Biochemical methods for monitoring protein thiol redox states in biological systems

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ARTICLE INFO

Article history:
Received 16 May 2014
Received in revised form
5 June 2014
Accepted 9 June 2014
Available online 13 June 2014

Keywords:
Redox state
Disulfide
S-nitrosation
Hyperoxidation
Keap1

Methods

ABSTRACT

Oxidative post-translational modifications of proteins resulting from events that increase cellular oxidant levels play important roles in physiological and pathophysiological processes. Evaluation of alterations to protein redox states is increasingly commonplace because of methodological advances that have enabled detection, quantification and identification of such changes in cells and tissues. This mini-review provides a synopsis of biochemical methods that can be utilized to monitor the array of different oxidative and electrophilic modifications that can occur to protein thiols and can be important in the regulatory or maladaptive impact oxidants can have on biological systems. Several of the methods discussed are valuable for monitoring the redox state of established redox sensing proteins such as Keap1.

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1. Introduction

Reactive oxygen species (ROS) and reactive nitrogen species (RNS) are catch-all terms encompassing a broad range of molecular entities that have potential to chemically oxidize biological molecules. Although in common use, we should be mindful that the individual molecular species that comprise ROS and RNS can be very different in their physicochemical properties that underlie biological responses occurring when this diverse array of species change their concentration. Thus, although we often classify this broad array of molecules together, this may not always be helpful when trying to understand the molecular events that underlie biological responses to these distinct entities. Cells
utilize a diverse collection of oxidase enzymes to catalyze reduction—oxidation (redox) reactions in which electrons are passed from an electron donor source to molecular oxygen, so reducing it to form the ROS species superoxide, which can dismutate to form hydrogen peroxide (H2O2). Many contemporary studies focus on ROS generated from NADPH oxidase enzymes [1,2], as these enzymes have evolved to specifically generate superoxide that is functionally important. Whereas oxidant production by other oxidase enzymes involved in cell metabolism [3] can be secondary by-products which may not impact on protein function. ROS are also generated by uncoupled nitric oxide synthase (NOS) enzymes [4,5] and macrophages that utilize it in host defense [6], as well as by mitochondria when electrons become uncoupled from their electron transport chain and combine with molecular oxygen to generate superoxide [7,8].

An elevated level of oxidants within the cell (due to their increased synthesis or decreased antioxidant capacity that limits ROS scavenging) is often referred to as ‘oxidative stress’. To many the term oxidative stress implicitly conveys the idea that ROS simply exert a detrimental impact on biological function. However oxidants are now known to have biological functions that are not injurious, and can be considered crucial to maintenance of homeostasis or adaptive signaling events that can limit injury. These biological responses triggered by changes in cellular oxidant concentration are commonly referred to as ‘redox signaling’. In terms of oxidants causing oxidative stress, this was once considered to occur via uncontrolled oxidation of cellular biomolecules. However, another important factor in the pathogenesis of oxidant-mediated injury involves ROS dysregulating basal redox signaling pathways crucial for homeostasis, thus interfering with regulatory pathways important for the maintenance of health.

Cysteine residues are relatively uncommon in proteins compared to other amino acid, comprising only about ~2.3% of the human proteome [9]. The thiol (also known as mercaptan or sulfhydryl) -SH side chain of cysteine is susceptible to reaction with ROS or RNS species, giving rise to a range of oxidative post-translational oxidative modifications, as schematically presented in Fig. 1, that in some cases can functionally regulate the protein. At first glance this would perhaps be considered an unlikely mechanism of regulation, as an elevation in cellular ROS might be anticipated to non-selectively oxidize all manner of protein thiols, potentially triggering uncoordinated functional changes that manifest as dysfunction and development of disease. However, this concept ignores the fact that there is selectivity in the oxidative modification of protein thiols induced by ROS. This is because ROS, such as H2O2, are selective in the thiols they oxidize as a result of oxidants preferentially reacting with deprotonated (–S–), nucleophilic thiolates with a low acid dissociation constant (pKa). Most cysteines thiols however have a pKa of 8–9, and so are almost fully protonated at physiological pH, making them much less reactive with oxidants and so not susceptible to oxidative modification and regulation in this way. In addition to the protein thiol pKa, which is lowered by proximity to proton accepting amino acids (histidine, lysine, arginine) or an increase in cytosol pH, susceptibility to oxidation may be controlled by their vicinity to oxidase enzymes.

