Coupling oxidative signals to protein phosphorylation via methionine oxidation in Arabidopsis

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The mechanisms involved in sensing oxidative signalling molecules, such as H₂O₂, in plant and animal cells are not completely understood. In the present study, we tested the postulate that oxidation of Met (methionine) to MetSO (Met sulfoxide) can couple oxidative signals to changes in protein phosphorylation. We demonstrate that when a Met residue functions as a hydrophobic recognition element within a phosphorylation motif, its oxidation can strongly inhibit peptide phosphorylation in vitro. This is shown to occur with recombinant soybean CDPKs (calcium-dependent protein kinases) and human AMPK (AMP-dependent protein kinase). To determine whether this effect may occur in vivo, we monitored the phosphorylation status of Arabidopsis leaf NR (nitrate reductase) on Ser⁵³⁴, using modification-specific antibodies. NR was a candidate protein for this mechanism because Met⁴⁰⁸, located at the P+4 position, serves as a hydrophobic recognition element for phosphorylation of Ser⁵³⁴ and its oxidation substantially inhibits phosphorylation of Ser⁵³⁴ in vitro. Two lines of evidence suggest that Met oxidation may inhibit phosphorylation of NR-Ser⁵³⁴ in vivo. First, phosphorylation of NR at the Ser⁵³⁴ site was sensitive to exogenous H₂O₂ and secondly, phosphorylation in normal darkened leaves was increased by overexpression of the cytosolic MetSO-repair enzyme PMSRA3 (peptide MetSO reductase A3). These results are consistent with the notion that oxidation of surface-exposed Met residues in kinase substrate proteins, such as NR, can inhibit the phosphorylation of nearby sites and thereby couple oxidative signals to changes in protein phosphorylation.

Key words: calcium-dependent protein kinase (CDPK), hydrogen peroxide (H₂O₂), methionine oxidation, oxidative signalling, phosphorylation motif.

INTRODUCTION

ROS (reactive oxygen species) are generated during normal aerobic metabolism by a variety of reactions, including mitochondrial and thylakoid electron transport, and enzymes such as glycolate oxidase and cellular peroxidases [1]. Numerous mechanisms exist to detoxify the most highly reactive ROS, such as superoxide, to form H₂O₂, which is relatively stable and found at significant levels in plant cells [2]. An emerging concept in both plants and animals is that ROS such as H₂O₂ play an important role in cellular signalling pathways [3,4]. In plants, H₂O₂ can alter gene expression [5,6], induce stomatal closure mediated by abscisic acid [7], and at higher concentrations can initiate programmed cell death [8]. However, it is not entirely clear how mild oxidants such as H₂O₂ are sensed at the molecular level.

Perception of H₂O₂ could involve chemical modification of amino acid side chains in cellular proteins. Cysteine residues are generally considered to be the archetypal redox-regulatory amino acid as reversible oxidation of the thiol group to form disulphides, or in some cases sulfenic acid (-SOH) or sulfenic acid (-SO₂H), is recognized to operate in redox signalling of a variety of proteins [9,10]. However, Met (methionine) residues in proteins can also be highly susceptible to modification by mild oxidants [11]. It has been demonstrated that Met oxidation occurs primarily on the surface of proteins and correlates closely with the solvent-exposed sulfur surface area [12,13]. Consequently, small changes in protein conformation influence the susceptibility of specific Met residues to oxidation, and static structural analysis can often not accurately predict the degree to which specific residues are solvent exposed [14]. Interestingly, oxidation of Met to the sulfoxide MetSO converts the side chain of this amino acid from hydrophobic to polar and hydrophilic [11,15]. This dramatic change in the chemical nature of the residue, coupled with the fact that oxidation is reversible [16,17], makes this modification of Met of potential regulatory significance. However, whether the rate of Met oxidation is rapid enough to function in redox signalling has been questioned [9]. Although there is some debate about whether the rate of Met oxidation is rapid enough to function in redox signalling, it is clear that MetSO exists in vivo. Thus, MetSO is likely to play a role in ROS responses, but perhaps other mechanisms also play a role in the early and rapid responses. Nonetheless, it is important to understand the impact of MetSO on protein function to develop a fuller understanding of ROS responses.

Much of what is known about the significance of Met oxidation has come from animal studies. While the effect of oxidation is to decrease the hydrophobicity of the residue’s side chain [15], structural changes can be induced as a result of Met oxidation that paradoxically result in an overall increase in the surface hydrophobicity of the protein [18]. Regardless of structural changes, Met oxidation can have a large impact on protein functionality, including interactions with other proteins. For example, the oxidation of specific Met residues attenuates binding of calmodulin to...
clients and is thought to play a role in downregulation of energy metabolism during stress and in altered calcium metabolism in aging animals [19]. Furthermore, a link between Met oxidation and post-translational modification of proteins involving phosphorylation has recently emerged with the demonstration that Met oxidation can alter the activities of protein phosphatases and protein kinases. The activity of calcineurin, a calcium/calmodulin-activated Ser/Thr protein phosphatase, is reduced by oxidation of Met residues in the calmodulin-binding domain of the enzyme [20], whereas CaMKII (Ca2+/calmodulin-dependent protein kinase II), a Ser/Thr protein kinase, is activated by oxidation of Met residues in the autoinhibitory domain of the enzyme [21].

