BYC 744 Proteomics-Mass Spectrometry

Qualitative and quantitative burrowing of the proteome

Stephen Barnes, PhD

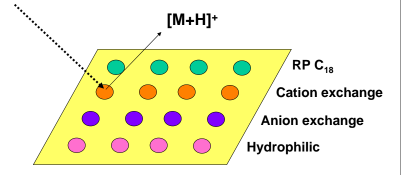
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Topics

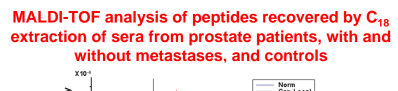
- Qualitative changes in the predicted proteome
 - Digging deeper SELDI/MALDI
 - Posttranslational modifications
 - · Isolation and characterization
 - · Making using of the chemistry of the modification
- Quantitative aspects
 - Isotope labeling
 - ICAT, ¹⁸O/¹⁶O, ¹⁵N/¹⁴N
 - · Chemical labeling
 - Absolute measures

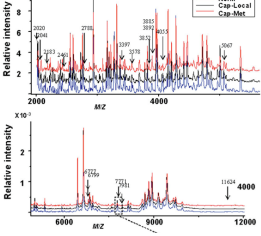




SELDI chip (plate) with local chemistry on surface causing selective binding of peptides

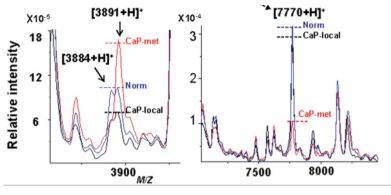
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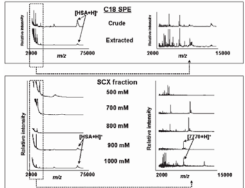




But what is it?

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Purification of prostate peptide by SPE extraction and strong cation exchange



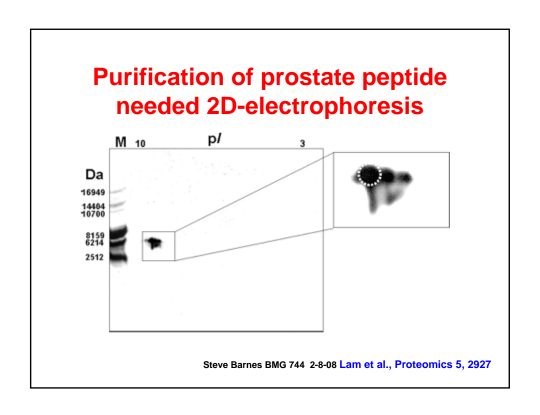
Serum was first extracted with a C₁₈ SPE cartridge

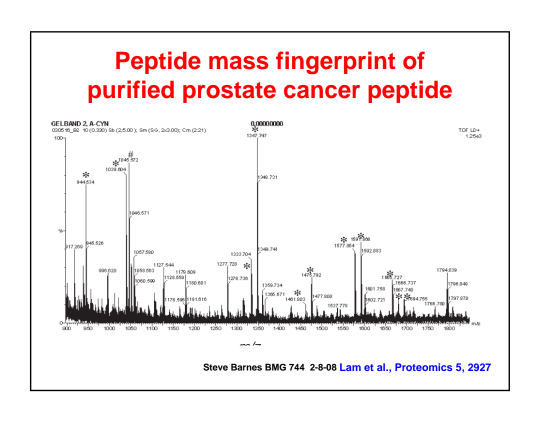
SPE fraction was passed over a [H+]-form cation exchange resin. Bound peptides were eluted with increasing step gradients of NaCl

Analyses of fractions were performed by MALDI-TOF MS

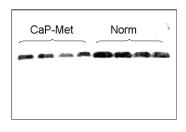
The most enriched fraction was contaminated with albumin

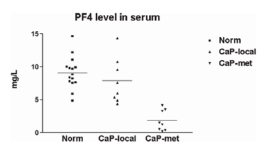
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Non-mass spec verification of prostate peptide





Western blot analysis for PF4

ELISA analysis for PF4

Moral: proteomics is a serious business that requires multiple dimensions of separation - glib methods don't work

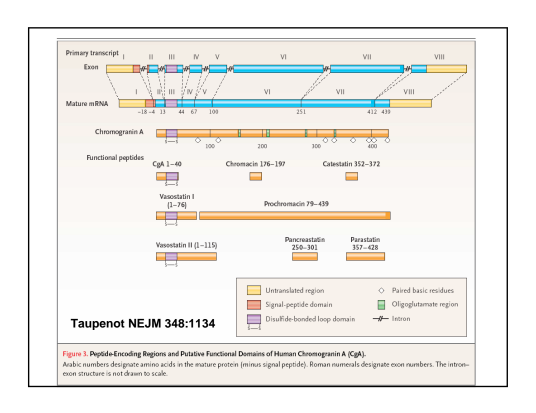
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General classes of modification