Although protein cysteine oxidation is often mechanistically rationalized by the oxidant directly reacting with the thiol, this may not always be the case. For example, although protein-tyrosine phosphatase 1B (PTP1B) is susceptible to oxidation [10], it has been questioned how this can happen in the cellular setting [11]. This is because although the target thiol in PTP1B has a pKa of ~6.8 and is clearly susceptible to oxidation, it may be difficult to reconcile this happening when peroxiredoxin (Prx) proteins with very reactive low pKa thiols (~5.5) are present in high abundance. One possibility is that PTP1B (or other targets) only become oxidized after the Prx are oxidant-inactivated by hyper-oxidation of their peroxidatic thiols [12]. In addition, it is conceivable that a low pKa protein thiol such as those in Prx or thioredoxin (Trx) may become oxidized and then react with the less reactive target protein cysteine to ‘pass on’ the oxidation [3], as depicted in Fig. 2.

Diverse arrays of oxidative modifications are crucial to redox signaling events and are integral to all manner of cellular and

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**Fig. 1.** Summary of the oxidative modifications formed in protein thiols. Protein thiols can form a variety of oxidative modifications, including reversible (intra-protein disulfides, inter-protein disulfides, S-sulfenation, S-nitrosation, S-thiolation, S-sulfhydration, S-sulfonamidation and non-reversible hyper-oxidized (S-sulfination, S-sulfonation) redox states. Some redox states, such as S-sulfenation, S-nitrosation or S-sulfhydration, can be intermediates that transition to disulfides. Prolonged exposure to oxidants can result in irreversible modifications such as S-sulfination or S-sulfonation.

**Fig. 2.** Alternate potential mechanisms leading to disulfide formation. A protein thiol may be oxidized via another redox sensitive protein, in this example thioredoxin (Trx), first becoming oxidized. Trx has a lower pKa than most other proteins and so is more likely to be preferentially oxidized by H2O2 to form an intra-molecular disulfide. The Trx disulfide is then attacked by a thiol of a second protein with a higher pKa which is then reduced by a second thiol. Trx essentially picks-up and passes on the oxidation state to the less reactive target protein thiol.
physiological processes [3,13–15]. Establishing the importance of protein redox regulation in these varied biological processes has been increasingly possible because of advances in biochemical analysis methods allowing alterations in protein thiol redox state to be monitored. These methods have also increased our knowledge of the number proteins regulated by post-translational modifications. In this mini-review we discuss contemporary biochemical methods that allow redox-sensing proteins to be studied in biological systems.

2. Monitoring reduced protein thiol status

Many methods exist for measuring the total reduced status of thiols within biological samples. Many functionalized cysteine-labeling reagents based on the selective thiol reactivity of mal- eimide, iodoacetate or disulfide moieties are commercially available. Such reagents are commonly available coupled with several different reporter functions or ‘handles’ (e.g. radiolabel, chromophore, fluorophore or affinity tag) which enable a readout of reduced thiols status [16]. With these thiol-labeling methods, a generalized rationale is that the samples under oxidative stress will have lower reduced thiol content than unstressed controls. The Ellman assay is a classic example of such an assay, providing a colorimetric readout of total reduced thiol content of a sample. As depicted in Fig. 3A, it is based on the ability of thiols in a sample to chemically reduce Ellman’s reagent 5,5’-DiThiobis-2-NitroBenzoi acid (DTNB), which possesses a reactive disulfide bond susceptible to reduction. In a stoichiometric reaction DTNB is reduced by free thiols by an exchange reaction in which a mixed disulfide and a yellow-colored 5-Thio-2-NitroBenzoic acid (TNB) is formed [17]. The intensity of yellow color (measured spectrophotometrically at 412 nm) increases proportionally with the reduced thiol content. The Ellman assay can be used quantitatively by employing external standards of cysteine or reduced glutathione (GSH), normalizing to sample protein content which allows direct comparison to results from other studies. The Ellman reagent reacts with both reduced protein and low molecular weight thiols, but trichloroacetic acid can be used to precipitate proteins to allow the latter component, which is often regarded principally as reduced GSH, to be measured independently. A major issue with the Ellman assay is its limited ability to detect subtle changes in thiol redox state, for example those alterations with physiological redox signaling. Ellman’s, as with many of the functionalized thiol-labeling reagents mentioned above, reacts with all reduced thiols in the sample. However, as also explained above, the proteins susceptible to oxidation are relatively select, typically with a low pKa. Thus most protein thiols in the sample are not anticipated to be susceptible to oxidation and so in the absence of harsh oxidant conditions or chronic oxidative stress, then the Ellman assay may not detect an alteration in thiol redox state because the net change is subtle, despite key sensor proteins becoming significantly oxidized. This issue is common to all the methods where the reactive reporter molecule reacts with thiols independently of their ionization state. One way of overcoming this issue is to use a labeling reagent, such as biotin-i-odoacetamide (BIAM), which selectively reacts with undefined thiols, namely those that are also susceptible to oxidation.

