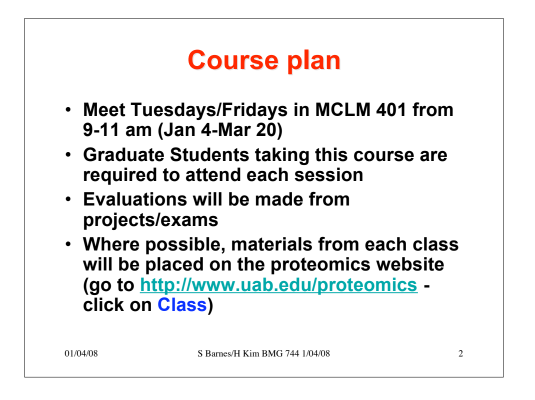
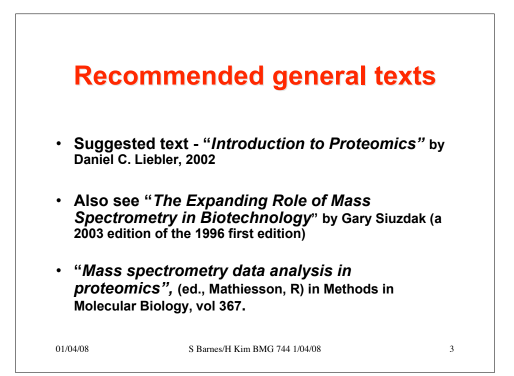
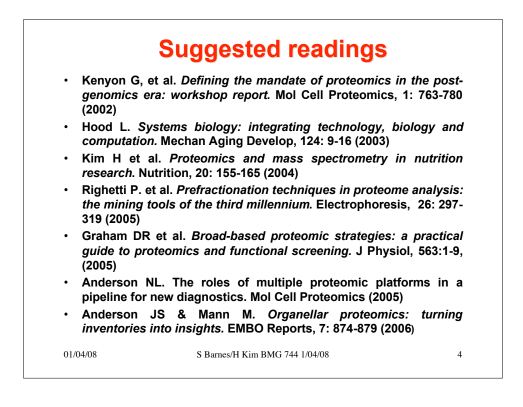
	teomics ectrom		
Stephen B 4-7117, MC <u>sbarnes@</u>		4-388	i Kim, PhD 0, MCLM 460A <u>kim@uab.edu</u>
Jim Mobley, PhD 6-6363, THT 513 mobleyja@uab.edu	Matt Renfrow, 6-4681, MCLM Renfrow@uab	570	Jeevan Prasain, PhD 6-2612, MCLM 456 jprasain@uab.edu
01/04/08	Peter E. Preve 5-5327, BB prevelig@u s Barnes/H Kim BM	elige, PhD RB 416 Iab.edu	







## **BMG/PHR 744 - section 1**

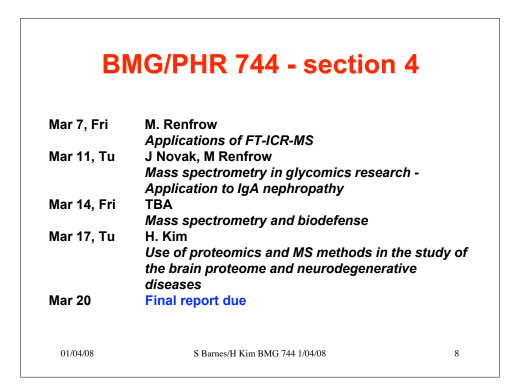
Jan 4, Fri	S. Barnes/H. Kim The world of biomolecules. The proteome, proteomics and other, omics and where to start			
Jan 8, Tue	proteomics and other –omics and where to start S. Barnes Methods for the identification of proteins: MALDI- TOF of proteins and peptide mass fingerprinting; LC analysis and 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	5		
Jan 22, Tu S. Barnes Separating your samples from the matrix				
01/04/08	S Barnes/H Kim BMG 744 1/04/08	5		

