Mass spectrometry of proteins, peptides and other analytes: principles and principal methods

Matt Renfrow January 11, 2008

# **Objectives of the Lecture**

1. Make ions

- 2. Separate/Analyze/Detect ions
- 3. What is mass resolution and mass accuracy?

#### What does a mass spectrometer do?

- 1. It measures mass better than any other technique.
- 2. It can give information about chemical structures.

#### What are mass measurements good for?

To identify, verify, and quantitate: metabolites, recombinant proteins, proteins isolated from natural sources, oligonucleotides, drug candidates, peptides, synthetic organic chemicals, polymers

### **Applications of Mass Spectrometry**

#### Pharmaceutical analysis

Bioavailability studies

Drug metabolism studies, pharmacokinetics

Characterization of potential drugs

Drug degradation product analysis

Screening of drug candidates

Identifying drug targets

#### Biomolecule characterization

Proteins and peptides

Oligonucleotides

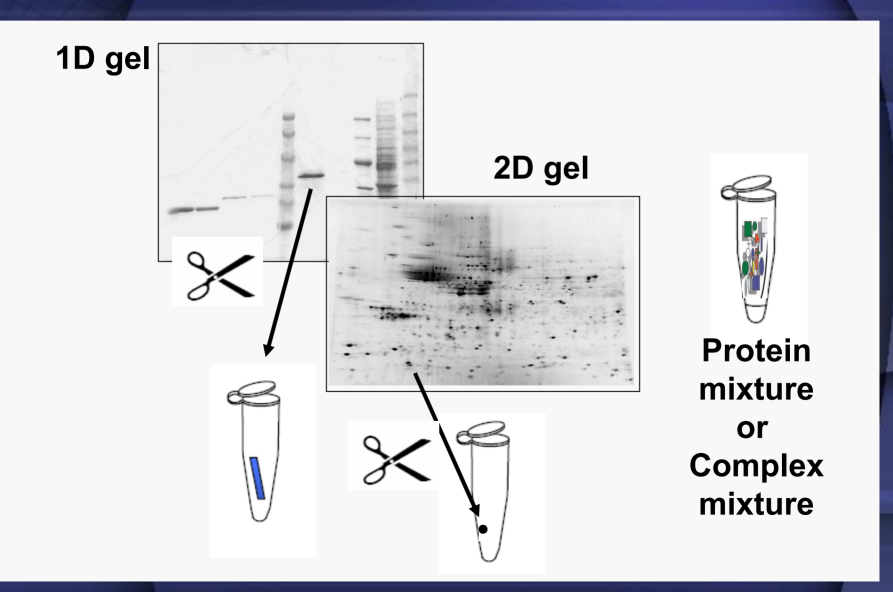
#### **Environmental analysis**

Pesticides on foods

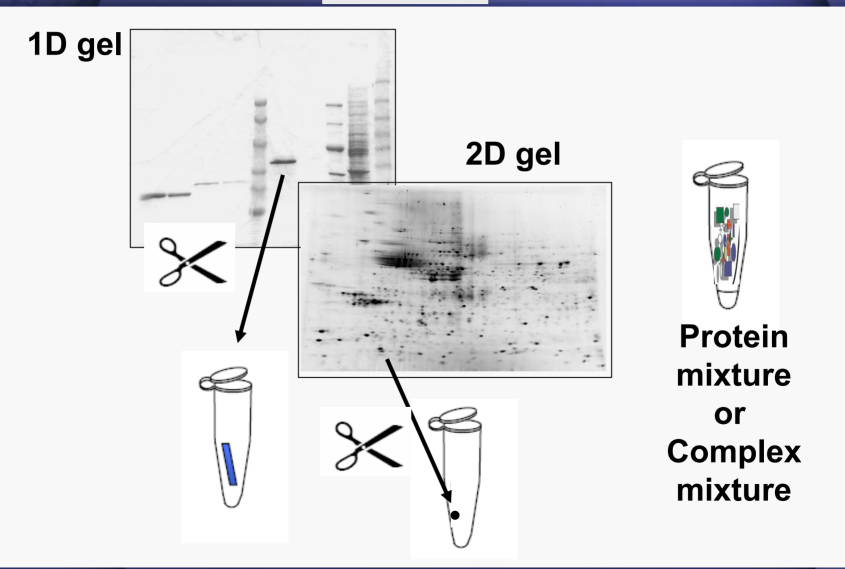
Soil and groundwater contamination

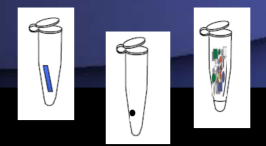
Forensic analysis/clinical

# MS of Proteins and Peptides



# Put it in your machine and tell me the RIGHT answer





## Put it in

- We need lons (+ or )
- In the gas phase

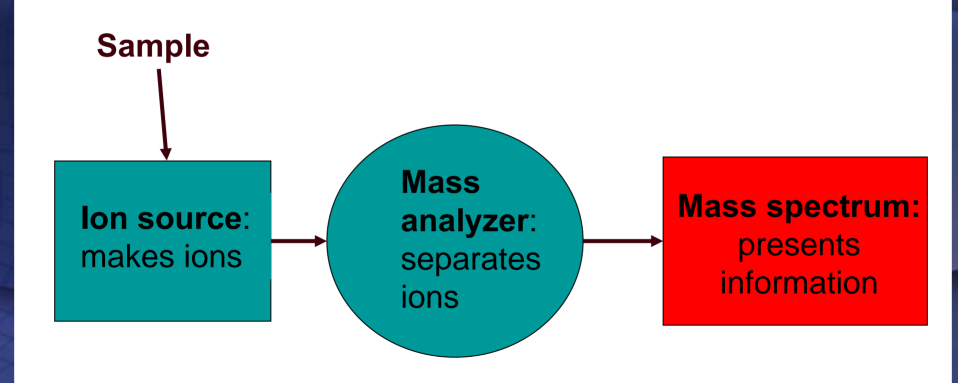
### Your machine

- Tof, Tof Tof
- Quadrupole, Ion trap
- FT-ICR, Orbitrap (high resolution)
- Hybrids

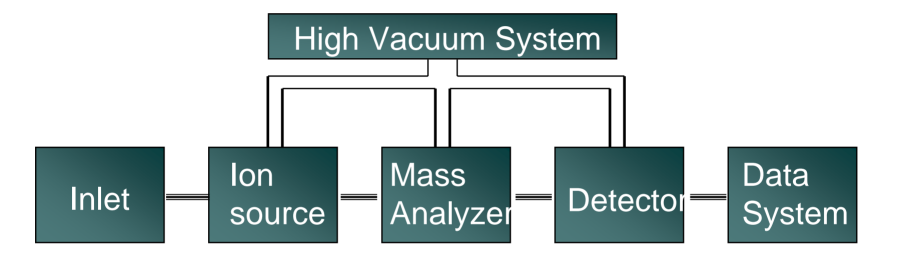
# Tell me the RIGHT answer

How right is it?
 mass resolution and accuracy

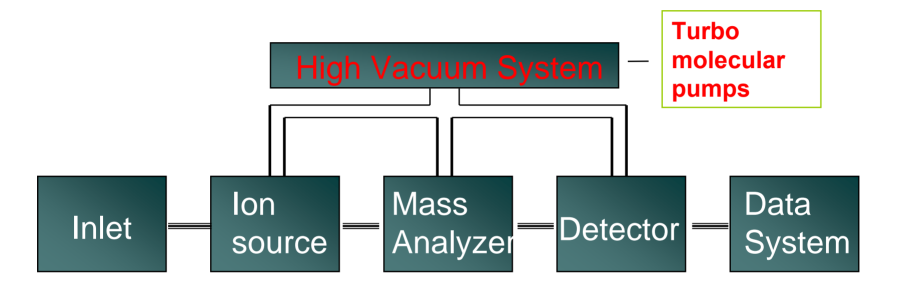
## How does a mass spectrometer work?