When samples are prepared at low pH (typically 6 or lower), most protein in the sample will be fully protonated as they have a pKa of 8–9 [8]. Under such conditions most reduced protein thiols will not label with BIAM as it selectively reacts with the thiolate state. However, low pKa thiols will remain at least partially deprotonated under these conditions. As oxidants also selectively target these very same low pKa reactive thiols, this low pH BIAM-labeling allows oxidant-sensitive thiols to be selectively studied.

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**Fig. 3.** Detection of the reduced (free) thiol content. (A) The Ellman assay is based on the susceptibility of a double bond in Ellman’s reagent 5,5’-dithiobis-2-nitrobenzoic acid (DTNB) to be chemically reduced by a free thiol group. The reaction products are a mixed disulfide and a yellow-colored 5-Thio-2-NitroBenzoic acid (TNB). The amount of TNB generated correlates with the reduced thiol content and can be measured spectrophotometrically. (B) Monobromobimane (mBrB) reacts with the thiols resulting in a fluorescent thiol-mBrB adduct formation which can be measured.
Another way of overcoming the inability of many global cysteine-labeling strategies to detect subtle changes in redox state is by focusing in on a single species of thiol-containing molecule. This can be achieved for low molecular weight thiols (e.g. cysteine, homocysteine, homocystine, GSH) by combining their labeling with pre- or post-chromatographic separation, typically high-performance liquid chromatography (HPLC). Samples can also be analyzed after chemical reduction to convert oxidized thiols present (e.g. cystine, homocystine, glutathione disulfide, S-thiolated proteins) to their reduced state, so enabling quantitation and more sensitive than DTNB and have not only been used for other methods which enable detection and quantitation of inter-

3. Monitoring intra-protein disulfides

Intra-protein disulfides are those formed between vicinal cysteine residues in a protein. The two thiols can be close enough to form a disulfide either by being adjacent in the primary sequence, or if not, as a result of their proximal orientation in the folded protein structure [23]. Intra-protein disulfides can induce faster migration on non-reducing SDS-PAGE in some proteins, but an additional validation study to corroborate the faster gel migration truly indicates a disulfide may be required. In addition, band shifts can be small and also are not guaranteed in many proteins of interest. However, as explained below, there are other methods which enable detection and quantitation of inter-protein disulfide formation.

The generic reductive switch-labeling method outlined above in Fig. 4 has been used to monitor intra-protein disulfide formation, (step 3) and their detection (step 4). This generalized approach is exploited by the so-called biotin-switch method, discussed below in the section on protein S-nitrosation. The blocking alkylation step is commonly performed with maleimide, N-ethylmaleimide (NEM), iodoacetamide, iodoacetate or methyl methane thiosulfonate (MMTS). The reduction step can be done with a reducing agent such as dithiothreitol (DTT) which is capable of reversing all of the oxidative modifications shown in Fig. 1, apart from the sulfenic or sulfonic states. Ascorbate and arsenite can be used to selectively reduce S-nitrosated or S-sulfonated proteins respectively. The nascent thiols generated by the reduction step can then be labeled with a thiol reactive reagents such as biotin-maleimide (as in the biotin switch method) [21,22], or polyethylene glycol-maleimide (PEG-switch method).

