Open Access

Method

Analysis of the Saccharomyces cerevisiae proteome with PeptideAtlas

Nichole L King*, Eric W Deutsch*, Jeffrey A Ranish*, Alexey I Nesvizhskii[†], James S Eddes*, Parag Mallick[‡], Jimmy Eng^{*§}, Frank Desiere[¶], Mark Flory[¥], Daniel B Martin*[#], Bong Kim*, Hookeun Lee**, Brian Raught*^{††} and Ruedi Aebersold***

Addresses: *Institute for Systems Biology, N 34th Street, Seattle, WA 98103, USA. †Department of Pathology, University of Michigan, Catherine Road, Ann Arbor, MI 48109, USA. †Louis Warschaw Prostate Cancer Center, Cedars-Sinai Medical Center, W. Third St, Los Angeles, CA 90048, USA. §PHSD, Fred Hutchinson Cancer Research Center, Seattle, WA 98109, USA. *Nestlé Research Center, 1000 Lausanne 26, Switzerland. *Department of Molecular Biology and Biochemistry, Wesleyan University, Middletown, CT 06459, USA. *Divisions of Human Biology and Clinical Research, Fred Hutchinson Cancer Research Center, Seattle, WA 98109-1024, USA. *IMSB, ETH Zurich and Faculty of Science, University of Zurich, Zurich, Switzerland. †University Health Network, Ontario Cancer Institute and McLaughlin Centre for Molecular Medicine, College Street, Toronto, ON M5G 1L7, Canada.

Correspondence: Nichole L King. Email: nking@systemsbiology.org

Published: 13 November 2006

Genome Biology 2006, 7:R106 (doi:10.1186/gb-2006-7-11-r106)

The electronic version of this article is the complete one and can be found online at http://genomebiology.com/2006/7/11/R106

Received: 5 July 2006 Revised: 2 October 2006 Accepted: 13 November 2006

© 2006 King et al.; licensee BioMed Central Ltd.

This is an open access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

We present the *Saccharomyces cerevisiae* PeptideAtlas composed from 47 diverse experiments and 4.9 million tandem mass spectra. The observed peptides align to 61% of *Saccharomyces* Genome Database (SGD) open reading frames (ORFs), 49% of the uncharacterized SGD ORFs, 54% of *S. cerevisiae* ORFs with a Gene Ontology annotation of 'molecular function unknown', and 76% of ORFs with Gene names. We highlight the use of this resource for data mining, construction of high quality lists for targeted proteomics, validation of proteins, and software development.

Background

The field of genomics is slowly reaching maturity. The genomes of many organisms have now been sequenced and the effort to annotate these genomes is now well underway. Transcriptomes are routinely investigated, as mRNA expression can be measured with highly sensitive microarrays and other methods. In contrast, the measurement and annotation of proteomes remains challenging. Proteome analysis is primarily based on mass spectrometry (MS) and is not as mature as gene expression analysis. However, proteomic measurements are preferable in some situations because, while mRNA expression studies indicate the potential for protein expression, they do not directly measure proteome character-

istics. For example, mRNA expression levels do not always correlate well with protein expression levels due to variations in translation efficiencies [1] and targeted degradation of proteins in the cell [2,3]. Additionally, proteins are subjected to numerous post-transcriptional modifications that alter the chemical composition of the protein. Proteins also interact with other proteins in a highly dynamic way.

Proteomics by MS has emerged as an effective tool for probing those properties of expressed genes that are not directly apparent from the mRNA sequence or transcript abundance, including the subcellular location of a protein of interest [4-6], the identification of post-translational modifications

[7,8], the characterization of interacting proteins or ligands [9], and the measurement of changes in these protein properties throughout the cell cycle or in response to a given stimuli or stress [10-12]. Coupled with various types of isotopic labeling reagents, MS can also be used to directly determine relative and absolute protein abundances [13]. Abundance measurements in proteomics are difficult, however, compared to mRNA studies as there are no amplification strategies such as PCR to increase the concentration of low abundance analytes.

In the present study, we attempt to characterize the Saccharomyces cerevisiae proteome using MS based proteomics. S. cerevisiae is a widely used and important model organism with a relatively large, but structurally simple, genome for which a high quality and well annotated sequence is available. It exhibits many of the same pathways and cellular functions as higher Eukaryotes. The largest published S. cerevisiae protein expression study used epitope tagging to detect 73% of the annotated Saccharomyces Genome Database (SGD) open reading frames (ORFs), which is 83% of SGD ORFs with Gene names [14,15]. Another recent study identified 72% of the predicted yeast proteome [16]. In this paper we combine the data from 47 different MS experiments that collectively generated 4.9 million spectra, into a single structure, the Saccharomyces cerevisiae PeptideAtlas.

The PeptideAtlas Project provides software tools and an infrastructure for the integration, visualization and analysis of multiple MS datasets [17-19]. This resource can be used to design future, more efficient experiments, to assist in the exploration of the proteome, and to support the development of proteomics software by making the data publicly accessible. We additionally demonstrate how this resource can be used in the construction of high quality lists of observable peptides for synthesis as reference molecules for targeted proteomics. This novel resource improves as more researchers contribute datasets.

Results and discussion PeptideAtlas construction

The S. cerevisiae PeptideAtlas is composed of 47 datasets (Table 1) from many different sources that were generated by using a variety of protocols and separation techniques. All samples in this atlas were proteolytically digested with trypsin, and many were treated with one of the isotope-coded affinity tagging (ICAT) reagents [10] or iodoacetamide. All samples were acquired using LC-ESI instruments (liquid chromatography separation, and electrospray ionization coupled with MS) - no matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) instrument datasets were available for inclusion. The PeptideAtlas can, however, accept and be expanded with data from any type of instrument using the mzXML data format (see below). A variety of protein or peptide separation techniques were employed in these experiments, including SDS-PAGE, free-flow electrophoresis and strong cation exchange chromatography, to generate fractions that were then subjected to reversed-phase HPLC separation prior to mass spectrometric analysis.

Processing of the acquired spectra and construction of a PeptideAtlas are briefly summarized in Figure 1 and described in more detail in previous publications [17-19]. For each submitted yeast dataset, all spectra were first converted to the mzXML format [20] irrespective of the original file format and then searched using SEQUEST [21] against a non-redundant S. cerevisiae reference protein database (the union of the SGD, Ensembl, NCI, and GenBank databases as detailed in Additional data file 1, plus keratin and trypsin). Redundant ORF sequences were coalesced to single entries with combined description fields. The union of the five protein sequence files yielded 13,748 distinct ORF sequences. Many ORF sequences differed by only a few amino acid residues, but all differences were retained in order to maximize the number of sequence assignments.

For each experiment the primary database search results were assigned statistical probabilities using the Peptide-Prophet program [22] implemented in the Trans-Proteomic-Pipeline [23]. Table 2 lists the number of spectra per experiment in total and with a PeptideProphet assignment $P \ge 0.9$, and the number of distinct peptide identifications cumulatively added by each experiment. The experiments are sorted approximately in the order submitted; the latter experiments will naturally make a smaller contribution to the total list of distinct peptides as many of the peptides identified in the latter experiments were also identified in earlier experiments.

Search results and spectra were stored in the Systems Biology Experiment Analysis Management System (SBEAMS), and all files were retained in an archive area.

To create the S. cerevisiae PeptideAtlas, all peptide information in contributed datasets was filtered to retain only those identifications with a PeptideProphet probability above 0.9. The remaining peptide sequences were then aligned with the SGD reference protein database using the NCBI program BLASTP [24] with arguments to achieve the highest scores for identities of 100% without gaps. Chromosomal coordinates were then calculated using the BLAST-provided coding sequence (CDS) coordinates of the peptide combined with the chromosomal coordinates of the protein features (where features are the contents of the SGD_features table that contain elements present in the GFF3 guidelines [25]). Peptide information along with the chromosomal coordinates were loaded into the PeptideAtlas database.

