Short communications

Proteome profiling of s-nitrosylated synaptosomal proteins by isobaric mass tags

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\textbf{HIGHLIGHTS}

- Protocol for quantitative proteomics of nitrosylation on synaptosomal proteins.
- Identification of endogenous nitrosylation independent of induction by NO donors.
- Use of iodoTMT sixplex mass tags for stable labeling, enrichment, identification, and multiplex quantitation.
- Applicable on low amounts of sample material of mouse and human brain tissue.

\textbf{GRAPHICAL ABSTRACT}

A \textbf{R} \textbf{T} \textbf{I} \textbf{C} \textbf{L} \textbf{E} \textbf{I} \textbf{N} \textbf{F} \textbf{O}

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1. Introduction

Nitric oxide (NO) is a free radical that is biosynthesized by nitric oxide synthases. NO contributes to numerous physiologic responses such as vasculature vasodilation, synaptic neurotransmission, respiration, inflammation, and apoptosis (Swomley et al., 2014). One of the important biological modifications induced by NO is S-nitrosylation of cysteine (SNO), which functions in regulating signal transduction. In neurodegenerative disorders, aberrant SNO can propagate protein misfolding, mitochondrial dysfunction, synaptic injury, and neuronal death (Nakamura et al., 2013). For example, increased levels of nitrosylated proteins have been reported in the brain of Alzheimer disease (AD) patients (Heneka et al., 2015). As one of the earliest pathological characteristics of AD is synapse loss, our aim was to establish a robust protocol for the analysis of synaptosomal S-nitrosylation in mouse and human brain.

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samples. Synaptosomes consist of the presynaptic terminal (which contains synaptic vesicles and mitochondria), as well as the postsynaptic density and postsynaptic membrane. They are preferably obtained by homogenization of brain tissue and density gradi-
tent centrifugation for enrichment of this specific compartment. (Swomley et al., 2014). Numerous proteomics techniques have been developed for detection and quantification of SNO (Qu et al., 2016). However, many of these require high amounts of starting materials in the range of 100–500 µg of protein lysate (Wojdyla and Rogowska-Wrzesinska, 2015). Furthermore, a common procedure applied to analyze SNO modification sites is induction of nitrosyla-
tion by use of NO-donors. It is difficult to compare the results from studies using NO donors to the in vivo situation, where cysteine S-nitrosylation depends on the undetermined concentrations of nitrosylating compounds (Zareba-Kozioł et al., 2014). Hence, iden-
tification of endogenous SNO levels derived from sub-fractionated compartments such as synaptosomes requires high sensitivity and specificity of the detection strategy. Typical approaches for quan-
titative SNO proteomics are two-dimensional gel electrophoresis or liquid chromatography (LC) followed by mass spectrometry. Two-dimensional difference in-gel electrophoresis (2D-DIGE) has been described using a modified labeling procedure (NitroDIGE) (Qu et al., 2014b). More frequently used is the biotin switch tech-
nique (BST) introduced by Jaffrey and Snyder (2001). It utilizes a replacement strategy to add a stable biotin into a SNO modifica-
tion, allowing their detection and identification by antibodies or MS (Jaffrey and Snyder, 2001). More recently, an MS-based, iso-
baric tag labeling strategy was introduced in which peptides are linked with different tags and then fragmented to produce reporter ions of specific masses (Dayon and Sanchez, 2012). Similar to BST, the commercially available labeling reagents (iodoTMT™) are used for replacement chemistry of SNO modifications in a sixplex multiplex (Qu et al., 2014a). Although introduced in 2012, this approach has been barely described in the literature for samples with low amounts of starting material and under basal, endoge-
nous conditions without NO donor treatment. This article provides detailed descriptions of the methodological workflow (Fig. 1) and efficiency of the iodoTMT™ protocol when applied to the analy-
sis of synaptosomal samples, discussing both technical variances of the method as well as the comparability of findings with results previously published on this field.

