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Strategies for profiling native S-nitrosylation

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Abstract

Cysteine is a uniquely reactive amino acid, capable of undergoing both nucleophilic and oxidative post-translational modifications. One such oxidation reaction involves the covalent modification of cysteine via the gaseous second messenger nitric oxide (NO), termed S-nitrosylation (SNO). This dynamic post-translational modification is involved in the redox regulation of proteins across all phylogenic kingdoms. In mammals, calcium-dependent activation of nitric oxide synthase triggers the local release of nitric oxide, which activates nearby guanylyl cyclases and cGMP-dependent pathways. In parallel, diffusible nitric oxide can locally modify redox active cellular thiols, functionally modulating many redox sensitive enzymes. Aberrant S-nitrosylation is implicated in the pathology of many diseases, including neurodegeneration, inflammation, and stroke. In this review, we discuss current methods to label sites of S-nitrosylation for biochemical analysis. The most popular method involves a series of biochemical steps to mask free thiols followed by selective nitrosothiol reduction and capture. Other emerging methods include mechanism-based phosphine probes and mercury enrichment chemistry. By bridging new enrichment approaches with high-resolution mass spectrometry, large-scale analysis of protein nitrosylation has highlighted new pathways of oxidative regulation.

Introduction

Sulfur is the lightest element that can produce stable exceptions to the octet rule due to the presence of "*d*" orbitals. Typical cysteine residues in proteins have a side chain pK_a values of 8.0¹, and thus ~10% of cysteine thiols are in their reactive thiolate form at physiological pH. However, many redox active or catalytic cysteine residues have dramatically reduced pK_a values. Such thiolates have evolved to promote catalysis or redox regulation. For example, the pK_a of the catalytic cysteine in methionine sulfoxide reductase is reduced to 5.7 upon substrate binding^{2,3}. Similarly the active site thiol of glutaredoxin has a low pK_a near $3.5^{4,5}$. Such altered acid-dissociation constants enhance thiol reactivity, which in turn promote reactions with electrophilic oxidants to produce distinct post-translational modifications.

In this review, we focus on the chemistry and dynamics of protein S-nitrosylation. This unique oxidative modification directly modulates the localization and activity of cellular proteins involved in cellular growth and regulation^{6,7}. In neurons, stimulus-dependent

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depolarization leads to calcium influx, calmodulin activation, and stimulation of nitric oxide (NO) synthases^{8–10}. Local nitric oxide release induces spatially restricted S-nitrosylation of channels, phosphatases, and other redox active thiols^{6,11,12}. Emerging proteomics studies implicate hundreds of endogenous sites of nitrosylation^{13–16}, although the stoichiometry and functional consequences of these post-translational remains an active research area.

There are several mechanistic routes leading to protein S-nitrosylation (Figure 1). Hemedependent nitric oxide synthases generate nitric oxide via a two-step, five-electron oxidation of L-arginine¹⁷⁻²⁰. This reaction uses two moles of molecular oxygen and 3/2 moles of NADPH per mole of nitric oxide formed²¹. Nitric oxide itself is not especially reactive towards protonated cellular thiols, particularly under aerobic conditions 2^{2-24} . In order to generate nitrosothiols, nitric oxide must first undergo secondary oxidation to nitrogen dioxide, which occurs via at least two distinct pathways²⁵. In the first pathway, nitric oxide reacts with a superoxide radical to make peroxynitrite. Peroxynitrite (pKa of 6.5^{26}) converts to peroxynitrous acid at physiological pH, which undergoes hemolytic cleavage to form the hydroxyl and the nitrogen dioxide radicals. The nitrogen dioxide radical can in turn react with nitric oxide to form dinitrogen trioxide, which reacts with thiolates to form nitrosothiols²⁴. In a second pathway, nitrogen dioxide reacts with a thiolate to generate nitrite and a thiyl radical²⁷. The resulting thiyl radical is the only species able to react directly with nitric oxide to generate nitrosothiols. Additionally, metal-dependent formation of thivl radicals²⁸ promotes S-nitrosylation by one-electron oxidation of thiols to thivl radicals, or through metal-nitrosyl complex intermediates^{29,30}. Furthermore, iron and nitric oxide spontaneously react to form dinitrosyliron complexes (DNIC), which can be intermediates in nitrosothiol formation $^{31-34}$. All of these routes generate diffusible reactive radicals with enhanced reactivity towards thiols with reduced pK_a values, such as catalytic or redox-active thiols.