We questioned whether the oxidation of Met residues in the vicinity of phosphorylation sites on the surfaces of proteins could alter recognition by the requisite protein kinase(s) and thereby serve as an additional mechanism to couple oxidative signals to changes in the post-translational modification of cellular proteins. The rationale behind this notion is that many Ser/Thr protein kinases target primary sequence motifs that involve hydrophobic residues as recognition elements. This includes the basophilic kinases, such as the CDPKs (calcium-dependent protein kinases) and SnRK1s (SNF1-related protein kinases) that are members of the same kinase superfamily [22], where a hydrophobic residue at the P – 5 position (i.e. five residues N-terminal to the phosphorylated residue) is often essential and a hydrophobic residue at P + 4 is stimulatory [23–25]. We speculated that when Met is the required hydrophobic residue, its oxidation to MetSO would inhibit phosphorylation of the target Ser/Thr residue because the sequence no longer contains the hydrophobic recognition element. To test this supposition, we initially examined the in vitro phosphorylation of synthetic peptides with Met residues at different positions. Dramatic effects of Met oxidation on phosphorylation in vitro were observed and prompted us to examine whether this might occur in vivo. To do this, we monitored the phosphorylation status of NR (nitrate reductase) at the regulatory SerThr site, which has a Met residue at the P + 4 position that serves as a hydrophobic recognition element. Observed changes in NR phosphorylation in response to: (i) exogenous H2O2, and (ii) overexpression of a cytosolic MetSO-repair enzyme (PMSRA3; peptide MetSO reductase A3) are consistent with the notion that Met oxidation may play a role in coupling oxidative signals to changes in protein phosphorylation in vivo.

**EXPERIMENTAL**

**Materials**

Synthetic peptides were produced as amides, and were purchased either from Research Genetics (Huntsville, AL, U.S.A.), Bethyl Laboratories (Montgomery, TX, U.S.A.) or GenScript (Piscataway, NJ, U.S.A.), and were > 90% pure. Activated MAPK (mitogen-activated protein kinase; no. 9101) and pSer (phospho-serine)-14-3-3-binding motif antibodies (no. 9606) and MAPK (mitogen-activated protein kinase; no. 9101) and pSer (phospho-serine)-14-3-3-binding motif antibodies (no. 9606) were purchased from Cell Signaling Technology (Danvers, MA,U.S.A.), and were purchased from Cell Signaling Technology (Danvers, MA, U.S.A.), and were purchased from Cell Signaling Technology (Danvers, MA, U.S.A.). Anti-peptide antibodies that recognize an N-terminal sequence of *Arabidopsis thaliana* NR (superscript number indicate residues in *Arabidopsis thaliana* Nia2: anti-N-terminal, RLEPPLNGVRVSYK138) and the regulatory phosphorylation site (anti-pSer534, 528SKSVpSTPFMT540) were produced by Bethyl Laboratories (Montgomery, TX, U.S.A.) and are described in [26]. Heterotrimeric AMPK (AMP-dependent protein kinase; no. PV4672) was purchased from Invitrogen (Carlsbad, CA, U.S.A.); PPKα (protein phosphatase 1α; no. 539493) was purchased from Calbiochem (San Diego, CA, U.S.A.), and soybean CDPKα and CDPKβ were expressed in *Escherichia coli* as previously described in [27].

**Plant growth**

*A. thaliana* (L.) Heynec ecotype Columbia (Col-0) was used as the wild-type. Plants for protein extraction and analysis were grown in a soil mixture with short days (8 h) in growth chambers (Conviron Model PGW36, Winnipeg, Canada) with a photosynthetic photon flux density of 100 μmol·m−2·s−1 at plant level and day/night temperature of 22/18°C. Fully expanded rosette leaves were harvested into liquid nitrogen from 3- to 4-week-old vegetative plants, as specified in the text.

**Peptide kinase assays**

The incorporation of radiolabel from [γ-32P]ATP into substrate peptides was measured using the phosphocellulose filter-binding assay. Each 40 μl reaction mixture contained recombinant CDPK in 50 mM Mops/NaOH, pH 7.5, 0.2 mM DTT (dithiothreitol), 0.2 mM CaCl2, and 4 μg of peptide as indicated. Where indicated, CaCl2 was replaced with 2 mM EGTA for determining Ca-independent kinase activity. Reactions were initiated with 0.1 mM [γ-32P]ATP (150 c.p.m.·pmol−1) plus 10 mM MgCl2, and stopped after 10 min at room temperature (22°C). Peptide assays with AMPK were performed in 25 μl reactions that contained 25 mM Hepes/NaOH, pH 7.4, 2.5 mM DTT, 10 mM MgCl2, 5 mM β-glycerophosphate, 0.5 mM EGTA, 0.01% (v/v) Triton X-100, 0.15 mM AMP, peptide as indicated and 0.1 mM [γ-32P]ATP (150 c.p.m.-pmol−1). As indicated, 1 mg/ml peptide solutions were pretreated with 150 mM H2O2 for 30 min at 25°C to oxidize Met residue(s) and then taken to dryness under vacuum before resuspending in H2O for use in experiments.

