# Repairing oxidized proteins in the bacterial envelope using respiratory chain electrons

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The reactive species of oxygen and chlorine damage cellular components, potentially leading to cell death. In proteins, the sulfur-containing amino acid methionine is converted to methionine sulfoxide, which can cause a loss of biological activity. To rescue proteins with methionine sulfoxide residues, living cells express methionine sulfoxide reductases (Msrs) in most subcellular compartments, including the cytosol, mitochondria and chloroplasts<sup>1–3</sup>. Here we report the identification of an enzymatic system, MsrPQ, repairing proteins containing methionine sulfoxide in the bacterial cell envelope, a compartment particularly exposed to the reactive species of oxygen and chlorine generated by the host defence mechanisms. MsrP, a molybdo-enzyme, and MsrQ, a haem-binding membrane protein, are widely conserved throughout Gram-negative bacteria, including major human pathogens. MsrPQ synthesis is induced by hypochlorous acid, a powerful antimicrobial released by neutrophils. Consistently, MsrPQ is essential for the maintenance of envelope integrity under bleach stress, rescuing a wide series of structurally unrelated periplasmic proteins from methionine oxidation, including the primary periplasmic chaperone SurA. For this activity, MsrPQ uses electrons from the respiratory chain, which represents a novel mechanism to import reducing equivalents into the bacterial cell envelope. A remarkable feature of MsrPQ is its capacity to reduce both rectus (R-) and sinister (S-) diastereoisomers of methionine sulfoxide, making this oxidoreductase complex functionally different from previously identified Msrs. The discovery that a large class of bacteria contain a single, non-stereospecific enzymatic complex fully protecting methionine residues from oxidation should prompt a search for similar systems in eukaryotic subcellular oxidizing compartments, including the endoplasmic reticulum.

The fact that no Msr had been identified in the cell envelope of important human pathogens, including Escherichia coli and Pseudomonas aeruginosa, was surprising as this compartment is particularly exposed to the oxidizing compounds present in the environment. We postulated that such a methionine sulfoxide (Met-O) reducing system had remained unidentified, and applied a genetic approach to uncover it, using E. coli as a model. We first constructed an E. coli Met auxotroph mutant lacking all cytoplasmic Msrs and found this strain (JB590) to be unable to use Met-O as the only Met source (Fig. 1a). We then searched for suppressor mutations conferring Met-O reducing capacity to JB590, which led to the isolation of strain BE100 (Fig. 1a). Genetic analysis of the suppressor revealed the presence of an insertion sequence element (IS2) within *yedV*, a gene coding for the histidine kinase of the uncharacterized YedV/YedW two-component system<sup>4</sup>. In close vicinity were two genes, *yedY* and *yedZ*, encoding, respectively, a periplasmic molybdopterin-containing oxidoreductase and its putative membrane redox partner<sup>5,6</sup>. YedY had been shown to reduce a variety of substrates *in vitro*, including trimethylamine *N*-oxide, and dimethyl, methionine and tetramethylene sulfoxides<sup>5</sup>. However, its physiological function had remained elusive, although a recent study in *Azospira suillum* suggested the homologous protein to



Figure 1 | The MsrPQ system reduces free Met-O and is induced by HOCl. a, JB590, a methionine auxotroph (Met<sup>-</sup>, numbered '1'), lacking all cytoplasmic Msrs (Met<sup>-</sup> Msr<sup>-</sup>, '2'), cannot grow on Met-O as the only Met source in contrast to suppressor BE100 (Met<sup>-</sup> Msr<sup>-</sup> Sup<sup>Met-O<sup>+</sup></sup>, '3'). Deletion of *yedY* (renamed *msrP*, '7'), *moeA* ('6') and *tatC* ('5'), but not of *trxA* ('4'), prevents the growth of BE100. **b**, The *yedYZ* operon is upregulated in BE100. The increase is observed both at the mRNA (quantitative PCR with reverse transcription (RT-qPCR), top; error bars, mean  $\pm$  s.e.m.; n = 3) and protein (western blot, bottom) levels. The RT-qPCR primers were designed to quantify the *yedY-yedZ* mRNA. **c**, Immunoblot analysis showing that HOCl (2 mM), but not H<sub>2</sub>O<sub>2</sub> (1 mM), induces YedY synthesis in a wild-type strain. Images in **a**-**c** are representative of experiments made in biological triplicate. Uncropped blots are in Supplementary Fig. 1.

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Figure 2 | MsrP non-stereospecifically reduces protein-bound Met-O. a, Oxidation of Met in calmodulin (CaM) by H<sub>2</sub>O<sub>2</sub> leads to a mobility shift of the oxidized protein (CaMox); compare lane 2 with lane 1. Incubation of CaMox with MsrP and a reducing system involving dithionite and benzyl viologen restores the mobility (lane 3). b, MsrP can reduce Met-O in CaMox. The oxidation state of peptides containing either Met36, Met109, Met124 or Met145-146 was determined by LC-MS/MS. Error bars, mean  $\pm$  s.e.m.; n = 4 for Met36, 109 and 145–146; n = 5 for Met124. Met-O residues were detected in the untreated and MsrP-treated samples owing to limitations inherent to the methodology applied and oxidation of the samples during analytical handling. c, Representation of the two diastereoisomers of Met-O, R-Met-O and S-Met-O. d, MsrP exhibits activity towards both diastereoisomers (left), contrary to the stereospecific enzymes MsrA and MsrB (right). Specific activities were assayed using 64 mM of either *R*- or S-Met-O. Error bars, mean  $\pm$  s.d.; n = 3. e, The suppressor BE100 is able to grow on both isoforms of Met-O. Images in a and e are representative of experiments made in biological triplicate. The uncropped gel is in Supplementary Fig. 2.

be important for hypochlorous acid (HOCl) resistance<sup>7</sup>. We found that insertion of the IS2 led to a 100-fold increase in the levels of the *yedYZ* messenger RNA (mRNA) in strain BE100 and to higher YedY protein levels (Fig. 1b). Deletion of either *yedY* or *yedZ* prevented BE100 from growing on Met-O (Fig. 1a and Extended Data Table 1), while the simultaneous overproduction of YedY and YedZ, but not of YedY or YedZ alone, rendered the parental strain JB590 able to use Met-O (Extended Data Table 1). Altogether, these results indicated that the ability of the suppressor strain BE100 to reduce Met-O resulted from the increased synthesis of YedY and YedZ, implying that these two proteins function together as an Msr system. Growth of the BE100 strain was dependent on moeA, a gene required for the synthesis of molybdopterin cofactors, and on *tatC*, encoding a protein required for the translocation of metalloenzymes across the inner membrane (Fig. 1a). Exposure of wild-type cells to HOCl, but not to  $H_2O_2$ , induced the synthesis of YedY to levels comparable to those observed in BE100 (Fig. 1c), indicating that these proteins are specifically expressed in response to bleach stress. Interestingly, induction by HOCl was dependent on the presence of a functional YedV/YedW system (Extended Data Fig. 1).

All previously identified Msrs rely on electrons derived from NADPH via the thioredoxin (Trx) system for activity<sup>1</sup>. This was not the case for YedYZ, as deletion of *trxA*, encoding the Trx responsible for Msr recycling<sup>8</sup>, had no effect on the ability of BE100 to reduce

Met-O (Fig. 1a). As YedZ contains a *b*-type haem<sup>9</sup>, a cofactor typically associated with the quinone-oxidizing cytochrome *b* of the respiratory chain complexes, we considered the respiratory chain as a potential electron source. Deletion of *menA* and *ubiE*, two genes required for quinone synthesis, prevented BE100 from using Met-O (Extended Data Table 1), supporting a model in which YedZ uses electrons derived from the inner membrane pool of mature quinones to provide reducing equivalents to YedY (Extended Data Fig. 2). From now on, YedY and YedZ will be referred to as MsrP (for periplasm) and MsrQ (for quinone), respectively.

