The persistence of the gastric pathogen *Helicobacter pylori* is due in part to urease and Msr (methionine sulfoxide reductase). Upon exposure to relatively mild (21% partial pressure of \(O_2\)) oxidative stress, a \(\Delta msr\) mutant showed both decreased urease specific activity in cell-free extracts and decreased nickel associated with the partially purified urease fraction as compared with the parent strain, yet urease apoprotein levels were the same for the \(\Delta msr\) and wild-type extracts. Urease activity of the \(\Delta msr\) mutant was not significantly different from the wild-type upon non-stress microaerobic incubation of strains. Urease maturation occurs through nickel mobilization via a suit of known accessory proteins, one being the GTPase UreG. Treatment of UreG with \(H_2O_2\) resulted in oxidation of MS-identified methionine residues and loss of up to 70% of its GTPase activity. Incubation of pure \(H_2O_2\)-treated UreG with Msr led to reductive repair of nine methionine residues and recovery of up to full enzyme activity. Binding of Msr to both oxidized and non-oxidized UreG was observed by cross-linking. Therefore we conclude Msr aids the survival of *H. pylori* in part by ensuring continual UreG-mediated urease maturation under stress conditions.

Key words: accessory protein, amino acid modification, GTPase, nickel (Ni), oxidative stress, protein oxidation.

INTRODUCTION

*Helicobacter pylori* is a Gram-negative microaerophilic bacterium that colonizes the gastric mucosa [1] of approximately one-half of the world’s population [2]. The World Health Organization classifies *H. pylori* as a carcinogen and it is the causative agent of most peptic ulcer diseases, chronic gastritis and gastric cancer in humans [3]. The pathogenesis and persistence attributes of *H. pylori* rely on its ability to combat harsh conditions; these include both the acidic environment of the gastric lumen and chronic exposure to ROS (reactive oxygen species) [3]. During colonization, *H. pylori* rely on a superoxide dismutase (SodB), catalase (KatA), AhpC (alkyl hydroperoxide reductase), a neutral-philactivating protein (NapA) and an NADPH quinone reductase (MdaB) [9,11].

In bacteria and other organisms, two types of Mr proteins have been described: MsrA and MsrB. These two forms of Mr reduce both the two isomers Met(S)O and Met(R)O of methionine sulfoxide respectively [12,13]. In *H. pylori*, MsrA and MsrB are fused to constitute a 42 kDa protein [14]. An *H. pylori* msr-deficient strain has been shown to be highly sensitive to oxidative stress, and it has a greatly diminished ability to colonize the stomach [15]. *H. pylori* Mr has also been shown to play a role in protecting catalase from oxidative damage [16]. However, only a few methionine-rich proteins have been identified as Msr-interacting [16,17] and the full extent of the physiological roles of Mr remain unknown.

*H. pylori* resists the acidic environment of the gastric region by producing urease, which hydrolyses urea to bicarbonate and ammonia [18,19]. Urease is the most abundant protein made by *H. pylori*, as it accounts for more than 10% of the total protein synthesized by the bacterium [20,21]. Urease is composed of only two subunits, UreA and UreB [19], unlike other bacterial ureases that are composed of UreA, UreB and UreC subunits [25]. The maturation of urease requires incorporation of Ni (nickel) into the active site, which is accomplished by several accessory proteins [22,23]. In *H. pylori*, these include UreE, UreF, UreG and UreH [24]. On the basis of studies in *Klebsiella aerogenes*, it is generally accepted that UreD (UreH in *H. pylori*), UreF and UreG drive protein conformational change, lysine carboxylation and GTP hydrolysis respectively, whereas UreE functions as a metallochaperone of the maturation system [25–28].

Of the accessory proteins, UreG is the most highly conserved and shares sequence homology with ATP- and GTP-binding proteins [29]. UreG belongs to the group of homologous P-loop GTPases [26]. Loss of all urease activity occurs upon introduction of site-directed mutations at the nucleotide-binding domain for both *H. pylori* [30] as well as *K. aerogenes* UreG [26]. In addition to the nucleotide-binding domain, UreG is a methionine-rich protein with methionine comprising ~4.5% of the primary amino acid sequence. From a tandem affinity purification (TAP) approach with UreG as the bait protein, UreG was proposed to interact with up to 33 different proteins, of which one was Mr [27]. However, the role of Mr in this possible interaction has not been studied. The TAP results combined with the high proportion of...
methionine residues in UreG caused us to examine a role for Msr in urease maturation. The role of Msr was addressed by studying a Δmsr mutant and by assessing the ability of Msr to repair the oxidized methionine residues of UreG. Finally, we demonstrated the intimate interaction between purified UreG and Msr.