2. Materials and methods

2.1. Tissue samples

Three 9-months-old, male wild type mice were euthanized by iso-
flurane and transcardially perfused with ice-cold normal saline (0.9% w/v NaCl). Mouse brain hemispheres without brain-
stem and cerebellum were prepared and immediately used for protein extraction. The method was transferred to two human frontal cortex brain samples from non-demented, non-diseased donors. Human brain samples were obtained from unfixed, frozen tissues with a postmortem interval of less than 4 h from the Banner Sun Health Research Institute Brain and Body Donation Program in Sun City, Arizona (Beach et al., 2008). All subjects participating in the Banner Brain and Body Donation Program signed the writ-
ten informed consent approved by the Banner Health Institutional Review Board.

2.2. Synaptosome isolation

For preparation of synaptosomes, 200 mg tissue where used with equal amounts of white and grey matter. Tissues were homog-
enized in 9-fold volume of tissue weight in 0.32 M sucrose buffer (50 mM Tris–acetate, 1 mM EDTA, 1 mM EGTA, 5 mM disodium pyrophosphate, 5 mM NaF, 2 mM Na3VO4, 1 mM AEBSF, cComplete protease inhibitor cocktail (Sigma-Aldrich, Munich, Germany), pH 7.4). Homogenization was performed manually by 15 slow up and down strokes using a Teflon-glass homogenizer (0.1–0.15 mm clearance). Samples were centrifuged to remove nuclei and cellular debris at 800g for 5 min at 4°C. The collected supernatant was lay-
ered on top of a discontinuous sucrose gradient consisting of two layers of sucrose buffer (1.0 M and 1.4 M). Gradient centrifugation was performed using a MLS-50 swinging bucket rotor (Beckman,
Krefeld, Germany) at 54000 g for 90 min at 4°C. The synapto-
somal fraction was collected from the interface between 1.0 M and 1.4 M sucrose layers. The fraction was diluted with 4-fold volume of HPLC-grade water and sedimented at 54000 g for 15 min at 4°C. The supernatant was discarded and the synaptosomal pellet was stored at −80°C. Enrichment of pre- and post-synaptic proteins was evaluated using electron microscopy and Western blotting against synaptophysin (1:1000, Abcam, ab52636), post-synaptic density protein (PSD95, 1:2000, Thermo Fisher Scientific, MA1-064), NMDA receptor 2B (NMDAR2B, 1:1000, Millipore, ab1557p), the nuclear marker Lamin B1 (1:500, Proteintech, 66095-1-lg) and alpha tubul-
in (1:2000, Thermo Fisher Scientific, 62204) as a loading control.

2.3. iodoTMT workflow for synaptosomal proteins

2.3.1. Protein labeling with iodoTMT

Three technical replicates were used to evaluate this approach. Synaptosomal peptides were resuspended in 1 µg µL−1 HENS lysis buffer (Thermo Fisher Scientific). Each synaptosome sample was prepared for six aliquots with an equal amount of proteins (50 µg), which were then individually labeled with iodoTMT sixplex labeling reagent followed by the optimized protocol. Subsequently, samples were digested, desalted, enriched, and analyzed by MS.

To avoid rearrangements of thiol-modifying groups, the protein mixture was treated with MMTS (20 mM final concentration, stock diluted in dimethyl formamide) at room temperature for 30 min with frequent vortexing (protected from light). Excess MMTS was removed by acetone precipitation at −20°C for 1 h. The pellets were resuspended in 50 µL of HENS buffer and reduced with 1 µL of 1 M sodium ascorbate in water. Each sample was immediately labeled with 1 µL of iodoTMT labeling reagent (0.2 mg in 10 µL of LC/MS grade methanol) at 37°C for 1 h, protected from light. Excess iodoTMT reagent was quenched with 0.5 M dithiothreitol (20 mM final concentration) at 37°C for 15 min. Up to six different samples labeled with iodoTMT were combined and alkylated with 10 µL of 0.5 M iodoacetamide in 300 µL of HENS buffer at 37°C for 1 h.

2.3.2. Tryptic digestion and peptide desalting

Alkylated samples were precipitated with acetonitrile precipi-
tation and dissolved in 50 mM ammonium bicarbonate buffer, pH 8 for overnight digestion using trypsin (protein: trypsin ratio = 1:40 w/w) at 37°C. The peptides were acidified with 10% tri-
fluoroacetic acid to final concentration of 0.5%, purified using C18 spin columns (Thermo Fisher Scientific) and eluted with 70% (v/v) acetonitrile (ACN).