We then tested whether MsrPQ, in addition to free Met-O, also rescued Met-O residues present in proteins. Purified MsrP was shown to reduce N-acetyl-Met-O, a substrate mimicking protein-bound Met-O (Extended Data Fig. 3a), with a Michaelis constant  $(K_m)$  of  $3.8 \pm 1.2$  mM, in line with the values reported for other Msrs<sup>10</sup>. Note that in the experiments involving purified MsrP, electrons were provided to the oxidoreductase by an inorganic system reducing molybdoenzymes<sup>11</sup>. Next, we tested the ability of MsrP to reduce oxidized calmodulin (CaMox), a substrate commonly used to assess Msr activity. We used a gel shift assay based on the reduced mobility exhibited in SDS-polyacrylamide gel electrophoresis (SDS-PAGE) by proteins containing Met-O<sup>12</sup>. Incubation of CaMox with MsrP restored its mobility, suggesting that MsrP was able to reduce Met-O residues in CaMox (Fig. 2a). This was confirmed by showing with liquid chromatography-tandem mass spectrometry (LC-MS/MS) that the oxidized Met residues that could be detected in CaMox were reduced back to levels similar to those observed in CaM after incubation with MsrP (Fig. 2b). Altogether, these results indicated that MsrP is able to reduce protein-bound Met-O.

Upon oxidation, two diastereoisomers of Met-O can form, referred to as *R* and *S*, owing to the asymmetric position of the oxidized sulfur atom in the lateral chain (Fig. 2c). All Msrs described so far exhibit stereospecificity, specifically reducing either the *R* (MsrB, MsrC) or the *S* isoform (MsrA, BisC). Using highly pure diastereoisomers (Extended Data Fig. 4), we found MsrP to exhibit activity towards both (Fig. 2d), with  $K_m$  values of  $25.7 \pm 4.7$  mM and  $8.0 \pm 2.7$  mM for *R*- and *S*-Met-O, respectively (Extended Data Fig. 3b). Accordingly, the BE100 suppressor strain was able to use *R*- and *S*-Met-O (Fig. 2e), in contrast to strains expressing single stereospecific Msrs (Extended Data Fig. 3c). Thus, MsrP is a new type of Msr with no stereospecificity.

To search for the physiological substrates of MsrP, periplasmic proteins from a  $\Delta msrP$  mutant were oxidized with HOCl, incubated with MsrP and subjected to a semi-quantitative two-dimensional LC–MS/MS analysis. Twenty proteins that had one or more HOCl-oxidized Met residues that MsrP could reduce were identified (Extended Data Table 2). Using gel shift assays in combination with LC–MS/MS analysis, we confirmed the ability of MsrP to reduce the chaperone SurA and the lipoprotein Pal (Fig. 3a, b). Altogether, these results established that MsrP is able to repair a wide panel of structurally and functionally diverse periplasmic proteins *in vitro*.

SurA is the primary periplasmic chaperone, escorting most  $\beta$ -barrel proteins to the outer membrane<sup>13,14</sup>. As HOCl-oxidized SurA loses its chaperone activity (Fig. 4a), we used this property to probe the physiological importance of the MsrPQ system. First, we showed that SurA could be oxidized *in vivo* by HOCl and that expression of the MsrPQ system, but not of MsrP alone, restored its mobility (Fig. 3c). Similar results were obtained for Pal (Fig. 3d), confirming that MsrP and MsrQ collaborate in the protection of SurA and Pal from oxidative damage. We then tested if the repair of SurA by MsrP, which restores the activity of the chaperone *in vitro* (Fig. 4a), was important to keep SurA active under HOCl stress. For this, we used a mutant strain lacking the chaperone Skp, in which SurA becomes essential<sup>15,16</sup>. We found that deleting *msrP* rendered the  $\Delta skp$  strain hypersensitive to HOCl (Fig. 4b), suggesting that oxidized, inactive SurA accumulates in the absence of MsrP. In agreement with this, the sensitivity of the  $\Delta skp \Delta msrP$ 



Figure 3 | The MsrPQ system rescues oxidized Met residues in SurA and Pal. a, b, Oxidation of SurA (SurA ox) and Pal (Pal ox) by H<sub>2</sub>O<sub>2</sub> leads to a mobility shift resulting from Met-O formation. Incubation with MsrP and the inorganic reducing system restores their mobility (top). The percentages of Met-O in the various samples were determined by LC-MS/MS analysis, confirming that MsrP reduces Met-O in SurA and Pal (bottom). Error bars, mean  $\pm$  s.e.m.; n = 3. Met-O residues were detected in the untreated and MsrP-treated samples owing to limitations inherent to the methodology applied and oxidation of the samples during analytical handling. c, d,  $\Delta msrPQ$  cells carrying msrP either alone or with *msrQ* under an isopropyl  $\beta$ -D-1-thiogalactopyranoside (IPTG)inducible promoter on a plasmid (pAG192 and pAG195, respectively) were grown with IPTG (100  $\mu$ M). Cells were treated with chloramphenicol  $(300\,\mu g\,ml^{-1})$  at an absorbance  $(A_{600\,nm})$  of 0.5 to block new protein synthesis and HOCl (3.5 mM) was added. Synthesis of MsrP and MsrQ together (top), but not of MsrP alone (bottom), restores SurA and Pal mobility. Images in **a**-**d** are representative of experiments made in biological triplicate. The small shift exhibited by SurA over time in the absence of MsrPQ could be due to a residual Msr activity, possibly an NADPH-dependent membrane-bound Msr activity previously detected<sup>21</sup>. Uncropped gels and blots are in Supplementary Fig. 3.

mutant to HOCl was suppressed by overexpression of SurA (Fig. 4c). Further highlighting the need to protect Met residues in periplasmic proteins, HOCl-pretreated  $\Delta msrP$  mutants were found to be more sensitive to SDS, a phenotype indicative of defects in the outer membrane (Fig. 4d)<sup>17</sup>.

The conservation of MsrPQ throughout Gram-negative bacteria (Extended Data Figs 5 and 6) illustrates the importance of having a Met-O reducing system in the periplasm. *Neisseria* species stand out as an exception in lacking MsrPQ. However, in these bacteria, evolutionary tinkering generated an envelope hybrid protein combining two classic stereospecific Msr domains<sup>18</sup>. A remarkable feature of MsrPQ is that its rescue activity depends on electrons provided by the respiratory chain. This represents an entirely novel way to provide reducing power for protein quality control in the envelope. Indeed, known reducing systems functioning in the periplasm use electrons provided by the inner membrane protein DsbD and Trx<sup>19</sup>. Hence, diverting electrons from the respiratory chain to control extracytosolic protein quality is an unprecedented link between metabolism and cellular integrity.

The chaperone SurA is one of the targets of the MsrPQ system. Having a protein folding helper under the control of a repair system reveals an additional layer in the complex control network of periplasmic protein quality. Testing if this system is an attractive target for antimicrobial development, as suggested by the colonization defect exhibited by the *msrP* mutant in *Campylobacter jejuni*<sup>20</sup>, will be the



Figure 4 | The reducing activity of the MsrPQ system is important for envelope integrity. a, Repair of oxidized SurA (SurA ox) by MsrP (SurA ox + MsrP) restores the ability of SurA to protect thermally unfolded citrate synthase from aggregation. The graph is representative of experiments made in biological triplicate (a.u., arbitrary units). b, While the wild-type (WT), the  $\Delta skp$  and the  $\Delta msrP$  strains are only moderately affected by exposure to HOCl (2 mM), the viability of the  $\Delta skp \Delta msrP$ mutant (in which SurA is essential) is decreased. Error bars, mean  $\pm$  s.e.m.; n=3. c, The sensitivity of the  $\Delta skp \Delta msrP$  mutant to HOCl is suppressed by SurA overexpression. Error bars, mean  $\pm$  s.e.m.; n=4. d, Pre-treatment with HOCl renders the  $\Delta msrP$  mutant hypersensitive to SDS, indicative of envelope defects. Error bars, mean  $\pm$  s.e.m.; n=3.

subject of future research. By highlighting the importance of protecting proteins targeted to oxidizing compartments, our work calls for a detailed investigation of the process of Met-O reduction in the endoplasmic reticulum, where only an *R*-Met-O-specific MsrB has been identified<sup>3</sup>. As has long been speculated, a possibility would be that the endoplasmic reticulum contains an epimerase catalysing the interconversion of *R*- and S-Met-O. Alternatively, in light of the present study, the endoplasmic reticulum could contain a novel Met-O reducing system yet to be discovered.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to F.B. (barras@imm.cnrs.fr) or J.-F.C. (jfcollet@uclouvain.be).

#### **METHODS**

No statistical methods were used to predetermine sample size. The experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment.

