- Can be placed in a temperaturecontrolled environment

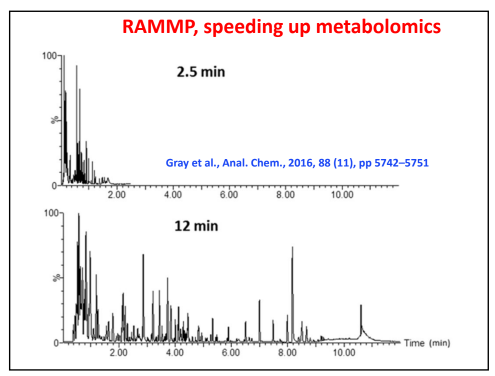
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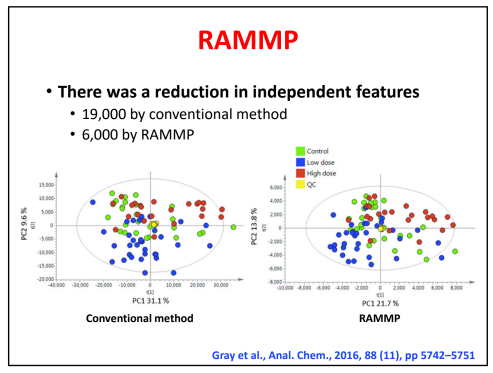






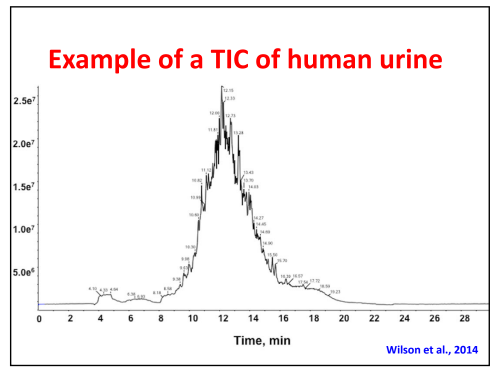


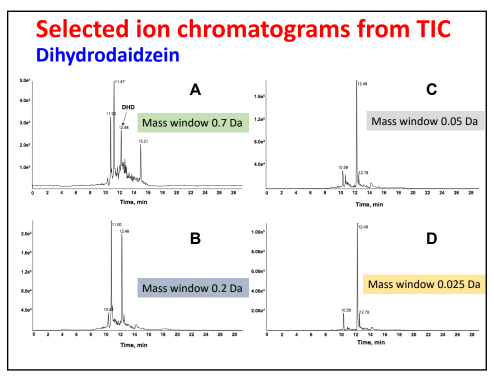


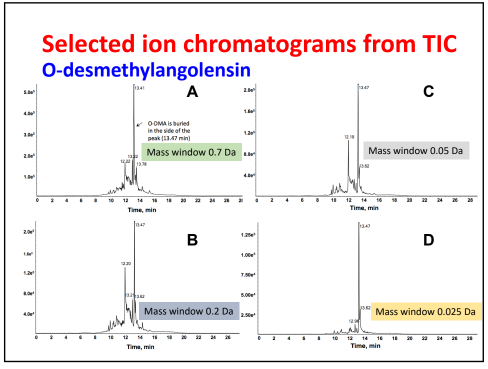


Selected ion monitoring

- The summation of all the ions collected in a GC or LC analysis is called the total ion current (TIC) and produces a total ion chromatogram
- By selecting a particular mass-to-charge ratio (m/z) value, one can see where a metabolite's molecular ion elutes from the column
 - This produces a selected ion chromatogram (SIC or XIC)
 - The quality of the SIC depends on the mass accuracy and resolution of the collected data

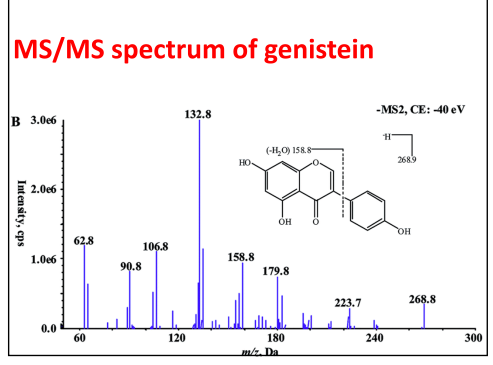






MS/MS

- A second mass spectrum (MSMS) that is informative arises from isolating the molecular ion
- The molecular ion is heated, either by collision with neutral gas (quadrupole, ion traps) or by using IR radiation (FT-ICR)
 - The extra energy increases the stretching of critical bonds, leading to dissociation of the molecular precursor ion into charged product ions
 - These generate the MS/MS spectrum for a metabolite
 - Ion traps can also isolate a product ion and create MSⁿ spectra