S-nitrosylation is reversible, either by nitric oxide release or by direct transfer to other cellular thiols^{35,36}. Such trans-nitrosylation reactions mobilize the exchange of nitric oxide from one protein to another, relaying nitroso-oxidation through multiple carriers 37-40. Trans-nitrosylation provides another route for the dynamic exchange of nitrosothiols. Millimolar glutathione levels maintain an intracellular reducing environment that protects proteins from oxidative modifications⁴¹. Abundant glutathione scavenges nitrosothiols by trans-nitrosylation, yielding a reduced protein thiol and nitrosoglutathione, which is reduced either by S-nitrosoglutathione reductase (GSNOR) or thioredoxin cascades. Such thiol exchange reactions are prevalent at physiological pH and predominate in comparison to hydrolysis reactions⁴², and are driven by the levels of reduced thiols in a given environment⁴³. Several studies have demonstrated trans-nitrosylation cascades relay the nitroso adduct from one protein thiol to another, eventually nitrosylating and inactivating select enzymes, for example nuclear chromatin-modifying enzymes⁴⁴. This model suggests stable S-nitrosylation sites are protected from the cellular environment, masked inside proteins or membranes. These studies demonstrate the surprising resilience of certain nitrosylated proteins in face of millimolar glutathione, and hints at orchestrated pathways of nitrosothiol transfer in cellular regulation. Indeed, hydrophobicity does enhance the rate of reaction between NO and oxygen by several fold⁴⁵, suggesting that thiols in hydrophobic

environments, either in membranes or in hydrophobic protein domains, may be more prone to stable S-nitrosylation⁶. Overall, protein S-nitrosylation is modulated by thiol pK_a , vicinal hydrophobicity⁶, proximity to NOS enzymes and by activities of redox enzymes such as thioredoxin, GSNOR, and accessibility to reduced glutathione.

Protein nitrosylation functionally regulates protein activities by transiently occupying thiol residues. Functional cysteines often reside in the active sites of enzymes, such as phosphatases, proteases, acyl-transferases, and ubiquitin ligases⁴⁶. These thiols reside in environments that promote thiolate formation by reducing the side-chain pK_a , leading to a more redox-active cysteine. Importantly, the active site of any enzyme is more likely to be protected from the environment, which likely prevents exchange with bulkier thiols. Accordingly, more stable S-nitrosylation is inversely correlated with thiolate exposure, and stabilized in protected environments. Indeed, nitrosylation of cellular phosphatases potentiates kinase cascades to promote cell growth, or inactivates lipid phosphatases during ischemic stroke^{47–49}. Due to the selective targeting of functional cysteines, nitrosylation may serve as a general redox switch important for the reversible inactivation of functional cysteine residues. For example, nitrosylation of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) recruits the ubiquitin ligase Siah1, and the GAPDH-Siah1 complex translocates to the nucleus where Siah1 targets nuclear proteins for degradation 50-52. Similarly, argininosuccinate synthase, the enzyme that generates arginine from citrulline⁵³, is inhibited by S-nitrosylation at Cys132⁵⁴. Elevated nitric oxide levels induce argininosuccinate synthase nitrosylation and inactivation, providing an autoregulatory loop that prevents excess oxidant production.

S-nitrosylation can also directly compete other reversible cysteine post-translational modifications, such as protein palmitoylation. Upon synaptic stimulation, protein palmitoyl thioesterases remove the membrane anchoring lipids from the neuronal scaffolding protein PSD-95, promoting egress from the post-synaptic density⁵⁵. Simultaneous activitydependent calcium influx activates neuronal nitric oxide synthase (nNOS), which is stably associated with PSD-95 via PDZ-dependent interactions. This locally generated flux of NO directly nitrosylates the newly de-acylated thiols, thus blocking further palmitoylation and membrane association. The mutually competitive modification of PSD-95 cysteines orchestrates synaptic release through an exchange of specific cysteine post-translational modifications⁵⁵. Furthermore, S-nitrosylation of the NMDA receptor leads to channel desensitization and channel closing, preventing excitotoxicity⁴⁸. This cascade of posttranslational events ensures proper membrane release of PSD-95, and ensures proper channel desensitization. This example raises the question if such mutually competitive modifications are unique for PSD-95, or if many proteins undergo dynamic exchange of cysteine PTMs. Furthermore, PSD-95 nitrosylation is channeled by protein-protein interactions, which provides selectivity for the deacylated thiols. The breadth of cysteine modifications competition is unknown. Interestingly, several palmitoylated proteins, including Ras and G-proteins, are rapidly de-palmitoylated after receptor stimulation^{56–58}, which coincides activation of NADPH oxidases and NO synthases.