Strains and microbial techniques. The strains used in this study are listed in Supplementary Table 1. Unless otherwise specified, for all deletion mutants, the corresponding alleles from the Keio collection<sup>22</sup> were transferred into the MC4100 wild-type strain using P1 transduction standard procedures<sup>23</sup> and checked by PCR. To excise the resistance cassette, we used pCP20 (refs 22, 24). Strain AG227, deleted for the entire yedYZ operon, was constructed as follows. First, a cat-sacB cassette, encoding chloramphenicol acetyl transferase and SacB, a protein conferring sensitivity to sucrose, was amplified from strain CH1990 using primers yedYZ::cat-sacB\_Fw and yedYZ::cat-sacB\_Rv. The resulting PCR product shared a 40-base-pair (bp) homology to the 5' untranslated region of yedY (msrP) and to the 3' untranslated region of yedZ (msrQ) at its 5' and 3' ends, respectively. After purification, the PCR product was transformed by electroporation into CH1940. These cells harbour the pSIM5-tet vector, which encodes the Red recombination system proteins Gam, Beta and Exo under the control of the temperature-sensitive repressor cI859, encoded by the same vector. Induction of the Gam, Beta and Exo proteins was induced by shifting the cells to 42 °C for 15 min before making them electrocompetent. Recombinant cells were selected on chloramphenicolcontaining plates (25µg ml<sup>-1</sup>) at 37 °C for 16 h. At this temperature, the pSIM5-tet vector, which has a temperature-sensitive origin of replication, is lost. Colonies were also tested for the presence of the *cat-sacB* cassette by negative selection on sucrose-containing media (5% sucrose, no NaCl). Finally, we verified that the cat-sacB cassette replaced the msrPQ operon in the resulting strain (AG219) by sequencing across the junctions. The *cat-sacB* cassette was subsequently moved from AG219 to TP1004 by P1 transduction. The cat-sacB cassette was eliminated from the resulting strain (AG220) by transforming it with the pSIM5-tet plasmid, electroporating it with the oligonucleotide Delta\_yedYZ (300 ng) and performing lambda red recombination as described above. Recombinants were selected on sucrose-containing media at 30 °C for 16 h. To eliminate the plasmid, the selected colonies were grown at 37 °C for 16 h. Loss of the cassette in the resulting AG227 strain was verified by positive (sucrose resistance) and negative (chloramphenicol sensitivity) selection and by PCR.

The *msrQ* deletion mutant (strain BE105) was generated using the PCR knockout method developed in ref. 24. Briefly, a DNA fragment containing the *cat* gene flanked with the homologous sequences found upstream and downstream of the *yedZ* gene was PCR-amplified using pKD3 as template and the oligonucleotides P1\_Up\_YedZ and P2\_Down\_YedZ. Strain BE100, carrying plasmid pKD46, was then transformed by electroporation with the amplified linear fragment. Chloramphenicol-resistant clones were selected and verified by PCR.

The *msrP::lacZ* fusion was constructed using the method described in ref. 25. Briefly, the *msrP* promoter region lying between nucleotide -797 and nucleotide +63, using the A nucleotide within the initiation triplet as a reference, was amplified by PCR with the appropriate oligonucleotides (*lacI-msrP*<sub>forward</sub> and *lacZmsrP'*<sub>reverse</sub>). Using mini-lambda-mediated recombineering, the PCR product was then directly recombined with the chromosome of a modified *E. coli* wild-type strain (PM1205), carrying a P<sub>BAD</sub>-*cat-sacB* cassette inserted in front of *lacZ*, at the ninth codon. Recombinants were selected for loss of the *cat-sacB* genes, resulting in the translational fusion of *msrP* to *lacZ*.

Plasmid construction. The plasmids and primers used in this study are listed in Supplementary Tables 2 and 3, respectively. The YedY-His<sub>6</sub> (MsrP-His<sub>6</sub>) expression vector was constructed as follows. Site-directed mutagenesis using primers pTAC\_NdeI\_Fw and pTAC\_NdeI\_Rv was performed using pTAC-MAT-Tag-2 as template to introduce an NdeI restriction site in the vector, yielding vector pAG177. yedY (msrP) DNA was amplified from the chromosome (MC4100) using primers pTAC\_yedY\_Fw and pTAC\_yedY-His6\_Rv, which resulted in the fusion of a His6 tag coding sequence at the 3' end. The PCR product was subsequently cloned into pAG177 using NdeI and BglII restriction sites, generating plasmid pAG178. To construct IPTG-inducible pTAC-MAT-Tag-2 vectors expressing either MsrP (without tag) or both MsrP and MsrQ, we first amplified the corresponding coding DNA sequences (msrP or the msrPQ operon) from the chromosome of strain MC4100 using primer pairs pTAC\_yedY\_Fw/ pTAC\_yedY\_Rv and pTAC\_yedY\_Fw/ pTAC\_yedZ\_Rv, respectively. The PCR products were then cloned into pAG177 using restriction sites NdeI and BglII, yielding pAG192 (MsrP) and pAG195 (MsrPQ). The complementation pAM238 vectors constitutively expressing either MsrP or MsrQ alone (without tag) or both MsrP and MsrQ were constructed as follows. We first amplified the corresponding coding DNA sequences (msrP, msrQ or the msrPQ locus) in addition to a 50 bp upstream region from each start codon (to include a ribosomal binding site) from the chromosome of strain MG1655 using primer pairs pAM238\_yedY\_Fw/ pAM238\_yedY\_Rv, pAM238\_yedZ\_Fw/ pAM238\_yedZ\_Rv and pAM238\_yedY\_Fw/ pAM238\_yedZ\_Rv, respectively.

The PCR products were then cloned into pAM238 using restriction sites KpnI and PstI, yielding pAG264 (MsrP), pAG275 (MsrQ), and pAG265 (MsrPQ).

The vector allowing the arabinose-inducible expression of SurA was constructed as follows. The *surA*-encoding DNA and its 50 bp upstream region (to include a ribosomal binding site) were amplified from the chromosome of strain MG1655 using the primer pair *surA\_Fw/surA\_Rv*. The PCR product was then cloned into pBAD33 using restriction sites KpnI and XbaI, yielding vector pAG290.

Analysis of the yedYZ operon expression by RT-qPCR. Expression levels of the yedYZ (msrPQ) mRNA were assessed in M63 minimal medium supplemented with 0.5% glycerol, 0.15% casamino acids, 1 mM MgSO<sub>4</sub>, 1 mM MoNa<sub>2</sub>O<sub>4</sub>,  $17 \, \mu M \ Fe_2(SO_4)_3$  and vitamins (thiamine  $10 \, \mu g \, ml^{-1}$ , biotin  $1 \, \mu g \, ml^{-1}$ , riboflavin  $10 \mu g \,ml^{-1}$  and nicotinamide  $10 \mu g \,ml^{-1}$ ). Overnight cultures of MG1655 were diluted to  $A_{600\,\text{nm}} = 0.04$  in fresh M63 minimal medium (100 ml) and cultured aerobically at 37 °C until  $A_{600 \text{ nm}} = 0.8$ . Cells (10 ml) were then pelleted, resuspended in TriPure (Roche) and homogenized. After mixing with chloroform, RNA was isolated by centrifugation (15 min, 15,700g, 4 °C), precipitated with isopropanol, washed with ethanol 70%, dried and finally resuspended in DEPC water. Any residual DNA was eliminated by treatment of the sample with DNase (Turbo DNA-free Kit, Ambion). A RevertAid RT kit (Thermo Scientific) was used to generate complementary DNA (cDNA) from 1 µg RNA extracted from each of the cultured strains. cDNAs were then diluted 1/10 and submitted to qPCR, using a qPCR Core kit for SYBR Green I No ROX (Eurogentec) and a MyiQ Single-Colour Real-Time PCR Detection System (Bio-Rad). Expression levels of yedYZ were normalized to the expression of gapA. Primers used for qPCR analysis were qPCR\_yedYZ\_Fw and qPCR\_yedYZ\_Rv for yedYZ, and qPCR\_gapA\_Fw and qPCR\_gapA\_Rv for gapA (Supplementary Table 3).