**MALDI–TOF (matrix-assisted laser-desorption ionization–time-of-flight)–MS analysis**

Peptides (10 μg/ml) were diluted 1:20 into 0.1% TFA (trifluoroacetic acid), and then mixed 1:1 with saturated HCCA (α-cyano-4-hydroxycinnamic acid). An aliquot (usually 0.3 μl) was spotted onto a GE Healthcare probe and allowed to air dry. MS analysis was performed with an Amersham Ettan™ MALDI–TOF/Pro spectrometer, operated in the linear mode.

**Immunoblot analysis**

Standard SDS/PAGE gels were run using 10 μg of total protein and transferred to PVDF. For the custom antibodies, membranes were blocked with 2% fish gelatin in PBS and then probed with the indicated primary antibody. Detection was completed by using secondary antibodies labelled with IR dyes (Molecular Probes or Rockland Immunochemicals) and scanned and quantitated using a Li-Cor Odyssey scanner and software (Li-Cor Biosciences). Commercial antibodies were used similarly and the recommended protocols were followed.

**Feeding exogenous H2O2 to Arabidopsis seedlings in liquid culture**

Approx. 50 Arabidopsis seedlings were grown in standard 8 cm plastic Petri plates using half-strength Murashige and Skoog salts supplemented with 0.25% sucrose for 12–14 days in 16 h light, 22°C. At 2 h into the photoperiod, the seedlings were moved to the dark for 1 h, at which time the indicated concentration of H2O2 was added. Treatments were continued in the dark for an additional 60 min, and then the seedlings were harvested, quickly rinsed in distilled water and frozen in liquid nitrogen. Protein extracts were completed using direct extracts in SDS sample buffer.
CDPK on blot treatment

Standard SDS/PAGE gels were run using 10 μg of total protein and transferred to PVDF membranes. The membranes were blocked with 2% BSA in TBST (20 mM Tris/HCl, pH 7.5, 140 mM NaCl and 0.1% Tween 20) and then transferred to kinase buffer (50 mM Mops, 0.2 mM CaCl₂ and 10 mM MgCl₂). Recombinant Soybean CDPKβ was added at 15 μg/ml in kinase buffer containing 50% glycerol, 100 mM NaCl, 2.5 mM DTT and 0.2 mM ATP for 1 h at room temperature. Control reactions were run in parallel, but without ATP. After treatment, the standard procedure for immunoblots was completed.

RESULTS AND DISCUSSION

Methionine oxidation in peptide substrates can block phosphorylation in vitro

Recombinant CDPKs were tested for the ability to phosphorylate synthetic peptide substrates that in most cases contained a single Met residue that was either reduced (untreated) or oxidized by preincubation with H₂O₂ prior to the phosphorylation reaction. Initial experiments used the SP49M peptide, which corresponds to the Ser⁴⁶⁰ regulatory site of spinach SPS (sucrose-phosphate synthase) [28] and fits the canonical motif targeted by CDPKs and SnRK1 with hydrophobic recognition elements at the P−5 and P+4 positions. The sequence surrounding spinach SPS Ser⁴⁶⁰ contains Met residues at both of these positions as well as the P+3 position. However, the SP49M peptide sequence was modified to contain norleucine (J), a non-oxidizable homologue of Met, at the P+3 and +4 positions. As expected, the reduced form of the SP49M peptide was readily phosphorylated by CDPKβ, as monitored by an increase in the peptide molecular mass of 80 units using linear mode MALDI–TOF-MS analysis (Figures 1A and 1B). Preincubation of the SP49M peptide with 150 mM H₂O₂ increased the molecular mass by 16 units, corresponding to oxidation of the single Met residue at the P−5 position (designated SP49Mox; Figure 1C). Unlike the reduced peptide, SP49Mox was a poor substrate for CDPKβ, as after 10 min of incubation, there was no increase in peptide mass suggestive of phosphorylation (Figure 1D). It should be noted that H₂O₂ was effectively removed following the peptide treatment, and thus direct inhibition of CDPK is not the basis for the effect observed. In a separate experiment, the impact of Met oxidation on phosphorylation of the SP49M peptide was confirmed using the more quantitative analysis of peptide kinase activity with [γ-³²P]ATP (Table 1, motif A). Thus, oxidation of Met at the P−5 position in a canonical CDPK motif can strongly inhibit phosphorylation by soybean CDPKβ. The SP49 peptide, which contains three Met residues, was a better substrate for CDPKβ than SP49M and, as expected, oxidation of the Met residues resulted in nearly complete inhibition of peptide phosphorylation (97% inhibition; Table 1, motif A). The Ser⁵³⁴ regulatory phosphorylation site of Arabidopsis NADH:NR [23,29,30] also conforms to the canonical motif and has a Met residue at the P+4 position, which is stimulatory but not essential for phosphorylation by CDPK and SnRK[1,23]. As expected, phosphorylation of a synthetic peptide based on the Ser⁵³⁴ site of NR (the SS34 peptide, Table 1, motif A) was substantially reduced when the Met residue was oxidized, which is consistent with the known stimulatory role of a hydrophobic residue at this position.

