

Noncatalytic Antioxidant Role for Helicobacter pylori Urease

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ABSTRACT The well-studied catalytic role of urease, the Ni-dependent conversion of urea into carbon dioxide and ammonia, has been shown to protect Helicobacter pylori against the low pH environment of the stomach lumen. We hypothesized that the abundantly expressed urease protein can play another noncatalytic role in combating oxidative stress via Met residue-mediated quenching of harmful oxidants. Three catalytically inactive urease mutant strains were constructed by single substitutions of Ni binding residues. The mutant versions synthesize normal levels of urease, and the altered versions retained all methionine residues. The three sitedirected urease mutants were able to better withstand a hypochlorous acid (HOCI) challenge than a $\Delta ureAB$ deletion strain. The capacity of purified urease to protect whole cells via oxidant quenching was assessed by adding urease enzyme to nongrowing HOCI-exposed cells. No wild-type cells were recovered with oxidant alone, whereas urease addition significantly aided viability. These results suggest that urease can protect H. pylori against oxidative damage and that the protective ability is distinct from the well-characterized catalytic role. To determine the capability of methionine sulfoxide reductase (Msr) to reduce oxidized Met residues in urease, purified H. pylori urease was exposed to HOCI and a previously described Msr peptide repair mixture was added. Of the 25 methionine residues in urease, 11 were subject to both oxidation and to Msr-mediated repair, as identified by mass spectrometry (MS) analysis; therefore, the oxidant-quenchable Met pool comprising urease can be recycled by the Msr repair system. Noncatalytic urease appears to play an important role in oxidant protection.

IMPORTANCE Chronic *Helicobacter pylori* infection can lead to gastric ulcers and gastric cancers. The enzyme urease contributes to the survival of the bacterium in the harsh environment of the stomach by increasing the local pH. In addition to combating acid, *H. pylori* must survive host-produced reactive oxygen species to persist in the gastric mucosa. We describe a cyclic amino acid-based antioxidant role of urease, whereby oxidized methionine residues can be recycled by methionine sulfoxide reductase to again quench oxidants. This work expands our understanding of the role of an already acknowledged pathogen virulence factor and specifically expands our knowledge of *H. pylori* survival mechanisms.

KEYWORDS gastric pathogen, gastritis, methionine, oxidation, oxidative stress, ulcer

Persistent *Helicobacter pylori* infections can cause chronic gastritis, peptic ulcer disease, and gastric cancer (1–4). Decades of immune response to the infection can lead to chronic inflammation and tissue damage (5, 6). *H. pylori* must first survive the harsh conditions of stomach gastric acid and then a prolonged host immune response after its colonization of the gastric mucosa (5). To combat the low pH of the stomach lumen, *H. pylori* expresses urease, which catalytically converts urea into ammonia and carbon dioxide, the former enabling *H. pylori* to resist the acidic gastric environment (7–9). Urease has been shown to be the most highly expressed protein in *H. pylori*,

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making up to 10% of the total protein content of the gastric pathogen (10). The enzyme consists of two structural proteins, UreA and UreB, with 12 subunits each that form a 1.1-MDa dodecamer (11). The UreAB heterodimer contains 25 methionine (Met) residues, 3% of the total amino acid content (12). Enzyme activity requires the addition of two nickel atoms per UreB monomer, and such maturation is facilitated by the urease accessory proteins UreEFGH and the hydrogenase maturation proteins HypAB (10, 13–15).

Although the catalytic activity of urease is required for initial colonization, (10, 16–18) several studies also indicate that non-acid-neutralizing roles may exist for urease. For instance, urease has been shown to be required for persistence in the mouse gastric mucosa, where the pH approaches neutrality (16, 19) and a urease-negative strain was unable to colonize a pH neutral piglet stomach (20). In addition, using a tetracycline-inducible system to turn on or off urease expression, Debowski et al. were able to show that urease-expressing bacteria were selected over time in a murine infection model, indicating that urease is required for chronic infection (16). Additionally, *H. pylori* urease has been shown to play multiple roles in modulating the host immune response. Indeed, it was shown to decrease opsonization (21), stimulate the chemotaxis of neutrophils and monocytes (22), induce apoptosis in gastric epithelial cells after binding to class II major histocompatibility complex (MHC) receptors (23), and induce proinflammatory cytokines (24). Recently, urease has been linked to the formation of gastric carcinoma through an ability to promote angiogenesis (25).