2.3.3. Enrichment of iodoTMT-labeled peptides

For the enrichment step, the labeled peptides were dried by vac-
uum centrifugation and resuspended in 100 µL TBS buffer (150 M NaCl, 50 mM Tris, pH 8.0). The peptide samples were incubated with 30 µL Anti-TMT resin for 2.5 h at room temperature with end-over-end mixing. The resin was washed three times with 100 µL TBS followed by three washing steps with 100 µL HPLC-grade water. Peptides were eluted twice with 30 µL TMT elution buffer (Thermo Fisher Scientific). Pooled eluates were dried by vacuum centrifu-
Fig. 1. The isobaric tag labeling procedure.

(A) Schematic image of the isobaric tag labeling procedure. After synaptosome isolation, SNO proteins are labeled by (i) blocking of free thiols using MMTS; (ii) reduction of cysteine S-nitrosylation with ascorbate, producing new free thiols on previous SNO sites; and (iii) isobaric tag labeling of the new thiols using iodoTMT. iodoTMT-labeled proteins are subjected to in-solution protein digestion. Derived peptides are enriched using affinity-capture on an anti-TMT resin. Eluted peptides are identified by nanoHPLC–MS/MS and quantified using the TMT sixplex.

(B) A representative MS/MS spectrum of the iodoTMT-labeled peptide SICDTSNFSDYIR derived from ubiquitin-like modifier-activating enzyme 1. Peaks of the 6 reporter ions are found in the enlarged section of the lower mass range.

2.3.4. LC–Mass spectrometry analysis

A C18 trap column (20 mm length, 100 μm inner diameter) was coupled to a C18 analytical column (200 mm length, 75 μm inner diameter) packed in-house with 1.9 μm ReproSil–Pur 120 C18-AQ particles (Dr. Maisch, Ammerbuch, Germany). Solvent A consisted of 0.1% formic acid. Solvent B consisted of 90% acetonitrile, 0.1% formic acid. Separation of the peptides used a linear gradient from 4% to 35% solvent B within 240 min at a flow rate of 300 nl/min. The nanoHPLC was coupled online to an LTQ Orbitrap Velos mass spectrometer (Thermo Fisher Scientific, Bremen, Germany). Peptide ions between 330 and 1700 m/z were scanned in the Orbitrap detector with a resolution of 30,000 (maximum fill time 400 ms, AGC target 106). The 20 most intense precursor ions (threshold intensity 5000) were subjected to higher energy collision induced dissociation (HCD) and fragments were again analyzed in the Orbitrap. Fragmented peptide ions were excluded from repeated analysis for 15 s.

2.4. Data analysis and protein identification

For data processing and analyses of database searches, Proteome Discoverer software 2.1.0.81 (Thermo Fisher Scientific) was used. Peptide identification was done on an in-house Mascot server,
version 2.5.1 (Matrix Science Ltd, London, UK). MS2 data (including a-series ions) were matched against mouse/human sequences from SwissProt (release 2015_04). Precursor ion m/z tolerance was 10 parts per million (ppm), fragment ion tolerance was 20 milli-
mass units (mmu). Up to two missed cleavages were allowed for
tryptic peptides. Low scoring spectrum matches were re-analyzed
with semitryptic specificity with up to one missed cleavage. Car-
bamidomethylation (Cys), Oxidation (Met), acetylation (protein
N-terminus), and iodoTMT sixplex (on Cys) were set as dynamic modifications. Mascot results from searches against SwissProt were
sent to the percolator algorithm version 2.05 as implemented in
Proteome Discoverer. Peptides were identified with a posteriori
error probability of 1% or better. Proteins were reported with at
least one matching peptide because of the experimental design: In
cases where only one site in a protein is nitrosylated, only one
labeled peptide can be identified after enrichment.

3. Results and discussions

3.1. Method performance

Technical characterization of the method was tested first on
wild-type mouse samples (n = 3) and then transferred to human
brain samples (n = 2) from donors without CSN disorders. To
test reporter ion intensity and determine inter-sample variances,
aliquots of the same samples were labeled with iodoTMT™ sixplex
reagents, one for each aliquot, and then pooled and analyzed (same-
same experiment). A representative MS/MS spectrum with intense
reporter ions and a clear b- and y-series of assigned fragment ions
is shown in Fig. 1B. When comparing reporter ion intensities from
the same-same-experiment over all identified peptides between all
of the 6 labels, there was an average variance of 20% between single
reporter ion channels and the mean of all channels, quantified
over all of the identified peptides. This indicates that the overall
technical variance of the method, including preparative pipetting,
sample handling, digestion, iodoTMT™ labeling and enrichment
until pooling and analysis of the samples, was within this range.

