

Proteomics and Mass Spectrometry 2008

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Course plan

- Meet Tuesdays/Fridays in MCLM 401 from 9-11 am (Jan 4-Mar 20)
- Graduate Students taking this course are required to attend each session
- Evaluations will be made from projects/exams
- Where possible, materials from each class will be placed on the proteomics website (go to <http://www.uab.edu/proteomics> - click on **Class**)

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Recommended general texts

- **Suggested text - “*Introduction to Proteomics*”** by Daniel C. Liebler, 2002
- **Also see “*The Expanding Role of Mass Spectrometry in Biotechnology*”** by Gary Siuzdak (a 2003 edition of the 1996 first edition)
- **“*Mass spectrometry data analysis in proteomics*”,** (ed., Mathiesson, R) in *Methods in Molecular Biology*, vol 367.

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Suggested readings

- Kenyon G, et al. *Defining the mandate of proteomics in the post-genomics era: workshop report*. *Mol Cell Proteomics*, 1: 763-780 (2002)
- Hood L. *Systems biology: integrating technology, biology and computation*. *Mechan Aging Develop*, 124: 9-16 (2003)
- Kim H et al. *Proteomics and mass spectrometry in nutrition research*. *Nutrition*, 20: 155-165 (2004)
- Righetti P. et al. *Prefractionation techniques in proteome analysis: the mining tools of the third millennium*. *Electrophoresis*, 26: 297-319 (2005)
- Graham DR et al. *Broad-based proteomic strategies: a practical guide to proteomics and functional screening*. *J Physiol*, 563:1-9, (2005)
- Anderson NL. *The roles of multiple proteomic platforms in a pipeline for new diagnostics*. *Mol Cell Proteomics* (2005)
- Anderson JS & Mann M. *Organellar proteomics: turning inventories into insights*. *EMBO Reports*, 7: 874-879 (2006)

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BMG/PHR 744 - section 1

- Jan 4, Fri S. Barnes/H. Kim
The world of biomolecules. The proteome, proteomics and other -omics and where to start
- Jan 8, Tue S. Barnes
Methods for the identification of proteins: MALDI-TOF of proteins and peptide mass fingerprinting; LC analysis and peptide sequencing
- Jan 11, Fri M. Renfrow
Mass spectrometry – gas phase transfer and instrumentation
- Jan 15, Tu S. Barnes
Ion fragmentation in mass spectrometry; application to proteomics
- Jan 18, Fri J. Prasain
Ion Fragmentation and small molecules/metabolomics
- Jan 22, Tu S. Barnes
Separating your samples from the matrix

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BMG/PHR 744 - section 2

- Jan 25, Fri J. Mobley
Connecting proteomics into bioinformatics; MUDPIT and SEQUEST; false discovery rates in complex systems
- Jan 29, Tu C. Crasto
The bioinformatics of the proteome
- Feb 1, Fri H. Kim
Simplifying the proteome - techniques of protein purification; protein separation and analysis by electrophoresis and other 2D-approaches
- Feb 5, Tu J. Mobley/J. Prasain
Experimental design issues in proteomics and metabolomics
- Feb 8, Fri S. Barnes
Mass spectrometry in qualitative and quantitative burrowing of the proteome
- Feb 12 J. Prasain
Qualitative and quantitative validation in metabolomics

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BMG/PHR 744 - section 3

Feb 15, Fri	P. Prevelige <i>Mass Spectrometry as a Tool for Studying Protein Structure</i>
Feb 19, Tu	P. Prevelige <i>Study of macromolecular complexes</i>
Feb 22, Fri	S. Barnes <i>Enzymology and mass spectrometry</i>
Feb 26, Tu	M. Renfrow <i>Analysis of protein-protein interactions by affinity purification and mass spectrometry</i>
Feb 29, Fri	C-C. Wang <i>Tissue and body fluid proteomics</i>
Mar 4, Tu	J. Mobley <i>Tissue imaging by mass spectrometry</i>