After reaching the gastric epithelium, *H. pylori* triggers responses by the host innate immune cells, which respond to the infection by generating reactive oxygen species (ROS) such as superoxide anion (O_2^{-}) , hydrogen peroxide (H_2O_2) , hydroxyl radicals (OH), and hypochlorous acid (HOCl) (10, 26, 27). Indeed, the exposure of gastric cells (28) or phagocytes (29) to *H. pylori* increases host cell ROS production. Patients with *H. pylori* infections have been shown to have larger amounts of ROS in their gastric mucosa (30). ROS can damage protein, DNA, and lipids (31). With regard to proteins, the amino acids most susceptible to oxidation are Met and cysteine (Cys) due to their sulfur-containing ligands (32).

H. pylori has many mechanisms to protect itself from, as well as repair damage caused by, oxidative stress. For instance, catalase and superoxide dismutase act to convert H_2O_2 and O_2^- into less harmful products. Catalase was recently shown to protect *H. pylori* against oxidative damage via an oxidant-quenching mechanism of its Met residues (33). The reaction involves methionine sulfoxide reductase (Msr) to reduce Met-SO to Met (34, 35). Msr reduces oxidized methionines of damaged proteins and has been shown to restore function to the damaged protein (36, 37). Cross-linking and direct repair assays showed that *H. pylori* Msr has at least five repair target proteins (34, 38, 39). These include AhpC, UreG, GroEL, catalase, and a site-specific recombinase (SSR), but urease was not observed as a repair target in those studies. In the present study, we describe the evidence for a noncatalytic role for urease, which involves Met-S/Met-SO recycling. This role aids in protecting the pathogen against oxidation-mediated cell death.

(A preliminary account of this work was presented at the American Society for Microbiology Microbe, New Orleans, LA, 1 to 5 June 2017 [40].)

RESULTS

Construction of catalytically inactive urease. Strains that synthesize inactive urease (herein referred to as apo-urease) were constructed to determine if the catalytically inactive enzyme is sufficient to quench oxidants. A sucrose-kanamycin selectioncounterselection was used to make markerless single-amino-acid substitutions in the UreB subunit of urease. The three single substitutions (His136Ala, His138Ala, and Lys219Ala) were constructed to replace nickel binding residues known to be required for enzyme activity (41). As expected, the three site-directed strains (*ureB*^{H136A}, *ureB*^{H138A}, and *ureB*^{K219A} strains) lacked urease activity compared to the wild-type (WT) strain 43504 (Table 1). Next, we sought to verify that the lack of urease activity was not

TABLE 1 Urease	activity of H	I. pylori	wild-type,	urease	deletion,	and	site-directed	mutant
strains								

Strain	Urease activity (μ mol/min/mg) a
43504 (wild type)	4.65 ± 0.96^a
Δ <i>ureAB</i> strain	0.06 ± 0.01
ureB ^{H136A} strain	0.07 ± 0.02
ureB ^{H138A} strain	0.07 ± 0.01
ureB ^{K219A} strain	0.08 ± 0.01

^aUrease activity expressed in micromoles of NH_3 produced per minute per milligram of total protein. Results shown are the means \pm SDs from 3 to 8 independent experiments with assays performed in triplicate.

linked to a decrease or loss of urease, e.g., that the mutant strains were still making apo-urease. Since urease is the most abundant protein in *H. pylori*, both UreA (27 kDa) and UreB (62 kDa) structural subunits can readily be visualized on Coomassie-stained SDS-PAGE gels (Fig. 1A). UreAB levels in the three site-directed mutants were found to be similar or slightly less than WT levels (Fig. 1A). This was confirmed by immunoblot, using anti-UreA antiserum (Fig. 1B). No UreA-specific band was seen in the Δ *ureAB* deletion strain, our negative control. These results indicate that apo-urease is still synthesized in each of the three site-directed mutants.

Catalytically inactive urease confers resistance to HOCI. The three catalytically inactive strains, as well as the WT (positive control) and the *ΔureAB* deletion strain (negative control), were exposed to HOCI to determine their ability to quench an oxidant and survive. The cells were incubated with HOCI for 1 h, diluted, and the CFU were counted (Fig. 2). The catalytically inactive mutants were as resistant as the WT and significantly (P < 0.01) less sensitive to HOCI oxidative damage than the urease gene deletion strain (Fig. 2). This suggests that the presence of urease, independent of its catalytic activity, can play a role in preventing cell death due to oxidative damage.