Summary statistics of the current yeast PeptideAtlas build are presented in Table 3. The build was constructed with a lower limit threshold of peptide assignments with $P \ge 0.9$, but the build may be searched with higher thresholds and those sum-

Genome Biology 2006,

Table I

List of experiments Experiment Instrument Treatment/labeling Separation Strain Data contributors Affiliation Reference LCQ Classic **ICAT** SCXBY4741 J Ranish, T Ideker ISB [42] gricat BY4742 cdc15_cdc23_newICAT LCQ DECA cllCAT SCX B Raught ISB BY4742 cdc15 cdc23 oldlCAT LCO DECA **ICAT** SCX B Raught ISB cdc15_cdc23_ICAT SCX BY4742 LCQ DECA cIICAT B Raught ISB cdc23_amf_newICAT LCQ DECA cllCAT SCX BY4742 B Raught ISB contFFE2Murea LCQ DECA cllCAT BY4741 ? F Kregenow, R Aebersold FFE BY4741 ? F Kregenow, R Aebersold **FFEY I** LCQ DECA Unknown ISB **FFEY I Scx** LCQ DECA FFE SCX BY4741 ? F Kregenow, R Aebersold ISB FFEY2 LCQ DECA XP FFE Unknown BY4741? F Kregenow, R Aebersold ISB Unknown PeteryeastlcatstdFFE LCQ DECA XP cllCAT BY4741 ? F Kregenow, R Aebersold ISB TSAAT000c LCQ DECA XP cllCAT SCX BY2125 M Flory et al. ISB [43] TSAAT030c LCQ DECA XP dICAT SCX BY2125 M Flory et al. ISB **[431** TSAAT060c LCQ DECA XP dICAT SCX BY2125 M Flory et al. ISB [43] TSAAT090c LCQ DECA XP cllCAT SCX BY2125 M Flory et al. ISB [43] TSAAT120c LCQ DECA XP cllCAT SCX BY2125 M Flory et al. ISB [43] TSAAxT000old LCQ DECA ICAT SCX BY2125 M Flory ISB [43] T00 LCQ DECA XP **ICAT** SCX BY2125 M Flory ISB [43] T30 LCQ DECA XP **ICAT** SCX BY2125 M Flory ISB [43] T50 LCQ DECA XP ICAT SCX BY2125 M Flory ISB **[431** opd00034_YEAST LCQ DECA XP None SCX DBY8724 P Lu University of Texas [44] opd00035_YEAST LCQ DECA XP None SCX DBY8724 P Lu University of Texas [44] peroximalPrep0702 LCQ Classic **ICAT** SCX BY4743 M Marelli et al. ISB [45] Comp12vs12sizefrac LCQ DECA Iodoacetemide SCX BY4741 DB Martin ISB LCQ DECA cllCAT SCX BY4743 M Marelli et al. ISB [45] pxproteome Comp12vs12standSCX LCQ DECA Iodoacetemide SCX BY4741 DB Martin ISB YeastICAT LCQ Classic **ICAT** SCXDerivative of BY4741 J Ranish ISB P_VMI LCQ Classic **ICAT** SCX BY4743 M Marelli et al. ISB [45] LCQ Classic cllCAT SCX BY4743 M Marelli et al. ISB peroximal clICAT [45] KR Serikawa et al. Ac30 LCQ DECA XP dICAT SCX BY1782, BY2125 University of [46] Washington LCQ DECA yeast Iodoacetemide SCX **YPH499** Gygi et al. Harvard Medical School [47] LCO DECA Gel. SCX BY4742 Ho et al. MDS Proteomics Iodoacetemide **[30]** gel msms SDS-PAGE, SCX mudpit LCQ DECA XP PLUS lodoacetemide **YRP480** P Haynes et al. University of [48] Arizona SDS-PAGE, IEF LCQ DECA XP PLUS P Haynes et al. ipg_ief lodoacetemide **YRP480** University of [48] Arizona P Haynes et al. LCQ DECA XP PLUS YRP480 rp_int_selected lodoacetemide SDS-PAGE University of [48] Arizona rp_mass_selected LCQ DECA XP PLUS SDS-PAGE YRP480 P Haynes et al. University of Iodoacetemide [48] Arizona sdspage LCQ DECA XP PLUS Iodoacetemide SDS-PAGE YRP480 P Haynes et al. University of [48] YM_NI4NI5_DAYGly DBY8724 LCQ Classic None Unknown P Lu University of Texas [44] YM NI4NI5 DAYSer DBY8724 LCO Classic None Unknown P Lu University of Texas [44] YM_NI4NI5_SCYGly LCO Classic None Unknown DBY8724 P Lu University of Texas [44] YM_NI4NI5_SCYSer LCQ Classic None Unknown DBY8724 P Lu University of Texas [44] APEX_04-22 LCQ Classic None Unknown **DBY8724** P Lu University of Texas [44] APEX 04-23 LCO Classic Unknown DBY8724 P Lu University of Texas None [44] DBY8724 P Lu APEX 04-24 LCQ Classic None Unknown University of Texas [44] LCQ Classic Unknown **DBY8724** University of Texas APEX 04-28 P Lu [44] YeastSCXReps LCQ Classic Iodacetimide SCX BY4741 Maynard et al. NIH [49] FFE_nonICAT LCQ Classic lodacetimide FFE BY2125? Mingliang Ye ISB

cllCAT is acid cleavable ICAT D0227/D9236 and ICAT is the older ICAT reagent D0422/D8450. FFE, free flow electrophoresis; IEF, is isoelectric focusing; SCX, strong cation exchange; SDS-PAGE, sodium dodecyl (lauryl) sulfate-polyacrylamide gel electrophoresis.

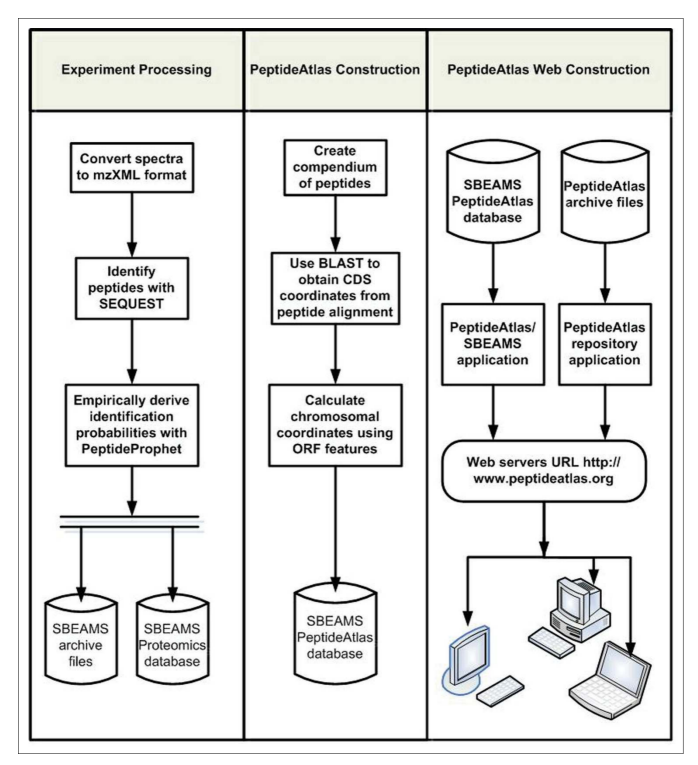


Figure 1
PeptideAtlas processing, creation, and interfaces. The first column outlines experiment level processing with SEQUEST [21] and PeptideProphet [22], the second column shows major stages in the construction of a PeptideAtlas [18] using BLAST [24] to obtain coding sequence (CDS) coordinates, and the third column shows the data, business logic, and presentation tiers for a PeptideAtlas.