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BMG/PHR 744 - section 4

Mar 7, Fri	M. Renfrow <i>Applications of FT-ICR-MS</i>
Mar 11, Tu	J Novak, M Renfrow <i>Mass spectrometry in glycomics research - Application to IgA nephropathy</i>
Mar 14, Fri	TBA <i>Mass spectrometry and biodefense</i>
Mar 17, Tu	H. Kim <i>Use of proteomics and MS methods in the study of the brain proteome and neurodegenerative diseases</i>
Mar 20	Final report due

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Course learning objectives

- **Introduction to the concepts and practice of systems biology**
- **Sample ionization and mass spectrometers**
- **Mass spectrometry and its principal methods**
 - **protein and peptide ID; ion fragmentation; stable isotope labeling; quantification**

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Course learning objectives

- **Informatics, statistics and quality control in mass spectrometry**
- **Importance of prefractionation in proteomics - 2DE, LC and arrays**
- **Applying mass spectrometry to protein modifications, function, structure and biological location, and to other biological molecules**

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Hopes and hazards of biomedical research

It boils down to whether having taken life apart into its distinct pieces, can we reassemble it in new ways? [*strong analogies to what have been the central quests of physics*]

Can we create a form of life that might live in a very hostile extra-terrestrial environment and thereby save humanity?

Or will we (*as well as our enemies*) instead create life forms that can terrorize or even eliminate us?
Will Einstein's and Oppenheimer's moral dilemmas surface in biomedical science?

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History of proteomics

- Essentially preceded genomics
- “Human protein index” conceived in the 1970’s by Norman and Leigh Anderson
- The term “proteomics” coined by Marc Wilkins in 1994
- Human proteomics initiative (HPI) began in 2000 in Switzerland - <http://www.hupo.org>
- Human Proteome Organization (HUPO) had meetings in 2002 in Versailles, France; 2003 in Montreal, Canada; 2004 in Beijing, China; 2005 in Munich, Germany; 2006 in Long Beach, CA; 2007 in Seoul, Korea. The 2008 meeting will be Amsterdam; 2009 in Toronto and 2010 in Sydney

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What proteomics is, what it isn't

**“Proteomics is not just a mass spectrum of
a spot on a gel”**

George Kenyon,
2002 National Academy of Sciences Symposium

**Proteomics is the identities, quantities, structures,
and biochemical and cellular functions of all
proteins in an organism, organ or organelle, and
how these vary in space, time and physiological
state**

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Collapse of the single target paradigm - the need for systems biology

Old paradigm

***Diseases are due
to single genes -
by knocking out
the gene, or
designing specific
inhibitors to its
protein, disease
can be cured***

But the gene
KO mouse
didn't notice
the loss of the
gene



New paradigm

***We have to
understand gene
and protein
networks -
proteins don't act
alone - effective
systems have
built in
redundancy***

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Research styles

- **Classical NIH R01**
 - A specific target and meaningful substrates
 - Emphasis on mechanism
 - Hypothesis-driven
 - **Linearizes locally multi-dimensional space**
- **Example**
 - Using an X-ray crystal structure of a protein to determine if a specific compound can fit into a binding pocket - from this “*a disease can be cured*” - this approach ignores whether the compound can get to the necessary biological site, whether it remains chemically intact, and where else it goes

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From substrates to targets to systems - a changing paradigm

- **Classical approach** - **one substrate/one target**
- **Mid 1980s** - use of a pure reagent to isolate DNAs from cDNA libraries (**multiple targets**)
- **Early 1990s** - use of a reagent library (**multiple ligands**) to perfect interaction with a specific target
- **2000+** - effects of specific reagents on cell systems using DNA microarrays (**500+ genes change, not just one**)
- **2008** - integration of transcriptomics, proteomics, peptidomics, metabolomics (**a single change affects everything, not just one thing, as in ecology**)

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Exploring information space - the *Systems Biology* approach

- **Systems biology means measuring everything about a system at the same time**
- **For a long time, it was deemed as too complex for useful or purposeful investigation**
- **But are the tools available today?**

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Systems Biology

“To understand biology at the system level, we must examine the structure and dynamics of cellular and organismal function, rather than the characteristics of isolated parts of a cell or organism.”