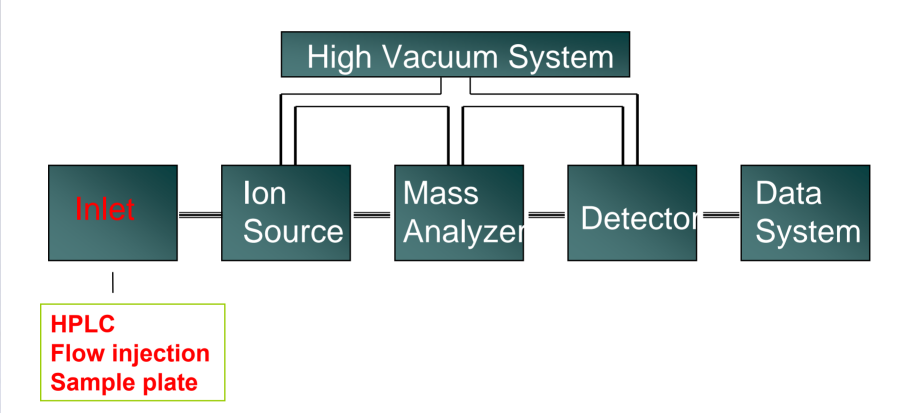
## **Mass Spectrometer Block Diagram**



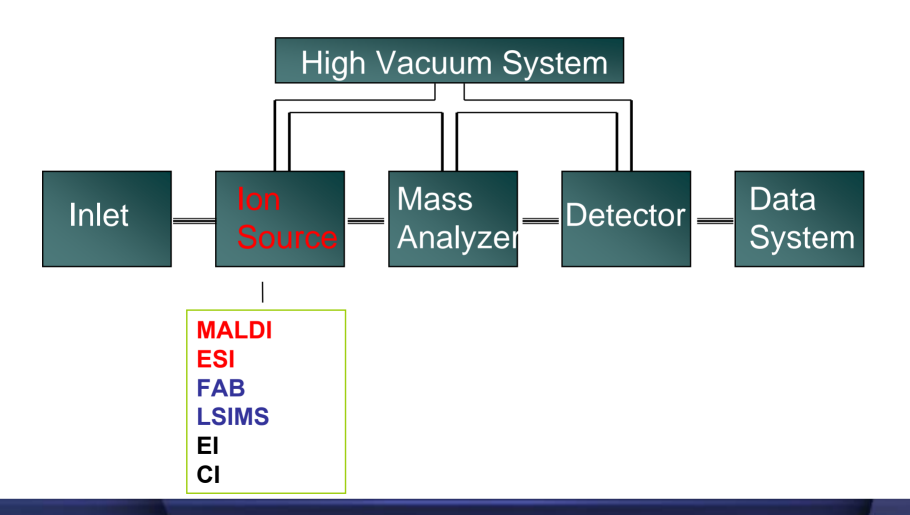
## Mass Spectrometer Block Diagram



## Sample Introduction

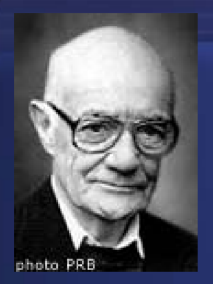


#### Ion Source



# **Nobel Prize in Chemistry- 2002**

For getting proteins and peptides into the gas phase



John Fenn



Koichi Tanaka

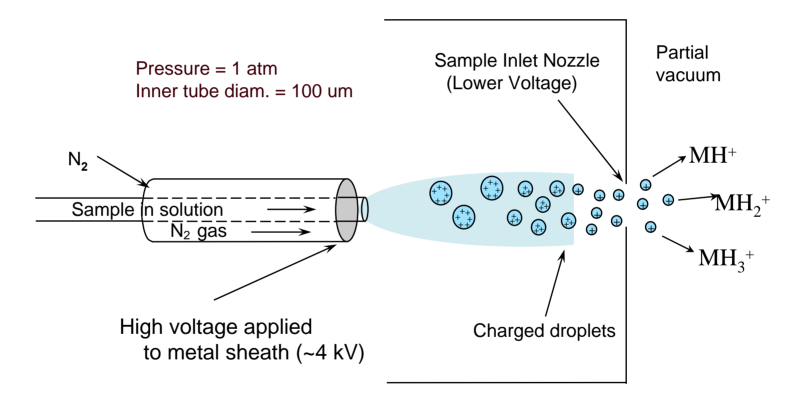
"for the development of methods for identification and structure analyses of biological macromolecules" and

"for their development of soft desorption ionisation methods for mass spectrometric analyses of biological macromolecules"

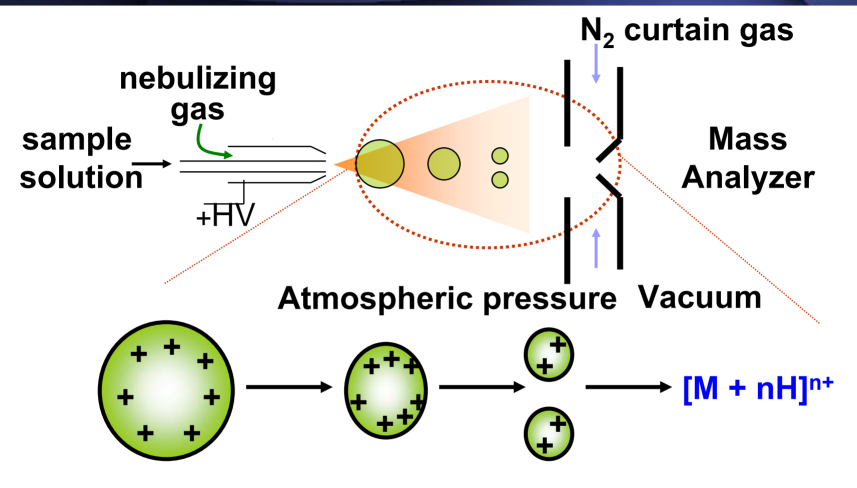
#### Ion Sources make ions from sample molecules

(lons are easier to detect than neutral molecules.)

**Electrospray ionization:** 

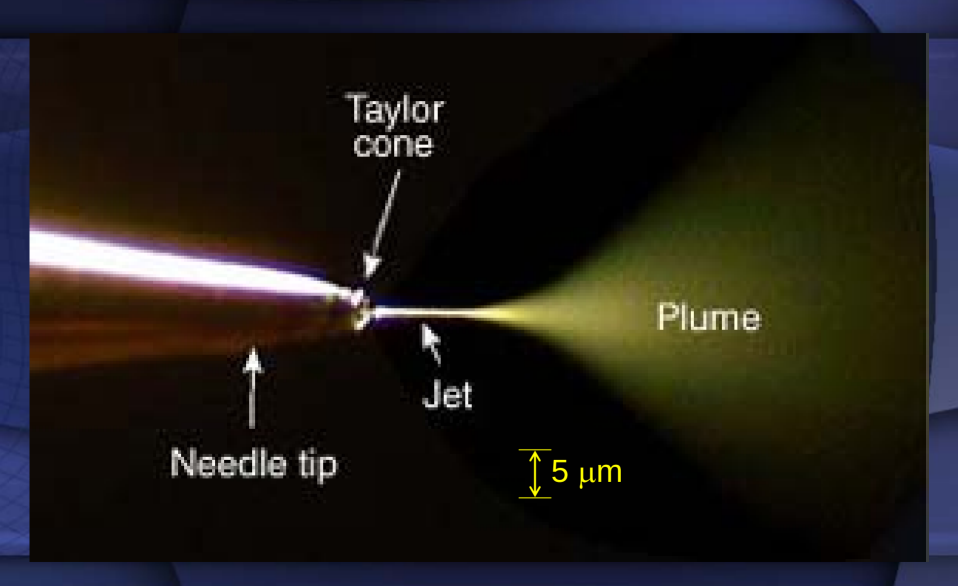


## Electrospray Ionization (ESI)

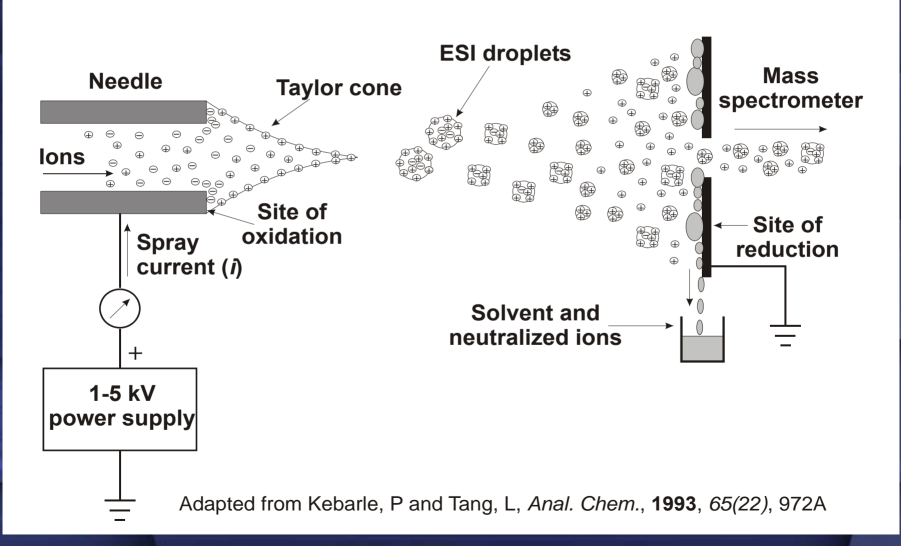


- 1. Solvent evaporation
- 2. Coulombic repulsion

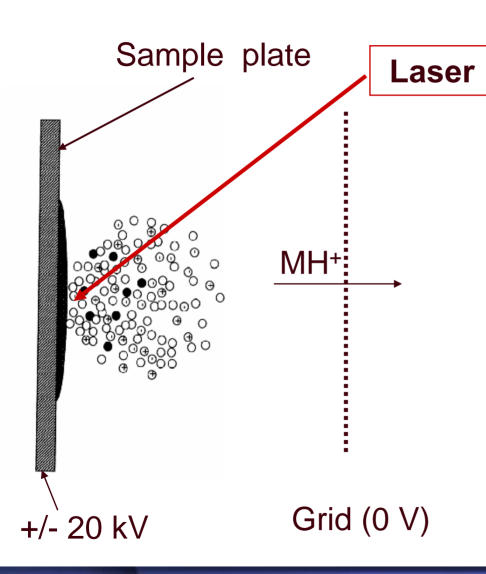
## **Electrospray Ionization (ESI)**



# Electrospray Ionization (ESI) Process (Positive Mode)



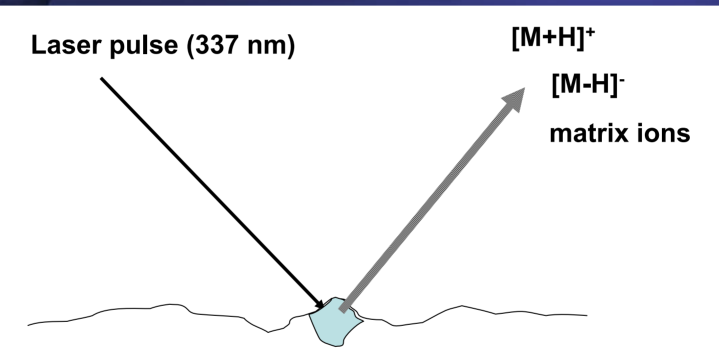
#### **MALDI: Matrix Assisted Laser Desorption Ionization**



- 1. Sample is mixed with matrix (X) and dried on plate.
- 2. Laser flash ionizes matrix molecules.
- 3. Sample molecules (M) are ionized by proton transfer: XH<sup>+</sup> + M → MH<sup>+</sup> + X.

## MALDI generation of ions

(Matrix-assisted laser desorption ionization)



Peptide/protein deposited on crystal surface

Sample mixed with a UV-absorbing matrix and is allowed to co-crystallize on the metal target.

# Matrices for MALDI analysis

#### Peptides/proteins

- 3,5-dimethoxy-4-hydroxycinnamic acid (sinapinic acid)
- α-cyano-4-hydroxycinnamic acid (CHCA)
- 2,5-dihydroxybenzoic acid (DHB)
- 2-(4-hydroxyphenylazo)-benzoic acid (HABA)

#### **Oligonucleotides**

- 2-aminobenzoic acid
- 3-hydroxypicolinic acid (3-HPA)
- 2,4,6-trihydroxyacetophenone (THAP)

The choice of matrix depends greatly on the solute to be analyzed.