Both holo-urease and apo-urease protect against HOCI-mediated killing. To determine if holo-urease (purified from WT strain 43504) and apo-urease (purified from either strain *ureB*^{H138A} or strain *ureB*^{K219A}) were able to protect against HOCI-mediated oxidative damage, each purified urease was preincubated with the oxidant for 15 min and the mixture was added to whole nongrowing WT or Δ *ureAB* cells. Cell survival was then monitored and compared to that with a HOCI-only treatment (no protein) (Fig. 3). While no cells were recovered from the HOCI treatment alone (detection limit, 10² CFU/mI), when HOCI was preincubated with purified urease (both holo and apo), the cell recovery levels reached those of cells without HOCI treatment (Fig. 3). This suggests



FIG 1 Urease in cell-free and whole-cell extracts of WT and *ureAB* mutants. (A) SDS-PAGE (12.5%) analysis of cell-free extract. (B) Immunoblot analysis. Whole-cell extracts were separated via 12.5% SDS-PAGE, transferred to a nitrocellulose membrane, and blotted with anti-UreA antibodies. Strains are indicated above each lane. UreA and UreB are indicated.



FIG 2 Apo-urease confers resistance to HOCI. Cells grown on BA plates with 10 μ M NiCl₂ were incubated with PBS or with PBS supplemented with 100 μ M HOCI for 1 h at 37°C, diluted, and plated. CFU were counted after 72 h at 37°C. Error bars indicate standard deviations from 4 to 10 independent experiments.

that purified urease itself can quench HOCI, irrespective of urease activity. Interestingly, the $\Delta ureAB$ strain, shown to be more sensitive than the WT to HOCI challenge (Fig. 2), was almost fully protected against HOCI that had been preincubated with purified urease (Fig. 3). An additional protein, UreE, with a low Met content (1%) was shown to not protect against the HOCI challenge, suggesting that Met content might contribute to the protective effect.

MS/MS identification of oxidized and reduced methionine residues in UreAB. Urease is abundantly expressed, is found outside cells as well as in the cytosol, and contains many Met residues (25 of 807 residues, accounting for 3% of the total residues) (10, 42, 43). To identify which Met residues of urease (UreAB) are oxidized and repaired, we used liquid chromatography-tandem mass spectrometry (LC-MS/MS). The urease holoenzyme was first oxidized with HOCl, and then a sample of oxidized urease was subsequently incubated with an Msr repair mixture. Oxidized, oxidized and repaired, and untreated samples were analyzed by LC-MS/MS. Of the 25 methionine residues in urease, 19 were detected by LC-MS/MS (Fig. 4). UreA Met¹ and UreB Met¹, Met¹⁰⁴, Met¹⁸², Met¹⁹¹, and Met⁴⁷⁸ were either not detected or could not be quantified reliably. An examination of the protein sequence shows that four of these methionines reside in peptides containing multiple basic residues that were digested by trypsin resulting in small hydrophilic fragments, making their detection by LC-MS/MS highly unlikely. A fifth methionine-containing peptide was not included in the data analysis, as the signal-to-noise ratio was far too low for reliable quantitation. Most of the detected Met residues in the unoxidized samples had low levels of oxidation (less than 0.15 average oxidation per peptide) except UreA Met⁷¹, Met⁷⁷, and Met⁸⁶ and UreB Met²⁶². HOCItreated Met residues UreA Met⁴⁴, Met⁷¹, Met⁷⁷, and Met⁸⁶ and UreB Met¹², Met²⁶²,



FIG 3 Holo- and apo-urease proteins protect against HOCI. Wild-type and Δ *ureAB* 43504 cells were incubated for 1 h in PBS, PBS with HOCI, or PBS with HOCI that had been previously incubated for 15 min at 37°C with either purified UreAB^{WT}, UreAB^{H138A}, UreAB^{K219A}, or UreE as indicated below each column. Final protein concentration was 0.25 μ M, and final HOCI concentration was 200 μ M in each reaction. Error bars indicate standard deviations from 3 to 9 independent experiments. ND, no CFU could be detected (detection limit, 10² CFU/mI).