mary statistics are also reported in Table 3. The number of distinct peptide sequences identified in these spectra (with P > 0.9) is 36,133. The number of SGD proteins with which

these peptides display perfect alignment is 4,063, which is 61% of all ORFs in the SGD protein database. If we apply a stricter criteria of removing identifications in which a peptide

Table 2

The number of spectra acquired and peptides identified per experiment

Experiment	numSpec($P > 0.0$)	numSpec(P > 0.9)	numDistinctPeptides(cumulative, P > 0.9)	
gricat	24,833	2,585		
cdc15_cdc23_newICAT	172,938	22,269	2,889	
cdc15_cdc23_oldICAT	156,353	8,281	3,184	
cdc15_cdc23_ICAT	21,392	3,337	3,293	
cdc23_amf_newICAT	127,673	25,861	4,096	
contFFE2Murea	105,686	5,473	4,430	
FFEYI	64,880	2,966	5,994	
FFEYIScx	7,045	796	6,396	
FFEY2	60,140	7,159	8,877	
PeteryeastlcatstdFFE	136,563	4,680	9,375	
TSAAT000c	217,919	43,628	11,423	
TSAAT030c	249,774	42,519	12,042	
TSAAT060c	214,851	28,089	12,424	
TSAAT090c	198,057	31,428	13,247	
TSAAT120c	179,408	26,182	13,588	
TSAAxT000old	101,166	6,311	13,728	
Т00	94,208	9,206	13,840	
T30	83,758	8,758	13,935	
T50	106,572	4,606	14,026	
opd00034_YEAST	24,049	4,554	14,557	
opd00035_YEAST	23,715	3,931	14,862	
peroximalPrep0702	24,209	3,197	15,236	
Comp 12vs 12sizefrac	28,926	11,703	17,553	
pxproteome	23,720	1,552	17,724	
Comp I 2 vs I 2 stand SCX	31,652	12,103	18,966	
YeastICAT	55,922	6,760	19,577	
PvMI	3,796	323	19,623	
peroximal_cllCAT	92,721	3,693	19,778	
Ac30	255,937	37,873	20,025	
yeast	140,567	29,411	24,548	
gel_msms	343,654	43,022	32,450	
mudpit	36,599	10,811	32,859	
ipg_ief	136,209	27,792	33,383	
rp_int_selected	53,358	14,669	33,684	
rp_mass_selected	71,151	4,881	33,857	
sdspage	92,802	19,208	34,734	
YM_N14N15_DAYGly	38,951	1,658	34,806	
YM_NI4NI5_DAYSer	39,315	1,862	34,843	
YM_NI4NI5_SCYGly	37,124	1,364	34,861	
YM_NI4NI5_SCYSer	37,658	2,238	34,902	
APEX_04-22	38,334	1,732	34,926	
APEX_04-23	38,175	1,775	34,949	
APEX_04-24	36,407	1,359	34,963	
APEX_04-28	40,612	2,248	34,995	
YeastSCXReps	118,083	60,568	35,446	
FFE_nonICAT	21,054	6,870	36,133	

Column 1 is the experiment name, column 2 is the number of spectra associated with PeptideProphet probabilities >0, column 3 is the number of spectra associated with PeptideProphet probabilities >0.9, and column 4 is the cumulative number of distinct peptides with PeptideProphet probabilities >0.9.

was only observed once in the entire *S. cerevisiae* PeptideAtlas, we then find that 43% of all SGD ORFs have been seen (with P > 0.9). If we also apply another criterion, that we count only the peptide to protein mappings that are not degenerate and that have $P \ge 0.9$, we observe 59% of SGD ORFs. The same criteria applied to peptides that have been seen more than once results in an observation of 41% of SGD ORFs. The number of peptides with perfect alignment to protein sequences in files other than SGD is 110 (Additional data file 1). Some of these identifications correspond to records that NCBI has discontinued or are identifications to appended contaminant sequences such as keratin or trypsin identified in the search, but are not present in the target genome (*S. cerevisiae* in this case).

Expected errors are calculated with equation 14 of the PeptideProphet paper using the summation of (1 - P_i) divided by N_i. This is applied to four cases, summarized as rows in Table 4, and three PeptideProphet probability limits, shown as columns in Table 4. The cases are: one, the assigned probability P of each MS/MS is used for all $P_i \ge P_{limit}$; two, the assigned probability P of each MS/MS is used for all $P_i \ge P_{limit}$ where the associated peptide has been seen in the S. cerevisiae Peptide-Atlas more than once; three, the best probability for each unique peptide sequence is used for all $P \ge P_{limit}$; and four, the best probability for each unique peptide sequence is used for all $P \ge P_{\text{limit}}$ when the peptide has been seen in the *S. cerevi*siae PeptideAtlas more than once. Note that cases three and four make an assumption that the peptide identifications can be represented by the best identification probability for that peptide. There are many methods to score groups of peptides, and we adopt the simplest in this paper for cases three and four as they represent the data as we have used it. For more detailed discussions on group scoring, please see [26,27]. Table 4 shows that the expected error rate for the S. cerevisiae PeptideAtlas as a whole is 9%. As an aside, one can construct

subsets of the build with smaller error rates by using a higher PeptideProphet probability threshold (see changes along a row in Table 4) or by reducing the number of peptides one is counting (see changes along a column in Table 4) by only using those peptides that have been observed more than once, and further removing information from multiple instances of those peptides.

In summary, the *S. cerevisiae* PeptideAtlas expected error rate is 9%, but the user is able to construct subset exports of the build with reduced error rates if desired.

To what extent do the peptides in the S. cerevisiae PeptideAtlas represent the S. cerevisiae proteome?

The coverage of the S. cerevisiae proteome by the PeptideAtlas is high, but not complete. Using only peptide identifications possessing a PeptideProphet score of P > 0.9, we have mapped to 61% of all SGD ORFs with at least one peptide hit. In Table 5 we present the observed ORFs categorized by feature type annotations. Of the 'uncharacterized' ORFs, 49% are represented in the S. cerevisiae PeptideAtlas (Additional data file 2). Uncharacterized ORFs are defined as putative gene products with homologs in another species that have, however, not been experimentally observed in S. cerevisiae. Of the 'verified' ORFs, 74% are represented in the PeptideAtlas. Verified ORFs are those that have been experimentally confirmed to exist in S. cerevisiae. A small percentage of ORFs are annotated as 'dubious'; only very few of these ORFs were found in the PeptideAtlas. Dubious ORFs are putative gene products that do not have homologs in other Saccharomyces species, and for which there is no experimental evidence of existence in S. cerevisiae. (Additional data file 3). Of the ORFs annotated as pseudogenes, 19% are represented in the PeptideAtlas. An SGD pseudogene has a functional homolog in another organism, and is predicted to no longer

Table 3
Statistics for the current S. cerevisiae PeptideAtlas

	$P_{limit} = 0.9, N_{obs} > 0$	$P_{limit} = 0.95, N_{obs} > 0$	$P_{limit} = 0.99, N_{obs} > 0$	$P_{limit} = 0.9, N_{obs} > 1$	$P_{limit} = 0.95, N_{obs} > 1$	$P_{limit} = 0.99, N_{obs} > 1$
# Experiments	47	47	47	47	47	47
# MS runs	2,579	2,579	2,579	2,579	2,579	2,579
# MS/MS	4.9 M	4.9 M	4.9 M	4.9 M	4.9 M	4.9 M
# MS/MS with $P \ge P_{limit}$	600,960	565,217	472,234	586,708	552,434	461,827
# Distinct peptides with $P \ge P_{limit}$	36,133	33,377	27,909	21,840	21,646	20,251
# Distinct peptides with perfect SGD alignment	35,434	32,790	27,499	21,469	21,281	19,942
# SGD ORFs seen in PeptideAtlas	4,249 (62%)	3,903 (57%)	3,476 (51%)	3,069 (45%)	3,049 (45%)	2,935 (43%)
# SGD ORFs unambiguously seen in PeptideAtlas	3,980 (59%)	3644 (54%)	3,224 (47%)	2,795 (41%)	2,778 (41%)	2,672 (39%)

The percentage of SGD ORFs seen in PeptideAtlas is shown as a function of lower limit PeptideProphet probabilities and number of times peptide has been observed above lower limit. Using the most generous parameters of the build, we see 62% of the SGD ORFs. As an aside, 68% of SGD ORFs have Systematic gene names and we observe 76% of those. This is comparable to the 83% of ORFs with Systematic gene names that Ghaemmaghami et al. [14] observed in their protein expression study.