Fig. 4. Overview of a generic tag-based reductive switch-labeling method for monitoring reversible protein thiol oxidation. This approach requires several steps. During Step 1 free thiols are blocked with an alkylating reagent (typically maleimide, N-ethylmaleimide, iodoacetamide, iodoacetate or methyl methane thiosulfonate). During Step 2 the reversibly oxidized thiol is reduced with the reducing agent (dithiothreitol, ascorbate or arsenite depending on the oxidation states under investigation), followed by the labeling of the nascent thiols with labeling agent (typically biotin-maleimide). Step 4 involves separation (SDS-PAGE, HPLC) and identification of the labeled protein nascent thiols (typically by immunoblotting or MS analysis).
but this requires prior knowledge that the protein oxidizes to this state. Thus, Trx redox state has been monitored using this approach, utilizing 4-acetamido-4-maleimidylstilbene-2,2'-disulfonic acid (AMS) as the labeling agent (step 3). AMS is an NEM variant that reacts with reduced thiols resulting in protein-AMS adducts which increases the mass by \(-0.5\) kDa. This additional molecular weight leads to a molecular shift on SDS-PAGE, generating an additional upper band on a gel. In contrast, if a protein thiol is oxidized it will not react with AMS and so no shift will be observed.

Fig. 5. Monitoring intra-protein disulfides by AMS addition. 4-acetamido-4-maleimidylstilbene-2,2'-disulfonic acid (AMS) reacts with reduced protein thiols resulting in protein-AMS adducts which increases the mass by \(-0.5\) kDa. This additional molecular weight leads to a molecular shift on SDS-PAGE, generating an additional upper band on a gel. In contrast, if a protein thiol is oxidized it will not react with AMS and so no shift will be observed.

Fig. 6. Monitoring intra-protein disulfides by PAO and dBrB addition. (A) Phenylarsine oxide (PAO) forms a stable protein-PAO adduct after reaction with reduced vicinal thiols. However, PAO cannot react with vicinal thiols that have formed an intra-disulfide. (B) Dibromobimane (dBrB) forms a fluorescent protein-dBrB adduct with vicinal cysteines, but this cannot be formed when the protein is oxidized to form an intra-protein. Thus, loss of PAO-labeling or PAO-dependent protein capture indicates intra-protein disulfide protein.

Phenylarsine oxide (PAO) selectively adducts reduced vicinal thiols to form a stable dithioarsine ring (as shown in Fig. 6A), but not when they form an intra-disulfide. Thus solid phase PAO can be used together with western immunoblotting analysis of candidate proteins to assess their intra-protein disulfide status or with mass spectrometry (MS) approaches for unbiased proteomics.
screening to identify novel targets [16,26]. Dibromobimane (dBrB) reacts with vicinal thiols to generate a fluorescent adduct that cannot form if an intra-protein is present (Fig. 6B). This provides a basis for monitoring this oxidation state in vitro with pure proteins, or if combined with chromatographic separation could perhaps be useful in analysis of complex biological samples [26,27].

4. Monitoring inter-protein disulfides

An inter-protein disulfide refers to a bond between cysteine thiols on two protein subunits, generating either homo- or heterodimers. As inter-protein disulfide bond substantively increases the protein molecular weight, this can readily be monitored using non-reducing Western immunoblotting analysis of candidates of interest. If the band shift is normalized when the samples are separately analyzed with a reducing agent, typically DTT or 2-mercaptoethanol (2-ME) present, this provides reasonable confidence the migration difference is indeed disulfide bond-mediated [28]. Of course this approach is a candidate based-approach and requires the antibody used detects both redox states of the protein, which in our experience can be an issue with some having significant selectivity for one state over the other.