**Immunoblot analysis of MsrP expression.** Synthesis of MsrP in strains JB590 and BE100 was assessed as follows. Overnight cultures were diluted to  $A_{600 \text{ nm}} = 0.04$  in fresh M63 minimal medium (100 ml) and cultured aerobically at 37 °C until  $A_{600 \text{ nm}} = 0.8$ . Nine hundred microlitres of each culture were then precipitated with 10% ice-cold trichloroacetic acid (TCA), pellets were washed with ice-cold acetone, dried, resuspended and heated at 95 °C in Laemmli SDS sample buffer (SB buffer) (2% SDS, 10% glycerol, 60 mM Tris-HCl, pH 7.4, 0.01% bromophenol blue), and loaded on an SDS–PAGE gel for immunoblot analysis. The protein amounts loaded were standardized by taking into account the  $A_{600 \text{ nm}}$  values of the cultures.

To monitor the MsrP expression levels after NaOCl or  $H_2O_2$  treatment, overnight cultures of wild-type cells (MG1655) were diluted to  $A_{600 nm} = 0.04$  in fresh lysogeny broth (LB) medium (100 ml) and grown aerobically at 37 °C to  $A_{600 nm} = 0.5$ . NaOCl (2 mM) or  $H_2O_2$  (1 mM) was then added to the cultures. Samples were TCA-precipitated, washed with ice-cold acetone, dried, suspended in SB buffer, heated at 95 °C and loaded on an SDS–PAGE gel for immunoblot analysis. The protein amounts loaded were standardized by taking into account the  $A_{600 nm}$  values of the cultures. The specificity of the anti-MsrP antibody was verified (Supplementary Fig. 5).

Preparation of pure diastereoisomeric forms of Met-O. L-Methionine sulfoxide  $([\alpha]_D^{24} = +14.3^{\circ} \text{ (water)})$ , triethylamine (>99%) and methanol (>99.6%) were obtained from Sigma-Aldrich, picric acid from Prolabo and D2O from SDS. Water was purified using Millipore Elix Essential 3 apparatus. <sup>1</sup>H and <sup>13</sup>C NMR were recorded on a Bruker Avance III Nanobay spectrometer (1H: 400 MHz; {1H}13C: 100 MHz). Chemical shifts ( $\delta$ ) were referenced to dioxane (<sup>1</sup>H:  $\delta$  = 3.75 p.p.m.;  $^{13}\text{C}: \delta = 67.19 \,\text{p.p.m.})^{26}$  , which was added as an internal reference; resonances are detailed as follows: <sup>1</sup>H,  $\delta$  in parts per million (multiplicity, *J*-coupling in hertz, integration, signal attribution);  ${}^{1}H{}^{13}C$ ,  $\delta$  in parts per million (signal attribution). For each diastereoisomer, chemical shifts are similar to those previously reported<sup>27</sup>. <sup>13</sup>C resonance assignments were confirmed by heteronuclear single quantum coherence experiments. Optical rotations were measured on an Anton Paar Modular Circular Polarimeter 200 instrument at 25 °C and 589 nm from aqueous solution containing 0.8-1.2 g per 100 ml of L-methionine sulfoxide. The values reported are the average and s.d. relative to three independent measurements recorded on distinct solutions.

The commercial mixture of diastereoisomers was separated following the previously reported method<sup>28</sup>. Briefly, 10 ml of water was added to L-methionine sulfoxide (1.333 g, 8.069 mmol) and picric acid (1.849 g, 8.071 mmol). The suspension was heated to reflux until complete dissolution and then slowly cooled to room temperature (~25 °C). The suspension was filtered on a sintered funnel and the solid was washed with cold water (10 ml in total). Both the solid (*dextro*) and filtrate (*levo*) were collected separately for further purification.

*Dextro*. To the dried solid, 20 ml of water were added and the mixture was heated to reflux then allowed to cool slowly to room temperature. The solid was filtered out, washed with 10 ml water and dried. Again, 11 ml of methanol were added to the resulting solid and the mixture heated to reflux. After slow cooling, the yellow crystals were filtered, washed with 5 ml methanol and dried. A portion was used for structure determination by X-ray analysis. To the dextrogyre picrate

salt (1.345 g, 3.42 mmol),  $\sim$ 1.1 equivalents of triethylamine were added as a dilute aqueous solution (22 ml, 175 mM, 3.85 mmol). Subsequently, 200 ml of acetone were added portion-wise to the above stirring suspension and a white solid precipitated. This was filtered, washed, triturated with acetone and finally dried in vacuum (533 mg, 80%).

*Levo.* The volume of the filtrate was reduced in vacuum at 40 °C to about 3–4 ml to obtain a saturated solution and a small amount of precipitate. Then, 1.5 ml of water were added, the suspension was filtered and the solid washed with minimal water (2 ml). The whole step was repeated once (reduce the volume, dilute, filter and wash), and the resulting solution was then completely dried in vacuum. To the resulting yellow residue, 15 ml of methanol were added and the suspension was heated to reflux. In our hands, no solid precipitated upon cooling (in contrast with the reported method<sup>28</sup>); therefore the solution was dried again in vacuum. Following the same protocol as before, to the levogyre-enriched picrate salt (1.354 g, 3.44 mmol), ~1.1 equivalents of triethylamine were added as a concentrated aqueous solution (3.8 ml, 1 M, 3.8 mmol). Afterwards, 200 ml of acetone were added portion-wise and a white solid precipitated. This was filtered, washed, triturated with acetone and finally dried in vacuum (515 mg, 77%).

*Dextro* (L-methionine-*S*-sulfoxide):  $[α]_D^{25} = +99.2 \pm 1.5^\circ$  (water); <sup>1</sup>H NMR (400 MHz, D<sub>2</sub>O pD = 6.5): 3.88 (t, <sup>3</sup>*J* = 6.3, 1H, Hα<sub>S</sub>), 3.02 (m, 2H, Hγ<sub>S</sub>), 2.74 (s, 3H, Hε<sub>S</sub>), 2.31 (dd, *J* = 14.4, 7.6, 2H, Hβ<sub>S</sub>); {<sup>1</sup>H}<sup>13</sup>C NMR (100 MHz, D<sub>2</sub>O): 173.8 (COO<sub>S</sub>), 54.0 (Cα<sub>S</sub>), 48.9 (Cγ<sub>S</sub>), 37.2 (Cε<sub>S</sub>), 24.4 (Cβ<sub>S</sub>). Literature values from ref. 28:  $[α]_D^{25} = +99^\circ$  (water), from ref. 27:  $[α]_D = +98.2^\circ$  (water, room temperature); <sup>1</sup>H NMR (300 MHz, D<sub>2</sub>O): 4.10 (m, 1H), 3.08–2.78 (m, 2H), 2.59 (s, 3H), 2.32–2.13 (m, 2H); <sup>13</sup>C NMR (75 MHz, D<sub>2</sub>O): 171.1, 52.0, 48.3, 37.0, 23.5.

 $\begin{array}{l} \label{eq:loss} Levo~(L-methionine-R-sulfoxide): $\left[\alpha\right]_D{}^{25} = -72.7 \pm 0.5^{\circ}~(water); {}^{1}H~NMR~(400~MHz, D_2O~pD = 6.5): 3.86~(t, {}^{3}J = 6.3, 1H, H\alpha_R), 3.12~(ddd, J = 13.4, 9.6, 7.0, 1H, H\gamma_{R1}~or~H\gamma_{R2}), 3.02~(m, 2H, H\gamma_S), 2.93~(ddd, J = 13.5, 9.1, 6.8, 1H, H\gamma_{R1}~or~H\gamma_{R2}), 2.74~(s, 3H, H\epsilon_R), 2.31~(m, 2H, H\beta_R); {}^{1}H{}^{13}C~NMR~(100~MHz, D_2O): 173.9~(COO_R), 54.2~(C\alpha_R), 54.0~(C\alpha_S), 48.9~(C\gamma_R), 37.2~(C\epsilon_S), 37.0~(C\epsilon_R), 24.4~(C\beta_R). Literature values from ref. 28: $\left[\alpha\right]_D{}^{26} = -71.6^{\circ}~(water), from ref. 27: $\left[\alpha\right]_D = -78^{\circ}~(water, room temperature); {}^{1}H~NMR~(300~MHz, D_2O): 4.10~(m, 1H), 3.08-2.78~(m, 2H), 2.59~(s, 3H), 2.32-2.13~(m, 2H); {}^{13}C~NMR~(75~MHz, D_2O): 171.1, 52.1, 48.4, 37.0, 23.7. \\ \end{array}$ 

In the <sup>1</sup>H NMR spectra, the resonance centred at 3.02 p.p.m. was attributed to the *S*- enantiomer. The relative integral values suggest that *R*-Met-O is contaminated by 3% of the *S*- diastereoisomer. Moreover, comparing the measured  $[\alpha]_D^{25}$  values with those reported in ref. 27, the data are consistent with the presence of 3% *S*- diastereoisomer as a contaminant. Such purity is in line with previous reports using the same separation method<sup>28,29</sup>. The absolute configuration of the L-methionine-*S*-sulfoxide was confirmed by X-ray structural analysis and matches previous assignments<sup>27,30</sup>.