- Biochemical events involving peptide processing
- Biochemical events stimulated by enzymes
- Chemical events driven by reactive species
- Chemical events determined by investigator

Examples of peptide processing of polypeptides

- Head groups of membrane proteins
- The family of proteins in HIV that are manufactured as one protein and then hydrolyzed by a protease
- Chromogranin a brain protein consisting of several bioactive peptides
- Formation of β-amyloid



Enzymatic modifications

- Phosphorylation/dephosphorylation
 - On serine, threonine, tyrosine
- Glycosylation
 - N-glycosylation (asparagine-linked)
 - O-glycosylation (serine-, threonine-linked)
- N-Acetylation/deacetylation
 - On lysines
- N-Methylation/demethylation
 - On lysines mono-, di- and trimethylation

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Phosphorylation of proteins

- In some cases, proteins are normally found in a stable, hyperphosphorylated state, e.g., casein
- In many cases, it is a transient event that causes 10-100 fold increase in enzyme activity. This is the way signals are propagated through a signal transduction pathway. However, the molar abundance of phosphorylation at an individual site may only be 1-2%.

Chemistry of phosphorylation

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Mass spectrometry of phosphorylated proteins

- Adds H₃PO₄ (+98)
- Eliminates water (-18)
- Net change +80
- if the phosphate ion is released (i.e., in negative ion spectra), it is seen as m/z 79 and/or m/z 63
- Phosphate can be confused with sulfate and bromide (sulfate, 79.9568 Da; phosphate, 79.9663 Da; and ⁸¹Br, 79.9083 Da)

Limitations of mass spectrometry

- Although it can deliver sensitivity in the low fmol range (similar to immunological methods), because it is a universal detection method, finding the needle in the "haystack" of all the other peptides is a challenge
- Recovering the phosphopeptides from the matrix of the sample is more important than the mass spectrometry measurement

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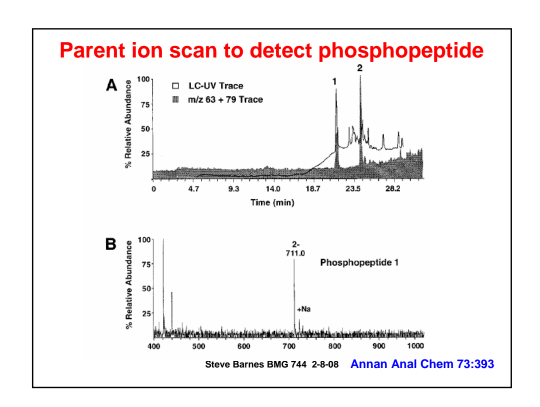
Finding a phosphate group

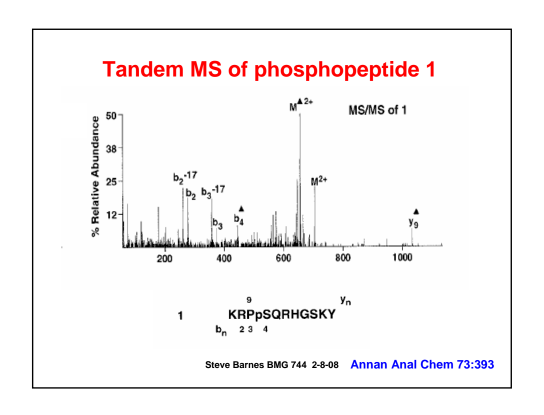
Several methods are in current use for detection of phosphopeptides

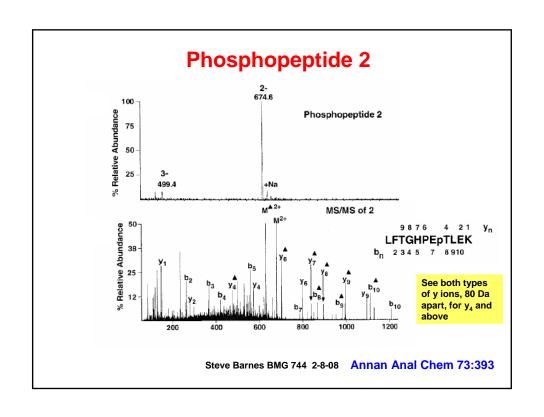
- use of parent ion or neutral loss scanning
- phosphatase sensitivity
- affinity methods for enrichment of phosphopeptides
 - anti-phospho-Ser/Thr/Tyr antibodies
 - metal ion affinity
 - chemical reaction/biotin affinity