#### Put it in

- We need lons (+ or -)
- In the gas phase

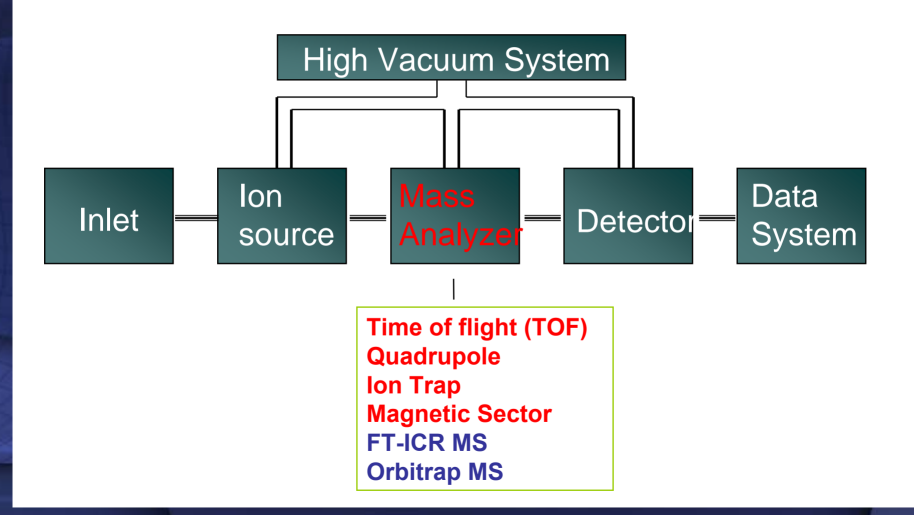
#### Your machine

- Tof, Tof / Tof
- Quadrupole, Ion trap
- FT-ICR, Orbitrap (high resolution)
- Hybrids

## Tell me the RIGHT answer

How right is it?
 mass resolution and accuracy

## **Mass Analyzer**



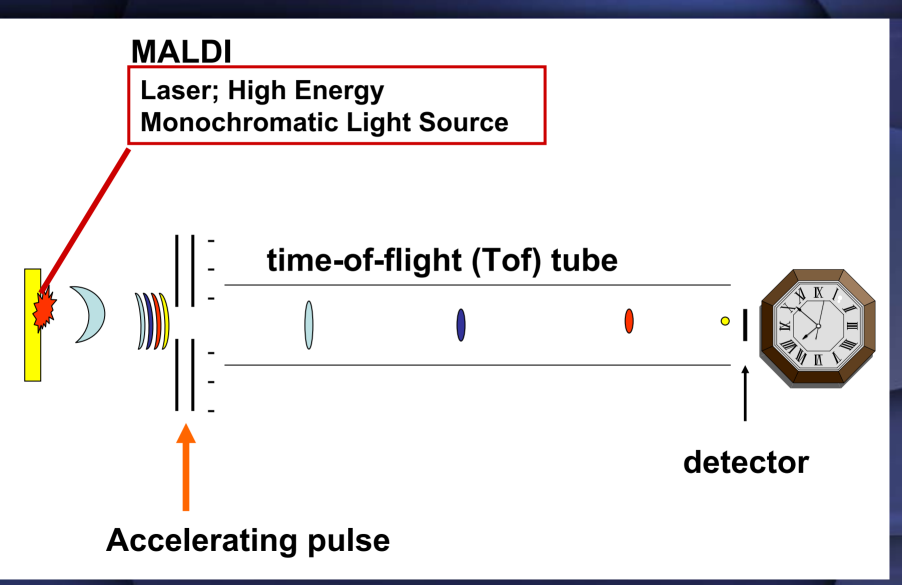
# Mass analyzers separate ions based on their mass-to-charge ratio (m/z)

- Operate under high vacuum
   (keeps ions from bumping into gas molecules)
- Actually measure mass-to-charge ratio of ions (m/z)

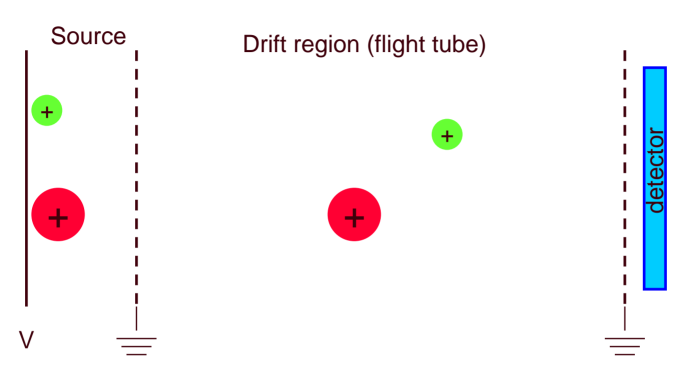
The importance of the mass-to-charge ratio is that according to classical electrodynamics two particles with the same mass-to-charge ratio move in the same path in a vacuum when subjected to the same electric and magnetic fields.

- •F=ma (Newton's second law of motion)
- • $\mathbf{F} = q(\mathbf{E} + \mathbf{v} \times \mathbf{B})$  (Lorentz force Law)
- $\bullet$ (m/q)a = **E** + v × **B**
- •Key specifications are <u>resolution</u>, <u>mass measurement</u> <u>accuracy</u>, and <u>sensitivity</u>.

# Time-of-flight (Tof) analyzer

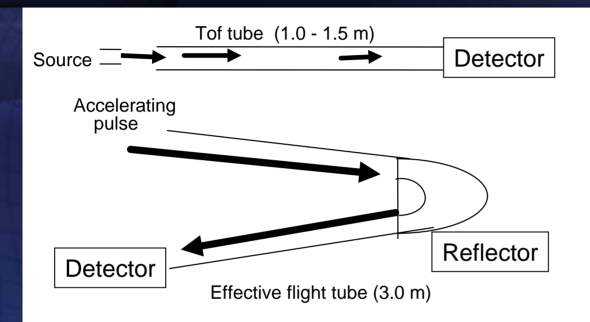


### Time-of-flight (TOF) Mass Analyzer



- lons are formed in pulses.
- The drift region is field free.
- Measures the time for ions to reach the detector.
- Small ions reach the detector before large ones.

# Time-of-flight (Tof) analyzer



Resolution 2 x 10<sup>4</sup>

No upper limit of mass

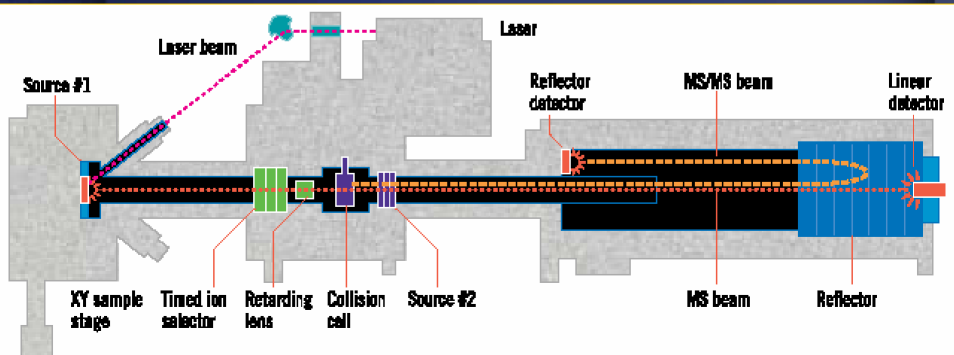
Scan times ~ 1 μsec, good for LC-MSMS

lons are accelerated so that they have equal kinetic energy. The ions "drift" down a 1 - 1.5 meter tube before striking a photomultiplier detector. "time of flight" (t) depends on the mass of the ion (m), where  $t = (m/2eV)^{1/2*}D$ 

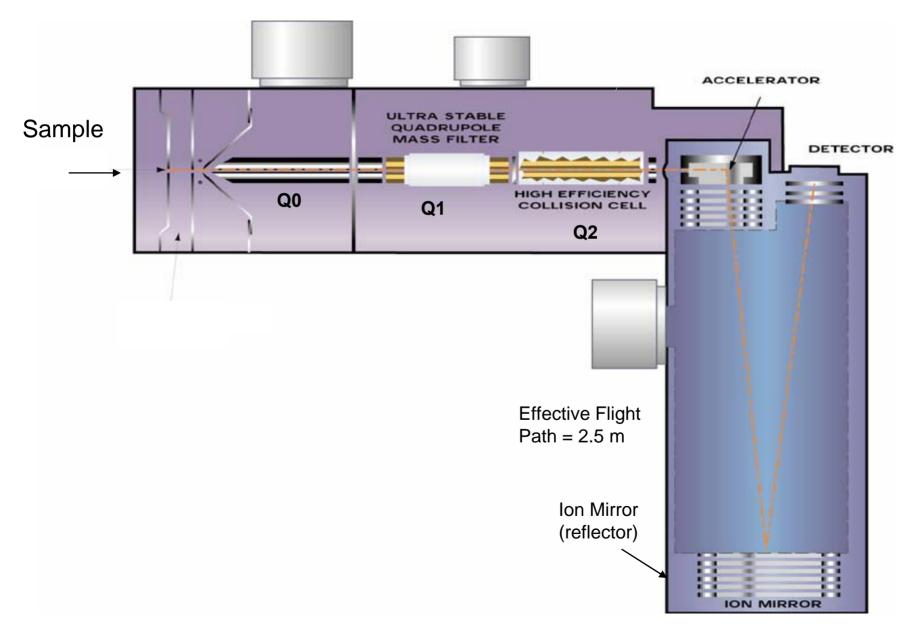
V is the applied potential and D is the flight tube distance. For a given instrument, the flight time varies as the square root of the mass of the ion.

# Principals of the MALDI-Tof/Tof 4700

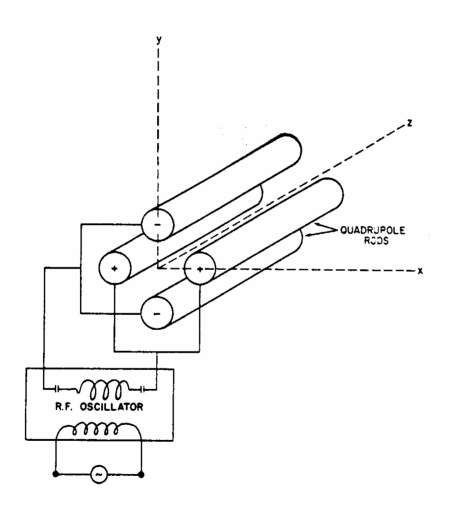




#### **QSTAR™ ESI QQ TOF or MALDI QQ TOF**



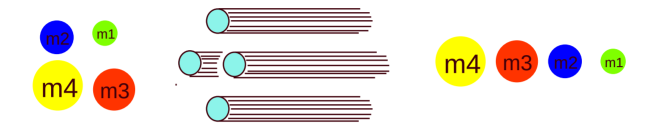
## Quadrupole Mass Analyzer



Uses a combination of RF and DC voltages to operate as a mass filter.