**Synthesis of N-acetyl-Met-O.** To synthesize N-acetyl-Met-O, Met-O (30 mg; Sigma-Aldrich) was solubilized in 2 ml 100% acetic acid. After addition of 2 ml of 97% acetic anhydride, the resulting mixture was incubated 2 h at 23 °C. Then, 2 ml of water were added and the mixture was lyophilized overnight. Finally, the lyophilized N-acetyl-Met-O was washed three times with 6 ml of water, re-lyophilized and suspended in 500 mM Na<sub>2</sub>HPO<sub>4</sub>, pH 9.0 to a final concentration of 1.5 M. The pH was then adjusted to 7 with NaOH.

Kinetic analysis of MsrP activity. The MsrP reductase activity was followed spectrophotometrically at 600 nm by monitoring the substrate-dependent oxidation of reduced benzyl viologen, serving as an electron donor. Reactions were performed anaerobically at 30°C in degassed and nitrogen-flushed 50 mM MOPS, pH 7.0 using stoppered cuvettes. Benzyl viologen was used at a final concentration of 0.4 mM (molar extinction coefficient,  $\varepsilon$ , of reduced benzyl viologen = 7,800 M<sup>-1</sup> cm<sup>-1</sup>) and reduced with sodium dithionite. The final reaction volume was kept constant, with the ordered addition of benzyl viologen, sodium dithionite, 1–32 mM *N*-acetyl-methionine sulfoxide (NacMet-O) and 10 nM MsrP-His<sub>6</sub>. The concentrations used for the *R*- and *S*-Met-O diastereoisomers were 1–64 mM. The Michaelis–Menten parameters (maximum velocity ( $V_{max}$ ) and  $K_m$ ) were determined using Graphpad Prism software.

Analysis of MsrA and MsrB activities. The reductase activities of MsrA and MsrB were followed spectrophotometrically at 340 nm by monitoring the substrate-dependent oxidation of NADPH ( $\varepsilon = 6,220 \, M^{-1} \, cm^{-1}$ ). Reactions were performed at 37 °C in HEPES–KOH 20 mM, pH 7.4, NaCl 10 mM, and the final reaction volumes were kept constant, with the ordered addition of 250 $\mu$ M NADPH (Roche), 2.6 $\mu$ M TrxR, 40 $\mu$ M Trx, 64 mM substrate and 1.5 $\mu$ M of either MsrA or MsrB.

Identification of the periplasmic proteins repaired by MsrP using twodimensional LC–MS/MS. The identification of the MsrP substrates was performed as follows. AG89 cells (2L) were grown aerobically at 37 °C in terrific broth to  $A_{600\,nm} = 0.8$ . Periplasmic extracts were prepared as described previously<sup>31</sup>. Briefly, cells were pelleted by centrifugation at 3,000g for 20 min at 4 °C and incubated on ice with gentle shaking for 30 min in 100 mM Tris-HCl, pH 8.0, 20% sucrose, 1 mM EDTA. This mixture also contained 20 mM *N*-ethylmaleimide to alkylate reduced cysteine residues in proteins to prevent their subsequent oxidation. Periplasmic proteins were then isolated by centrifugation of the cells at 3,000g for 20 min at 4 °C. The periplasmic extract was subsequently concentrated by ultrafiltration in an Amicon cell (3,000 Da cutoff, YM-3 membrane) and loaded on a PD-10 column (GE Healthcare) equilibrated with 50 mM NAPi, pH 8.0, 50 mM NACl. After concentration using a 5 kDa cutoff Vivaspin 4 (Sartorius) concentrator, the extract was finally separated in three samples. Two samples were incubated 10 min at 37 °C with 2 mM NaOCl whereas the third was left untreated to serve as reduced control. NaOCl was then removed by gel filtration using a NAP-5 column (GE Healthcare) equilibrated with 50 mM MOPS, pH 7.0. The untreated sample was also subjected to the NAP-5 gel filtration.

One of the NaOCl-oxidized fractions was then reduced in vitro by incubation for 1 h at 37 °C with 10  $\mu M$  MsrP, 10 mM benzyl viologen and an excess of sodium dithionite. The other NaOCl-oxidized fraction, used as an oxidized control, and the non-oxidized fraction were incubated with 10 mM benzyl viologen and an excess of sodium dithionite but without MsrP. The three samples were then de-salted by dialysis against 50 mM MOPS, pH 7.0 by using Slide-A-Lyzer 3,500 MWCO G2 cassettes (Thermo Scientific). The three samples (500µg) were precipitated by adding TCA to a final concentration of 10% w/v. The resulting pellets were washed with ice-cold acetone, dried in a Speedvac, suspended in 0.1 M NH<sub>4</sub>HCO<sub>3</sub>, pH 8.0, digested overnight at 30 °C with 3 µg sequencing-grade trypsin, and analysed by two-dimensional LC-MS/MS essentially as described<sup>32</sup>. Briefly, peptides were first separated on a first-dimension hydrophilic interaction liquid chromatography (HILIC) column with a reverse acetonitrile gradient and 25 fractions of 1 ml collected (2 min per fraction). After drying, peptides were analysed by LC-MS/MS on a C18 column. The MS scan routine was set to analyse by MS/MS the five most intense ions of each full MS scan; dynamic exclusion was enabled to assure detection of co-eluting peptides.

Protein identification by mass spectrometry. Raw data collection of approximately 230,000 MS/MS spectra per two-dimensional LC-MS/MS experiment was followed by protein identification using SEQUEST. All MS raw files have been deposited in the ProteomeXchange Consortium<sup>33</sup> via the PRIDE partner repository with the data set identifier PXD002804. In detail, peak lists were generated using extract-msn (ThermoScientific) within Proteome Discoverer 1.4.1. From raw files, MS/MS spectra were exported with the following settings: peptide mass range 350-5,000 Da; minimal total ion intensity 500. The resulting peak lists were searched using SequestHT against a target-decoy E. coli protein database (release 07.01.2008, 8,678 entries comprising forward and reverse sequences) obtained from Uniprot. The following parameters were used: trypsin was selected with proteolytic cleavage only after arginine and lysine, number of internal cleavage sites was set to 1, mass tolerance for precursors and fragment ions was 1.0 Da, and considered dynamic modifications were +15.99 Da for oxidized methionine and +125.12 Da for N-ethylmaleimide on cysteines. Peptide matches were filtered using the q value and posterior error probability calculated by the Percolator algorithm ensuring an estimated false positive rate below 5%. The filtered SEQUEST HT output files for each peptide were grouped according to the protein from which they were derived using the multiconsensus results tool within Proteome Discoverer. Then the values of the spectral matches of only Met-containing peptides were combined from the three two-dimensional LC-MS/MS experiments and exported in a Microsoft Excel spreadsheet, with the rows referring to the peptide sequences and the columns to the fractions of the HILIC column. Oxidation of Met residues to Met-O by NaOCl causes a hydrophilic shift, which influences their retention time and makes them elute later (4-8 min) than their reduced counterpart on a HILIC column. If these Met-O are reduced by MsrP, they will then show a hydrophobic shift and elute at the same retention time on the HILIC column as in the control sample. By comparing the retention times and the number of peptide spectral matches of the Met-O-containing peptides in a periplasmic extract under three experimental conditions (control, oxidized by NaOCl with and without MsrP), one can identify 'bona fide' potential MsrP substrates.

**Protein expression and purification.** TP1004 cells harbouring plasmid pAG178 and overexpressing MsrP-His<sub>6</sub> protein were grown aerobically at 30 °C in terrific broth (Sigma-Aldrich) supplemented with sodium molybdate (1.5 mM) and ampicillin ( $200 \,\mu$ g ml<sup>-1</sup>). When cells reached  $A_{600\,nm} = 0.8$ , expression was induced with 0.1 mM IPTG for 3 h. Periplasmic proteins were then extracted as in ref. 32. MsrP-His<sub>6</sub> was then purified by loading the periplasmic extract on a 1 ml HisTrap FF column (GE Healthcare) equilibrated with buffer A (NaPi 50 mM, pH 8.0, NaCl 300 mM). After washing the column with buffer A, MsrP-His<sub>6</sub> was eluted by applying a linear gradient of imidazole (from 0 to 300 mM) in buffer A. The fractions containing MsrP-His<sub>6</sub> were pooled, concentrated using a 5 kDa cutoff Vivaspin

15 (Sartorius) device and de-salted on a PD-10 column (GE Healthcare) equilibrated with 50 mM NaPi, pH 8.0, 150 mM NaCl.