“Properties of systems, such as robustness, emerge as central issues, and understanding these properties may have an impact on the future of medicine.”

*“However, many breakthroughs in experimental devices, advanced software, **and analytical methods** are required before the achievements of systems biology can live up to their much-touted potential.”*

Kitano, 2002

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Biological Data of the Future

- Destructive
- Qualitative
- Uni-dimensional

- Low temporal resolution
- Low data density
- Variable standards
- Non cumulative

- Non-destructive
- Quantitative
- Multi-dimensional and spatially resolved
- High Temporal resolution
- High data density
- Stricter standards
- Cumulative

CURRENT NATURE of DATA

Elias Zerhouni, FASEB 2004

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Techniques in Systems Biology

- DNA microarrays to describe and *quantify* the transcriptosome
- Large scale and small scale proteomics
- Protein arrays
- Protein structure
- Metabolomics
- Integrated computational models

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Papers on systems biology - 2006

<http://www.nature.com/focus/systemsbiology/userguide/index.html>

This is a cluster of papers in Nature Cell Biology, Nature Reviews in Molecular and Cell Biology, and Molecular Systems Biology that was published in October, 2006

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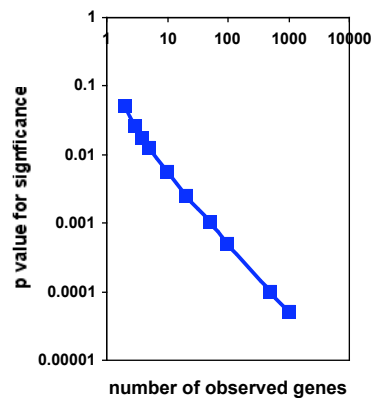
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High dimensionality of microarray or proteomics data means you must understand and deal with statistics

While reproducible data can be obtained, the large numbers of parameters (individual genes or proteins) require large changes in expression before a change can be regarded as significant

use of the Bonferroni correction
A conservative correction



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Properties of a system and fold-change

- The primary assumption of most users of DNA microarray data (and proteomics data) has been that the cut-off for assessing biologically meaningful change is two-fold
- This is a very naïve view of properties of a system

“Fold-change is inversely related to biological importance”

Barnes’ law

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Properties of a system and fold-change

- For a system, items that are important are the least likely to change
 - when they do, then catastrophic events will occur
 - Proliferation vs apoptosis (PTEN < 50% change)
- A system may tolerate a lot of variance in molecules that are “unimportant” to the system”
- How can we perceive “importance”?
 - Re-weight the data by dividing by the variance
 - Need to have enough information about each item to calculate its variance ($n > 5$)

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Vulnerability of a system

- **To really understand biological systems, you have to appreciate their dynamic state**
 - Read about control theory
 - Realize that systems are subject to rhythms
 - Subject them to fourier transform analysis to detect their resonance (requires far more data than we can currently collect)
- **A small signal at the right frequency can disrupt the system**
 - Analogies “the small boy in the bath” and “the screech of chalk on a chalk board”

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Hazards of interpreting microarray (proteomic) data

- **“Expression patterns are the place where environmental variables and genetic variation come together. Environmental variables will affect gene expression levels.”**
- **“Don’t we need to be very careful to understand the environmental inputs that might have an impact on that expression? Perhaps an over-the-counter herbal supplement might cause an expression pattern that looks like that of a very aggressive tumor.”**

Abridged from Karen Kline, 2002

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Why study the proteome when we can do DNA microarrays?