FIG 4 LC-MS/MS analysis of oxidized and Msr-repaired urease. Purified urease was incubated with a 60-fold molar excess of HOCI for 15 min. Excess HOCI was quenched with 15 mM Met and removed via dialysis. After dialysis, oxidized urease was incubated with purified Msr, Trx1, TrxR, NADPH, and DTT for 2 h at 37°C. Urease samples were digested by trypsin, and oxidation levels of methionine residues were quantified by LC-MS/MS. The oxidation level for the untreated sample was below the limit of detection for UreA Met⁵⁸. White bars, untreated; black bars, HOCI treated; gray bars, treated with HOCI plus Msr repair mixture. Error bars indicate standard deviations from 2 independent experiments with 3 replicates each.

Met³¹⁵, Met³¹⁷, Met³¹⁹, Met³⁵³, and Met³⁶⁶ had greater than 0.8 average oxidation per peptide. UreA Met¹² and UreB Met⁴⁴⁸ showed approximately 0.5 average oxidation per peptide, and all other residues showed less than 0.5 average oxidation per peptide. After HOCl-oxidized urease was treated with an Msr repair mixture, most Met residues were reduced to levels similar to that observed for the unoxidized sample, with the exception of UreB Met¹², Met²⁶², Met³⁵³, and Met³⁶⁶. Most of the digested peptides contained a single Met residue, the exceptions being UreA peptide 63 to 92 and UreB peptide 289 to 326, which contain three Met residues each (Met⁷¹, Met⁷⁷, and Met⁸⁶ and Met³¹⁵, Met³¹⁷, and Met³¹⁹, respectively). Likewise, UreB peptide 339 to 368 contains two Met residues (Met³⁵³ and Met³⁶⁶). For peptides with more than one Met residue, the oxidation of each Met cannot be accurately quantified by collision-induced dissociation (CID) MS/MS (44, 45). In summary, the Msr repair mixture reduced 11 of the oxidized urease can be reduced by Msr and thus probably recycles the oxidant-quenchable pool of urease.

DISCUSSION

Previous studies have focused on the catalytic by-products of urea hydrolysis by urease (NH_3 and CO_2) and their contribution to pathogen virulence. The ammonia produced by the enzyme supplements nitrogen metabolism, promotes alkalinization, and contributes to damage of host cell tissue (46). Additionally, carbon dioxide has been shown to protect the bacterium against highly toxic peroxynitrite during *H. pylori* infection (47). Now, there is growing evidence that urease plays noncatalytic roles in *H. pylori* infection. Indeed, urease is required for persistence in the mouse gastric epithelium, where the pH approaches neutrality (16, 19). Urease has been shown to play roles in modulating the immune response to infection (21–24), and *ureA* was found to be upregulated in response to oxidative stress (48).

We demonstrate that urease has a previously undescribed role as an antioxidant, thus adding to the growing number of noncatalytic roles that urease can play. Indeed, some of the observations cited above (e.g., urease requirement for persistence even at neutrality) may be due to an antioxidant role. This role depends at least in part on the primary sequence of the protein (e.g., Met residues). Although gastric acid exposure is an impressive obstacle to overcome, *H. pylori* also survives a barrage of host-produced ROS, including HOCI, which can reach levels as high as 5 mM at inflamed sites (49). *H. pylori* survives this via multiple mechanisms of detoxification and repair, enabling the infection to oftentimes persist throughout the life span of the host. Nickel-deficient mutants that lacked catalytic activity were as resistant as the WT to HOCI-mediated oxidative stress. In contrast, a urease deletion mutant was much more sensitive than the WT to HOCI-mediated oxidative stress. Thus, catalytic activity is not required for combating the oxidant. Rather, we propose that the Met residues of urease can quench oxidants, forming Met-SO in the process. Catalase, another abundant protein, was similarly shown to quench oxidants to aid bacterial survival, and the mechanism was independent of catalytic activity (33). A similar antioxidant mechanism of relying on Met residues was assigned to pure *E. coli* glutamine synthetase; however, the system was not shown to confer oxidative stress resistance to cells (50).