VU1 CaM, MsrA and MsrB were expressed and purified as described previously<sup>34,35</sup>.

Trx was expressed and purified as follows. BL21 (DE3) cells harbouring plasmid pMD205, overexpressing Trx with a carboxy (C)-terminal His<sub>6</sub> tag, were grown aerobically at 37 °C in LB supplemented with kanamycin ( $50 \mu g ml^{-1}$ ). Expression was induced at  $A_{600 nm} = 0.6$  with 1 mM IPTG for 3 h. Cells were then pelleted, resuspended in buffer A (NaPi 50 mM, pH 8.0, NaCl 300 mM) and disrupted by two passes through a French pressure cell at 12,000 psi. The lysate was then centrifuged at 30,000g and at 4 °C for 45 min, to remove cell debris, and Trx was purified as described for MsrP-His<sub>6</sub>. Ni-NTA-purified Trx was then loaded on a 120 ml HiLoad 16/60 Superdex 75 PG column (GE Healthcare) previously equilibrated with HEPES–KOH 50 mM, pH 7.4, NaCl 100 mM. The resulting Trx-containing fractions were pooled and concentrated using a 5 kDa cutoff Vivaspin 15 device.

Thioredoxin reductase (TrxR) was expressed and purified as follows. BL21 (DE3) cells harbouring plasmid pPL223-2, overexpressing TrxR with an amino (N)-terminal His<sub>6</sub> tag, were grown aerobically at 37 °C in LB supplemented with ampicillin ( $200 \,\mu g \, ml^{-1}$ ). Expression was induced at  $A_{600 \, nm} = 0.6$  with 1 mM IPTG for 3 h. Protein extraction was performed as described for Trx and purification was performed as described for MsrP-His<sub>6</sub>.

BL21 (DE3) cells harbouring plasmid pKD11, overexpressing SurA with a C-terminal His<sub>6</sub> tag, were grown aerobically at 37 °C in LB supplemented with kanamycin (50 $\mu$ g ml<sup>-1</sup>). Expression was induced at  $A_{600 \text{ nm}} = 0.6$  with 1 mM IPTG for 3h. Protein extraction and purification were performed as described for MsrP-His<sub>6</sub>.

MG1655 cells harbouring plasmid pKD84, overexpressing SurA with a C-terminal *Strep*-tag, were grown aerobically at 37 °C in LB supplemented with ampicillin ( $200 \,\mu g \, ml^{-1}$ ). Expression was induced at  $A_{600 \, nm} = 0.7$  with a final concentration of  $200 \,\mu g \, l^{-1}$  anhydrotetracycline (AHT) for 5 h. Protein extraction was performed as described for MsrP-His<sub>6</sub>. SurA-*Strep* was then purified by loading the periplasmic extract on a 5 ml Strep-Tactin Superflow cartridge H-PR (IBA) equilibrated with buffer A (Tris-HCl 100 mM, pH 8.0, NaCl 150 mM, EDTA 1 mM). After washing the column with buffer A, SurA-*Strep* was eluted by applying a linear gradient of desthiobiotin (from 0 to 2.5 mM) in buffer A. The fractions containing SurA-*Strep* were pooled, concentrated using a 5 kDa cutoff Vivaspin 15 (Sartorius) device and de-salted on a PD-10 column (GE Healthcare) equilibrated with 50 mM NaPi, pH 8.0, 150 mM NaCl.

A modified version of Pal lacking the signal sequence and in which the first cysteine of the lipobox was replaced by an alanine (Pal<sub>C1A</sub>) was expressed with an N-terminal His<sub>6</sub> tag from the pEB0513 vector in BL21 (DE3) cells. Cells were grown aerobically at 37 °C in LB supplemented with ampicillin ( $200 \mu g m l^{-1}$ ). Expression was induced at  $A_{600 nm} = 0.6$  with 1 mM IPTG for 3 h. Protein extraction was performed as described for Trx and purification was performed as described for MsrP-His<sub>6</sub>.

*In vitro* repair of oxidized CaM, SurA and Pal by MsrP. CaM was oxidized *in vitro* as described previously<sup>36</sup>. SurA-His<sub>6</sub> and Pal were oxidized *in vitro* by incubating the purified proteins ( $50\mu$ M) for 2 h 30 min at 30 °C with 100 mM H<sub>2</sub>O<sub>2</sub> in a buffer containing 50 mM NaPi, pH 8.0, 50 mM NaCl. H<sub>2</sub>O<sub>2</sub> was then removed by gel filtration using a NAP-5 column (GE Healthcare) equilibrated with 50 mM NaPi, pH 8.0, 150 mM NaCl.

In vitro repair of oxidized CaM (CaMox), SurA (SurA ox) and Pal (Pal ox) was assessed by incubating the oxidized proteins (2µM of CaMox and SurA ox, 5µM of Pal ox) with purified MsrP-His<sub>6</sub> (2µM for CaMox and SurA ox, 5µM for Pal ox), 10 mM benzyl viologen and an excess of sodium dithionite at 37 °C for 1 h. As controls, the oxidized proteins were incubated separately with either MsrP-His<sub>6</sub> or the inorganic reducing system (benzyl viologen and sodium dithionite). The reactions were stopped by adding SB buffer and heating at 95 °C for the CaM and SurA samples or by adding 0.1% trifluoroacetic acid for the Pal samples. The CaM and SurA samples were then loaded on an SDS-PAGE gel and the proteins visualized with the PageBlue Protein Staining Solution (Fermentas). For the Pal samples (20µg), proteins were separated by reverse-phase high-performance liquid chromatography on a C4 column (Vydac 214TP54, 4.6 mm × 250 mm) at a flow rate of  $400\,\mu l\,min^{-1}$  with a linear gradient of acetonitrile in 0.1% trifluoroacetic acid (0-70% acetonitrile in 90 min). Absorbance was monitored at 214 nm and the peaks were collected. The fractions were dried in a Speedvac and the proteins resupsended in  $25\,\mu l$  of  $100\,mM$   $NH_4HCO_3$  before overnight digestion at 30 °C with 0.5 µg of trypsin or EndoGlu-C. The peptides were then analysed as described below.

For CaM and SurA, the gel bands corresponding to the different oxidation states were in-gel digested with trypsin and the resulting peptides analysed by LC–MS/MS on a C18 reverse-phase column as described above. Relative abundances of every Met-containing peptide in its different oxidation state were obtained by integration of peak area intensities, taking into account the extracted ion chromatogram of both doubly and triply charged ions.

*In vivo* repair of oxidized SurA and Pal by MsrP. The *in vivo* repair of SurA ox and Pal ox by the MsrPQ system or MsrP alone expressed from plasmids pAG195 and pAG192, respectively, was performed as follows. Overnight cultures of AG233 (containing the empty pAG177 vector), AG234 (containing the pAG195 plasmid) and AG289 (containing the pAG192 plasmid) were diluted to  $A_{600 \text{ nm}} = 0.04$  into fresh LB medium (100 ml) and cells were grown aerobically at 37 °C in the presence of 0.1 mM IPTG and 200 µg ml<sup>-1</sup> ampicillin. At  $A_{600 \text{ nm}} = 0.5$ , cells were subjected to NaOCl treatment (3.5 mM) and protein synthesis was blocked by the addition of chloramphenicol (300 µg ml<sup>-1</sup>). Samples were taken at different time points after NaOCl addition and precipitated with TCA. The pellets were then washed with ice-cold acetone, suspended in SB buffer, heated at 95 °C and loaded on a SDS–PAGE gel for immunoblot analysis using anti-Pal<sup>37</sup> and anti-SurA antibodies. The specificity of the anti-SurA antibody was verified (Supplementary Fig. 6). The protein amounts loaded were standardized by taking into account the  $A_{600 \text{ nm}}$ values of the cultures.