EXPERIMENTAL

Bacterial strains and growth conditions

H. pylori strain SS1 was used as the parental strain for all studies. H. pylori were routinely grown on Brucella agar (Oxoid) plates containing 10% defibrinated sheep blood (BA plates) (QuadFive) and maintained at 37°C under 5% CO2, 4% O2, balanced with N2, in a microaerobic humidified chamber. The Δmsr mutant was described previously [11,15]. Escherichia coli cultures were grown aerobically in LB (Luria–Bertani) broth or plates containing 10% defibrinated sheep blood (BA plates) (QuadFive) and maintained at 37°C under 5% CO2, 4% O2, balanced with N2, in a microaerobic humidified chamber. The Δmsr mutant was described previously [11,15]. Escherichia coli cultures were grown aerobically in LB (Luria–Bertani) broth or agar and ampicillin, kanamycin and chloramphenicol were added when needed at a final concentration of 100, 30 and 30 μg/ml respectively.

Protein purification

H. pylori UreG was expressed as a His6-tagged protein in E. coli BL2(DE3)-RIL (Novagen). Briefly, the ureG gene was amplified by PCR using genomic DNA from strain 43504 as a template and primers NdeUreG (5′-ACGGCCCTATAGGTTA-AATTGAGG-3′) and XhoUreG (5′-CGTGAAGCTCGAGA TTCCAAATAAAGCGTTG-3′, designed to amplify ureG without its stop codon). The resulting 0.6 kb DNA fragment was digested with Ndel and XhoI, and ligated into the similarly digested expression vector pET21b (Novagen). The recombinant plasmid was sequenced at the Georgia Genomics Facility (University of Georgia, Athens, GA, U.S.A.) to ensure that no error was introduced by PCR. E. coli cultures harbouring the recombinant plasmid were grown to a D600 of 0.6 at 37°C in 500 ml of LB medium with chloramphenicol and ampicillin. Cultures were then induced with 0.5 mM IPTG (isopropyl β-D-thiogalactopyranoside) at 37°C for 3 h. Cells were harvested by centrifugation and the pellet was resuspended in buffer A (50 mM Tris/HCl, pH 7.5, 500 mM NaCl and 40 mM imidazole), then broken by four passages through an ice-cold French pressure cell. The cell debris was removed by centrifugation at 15 000 g for 20 min at 4°C. The supernatant was loaded onto a Q-Sepharose Fast Flow column (GE Healthcare) with a 0–600 mM NaCl gradient. The peak urease-containing fractions were pooled and dialysed overnight against 20 mM NaCl (atomic absorption grade), assayed for protein concentration (BCA protein assay kit) and Ni levels were measured by graphite furnace atomic absorption spectrometry (Shimadzu AA-6701F) as described previously [32].

GSTase assay

GSTase assays were performed using a Malachite Green-based kit (Innova Biosciences) following the manufacturer’s instructions. Briefly, 6 μM untreated, oxidized or repaired (see section on Msr repair) UreG was incubated with 50 mM Tris, pH 7.5, 2.5 mM MgCl2 and 0.5 mM GTP for 30 min. Next, Gold mix (PColorLock™ Gold plus accelerator) was added to the mixture and incubated for an additional 2 min. The stabilizer buffer was then incubated with the mixture for 30 min. Finally, absorbance was determined at 595 nm using a Molecular Devices plate reader. The readings obtained at A0.5 were compared with a standard curve with known amounts of phosphate. All steps were carried out at room temperature (22°C).