Table 4

Expected errors

Case	Expected errors P _{limit} = 0.9	Expected errors P _{limit} = 0.95	Expected errors P _{limit} = 0.99
$MS/MS P_i \ge P_{limit}$	0.00915 (9%)	0.00517 (5%)	0.00137 (1%)
MS/MS $P_i \ge P_{limit}$, $N_{peptide observed} > 1$	0.00884 (9%)	0.00506 (5%)	0.00136 (1%)
Consensus peptide best P_i , $P_i \ge P_{limit}$	0.01027 (10%)	0.00510 (5%)	0.00120 (1%)
Consensus peptide best P_i , $P_i \ge P_{limit}$, $N_{peptide\ observed} > I$	0.00272 (3%)	0.00215 (2%)	0.00078 (1%)

Genome Biology 2006,

Expected errors in the Saccharomyces PeptideAtlas are calculated for four cases and lower probability limits ($P_{limit} = 0.9, 0.95, 0.99$): the assigned probability P of each MS/MS is used for all $P_i \ge P_{limit}$; the assigned probability P of each MS/MS is used for all $P_i \ge P_{limit}$; where the associated peptide has been seen in the S. cerevisiae PeptideAtlas more than once'; the best probability for each unique peptide sequence is used for all $P_i \ge P_{limit}$; the best probability for each unique peptide sequence is used for all $P_i \ge P_{limit}$ when the peptide has been seen in the S. cerevisiae PeptideAtlas more than once.

function because mutations prevent its transcription or translation. The pseudogene classification is based upon observations of ORFs from the S288C strain. Of the ORFs annotated as transposable elements, 20% are present in the PeptideAtlas (Additional data file 4). Note that the coverage of the ORFs in these categories decreases if we remove those peptides that have only been observed once in the entire atlas. With the single hit peptides removed, we see 56% of SGD verified ORFs in PeptideAtlas, 29% of the uncharacterized ORFs, none of the ORFs from verified/silenced_genes, 2% of the dubious ORFs, 5% of ORFs from pseudogenes, and 15% of ORFs from genes categorized as transposable element genes. The assignments to dubious ORFs should be viewed skeptically as the numbers of observations are extremely small and within error of the atlas.

A histogram of ORF sequence coverage is shown in Figure 2; the PeptideAtlas distribution is represented by the shaded distribution on the left, while an *in silico* digested SGD reference ORF distribution filtered to retain peptides with average molecular weights between 500 and 4,000 Da is seen on the right. About 40% of the ORFs in PeptideAtlas have sequence coverage greater than 20%. Importantly, the entire yeast proteome is not expected to be observable by current tandem MS techniques. This is not an impediment to protein identification, as the entire set of measurable peptides for a given pro-

tein is not necessary for an unambiguous identification of the protein. Some of the reasons that perfect sequence coverage is not possible are inherent in the instruments (discussed further in the next section) and in the search techniques. For example, we may miss identifications of sequences from post-translationally modified proteins in the search strategy applied.

Biases in the S. cerevisiae PeptideAtlas

Since all the data in the PeptideAtlas were acquired using LC-ESI-MS/MS, we examined peptide hydrophobicity, mass and charge distributions to characterize any inherent biases in the peptide dataset.

Using the Guo [28] and Krokhin *et al.* parameters [29] we created hydrophobicity histograms for the *S. cerevisiae* PeptideAtlas peptides (darker hatched bars), overlaid on an *in silico* digest of the entire SGD database (lighter hatched bars) allowing one missed tryptic cleavage (Figure 3). While peptides of moderate hydrophobicity were efficiently observed, the *S. cerevisiae* PeptideAtlas is clearly lacking in the most hydrophilic peptides - presumably because these peptides do not efficiently bind to standard HPLC columns and proceed to waste instead of entering the mass spectrometer. Other types of upstream separation techniques or modification of HPLC solvent conditions will most likely be required to improve the

Table 5

Numbers of proteins matching SGD ORF annotation categories

ORF annotation	SGD	PeptideAtlas P > 0.9, N _{obs} > 0	PeptideAtlas P > 0.9, N _{obs} > I	
Uncharacterized	1,414	695 (49%)	405 (29%)	
Verified	4,366	3,250 (74%)	2,449 (56%)	
Verified silenced gene	4	I (25%)	0 (0%)	
Dubious	820	95 (12%)	13 (2%)	
Pseudogene	21	4 (19%)	I (5%)	
Transposable element gene	89	18 (20%)	13 (15%)	
Total	6,714	4,063 (61%)	2,881 (43%)	

The percent of SGD ORFs seen in the PeptideAtlas are shown in columns 3 and 4. Column 3 uses all peptides with probabilities \geq 0.9, while column 4 excludes peptides that have only been seen once in the S. cerevisiae PeptideAtlas.

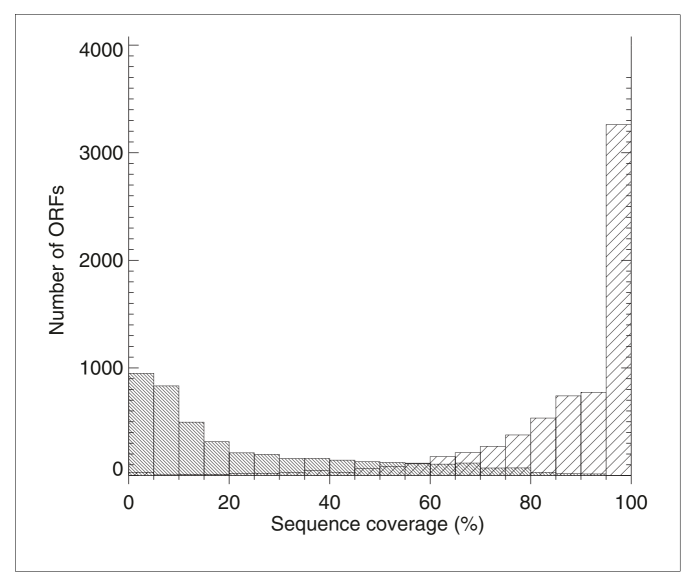


Figure 2
Histrograms of protein sequence coverage. A histogram of the sequence coverage of the S. cerevisiae ORFs by PeptideAtlas (darker filled bars) and an in silico tryptic digestion of the reference protein database with a mass range of 500 to 4,000 Da (lighter diagonal pattern filled bars) is shown. Of the PeptideAtlas ORFs, 61% have coverage below 20%, while 39% have a coverage above 20%.

detection of these hydrophilic peptides. Similarly, the most hydrophobic peptides are also not as efficiently observed as peptides with more moderate hydrophobicity scores, presumably because these peptides do not elute efficiently under standard LC gradient conditions.