Ascorbate-dependent enrichment strategies

Sensitive and selective labeling tools are critical for the precise detection and annotation of S-nitrosylation. Despite the acceptance of S-nitrosylation as an important protein regulatory modification, there remains a lack of direct methods to study this reversible redox modification. Early approaches used chemiluminescent, colorimetric or electrochemical methods to detect total nitric oxide liberated from nitrosylated thiols in a sample⁵⁹. These methods measure bulk release, and do not distinguish between heme, metabolite, or protein sources. Furthermore, these approaches eliminate any information about the sites and dynamics of nitrosylation on select proteins.

The "biotin-switch" technique was a major advance in the study and annotation of Snitrosylated proteins⁶⁰. This widely adopted method involves a series of biochemical steps, beginning with addition of the alkylating agent 2-iodacetamide, or by methyl methanethiosulfonate (MMTS) to block all free thiols. After removing the thiol capture reagents, the sample is treated with ascorbate, which reduces nitrosothiols to generate free sulfhydryl groups. This approach is selective for S-nitrosylation over other oxidative modifications, largely due to the unique mechanism of indirect reduction (Figure 2) 61,62 . In the presence of nitrosothiols, ascorbate undergoes a trans-nitrosation reaction involving nitrosonium (NO⁺) transfer to generate nitrosoascorbate, which decomposes to nitric oxide and the semi-dehydroascorbate radical. Therefore, ascorbate does not directly donate an electron for nitrosothiol reduction. This distinct mechanism is thought to provide chemical orthogonality to other oxidative modifications, making ascorbate an ideal nitrosothiolselective reducing agent⁶². Following ascorbate-mediated reduction, the newly unmasked thiols groups are then captured using a pyridyldithiol-activated, sulfhydryl-reactive biotinlinked probes for affinity enrichment⁶⁰. While these reagents are commonly available. several commercial kits are available that include ascorbate, metal-chelating buffers, MMTS, and iodoacetamide detection reagents. Recent adaptations for quantitative proteomics labeling strategies are also commercially available, and have been used to quantify individual sites of S-nitrosylation⁶³. Thiol resin assisted capture (RAC) replaces biotin and streptavidin purification with direct disulfide capture to activated thiol resin immediately after ascorbate treatment^{64,65}. This approach simplifies the procedure, and eliminates non-specific enrichment of endogenous biotinylated carboxylases.

Ultimately, the biotin-switch purification method is indirect, and highly dependent on the complete alkylation of all free thiols. Protein nitrosylation is a low abundance modification, so even low levels of uncapped thiols can lead to a high false positive rate. Interestingly, a heating step is generally included during thiol capping, often to 50° C. S-nitrosothiols are known to undergo thermal decomposition via homolytic cleavage of the S-N bond to yield the corresponding disulfides and nitric oxide, the latter is then oxidized to nitrogen dioxide^{66,67}. This step may introduce later complications, as the thermal stability of distinct nitrosothiols has not been thoroughly evaluated. Clearly, prolonged heating will promote nitric oxide release, and potentially suppress detection of labile nitrosothiols. Furthermore, after ascorbate reduction, newly free thiols are able to exchange with existing disulfide bonds, scrambling native sites of nitrosylation. Resin assisted capture is likely to reduce the extent of scrambling by providing high excess of activated disulfides for immediate

capture⁶⁵. Another source of false-positives is sunlight driven disulfide reduction, which can be eliminated by performing all procedures in complete darkness⁶⁸. This unfortunate restraint makes sample preparation more tedious, but is essential to eliminate non-specific disulfide reduction. Overall, biotin-switch technique is the current standard for nitrosothiol labeling, enrichment, and analysis.