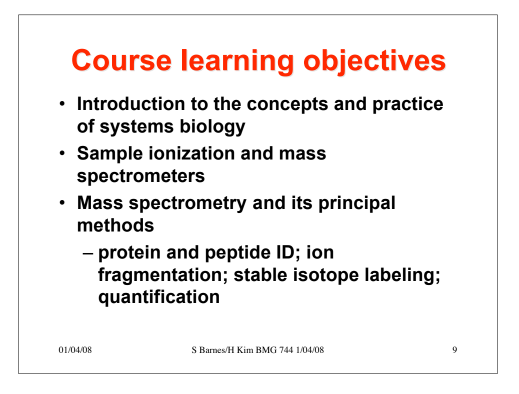
## BMG/PHR 744 - section 2

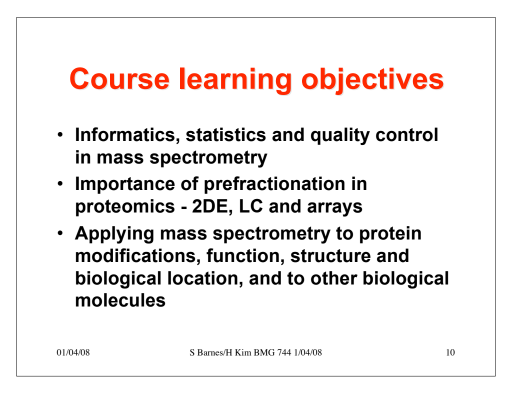
Jan 25, Fri	J.Mobley Connecting proteomics into bioinformatics; MUDPIT and SEQUEST; false discovery rates in complex systems
Jan 29, Tu	C. Crasto The bioinformatics of the proteome
Feb 1, Fri	H. Kim Simplifying the proteome - techniques of protein purification; protein separation and analysis by electrophoresis 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
01/04/08	S Barnes/H Kim BMG 744 1/04/08 6

## **BMG/PHR 744 - section 3**

Feb 15, Fri	P. Prevelige Mass Spectrometry as a Tool for Studying Protein Structure	
Feb 19, Tu	P. Prevelige	
	Study of macromolecular complexes	
Feb 22, Fri	S. Barnes	
	Enzymology and mass spectrometry	
Feb 26, Tu	M. Renfrow	
	Analysis of protein-protein interactions by affinity purification and mass spectrometry	
Feb 29, Fri	C-C. Wang	
	Tissue and body fluid proteomics	
Mar 4, Tu	J. Mobley	
·	Tissue imaging by mass spectrometry	
01/04/08	S Barnes/H Kim BMG 744 1/04/08	7







## 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]

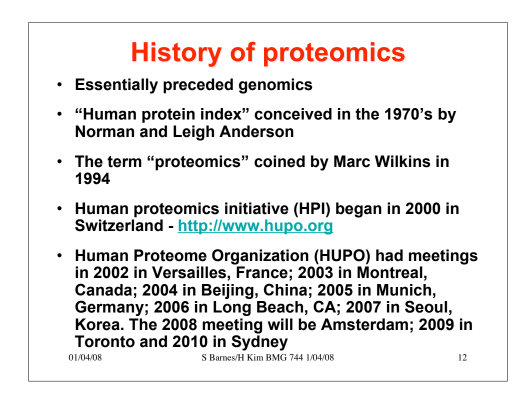
Can we create a form of life that might live in a very hostile extra-terrestial environment and thereby save humanity?