The recombinant CDPKβ will also readily phosphorylate certain peptides that do not match the canonical motif. For example, a peptide based on Ser⁴⁵ of the Ca²⁺-ATPase, ACA2 [31], can be phosphorylated by CDPKs in vitro but lacks a basic residue at the P−3 position (See Table 1, motif B). The ACA2 motif is defined by basic residues at P−6 (and beyond) and P+2 and with a hydrophobic residue at P−5 [32]. Similar to the peptides conforming to the canonical motif, the ACA2A-5 peptide was a good substrate when the Met residue was reduced, but a very poor substrate when the Met at P−5 was oxidized (Table 1, motif B).

The second non-canonical motif identified for CDPKs was based on the Ser⁶⁰ regulatory phosphorylation site of tomato ACS (1-amino-cyclopropane-1-carboxylate synthase) 2 [33]. The sequence around Ser⁶⁰ does not conform to the canonical motif for CDPK substrates, because it lacks both the basic residue at P−3 and the hydrophobic residue at P−5, but nonetheless was shown to be efficiently phosphorylated by crude tomato extracts [33] and also recombinant CDPKs [27]. Analysis of peptide variants defined the ACS motif as Φ-Basic-Φ-Ser/
Table 1 Oxidation of Met residues in substrate peptides inhibits phosphorylation in vitro by soybean CDPKβ

| Motif | Peptide       | Sequence           | Kinase activity with peptide (c.p.m. × 10⁻³ incorporated) | % Inhibition |
|-------|---------------|--------------------|-----------------------------------------------------------|--------------|
|       | Reduced       | Oxidized           |                                             |              |
| (A) ϕ-B-x-B-x-x-{S/T}-x-x-x-ϕ | SP49 | KGRMQRISSEVMK       | 51.3 ± 3.4                                         | 1.4 ± 0.1  | 97          |
|       |               |                    | SP49M                                      | 21.9 ± 0.6  | 2.2 ± 0.3  | 90          |
|       |               |                    | NR-S34                                     | 212 ± 2.0   | 85.4 ± 0.4 | 60          |
| (B) B-B-x-B-x-(4) -S/T/-x-B | ACA2M-5 | RRFMTANLSKR       | 161 ± 8.1                                         | 9.6 ± 1.4   | 94          |
| (C) ϕ-B-ϕ-{S/T}-ϕ-x-B-B | ACSM-3 | KNNRLSFGRM       | 39.1 ± 0.3                                         | 40.4 ± 0.7  | (3)         |
|       | ACSM-1        | NNLMRSFKR         | 106.4 ± 4.6                                         | 68.2 ± 2.8  | 36          |
|       | ACSM+1        | NNMLRSFKR         | 47.0 ± 2.5                                         | 5.3 ± 0.1   | 89          |
|       | ACSM-3        | NNLMRSFKR         | 96.3 ± 3.1                                         | 33.8 ± 0.7  | 67          |

Thr-Φ-X-Basic-Basic, where Φ is a hydrophobic residue and X is any amino acid [27]. The ACS3 peptide (KKNLRLSFGRM) is based on the tomato ACS2 protein sequence and contains a single Met residue at the P + 5 position (underlined); however, there is no recognized requirement for a hydrophilic residue at this position. Accordingly, treatment of the ACS3 peptide with H₂O₂ oxidized the single Met residue and increased the molecular mass by 16 Da to form the ACS3ox peptide, which was readily phosphorylated as monitored qualitatively by MALDI–TOF-MS (results not shown) or quantitatively by radiolabel incorporation (Table 1, motif C). Thus oxidation of Met residues at positions that are not essential as hydrophilic recognition elements does not inhibit phosphorylation. These results also indicate that the mild oxidant was effectively removed from the peptide preparations during processing, so that there was no H₂O₂ remaining that could directly inhibit the protein kinase.

CDPKs will also phosphorylate a shorter 10-residue peptide based on the non-canonical ACS motif [27]. Variants of the native peptide sequence (NNLRLSFGRM) were produced with a single Met substituted at each of the three positions (P − 3, −1 and +1; underlined in the sequence) thought to be important for hydrophilic interactions with the protein kinase. All three of the Met-containing peptides were readily phosphorylated by CDPKβ when the Met was reduced, and in each case, oxidation of the Met residue resulted in substantial inhibition of phosphorylation (Table 1, motif C). Complete conversion of Met to MetSO was verified for each peptide by MALDI–TOF-MS analysis (results for the ACSM-1 peptide are shown in Supplementary Figure S1 at http://www.BiochemJ.org/bj/442/bj44220305add.htm), but the impact on peptide phosphorylation clearly varied with position, with MetSO at the P + 1 position having the greatest effect and the P − 1 position having the least effect. It is important to note that the relative importance of the hydrophilic residues at the three different positions was not established in the original report. Rather, it was simply determined that combined substitution of alanine residues at all three positions prevented phosphorylation [27]. Consistently, oxidation of a single Met residue at each of the three positions had a significant effect on peptide phosphorylation.