Diagonal electrophoresis is a sequential non-reducing followed by reducing gel analysis procedure that can allow the unbiased (i.e. non-candidate based) identification of proteins that form inter-protein disulfides [16,26,29]. This method, which is described in detail elsewhere [16,29,30], involves running a non-reducing SDS-PAGE to separate all proteins, including those with disulfide bonds. After electrophoresis, the entire lane containing the separated proteins is excised and placed horizontally on a second SDS-PAGE gel and separated again, but under reducing conditions by adding SDS sample buffer with DTT or 2-ME. When the gel is stained for total protein, the dominant feature is a diagonal line caused by most protein species present in the total protein, the dominant feature is a diagonal line caused by most protein species present in the total sample. This protein can then be used as the samples are together, a specific signal relative to another allows identifying both redox states of the protein. A protein microarray-based method has also developed that allows unbiased, high-throughput identification of protein thiols, as well as a resin-assisted capture method for fluorescent analysis methods where screening to identify novel targets can be multiplexed as variants of the CyDye that have unique fluorescent properties can be used to label different samples, which can then be analyzed by difference gel electrophoresis (DIGE). DIGE analysis typically involves mixing samples (that have been separately labeled with different fluorophores) together before electrophoretic separation, most often with two-dimensional sequential isoelectric focussing followed by SDS-PAGE analysis [16,52]. Combining the samples overcomes issues with related fluorescent analysis methods where samples can be compared after running them separately on different gels and looking for alterations in fluorescent spot patterns, with a loss of labeling indicating oxidation. With DIGE, as the samples are together, a specific protein species present in each sample will run precisely on top of each other as a single spot and overcome issues with using separate gels for each sample. Using ratiometric analysis and looking for loss of one fluorescent signal relative to another allows identification of a protein in a sample which is more thiol-oxidized. This protein can then be excised and identified using MS.
6. Monitoring protein S-sulfenation

Protein S-sulfenation refers to oxidation of a thiol to the sulfenic acid (−SOH, see Fig. 1) state, and can be induced by molecular oxygen or hydrogen peroxide and related species. S-sulfenation is generally considered a labile modification that like S-nitrosation will rapidly react with other thiols on contact to form more stable intermediates. Sulfenic acids can be monitored or at least observed in structural datasets using X-ray crystallography or nuclear magnetic resonance, or alternately by MS [53]. The Ellman assay described above for monitoring reduced thiols can be used indirectly to monitor sulfenates in vitro. This is achieved by monitoring consumption of an added thiol, which serves as a reporter because it reacts with the sulfenic acid to form a disulfide [54].

Generally, most of the techniques utilizing the reaction of the sulfenic acid with chemical probes are based on the electrophilic character of the sulfur atom and its weak nucleophilic properties [55]. 4-Chloro-7-nitrobenzo-2-oxa-1,3-diazole has been used to monitor sulfenation in vitro [56,57]. 5,5-Dimethyl-1,3-cyclohexane-dione (dimedone) stably derivatizes protein sulfenates (Fig. 7), which can be monitored colorimetrically or with MS in vitro with purified proteins [58,59]. However, antibodies that pan-specifically detect protein sulfenates derivatized by dimedone have been developed. This allowed protein sulfenation in cells and tissues to be monitored more easily [55,60,61]. Functionalized biotinylated or fluorescent derivatives of dimedone have also been synthesized, and have allowed protein sulfenation in cells to be studied, allowing target proteins to be captured and identified [58,59]. Another dimedone synthetic derivative, DYn-2, appears more efficient in labeling protein sulfenic acids in live cells. DYn-2, which is functionalized with a small azide, allows conjugated protein sulfenates to be captured via a Staudinger ligation click-chemistry reaction [62].

A modified biotin-switch method (see Fig. 4) can be used for the detection, purification and identification of protein sulfenic acids [21,57]. This method relies on the selective reduction of sulfenates by arsenite, with post-labeling of the nascent generated thiols with a biotinylated alkylating reagent. One potential issue with this, as with the ascorbate-dependent detection of S-nitrosated proteins, is that the analysis is carried out under SDS denaturing conditions which may result in loss of many protein sulfenates due to their destabilization during protein unfolding.