Measuring a mass transition

- Instead of measuring the full MS/MS spectrum, ions from the MS/MS can be individually measured
- This is referred to as a mass transition from the molecular or precursor ion to a specific product ion
- It is also known as reaction ion monitoring

Targeted vs untargeted methods

- If we know what the metabolites to be measured are (from previous untargeted analyses, or prior knowledge), then a multiple reaction monitoring (MRM) approach is the best way to go since allows quantitative analysis of possibly 100s of metabolites
- If there is no hypothesis, but instead you want to generate hypotheses, then the untargeted approach is better.

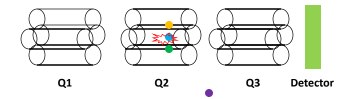
49

Multiple reaction ion monitoring



Ionizer

Quantitative analysis of metabolites in a complex mixture carried out using a triple quadrupole instrument

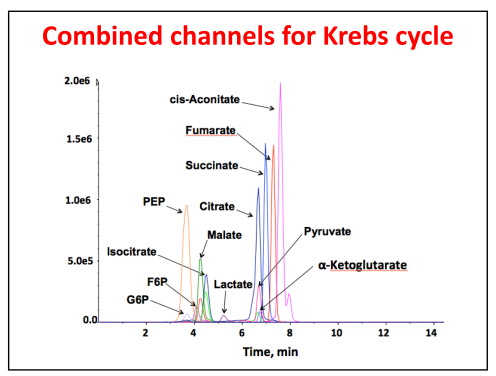


Based on precursor ion/product ion pair(s)

Courtesy, John Cutts

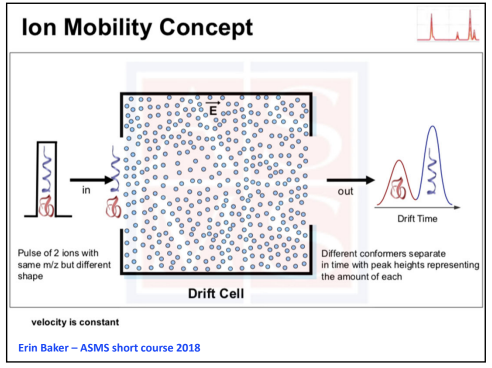
How many MRM transitions?

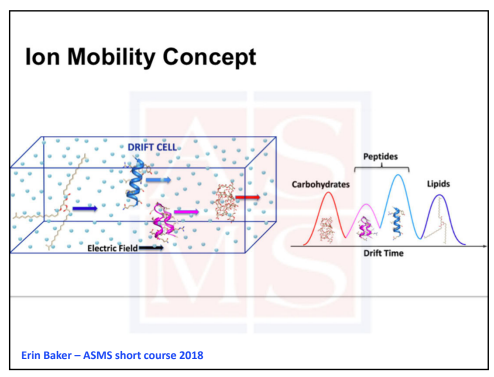
- Acquisition can be as little as 2 msec, but acquisition time determines sensitivity
- Fast switching electronics can measure as many as 500 different transitions per second
- Since measuring the area under a peak requires 10 data points, the number of transitions measured has to be matched against the shape and width of the chromatographic peaks – to be discussed in more detail later