In theory, all identified peptides should contain a TMT™ label
attached to a previously nitrosylated cysteine. Within our dataset,
an average of 72% of identified peptides contained cysteine, among
which 93%–98% were labeled with iodoTMT™. In conclusion,
cysteine-targeted labeling was highly specific, whereas the anti-
TMT-antibody based enrichment and purification strategy could
not completely remove non-modified peptides. There was barely
any unspecific TMT-labeling (<1% of peptides) on the peptide N-
terminus (which is the target of the regular TMT™ protocol). Nor-
worthy, it was not possible to identify nitrosylated peptides form
the synaptosome samples without application of the label-
ing/enrichment strategy.

3.2. Proteome analysis

In murine synaptosomes 1462 unique peptides were identified
representing 617 proteins, of which 42% were identified for the first
time. In human synaptosomes, 570 unique peptides were identified
representing 337 proteins, of which 73% were identified for the first
time (by comparing to the SNO database http://disSNO.mbc.
tcu.edu.tw). In comparison, the absolute number of SNO-modified
proteins was 2.6-fold higher in mouse proteins compared to human
proteins. This matches data found on the SNO database (http://140.
138.144.145/~disSNO/statistics.php), which at this point lists 1494
murine and 720 human SNO proteins. The lower number of SNO
proteins found in human tissue by our approach and described
in the database could be attributed to both technical as well as
biological variances: Technically, murine brain samples can be
prepared directly after euthanizing the animal, which is impossible
for human brain samples. Furthermore, due to perfusion with
cold saline solution, mouse brain samples are cooled faster. Con-
trolled cold temperature can be used to reduce the activity of
nitric oxide synthases, as it has been demonstrated in example
for NOS2 (Venturini et al., 1999). For human samples, the post
mortem interval was limited to a maximum of 4 h for this study,
but this time window might already impact the yield of nitro-
sylated proteins. Additionally, synaptosomes from mouse brains
could be prepared without freeze-thaw cycles, while human brain
samples were obtained as frozen material from a biobank, which
did not prepare its specimen specifically for this type of analysis.
Supposable, the SNO proteome could be further preserved by use
of appropriate chemical stabilizers or inhibitors additional to cool-
ing and protease inhibitors, as it is common practice in example
for phosphoproteomics. To our knowledge, such substances have
not found widespread use yet, although some studies reported use
of inhibitors (Koo et al., 2016; Foster et al., 2012). Lastly, there
might be substantial species-derived differences between murine
and human samples, which cannot be entirely distinguished from
method-derived differences to this point.

The composition of the SNO-proteome was analyzed using
Gene Ontology enRichment analysis and visualIAtion (Gorilla)
search engine (Eden et al., 2007) and Uniprot/SwissProt based
on the localization classification (UniProt, 2013). The Gorilla
search engine result (illustrated in Fig. 2A) indicates enriched
terms in the Cellular Components of organelle membrane
(GO:0031090), oxidoreductase complex (GO:1990204), cytoplasmic
part (GO:0044444), organelle inner membrane (GO:0019866),
intracellular organelle (GO:0043229), intracellular membrane
bounded organelle (GO:0043231) and components of mitochondr-
ial (GO:0044429; GO:0005739; GO:0031966; GO:0044455; GO:0005743). Definitions of the specific components are available
from the Gene Ontology Consortium (http://www.geneontology.
org/) database. This result revealed the enrichment of synaptosome
components such as cytoplasmic part, cell membrane and mito-
ochondria that are enriched in the presynaptic terminal (Swonley
et al., 2014).