We predict that Met-SO residues of urease could be reduced by Msr and then used again to quench oxidants. Indeed, Msr was previously shown to reduce and repair several H. pylori proteins such as AhpC, UreG, GroEL, catalase, and site-specific recombinase (SSR) (34, 38, 39). However, it was previously unknown whether Msr could reduce urease Met residues. Through mass spectrometry analysis, we found that of the 25 Met residues in urease, 11 were shown to be susceptible to oxidation and were reduced (i.e., repaired) by Msr. Thus, urease can be added to the list of Msr repair targets. Upon PDBePISA analysis of surface accessibility, using the X-ray crystal structure of H. pylori urease (Protein Data Bank code 1E9Z), 9 of the 11 Met residues that were oxidized and then subsequently reduced are surface exposed (11, 51). This is consistent with previous reports of surface methionine residues being subject to oxidation and subsequent repair by Msr (26, 38). H. pylori Msr deletion mutants showed lowered urease activity when exposed to oxidative stress $(21\% O_2)$ (38). This was attributed to the inability of the strain to repair UreG, a urease accessory protein required for urease activity; however, we can now hypothesize the lowered activity could also be due to the lack of reduction of oxidized Met residues of the structural proteins of urease (UreAB).

An Msr deletion mutant has been shown to be more sensitive to HOCl challenge than its parent strain (SS1) (39). Here, we show a high HOCl sensitivity of a strain lacking one of the described repair targets of Msr, namely, urease. Conclusions based on comparing the phenotypes of the $\Delta ureAB$ strain to that of the Δmsr strain (39) are not possible due to the use of different parent strains. Comparisons are further complicated by the fact that Msr has multiple repair targets besides urease.

While urease lacks a described secretion pathway in *H. pylori*, it has been found outside the cell, possibly due to cell lysis (42, 52). We show that purified urease added exogenously and whole cells expressing urease quench HOCl and survive HOCl challenge, respectively. Regardless of location, urease activity is not required for the described oxidant quenching effect. Seemingly, both extracellular and intracellular ureases likely act to quench oxidants. However, intracellular urease may play a larger role in protecting against oxidants because of its Met turnover—Msr is unlikely to be active extracellularly. Although this study focused on the role of Met residues in urease oxidant quenching, the Cys residues of urease might also contribute to the antioxidant ability we describe. While Cys residues are very sensitive to oxidation, the UreAB heterodimer only contains 4 Cys residues compared to 25 Met residues. Oxidized Cys residues could be reduced enzymatically through the thioredoxin/ thioredoxin reductase (Trx/TrxR) system (32). *H. pylori* contains two thioredoxins (Trx1 and Trx2), which have been shown to play a role in preventing macromolecule damage, and one TrxR; however, it is not known if the Trx system can repair oxidized Cys residues in *H. pylori* (53, 54).

Future site-directed mutagenesis studies of individual Met residues in urease could elucidate the role of individual residues in the quenching. Also, the use of the mouse model to gauge colonization by the catalytically inactive strains is likely to be a promising approach to assess *in vivo* function, although the acid barrier would have to be minimized to observe colonization by urease-activity-negative strains. Catalytically

TABLE 2 Strains and plasmids used in this study

		Source or
Strain or plasmid	Relevant characteristic(s) ^a	reference
E. coli TOP10	Cloning strain	Invitrogen
H. pylori		
43504	Parental strain	ATCC ^b
∆ <i>ureB</i> mutant	∆ <i>ureB</i> ::KSF Kan ^r Suc ^s	This study
∆ <i>ureAB</i> mutant	Δ <i>ureAB::cat</i> Cm ^r	15
ureB ^{H136A} strain	Codon for His136 replaced by Ala	This study
ureB ^{H138A} strain	Codon for His138 replaced by Ala	This study
ureB ^{K219A} strain	Codon for Lys219 replaced by Ala	This study
Plasmids		
pBluescript KS(+)	Cloning vector; Ap ^r	Stratagene
pKS- <i>ureB</i>	ureB cloned into the KpnI-SacI site of pBluescript KS(+)	This study
pKS- <i>ureB</i> ::KSF	KSF cassette cloned into the BamHI site of pKS-ureB	This study
рKS- <i>ureB</i> ^{H136A}	pKS-ureB with His136 replaced by Ala	This study
рКS- <i>ureB</i> ^{н138A}	pKS-ureB with His138 replaced by Ala	This study
pKS- <i>ureB</i> ^{K219A}	pKS-ureB with Lys219 replaced by Ala	This study

^aCm, chloramphenicol; Ap, ampicillin; Kan, kanamycin; Suc, sucrose; KSF, Kan^r-sacB-P_{flaA} (62).

