Targeted LC-SRM/MS Quantification of Mammalian Synaptic Proteins with Mouse Express® Brain Tissue, a New Isotopically Labeled Proteome Standard

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Maintenance and reshaping of synaptic connections in the brain underlies the ability to learn and remember. Complex protein machinery at the synapse is constantly seeking out, strengthening and pruning neuronal connections, literally rewiring in response to experience. The process by which neurons accomplish this feat has been termed “neuroplasticity.” Elucidating the molecular mechanics of neuroplasticity is critical to understanding how the human brain interacts with the world around it, as well as how neuropsychiatric diseases assault our memories, thoughts and motor function.

Trafficking of synaptic proteins is critical to neuroplasticity, and many of these trafficked proteins have been implicated in a broad spectrum of neuropsychiatric disorders.1,2 An in-depth understanding of how these trafficking events function could hold the key to developing new therapeutics. It has become clear that targeted multiplexed quantification of proteins across neuronal subcellular domains is essential to probe these normal and aberrant trafficking events. Mass spectrometry (MS)-based proteomics has emerged as a valuable tool for studying these phenomena. Stable isotope labeling with amino acids in cell culture (SILAC)-based methodologies have been successfully applied to cellular models of many diseases.3 However, this methodology has found limited utility in neuroscience research as complex psychiatric diseases are difficult to model in animals and are impossible to model in culture. Isobaric-labeling methodologies, such as iTRAQ, have proven more useful in assaying trafficking in humans and animal models,4,5 but peptide chemical-tagging methodologies suffer from compression of the quantitative signal and have difficulty repetitively quantifying a targeted subset of peptides. The availability of protein standards, from Cambridge Isotope Laboratories, Inc. (CIL), generated from stable isotope labeling in mammals (SILAM) with L-lysine·2HCl (13C6, 99%) (CLM-2247-H) allows multiplexed targeted quantitative analysis of protein trafficking in brain tissue without the costly and untimely synthesis of individual peptide or protein standards.

(continued)
“Heavy” (labeled) membrane fractions were prepared from 40 mg centrifugation and pH-specific Triton X-100 precipitation (prepared from the cortex of two mice by sucrose density gradient human sequences. Peptide homology allows this method to be containing at least one lysine with homology between mouse and Pinpoint™ (ThermoFisher Scientific) was utilized to select peptides synaptic fractions from mouse and human tissue were mined SRM design

**Experimental Design**

**SRM design:** Discovery data from 2D LC-MS/MS analyses of synaptic fractions from mouse and human tissue were mined for tryptic peptides from 200 synaptic proteins of interest. Pinpoint™ (ThermoFisher Scientific) was utilized to select peptides containing at least one lysine with homology between mouse and human sequences. Peptide homology allows this method to be utilized in clinical subjects as well as additional animal models.

**Fractionation and sample preparation:** “Light” (unlabeled) vesicular, presynaptic and postsynaptic density fractions were prepared from the cortex of two mice by sucrose density gradient centrifugation and pH-specific Triton X-100 precipitation (Figure 1). “Heavy” (labeled) membrane fractions were prepared from 40 mg Mouse Express® Brain Tissue (male) L-Lysine (13C6, 97%) (MT-LYSC6) using an abbreviated version of the fractionation method (Figure 1). “Light” fractions were mixed with Mouse Express® brain tissue preparations at a ratio of 2:1 (Figure 1). The mixed proteomes were separated on a 4-12% bis-Tris gel, divided into two fractions: (i.e. <64 kDa and between 64 – 250 kDa, reduced, alkylated and digested with trypsin (Figure 1)).

**LC-SMR/MS:** The resulting peptides were desalted offline and analyzed on a TSQ Vantage QqQ (ThermoFisher Scientific) interfaced to a NanoLC-Ultra 2D (Eksigent) and CaptiveSpray source (Michrom). A 5 μL peptide sample was loaded onto a 150 mm x 100 μm Magic C18 column (Michrom) at 1 μl/min 97% mobile phase A (H2O with 0.1% (v/v) formic acid) 3% mobile phase B (acetonitrile with 0.1% (v/v) formic acid) for 12 min, and eluted at 750 nl/min over a 25 min gradient from 3-35% mobile phase B. Retention windows were 1 – 1.5 min, depending on the number of SRMs to be analyzed. In this manner, SRM Dwell times are dynamic, based on the number of analytes eluting at a given time, and range from 5 to 60 msec per transition monitored in these experiments.

**Data Analysis**

First, light/heavy ratios were calculated for each peptide in Pinpoint™ using the integrated area under the curve (Figure 2). Where multiple peptides from a single protein were monitored, Pinpoint™ averaged the ratio values. To gain a global view of the data, the proteins were grouped into functional categories by gene ontology (GO) terms within ProteinCenter™ (ThermoFisher Scientific). Relative amounts of representative proteins from four of these categories are reported in Figure 3.