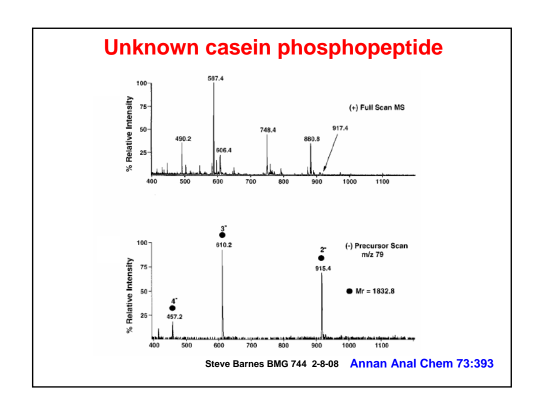
Parent ion scanning to detect phosphopeptides

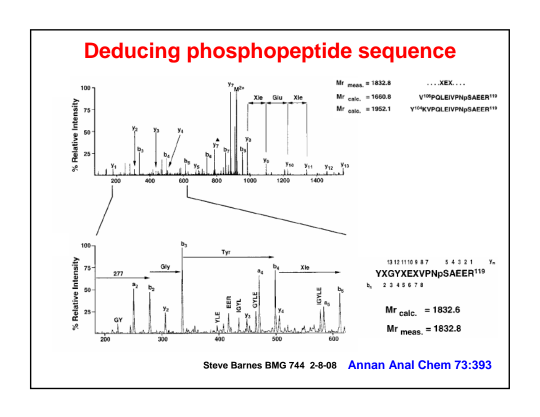
- The procedure depends on the detection of the m/z 79 ion fragment (PO₃-) during collision-induced dissociation in a triple quadrupole instrument operating in the negative ion mode
- Parent ion scanning is a reversal of the more familiar daughter ion MS-MS where the parent ion is selected (in Q1) and a mass spectrum of the daughter ion fragments is obtained by scanning in Q3
- In parent ion scanning, the daughter ion fragment (in this case m/z 79) is held constant in Q3 and a mass spectrum of parent ions that give rise to the daughter ion obtained by scanning in Q1.
- Having identified the phosphopeptides, the sample can be reanalyzed to obtain daughter ion MS-MS spectra on selected ions in the positive ion mode











Recovery and enhancement of phosphopeptides

The biggest problem in the detection of phosphopeptides is how to convert the initial sample matrix into a form suitable for mass spectrometry analysis.

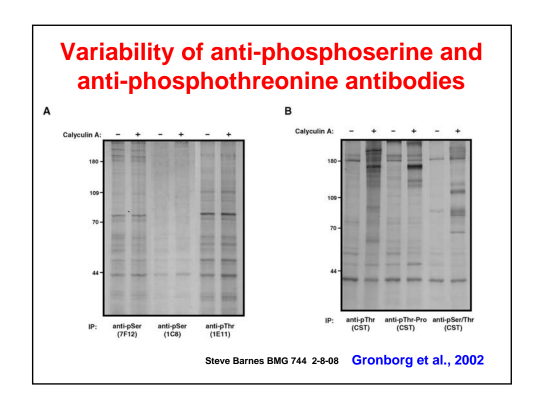
- how to handle minute samples with minimal losses
- how to recover and detect all the phosphopeptides
- how to recover and detect the non-phosphorylated proteins to determine the extent of phosphorylation at individual sites

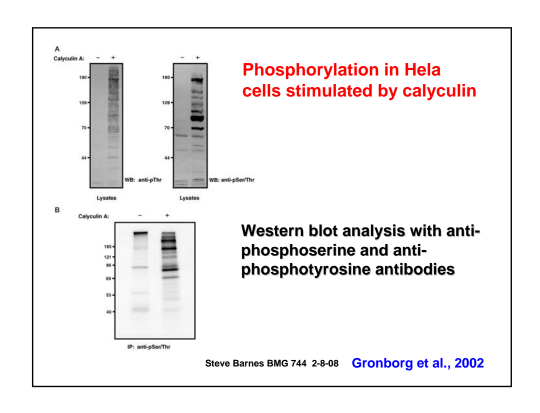
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Antibodies and phosphopeptides

- In this approach, both the phosphorylated and nonphosphorylated forms of a protein may be recovered from the sample matrix
- This can most easily achieved by immunoprecipitation of the protein with an antibody that recognizes epitope(s) that is(are) in common with both forms

NOTE that as with all immunoprecipitation methods, the best results will be achieved if the antibody is coupled to agarose beads. This allows selective immuno-absorption of the antigen, washing free of contaminating proteins, AND following elution, minimization of the amount of antibody that is in the eluate and therefore would be analyzed by mass spectrometry





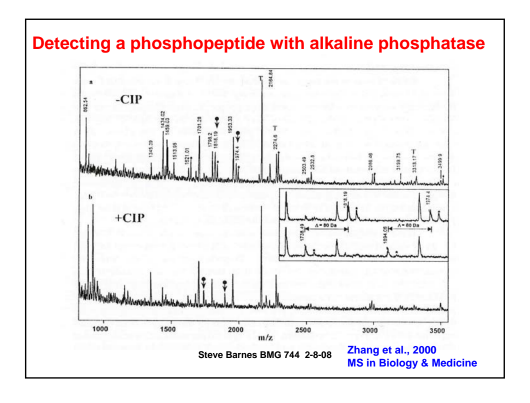
Detection of phosphopeptides based on their sensitivity to phosphatase

