# Counting Strategies for Differential Characterization Of Post-Translational Modifications

# McComb ME<sup>1</sup>, Spencer JL<sup>1</sup>, Bhatia VN<sup>1</sup>, Whelan SA<sup>1</sup>, Kehasse A<sup>1</sup>, Perlman DH<sup>2</sup>, Heckendorf CF<sup>1</sup>, and Costello CE<sup>1</sup>

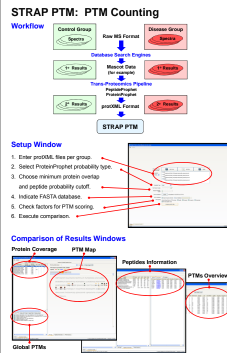
<sup>1</sup>Boston University School of Medicine, Boston, MA 02118

<sup>2</sup>Princeton University, Princeton, NJ 08544

## 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



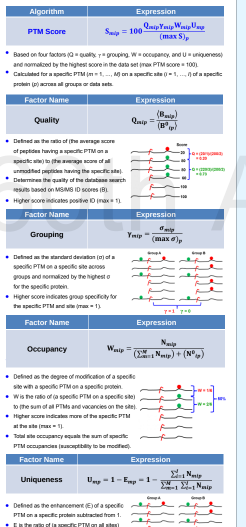
# Methods

to (all PTMs on all sites).

PTM (max = 1).

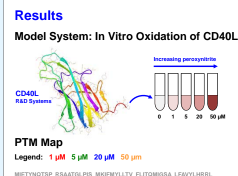
Higher score indicates more specificity in the

## STRAP PTM: PTM Scoring



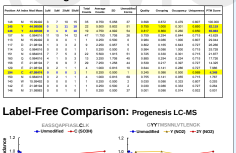
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E=3/9, U=8/9 E=6/9, U=3/9



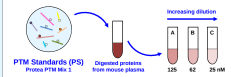
#### DKIEDERNLH EDFVFMIKTIQ RCNTGERSLS LLNCEEIKSQ FEGFVKDIML NKEETKKENS FEMAKGOONP QIAAHVISEA SSKTTSVLQW AEKGYYTMSN NLVTLENGKO LTVKRQGLYY IYAQVTFCSN REASSQAPFI ASLCLISPGR FERILLRAAN THSSAKPCGQ QSIHLGGVFE LQPGASVFVN TDPSQVSHG TGFTSFGLLK L (Y145, Y146) NO2 (C194) SO3H

## PTM Scoring



# Results

### Model System: PTM Peptides in Plasma

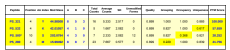


## PTM Map

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

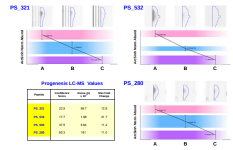
PS_280	SYSMEHFRWG	(M4) O
PS_321	DRVYIHPFHL	(Y4) NO2
PS_500	EAISPPDAAS AAPLR	(S4, S10) GalNAc
PS 532	DENKEHTEPQ TAIGV	(K4) Ac

### PTM Scoring



\* Peptide PS\_500 was also observed as a native mouse plasma peptide and thus was scored with unmodified forms.

## Label-Free Comparison: Progenesis LC-MS



# Conclusions

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Peroxynitrite (uM)

- > 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.

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Peroxynitrite (uM)

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