Exposure of SS1 wild-type and Δmsr strains to oxygen stress and measurement of urease activity

H. pylori strain SS1 wild-type or the Δmsr mutant was grown for 48 h, resuspended in BHI (brain heart infusion) broth with 0.4% β-cyclodextrin, pH 7.0, and exposed to 21% O2 (air) for 2 h while shaking (200 rev/min) at 37°C. Cells were harvested by centrifugation at 10 000 g for 10 min and broken by sonication (Heat Systems Ultrasones sonicator, 10 s at 4 W output power and 40% duty cycle). Cell debris was removed via centrifugation at 14 000 g for 10 min, and the supernatant was assayed for urease activity according to the method of Weatherburn [31].

Partial purification of urease and Ni determination

Urease was partially purified from the oxygen-stressed SS1 wild-type and Δmsr strains as described previously [32] with the following modification. Briefly, 20 mM Hepes, pH 7.2, was used as the starting buffer and the protein was washed from the Q-Sepharose HiTrap column (GE Healthcare) with a 0–600 mM NaCl gradient. The peak urease-containing fractions were pooled and dialysed overnight against 20 mM NaCl (atomic absorption grade), assayed for protein concentration (BCA protein assay kit) and Ni levels were measured by graphite furnace atomic absorption spectrometry (Shimadzu AA-6701F) as described previously [32].

Immunoblot and quantification of urease in H. pylori crude extracts

H. pylori SS1 wild-type or the Δmsr mutant was exposed to 21% O2 as described above. Cells were then resuspended in BHI broth with 0.4% β-cyclodextrin and broken by sonication. Approximately 10 μg of crude extract from both strains was then loaded on to SDS/PAGE (12.5% gels) to assess purity. Proteins were visualized with Coomassie Brilliant Blue R-250. Pure UreG was then concentrated using an Amicon Ultra Centrifugal filter (Millipore) and protein concentration was determined with the BCA (bicinchoninic acid) protein kit (Pierce) and dialysed overnight against 20 mM NaCl (atomic absorption grade), assayed for protein concentration (BCA protein assay kit) and Ni levels were measured by graphite furnace atomic absorption spectrometry (Shimadzu AA-6701F) as described previously [32].

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Oxidation of purified UreG

Purified UreG at 27 μM was treated with 25–50 mM H₂O₂ for 3 h in the dark and at room temperature under aseptic conditions. Excess H₂O₂ was removed by overnight dialysis against 50 mM Tris/HCl, pH 7.5.

Repair of oxidant-damaged UreG

UreG or H₂O₂-treated (50 and 100 mM) UreG was repaired with Mrp (equimolar concentration relative to UreG) or buffer (50 mM Tris/HCl, pH 7.5) in the presence of 400 μM NADPH (Sigma), 5 μM Trx and 100 nM TrxR for 1 h at 37 °C. GTPase activities were then measured.

Repair of oxidant-damaged urease

The Δmrp mutant was exposed to 2 h of oxygen stress and then the urease was partially purified from the mutant as described above. Partially purified urease (14 μg of total protein) was incubated with 10 mM DTT (dithiothreitol) alone or with 10 mM DTT and 14 μg Msr at 37 °C for 1 h. Urease activity was then measured according to the method of Weatherburn [31].

Asp-N endoproteinase (Asp-N) digestion and pseudo-MRM (multiple reaction monitoring) LC (liquid chromatography)–MS/MS (tandem MS) analysis

UreG samples were digested with Asp-N and analysed by C₁₈ reverse-phase LC coupled with pseudo-MRM MS/MS as described previously [16]. Briefly, samples were heated to 60 °C in 50 mM Tris/HCl, pH 7.5. The samples were then exposed to cool to room temperature and Asp-N protease (Thermo Scientific Pierce) was added at 1:100 (protease/UreG) and digested overnight at 37 °C. Samples were stored at −20 °C until analysis.

After digestion, samples were analysed using the LTQ front-end of an LTQ-FIT mass spectrometer (Thermo Scientific) coupled to an Agilent 1100 HPLC system with CaptiveSpray ionization using an Advance Ion Source for Thermo MS (Michrom Bioresearch) samples were autoinjected on to a C₁₈ reverse-phase LC coupled with pseudo-MRM MS/MS as described above. The samples were heated to 60 °C in 50 mM Tris/HCl and 0.5 mM zinc acetate at pH 8.0 in the presence of 10 mM DTT for 45 min. They were then allowed to cool to room temperature and Asp-N protease (Thermo Scientific Pierce) was added at 1:100 (protease/UreG) and digested overnight at 37 °C. Samples were stored at −20 °C until analysis.