- An alternative source of potentially phosphorylated proteins are individual spots on 2D-IEF/SDS gels. The protein preparation so isolated is either hydrolyzed by trypsin in solution (or in the gel piece) or using solidphase trypsin
- One portion of the resulting tryptic peptides (in 50% acetonitrile:water) is analyzed by MALDI-TOF-MS. A second portion is diluted into 50 mM NH₄HCO₃ buffer and reacted with 0.5 U calf intestinal alkaline phosphatase at 37°C for 30 min. Sample is dried with a SpeedVac, redissolved in 50% acetonitrile:water, and reanalyzed by MALDI-TOF-MS

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Identifying phosphopeptides using alkaline phosphatase

- Phosphopeptides shift down by m/z 80 (or units of 80 in the case of multiply phosphorylated peptides)
- The peaks identified as phosphopeptides can then be analyzed in a nanoelectrospray experiment where collision-induced dissociation is used to determine the identity of the peptide and the phosphorylation site in the sequence



Selective enhancement of phosphopeptides in tryptic digests

- Immobilized metal affinity chromatography (IMAC). Similar to Niaffinity resins used in the purification of 6xHis-tagged proteins.
 The affinity phase can be charged with different metal ions (as their chlorides)
- Fe(III) and Ga(III), and to a lesser extent Zr(IV), were the most effective for the recovery of two synthetic phosphopeptides
- A tryptic digest containing both phosphorylated and nonphosphorylated peptides is passed over the IMAC column at acid pH (pH 2.5-3)
- The column is washed with 0.1 M acetic acid to remove unbound peptides
- Elute with sodium phosphate (have to desalt) or with NH₄OH
- Esterification may prevent Asp- or Glu-containing peptides from binding

Selective biotinylation of phospho-groups

How to identify phosphorylated peaks by searching databases

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- The databases you have used earlier in this class have some ability to predict the expected masses for a limited number of posttranslational modifications
 - MASCOT (http://www.matrixscience.com)
 - PROTEIN PROSPECTOR (http://prospector.ucsf.edu/)

How to identify posttranslational modifications

FindMod at

http://www.expasy.org/tools/findmod/

It examines mass fingerprinting data for mass differences between empirical and theoretical peptides. If the mass difference corresponds to a known modification, it also makes intelligent guesses as to the site of modification.

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FindMod

Some of the modifications considered are:

acetylation C-mannosylation farnesylation γ-carboxyglutamic acid methylation O-GIcNac pyridoxal phosphate sulfation

amidation deamidation formylation hydroxylation myristoylation palmitoylation

biotinylation flavinylation geranyl-geranylation lipoylation N-acyl diglyceride phosphorylation phospho-pantetheine pyrrolidone-carboxylic acid

NOTE that none of the common chemical modifications (alkylation of sulfhydryl groups with iodoacetic acid, iodoacetamide, 4-vinylpyridine, and acrylamide) were included.

Wilkins M.R., Gasteiger E., Gooley A., Herbert B., Molloy M.P., Binz P.A., Ou K., Sanchez J.-C., Bairoch A., Williams K.L, Hochstrasser D.F. High-throughput Mass Spectrometric Discovery of Protein Post-translational Modifications. Journal of Molecular Biology, 289, p. 645-657 (1999)

Gasteiger E., Hoogland C., Gattiker A., Duvaud S., Wilkins M.R., Appel R.D., Bairoch A.; Protein Identification and Analysis Tools on the ExPASy Server; (In) John M. Walker (ed): The Proteomics Protocols Handbook. Humana Press (2005).

Site for compilation of PTMs

https://www.abrf.org/index.cfm/dm.home

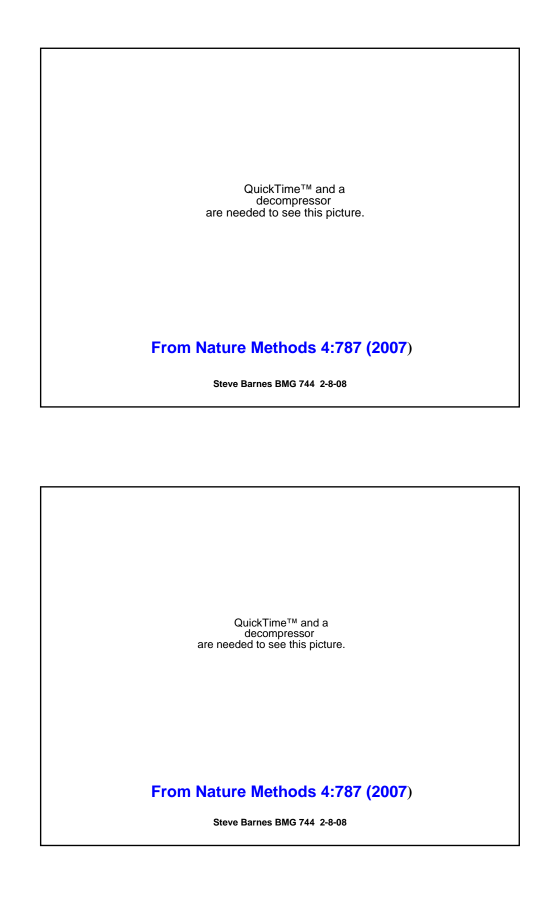
This site was put together by Ken Mitchelhill, Len Packman and friends