- Has four parallel metal rods.
- Lets one mass pass through at a time.
- Can scan through all masses or sit at one fixed mass.

#### **Quadrupoles have variable ion transmission modes**

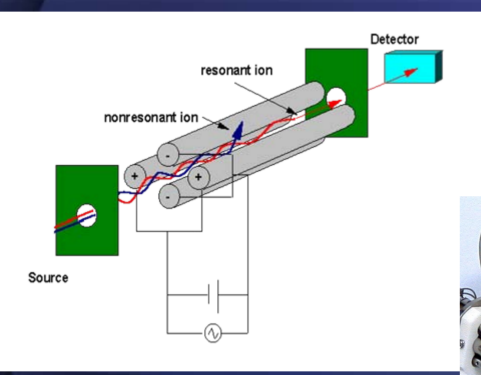


mass scanning mode



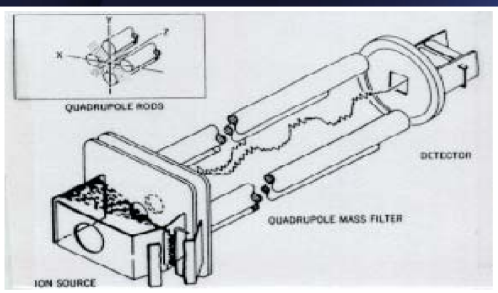
single mass transmission mode

# Quadrupole mass filter / ion guide



Octapole

# Quadrupole analyzer



Mass resolution 2 x 10<sup>3</sup>

Tolerant of relatively high pressure (10<sup>-4</sup> torr)

Upper limit for *m/z* is 3,000-4,000

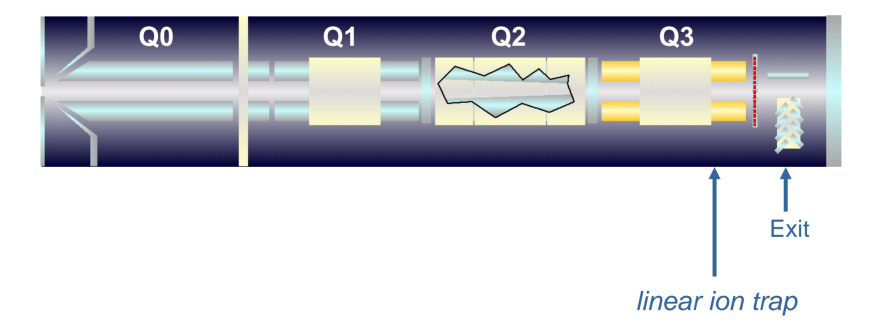
lons are accelerated electrically (5-15V) and passed along the long central axis of four rods arranged symmetrically.

By applying combined DC and oscillating RF potentials, the ions drift along irregular flight paths along the rod axis. The DC/RF ratio is held constant and the absolute values of DC and RF are varied.

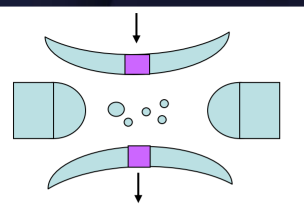
Only ions with a particular *m/z* value have stable trajectories for a given value of DC and RF.

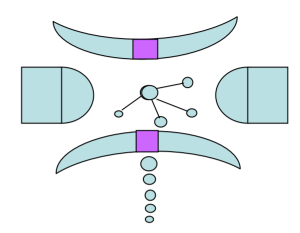
If DC is set to 0, then all ions have stable trajectories. A scan can be accomplished over a period of 10-1000 msec.

#### QTRAP: Linear Ion Trap on a Triple Quadrupole



# Ion Traps



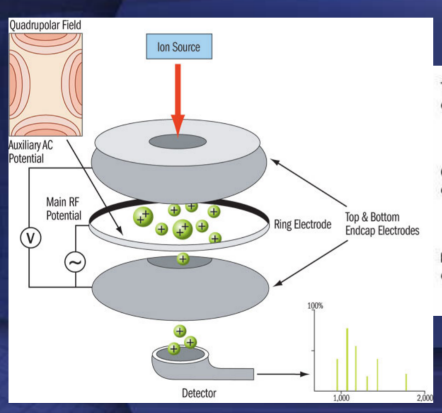


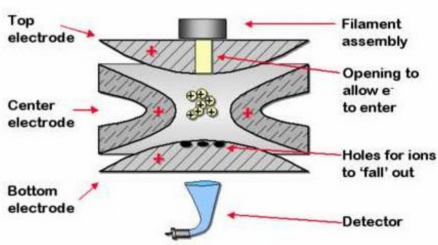
The ion trap is an energy well - ions with sufficient energy to enter the trap are retained by an energy barrier on the exit side of the trap. The advantage of the ion trap is that it accumulates selected ions prior to their analysis giving it high initial sensitivity (detection limit of approx. 20 fmol).

lons are fragmented by collision with helium gas and their daughter ions analyzed within the trap. Selected daughter ions can undergo further fragmentation, thus allowing MS<sup>n</sup>.

The ion trap has a high efficiency of transfer of fragment ions to the next stage of fragmentation (unlike the triple quadrupole instrument).

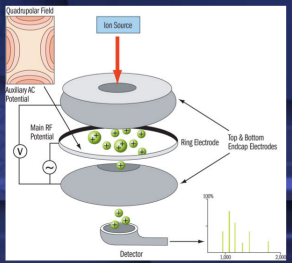
# **Expanded view of 3D ion trap**



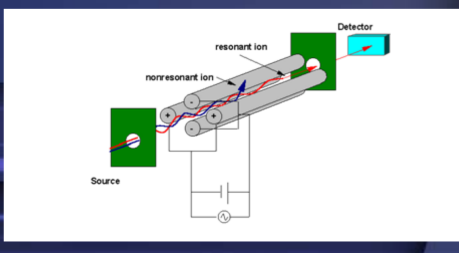


# 3D ion trap and 2D ion trap

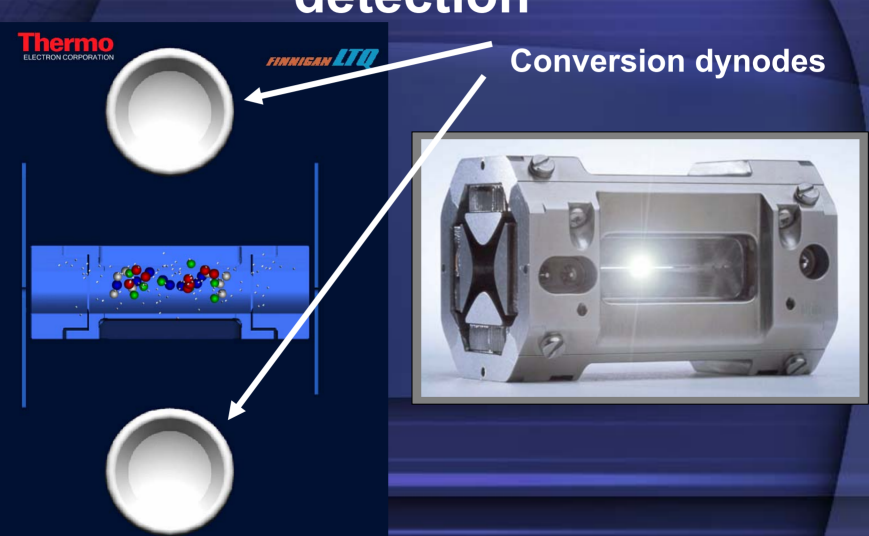








## 2D ion trap detection

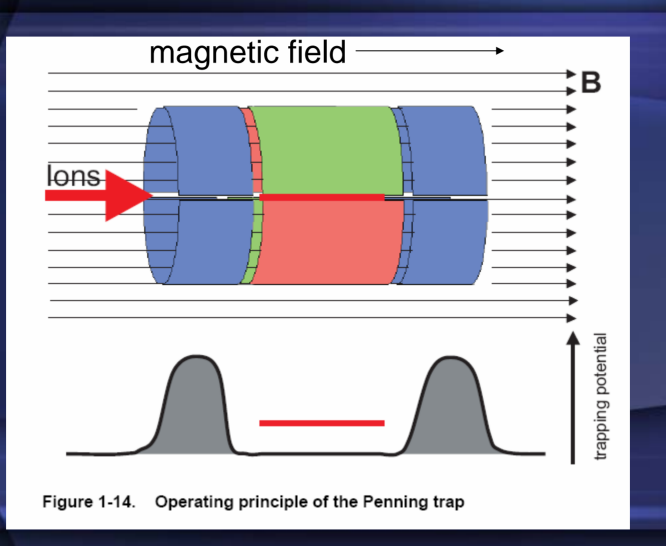


## Penning Trap (ICR cell)

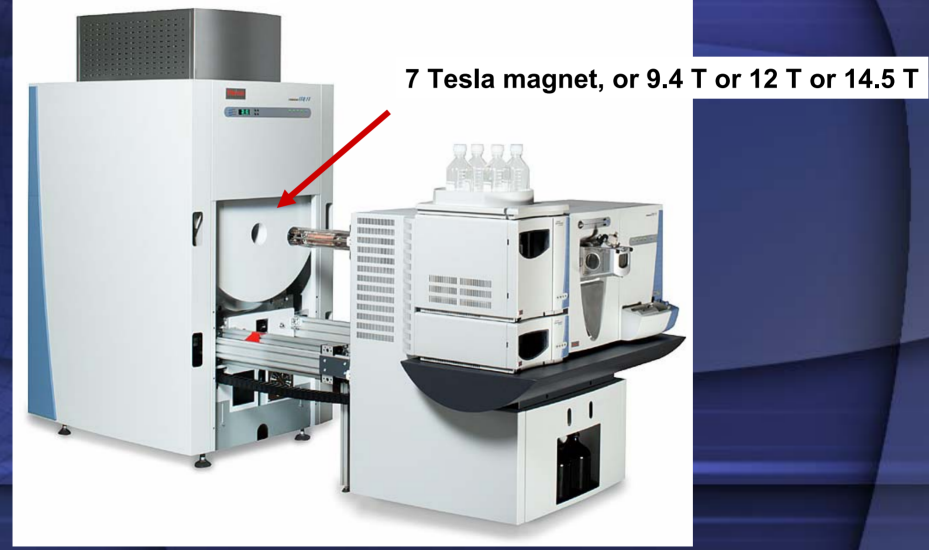




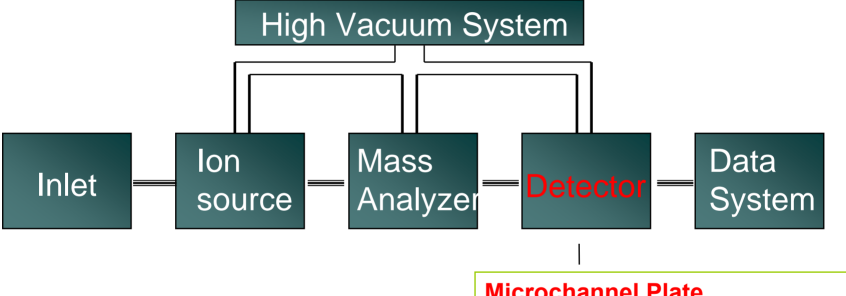
## Penning Trap (ICR cell)