RESULTS

Statistical significance

Results are presented as the means ± S.D. All data comparisons were performed using Student’s t test. These values were calculated using the Graphpad QuickCalcs website (http://graphpad.com/quickcalcs/).

RESULTS

Purification of UreG

To study H. pylori UreG biochemically, the protein was overexpressed and purified as a recombinant His₆-tagged protein from E. coli. The purity of the protein was assessed via SDS/PAGE analysis. The purified UreG migrated at a mass of approximately 24 kDa (Figure 1). The Authors Journal compilation © 2013 Biochemical Society

Interaction between UreG and Msr

To analyse the direct binding of Msr to UreG we used a Sulfo-SBED trifunctional cross-linking reagent. Sulfo-SBED contains biotin, a sulfo-NHS (sulfo-N-hydroxysuccinimide) active ester, and a photoactivatable aryl azide. Upon incubation of untreated ‘as-purified’ UreG and separately H₂O₂-treated UreG (oxUreG) with Sulfo-SBED, the sulfo-NHS ester reacted with the amine

Figure 1 SDS/PAGE of purified UreG

Lane 1, molecular mass marker with values indicated to the left in kDa; lane 2, cell extract from non-induced E. coli BL21(DE3)-RIL harbouring pET21b-ureG; lane 3, cell extract from IPTG-induced E. coli BL21(DE3)-RIL harbouring pET21b-ureG; lane 4, purified UreG after Ni-NTA extraction; lane 5, concentrated pure UreG. The arrowhead to the right indicates UreG.

Scientific). Sulfo-SBED was dissolved in dimethylformamide at 40 μg/ml and then added to UreG in PBS at a 5-fold molar excess. The mixture was incubated at room temperature for 30 min in the dark to preserve the aryl azide group. Excess Sulfo-SBED was removed via overnight dialysis against 50 mM Tris, pH 7.5. UreG or oxUreG conjugated to Sulfo-SBED (4 μM) was then incubated with equal molar concentrations of Msr, lysozyme or alone in a final volume of 40 μl for 2 h at room temperature in the dark. The binding of Msr to UreG was captured upon UV photoactivation of the aryl azide moiety. UV photoactivation was carried out using a 365 nm UV lamp held 5 cm from the mixture for 15 min on ice. The cross-linked mixture (20 μl) was then reduced with 0.5 M DTT in the dark for 10 min. The proteins were then resolved via SDS/PAGE followed by transfer on to a nitrocellulose membrane. The biotin label was detected by incubating the membrane-bound proteins with streptavidin–HRP (horseradish peroxidase) for 1 h. The membrane was developed by enhanced chemiluminescence (GE Healthcare).

Cross-linking and biotin label transfer

Purified UreG or 50 mM H₂O₂-treated UreG (oxUreG) was conjugated to Sulfo-SBED [sulfosuccinimidyl-2-[6-(biotin-amido)-2-(p-azidobenzamido)hexanoamido]ethyl-1,3-dithioproionate] using the Sulfo-SBED biotin label transfer kit (Thermo Scientific). Sulfo-SBED was dissolved in dimethylformamide at 40 μg/ml and then added to UreG in PBS at a 5-fold molar excess. The mixture was incubated at room temperature for 30 min in the dark to preserve the aryl azide group. Excess Sulfo-SBED was removed via overnight dialysis against 50 mM Tris, pH 7.5. UreG or oxUreG conjugated to Sulfo-SBED (4 μM) was then incubated with equal molar concentrations of Msr, lysozyme or alone in a final volume of 40 μl for 2 h at room temperature in the dark. The binding of Msr to UreG was captured upon UV photoactivation of the aryl azide moiety. UV photoactivation was carried out using a 365 nm UV lamp held 5 cm from the mixture for 15 min on ice. The cross-linked mixture (20 μl) was then reduced with 0.5 M DTT in the dark for 10 min. The proteins were then resolved via SDS/PAGE followed by transfer on to a nitrocellulose membrane. The biotin label was detected by incubating the membrane-bound proteins with streptavidin–HRP (horseradish peroxidase) for 1 h. The membrane was developed by enhanced chemiluminescence (GE Healthcare).