Collectively, the results suggest that Met residues must be in the reduced form in order to serve as hydrophilic recognition elements in phosphorylation motifs. This requirement is documented with three distinct motifs (canonical and non-canonical) and two protein kinases: CDPKβ (Table 1) and CDPKα (Supplementary Table S1 at http://www.BiochemJ.org/bj/442/bj4220305add.htm). This effect can also be extended to positions not recognized as hydrophilic recognition elements and to another distinct protein kinase, AMPK. One well known physiological target of human AMPK is Ser60 of ACC1 (acetyl-CoA carboxylase 1) [34], which contains a Met residue at the P − 1 position. A 15-residue synthetic peptide based on this site (LHIRSSMSGLHLY) conforms to the canonical motif and is a good substrate for human AMPK and soybean CDPKβ (Supplementary Figure S2 at http://www.BiochemJ.org/bj/442/bj44220305add.htm). Oxidation of the Met at the P − 1 position strongly inhibited phosphorylation by both kinases, suggesting that while a hydrophilic residue at this position is not essential, a large polar residue is not well tolerated. Hence, the impact of Met oxidation at a specified position is motif-specific and can have a significant effect even at positions not thought to serve as hydrophilic recognition elements.

In principle, the use of phosphate containing oxidized Met residues could be the result of a lack of peptide binding to the protein kinase or an inability of the kinase to phosphorylate the oxidized peptide. In order to distinguish between these two possibilities, increasing amounts of the SP49Mox peptide were added to reaction mixtures containing the untreated (reduced) SP49M peptide as substrate. As shown in figure 2, the oxidized peptide does not inhibit phosphorylation of the reduced SP49M peptide by CDPKβ.

Increasing amounts of the SP49Mox peptide (KGR[Met]SO[RISSE]VEJK) were added to complete phosphorylation reaction mixtures containing the SP49M peptide (KGR[RISSE]VEJK). Total incorporation of [32P] was monitored with the SP49M peptide alone (white bar) or with an equivalent amount or 2-fold excess of the SP49Mox peptide (grey bars), or with the SP49Mox peptide alone (black bars). Values are means ± S.E.M. (n = 3) from a representative experiment.
Methionine oxidation and protein phosphorylation

Figure 3 Effect of exogenous H₂O₂ on the phosphorylation status of cellular proteins

Liquid cultured Arabidopsis seedlings were treated with various concentrations of H₂O₂ for 1 h in the dark. (A) MAPK activation as monitored by immunoblotting with activation loop antibodies. (B) Integrated signal for ProQ Diamond phosphoprotein staining of cellular proteins greater than 50 kDa. (C) Relative phosphorylation of NR at the Ser⁵³⁴ site, expressed as the pSer⁵³⁴/NR ratio, where NR is total enzyme protein detected with an antibody directed against the N-terminus of NR. (D) NR protein content by immunoblotting. At 2 h into the photoperiod, the seedlings were moved to the dark for 1 h, at which time the indicated concentration of H₂O₂ was added. Treatments were completed in the dark, and after 1 h the seedlings were harvested. Values shown are means ± S.E.M of three biological replicates, and within each panel values are normalized to the 0 mM H₂O₂ control, except in (A), which was normalized to the 5 mM H₂O₂ treatment.

Figure 2, addition of the SP49Mox peptide to reaction mixtures containing the SP49M peptide resulted in a slight increase in total [³²P] incorporation equivalent to the activity supported by the oxidized peptide alone. The clear lack of competition suggests that the oxidized peptide is not recognized and does not bind to the kinase active site.

In contrast, dephosphorylation of phosphorylated SP49M by lambda protein phosphatase or recombinant PP1α of rabbit skeletal muscle was relatively unaffected by oxidation of the Met residue at the P−5 position. In these experiments, MALDI–TOF-MS was used to monitor the dephosphorylation of the synthetic phosphopeptides (designated phospho-SP49M or phospho-SP49Mox) during a 30 min incubation with the recombinant protein phosphatases. As shown in Supplementary Figure S3 (at http://www.BiochemJ.org/bj/422/bj4220305add.htm), the progress curves for dephosphorylation catalysed by PPlα were generally similar for the phosphorylated reduced and oxidized peptides, and quite distinct from the large effect Met oxidation had on peptide phosphorylation. Similar results were obtained with lambda protein phosphatase (results not shown). Collectively, these results suggest that peptidyl Met oxidation may potentially have more impact on phosphorylation rather than dephosphorylation, but this certainly needs to be studied further.