^bAmerican Type Culture Collection, Manassas, VA.

inactive urease may contribute to the persistence of *H. pylori* infections of the host gastric epithelium, where the pathogen is exposed to multiple oxidants (30).

Although the role of urease in *H. pylori* pathogenesis has been extensively studied, urease is also a virulence factor for several pathogens, including *Proteus mirabilis* and *Klebsiella pneumoniae* (55). During *P. mirabilis* infection of the urinary tract, urease is required for mineral precipitation to form urinary stones, which impede the elimination of the infection (56). During mice gastrointestinal tract infections, urease-negative strains of *K. pneumoniae* were outcompeted by WT strains, and so it was suggested that the ammonia produced by urease serves as a supplementary nitrogen source (57). *K. pneumoniae* and *P. mirabilis* must also defend against HOCI produced by neutrophils as a part of the innate immune response to infection (56, 58). These species also contain Msr, enabling the possibility that a similar method for urease methionine oxidation and repair by Msr would be found in additional pathogens. Additionally, other highly expressed proteins that are Met rich might play a similar antioxidant role. Indeed, catalase, a Met-rich and abundant *H. pylori* protein, has been shown to have similar noncatalytic antioxidant properties (33).

Urease and Msr can be found in plants, fungi, archaea, and bacteria (59, 60). Throughout the three domains of life, urease is increasingly seen as a "moonlighting protein" with a variety of noncatalytic roles, including fungicidal, insecticidal, and proinflammatory effects (61). For *H. pylori*, an antioxidant mechanism based on urease Met residue recycling is expected to explain at least part of the *in vivo* requirement for this important enzyme.

MATERIALS AND METHODS

Growth conditions. *Escherichia coli* cells were cultured at 37°C aerobically in Luria-Bertani (LB) broth or on agar LB plates with 100 μ g/ml ampicillin and 30 μ g/ml chloramphenicol as needed. *H. pylori* cells (ATCC 43504) were routinely grown at 37°C on brucella agar plates supplemented with 10% defibrinated sheep blood (BA) under microaerophilic conditions (4% O₂, 5% CO₂, and 91% N₂). As needed, *H. pylori* was grown on BA supplemented with 5 or 10 μ M NiCl₂, 5% sucrose, 20 μ g/ml kanamycin, or 30 μ g/ml chloramphenicol. For urease protein purification, *H. pylori* cells were grown on BA plates supplemented 5 μ M NiCl₂.

Strain construction. The *E. coli* and *H. pylori* strains and plasmids used in this study are listed in Table 2. Site-directed mutants were constructed using a kanamycin-sucrose selection-counterselection system described previously (33, 39, 62). Briefly, a 1,400-bp DNA fragment containing a partial *ureB* gene was amplified by PCR using primers UreBF1 and UreBR1 (Table 3) from the chromosomal DNA of *H. pylori* 26695 (12) and cloned into pBluescript KS(+) to obtain pKS-*ureB*. The Kanr-*sacB*-P_{*haA*} (KSF) cassette (62) was inserted into *ureB* resulting in pKS-*ureB*::KSF, which was then naturally transformed into WT strain 43504, thereby replacing parent *ureB* with *ureB*::KSF. The resulting strain ($\Delta ureB$::KSF strain) was found to be urease negative and kanamycin resistant. Site-directed mutations were introduced to pKS-*ureB* by

Name	Sequence $(5' \rightarrow 3')^a$	Restriction site
UreBF1	GCTTCGGTACCCCTGAGAGAAGGCATGAG	Kpnl
UreBR1	GCATCGAGCTCCCGGCAACACTTGTCT	Sacl
UreB:H136AF	GTGGTATTGACACAGCTATCCACTTCATTTCAC	
UreB:H136AR	GTGAAATGAAGTGGATAGCTGTGTCAATACCAC	
UreB:H138AF	ATTGACACACACATCGCCTTCATTTCACCCC	
UreB:H138AR	GGGGTGAAATGAAGGCCATGTGTGTGTCAAT	
UreB:K219AF	GGTGCGATTGGCTTTGCAATCCACGAAGACTGG	
UreB:K219AR	CCAGTCTTCGTGGATTGCAAAGCCAATCGCACC	

TABLE	3	Primers	used	in	this	paper
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^aPrimers were purchased from Integrated DNA Technology, Coralville, IA.