**Results**

We observed a significant interaction between microdomain partitioning and functions performed in those domains. A subset of the data for four families is reported in Figure 3. The postsynaptic density (PSD) fraction is highly enriched in kinases and scaffolding proteins, such as PSD-95. Scaffolding proteins link glutamate receptors to each other and kinases in PSD facilitating amplification and propagation of neurotransmitter signals. The synaptic vesicles, on the other hand, are highly enriched for soluble N-ethylmaleimide sensitive factor attachment protein receptor (SNARE) complex proteins, responsible for mediating vesicular fusion, and display a moderate enrichment of G-Protein subunits. The SNARE complex is responsible for fusing vesicles to the presynaptic membrane for neurotransmitter release.

**Discussion**

Quantitative analyses of subcellular proteomes has become increasingly important in neuroscience. Intracellular signaling pathways in the brain are no longer seen as a linear process from a molecule to the next, but are thought to occur in the context of numerous other protein interactions. This shift in paradigm has begun to impact the study of neuropsychiatric illnesses and thus, it is critically important to have the ability to quantify many proteins simultaneously with microdomain specificity.
Chromatogram for Synaptotagmin I peptide

LTVVILEAK was monitored in the endogenous “light” form and labeled “heavy” form. Chromatograms represent the signal from three SRM transitions representing the precursor MH+2 ion to the y5, y6 and y7 fragments in vesicular, presynaptic and PSD fractions.

A) Chromatograms for the “light” endogenous peptide monitored by three SRM m/z transitions in vesicular, presynaptic and PSD fractions.

B) Chromatograms for the “heavy” Mouse Express® peptides monitored by three SRM m/z transitions for the same fragment ions in the same fractions.

Figure 3. Reciprocal Enrichment of Synaptic Proteins: Relative amounts of select scaffold proteins, kinases, G protein subunits and SNARE complex proteins in vesicular (A), presynaptic (B) and PSD (C) biochemical fractions are reported. SNARE complex proteins are enriched in the vesicular fraction over other proteins, while scaffold proteins and kinases are moderately enriched in the presynaptic fraction and substantially enriched in the PSD. (D) A depiction of the synaptic biochemical fractions.
The neuronal protein standards prepared from SILAM tissue facilitated the development of a multiplexed assay for the targeted quantification of synaptic proteins. This method represents a significant advance over existing methodologies. Stable isotope-labeled peptide standards may allow for absolute quantification of target proteins and PTMs, but assaying large numbers of proteins can be prohibitively expensive. Furthermore, digestion efficiency and peptide recovery can be difficult to control for using peptide standards. We have found the reproducibility of the LC-SRM/MS with the Mouse Express® brain tissue SILAM standard to be superior to quantification with AQUA peptides when samples incur lengthy predigestion sample preparation. Isobaric-labeling methods allow for the quantification of large numbers of proteins, but performing targeted analysis can be difficult and tissue requirements can be high, which is problematic when assaying proteins in discreet rodent brain areas or in precious human samples. Sensitivity of the triple-stage quadrupole mass spectrometer is an important parameter when analyzing low abundance proteins in the experiments described above. Additionally, the Mouse Express® brain tissue SILAM standard acts as a carrier, preventing non-specific losses of low abundance proteins in precious samples. Finally, due to the highly conserved nature of neuronal proteins across mammalian species, Mouse Express® brain tissue can be used to quantify synaptic proteins in many animal models, and, perhaps most importantly, human tissue.

We have successfully utilized the method described above to quantify synaptic proteins in five subcellular fractions prepared from human brain tissue. Currently, the method has been validated for 120 proteins. This number will continue to expand as new data sets are continuously being published and online databases, such as SRM Atlas, grow. The addition of SILAM standards prepared from Mouse Express® brain tissue to subcellular fractionation-targeted LC-SRM/MS methodologies represents an essential powerful new tool for the study of neuropsychiatric illness.

### Related Products

**Mouse Express® Mouse Tissue**

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<th>Catalog No.</th>
<th>Description</th>
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<tr>
<td>MFLYS6-MB</td>
<td>Mouse Express® Brain Tissue (M) L-Lysine (13C6, 97%)</td>
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<tr>
<td>MFLYS6-FB</td>
<td>Mouse Express® Brain Tissue (F) L-Lysine (13C6, 97%)</td>
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**Mouse Express® Mouse Feed**

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<td>MLK-LYS-C</td>
<td>Mouse Express® L-Lysine (13C6, 99%) Mouse Feed Kit: 1 kg of L-lysine 13C6 feed and 1 kg of (unlabeled) feed</td>
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### References