7. Monitoring protein S-sulfhydration

The disulfide bonding of H2S to a protein has been referred to as sulfhydration [63], although sulfuration is perhaps more appropriate terminology [64]. Due to the low (−2) oxidation state of H2S it is perhaps unlikely to directly react with proteins to form the disulfide. Consequently, intermediate polysulfides (and/or sulfane sulfur) have been suggested as species with oxidant properties mediating S-sulfhydration [65]. Polysulfides may be formed from H2S via autoxidation of HS− in the presence of molecular oxygen at neutral or slightly alkaline pH. S-sulfhydration could occur via the interaction of the sulfenic acid with HS− or H2S to form a persulfide bond or via a sulfenamide intermediate (reviewed in [66]). Protein S-sulfhydration may follow after H2S first forms persulfide or polysulfides, which could then undergo thiol-disulfide exchange. Perhaps the first study demonstrating protein S-sulfhydration in vivo utilized a cystathionine γ-lyase-deficient mouse model and a modified biotin switch assay [67]. A variant of the switch method shown in Fig. 4 was used to monitor S-sulfhydrated proteins, this included blocking free (unmodified) thiol groups with MMTS, with removal of unincorporated MMTS using acetone, followed by labeling of sulfhydrated (−S−SH) thiols with biotin-HPDP [67]. In the original study biotinylated proteins were captured using solid-phase avidin, eluted by SDS-PAGE and immunooblotted using biotin-conjugated antibody or targets identified using mass-spectrometry [67]. Quantification of S-sulfhydration can be achieved by comparing blots of the biotin-switch samples to the blots of total lysates (not subjected to the biotin-switch). The question however remains as how MMTS while alkylating free protein thiols does not also modify –S−SH group via thiol disulfide exchange. Recently Snyder’s group reported a modification of this method with maleimide which alkylates free thiol groups of proteins but does not affect nitrosated or other oxidized thiols. This approach allowed simultaneous measurement of sulfhydration and nitrosylation of NFκB in the same sample [68].

8. Monitoring protein S-thiolation

Protein S-thiolation is an umbrella term for disulfides between a protein and small thiol-containing molecules such as glutathione or cysteine, generating S-glutathiolated or S-cysteinylated proteins respectively. Radiolabeled glutathione or other small thiols are sensitive tools for the quantitative detection of protein S-thiolation. Tritiated GSH with quantitation of incorporation into protein by liquid scintillation counting [69], or providing 35S-cysteine in the presence of a protein synthesis inhibitor to label the GSH pool followed by autoraphy of samples resolved by SDS-PAGE [70], enabled monitoring of S-thiolation.

A commonly used non-radioactive approach is to use biotinylated glutathione ethyl ester (BioGEE), a reduced form of glutathione which is cell-permeant due to the acetyl group [70]. Once in the cell, esterases remove the acetyl group and the glutathione which is cell-permeant due to the acetyl group and the biotinylated GSH can participate in redox reactions including protein S-glutathiolation. Proteins carrying biotinylated glutathione can be detected, identified, quantified and purified following avidin-capture and MS analysis [70]. Glutathione N,N-biotinyl glutathione disulfide (biotin–GSSG–biotin) which S-glutathiolates

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**Fig. 7.** Identification of S-sulfonated proteins with dimedone. Dimedone (5,5-dimethyl-1,3-cyclohexanedione) forms a stable protein-dimedone adduct which can be identified either colorimetrically or with MS or using antibody-based approach.
proteins via thiol-disulfide exchange has also been used to study protein S-glutathiolation in cells and organs [71]. Similarly, protein S-homocysteinylation has been monitored using homocysteine labeling with a fluorescent or biotin tag [72]. Such approaches, however, do not readily permit direct detection of endogenous S-thiolated proteins, for example in stored tissue samples. In this case, antibodies to S-glutathiolated or S-homocysteinylated proteins can be utilized [71]. Candidate protein S-glutathiolation can be monitored by its immunoprecipitation, followed by immunoblotting with a pan-specific anti-glutathione antibody, as reported for detection of this modification in NOS [4].

The generic biotin-switch methods, as outlined in Fig. 4, can be adapted for selective monitoring of protein S-glutathiolation. This can be achieved on tissue sections or in homogenates by using the disulfide oxidoreductase glutaredoxin to reduce the target proteins back to the free thiol state, with subsequent labeling with a thiol-reactive functionalized (e.g. biotin, fluorophore, etc.) reagent [73,74].