overlapping PCR using internal primers UreB:H136AF, UreB:H136AR, UreB:H138AF, UreB:H138AR, UreB: K219AF, and UreB:K219AR and external primers UreBF1 and UreBR1 (Table 3). The resulting plasmids (pKS-*ureB*^{H136A}, pKS-*ureB*^{H138A}, and pKS-*ureB*^{K219A}) were introduced into the Δ *ureB*::KSF strain by natural transformation. UreB site-directed mutants were selected on BA plates containing 5% sucrose. The resulting strains were sucrose resistant and urease negative. The Δ *ureAB* strain was constructed as described previously (15). All plasmids and PCR products were sequenced at the Georgia Genomics Facility, University of Georgia, Athens, GA, and compared with DNA sequences from strain 26695 (12) to ensure that no error had been introduced following PCR amplification, as well as to verify the presence of engineered site-directed mutations within *ureB*.

Urease assays. Whole cells grown on BA plates under microaerophilic conditions at 37° C were resuspended in 50 mM HEPES (pH 7.5), lysed by sonication (Heat Systems Ultrasonics sonicator: 30 s at 4-W output power and 50% duty cycle), and centrifuged at 21,100 \times *g* for 5 min. The cell-free supernatant was assayed for urease activity following the phenol-hypochlorite (Weatherburn) method (63). Protein concentration was determined using the bicinchoninic acid (BCA) kit (Thermo Scientific Pierce).

SDS-PAGE and immunoblotting. Whole cells grown on BA plates supplemented with 10 μ M NiCl₂ were resuspended in phosphate-buffered saline (PBS) and treated as above to obtain cell-free extracts (CFE). Total protein from CFE (0.4 μ g protein per lane) was loaded onto 12.5% SDS-PAGE gels and stained with Coomassie brilliant blue G-250 or transferred to a nitrocellulose membrane for immunoblotting as previously described with modifications (33, 38). Briefly, the nitrocellulose membrane was blocked with 3% gelatin in Tris-buffered saline (TBS; 50 mM Tris [pH 7.6] and 150 mM NaCl [pH 7.6]). After blocking, the membrane was incubated with anti-UreA antibody (rabbit polyclonal; Santa Cruz Biotechnology) diluted 1:2,000 in TBS with 0.1% Tween 20 (TTBS) containing 1% gelatin for 1 h. The membrane was then washed with TTBS and incubated with goat anti-rabbit IgG conjugated to alkaline phosphatase (Bio-Rad) diluted 1:2,000 in TTBS with 1% gelatin. After 1 h of incubation, the blot was developed with Nitro Blue Tetrazolium–5-bromo-4-chloro-3-indolyl phosphate in 10 mM Tris-HCl (pH 9.5) and 150 mM NaCl.

Whole-cell HOCI tolerance assay. Whole-cell HOCI tolerance assays were performed as previously described (33) with the following modification. Briefly, cells grown on BA plates with 10 μ M NiCl₂ were resuspended in PBS to an optical density at 600 nm (OD₆₀₀) of approximately 1.1 and then mixed with either PBS or PBS with NaOCI (100 μ M final concentration, 10 to 15% available chlorine [referred to as HOCI]) for 1 h at 37°C. The HOCI-challenged cells were diluted in PBS and plated on BA plates, and the CFU were counted after 72 h of incubation at 37°C under microaerophilic conditions.

Purified protein HOCI tolerance assay. HOCI tolerance assays with purified proteins were performed as previously described (33). Briefly, wild-type and $\Delta ureAB$ 43504 cells (OD₆₀₀ of 1.3) were incubated for 1 h in PBS, PBS with 200 μ M HOCI, or PBS with HOCI that had been previously incubated with 0.25 μ M UreAB or UreE for 15 min at 37°C. After the PBS or HOCI challenge, the cells were plated on BA plates and the CFU were counted after 72 h of incubation at 37°C under microaerophilic conditions.