# Put the trap in a high magnetic field lon cyclotron resonance



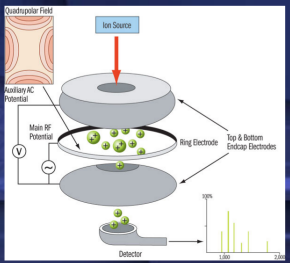
#### **Detector**

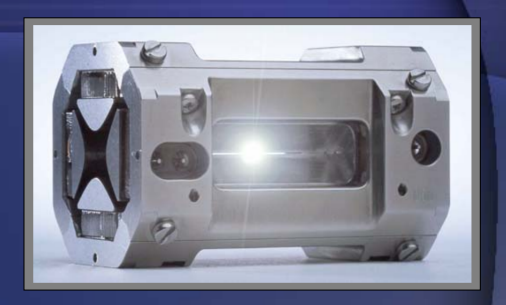


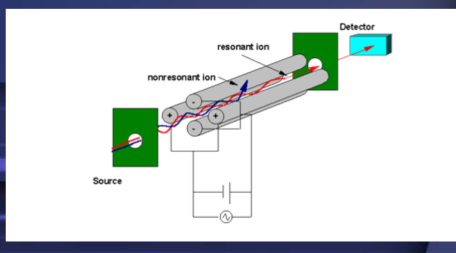
Microchannel Plate Electron Multiplier Mass analyzer/ion trap AC image

## 3D ion trap and 2D ion trap

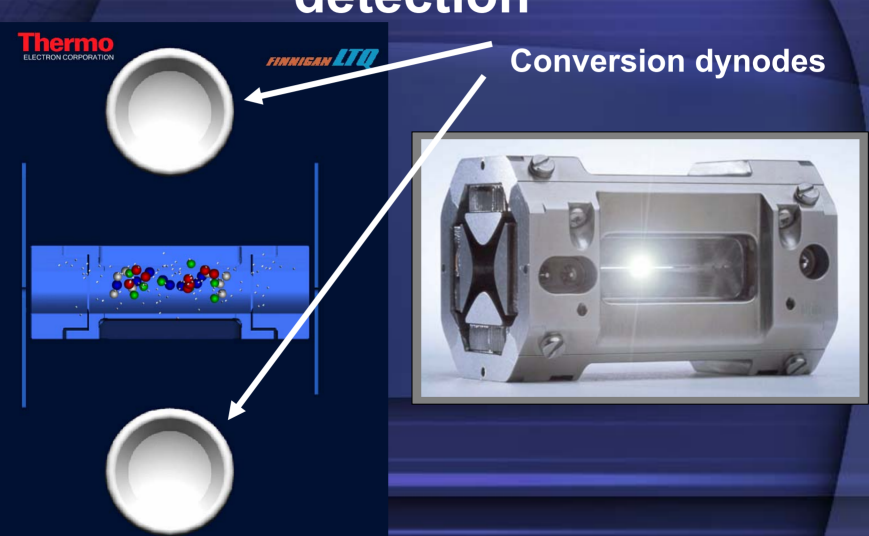




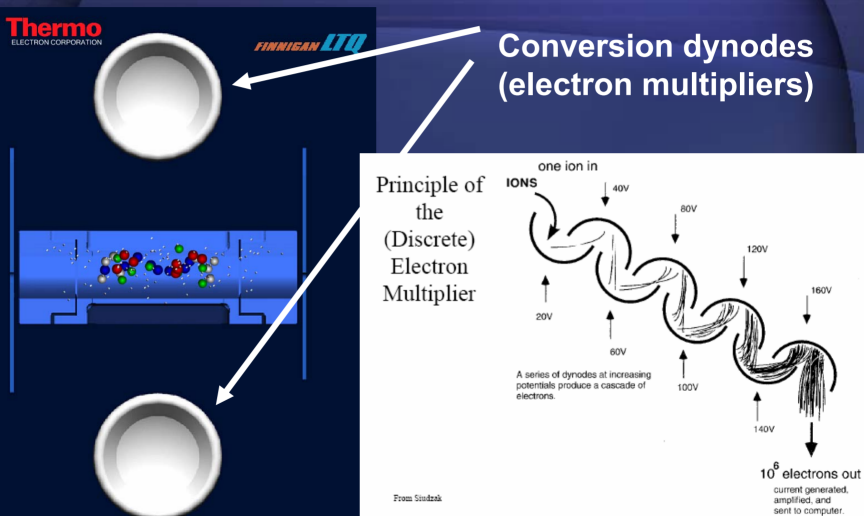




## 2D ion trap detection



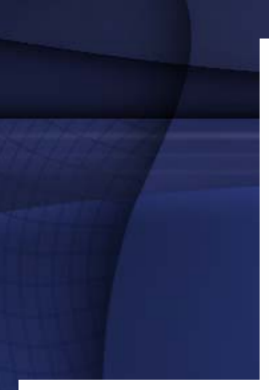
## 2D ion trap detection



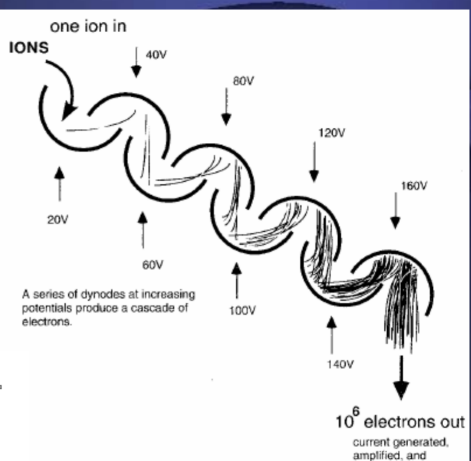
160V

current generated.

amplified, and sent to computer.



Principle of the (Discrete) Electron Multiplier



sent to computer.

#### Continuous Dynode Electron Multiplier

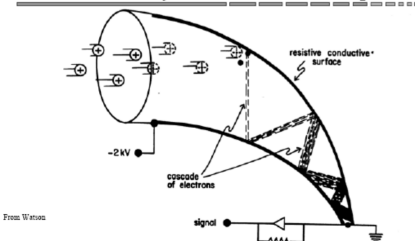
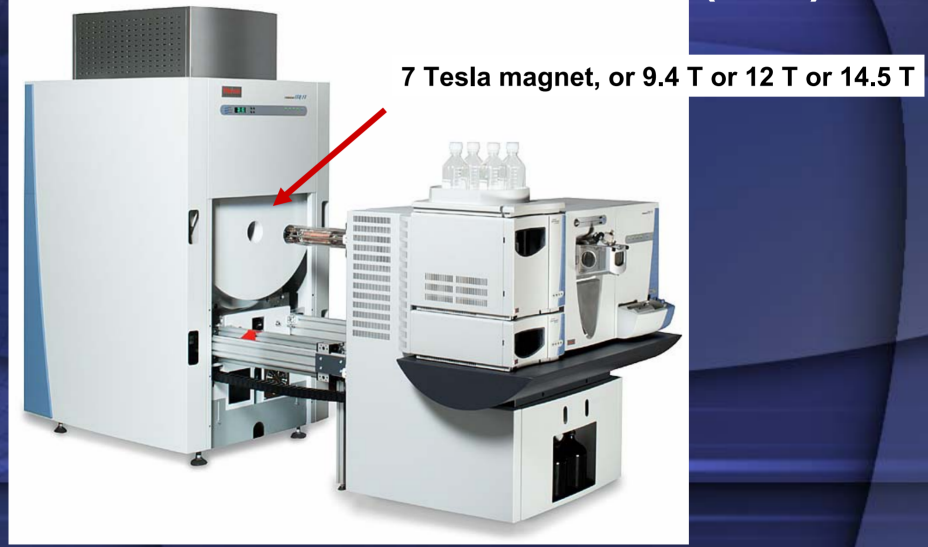
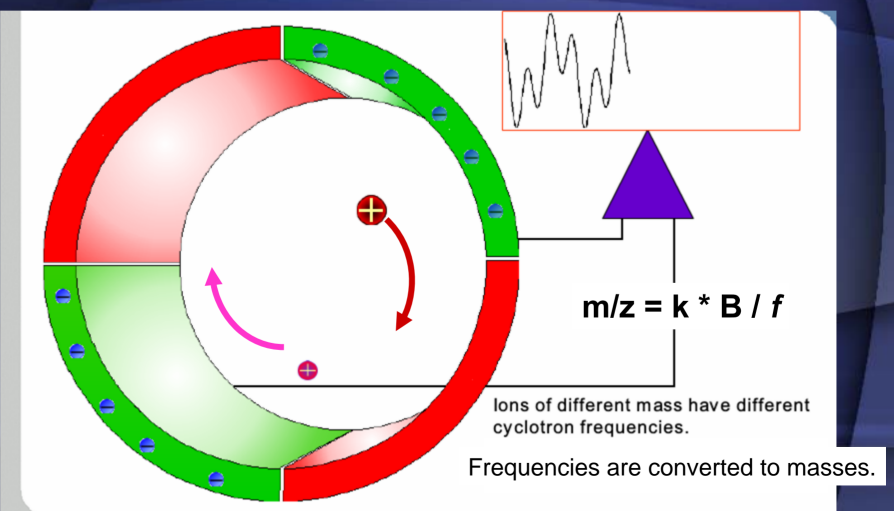


FIG. 13.3. Conceptual diagram of a nonmagnetic electron multiplier; the field gradient along the resistive conductive internal surface of the cornucopia attracts the cascading electrons toward the preamplifier.

