Mass spectra of peptides and proteins - and LC analysis of proteomes

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Overview

• A mass spectrum
• Electrospray MS
  – Analysis of intact proteins
  – Molecular weight calculations
  – Max Entropy
• Peptides
  – Purity
• Integration of MS with LC and CE
  – Multidimensional LC
• Tandem MS
  – Identifying modification sites
A mass spectrum of several peptides from a tryptic digest

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Isotope profile of individual peptide ion
How to represent the mass of compound?

• At low mass resolution (where the isotope peaks cannot be resolved) what is observed is the *average mass*

• At high resolution where the isotopic peaks are fully resolved, then we can determine the *monoisotopic mass* for each one
Take home question (due Feb 6th)

1. What is the monoisotopic mass of human cytochrome C?
   - Hint: workout the empirical formula of hCyt C

2. What is the molecular weight of the most abundant species of human Cyt C?
   - Hint: assume that the abundance of $^{13}$C is 1.00% of total carbon atoms - do not worry about $^2$H or other isotopes
ESI-MS and purity of peptides

Guarantees of purity based on observation of “a single peak by reverse-phase HPLC” and by “it gave the correct sequence when analyzed by Edman degradation” are hollow. The lower spectrum was of a “pure” HPLC peak. The method of purification was amended and the upper spectrum was obtained.
Ionizing proteins and peptides

- \( ^+\text{H}_3\text{NCHR}_1\text{CO(NHCHR}_n\text{CO)}_n\text{NHCHR}_2\text{COOH} \) is the ion that’s found in dilute acid solution

- If there are internal basic residues, then the ions will be of the form \([\text{M}+\text{nH}]^{n+} \), where \( n = 1, 2, \text{etc.} \)

- A tryptic peptide will have a N-terminal amino group and an amino group from Arg or Lys
  - If the peptide has a mol. wt. of 1000 Da, then the singly charged ion will have a \( m/z \) of 1001, whereas the doubly charged ion has a \( m/z \) of 501
**ESI mass spectrum of ribonuclease**

<table>
<thead>
<tr>
<th>Peak (m/z)</th>
<th>Intensity</th>
<th>Charge (est.)</th>
<th>Mol. Wt. (Est.)</th>
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</table>

Cumulative MW estimate = 13,680.29

SD = 2.94

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Calculation of molecular weights and ion states

• For two ions in a series for a peptide of molecular weight M, the lower m/z value (x) will be for the n+1 ion state and the larger m/z value (y) will be for the n+ ion state.
  - (1) \( \frac{M+n}{n} = y \)
  - (2) \( \frac{M+n+1}{n+1} = x \)

• Hence
  - (3) \( M+n = ny \) and \( M = ny-n \)
  - (4) \( M+n+1 = (n+1)x \) and \( M = (n+1)x-(n+1) \)

• Hence
  - \( ny-n = (n+1)x - (n+1) \)
  - \( ny-n-xn+n = x-1 \)
  - \( n(y-x) = x-1 \)
  - \( n = (x-1)/(y-x) \)

• The value of \( n \) can then be substituted in equation (1) to obtain the molecular weight of the peptide

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Oxidized Aprotinin ESI mass spectrum

Control Aprotinin ESI mass spectrum

Deconvoluted mass spectra

Oxidized

Control

Junlong Shao

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Deconvolution of oxidized forms of β-lactoglobulin

Junlong Shao
LC/MS of 4HNE-Modified Cytochrome C

12356 Native Cytochrome C
12512 Cytochrome C + One 4HNE Michael Addition (+156)
12494 Cytochrome C + One 4HNE Schiff Base (+138)
12668 Cytochrome C + Three 4HNE Michael Addition (+467)
12823 S. Barnes-UAB 1/30/04

Courtesy of Amanda Foxwell
ESI spectrum of bacterially expressed protein

Each ion is \([\text{M+nH}]^{n+}\)

For 50+ charge state of a 50 kDa protein,

\[ m/z = \frac{[50,000 + 50]}{50} = 1,001 \]
Deconvolution of MS data

• When several proteins are present, then their multiply charged ion clusters overlap

• Can this be overcome? - yes, use the MaxEntropy program provided by Micromass
MaxEnt deconvolution of MWs

15,157.6 + 0.9 Da

50,011.3 + 18.3 Da

6xHis-tag BAT

57,195.2 + 12.9 Da

E. coli GRoEL

100

%}

mass

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Courtesy of Mindan Sfakianos
Summary of determining MW by ESI

• The multiple charge states of a protein allow:
  – Mol Wt of large proteins to be estimated
  – accurate estimation of mol wt (super SDS-PAGE gel)

• Important to remember that the protein sample must be free of salt
  – Typically, a sample is cleaned up on a short reverse-phase column prior to electrospray
  – Alternative, use ammonium acetate as buffer
Use of FT-MS in ESI of proteins