Effect of oxidative signals on phosphorylation of NR in vivo

In order to determine whether Met oxidation affects protein phosphorylation in vivo, we focused on NR, which has a Met residue at the P+4 position that, when oxidized, can reduce phosphorylation of the regulatory Ser⁵³⁴ site (Table 1, motif A). The phosphorylation status of NR at the regulatory site can be monitored using modification-specific antibodies (anti-pSer⁵³⁴ antibodies), and total NR protein can be monitored using antibodies directed against an N-terminal sequence (anti-NT antibodies) [26]. Initially, we examined the effect of exogenous H₂O₂ on NR phosphorylation in Arabidopsis seedlings grown in liquid culture [35]. As a marker for ROS signalling, we used the activation of MAPKs, which is well known to respond to exogenous H₂O₂ [36–38]. In our system, MPK6 [38] activation was the most pronounced and was maximal with 5 mM H₂O₂. At higher concentrations of H₂O₂, MPK6 activation was reduced slightly and only decreased appreciably at 100 mM H₂O₂ (Figure 3A and Supplementary Figure S4A). Because the exposure time to H₂O₂ was relatively short (60 min), there was no apparent phytotoxicity or cell death in our seedlings, which can occur in response to high concentrations of H₂O₂ (50 to 100 mM) but requires longer periods of exposure. For example, infiltration of 70 mM H₂O₂ into Arabidopsis leaves resulted in cell death after 7 days [39], and treatment of the more sensitive suspension cells with 50 mM H₂O₂ produced little cell death within the first 3 h [40]. Accordingly, the seedlings in our system remained fully viable as evidenced by maintenance of phosphorylation of a subset of the plant proteome (arbitrarily set as proteins greater than 50 kDa; Figure 3B and Supplementary Figure S4D). In marked contrast, NR-Ser⁵³⁴ phosphorylation was increased by 5 mM H₂O₂ and then progressively decreased at higher concentrations (Figure 3C and Supplementary Figure S4B). Undoubtedly, many factors contribute to these changes in NR phosphorylation, with increased cytosolic [Ca²⁺] in response to exogenous H₂O₂ [41], potentially increasing phosphorylation of NR by CDPKs, whereas oxidation of NR-Met⁵³⁸ would be expected to cause.
inhibition of phosphorylation. The observed changes in relative NR phosphorylation (Figure 3C) are consistent with the notion that low concentrations of H₂O₂ are sufficient to trigger a rise in cytosolic [Ca²⁺], whereas higher concentrations are required for protein oxidation. It is also interesting to note that exogenous H₂O₂ increased total NR protein in the seedlings (Figure 3D and Supplementary Figure S4C). The steady-state level of Nia2 transcript is only increased 1.11-fold by H₂O₂ (20 mM for 1 h; Genevestigator, http://www.genevestigator.ethz.ch/gv), and so increased translation and/or decreased proteolytic degradation of NR protein may be responsible for the rapid changes in content observed. The upregulation of NR protein even at the highest concentration of H₂O₂ is also consistent with the notion that the seedlings were completely viable under the conditions tested.

The concentrations of exogenous H₂O₂ supplied to intact seedlings in the present study (up to 100 mM) are certainly in excess of reported values for the leaf content of H₂O₂, which range from roughly 0.5 to 5 μmol·g⁻¹ fresh weight for different species (corresponding to 0.5 to 5 mM if distributed uniformly) [2,42]. The extent to which the internal H₂O₂ concentration was increased in our experiments is not known, but tissue barriers to uptake coupled with various mechanisms to remove H₂O₂ almost certainly reduced the internal concentration to well below the external concentration. Consequently, we applied exogenous H₂O₂ over a concentration range that was empirically determined to enhance ROS signalling but without loss of tissue viability. Hence, the results obtained should be physiologically relevant.

We developed an on-blot CDPK phosphorylation assay to determine whether Met oxidation might contribute to the reduced phosphorylation of NR-Ser⁵³⁴ in vivo in response to exogenous H₂O₂. In the experiment presented in Figure 4, exogenous H₂O₂ resulted in a progressive inhibition of NR phosphorylation (Figure 4A). Three samples of NR, differing in phosphorylation status at the Ser⁵³⁴ site, were then compared as substrates for CDPK phosphorylation on the PVDF membrane. The results are presented as the relative increase in the pSer⁵³⁴ signal in response to incubation with recombinant CDPKβ (Figure 4B). The sample prepared from seedlings exposed to 5 mM H₂O₂ had an initially high phosphorylation status and, as expected, no increase in phosphorylation was observed when incubated with CDPKβ. In contrast, the light control sample (obtained before seedlings were darkened) had an initial pSer⁵³⁴/NR ratio of approx. 0.35 and was rapidly phosphorylated in the on-blot assay. However, the extract from seedlings treated with 100 mM H₂O₂ also had a low pSer⁵³⁴/NR ratio and was phosphorylated on the blot, but at a lower rate compared with the light control sample. The difference in rate of on-blot phosphorylation of these two samples is consistent with the notion that oxidation of NR-Met⁵³⁸ occurred in vivo in response to exogenous H₂O₂ and contributed to the reduction in phosphorylation at the Ser⁵³⁴ site.

Another approach to determine whether Met oxidation affects NR phosphorylation in vivo was undertaken with plants overexpressing the MetSO-repair enzyme PMSRA3. A. thaliana MSRA3 encodes a cytosolic enzyme involved in the repair of S-MetSO diastereoisomers, and plants overexpressing this gene have increased tolerance to acute oxidative stress (treatment with methyl viologen) [43]. As shown in Figure 5(A), the relative phosphorylation status of NR at the Ser⁵³⁴ site was greater in darkened leaves compared with illuminated leaves [26,44], but the striking result was that overexpression of PMSRA3 increased NR-Ser⁵³⁴ phosphorylation in the dark, which is consistent with the notion that Met oxidation may inhibit NR phosphorylation in vivo. The increased relative phosphorylation of NR was also associated with increased absolute levels of NR protein in the transgenic plants relative to wild-type (Figure 5B). These results suggest that under normal dark conditions Met oxidation can attenuate NR phosphorylation.