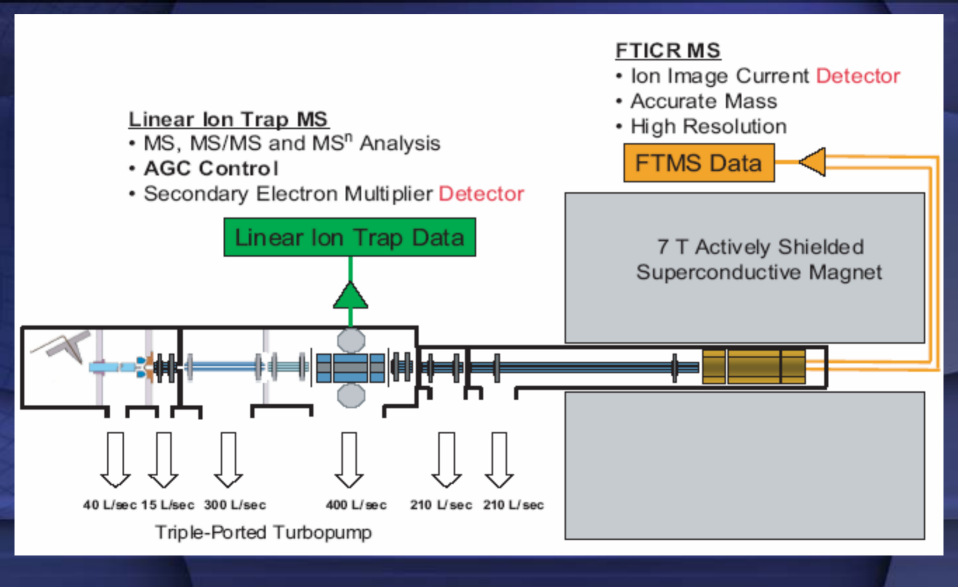
# Detecting in the ion trap lon cyclotron resonance (ICR)



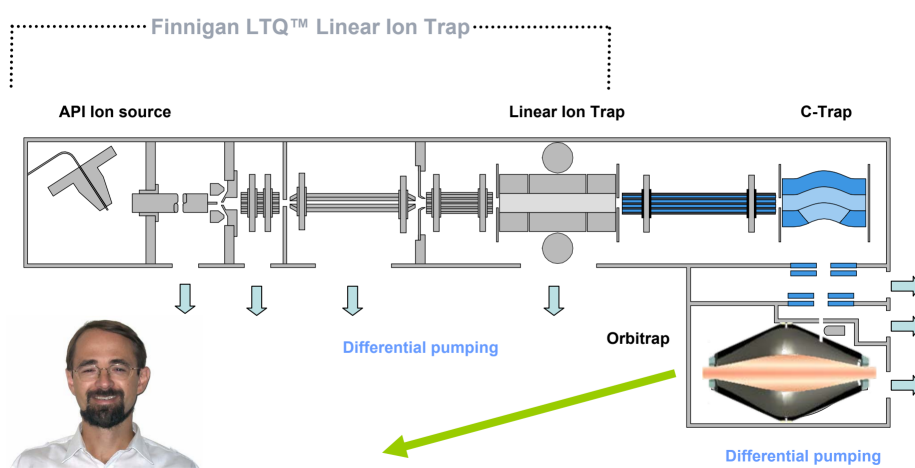
## Fourier transformlon cyclotron resonance FT-ICR MS



## ThermoFinnigan LTQ-FT

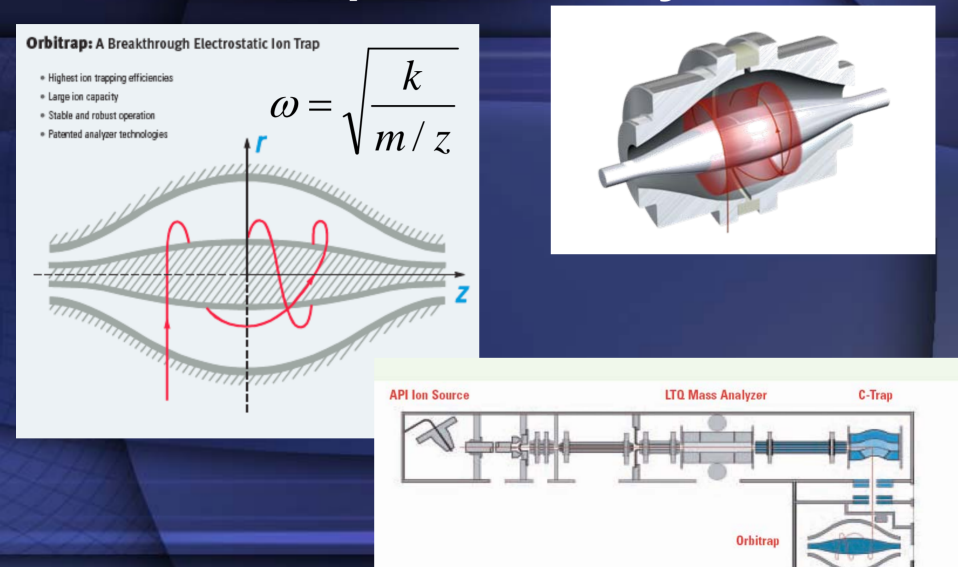


## LTQ Orbitrap™ Hybrid Mass Spectrometer

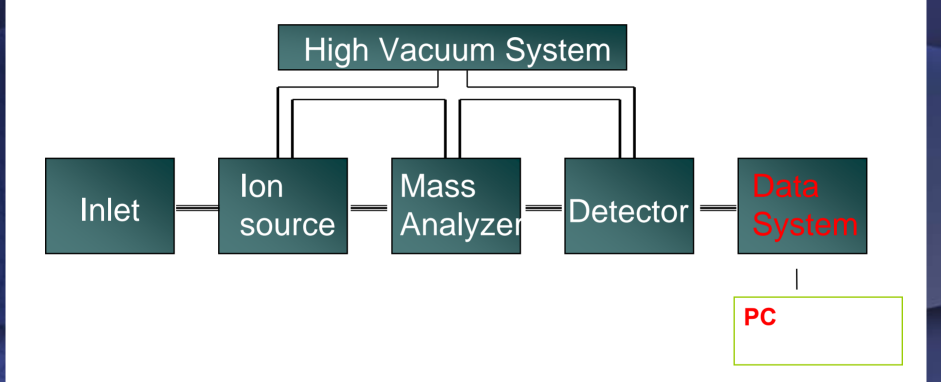


Inventor: Dr. Alexander Makarov, Thermo Electron (Bremen)

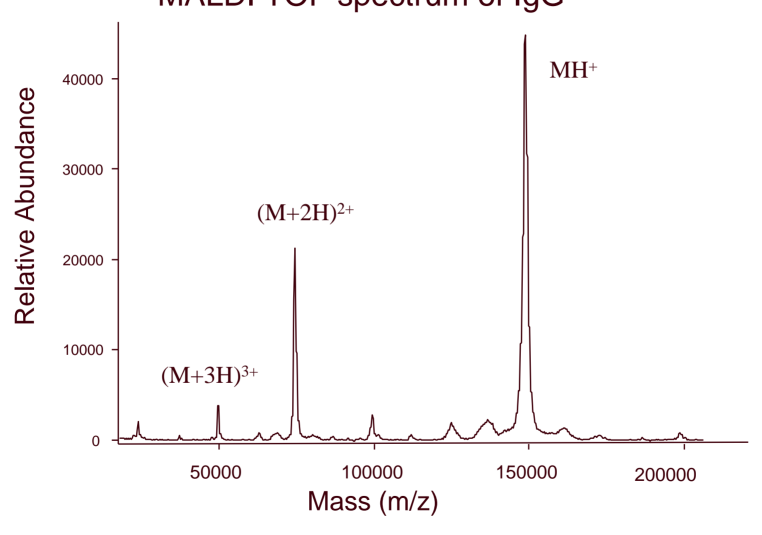
## Orbitrap Mass Analyzer



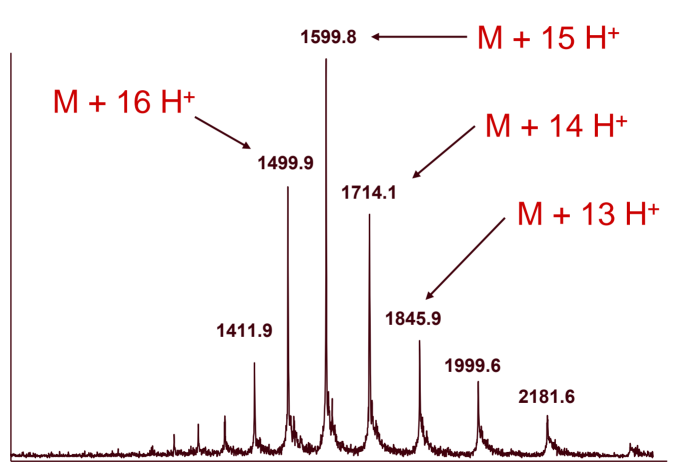
## **Data System**



## The mass spectrum shows the results MALDI TOF spectrum of IgG



#### ESI Spectrum of Trypsinogen (MW 23983)



m/z - Mass-to-charge ratio

#### How do mass spectrometers get their names?

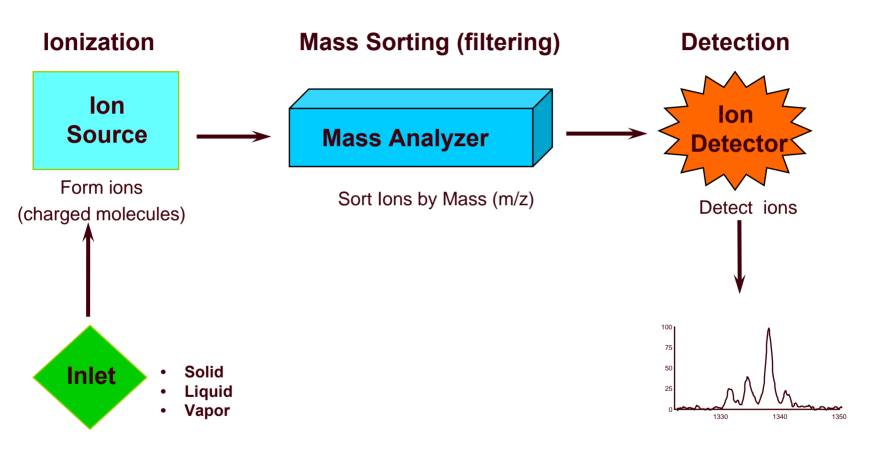
#### Types of ion sources:

- Electrospray (ESI)
- Matrix Assisted Laser Desorption Ionization (MALDI)

#### Types of mass analyzers:

- Quadrupole (Quad, Q)
- Ion Trap
- Time-of-Flight (TOF)
- -Either source type can work with either analyzer type: "MALDI-TOF," "ESI-Quad."
- -Analyzers can be combined to create "hybrid" instruments. ESI-QQQ, MALDI QQ TOF, Q Trap

### Summary: acquiring a mass spectrum



**Mass Spectrum** 

### Put it in

- We need lons (+ or -)
- In the gas phase

### Your machine

- Tof, Tof / Tof
- Quadrupole, Ion trap
- FT-ICR, Orbitrap (high resolution)
- Hybrids

## Tell me the RIGHT answer

How right is it?
 mass resolution and accuracy

### How is mass defined?

Assigning numerical value to the intrinsic property of "mass" is based on using carbon-12, <sup>12</sup>C, as a reference point.