A bias is also present in the distribution of peptide masses. Figure 4 shows histograms of *S. cerevisiae* PeptideAtlas average molecular weights (solid bars) overlaid on an *in silico* digest of the entire SGD database (hatched bars). The acquisition settings for MS/MS instruments are typically in the range of 400 to 2,000 m/z which, accounting for charge

states, limits the peptide mass detection range to 400 to 6,000 Da. The database searches, however, have a range of roughly 600 to 4,200 Da. The *in silico* digest reference distribution suggests that there would be a peak at a mass of approximately 700 amu, but the observed peak is near 1,500 amu. In the PeptideAtlas, we appear to be missing many of the peptides with masses less than 1,400 Da, largely because these smaller peptides are more difficult to identify using standard database search tools. Importantly, however, smaller peptides are often not as useful in protein identification as the longer peptides, since the short amino acid sequences are less likely to be unique to a single protein.

Genome Biology 2006,

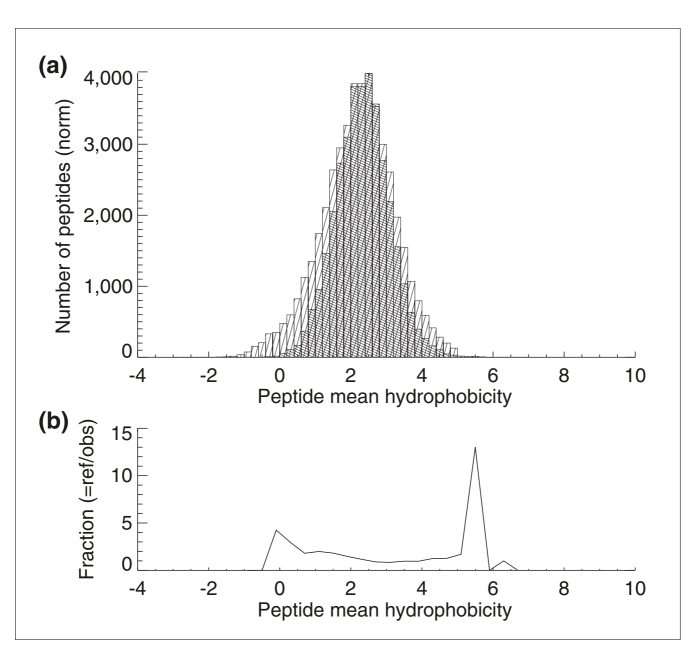


Figure 3 Hydrophobicity histrogams. (a) Mean hydrophobicity histogram for peptides in the S. cerevisiae PeptideAtlas (darker hashed bars) and an in silico tryptic digest of the SGD reference protein database allowing one missed cleavage (lighter hashed bars). (b) The reference peptides' hydrophobicities divided by the observed peptides' hydrophobicities. The lowest hydrophobicity peptides are generally washed off the column in the reverse phase stage of the HPLC process and hence not measured.

Additionally, larger peptides tend to have more than one chance of being observed, as charge states of +2 or +3 can put the peptides within range of the acquisition settings of the instrument. Figure 4 shows that there are a larger number of peptides in charge states of +2 and +3, with only a small percentage of peptide identifications derived from a +1 charge state. This is expected given that the datasets in this version of the PeptideAtlas are from ESI instruments. We do not currently search the spectra for ions with charge states larger than +3 and it could be expected that many of the missed

larger peptides might generate higher charge state ions. The addition of MALDI-TOF datasets to the atlas will populate the database with identifications from ions in the +1 charge state. Krogan et al. [16] find high protein discovery rates using MALDI, so this approach is promising.

In summary, a sizable fraction of the yeast proteome has been identified using LC-ESI MS/MS. The smallest peptides are not well represented in the PeptideAtlas. Additionally, the most hydrophilic peptides and hydrophobic peptides are

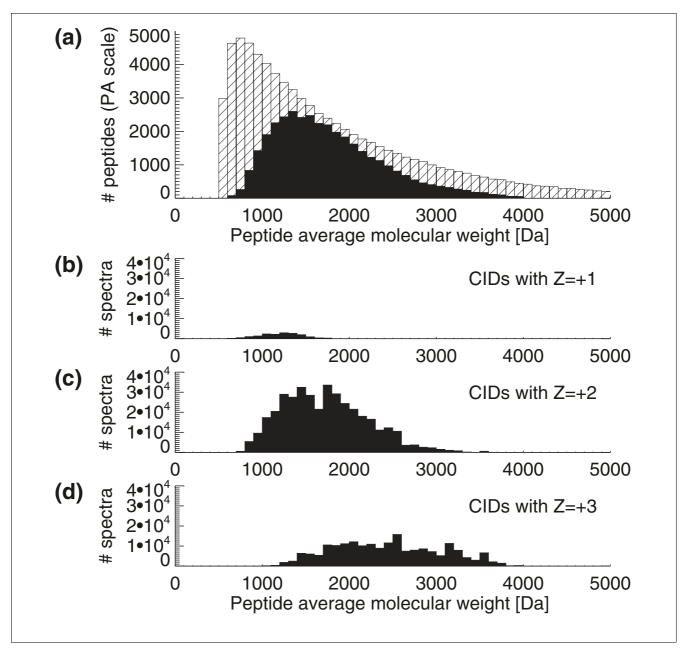


Figure 4 Mass histograms. (a) Average molecular weight of unique peptide sequences in PeptideAtlas (solid filled bars) and the in silico tryptically digested SGD protein database (hashed bars) allowing one missed cleavage. (b-d) Mass histograms of spectra, separated by charge. The large number of peptides with masses less than 1,000 Da are difficult to identify in database searches, and hence are not present in the PeptideAtlas. CID, collision-induced dissociation.

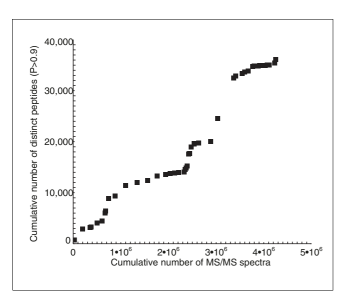
under represented. It will be interesting to determine how these distributions change as more diverse types of data are added to the atlas.

The relationship between number of spectra and number of identified peptides

As the PeptideAtlas is continually populated with new datasets, we expect that, at some stage, the addition of new spectra will produce few new peptide identifications. We are thus tracking the number of unique peptides contributed by each

additional experiment. From a total of 4.9 million MS/MS we have 36,133 distinct peptides spectra, PeptideProphet scores >0.9 (Table 3). As an aside, the number of distinct peptides from an in silico tryptic digestion of the SGD protein database, allowing one missed cleavage, and counting those peptides whose masses are in the range 500 to 4,000 Da, is 436,445, so we currently observe roughly 10% of what we might expect if all peptides had equal possibility of being observed. The current rate of inclusion of unique peptide identifications with P > 0.9 is shown in Figure

King et al. R106.11



http://genomebiology.com/2006/7/11/R106

Figure 5 Cumulative number of MS/MS spectra versus the number of unique peptide identifications with P > 0.9. The slope is nearly horizontal in regions of the curve where similar experiments were performed. The curve is expected to show saturation when additional spectra provide no new peptides above P > 0.9. The number of distinct peptides from an in silico trypic digestion of the SGD protein database, allowing one missed cleavage, and counting those peptides whose masses are in the range 500 to 4,000 Da, is 436,445.

5. In general, there is one peptide added for every 125 spectra. Flattened areas of the curve are due to overlapping identification of peptides from similar experiments (and instrument), rather than the expected final trend of saturation of the proteome sequences. Remarkable increases in distinct peptide yields are seen from the Gygi et al. [10] yeast dataset and the Ho et al. [30] gel_msms dataset. In summary, we have identified roughly ten percent of the peptides predicted from an in silico digested protein database, and have not yet reached saturation of novel additions. Novel additions are expected with the inclusion of results from new experiment designs and instrument platforms.