Oxidation, repair and purification of SurA for analysis of chaperone function. SurA-Strep was oxidized in vitro by incubating the purified protein (200 µM) for 3 h at 30 °C with 100 mM H<sub>2</sub>O<sub>2</sub> in a buffer containing 50 mM NaPi, pH 8.0, 150 mM NaCl. H<sub>2</sub>O<sub>2</sub> was then removed by gel filtration using a NAP-5 column (GE Healthcare) equilibrated with 50 mM NaPi, pH 8.0, 150 mM NaCl. For the in vitro repair of oxidized SurA (SurA ox), the oxidized protein (30 µM) was incubated with purified MsrP-His<sub>6</sub> (30µM), 10 mM benzyl viologen and 10 mM of sodium dithionite at 37 °C for 1 h. Following repair, SurA was purified by passing the sample through a gravity flow column containing 200 µl Strep-Tactin Sepharose beads (from a 50% suspension, IBA), previously equilibrated with buffer A (Tris-HCl 100 mM, pH 8.0, NaCl 150 mM, EDTA 1 mM). After washing with buffer A, repaired SurA was eluted using buffer A containing 2.5 mM desthiobiotin. The elution fractions were pooled and submitted to buffer exchange using a NAP-5 column (GE Healthcare) equilibrated with 50 mM NaPi, pH 8.0, 150 mM NaCl. To check for the correct oxidation, repair and purification of SurA, samples were loaded on an SDS-PAGE gel and the proteins visualized with the PageBlue Protein Staining Solution (Fermentas).

Analysis of chaperone function. The ability of SurA to act as a chaperone preventing the thermal aggregation of citrate synthase (Sigma, reference C3260) was assessed as follows. The aggregation of citrate synthase  $(0.15 \,\mu\text{M})$  was monitored at 43 °C in 40 mM HEPES–KOH, pH 7.5, in the absence or in the presence of  $0.6 \,\mu\text{M}$  SurA, SurA ox or MsrP-repaired SurA ox using light-scattering measurements. To avoid effects that might have been caused by the protein buffer, all samples were added to the assay in constant volume. SurA ox and MsrP-repaired SurA ox were obtained as described above. Light-scattering measurements were made using a Varian Cary Eclipse spectrofluorometer both with excitation and with emission wavelengths set to 500 nm at a spectral bandwidth of 2.5 nm. Data points were recorded every 0.1 s.

**Genetic analysis of Met-O assimilation.** The ability of various *E. coli* strains (BE100, JB08, CH193, BE104) to assimilate Met-O was assessed on M9 minimal medium supplemented with either Met or Met-O at  $20 \,\mu g \, ml^{-1}$ . Plates were incubated at 37 °C for 72 h. Overnight cultures of strains AG272, AG273, AG279 and AG274 were diluted to  $A_{600 \, nm} = 0.04$  into fresh M63 minimal medium (100 ml) supplemented with 0.5% glycerol,  $150 \,\mu g \, ml^{-1}$  of each amino acid, 1 mM MgSO4, 1 mM MoNa<sub>2</sub>O<sub>4</sub>, 17  $\mu$ M Fe<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>, vitamins (thiamine  $10 \,\mu g \, ml^{-1}$ ) spectinomycin, and grown aerobically at 37 °C. When  $A_{600 \, nm}$  reached 0.5, cells (5 ml) were washed three times with M63 medium containing  $150 \,\mu g \, ml^{-1}$  Met-O instead of methionine, and serially diluted in the same medium. Five microlitres of each dilution were then spotted on M63 plates containing either Met or Met-O at  $150 \,\mu g \, ml^{-1}$ , and plates were subsequently incubated at 37 °C for 40 h.

**HOCl induction assays.** The *msrP::lacZ*-containing strains (CH183, CH186 and CH187) were grown at 37 °C with shaking in M9 minimal medium. When cells reached  $A_{600\,nm} \approx 0.2$ , cultures were split into two plastic tubes, one of them containing HOCl ( $200\,\mu$ M). These tubes were then incubated with an inclination of 90° with shaking at 37 °C. After 30 min of incubation, 1 ml was harvested and the bacteria were resuspended in 1 ml of  $\beta$ -galactosidase buffer. Levels of  $\beta$ -galactosidase were measured as described<sup>38</sup>.

**HOCI survival assays.** NR744, NR745, CH0127 and AG190 cells were grown aerobically at 37 °C with shaking in 50 ml of LB medium in 500 ml flasks. When cells reached  $A_{600 \text{ nm}} \approx 0.45$ , 5 ml samples were transferred to conical polypropylene centrifuge tubes (50 ml; Sarstedt) and HOCl (2 mM) was added. Cells were then incubated at 37 °C with shaking (150 r.p.m.) at 90° inclination. Samples were taken at various time points after stress, diluted in PBS buffer, spotted on LB agar and incubated at 37 °C for 16 h. Cell survival was determined by counting colony-forming units (c.f.u.) per millilitre. The absolute c.f.u. at time-point 0 (used as 100%) was  ${\sim}10^8$  cells per millilitre in all experiments. For strains CH194, CH196 and CH197, the same protocol was used with chloramphenicol (25  $\mu g\,ml^{-1}$ ) and arabinose (0.2%) added to the cultures.

SDS survival assays. Cells (MG1655 and BE107) were grown at 37 °C with shaking in 10 ml of LB (in 100 ml flasks). When cells reached  $A_{600\,\text{nm}} \approx 0.8, 5 \text{ ml samples}$ were transferred to conical polypropylene centrifuge tubes (50 ml, Sarstedt) and HOCl (2 mM) was added. After 5 min of incubation, samples were taken and diluted in PBS buffer to  $\sim 2 \times 10^3$  cells per millilitre. Aliquots (100 µl) were then spread on LB agar plates containing SDS (1%). Colonies were counted the next day. Data set construction and phylogenetic analyses. A non-redundant local protein database containing 1,342 complete prokaryotic proteomes available in NCBI (http://www.ncbi.nlm.nih.gov/) as of 30 July 2014 was built. This database was queried with the BlastP program (default parameters)<sup>39</sup>, using YedY (NP\_416480) and YedZ (NP\_416481) of E. coli strain K-12 substrate MG1655 as a seed. Distinction between homologous and non-homologous sequences was assessed by visual inspection of each BlastP output (no arbitrary cut-off on the E value or score). To ensure that we did not overlook divergent YedY or YedZ proteins, iterative BlastP queries were performed using homologues identified at each step as new seeds. The list of YedY and YedZ homologues is provided in Supplementary Data 1. The retrieved sequences were aligned using MAFFT version 7 (default parameters<sup>40</sup>; Supplementary Data 2 and 3). Each alignment was visually inspected and manually refined when necessary using the ED program from the MUST package<sup>41</sup>. Regions where the homology between amino-acid positions was doubtful were removed by using BMGE software (BLOSUM30 similarity matrix<sup>42</sup>).

For each homologue, the genomic context was investigated using MGcV (Microbial Genomic context Viewer<sup>43</sup>). The domain composition and protein location of each homologue was also analysed using pfam version 27.0 (ref. 44), SignalP version 4.1 (ref. 45) and TMHMM server version 2.0 (ref. 46), respectively.

For the YedY protein, preliminary phylogenetic analysis used FastTree version 2 and a gamma distribution with four categories<sup>47</sup>. On the basis of the resulting tree, the subfamily containing the sequence from *E. coli* was identified and selected for further phylogenetic investigations. The corresponding sequences were realigned using MAFFT version 7. The resulting alignment was trimmed with BMGE as previously described.

Maximum likelihood trees were computed using PHYML version 3.1 (ref. 48) with the Le and Gascuel model (amino-acid frequencies estimated from the data set) and a gamma distribution (four discrete categories of sites and an estimated alpha parameter) to take into account variations in evolutionary rate across sites. Branch robustness was estimated by the non-parametric bootstrap procedure implemented in PhyML (100 replicates of the original data set with the same parameters). Bayesian inferences were performed using MrBayes 3.2 (ref. 49) with a mixed model of amino-acid substitution including a gamma distribution (four discrete categories). MrBayes was run with four chains for one million generations and trees were sampled every 100 generations. To construct the consensus tree, the first 2,000 trees were discarded as 'burn in'.