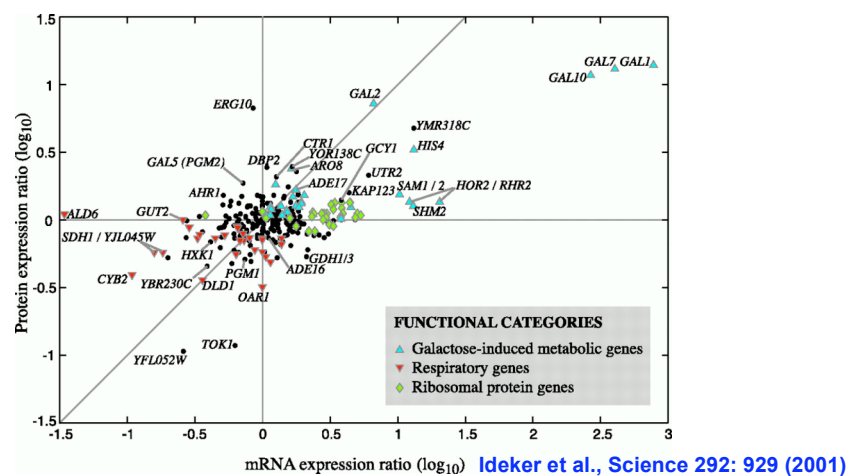
- DNA microarray analysis allows one to examine the mRNA levels of thousands and thousands of genes
- However, the correlation between gene expression and protein levels is poor at best
- Is this a new finding? No, before the age of genomics, it was well known

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Apparent poor relationship between gene expression and protein content



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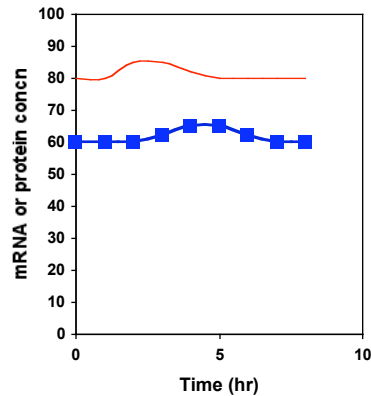
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Housekeeping genes and proteins are related

This is the relationship between mRNA (red) and protein (blue) levels expression of a house-keeping gene/protein, i.e., one that has to be expressed at all times

- Even with the small perturbation, the amounts of mRNA and protein are well correlated to each other



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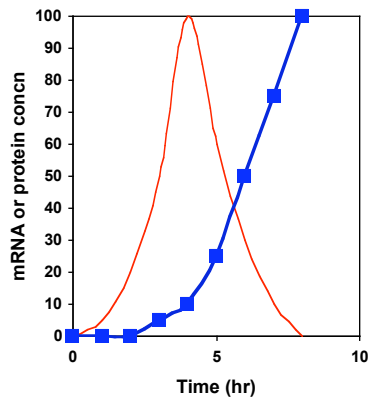
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(Barnes & Allison, 2004)

Sampling time affects interpretation of correlation between mRNA and protein expression for important proteins

Sometimes, determining the relationship between mRNA (red) and protein (blue) levels depends totally on when you measure them - for the figure opposite, the ratio at 2.5 hr is 10:1, whereas at 7.5 hr it's 1:100

- better to measure the ratio over time and integrate the area under the curve



(Barnes & Allison, 2004)

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Defining disease from the proteome

- Numerous examples of a revised picture of disease from analysis of the proteome
 - Aging
 - Cancer
 - Cardiovascular disease
 - Neurodegeneration
- Infectious disease and the microbial proteome

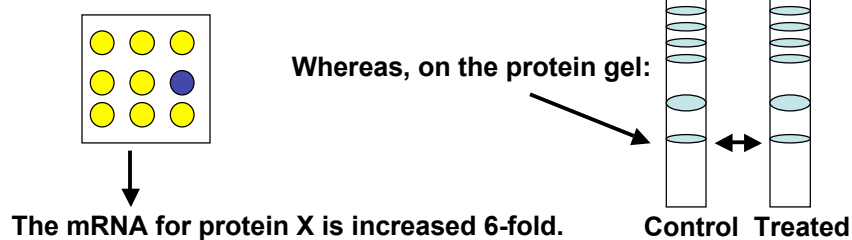
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Upregulation of a gene for a crucial protein may have poor correlation at the protein level: example A.

- May be manifested as little or no change in amount, because the cell “damps” out the change.