What new information does the PeptideAtlas contribute about the S. cerevisiae proteome and genome?

Having characterized some observational biases in peptide content of the PeptideAtlas, we now briefly examine characteristics of the identifications in relation to predictions based on codon bias and gene ontology categories. Figure 6 illustrates the codon enrichment correlation (CEC) for all ORFs in the S. cerevisiae PeptideAtlas and all ORFs in Ghaemmaghami et al. [14]. CEC is a parameter constructed by Ghaemmaghami et al. [14] to represent a measurement of the deviation of observed protein codon-usage from codon-usage in a randomly generated ORF. These distributions are both skewed toward high positive values, typically greater than

0.25, signifying that their sequences deviate significantly from that of ORFs derived from random codons; these are thus likely to be true ORFs and not spurious predictions. Figure 6b is a histogram of CEC for ORFs not observed in the Ghaemmaghami et al. expression sets, and for all ORFs not present in PeptideAtlas. Their distributions are evenly distributed around the origin, suggesting that both datasets are missing bona fide ORFs as well as ORFs that are not likely to code for proteins.

The proteins identified in the yeast PeptideAtlas are generally evenly distributed with respect to Gene Ontology (GO) molecular function categories (Figure 7). Interestingly, we observe 52% of yeast ORFs annotated as 'molecular function unknown'. If we filter out peptides that have been observed only once in the PeptideAtlas, the percent of 'molecular function unknown' genes we see is 32% (approximately 659 genes). The same trend is seen in the GO 'cellular components' and the GO 'biological processes' categories. Figures 8 and 9 show that a significant fraction of ORFs in PeptideAtlas are present in the catgories 'cellular component unknown' and 'biological process unknown'.

The S. cerevisiae PeptideAtlas, by verifying that these unannotated ORFs produce identifiable proteins, could stimulate interest in determining their function.

PeptideAtlas user interface

Genome Biology 2006,

Summary statistics and query interfaces for the PeptideAtlas are available at [31]. Perl cgi-bin programs and Java servlets and JSP pages enable public queries of the database and public archive area flat files. From the PeptideAtlas front page [31], atlas data may be accessed by links on the navigation bar. The 'Data Repository' link leads to a form where the user can retrieve all publicly available datasets and search results. The 'Search Database' link leads to several pages where the user can either search the atlas for keywords, find the information summary for a given peptide in the atlas, or produce a tailored list of peptides from the atlas by specifying a variety of constraints.

The S. cerevisiae PeptideAtlas user interface thus allows for an extremely large number of diverse tandem MS datasets to be searched, processed, and combined in a user-specifiable fashion.

Uses of the S. cerevisiae PeptideAtlas

The S. cerevisiae PeptideAtlas represents an extremely useful data mining tool. Here, we present two examples of how the database may be used. The first example demonstrates the creation of a list for construction of synthetic peptides, and the second example demonstrates validation of predicted SGD introns.

Using quantitative MS techniques, synthetic peptides may be utilized to determine subunit stoichiometry within a given

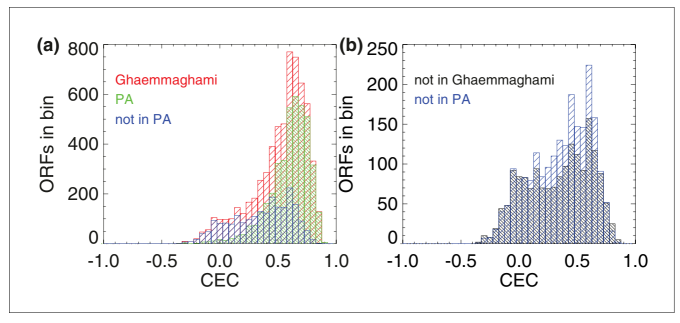


Figure 6
Comparison of codon enrichment correlation (CEC) distributions. (a) Histogram of CEC for all ORFs listed in Ghaemmaghami et al. [14], for all ORFs seen in PeptideAtlas (PA), and for all ORFs not seen in PeptideAtlas; and (b) histogram of CEC for ORFs not seen in the Ghaemmaghami et al. expression sets and for all ORFs not seen in PeptideAtlas. Observed proteins in PeptideAtlas and in Ghaemmaghami et al. show a positive skew in CEC showing that they deviate significantly from that expected from a random sequence of codons, while the unobserved proteins show more uniformly distributed CEC values.

multi-protein complex [32] or to determine the absolute quantity of a protein in a sample [33,34]. To design such

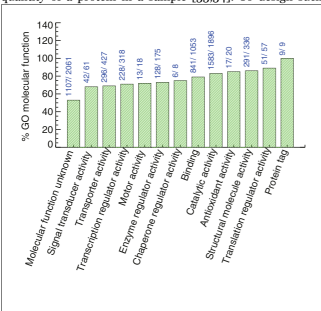


Figure 7
The number of genes identified in GO Molecular Function terms (more specifically, the first level children of molecular function). The bars are annotated with the number of SGD genes annotated in that term and the number of SGD genes seen in PeptideAtlas for that term. Many annotated as unknown are present in the PeptideAtlas.

peptides, one could use the S. cerevisiae PeptideAtlas to: identify those peptides of member proteins that have been observed most often in the mass spectrometer; identify those that are specific for a single protein; and determine which of these peptides contain amino acids necessary for a given type of labeling reagent and suitable for peptide synthesis. For example, if the stoichiometry of the general transcription factor TFIIF complex was to be determined, the user would go to the PeptideAtlas URL [31], enter %tfIIf% into the search box and select 'GO'. The results are links to PeptideAtlas pages corresponding to the three SGD ORFs that are part of the TFIIF transcription factor complex in this organism [35,36]. One may either follow the links to the individual pages for the three ORFs, or open a new 'Browse Peptides' page using the tab located near the top of the page to query the atlas in more detail. In the new 'Browse Peptides' page, the user can enter the three SGD ORFs into the 'Protein Name Constraint' text box (separated by semi-colons: YPL129W; YGR186W; YGRoo5C). Further useful constraints to enter are '>0.9' for 'Best Probability Constraint', '>1' for 'Number of Observations Constraint', and ' = 1' for 'Number of Proteins Mapped Constraint'. Following selection of 'Query' near the bottom of the page, results are returned below the form. To tailor the list to ICAT experiments, one could also enter %C% in the 'Peptide Sequence Constraint' text box above and select 'Query' again. The returned list consists of five cysteine-containing peptides whose sequences have been observed more than once with high confidence, and which are present in only one

Genome Biology 2006,

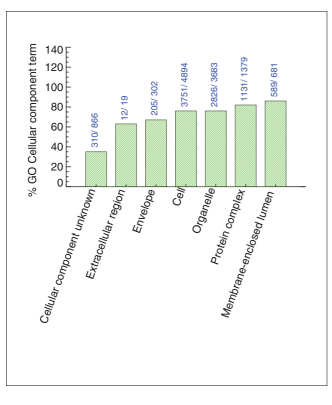


Figure 8
The number of genes identified in GO Cellular Component terms (more specifically, the first level children). The bars are annotated with the number of SGD genes and the number of SGD genes in PeptideAtlas for that term. Many annotated as unknown are present in the PeptideAtlas.

protein. The resulting list may be exported to the user's desktop in Excel, xml, tsv, or csv file formats.