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UreG has previously been shown to be negligible. H. pylori exert a cumulative protein oxidation affect. The GTPase activity of are subject to a variety of oxidants simultaneously that are likely to oxidize repair targets. It is also important to note that pathogens oxidant levels seem high, but are the amounts sometimes used damage, we incubated purified UreG with H2O2 and measured.

Figure 2 Interaction between UreG and Msr identified by biotin transfer

Sulfo-SBED-conjugated UreG or 50 mM H2O2-oxidized UreG was incubated with Msr, lysozyme or alone and then the mixture was subjected to UV-cross-linking. Samples were taken before and after exposure to UV light. The conjugated samples were reduced with 0.5 M DTT for label transfer and the proteins were resolved via SDS/PAGE, transferred on to a nitrocellulose membrane, and probed with streptavidin–HRP. Lane 1, UreG; lane 2, oxidized UreG; lane 3, UreG and lysozyme; lane 4, oxidized UreG and lysozyme; lane 5, UreG and Msr; lane 6, oxidized UreG and Msr; lane 7, UreG; lane 8, oxidized UreG; lane 9, UreG and Msr; lane 10, oxidized UreG and Msr. Samples in lanes 7–10 were not exposed to UV light. The molecular masses are displayed to the left with values in kDa. The arrowheads to the right indicate Msr or oxidized UreG and Msr. Samples in lanes 7–10).

UreG activities in oxidized and repaired samples

To determine the susceptibility of H. pylori UreG to oxidant damage, we incubated purified UreG with H2O2, and measured the resulting GTPase activity. Incubation of purified UreG with H2O2 resulted in a dose-dependent decrease in GTPase activity (Figure 3). After incubation of 6 μM UreG with 25, 50 and 100 mM H2O2, the GTPase activity was significantly decreased to ∼44, 35 and 28 % of the untreated sample respectively. These oxidant levels seem high, but are the amounts sometimes used to oxidize repair targets. It is also important to note that pathogens are subject to a variety of oxidants simultaneously that are likely to exert a cumulative protein oxidation affect. The GTPase activity of H. pylori UreG has previously been shown to be negligible [30,35]. However, the calculated kcat of 0.032 min−1 for the untreated H. pylori UreG used in the present study is comparable with the measured GTPase activities of UreG in other organisms (Bacillus pasteurii UreG kcat = 0.04 min−1, Mycobacterium tuberculosis UreG kcat = 0.01 min−1, Glycine max UreG kcat = 0.01 min−1) [36–38]. To determine whether the decrease in GTPase activity after UreG incubation with H2O2 was due to methionine oxidation and also whether Msr is capable of restoring activity, oxidized UreG samples were incubated with Msr (plus repair system) and the resulting GTPase activities were measured. As a control, UreG was incubated with the repair components (Trx, TrxR and NADPH) without Msr. The latter sample is considered the untreated sample and is given 100 % activity (in earlier experiments it had the same activity as untreated UreG with none of the repair mixture components added). Upon incubation with Msr-containing repair components (Msr, Trx, TrxR and NADPH), the activity of the oxidant-damaged UreG was restored (Figure 4); it achieved full (non-oxidant-treated) levels for the 50 mM H2O2-oxidized sample and ∼80 % of the non-oxidant-treated levels for the 100 mM H2O2-treated sample. GTPase restoration (i.e. repair) was never seen when the oxidized UreG samples were incubated with the repair system without Msr. These results indicate that Msr can repair oxidatively damaged UreG and restore enzyme activity. Interestingly, a 20 % increase in activity with the addition of Msr to the untreated sample was sometimes observed, indicating some spontaneous UreG oxidation during preparation and storage.

Figure 3 H2O2 inactivation of UreG GTPase activity

Purified UreG (6 μM) was incubated with various concentrations of H2O2 (0, 25, 50 or 100 mM) for 3 h. Excess oxidant was removed via overnight dialysis. GTPase activities were measured using a colorimetric assay to detect the release of Pi, and are presented as percentage activity of the untreated sample. The untreated sample (100 %) is 0.186 μmol P1/min per mg of UreG. Each concentration is statistically significantly less than every higher concentration shown in the Figure at P < 0.05. Results are means ± S.D. (n = 8, two independent experiments were each sampled four times).