Classifications based on Uniprot/SwissProt localization entries
showed comparable enrichments of typical synaptosome compo-
ments between the unlabeled synaptosome (2B) and the iodoTMT
labeled SNO synaptosome (2C), both obtained from our protocols
and results published by others on regular TMT labeling of synaptosome proteomes (2D). The synaptosome SNO-proteome
(geroTMT-labeled) contained a large fraction of mitochondrial pro-
teins (30%), presumably originating from the high amounts of
mitochondria located in synapses (Fig. 2C). Additionally, large fractions of cytoplasmic (25%) and cell membrane (20%) associated proteins as well as a small fraction of concrete synapse associated proteins (6–8%) were found, all well in accordance with the expected composition of synaptosomes. The distribution of the identified synaptosomal proteins according to their localization
between unlabeled, and labeled synaptosome (iodoTMT or
TMT) shows analogous enrichment results with a majority of cyto-
plasm 24–28%, mitochondria 24–34%, cell membrane 14–22%, and
synapse 6–11%. We also found nuclear and endoplasmic reticu-
lar associated proteins resulting from small fragments of these
organelles with similar density as synaptosomes.

For the murine samples, these numbers are comparable with,
if not higher than, those described in previous reports on synap-
osome samples: In example, SNOsID (SNO side identification)
analysis using BST identified 138 SNO proteins (Zareba-Kozio1 et al.,
2014) of which 85% were found to overlap with our results (see
Fig. 2G); and TMT (Tandem Mass Tag) labeling analysis identified
240 proteins (Fig. 2F), both from murine synaptosome. For
human samples, to our knowledge, there are no published data for
synaptosome-derived SNO-proteomes. We found that the overall composition of the detected proteome is congruent with previous publications on the synaptosome proteome labeled with TMT independent of their modifications (Fig. 2D) (Engmann et al., 2010) and with BST on SNOSID (Fig. 2E) (Zareba-Koziol et al., 2014).

3.3. Advantages and limitations

The iodoTMT approach showed a large number of identified proteins obtained from a small amount of starting material (50 μg of protein from synaptosome lysate). The number of identified SNO proteins was comparable or higher in comparison to previous studies. Further advantages include high labeling specificity and the use of one stable labeling for enrichment, identification and multiplex quantitation of modified proteins. Also, this protocol was functional for identification of endogenous levels of SNO proteins, without prior induction of nitrosylation by NO donor treatment. Finally, the iodoTMT protocol was easy to adapt and did not require extensive method optimization.
However, this approach also has limitations. The stability of the iodoTMT labeling reagent after reconstitution is limited to 1 week when stored at −20 °C. Therefore, it is recommendable to perform all labeling steps for a study within this time frame.

Another limitation stems from the nature of the labeling and enrichment procedure: As only modified peptides are identified and quantified, the procedure frequently depends on MS/MS-based information implied from a single peptide ion. This limits the number of peptides available for protein identification, which in consequence should be achieved from that single peptide. Therefore, protein identifications are only valid when derived from unique (protein-specific) modified peptides. If identified peptides are not unique, multiple proteins or protein classes have to be considered as potential origin of the peptide.

Although highly efficient, the entire workflow consists of multiple steps (such as blocking, labeling, enrichment, cleaning, digestion) which are time-consuming and can add up to inter-run variances, which were in an average range of 20% in the same experiments. To control this in quantitative studies, we suggest setting up an internal standard that contains pooled and aliquoted proteins of all representative samples. This standard should be added at the beginning of each preparation batch and be analyzed as one of the sixplex samples on each LC-MC-run, hence enabling to control for any technical failures or irregularities that interfere with quantification. Although this standard cannot reduce inter-run-variances, it could be used to characterize and monitor such variances for quality control purposes.

4. Conclusions

In summary, the protocol possesses many advantages in terms of enrichment, specificity, and quantitation, all based on a single labeling reaction. Additionally, the iodoTMT approach enables sample (6-plex) multiplexing. The protocol was sensitive enough to utilize low amounts of starting material while at the same time providing data comparable to previously published results. Given the increasing scientific interest on protein S-nitrosylation (Qu et al., 2016), we anticipate it to be useful for other researchers analyzing this instable protein modification, especially when dealing with limited amounts of samples such as in analysis of synaptosomes or other cellular compartments.

Conflict of interest

None.

Authors contribution

TSW performed initial method screening, animal and human brain preparations, synaptosome isolation and protein labeling and drafted the manuscript, MS and NBA performed the mass spectrometric measurements and MS data analysis, FB and MTH aimed the study, supervised experimental design and interpretation and drafted the manuscript, VG and MPK approved the study and drafted the manuscript.

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