As an example of a bioinformatics use of the S. cerevisiae PeptideAtlas, one could search for all experimentally observed splice forms predicted in the SGD database that are present in the S. cerevisiae PeptideAtlas. In this case, the user would select 'Search Database' from the navigation bar on the left, select the latest Saccharomyces atlas from the 'Atlas Build' list, then in the text box 'Does Mapping Span Exons Constraint' (which is to indicate that the peptide sequence is aligned to the exons on both sides of an intron) enter 'y', and select 'QUERY'. Rows of data are returned for the latest yeast build. That list can be saved and compared against a list of SGD introns to find that the atlas contains 13% of the 367 SGD yeast ORF introns. Two of these intron validations are for uncharacterized (experimentally unobserved) YPR063C and YBL059C-A. To visualize the resulting information in the Ensembl Genome browser, the user may click on the coordinates in the results table (instructions on enabling the visualization are present on the navigation bar of the PeptideAtlas web pages).

Reverse look-ups of PeptideAtlas peptides are also possible from the Ensembl Genome browser website. For example,

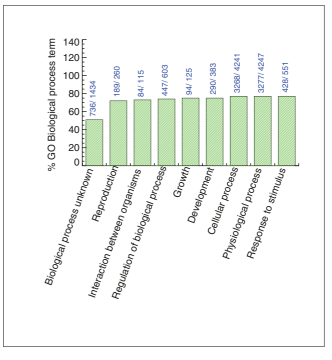


Figure 9The number of genes identified in GO Biological Process terms (more specifically, the first level children). The bars are annotated with the number of SGD genes and the number of SGD genes in PeptideAtlas for that term.

while browsing the genome view centered around the uncharacterized ORF YNL010W at [37] one can select the *S. cerevisiae* PeptideAtlas as a distributed annotation system (DAS) source, resulting in a large number of PeptideAtlas entries confirming portions of this predicted ORF sequence. Selecting any of these links can bring the user back to the PeptideAtlas summary pages for more detail.

Conclusion

We have demonstrated that MS/MS spectra from a large number of diverse sources can be uniformly processed and combined to create a large dataset useful for exploring and validating the yeast proteome. The public interfaces allow creation of high quality peptide lists that can be used to synthesize reference molecules for targeted proteomics. The datasets are also useful to examine MS based proteomics in general, can be used for software development, and can be used by other researchers to validate hypothetical proteins for examples. The *S. cerevisiae* PeptideAtlas possesses the highest degree of proteome coverage for any eukaryotic organism to date in a single public database offering entire datasets as validation, and, as such, is a growing resource that will continue to improve as more researchers contribute data.

Materials and methods Experiment processing

Some of the process level details of the experiment processing are provided here. The mzXML [20] conversions from vendor format files to mzXML files were performed using software available at our Sashimi software site [38]. The SEQUEST parameters used in MS/MS assignment were for semi-tryptic digestion and one static modification of methionine due to oxidation. Additional parameters were required for some datasets, such as those labeled with ICAT or samples treated with iodoacetemide. All sequest params files can be found in the searched archive files at our repository [39] for the public datasets. (Datasets that researchers have requested to be kept as private until they have published are not present in the data repository, but are included in the PeptideAtlas database with minimum sample annotation.)

The PeptideProphet software is available at our Sashimi software site [23]. The SBEAMS database application and the Proteomics module and PeptideAtlas modules within it are available at our SBEAMS software site [40] as downloads from a subversion code repository. The code is browsable from the worldwide web [41]. The BLAST algorithm was used with the parameters for 100% identity matches to a small peptide in a protein reference database:

blastp -F F -W 2 -M PAM30 -G 9 -E 1 -e 10 -K 50 -b 50

Additional data files

The following additional data are available with the online version of this paper. Additional data file 1 is a comma-separated text file containing a table of S. cerevisiae ORF fasta files used to create the reference database. The table includes the ftp addresses of the datasets and the number of ORFs unique to each dataset. Additional data file 2 is a comma-separated text file containing a table of the uncharacterized ORFs seen in S. cerevisiae PeptideAtlas. Included are the number of times their peptides were observed and the best Peptide-Prophet probability associated with their peptides. Additional data file 3 is a comma-separated text file containing a table of the dubious ORFs seen in S. cerevisiae PeptideAtlas. Included are the number of times their peptides were observed, and the best PeptideProphet probability associated with their peptides. Additional data file 4 is a comma-separated text file containing a table of the ORFs from transposable element genes seen in S. cerevisiae PeptideAtlas, the number of times their peptides were observed, and the best PeptideProphet probability associated with their peptides.

Acknowledgements

The research reported in this article was supported in part by contract No. No1-HV-28179 from the National Heart, Lung, and Blood Institute. We thank Olga Vitek and Julian Watts for their advice and consultation. We thank Steve Stein (NIST) and are grateful to all of the researchers who have made their datasets publicly available, specifically, Marcello Marelli and collaborators (ISB), Peng Lu (OPD, University of Texas), P Haynes and collaborators

orators (University of Arizona), KR Serikawa and collaborators (University of Washington), S Gygi and collaborators (Harvard Medical School), Ho and collaborators (MDS Proteomics, Samuel Lunenfeld Research Institute, University of Toronto, Kings College Circle), and Trey Ideker (ISB).

References

- Sonenberg N, Hershey WB, Mathews MB: Translational Control of Gene Expression (Cold Spring Harbor Monograph) Cold Spring Harbor: Cold Spring Harbor Laboratory Press; 2001.
- Hershko A, Ciechanover A: The ubiquitin system. Annu Rev Biochem 1998, 67:425-479.
- Pickart CM: Mechanisms underlying ubiquitination. Annu Rev Biochem 2001, 70:503-533.
- Dreger M: Subcellular proteomics. Mass Spectrom Rev 2003, 22:27-56.
- Washburn MP, Wolters D, Yates JR 3rd: Large-scale analysis of the yeast proteome by multidimensional protein identification technology. Nat Biotechnol 2001, 19:242-247.
- Gavin AC, Aloy P, Grandi P, Krause R, Boesche M, Marzioch M, Rau C, Jensen LJ, Bastuck S, Dumpelfeld B, et al.: Proteome survey reveals modularity of the yeast cell machinery. Nature 2006, 440:631-636.
- Gruhler A, Olsen JV, Mohammed S, Mortensen P, Faergeman NJ, Mann M, Jensen ON: Quantitative phosphoproteomics applied to the yeast pheromone signaling pathway. Mol Cell Proteomics 2005, 4:310-327.
- 8. Jensen ON: Modification-specific proteomics: characterization of post-translational modifications by mass spectrometry. Curr Opin Chem Biol 2004, 8:33-41.
- Gingras AC, Aebersold R, Raught B: Advances in protein complex analysis using mass spectrometry. J Physiol 2005, 563:11-21.
- Gygi SP, Rist B, Gerber SA, Turecek F, Gelb MH, Aebersold R: Quantitative analysis of complex protein mixtures using isotope-coded affinity tags. Nat Biotechnol 1999, 17:994-999.
- Blagoev B, Kratchmarova I, Ong SE, Nielsen M, Foster LJ, Mann M: A proteomics strategy to elucidate functional protein-protein interactions applied to EGF signaling. Nat Biotechnol 2003, 21:315-318.
- Brand M, Ranish JA, Kummer NT, Hamilton J, Igarashi K, Francastel C, Chi TH, Crabtree GR, Aebersold R, Groudine M: Dynamic changes in transcription factor complexes during erythroid differentiation revealed by quantitative proteomics. Nat Struct Mol Biol 2004, 11:73-80.
- 13. Von Haller PD, Yi E, Donohoe S, Vaughn K, Keller A, Nesvizhskii Al, Eng J, Li XJ, Goodlett DR, Aebersold R, Watts JD: The application of new software tools to quantitative protein profiling via isotope-coded affinity tag (ICAT) and tandem mass spectrometry: I. Statistically annotated datasets for peptide sequences and proteins identified via the application of ICAT and tandem mass spectrometry to proteins copurifying with T cell lipid rafts. Mol Cell Proteomics 2003, 2:426-427.
- Ghaemmaghami S, Huh WK, Bower K, Howson RW, Belle A, Dephoure N, O'Shea EK, Weissman JS: Global analysis of protein expression in yeast. Nature 2003, 425:737-741.
- Huh WK, Falvo JV, Gerke LC, Carroll AS, Howson RW, Weissman JS, O'Shea EK: Global analysis of protein localization in budding yeast. Nature 2003, 425:686-691.
- Krogan NJ, Cagney G, Yu H, Zhong G, Guo X, Ignatchenko A, Li J, Pu S, Datta N, Tikuisis AP, et al.: Global landscape of protein complexes in the yeast Saccharomyces cerevisiae. Nature 2006, 440:637-643
- Desiere F, Deutsch EW, King NL, Nesvizhskii AI, Mallick P, Eng J, Chen S, Eddes J, Loevenich SN, Aebersold R: The PeptideAtlas project. Nucleic Acids Res 2006, 34:D655-658.
- 18. Desiere F, Deutsch EW, Nesvizhskii Al, Mallick P, King NL, Eng JK, Aderem A, Boyle R, Brunner E, Donohoe S, et al.: Integration with the human genome of peptide sequences obtained by high-throughput mass spectrometry. Genome Biol 2005, 6:R9.
- Deutsch EW, Eng JK, Zhang H, King NL, Nesvizhskii Al, Lin B, Lee H, Yi EC, Ossola R, Aebersold R: Human plasma PeptideAtlas. Proteomics 2005, 5:3497-3500.
- Pedrioli PG, Eng JK, Hubley R, Vogelzang M, Deutsch EW, Raught B, Pratt B, Nilsson E, Angeletti RH, Apweiler R, et al.: A common open representation of mass spectrometry data and its

