

Counting Strategies for Differential Characterization Of Post-Translational Modifications

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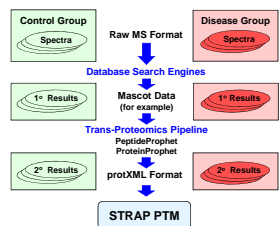
Introduction

There is increasing interest in the identification and characterization of protein post-translational modifications (PTMs) as the field of mass spectrometry-based proteomics begins to mature. However, the vast amount of information obtained within a typical differential proteomics study makes the facile measurement of PTMs quite challenging. In addition to label-free approaches, we have begun to explore counting methods for differential analysis of PTM changes in proteomes. Our novel approach has resulted in the development of a software program, the **Software Tool for Rapid Annotation of Proteins: Post-Translation Modification edition (STRAP PTM)**. STRAP PTM uses a new counting-based PTM scoring algorithm to facilitate multi-sample PTM comparison through collation and visualization. Here we demonstrate the utility of STRAP PTM across different PTM/proteomics experiments.

Methods

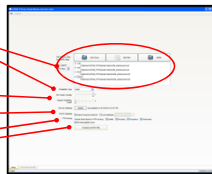
STRAP PTM: PTM Counting

Workflow

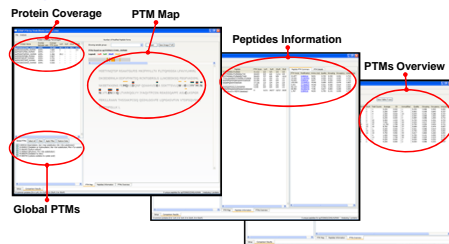


Setup Window

1. Enter proXML files per group.
2. Select ProteinProphet probability type.
3. Choose minimum protein overlap and peptide probability cutoff.
4. Indicate FASTA database.
5. Check factors for PTM scoring.
6. Execute comparison.



Comparison of Results Windows



Methods

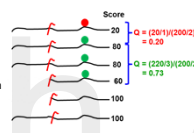
STRAP PTM: PTM Scoring

Algorithm	Expression
PTM Score	$S_{mip} = 100 \frac{Q_{mip} \gamma_{mip} W_{mip} U_{mip}}{(\max S)_p}$

- Based on four factors (Q = quality, γ = grouping, W = occupancy, and U = uniqueness) and normalized by the highest score in the data set (max PTM score = 100).
- Calculated for a specific PTM ($m = 1, \dots, M$) on a specific site ($i = 1, \dots, I$) of a specific protein (p) across all groups or data sets.

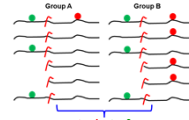
Factor Name	Expression
Quality	$Q_{mip} = \frac{(B_{mip})}{(B^0_{ip})}$

- Defined as the ratio of (the average score of peptides having a specific PTM on a specific site) to (the average score of all unmodified peptides having the specific site).
- Determines the quality of the database search results based on MS/MS ID scores (B).
- Higher score indicates positive ID (max = 1).



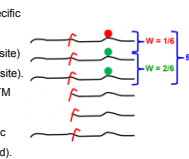
Factor Name	Expression
Grouping	$\gamma_{mip} = \frac{\sigma_{mip}}{(\max \sigma)_p}$

- Defined as the standard deviation (σ) of a specific PTM on a specific site across groups and normalized by the highest σ for the specific protein.
- Higher score indicates group specificity for the specific PTM and site (max = 1).



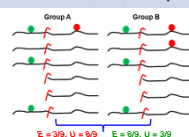
Factor Name	Expression
Occupancy	$W_{mip} = \frac{N_{mip}}{(\sum_{m=1}^M N_{mip}) + (N^0_{ip})}$

- Defined as the degree of modification of a specific site with a specific PTM on a specific protein.
- W is the ratio of (a specific PTM on a specific site) to (the sum of all PTMs and vacancies on the site).
- Higher score indicates more of the specific PTM at the site (max = 1).
- Total site occupancy equals the sum of specific PTM occupancies (susceptibility to be modified).



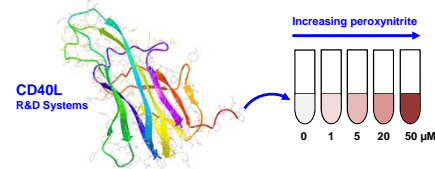
Factor Name	Expression
Uniqueness	$U_{mp} = 1 - E_{mp} = 1 - \frac{\sum_{i=1}^I N_{mip}}{\sum_{m=1}^M \sum_{i=1}^I N_{mip}}$

- Defined as the enhancement (E) of a specific PTM on a specific protein subtracted from 1.
- E is the ratio of (a specific PTM on all sites) to (all PTMs on all sites).
- Higher score indicates more specificity in the PTM (max = 1).