Currently ranges from dephospho (-79) to (Hex)3-HexNAc-(dHex)HexNAc (+1,039)

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QuickTime™ and a decompressor are needed to see this picture.

From Nature Methods 4:787 (2007)



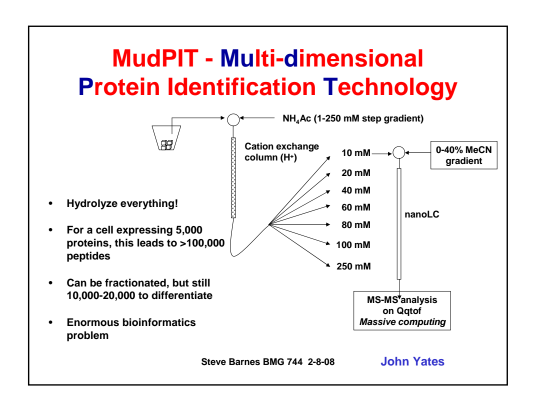
Detecting PTMs

- If we don't have any idea about what the modification(s) is(are), how do we proceed?
- We won't have the modification on MASCOT or Protein Prospector
- No antibodies or convenient affinity phases

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Can we approach this globally?

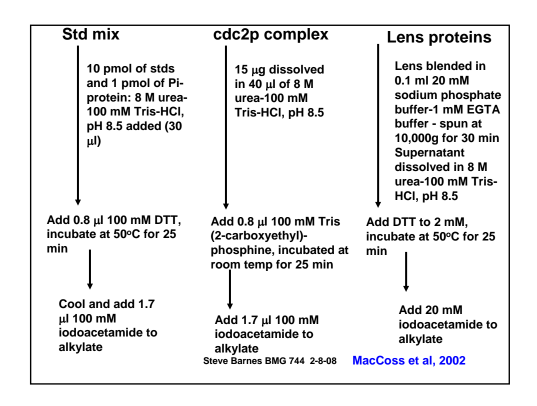
- It's asking a lot
 - Too many degrees of freedom
 - Endless modifications
- Should we try?
 - John Yates' group has
 - MacCoss et al. PNAS 99:7900 (2002)

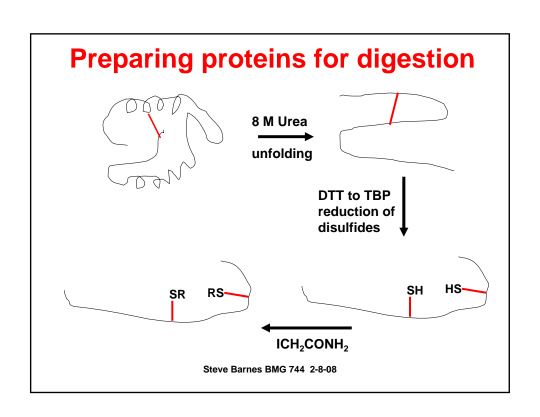


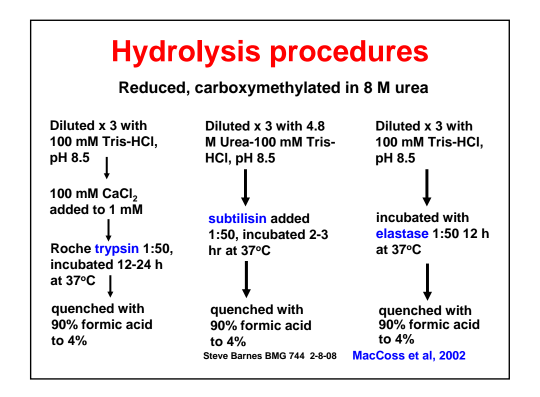
The Yates approach - selection of "proteomes"