Protein purification. *H. pylori* 43504 cells were grown on BA plates supplemented with 5 μ M NiCl₂ under microaerophilic conditions (4% O₂, 5% CO₂, 91% N₂) at 37°C and broken by passage through a French pressure cell at 18,000 lb/in² or by cell disruption at 20,000 lb/in² (Constant Systems One Shot). Cell debris was removed by centrifugation at 4,500 × *g*. Urease was purified from the resulting supernatant (cell-free extract) using fast protein liquid chromatography (AKTA; GE Healthcare). Cell-free extract was loaded onto a Q-Sepharose HiTrap column that had been equilibrated with 50 mM NaPO₄ (pH 7.2) and 25 mM NaCl (buffer A). The protein was eluted with a linear gradient of 0.025 to 1 M NaCl in buffer A. Urease-containing fractions, as determined by the phenol red urease assay (64), were concentrated with Amicon Ultra-4 devices with a 10-kDa molecular mass cutoff (Merck Millipore). The concentrated urease was further purified with a HiLoad 16/60 Superdex 75 column in buffer A with 0.35 M NaCl. Urease-containing fractions, were identified and concentrated as above, and the final protein concentration was determined using the BCA kit (Thermo Scientific Pierce). UreAB proteins were purified to near homogeneity (>90%), as determined by SDS-PAGE. UreE was purified as previously described (65).

Urease oxidation and Msr repair. Urease oxidation and subsequent Msr repair were performed as previously described with modifications (26). Briefly, purified urease (10 μ M) was incubated with a 60-fold molar excess of HOCI for 15 min at room temperature in the dark. HOCI was quenched with excess Met (15 mM final) for 30 min on ice. Excess Met and HOCI were removed via dialysis (Spectra/Por; molecular weight cutoff [MWCO], 12,000 to 14,000) in 50 mM sodium phosphate buffer.

Oxidized and quenched urease was incubated with 6.2 μ M Msr, 5 μ M Trx1, 0.5 μ M TrxR, 400 μ M NADPH, and 100 μ M dithiothreitol (DTT) for 2 h at 37°C. The reaction was stopped by incubating at 95°C for 5 min.

Sample preparation and mass spectrometry analysis. Concentrated ammonium bicarbonate buffer was added to the protein to adjust to a final buffer concentration of 50 mM (pH 8.0). The samples were denatured by heating at 95°C for 20 min in the presence of 10 mM DTT. After cooling the sample, 20 mM iodoacetamide was added and the sample was kept at room temperature in dark for 45 min. Next, 20 mM DTT was added to quench the excess iodoacetamide and the mixture was incubated for 20 min at room temperature. Finally, the samples were digested with sequencinggrade trypsin at a 1:20 ratio of protease to protein by incubating at 37°C overnight with rotation. The samples, after trypsin digestion, were acidified with 0.1% formic acid and spun briefly, and the supernatant was transferred to sample vials. An Orbitrap Fusion mass spectrometer (Thermo Fisher Scientific) coupled to an Ultimate 3000 high-performance liquid chromatography (HPLC) system was used for the nano-LC-MS/MS analysis. The samples were first loaded onto a trap cartridge containing C_{18} stationary phase to desalt the samples. After 4 min, the samples were back-eluted from the trapping cartridge onto a nano PepMap column (75-μm inner diameter, 150-mm length, 3-μm particle size). Mobile phase A consisted of water with 0.1% formic acid, and mobile phase B consisted of acetonitrile with 0.1% formic acid. The peptide separation was achieved with a linear binary gradient from 0 to 40% phase B for 25 min at 0.3 μ l/min. The peptides were fragmented by collision-induced dissociation, and the normalized collision energy was set to 35%. The MS scan range was m/z 150 to 2,000, and the top five peaks were selected in precursor scan for the data-dependent CID fragmentation in each cycle. The data files were searched against the protein sequences of UreA and UreB of H. pylori strain 26695. The peak intensities of the unoxidized peptides and their corresponding methionine oxidation products observed in LC-MS were used to calculate the average oxidation events per peptide in the sample. The ion intensity of the oxidized peptides was multiplied by the number of oxidation events required for the mass shift (e.g., one oxidation on methionine for +16, two events for +32) and then divided by the sum of the ion intensities of all unoxidized and oxidized peptide masses as represented by the equation below.

 $P = [I(+16) \text{ oxidized } \times 1 + I(+32) \text{ oxidized } \times 2 + I(+48) \text{ oxidized } \times 3 + \dots]$

[*I*unoxidized + I(+16)oxidized + I(+32)oxidized + I(+48)oxidized ...]

In this equation, *P* represents the average oxidation level of the peptide and *I* represents the peak intensities of oxidized and methionine unoxidized peptides.

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