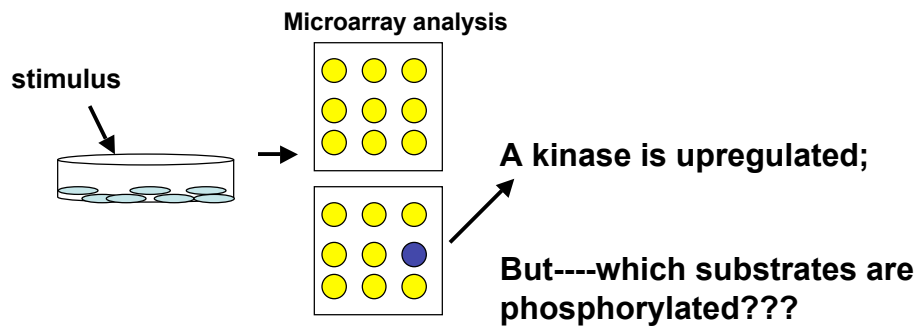


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Gene expression data may predict the nature of protein modifications, *but not which proteins are modified.*



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Summary: Why do we need to do broad scale “proteomics”?

- **When you don’t have a clue;**
- **When you have only a very small clue;**
i.e. you’ve done a microarray experiment, and you have a list of 3,284 genes that are differentially regulated in your system;
- **When you knock out a gene (and hence a protein) that you’re convinced is essential for life and health, and the animal pees as usual.**

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Rationales for proteomics approaches in today's research

- **Identify a biomarker(s), as an end unto itself;**
 - Cancer detection/Monitor response to chemotherapy
 - Identify one pathogen from others;
 - Distinguish a virulent strain of pathogen from nonvirulent.
- **Characterize protein expression profiles between diseased versus healthy tissues--**
 - For understanding the disease process;
 - To develop drug targets;
- **In cancer, there may be novel proteins due to chromosome instability (ETV6-ABL and BCR-ABL), or inappropriate expression may occur (proteins from embryonic or fetal stages of development)**

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Where there is pathology, but the genetic basis unknown, proteomics can have critical role in identifying proteins to target for therapeutic intervention

Two major disease examples:

- **HIV: protease is targeted today;**
 - are there other proteins, either viral or host, that could be targeted to better deal with the disease?
- **Alzheimer's disease: 3 known mutations (APP, PS1, PS2) and risk factors (ApoE, estrogen loss);**
 - 50% of AD patients do not have any of the known genetic abnormalities, yet all become demented, all have amyloid plaques and NFT in their brains.
 - Remember, every AD patient has AD 100%.

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Challenges in proteomics

- So many protein forms! How many 100-mers are possible using 20 different amino acids?
- Genomic sequence doesn't always predict the encoded protein:
 - Posttranslational modifications
 - Differential splicing
 - Unknown transcriptional mechanisms
- The dynamic range ($>10^9$)
- No equivalent to the PCR reaction
 - Avogadro's number

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Avogadro's number comes back to haunt us: a reality check for proteomics

- One gram-mole of anything is 6.02×10^{23} molecules
- For mass spec detection and analysis, we need at least 100 femtomole (10^{-13} moles) or 6.02×10^{10} molecules

<u>protein expression per cell</u>	<u>need # cells for 100 fmoles</u>
10	6.02×10^9
100	6.02×10^8
1,000	6.02×10^7
10,000	6.02×10^6
100,000	6.02×10^5

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Central issue in proteome complexity: dynamic range

Cellular proteomes involve a very wide *dynamic range*: proteins can differ in their amounts by nine (9) orders of magnitude;

- In blood, albumin is 3.5 g/100 ml (35 g/L = 0.5 mM) (10^{-3} M), whereas cytokines are pM (10^{-12} M)
- A 2D gel that is overloaded with respect to an abundant protein, may have *barely detectable* amounts of a low abundance protein.....And if you can't see it, you can't analyze or quantify it!!!!