The anti-pSer⁵³⁴ antibodies used in the experiments presented in Figures 4 and 5 were produced using the phosphopeptide SLKKSvpSTPFMNT as the antigen [26], raising the possibility that oxidation of Met⁵³⁸ may directly interfere with antibody cross-reactivity. In that case, a reduced immunoblot signal may not be the result of decreased phosphorylation but rather oxidation of the nearby Met residue. However, this seems unlikely for several reasons. First, pre-absorption of antibodies on a column with an immunobilized phosphopeptide corresponding to the C-terminal portion of the antigen peptide (SvpSTPFMNT) did not reduce the intensity of immunoblot signals, suggesting that the Met residue lies outside of the epitope (results not shown). Second, treatment of a blot with 1% H₂O₂ to oxidize Met residues on immobilized proteins had no effect on detection of total NR protein on the blot (using the anti-NT antibodies), whereas the anti-pSer⁵³⁴ immunoblot signal was increased slightly (results not shown). That inhibition was not observed strongly suggests that Met
oxidation is not affecting our ability to monitor phosphorylation of Ser$^{534}$. Finally, equivalent results for phosphorylation of NR at the Ser$^{534}$ site were obtained with our antibodies compared with commercial antibodies against the module 1 14-3-3 binding motif ([KR]-x-x-pSer-x-P; where x is any amino acid) that does not include the Met residue (Supplementary Figure S5 at http://www.BiochemJ.org/bj/422/bj4220305add.htm). Hence, reduced phosphorylation of NR-Ser$^{534}$ at high exogenous H$_2$O$_2$ concentrations cannot be attributed to failure of the antibodies to bind to the phosphorylated Ser$^{534}$ sequence when the nearby Met is oxidized.

Concluding remarks

The most important result of the present study is the demonstration that Met oxidation can act as a redox switch coupling oxidative signals to protein phosphorylation. This is shown to occur with synthetic peptides phosphorylated in vitro with CDPKs and human AMPK, and we suggest may apply broadly to protein kinases that target motifs with essential hydrophobic recognition elements. Identifying proteins where this mechanism may be important in vivo is a challenge for future research. Because phosphorylation sites are typically on the surface of proteins, Met residues in the vicinity are also likely to be solvent accessible and therefore susceptible to oxidation. We are speculating that the propensity for oxidation of Met residues has been exploited in Nature to serve as a redox switch to regulate phosphorylation of nearby sites. Identifying proteins with conserved Met residues in the vicinity of known phosphorylation sites is certainly one approach to catalogue possible candidate proteins that might be regulated in this manner.

We provide evidence in the present study that phosphorylation of NR may be regulated by oxidative signals involving the Met redox switch. The regulatory Ser$^{534}$ phosphorylation site on NR controls binding of a 14-3-3 protein, to form a catalytically inactive complex (NR-pSer$^{534}$–14-3-3) that modulates NR activity in response to light and other environmental signals [45,46]. The phosphorylation site occurs in a region of the NR molecule known as hinge 1, which contains −60 amino acids that connect the molybdenum cofactor domain and cytochrome $b_5$ domain [47]. Based on limited structural information [48], this region is suggested to be surface exposed, which would certainly be necessary to allow phosphorylation and binding of a 14-3-3 protein, and also suggests that Met$^{538}$ would be solvent exposed and therefore susceptible to oxidation. NR phosphorylation is sensitive to oxidative signals and two lines of evidence are consistent with the notion that Met$^{538}$ oxidation may play a role. Certainly more work needs to be done to confirm and extend these results, but NR appears to be a good candidate protein to examine the role of Met oxidation as a redox switch controlling phosphorylation. It is possible that this mechanism may be involved in the activation of NR in vivo in response to hypoxia/anoxia [49], which paradoxically involves increased ROS production [50] and is beneficial for plant tolerance to root flooding [51]. This work just begins to explore the role Met oxidation plays in modulating protein phosphorylation of both plant and animal systems and future work will probably uncover many additional aspects in which the Met oxidation redox switch plays critical roles.

AUTHOR CONTRIBUTION

Shane Hardin, Clayton Larue, Man-Ho Oh and Vanita Jain performed research and analysed data. Steven Huber analysed data and wrote the paper.

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241665–1672

25405–410

26489–493

274155–167

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292165–2172

302805–28057

31293–298

32432–438

3328051–28057

34315–319

35307–313

36330–334

3728–29

38449–456

3921685–1703

40281–285

41205–212

42203–207

43103–110

441019–1029

4572–82

46407–412

47353–356

4838697–38704

49135–140

502335–339

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SUPPLEMENTARY ONLINE DATA

Coupling oxidative signals to protein phosphorylation via methionine oxidation in Arabidopsis

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Figure S1 Met oxidation of the ACSM-1 synthetic peptide partially inhibits phosphorylation by recombinant soybean CDPKβ in vitro

Peptides were analysed by linear mode MALDI–TOF-MS analysis. (A) The ACSM-1 peptide (NNLRM$\text{S}$FGKR, where the phosphorylated serine residue is in bold) at zero time or (B) after 10 min incubation with CDPKβ and ATP. Note the increase in mass of 80 units corresponding to single site phosphorylation. (C) Oxidation of the ACSM-1 peptide by treatment with 0.5% H$_2$O$_2$ followed by lyophilization to remove the peroxide; note the increase in mass of 16 units corresponding to conversion of Met to MetSO in the ACSM-1ox peptide. (D) The ACSM-1ox peptide after incubation with CDPKβ and ATP for 10 min showing only partial conversion to the phosphorylated form.