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?* 

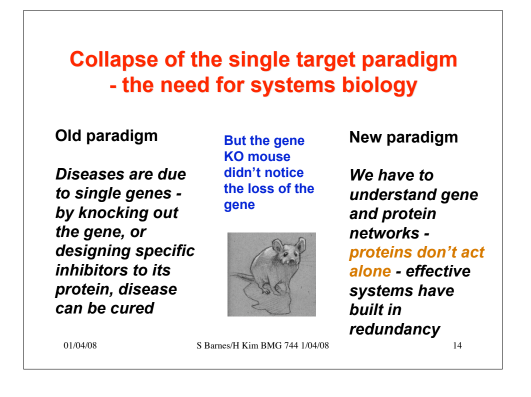
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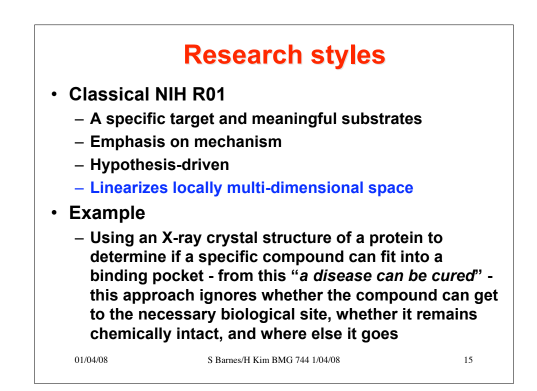
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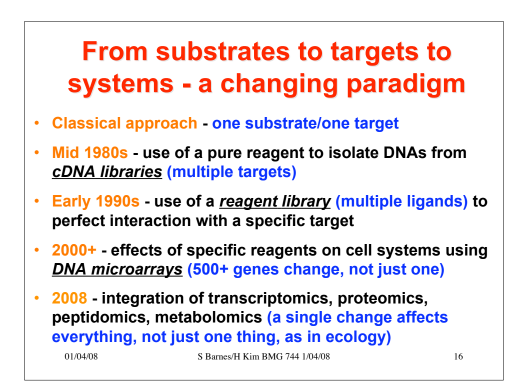
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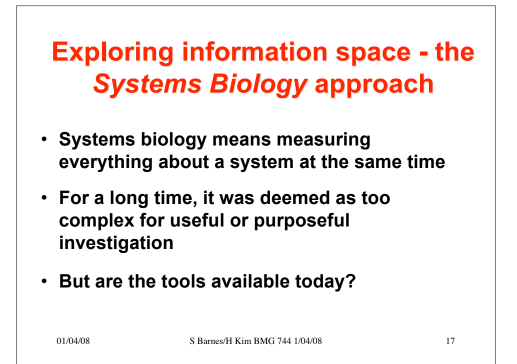


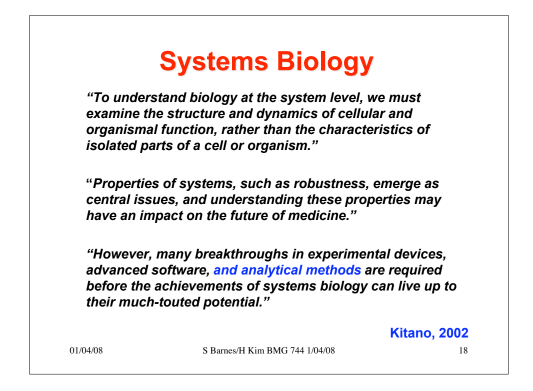
W	hat proteomics is,	ı.
	what it isn't	
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	George Kenyon,	
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and biochem proteins in a	is the identities, quantities, str nical and cellular functions of a in organism, organ or organella ary in space, time and physiolo	all e, and
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E	Biological Data of the Future
Destructive	Non-destructive
Qualitative	Quantitative
Uni-dimensional	<ul> <li>Multi-dimensional and spatially resolved</li> </ul>
<ul> <li>Low temporal resolution</li> </ul>	<ul> <li>High Temporal resolution</li> </ul>
<ul> <li>Low data density</li> </ul>	High data density
<ul> <li>Variable standards</li> </ul>	Stricter standards
Non cumulative	Cumulative
CURRENT NATURE of DATA	Elias Zerhouni, FASEB 200
01/04/08 S Barnes/H Kin	n BMG 744 1/04/08 19