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The need to isolate the lower abundant proteins

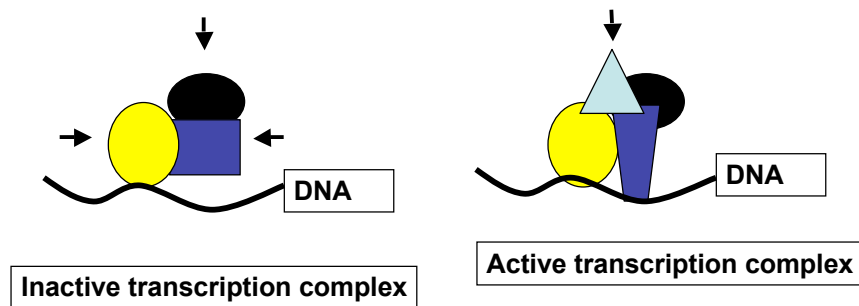
- See lecture on February 1 for expanded discussion of the topic of protein purification
- Parameters of proteins that are the basis for purifying them from each other:
 - Biological properties of proteins:
 - Intracellular location; Protein-protein interactions; Posttranslational modifications
 - Intrinsic properties:
 - Net charge; Size; Extent of tertiary structure; Hydrophobicity

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A valuable role of the proteomics approach:
Discovery & analysis of protein-protein interactions



(The arrows indicate the proteins that could be antigens for immunoaffinity purification of the complex.)

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Experimental design and quality control issues

- **How do we carry out a proteomics experiment?**
 - Randomize sample analysis
 - Process samples blinded to identities
 - Standardize procedures and vendors of disposable plastics used in experiment-- minimize variation where possible
 - Consult with statistician before experiment; ensure enough “power” for the experiment so that statistical analysis yields significant data.

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Take home lessons in analyzing proteins with proteomics methods

- The fewer proteins in the proteome you analyze, the better the chances of detecting the ones that “matter.”
- Genomics data can complement proteomics data.
- Understanding the biological properties of the proteins of interest can enhance proteomics analysis.
- Intrinsic properties of proteins form the basis of invaluable prefractionation prior to proteomics analysis.
- Quality control is an issue that becomes increasingly important with large datasets and measurement of small changes

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Predicting the proteome

- *Bioinformatics* is the basis of high throughput proteome analysis using mass spectrometry. Protein sequences can be computationally predicted from the genome sequence
- However, *bioinformatics* is not able to predict with accuracy the sites or chemistry of posttranslational modifications - these need to be defined chemically (using mass spectrometry)

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Predicting the proteome

- ***Predicting the proteome has elements of a circular argument***
 - protein sequences were initially determined chemically and were correlated with the early gene sequences. It then became easier to sequence a protein from its mRNA (captured from a cDNA library). This could be checked (to a degree) by comparison to peptide sequences. Now we have the human genome (actually two of them).
- ***So, is it valid to predict the genes (and hence the proteome) from the sequence of the genome?***
 - We're doing this in current research. But as we'll see, the mass spectrometer is the ultimate test of this hypothesis -
 - why? because of its mass accuracy

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Protein structure

- **Determined by folding - folding rules not yet defined - cannot predict structure *de novo***
- **X-ray crystallography has been used to produce elegant structural information**
- **NMR and H-D exchange combined with mass spec enable the in solution structure to be determined (see Peter Prevelige's talks on February 15/19)**

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Protein informatics

- **The predicted sequences of the proteins encoded by genes in sequenced genomes are available in many publicly available databases (subject to the limitations mentioned earlier)**
- **The mass of the protein is less useful (for now) than the masses of its fragment ions - as we'll see later, the masses of tryptic peptides can be used to identify a protein in a matter of seconds**

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So, what do we do with all these data?

- **Management of the data generated by DNA microarray and proteomics/protein arrays**
 - High dimensional analysis
- **Beyond the capabilities of individual investigators**
- **Urgent need for visualization tools**
- **The importance of new statistical methods for analysis of high dimensional systems**

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PROTIG and Videocast

- **There is an NIH-based proteome special interest group (PROTIG)**
 - Sign up at <http://proteome.nih.gov>
- **Proteomics and mass spec talks are available for viewing (using Real Player)**
 - Log on at <http://videocast.nih.gov>
 - Podcasts are also available