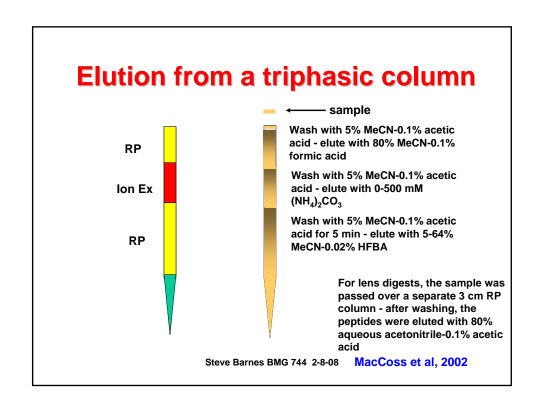
- BioRad mixed molecular weight standards (10 pmoleach) plus 1 pmol of a phosphorylated glycogen phosphorylase
- Cdc2 protein complex isolated with TAP (15 μg)
- · Lens proteins from 4-yr old

Steve Barnes BMG 744 2-8-08 MacCoss et al, 2002





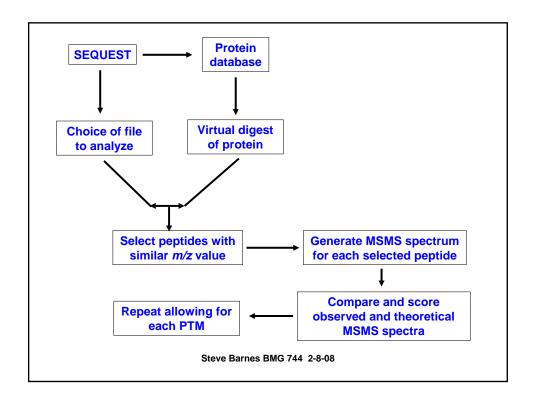




Automated MS-MS analysis

- Limit analysis to 2+ or 3+-charged peptides
- Delete poor quality spectra
- Identify peptides with 98-Da neutral loss
- Analyze the remaining spectra with SEQUEST (operating on a 31-node beowulf computer cluster) to identify proteins
- Search sequences of identified proteins for PTMs by 80 (STY phosphorylation), 42 (K acetylation), 16 (MWY oxidation) and 14 (K methylation)

Steve Barnes BMG 744 2-8-08 MacCoss et al, 2002



Results

Protein standard mixture:

- The three digests were combined 83.7% and 95.4% coverage for glycogen phosphorylase and BSA
- Identified the glycogen phosphorylase phosphorylation site as well as the two known sites (S69/S345) in ovalbumin
- N-acetylation found plus many sites of methionine oxidation (? due to work up or real)
- New sites found phosphorylation at S237/S241 in ovalbumin and methylation at R652

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Results

Cdc2p complex:

- 200 proteins 20 showed >40% sequence coverage
- Expected Y15 and T167 phosphorylation sites on Cdc2p found
- New phosphorylation sites found on cyclin partners Cdc13 and Cig1p
- Multiple methylation sites on Cdc2p

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Results

Lens proteins:

- Found 272 proteins 52 had >40% sequence coverage
- 90% are crystallins
- PTMs accumulate over your lifetime
- Used 18-step MudPIT because of complexity
- 73 different PTMs found on the 11 crystallins
- Found the 13 of 18 PTMs previously described in all species
- Found 60 other new PTMs in phosphorylation, oxidation, acetylation and methylation

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Quantitative proteomics

Use of isotopes

- ICAT (d_o/d₈) and ICAT 13 C₀/ 13 C₈
- d₀/d₁₀ propionic anhydride (N-terminal labeling)
- ¹⁵N/¹⁴N (whole cell labeling)
- ¹⁸O/¹⁶O (trypsin)
- iTRAQ labeling
- Non-isotope methods
 - Peptide coverage
 - Classical triple quadrupole methods

Quantification methods in mass spectrometry

From review in Nature Methods 4-79 (2007)

QuickTime™ and a decompressor are needed to see this picture.

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Isotope-coded affinity technology

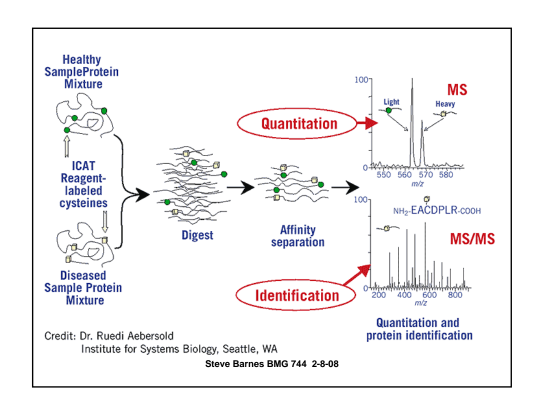
Isotope-Coded Affinity Tags

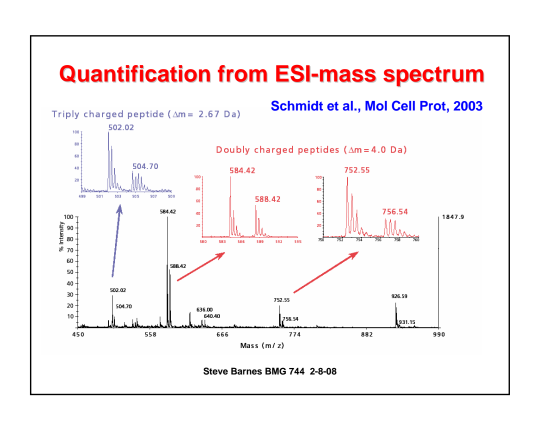
heavy reagent: D8-ICAT Reagent (X=deuterium)

light reagent: DO-ICAT Reagent (X=hydrogen)