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### LETTER RESEARCH



**Extended Data Figure 1** | **Induction of MsrPQ by HOCI is dependent on the presence of a functional YedVW two-component system.** Top, immunoblot analysis shows that the induction of MsrP synthesis by HOCI (0.2 mM) is *yedW*-dependent. The image is representative of experiments

made in biological triplicate. Bottom, an *msrP::lacZ* fusion was used as a read-out for *msrP* expression. Deletion of *yedV* upregulates *msrP* expression, while deletion of *yedW* prevents its induction by HOCl. Error bars, mean  $\pm$  s.e.m.; n = 4. The uncropped blot is shown in Supplementary Fig. 4.



**Extended Data Figure 2** | **Respiratory chain-powered, nonstereospecific reduction of Met-O in periplasmic proteins by the MsrPQ system maintains envelope integrity.** Upon exposure to reactive species of chlorine (RCS) and/or reactive species of oxygen (ROS), methionine residues (Met) in periplasmic proteins such as SurA and Pal get oxidized and randomly form either the *R*- or the *S*- diastereoisomer of Met-O. This results in the loss of function of some proteins important for maintaining the integrity of the envelope, such as SurA, giving rise to envelope defects. MsrP catalyses the reduction of both diastereoisomers of Met-O with the help of its molybdenum-molybdopterin (Mo-MPT) cofactor. Electrons for reduction are provided by the quinone (Q) pool of the respiratory chain through MsrQ, the inner membrane haem *b*-containing partner of MsrP. PG, peptidoglycan.





Extended Data Figure 3 | MsrP non-stereospecifically reduces Met-O. a, MsrP reduces *N*-acetyl-Met-O (NacMet-O), a substrate mimicking protein-bound Met-O, with  $K_m = 3.8 \pm 1.2$  mM, turnover number  $(k_{cat}) = 30.5 \pm 3.1$  s<sup>-1</sup> and  $V_{max} = 56.3 \pm 5.8 \,\mu$ mol min<sup>-1</sup> per milligram protein (error bars, mean  $\pm$  s.d.; n = 3). b, MsrP is a non-stereospecific Msr, being able to reduce both *S*-Met-O (with  $K_m = 8.0 \pm 2.7$  mM,  $k_{cat} = 36.0 \pm 3.6$  s<sup>-1</sup> and  $V_{max} = 67.2 \pm 6.4 \,\mu$ mol min<sup>-1</sup> per milligram protein) and *R*-Met-O (with  $K_m = 25.7 \pm 4.7$  mM,  $k_{cat} = 168.3 \pm 15.0$  s<sup>-1</sup> and  $V_{max} = 313.4 \pm 27.6 \,\mu$ mol min<sup>-1</sup> per milligram protein). Error bars, mean  $\pm$  s.d.; n = 3. c, Strain JB08 (Met<sup>-</sup> MsrA<sup>-</sup> MsrB<sup>-</sup> BisC<sup>-</sup>, producing MsrC) is able to grow only on *R*-Met-O, whereas strain CH193

(Met<sup>-</sup> MsrA<sup>-</sup> MsrB<sup>-</sup> MsrC<sup>-</sup>, producing BisC) in only able to grow on S-Met-O. Deletion of *msrP* in strain BE100 (Met<sup>-</sup> Msr<sup>-</sup> Sup<sup>Met-O<sup>+</sup></sup>) prevents its growth on *R*- and S-Met-O (strain BE104 = Met<sup>-</sup> Msr<sup>-</sup> Sup<sup>Met-O<sup>+</sup></sup>  $\Delta$ *msrP*, compare with growth of BE100 in Fig. 2e). Images are representative of experiments made in biological triplicate. **d**, The periplasmic chaperone SurA was treated with H<sub>2</sub>O<sub>2</sub>, giving rise to SurA ox, a sample of which was subsequently incubated with MsrP and the inorganic reducing system *in vitro*. The oxidation state of specific Met residues (Met 136, 231 and 298) in the various samples was determined by LC-MS/MS analysis. Error bars, mean  $\pm$  s.e.m.; *n* = 4.





of L-methionine sulfoxide in D<sub>2</sub>O pD 6.5, either as a mixture of *R*- and *S*- diastereoisomers (top), isolated *S*- (middle) or isolated *R*- (bottom) (containing 30 mM dioxane as an internal reference). **d**, Zoom on the <sup>13</sup>C NMR spectra of ~150 mM solutions of L-methionine sulfoxide in D<sub>2</sub>O pD 6.5, either as a mixture of *R*- and *S*- diastereoisomers (top), isolated *S*- (middle) or isolated *R*- (bottom) (containing 30 mM dioxane as an internal reference).

## LETTER RESEARCH





**Extended Data Figure 5** | **Individual phylogenies of YedY.** Shown are unrooted Bayesian phylogenetic trees for YedY (b1971, 310 sequences, 260 positions). Numbers at nodes indicate posterior probabilities computed by MrBayes<sup>49</sup> and bootstrap values computed by PhyML<sup>48</sup>.

Only posterior probabilities and bootstrap values above 0.5 and 50%, respectively, are shown. Scale bars, average number of substitutions per site. In the phylogenetic tree, YedY from *E. coli* is highlighted in grey.

## LETTER RESEARCH





**Extended Data Figure 6** | **Individual phylogenies of YedZ**. Shown are unrooted Bayesian phylogenetic trees for YedZ (b1972, 369 sequences, 135 positions). Numbers at nodes indicate posterior probabilities computed by MrBayes<sup>49</sup> and bootstrap values computed by PhyML<sup>48</sup>.

Only posterior probabilities and bootstrap values above 0.5 and 50%, respectively, are shown. Scale bars, average number of substitutions per site. In the phylogenetic tree, YedZ from *E. coli* is highlighted in grey.

#### Extended Data Table 1 | The MsrPQ system uses electrons from the respiratory chain to reduce free Met-O

Strain description	Met	Met-O
Met	+	+
Met <sup>-</sup> Msr <sup>-</sup> (JB590)	+	-
Met Msr Sup <sup>Met-O+</sup> (BE100)	+	+
Met Msr Sup <sup>Met-O+</sup> Δ <i>yedZ</i> (BE105)	+	-
Met Msr empty vector (AG272)	+	-
Met Msr pyedY (AG273)	+	-
Met Msr p <i>yedZ</i> (AG279)	+	-
Met Msr pyedYyedZ (AG274)	+	+
Met Msr Sup <sup>Met-O+</sup> $\Delta menA \Delta ubiE$ (BE106)	+	-

This table shows the ability of the various strains to grow (+) or not (-) using Met-O as the sole Met source. Strains were grown for 40–72 h at 37 °C. The results are representative of experiments made in biological triplicate.



Extended Data Table 2   Lis	at of proteins identified as	potential MsrP substrates
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Protein	Function	Number and percentage of methionines in the protein*
SurA	Primary periplasmic chaperone	14 (3.4%)
LolA	Outer-membrane lipoprotein carrier protein	2 (1.1%)
Pal	Peptidoglycan-associated lipoprotein	6 (3.9%)
MlaC	Probable phospholipid-binding protein	4 (2.1%)
PpiA	Peptidyl-prolyl cis-trans isomerase A	4 (2.4%)
DsbA	Thiol:disulfide interchange protein	6 (3.2%)
CysP	Thiosulfate-binding protein	4 (1.3%)
PotD	Spermidine/putrescine-binding periplasmic protein	9 (2.8%)
МррА	Periplasmic murein peptide-binding protein	7 (1.4%)
ProX	Glycine betaine-binding periplasmic protein	6 (1.9%)
MalE	Maltose-binding periplasmic protein	6 (1.6%)
MgIB	D-galactose-binding periplasmic protein	6 (1.9%)
RbsB	D-ribose-binding periplasmic protein	4 (1.5%)
FecB	Fe3+ dicitrate-binding periplasmic protein	7 (2.5%)
RcnB	Nickel/cobalt homeostasis protein	2 (2.3%)
ZnuA	High-affinity zinc uptake system protein	6 (2.1%)
Ecotin	General inhibitor of pancreatic serine proteases	4 (2.8%)
lvy	Inhibitor of vertebrate lysozyme	5 (3.9%)
PspE	Thiosulfate sulfurtransferase	2 (2.4%)
YmgD	Uncharacterized protein	4 (4.4%)

\*Referring to the mature protein without its signal sequence

Semi-quantitative two-dimensional LC–MS/MS analysis was used to identify proteins that have one or more oxidized Met residues that MsrP could reduce. The first column indicates the name of the protein, the second describes its function and the third gives the number and percentage of methionine residues in the mature protein (excluding the signal sequence).