- application to proteomics research. Nat Biotechnol 2004, 22:1459-1466.
- 21. Eng JK, McCormack AL, Yates JR 3rd: An approach to correlate tandem mass spectral data of peptides with amino acid sequences in a protein database. | Am Soc Mass Spectrom 1994, **5:**976-989.
- 22. Keller A, Nesvizhskii Al, Kolker E, Aebersold R: Empirical statistical model to estimate the accuracy of peptide identifications made by MS/MS and database search. Anal Chem 2002, **74:**5383-5392
- 23. Keller A, Eng J, Zhang N, Li XJ, Aebersold R: A uniform proteomics MS/MS analysis platform utilizing open XML file formats. Mol Syst Biol 2005, 1:2005.0017
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ: Basic local alignment search tool. J Mol Biol 1990, 215:403-410.
- Generic Feature Format Version 3. [http://song.source forge.net/gff3.shtml]
- Nesvizhskii Al, Aebersold R: Interpretation of shotgun proteomic data: the protein inference problem. Proteomics 2005, 4:1419-1440.
- 27. Nesvizhskii Al, Keller A, Kolker E, Aebersold R: A statistical model for identifying proteins by tandem mass spectrometry. Anal Chem 2003, 75:4646-4658.
- Guo D: Prediction of peptide retention times in reversedphase high-performance liquid chromatography I. Determination of retention coefficients of amino acid residues of model synthetic peptides. J Chromatography A 1986, 359:499-518.
- 29. Krokhin OV, Craig R, Spicer V, Ens W, Standing KG, Beavis RC, Wilkins JA: An improved model for prediction of retention times of tryptic peptides in ion pair reversed-phase HPLC: its application to protein peptide mapping by off-line HPLC-MALDI MS. Mol Cell Proteomics 2004, 3:908-919
- 30. Ho Y, Gruhler A, Heilbut A, Bader GD, Moore L, Adams SL, Millar A, Taylor P, Bennett K, Boutilier K, et al.: Systematic identification of protein complexes in Saccharomyces cerevisiae by mass spectrometry. Nature 2002, 415:180-183.
- PeptideAtlas. [http://www.peptideatlas.org]
 Ranish JA, Yi EC, Leslie DM, Purvine SO, Goodlett DR, Eng J, Aebersold R: The study of macromolecular complexes by quantitative proteomics. Nat Genet 2003, 33:349-355
- Gerber SA, Rush J, Stemman O, Kirschner MW, Gygi SP: Absolute quantification of proteins and phosphoproteins from cell Proc Natl Acad Sci USA 2003, lysates by tandem MS. 100:6940-6945.
- 34. Lu Y, Bottari P, Turecek F, Aebersold R, Gelb MH: Absolute quantification of specific proteins in complex mixtures using visible isotope-coded affinity tags. Anal Chem 2004, 76:4104-4111.
- 35. Myer VE, Young RA: RNA polymerase II holoenzymes and subcomplexes. J Biol Chem 1998, 273:27757-27760.
- Rani PG, Ranish JA, Hahn S: RNA polymerase II (Pol II)-TFIIF and Pol II-mediator complexes: the major stable Pol II complexes and their activity in transcription initiation and reinitiation. Mol Cell Biol 2004, 24:1709-1720.
- 37. Ensembl: S. cerevisiae Contig View. [http://www.ensembl.org/ Saccharomyces_cerevisiae/contigview?c=XIV:6 | 4000.5;w=100]
- Sashimi. [http://sashimi.sourceforge.net]
- 39. PeptideAtlas Data Repository. [http://www.peptideatlas.org/ repository
 - Systems Biology Experiment Analysis Management System. [http://www.sbeams.org]
- SBEAMS CVS Trunk. [http://www.sbeams.org/cgi/viewcvs/ viewcvs.cgi/sbeams/trunk/]
- 42. Ideker T, Thorsson V, Ranish JA, Christmas R, Buhler J, Eng JK, Bumgarner R, Goodlett DR, Aebersold R, Hood L: Integrated genomic and proteomic analyses of a systematically perturbed metabolic network. Science 2001, 292:929-934.
- 43. MacKay VL, Li X, Flory MR, Turcott E, Law GL, Serikawa KA, Xu XL, Lee H, Goodlett DR, Aebersold R, et al.: Gene expression analyzed by high-resolution state array analysis and quantitative proteomics: response of yeast to mating pheromone. Mol Cell Proteomics 2004, 3:478-489.
- 44. Prince JT, Carlson MW, Wang R, Lu P, Marcotte EM: The need for a public proteomics repository. Nat Biotechnol 2004, 22:471-472.
- Marelli M, Smith JJ, Jung S, Yi E, Nesvizhskii Al, Christmas RH, Saleem RA, Tam YY, Fagarasanu A, Goodlett DR, et al.: Quantitative mass spectrometry reveals a role for the GTPase Rholp in actin organization on the peroxisome membrane. J Cell Biol 2004, 167:1099-1112.

- 46. Serikawa KA, Xu XL, MacKay VL, Law GL, Zong Q, Zhao LP, Bumgarner R, Morris DR: The transcriptome and its translation during recovery from cell cycle arrest in Saccharomyces cerevisiae. Mol Cell Proteomics 2003, 2:191-204.
- 47. Peng J, Elias JE, Thoreen CC, Licklider LJ, Gygi SP: Evaluation of multidimensional chromatography coupled with tandem mass spectrometry (LC/LC-MS/MS) for large-scale protein analysis: the yeast proteome. J Proteome Res 2003, 2:43-50.
- Breci L, Hattrup E, Keeler M, Letarte J, Johnson R, Haynes PA: Comprehensive proteomics in yeast using chromatographic fractionation, phase fractionation, gas protein electrophoresis, and isoelectric focusing. Proteomics 2005, **5:**2018-2028.
- Maynard DM, Masuda J, Yang X, Kowalak JA, Markey SP: Characterizing complex peptide mixtures using a multi-dimensional liquid chromatography-mass spectrometry system: Saccharomyces cerevisiae as a model system. J Chromatogr B Analyt Technol Biomed Life Sci 2004, 810:69-76.