9. Monitoring ‘hyperoxidation’ of protein thiols

As shown in Fig. 1, once a sulfenic acid forms it can react with additional oxidant molecules to stepwise transition into more stable sulfenic (PSO\textsubscript{H}) and then sulfonic (PSO\textsubscript{2}H) acid states. These are generally considered biologically irreversible modifications, perhaps proving evidence of damage associated with acute hyper-exposure to oxidants or chronic oxidative stress. However, the sulfination of the peroxidatic thiol in 2-Cys peroxiredoxins was shown to be reversible in cells [12], and mediated by sulfiredoxin through an ATP-dependent reaction [75]. PSO\textsubscript{2}H formation also occurs in glyceraldehyde-3-phosphate dehydrogenase (GAPDH) [76], PTP1B [10] and Dj-1 [77]. PSO\textsubscript{H} and PSO\textsubscript{2}H have been indexed by monitoring band shifts on isoelectric focussing gels [78]. Antibodies that detect sulfenic or sulfonic acid formation in peroxiredoxin [12,79], Dj-1 [80,81], or GAPDH [61] have been reported.

10. Monitoring lipid modifications of protein thiols

Protein thiols can undergo modification by a variety of lipid electrophiles, such as hydroxynonenal, malondialdehyde, acrolein, 15 d prostaglandin J\textsubscript{2} (15 D-PGJ\textsubscript{2}) or nitroalkenes via Schiff base or Michael addition reactions [82]. A general strategy involves labeling a lipid electrophile of interest, applying it exogenously to model systems to enable monitoring, purification and identification of target proteins. For example biotinylated or fluorescent BODIPY analogues of 15 D-PGJ\textsubscript{2} or PGA\textsubscript{2} have been utilized [83,84]. Biotinylated arachidonic acid can be added to cell or tissue models, after which it is metabolized endogenously to generate different products. Some of these products of arachidonic acid metabolism can be electrophiles that may adduct to proteins; thus how different stimuli modulate protein-electrophile adduction to potentially exert a functional impact can be studied [83–85]. As a label is incorporated to protein targets, this also enables subcellular localization of the modifications to be monitored using microscopy [83]. Using such approaches, 15 D-PGJ\textsubscript{2} was shown to covalently adduct to the Kelch like ECH-associated protein 1 (Keap1) in a concentration- and time-dependent manner. This modification results in activation of the nuclear factor (erythroid-derived 2)-like 2 (Nrf-2) pathways to enhance antioxidant synthesis [84]. Sulfuraphane is another electrophile that covalently modifies Keap1, so triggering elevated antioxidant synthesis. The targets of sulfuraphane have been studied by synthesizing ‘click chemistry’ analogues that enable target proteins to be monitored and identified [86,87].

Commercially available antibodies have been used extensively to monitor hydroxynonenal or malondialdehyde modification or protein with immunoblotting or cyto-imaging methods [29]. Nitroalkene modification has been monitored with a switch-type method (Fig. 4), in which the lipid adduct is reductively released from target proteins by 2-ME with monitoring by MS [88]. Protein S-palmitoylation can be similarly monitored by its enzymatic reversal using a thioesterase [89], or chemically with hydroxylamine [90]. Some modifications of proteins by lipid electrophiles result in introduction of a carbonyl moiety [82]. Protein carbonylation can be monitored via reaction with 2,4-dinitrophenylhydrazine, which can be measured colorimetrically with a spectrophotometer [91,92], by HPLC [93], or using immunoblotting with an antibody generated to this carbonyl-labeling derivatisation reagent [92,94].

11. Monitoring redox states within cellular compartments

Several of the methods described above potentially enable the redox state changes in a discreet cellular compartment (nuclei, mitochondria, endoplasmic reticulum, cytosol, etc.) to be monitored. Although this can be achieved by integrating subcellular fractionation where compatible with one of the redox analytical methods described above, a major problem is the redox state of the target proteins are readily altered, especially during lengthy sample preparations. Some of the protocols outlined are amenable to be combined with imaging methods, allowing redox state alteration in labeling to be monitored at defined subcellular locations. For example, antibodies to specific oxidation states commonly used for immunoblot analysis can also be valuable for immunocytochemistry. Cells can also be studied using a modified biotin-switch method (see Fig. 4) adapted for in situ analysis of fixed samples, again allowing the localization of the oxidized proteins to be determined. Subcellular localization of S-nitrosated proteins [95], or disulfide-containing proteins [96] has been achieved using this strategy.