Results

Model System: In Vitro Oxidation of CD40L



PTM Map

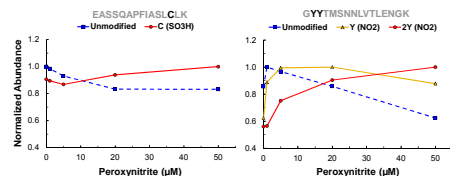
Legend: 1 μM 5 μM 20 μM 50 μM

MIETYNQTSF RAAATGLPIS MKIFMYLLTV FLITQMIGSA LFAVYLHRRLL
DKIEDERNLH EDFVFMKIQ RCTNGERSLS LLNCEEKISQ FEGFVKDML
NKEETKENS FEMQKGDQNP QIAAHVISEA SKSTTSVLGV AEGKYITMSN
NLVTLENGKQ LTVKRGLLY IYAQVTFCSN REASSQAPFI ASLCLSPGR
FERILLRAAN THSSAKPCGQ QSIHLGVFV LQPSAGVFN YLDFSQVSHG
TGFTSFLGL L
(Y145, Y146) NO2
(C194) SO3H

PTM Scoring

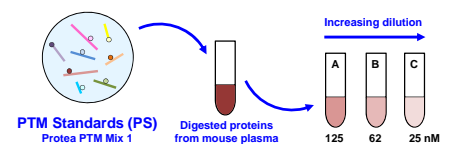
Protein	AA Index	Mod	Site	Total Counts	Average Counts	SD	Unmodified Forms	Quality	Grouping	Occupancy	Uniqueness	PTM Score				
146	M	15	15402	3	7	10	15	35	8.750	5.058	37	0.868	0.872	0.479	0.807	100.000
146	Y	14	15402	9	1	11	18	22	5.000	5.802	91	0.205	1.000	0.301	0.890	100.000
146	F	14	15402	8	1	8	18	19	4.750	4.902	54	0.817	0.990	0.280	0.890	100.000
113	M	15	15402	0	0	0	1	1	0.250	0.300	0	0.984	0.988	1.000	0.807	29.044
117	D	21	18134	1	3	2	2	9	2.250	2.907	5	0.802	0.185	0.643	0.727	28.286
114	Q	18	18421	0	0	1	1	1	0.250	0.300	0	0.984	0.988	1.000	0.715	25.728
151	N	18	18421	7	7	5	3	22	5.500	1.915	51	0.725	0.330	0.301	0.715	21.877
350	Q	18	18421	4	1	5	3	13	3.250	1.758	45	0.865	0.294	0.224	0.715	17.728
198	E	21	18134	7	6	9	7	29	7.250	1.258	44	0.539	0.217	0.387	0.727	14.325
159	C	21	18134	6	2	1	1	4	1.000	0.616	10	0.644	0.141	0.288	0.727	7.240
150	N	18	18421	0	2	1	1	4	1.000	0.616	69	0.755	0.141	0.055	0.715	1.787
140	W	15	15402	0	0	0	1	1	0.250	0.300	2	0.920	0.988	0.333	0.807	0.229
142	E	21	18134	0	0	0	1	1	0.250	0.300	2	0.920	0.988	0.333	0.727	0.264
145	M	15	15402	0	0	0	1	1	0.250	0.300	37	0.901	0.988	0.074	0.820	0.021

Label-Free Comparison: Progenesis LC-MS



Results

Model System: PTM Peptides in Plasma



PTM Map

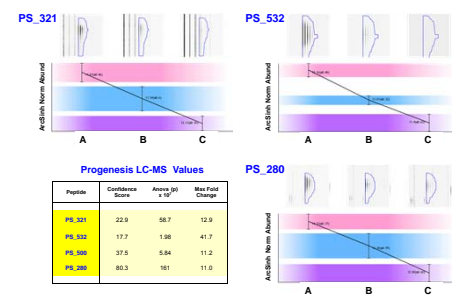
Legend: A (125 nM) B (62 nM) C (25 nM)

PS_280 SYSMEHFRWG (M4) O
PS_321 DRVYHFFHL (Y4) NO2
PS_500 EALSPDPAAS AAPLR (S4, S10) GalNAc
PS_532 DFNKHFHTFPQ TAIGV (K4) Ac

PTM Scoring

Peptide	Position	AA Index	Mod	A	B	C	Total Counts	Average Counts	SD	Unmodified Forms	Quality	Grouping	Occupancy	Uniqueness	PTM Score
PS_280	4	Y	14	125	62	25	16	5.333	2.517	0	0.890	1.000	1.000	0.885	100.000
PS_532	4	K	18	125	62	25	4	1.667	2.082	0	0.890	0.827	1.000	0.817	67.889
PS_321	10	R	18	125	62	25	7	2.333	2.082	12	0.890	0.827	0.288	0.821	68.292
PS_280	4	K	18	125	62	25	23	7.667	0.577	0	0.890	0.229	1.000	0.838	29.748

Label-Free Comparison: Progenesis LC-MS



Conclusions

- PTM counting is a powerful technique for rapid semi-quantitative integration of large mass spectrometry-based proteomics data sets.
- STRAP PTM represents a novel counting approach that uses a new scoring algorithm to rank PTMs in differential proteomics experiments.
- STRAP PTM software is easy to implement on a PC and provides fast turnaround for even large data sets (on the order of minutes).
- Select STRAP PTM results correlate well with label-free results from both simple and complex model data sets.

Acknowledgments:

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