• The very high resolving power of FT-MS enables a direct measure of charge state of an individual ion since each peptide or polypeptide will have several/many isotope peaks

• The distance in Da between successive isotope peaks of a multiply charged ion is the reciprocal of the number of charges
Bovine Serum Albumin (66 kDa)
4.7 T Actively Shielded Magnet

ESI: BSA

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Bruker Daltonics
Chemically modifying an antibody

Scheme 1

PTX-2'-OH + Succinic Anhydride → PTX-2'-O₂CCH₂CH₂CO₂H

PTX-SX

Scheme 2

MAb + PTX-SX \(\xrightarrow{\text{NHS/EEDQ}}\) DMF → NH-PTX

PTXMAb

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Structure of modified antibody

PTX-MAb

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Modification of an antibody by MALDI-TOF

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Example of MS to detect a amino acid modification

• Cytochrome c, a mitochondrial enzyme, was reacted with 4-hydroxynonenenal, an aldehyde formed by oxidation of long chain, unsaturated fatty acids

• Site of attachment believed to be on lysine groups (to form a Schiff’s base)

• However, increase in MW consistent with Michael addition

• Protein hydrolyzed with trypsin
Cytochrome C Modified by HNE
MALDI-TOF Mass Spectrum

MALDI spectra usually contain only the molecular ion $[\text{M+H}]^+$. This is an advantage since it maximizes the signal, but is a disadvantage in that it gives no clues as to structure.

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The quadrupole analyzer (Q3) is slow and insensitive - it’s a filter - thus throws away large amounts of data.

TOF detector collects all ions generated and yields fmol rather than pmol sensitivity.

Also gives far greater mass accuracy - from 1000 ppm on the triple quad to <20 ppm on the Q-tof.

Crucially important for automated interpretation of MS-MS spectra to yield amino acid sequence.
Tandem mass spectrometry on a triple quadrupole instrument

- Daughter ion spectra
- Parent ion spectra
- Multiple reaction ion scanning

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Daughter ion tandem MS

The molecular ion is selected in Q1, collided with Argon gas in Q2, and “daughters” analyzed in Q3.
Parent ion tandem MS

All molecular ions allowed into Q1, collided in Q2, and a selected daughter ion measured in Q3. This could be an O-GlcNAc fragment (m/z 204). The parent peptide ions containing an O-GlcNAc will be revealed. Very difficult to do this experiment on a Q-tof.

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Multiple reaction ion monitoring (MRM)

A single molecular ion selected in Q1, collided in Q2 and a selected daughter ion measured in Q3. Up to 8 pairs of parent/daughter ions can be measured. This is a quantitative analysis. Can’t be done on a Q-tof.
MRM analysis of isoflavones in dog plasma

The combination of parent ion and daughter ion allows specific detection of each of these isoflavonoids, even when they are not chromatographically separable.

The figure in the right hand corner is the full scale intensity for each channel.

Apigenin is an internal standard and is an isomer of genistein.
LC-MS of peptide mixtures

- Load sample
- Pre-column for desalting
- Waste

Analytical reverse phase column
75 µm i.d. x 15 cm

Flow rate 200 nl/min

Acetonitrile gradient

Collision gas

Q1
Q2

Electrostatic reflector

TOF detector

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Qtof Tryptic Digest of Control Peptide

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Amanda Foxwell
Qtot Tryptic Digest of Modified Peptide

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Conclusions of experiment

• A peptide was identified that increased its molecular weight by 156 Da

• Tandem MS revealed that the 4-HNE was attached to His-33 to form a Michael adduct
- Hydrolyze everything!
- For a cell expressing 5,000 proteins, this leads to >100,000 peptides
- Can be fractionated, but still 10,000-20,000 to differentiate
- Enormous bioinformatics problem

MUDPIT - MULTI-DIMENSIONAL PROTEIN IDENTIFICATION TECHNOLOGY

NH₄Cl (1-200 mM step gradient)
Cation exchange column (H⁺)

0-40% MeCN gradient
nanoLC

MS-MS analysis on Qqtof
Massive computing

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John Yates
Connecting CE and LC to MALDI analysis

- CE analysis
- nanoLC analysis

Creates 20 mm wide tracks that can be scanned by MALDI laser for MS analysis

Parallel capture of effluents of 8 nanoLC separations on Mylar - can be scanned simultaneously by fast laser

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Pros/Cons of laying down LC or EC separations on matrix plate

- Allows off-line analysis both in real time and then in a retrospective mode
- MALDI-TOF analysis is very fast
- Can do TOF-TOF MS-MS analysis
- BUT what happens chemically on the acidic environment on the surface of the plate during storage?
- Also, can the laser beam cause chemical changes?