MS/MS identification of methionine residues oxidized and repaired in UreG

UreG samples were digested with Asp-N and the resulting peptide mixture was then subjected to LC–MS/MS. All nine methionine residues of UreG could be identified and they all showed significant oxidation after treatment with 50 mM H2O2 (Figure 5). For all untreated samples, the oxidation for all methionine residues remained less than 10 %. Met1, Met25, Met47, Met77, Met85 and Met160 were all greater than 80 % oxidized after treatment with H2O2. Met1, Met53 and Met160 were ∼70–80 % oxidized. After incubation of the oxidized samples with Msr plus 10 mM DTT as
**Figure 4** Msr repair of H$_2$O$_2$-damaged UreG

H$_2$O$_2$-treated UreG (6 μM) was incubated with equimolar amounts of Msr or buffer, along with the Msr repair components (400 μM NADPH, 5 μM Trx and 100 nM TrxR) at 37 °C for 1 h. The samples were then assayed for GTPase activity spectrophotometrically. UreG GTPase activity of the untreated sample without the addition of Msr is considered as 100 % and is 0.136 μmol of P/min per mg of UreG. Results are means ± S.D. (n = 6, three independent experiments sampled in duplicate).

**Figure 5** MS/MS identification of methionine residues after oxidation and repair of UreG

UreG was incubated with buffer or 50 mM H$_2$O$_2$ in buffer for 3 h. Excess H$_2$O$_2$ was removed via overnight dialysis. Following dialysis, oxidized UreG samples were incubated with or without Msr in the presence of DTT at 37 °C for 1 h. DTT alone did not result in the repair of any methionine residues. No oxidized Met-SO$_2$ (methionine sulfoxide) repair of any residues. The results indicate that Msr is able to repair all methionine residues of UreG.

the reducing agent, all methionine residues remained less than 20 % oxidized (Figure 5). DTT alone did not cause Met-SO$_2$ (methionine sulfoxide) repair of any residues. The results indicate that Msr is able to repair all methionine residues of UreG.

**Urease activities in wild-type and Δmsr strains after exposure to mild oxidative stress**

Despite expressing a broad repertoire of stress-combating enzymes and being an obligate aerobe, *H. pylori* is sensitive to oxidative molecules, including high oxygen. All wild-type strains of the bacterium are routinely grown in controlled atmospheres maintained below 12 % partial pressure O$_2$ [10]. We thus addressed the physiological importance of UreG repair under a relatively mild stress condition, i.e. ambient O$_2$ levels. Since the UreG GTPase activity is susceptible to oxidation (Figure 3), and the activity is restored by Msr (Figure 4), we reasoned the importance of UreG repair could be assessed by comparing the Δmsr mutant with the parent in urease activity upon cell exposure to 2 h of ambient O$_2$. This mild oxidant treatment has been used previously to study roles of oxidative stress enzymes in whole *H. pylori* cells [39,40]; the 2 h treatment causes viability loss of some oxidative-stress-sensitive mutants, but the parent strain maintains full viability (but not growth) during this incubation [15,40]. A significant decrease in cell-free extract urease activity to 64 % of the parent strain level was observed in the Δmsr mutant (Figure 6A) after 2 h of air exposure. As a control, the wild-type and Δmsr strains were left at 4 % partial pressure oxygen and the Δmsr mutant showed no statistically significant difference from the wild-type in activity over the 2 h period (Figure 6A). If urease maturation is the deficiency associated with the Δmsr mutant, we would expect urease apo-protein levels to be equal in that mutant and the parent. Urease apo-protein levels indeed appear to be...
Urease activity from the oxygen-stressed during this treatment. To address this concern, partially purified that urease itself could be damaged by methionine oxidation after exposure to 21 % O2. Whole-cell extracts were resolved via SDS/PAGE, transferred on to by sonication and urease activity was measured in cell-free extracts. Results are means +− S.D (n = 12, based on four independent experiments each sampled in triplicate). *P < 0.01. (B) SDS/PAGE analysis of cell-free extract from SS1 wild-type (9 μg) and Δmsr (10 μg) after exposure to 21 % O2 (C) Immunoblot analysis of SS1 wild-type and Δmsr cell-free extract after exposure to 21 % O2. Whole-cell extracts were resolved via SDS/PAGE, transferred on to a nitrocellulose membrane and blotted with anti-UreA antibodies. Molecular masses in kDa are indicated to the left-hand side of (B) and (C).