Mitochondria selective thiol redox probes that combine a lipophilic triphenylphosphonium (TPP) motif with a thiol-reactive labeling moiety have been developed. The membrane potential of the mitochondrial causes the cationic, lipophilic TPP-labeled probe to selectively accumulate within these organelles where it differentially labels proteins thiols depending on their redox state [97]. The TPP motif has also been used to deliver other mitochondria-targeted cargoes, such as the antioxidants MitoVit E or MitoQ. Proteins thiols labeled with TPP derivatives can be assessed after electrophoresis using MS or antibodies developed to this targeting motif, which has allowed alteration in the mitochondrial respiratory chain complexes to be monitored [97,98]. Mitob is a mitochondria-targeted ratiometric MS probe that allows levels of oxidants such as H\textsubscript{2}O\textsubscript{2} to be measured [99].

During the last decade genetically encoded redox sensor probes that potentially allow quantitative, real-time, subcellular imaging of ROS production have been developed [100,101]. Such sensor probes typically comprise a fluorescent protein fused to a redox-sensitive protein domain that is reversibly oxidatively post-translationally modified depending on the cellular redox state. The redox state of the sensor alters the conformation of the probe protein to alter the efficiency of direct fluorescence or fluorescence resonance energy transfer which is quantified by various fluorescence-monitoring methods [100]. HyPer is a genetically encoded fluorescence probe for measuring H\textsubscript{2}O\textsubscript{2}, with enhanced HyPer 2 and HyPer 3 variant available [100,102,103]. HyPer is a yellow fluorescent protein (cpYFP) integrated into the conformation-changing region of the Escherichia coli transcription oxido-reductase.
regulatory domain (OxyR-RD), enabling it to sense H₂O₂. H₂O₂ induces an intraprotein disulfide within the HyPer probe, causing a profound conformational change that markedly alters the fluorescence emission peak at 516 nm [100]. A genetically encoded redox probe for monitoring the redox state of glutathione has also been designed by fusing a roGFP2 to human glutaredoxin-1 (roGFP2-Grx1) [104]. RexYFP, a genetically encoded redox probe for the NAD(H) redox state changes, was developed by introducing YFP into the T-REX redox sensor from Thermus aquaticus [105]. RexYFP, which is also equipped with a pH sensor to reduce pH-dependent artefacts, enables changes in the NAD+/NADH ratio in different cellular compartments to be monitored.

12. Present and future challenges

Methods and technologies for monitoring redox state in biological systems continue to emerge, driven by the intense contemporary research activity in this area. However, many of the methods are not suitable for monitoring ROS levels in cells or in tissues in vivo and so can provide only limited insight. Furthermore, many of the methods are also limited by only providing a snapshot of the redox state. Real-Time Imaging in vivo is already available to some extent, as discussed above, but we envisage these methods to evolve significantly in the near future. With the increasing prevalence of genetically engineered murine models, it is likely that such genetically encoded redox sensors will perhaps become widely utilized. For example a ‘HyPer zebrafish’ that enables in vivo monitoring of H₂O₂ has already been developed [102,106] as well as a Thy-mito-Grx1-roGFP2 transgenic mouse with a roGFP2 redox sensor expressed in neuronal mitochondria [107].

A remaining issue with methods for measuring specific ROS species, especially those allowing in vivo analysis, is that many of them are not absolutely quantitative. Quantitation is often relative to control, but the absolute concentration or amount of an oxidant produced basally or following an intervention of interest is important information that cannot readily or reliably be determined. Another issue that plagues redox analytical methods is that a steady state, net readout of the redox state of a target protein or concentration of an oxidant is provided. This misses important information about the turnover or redox cycling rates of redox active components. Monitoring of redox flux through a defined pathway of interest could again perhaps be achieved using optical sensors, but the temporal resolution of fluorescent probes currently limits this.

Acknowledgments

This work was supported by the British Heart Foundation (BHF), King’s BHF Centre of Research Excellence, Medical Research Council UK; Fondation Leducq, France, European Research Council, the Department of Health via the National Institute for Health Research comprehensive Biomedical Research Centre award to Guy’s & St. Thomas’ National Health Service Foundation Trust.

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