One unit of mass is defined as a Dalton (Da).

One Dalton is defined as 1/12 the mass of a single carbon-12 atom.

Thus, one <sup>12</sup>C atom has a mass of 12.0000 Da.

### Isotopes

#### **+**Most elements have more than one stable isotope.

For example, most carbon atoms have a mass of 12 Da, but in nature, 1.1% of C atoms have an extra neutron, making their mass 13 Da.

#### **+Why do we care?**

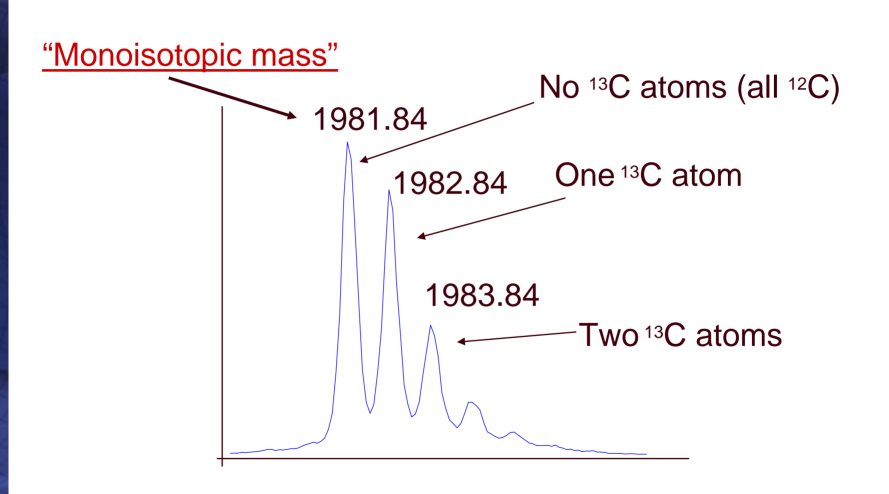
Mass spectrometers can "see" isotope peaks if their resolution is high enough.

If an MS instrument has resolution high enough to resolve these isotopes, better mass accuracy is achieved.

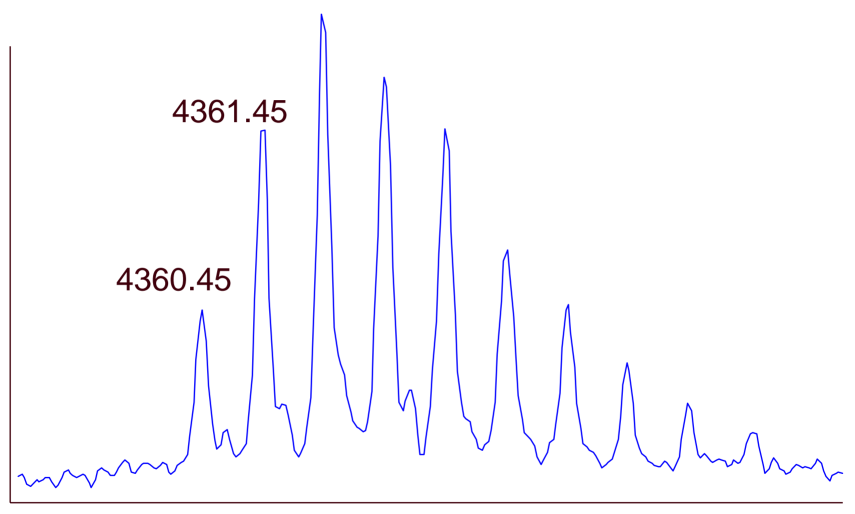
## Stable isotopes of most abundant elements of peptides

Element	Mass	Abundance
Н	1.0078	99.985%
	2.0141	0.015
C	12.0000	98.89
	13.0034	1.11
N	14.0031	99.64
	15.0001	0.36
O	15.9949	99.76
	16.9991	0.04
	17.9992	0.20

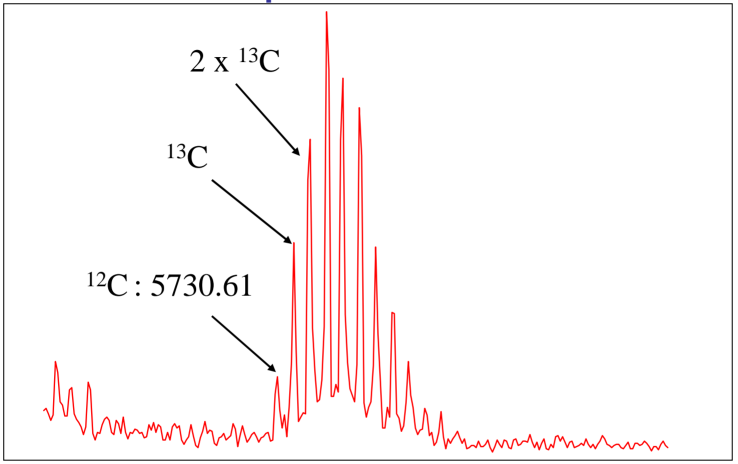
## Mass spectrum of peptide with 94 C-atoms (19 amino acid residues)



#### Isotope pattern for a larger peptide (207 C-atoms)

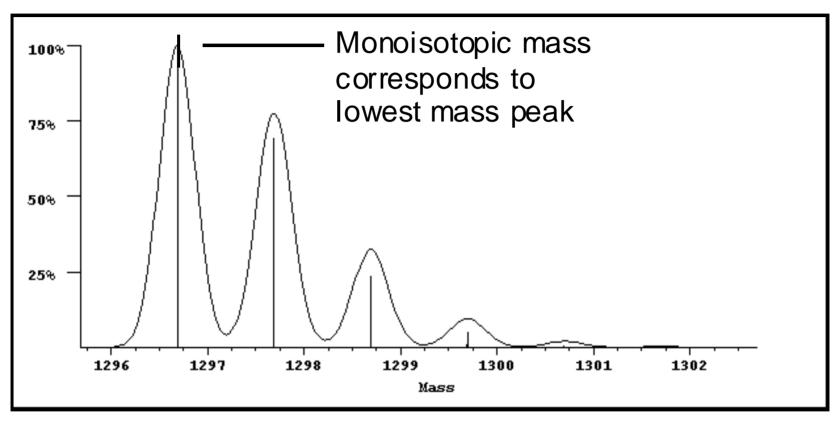


### Mass spectrum of insulin



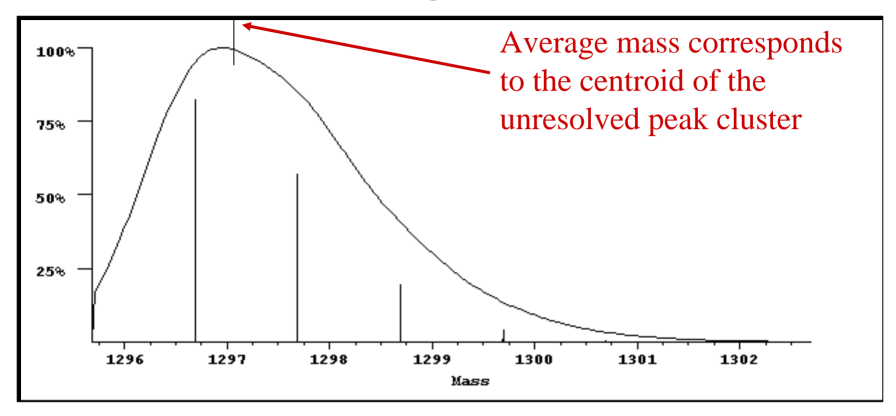
Insulin has 257 C-atoms. Above this mass, the monoisotopic peak is too small to be very useful, and the average mass is usually used.

## Monoisotopic mass



When the isotopes are clearly resolved the **monoisotopic mass** is used as it is the most accurate measurement.