This reagent reacts with cysteine-containing proteins (80-85% of proteome)

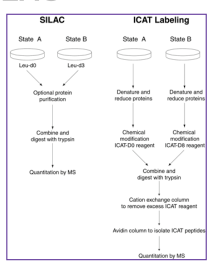
Labeling can be replacement of hydrogens (X) with deuterium, or better to exchange ¹²C with ¹³C in the linker region (this avoids chromatography issues)





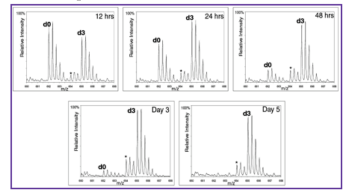
SILAC

 SILAC, stable isotope labeling by amino acids in cell culture, is being used to quantify proteins.



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Time-dependent leucine incorporation with SILAC



The cells are pre-labeled with leucine-d₀. Leucine-d₃ is added to the medium and cells sampled at various times later. The peaks annotated with d0 and d3 are the triply charged peaks of the peptide VAPEEHPVLLTEAPLNPK, which contains three leucines.

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¹⁸O-labeling

 Trypsin catalyzes the transfer of ¹⁸O in ¹⁸O-enriched water to both the carboxylate oxygens of the C-terminus of tryptic peptides

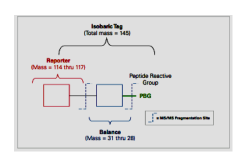
R-COOH
$$\longrightarrow$$
 R-C¹⁸O₂H

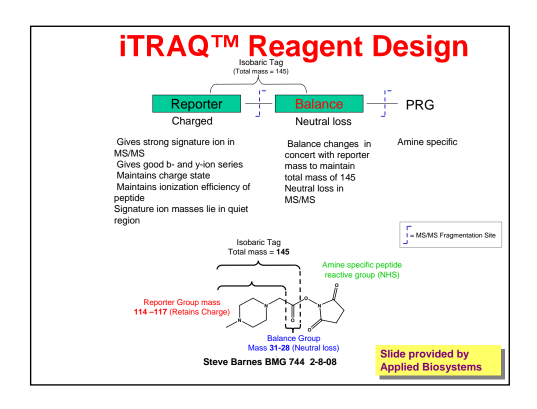
- The peptides have an increase in mass of 4 Da
- Generally not considered a large enough mass difference

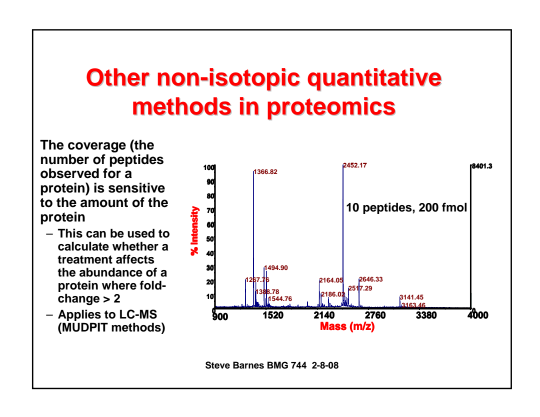
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iTRAQ quantification

- The iTRAQ™ reagents
 - React with Lys amino groups and each one adds 145 Da to the molecular weight of the peptide
 - Fragmentation produces reporter ions from m/z 114, 115, 116 and 117
 - New iTRAQ kit contains 8 forms with reporter fragment ions of m/z 114, 115, 116, 117, 118, 119 and 121



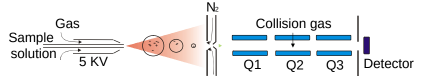




Triple quad MRM analysis

Peptides of interest can be analyzed like small molecules

 Choose the parent molecular ion, collide with argon gas and select a unique fragment



· Multiple reaction ion scanning

First filter the [M-H]- molecular ion of the analyte (Q1)

Fragment the molecular ion with N₂ gas (Q2)

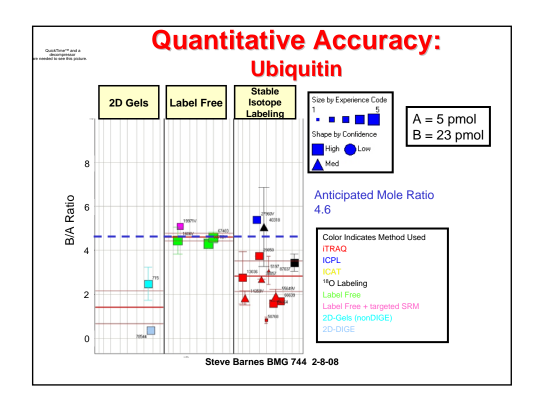
Select a specific (and unique) fragment ion (Q3)