Figure 6 Urease activity and expression after oxidant stress

(A) Urease activity after exposure to oxidative stress. H. pylori SS1 wild-type (WT) and Δmsr strains were exposed to 21 % O2 (air) for 2 h or left at 4 % O2 as a control. Cells were then lysed by sonication and urease activity was measured in cell-free extracts. Results are means ± S.D (n = 12, based on four independent experiments each sampled in triplicate). *P < 0.01. (B) SDS/PAGE analysis of cell-free extract from SS1 wild-type (9 μg) and Δmsr (10 μg) after exposure to 21 % O2. (C) Immunoblot analysis of SS1 wild-type and Δmsr cell-free extract after exposure to 21 % O2. Whole-cell extracts were resolved via SDS/PAGE, transferred on to a nitrocellulose membrane and blotted with anti-UreA antibodies. Molecular masses in kDa are indicated to the left-hand side of (B) and (C).

Evidence to suggest this decrease in activity is due to a deficiency of urease maturation was obtained by measuring the Ni content associated with partially purified urease of both parent and mutant strains. Urease was partially purified and the urease fractions were analysed for Ni levels. The Δmsr mutant contained 33 % less Ni content than the wild-type (30 ± 1.6 ng of Ni per mg of protein for the wild-type and 20 ± 0.8 ng of Ni per mg of protein for Δmsr). These data were based on 13 replicates for each strain and the difference between the two strains was statistically significant (P < 0.01). Still, as urease was only partially purified, some of the Ni measurement could be associated with other proteins in both strains. The decreased urease-specific activity shown in the oxygen-stressed Δmsr mutant raises the possibility that urease itself could be damaged by methionine oxidation during this treatment. To address this concern, partially purified urease from the oxygen-stressed Δmsr mutant was incubated with DTT, or with DTT and Msr. No gain in urease activity was seen in either case (results not shown). These results suggest that the decrease in urease activity observed in the oxidant-stressed strain is not due to methionine oxidation of urease. Rather, UreG and possibly other proteins involved in urease maturation are prone to methionine oxidation and can be repaired by Msr.

DISCUSSION

The long-term survivability of H. pylori in the host requires that the bacterium survive harsh conditions, including the host inflammatory responses. The pathogen’s persistence is key to the most severe disease symptoms, and is in part due to the battery of DNA- and protein-repair enzymes that confer oxidative stress resistance on the pathogen in vivo. Proteins are often the targets of ROS due to their abundance in cells and their ease of reactivity with oxidants, and methionine is one of the most sensitive amino acids to oxidation. It is of obvious benefit for the bacterium to repair damaged Met-SO-containing proteins rather than synthesize new proteins due to energy input costs for synthesis and in order to rapidly maintain key enzyme function.

Many oxidative-stress-combating enzymes have been described in H. pylori, but our knowledge of the significance and physiological role of Msr continues to expand. An H. pylori Δmsr mutant shows attenuated growth in the presence of chemical oxidants and the strain is severely deficient in its ability to colonize the mouse stomach [15]. In addition, carbonylated proteins have been shown to accumulate in a Δmsr mutant after exposure of cells to oxidant damage [16]. These proteins were not oxidized in the parent strain [15]. However, only a few specific targets of Msr-mediated repair in H. pylori have been identified. These include GroEL, SSR (site-specific recombinase), AhpC and catalase [16,17]. Msr-repair targets in other bacteria include GroEL and Ffh [41,42].