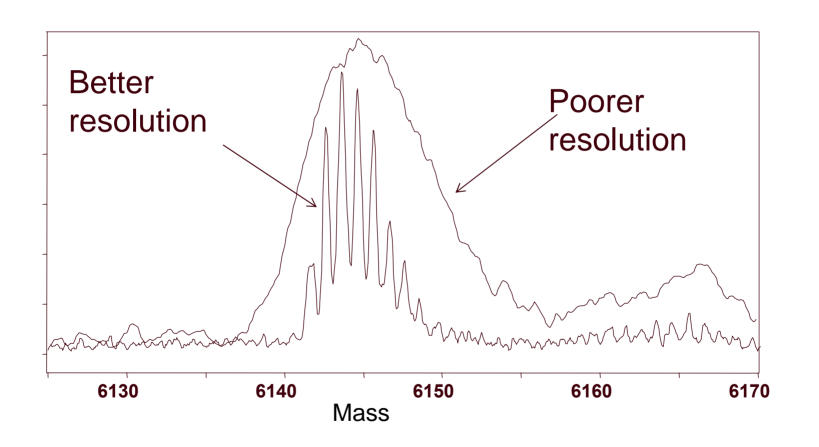
## Average mass

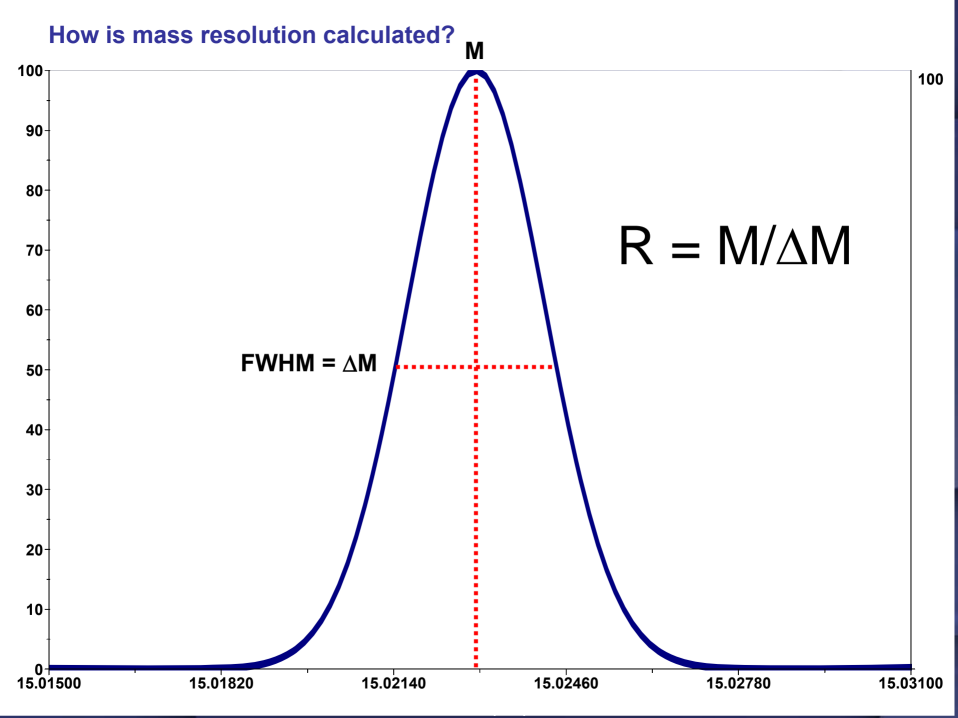


When the isotopes are not resolved, the centroid of the envelope corresponds to the weighted average of all the the isotope peaks in the cluster, which is the same as the average or chemical mass.

#### What if the resolution is not so good?

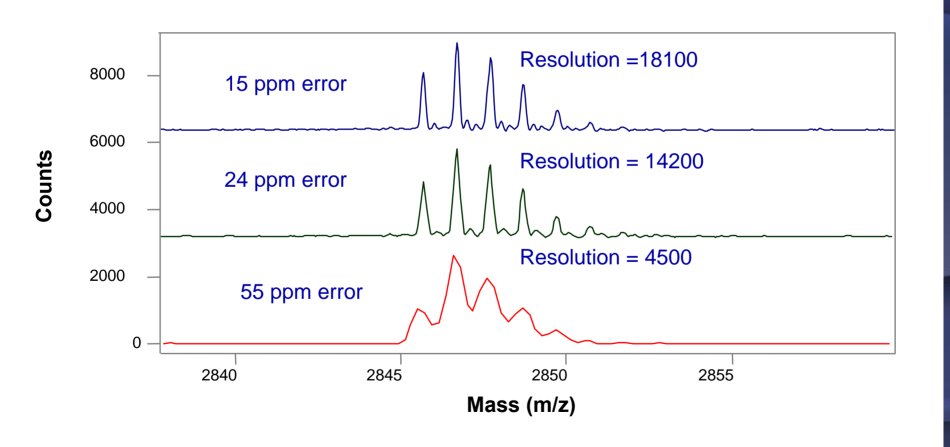
At lower resolution, the mass measured is the average mass.





#### Mass measurement accuracy depends on resolution

#### High resolution means better mass accuracy



# How do we achieve superior mass resolution?

Reflector TOF Mass Analyzer

Delayed Extraction on a MALDI source

Measure frequency

(FT-ICR MS, Orbitrap MS)

### Important performance factors

*Mass accuracy*: How accurate is the mass measurement?

**Resolution:** How well separated are the peaks from each other?

Sensitivity: How small an amount can be analyzed?

#### Two peptides - same nominal mass - simulation

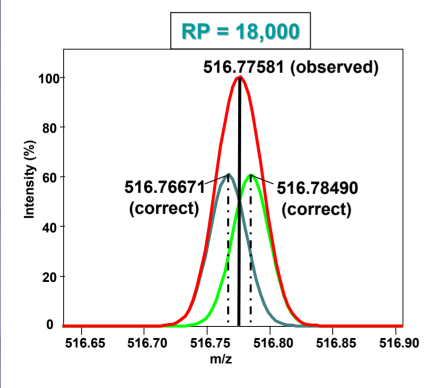
Peptide mixture: [Val<sup>5</sup>]-Angiotensin II Lys-des-Arg<sup>9</sup>-Bradykinin

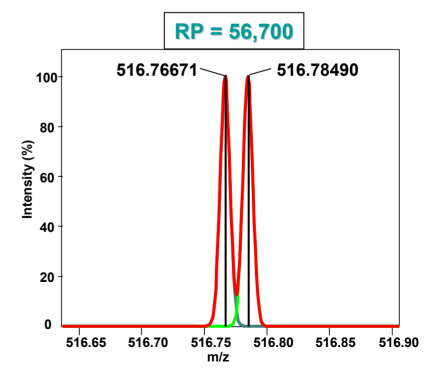
Sequence: DRVYVHPF KRPPGFSPF

Formula:  $C_{49}H_{69}N_{13}O_{12}$   $C_{50}H_{73}N_{13}O_{11}$ 

Exact mass:  $[M+2H]^{2+} = 516.76671$   $[M+2H]^{2+} = 516.78490$ 

∆m (mmu): 18.2 mmu

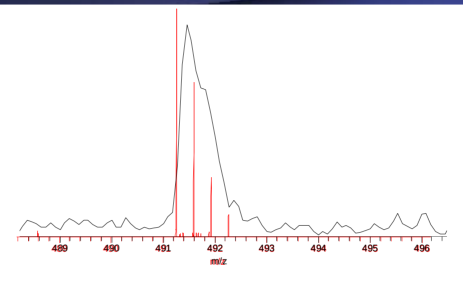


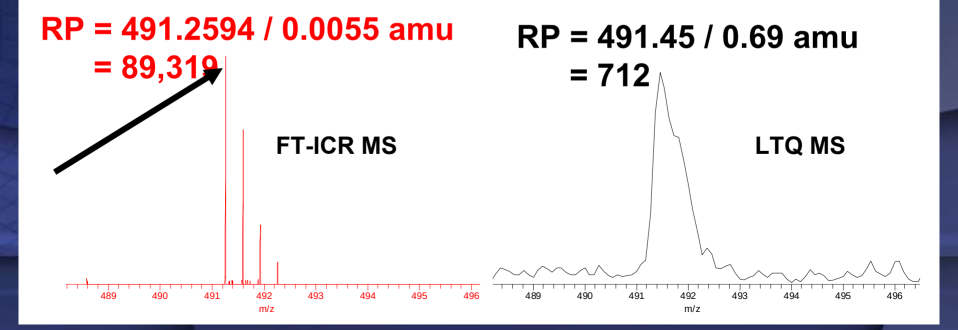


## Is Mass Accuracy Important? Results for error limit up to 5 ppm

	Theoretical Mass	Delta [ppm]	Delta [mmu]	RDB	Composition
1 ppm (4)	516.76671 516.76647 516.76638 516.76705	0.0 0.5 0.6 -0.7	0.0 0.2 0.3 -0.3	21.0 15.0 12.0 11.5	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
2 ppm (10)	516.76604 516.76738 516.76604 516.76580 516.76772 516.76773	1.3 -1.3 1.3 1.8 -2.0 -2.0	0.7 -0.7 0.7 0.9 -1.0 -1.0	16.0 20.5 21.5 15.5 16.5 11.0	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
5 ppm (23)	516.76805 516.76537 516.76807 516.76513 516.76513 516.76839 516.76872 516.76872 516.76874 516.76874 516.76897 516.76907	-2.6 2.6 -2.6 3.0 3.1 -3.3 3.7 -3.9 3.9 -3.9 4.3 -4.4 -4.6	-1.3 1.3 -1.4 1.6 1.6 -1.7 1.9 -2.0 2.0 -2.0 -2.0 -2.2 -2.3 -2.4	25.5 16.5 7.0 10.5 16.0 16.0 20.0 25.0 17.0 6.5 11.0 12.5 15.5	C <sub>52</sub> H <sub>69</sub> O <sub>9</sub> N <sub>14</sub> C <sub>46</sub> H <sub>73</sub> O <sub>15</sub> N <sub>12</sub> C <sub>38</sub> H <sub>79</sub> O <sub>14</sub> N <sub>15</sub> S <sub>2</sub> C <sub>46</sub> H <sub>81</sub> O <sub>14</sub> N <sub>8</sub> S <sub>2</sub> C <sub>45</sub> H <sub>75</sub> O <sub>9</sub> N <sub>15</sub> S <sub>2</sub> C <sub>46</sub> H <sub>75</sub> O <sub>12</sub> N <sub>13</sub> S <sub>1</sub> C <sub>52</sub> H <sub>75</sub> O <sub>11</sub> N <sub>9</sub> S <sub>1</sub> C <sub>54</sub> H <sub>71</sub> O <sub>10</sub> N <sub>11</sub> C <sub>44</sub> H <sub>71</sub> O <sub>14</sub> N <sub>15</sub> C <sub>40</sub> H <sub>81</sub> O <sub>15</sub> N <sub>12</sub> S <sub>2</sub> C <sub>46</sub> H <sub>79</sub> O <sub>13</sub> N <sub>11</sub> S <sub>2</sub> C <sub>40</sub> H <sub>73</sub> O <sub>16</sub> N <sub>16</sub> C <sub>48</sub> H <sub>77</sub> O <sub>13</sub> N <sub>10</sub> S <sub>1</sub>

## Mass Resulution = $m / \Delta m_{50\%}$





## **Objectives of the Lecture**

- 1. Make ions ESI, MALDI
- 2. Separate/Analyze/Detect ions
  Tof, ion trap, quadrupole,
  FT-ICR, Orbitrap
  Electron multipliers
- 3. What is mass resolution and mass accuracy?

Slide Acknowledgements
Thermo Electron (Fisher)
Bruker
ABI
Sandler Mass Spectrometry Group
David Agard group

http://www.msg.ucsf.edu/agard/

(Univ. of Calif Sanfrancisco)