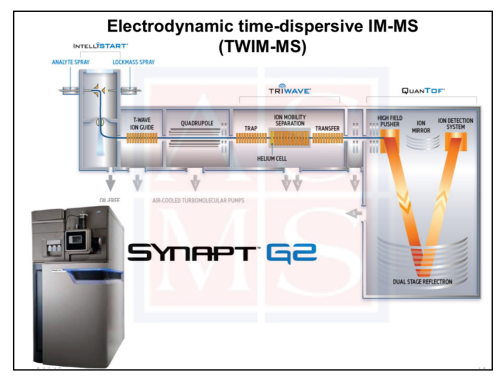


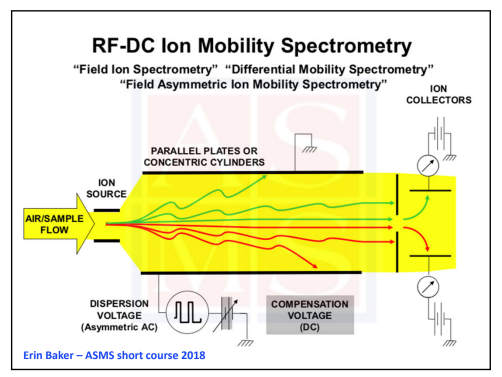
Ion mobility – another parameter

53



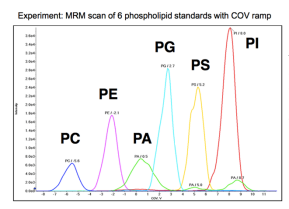






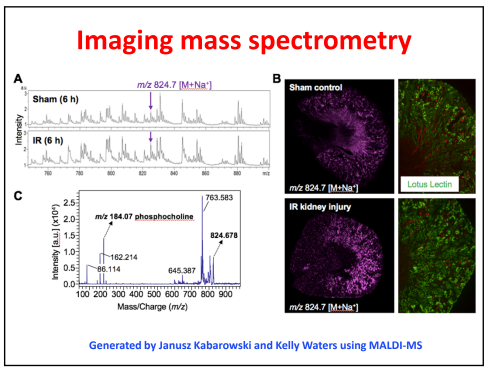
Ion mobility mass spectrometry

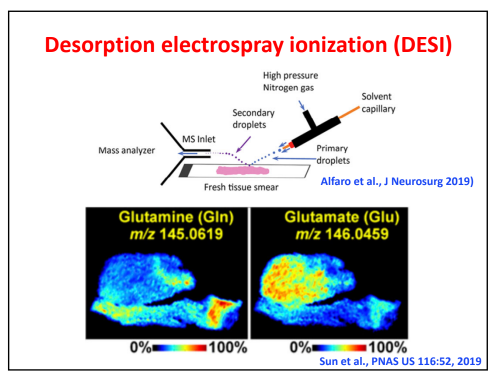
Another method of separating classes of compounds as well as compounds with the same molecular mass

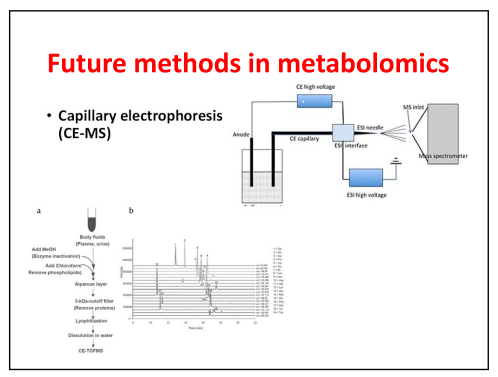


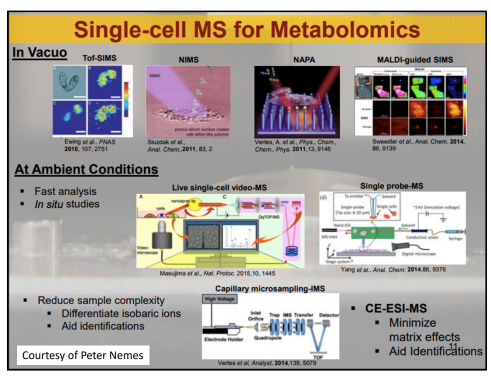
This is a gas-phase separation of these phospholipids, i.e., no chromatography.

SCIEX use a differential mobility process.









Imaging metabolites in real time

- In an ideal world, we want to measure metabolites without their degradation, spatially (preferably sub-cellularly) and with regard to time
 - MS has high qualitative mass resolution and sensitivity, but it is destructive and not subcellular. Has poor time resolution
 - NMR is non-destructive and quantitative, but is not sensitive and not subcellular. Poor time resolution
- Correlated anti-Stokes Raman Spectroscopy
 - https://bernstein.harvard.edu/research/cars-why.htm
 - Is nondestructive, has high sensitivity and spatial and time resolution, but poor qualitative resolution (distinguishing metabolites)