Figure S2 The oxidized form of the ACC1-Ser\textsuperscript{80} peptide is a poor substrate for human AMPK and soybean CDPKβ in vitro

Human AMPK or soybean CDPKβ were used to phosphorylate the untreated or oxidized ACC1-Ser\textsuperscript{80} peptide in vitro in the [\textsuperscript{32}P]ATP peptide kinase assay.

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Peptides were analysed by linear mode MALDI–TOF–MS analysis. (A) The phospho-SP49M peptide (KGRMRRISVEJJK, where J is norleucine and the phosphorylated serine residue is in bold) at zero time and after a 30 min incubation with PP1α. Note the decrease in mass of 80 units corresponding to single site dephosphorylation. (B) Oxidation of the phospho-SP49M peptide by treatment with 0.5 % H2O2 followed by lyophilization to remove the peroxide; note the increase in mass of 16 units corresponding to conversion of Met to MetSO in the phospho-SP49Mox peptide and subsequent effective dephosphorylation by PP1α. (C) Time course for dephosphorylation of the phospho-SP49M peptide (■) and phospho-SP49Mox peptide (○) by PP1α. Reactions contained 50 mM Hepes, pH 7.5, 100 mM NaCl, 2 mM DTT, 0.01 % Brij 35, 0.2 mM MnCl2 and peptide as indicated, and were initiated by the addition of 2.5 units of PP1α. At the times indicated, aliquots were removed, diluted 10-fold with 0.11 % TFA (trifluoroacetic acid), prior to mixing 1:1 with DHB (2,5-dihydroxybenzoic acid) matrix in 50 % acetonitrile containing 2 % phosphoric acid. Aliquots (0.3 μl) were spotted onto the MALDI target and linear spectra were acquired. Appearance of the dephosphorylated peptide was quantified by monitoring the product peak height, which is plotted as a percentage of the maximum achieved after a 30 min incubation. Equivalent amounts of the dephosphorylated forms of the oxidized and reduced peptides were produced during the experiment.
Table S1  Oxidation of Met residues in substrate peptides inhibits phosphorylation in vitro by soybean CDPKα

Residues at essential recognition positions are shown in italics, the phosphorylated residue is underlined, and Met residues are in bold; ϕ is a hydrophobic residue, B is a basic residue and x is any amino acid. Values are means ± S.E.M. (n = 3) from a representative experiment.

| Motif | Peptide | Sequence | Kinase activity with peptide (c.p.m. × 10⁻³ incorporated) |
|-------|---------|----------|----------------------------------------------------------|
|       |         |          | Reduced | Oxidized | % Inhibition |
| (A) ϕ-x-B-x-x-[S/T]-x-x-x-ϕ | SP49 | KGRMRRRSSVEMMK | 6.66 ± 0.6 | 0.32 ± 0.07 | 95 |
|       | SP49M | KGRMRRRSSVEJKK | 6.18 ± 0.6 | 0.43 ± 0.07 | 93 |
| (B) B-B-x-B-ϕ-x(4)-[S/T]-x-B | ACA2M-5 | RRFAMTANLSKRY | 28.2 ± 2.6 | 2.1 ± 0.22 | 93 |
| (C) ϕ-B-ϕ-[S/T]-ϕ-x-B-B | ACS-3 | KKNLRLSFGRKMY | 18.2 ± 1.3 | 22.3 ± 1.8 | (23) |
|       | ACSM-1 | NNLGMSFGKR | 24.6 ± 1.6 | 10.9 ± 0.7 | 56 |
|       | ACSM-1+1 | NNLRLSMGKR | 8.4 ± 1.0 | 1.3 ± 0.3 | 85 |
|       | ACSM-3 | NMRCSFGKR | 23.6 ± 0.9 | 4.5 ± 0.2 | 81 |

Figure S4  Representative immunoblots showing the effect of exogenous H₂O₂ on (A) MPK6 activation; (B) NR phosphorylation at the Ser⁵³⁴ site; (C) NR protein; and (D) ProQ Diamond phosphoprotein staining of cellular proteins.

The results shown are one of three independent biological replicates that were averaged to generate Figure 3 of the main text.

Figure S5  Met⁵³⁸ oxidation does not affect immunochemical detection of phosphorylation at the Ser⁵³⁴ site by the custom anti-NR-pSer⁵³⁴ antibodies

Blots containing proteins extracted from Arabidopsis seedlings in liquid culture treated with exogenous H₂O₂ were separately probed with custom anti-NR-pSer⁵³⁴ antibodies (pSS34 Ab) or commercial antibodies (Cell Signaling Technology) that recognize the mode 1 binding site for 14-3-3 proteins, [KR]-x-x-pS-x-P. Both antibodies detected phosphorylated NR in an equivalent manner, confirming that changes in apparent phosphorylation of Ser⁵³⁴ could not be attributed to oxidation of the nearby Met⁵³⁸ residue.