Caveats of nitric oxide donors

Thousands of S-nitrosylated proteins have been reported from ascorbate-dependent enrichment using the biotin-switch method and mass spectrometry¹³. Unfortunately, nearly all reported proteomics data was collected from biological samples after the addition of exogenous or physiological nitrosylating agents. Upon donor release, gaseous NO is oxidized by molecular oxygen to form a peroxynitrite radical, which then reacts with a second NO molecule to generate two molecules of nitrogen dioxide. This reaction is limited by an apparent third-order rate law ($k = 2.5 \times 10^6 \text{ M}^{-2} \text{s}^{-1}$)^{69–71}, which means the reaction rate depends on the product of square of the nitric oxide concentration and molecular oxygen⁶⁹. Therefore, at very low nitric oxide concentrations the reaction is very slow, but at high nitric oxide concentrations the reaction is extremely rapid. Thus, depending on the concentration of nitric oxide released, the half-life of the reaction can range from 0.5 seconds to 50 hours⁶⁹. When donors release nitric oxide at higher than physiological concentrations, it is likely that non-physiological nitrosothiols are formed at less activated thiols. Therefore, it is difficult to interpret the results of proteomics experiments performed on donor treated samples⁷².

In a recent study, two nitric oxide donors (spermine NONOate and CysNO) were compared to understand their ability to form nitrosothiols⁷³. Surprisingly, spermine NONOate ($t_{1/2}$ = 39 min and 230 min) produces a high amount of dinitrosyliron complexes (DNIC) and very low amount of nitrosothiols. Conversely, Cys-NO ($t_{1/2}$ 2min)^{73,74} efficiently produced nitrosothiols. Since the majority of nitrosylation proteomics experiments rely on nitric oxide donor-treated samples, much of the current literature should be carefully interpreted. While nitric oxide donors are important tools for many experiments, it is important in the future to focus on detecting endogenous nitrosothiols.

Emerging nitrosothiol enrichment strategies

Given the limitations of the biotin-switch method, several alternative approaches have recently been reported (Figure 3A). In a series of innovative reports, Xian presents triaryl-substituted phosphines as a novel chemoselective reaction for conversion of nitrosothiols to a stable substituted thiobenzamide^{75,76}. This reductive ligation reaction mechanism is similar to the Staudinger ligation, and is initiated by nitrosothiol reaction with the phosphine to form an azaylide, which then undergoes an intramolecular reaction and hydrolysis to yield the substituted thiobenzamide adduct. Additional variants of this reaction proceed by a similar azaylide intermediate, but undergo distinct rearrangements to yield varying products. The bis-ligation reaction uses phosphine-thioester probes to form disulfide-iminophosphorane products⁷⁷. Importantly, the nitrogen originating from nitric oxide transforms to the iminophosphorane, providing an analytical linkage to both the originating

nitric oxide and thiol. This methodology has been used to quantify the formation of Snitrosoglutathione in activated macrophages by mass spectrometry, and allowed sensitive profiling of other nitrosylated metabolites in cell lysates⁷⁸. The one-step disulfide method similarly uses a phosphine-thioester to first form the thiobenzamide adduct and thiolate, followed by intermolecular thioester exchange with the released thiolates to generate a disulfide linkage⁷⁷. The reaction results in disulfide formation and elimination of the phosphine oxide. Finally, alkyl-aryl phosphines react with SNO generation of the azaylide, followed by reductive elimination to generate dehydroalanine⁷⁹. Dehydrolalanine is an electrophilic Michael acceptor, and can be linked to a variety of tagged nucleophiles for straightforward chemoselective enrichment.

In cell lysates, excess phosphine reagent led to complete reduction of nitrosothiols to free thiols⁷⁵, suggesting these methods may require careful optimization. To avoid this pitfall, free thiols were first alkylated with NEM, followed by selective nitrosothiol reduction with triaryl-phosphines. The resulting free thiols were labeled using biotin or fluorophore-linked maleimide reagents for nitrosothiol detection. In gel-based experiments, phosphine reduction demonstrated superior selectivity to dithiothreitol, which non-selectively reduced disulfides and sulfenylated thiols⁷⁵. This methodology was used in fixed cells to detect nitrosothiols after lipopolysaccharide stimulation in macrophages. Overall, these mechanistic phosphine probes show early promise as an alternative to ascorbate-dependent enrichment and proteomic analysis.