In the host gastric region, H. pylori is in frequent contact with acid and oxidative molecules. Urease is essential for H. pylori to survive the acid stress and it is a highly abundant protein that requires Ni to function. Inside the host, H. pylori probably encounters various levels of Ni, so it employs many Ni-sequestering proteins (HspA, Hpn and Hpn-like) thought to gather the ion for storage and possible transfer to the metalloenzymes [43–45]. To achieve optimal urease activity, the active sites must be fully loaded with Ni and the Ni demand is thus high; Ni-saturated urease contains 12 Ni atoms per molecule [46]. As Mobley et al. [47] demonstrated, higher counts of Ni in recombinant urease correlated with higher urease activities. We observed a 31 % decrease in urease activity and a concomitant 33 % decrease in urease-associated Ni levels in the Δmsr mutant compared with the parent after exposure to mild oxidative conditions (e.g. air). Although modest, we propose that this is significant to the in vivo situation, where methionine damage could be much greater, and more potent oxidants exist. Indeed, Stingl and De Reuse [48] calculated that under in vitro conditions without added Ni, only a small proportion of the urease active sites are filled with Ni, but that is sufficient for full acid resistance. Under similar conditions (no added Ni), we still see a significant difference in the Ni load between the wild-type and Δmsr mutant, which further suggests a role for Msr in protecting urease maturation. In addition, HOCl is a highly potent oxidant (much more potent than H2O2) and it can achieve levels of 5 mM at

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from oxygen-stressed cells. Taken together, our results indicate that non-lethal elevated oxygen (∼21% partial pressure) caused a urease maturation defect.

Interestingly, the *H. pylori* urease maturation proteins have been reported to assemble at the cytoplasmic membrane in a pH-dependent manner when urease is also undergoing increased Ni-dependent maturation [51,52]. *H. pylori* Msr has also been shown to be membrane associated [15], so studies of Msr–UreG recognition (and perhaps roles of membrane proteins) as a function of pH may be informative. Although UreG complexes with UreE to function in Ni delivery [53,54], UreE does not seem to be a target for Msr; cross-linking approaches to identify such an interaction between the two proteins were negative [17]. Also, UreE is not a methionine-rich protein (∼1.8% methionine), and other Msr targets are rich in methionine. In the present study, we found that UreG is sensitive to inactivation by H₂O₂ and, given that UreG is rich in methionine residues (4.5%), we speculated that it was inactivated due to methionine oxidation. We found that H₂O₂ greatly inhibited UreG GTPase activity compared with the untreated UreG sample. It is likely that the bulk of inactivation of enzyme activity is due to Met-SO formation since the other residues susceptible to oxidation (tyrosine and cysteine) occur in small amounts within UreG (1.5% each) and Msr (known to repair methionine residues only) restored the oxidized protein’s GTPase activity.

In *H. pylori*, Msr was able to restore full UreG enzyme activity. This is similar to Msr-mediated catalase repair in which Msr restored up to 82% of full enzyme activity [16]. All nine methionine residues in UreG showed susceptibility to oxidation, and all were repaired with the addition of Msr. Interestingly, almost all (eight out of nine) methionine residues in UreG are predicted to be surface- or solvent-exposed. This is consistent with evidence that exposed methionine residues are more readily oxidized by oxidants and would be expected to be accessible to Msr [55–58]. A few prior accounts of Met-SO protein repair upon oxidative inactivation have been demonstrated in other systems such as *E. coli* L12 protein [59] and *H. pylori* catalase [16]. Oxidation and/or repair of every methionine residue within an enzyme has not been shown previously in any other system. For *E. coli* GroEL, 12 out of 23 methionine residues were converted into Met-SO [42]. Some activity (70% compared with untreated) of GroEL was restored by Msr when the lowest oxidant concentration was used, but harsher oxidation led to the production of some methionine-sulfone residues. In this case no activity could be restored [40]. Analysis of oxidized or Msr-repaired *E. coli* Ffh showed that four out of five methionine-containing peptides were oxidized and repaired [41]. Not all methionine residues of *H. pylori* catalase were susceptible to HOCl-mediated oxidation, but those that were indeed became targets of repair [16]. Our experiments expand our understanding of the unusual recognition flexibility of Msr, in that all nine methionine residues are repaired, and another target enzyme with a defined physiological role in pathogenesis is identified.

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**REFERENCES**


**AUTHOR CONTRIBUTION**

Lisa Kuhns, Manish Mahawar and Robert Maier conceived and designed the experiments. Joshua Sharp performed the MS. Stéphane Benoît constructed the UreG overexpression plasmid and assisted with atomic absorption spectrometry. Lisa Kuhns performed all the experiments. Lisa Kuhns and Robert Maier interpreted the data and wrote the paper.

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