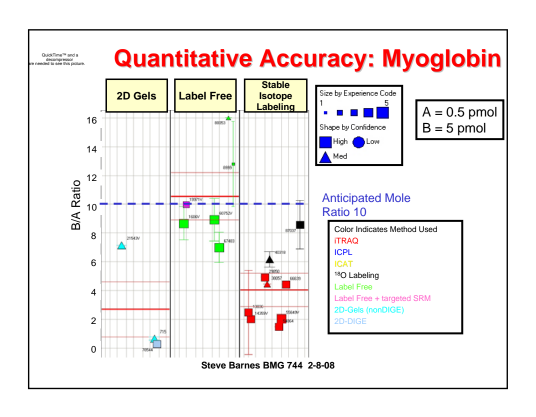
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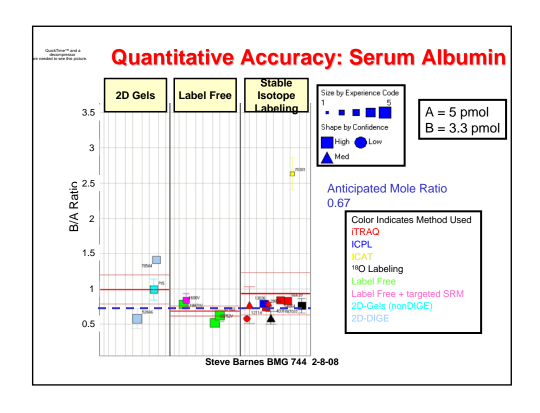
Quantitation experiment for biotinylated cytochrome c MRM analysis monitored in 50 channels 4.5e5 4.0e5 2.0e5

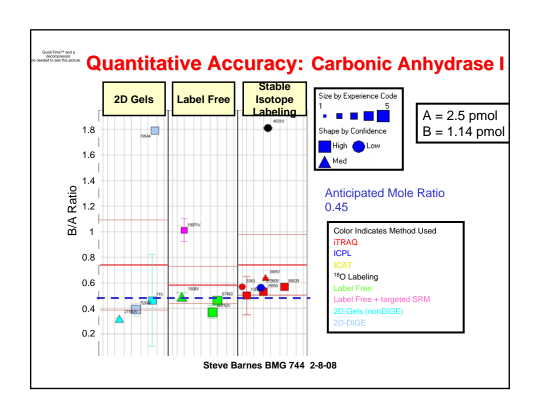
Each colored peak represents a different biotinylated peptide

Time, min







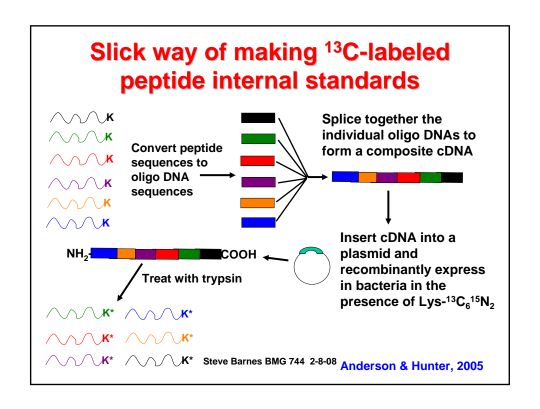


Application of LC-MRM-MS to the plasma proteome In silico, generate Identify the proteins of interest

Select the best combination of parent peptide and + fragment y ion

the tryptic peptides from each protein

Determine the expected y ions for each peptide and compare to y ions of all other tryptic peptides of known human proteins that have masses within + 1Da



Quantitative peptide MRM-MS

- The albumin-depleted plasma proteome is mixed with the composite ¹³C,¹⁵N-labeled protein internal standard and then treated with trypsin
- The molecular ions (doubly charged) and the specific y ions for each peptide and its labeled form are entered into the MRM script one channel at a time
- A single run may consist of 30 peptides in 60 channels
- Sensitivity is compromised by "sharing out" measurement time, but can be compensated for by carrying out nanoLC

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Advantage of a C-terminal labeled lysine

```
186 301 448 505 642 755 886 987 1115 bions

A D E F G H I M T K

1133 1062 948 833 686 629 492 379 248 147 yions
```

With the labeled lysine at the C-terminus, only the b₁₀ ion contains the isotope atoms

```
186 301 448 505 642 755 886 987 1123 bions

A D E F G H I M T K*

1141 1070 956 841 694 637 500 387 256 155 yions
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References for this talk (1)

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