Organo-mercury enrichment methods have emerged as an alternative approach for direct nitrosothiol labeling and enrichment^{80,81} (Figure 3B). Nitroso-cysteines react directly with phenylmercury to yield a stable thiol-mercury bond. This reaction is direct, selective, and highly efficient. By coupling the phenylmercury to biotin or agarose beads, nitrosothiols can be directly labeled and enriched from tissue lysates. Bound peptides are then released from the resin by mild performic acid treatment, which oxidizes the thiol to the sulfonic acid for selective detection by mass spectrometry⁸¹. The fate of MMTS-capped thiols is not reported, although they are presumably simultaneously oxidized. This suggests MMTS should be replaced with iodoacetamide reagent to block free thiols before analysis. Furthermore, methionine is similarly oxidized by perfomic acid to the sulfone, adding additional complexity to the proteomic analysis. Nonetheless, this method lead to the identification of nearly a thousand nitrosylation sites summed across mouse brain, heart, liver, kidney, lung, and thymus tissues. Furthermore, about half or more of these sites were absent in eNOS knockout mice, suggesting the majority of nitrosylation originates from nitric oxide synthases⁸⁰. Interestingly, more than 70% of the nitrosylation sites in heart tissue were found on mitochondrial proteins⁸⁰. Further analysis suggests widespread regulation of metabolic enzymes in involved in glycolysis, gluconeogenesis, pyruvate metabolism, the Kreb's cycle, oxidative phosphorylation, amino acid metabolism, ketone body formation, and fatty acid pathways. The simplicity of this method offers several advantages over ascorbate-dependent capture methods, particularly since it is a direct enrichment followed by a unique oxidation to sulfonate for direct proteomic annotation. Furthermore, this method is the first to report robust differences by comparative proteomic analysis of eNOS knockout mice. This data provides key evidence that nitric oxide synthase activation contributes to

Conclusions

broader implications on cellular metabolism.

Protein S-nitrosylation has emerged as an important oxidative post-translational modification. Several labeling methods have emerged that take advantage of the unique reactivity, either by reduction and capture, or by direct chemical labeling. Importantly, such methodologies have enabled proteomic analysis of S-nitrosylation in cells and tissues. In conjunction with isotopic labeling methods, quantitative proteomic profiling of S-nitrosylation will enable in-depth global profiling, independent of nitric oxide donors. With the development of selective methods for each distinct cysteine post-translational modification, multiplexed analysis has the potential to discover new cellular pathways orchestrated post-translational exchanges. Such analysis will integrate palmitoylation dynamics, and explore the mutual competition at distinct cysteine residues between nitric oxide and hydrogen peroxide. Importantly, each of these modifications are labile, and rapidly hydrolyzed in the presence of free thiols, such as glutathione. Therefore, we anticipate such tools will enable a greater understanding of the role of compartmentalization in thiol modification stability and function.

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Figure 1.

Formation of nitrosothiols from nitric oxide (NO) occurs through distinct oxidative pathways^{27,82–85}, each involving two molecules of NO for each nitrosothiol formed. Superoxide radical = $O_2^{-\bullet}$. Oxygen = O_2 .



Figure 2.

Biotin-switch technique for ascorbate-dependent reduction of nitrosothiols. Enriched proteins are either analyzed by SDS-PAGE or annotated by mass spectrometry.

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Figure 3.

Chemoselective nitrosothiol labeling methods. In both approaches, free thiols are first blocked by addition of MMTS. (A) Triaryl-phosphine ligation methods. Three reactions are shown that describe recent reports of nitrosothiol-selective phosphine reactions. The reductive ligation approach was demonstrated on fixed cells, but led to over-reduction to the free thiol. The one-step disulfide formation reaction was demonstrated on cell lysates after nitric oxide donor treatment. (B) Phenyl-mercury enrichment of nitrosothiols for proteomic annotation. Sepharose beads or biotin are linked to phenyl-mercury for nitrosothiol enrichment, followed by trypsin digestion. Nitrosylated peptides are released from the resin by perfomic acid oxidation to the sulfonic acid for mass spectrometry annotation.