Complex Responses to Hydrogen Peroxide and Hypochlorous Acid by the Probiotic Bacterium *Lactobacillus reuteri*

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**ABSTRACT** Inflammatory diseases of the gut are associated with increased intestinal oxygen concentrations and high levels of inflammatory oxidants, including hydrogen peroxide (H$_2$O$_2$) and hypochlorous acid (HOCl), which are antimicrobial compounds produced by the innate immune system. This contributes to dysbiotic changes in the gut microbiome, including increased populations of proinflammatory enterobacteria (*Escherichia coli* and related species) and decreased levels of health-associated anaerobic Firmicutes and Bacteroidetes. The pathways for H$_2$O$_2$ and HOCl resistance in *E. coli* have been well studied, but little is known about how commensal and probiotic bacteria respond to inflammatory oxidants. In this work, we have characterized the transcriptomic response of the anti-inflammatory, gut-colonizing Gram-positive probiotic *Lactobacillus reuteri* to both H$_2$O$_2$ and HOCl. *L. reuteri* mounts distinct but overlapping responses to each of these stressors, and both gene expression and survival were strongly affected by the presence or absence of oxygen. Oxidative stress response in *L. reuteri* required several factors not found in enterobacteria, including the small heat shock protein Lo18, polyphosphate kinase 2, and RsIR, an *L. reuteri*-specific regulator of anti-inflammatory mechanisms.

**IMPORTANCE** Reactive oxidants, including hydrogen peroxide and hypochlorous acid, are antimicrobial compounds produced by the immune system during inflammation. Little is known, however, about how many important types of bacteria present in the human microbiome respond to these oxidants, especially commensal and other health-associated species. We have now mapped the stress response to both H$_2$O$_2$ and HOCl in the intestinal lactic acid bacterium *Lactobacillus reuteri*.

**KEYWORDS** oxidative stress, probiotics, transcriptomics
changes in the microbiome and the symptoms of disease (1, 13), indicating that manipulating gut bacteria is an important element in controlling these diseases.

Probiotics are live microorganisms which, when consumed in sufficient quantities, have a measurable health benefit (14), and a variety of different probiotic bacteria have been shown to have anti-inflammatory effects in the gut (15, 16). The most commonly used probiotics are lactic acid bacteria of the genus *Lactobacillus* (17), which are able to both modulate the host immune system and outcompete enterobacterial pathogens (15) and some strains of which have been shown to improve outcomes for inflammatory bowel diseases in both humans and animal models (18–20). The effectiveness of probiotics for treating inflammation in the gut, however, may be limited by their ability to survive attack by the overactive host immune system, including the oxidative damage caused by ROS and RCS. While the general stress response physiology of lactic acid bacteria has been relatively well characterized (21), bacterial responses to oxidative stress are best understood for *E. coli* and related inflammation-enriched enterobacteria (22–25). This is especially true of RCS, including hypochlorous acid (HOCl) and reactive chloramines, which are extremely potent antimicrobial compounds produced by the neutrophil enzyme myeloperoxidase (22, 26–28). Relatively little is known about how health-associated probiotic and commensal bacteria sense and respond to inflammatory oxidants (21, 29–31).

*Lactobacillus reuteri* is a well-established model probiotic bacterium that is able to stably colonize the mammalian intestine (32, 33), where several strains have been shown to combat inflammation and enteric infections by different mechanisms, including anti-inflammatory histamine synthesis by strains ATCC PTA 6475 and ATCC PTA 5289 (34–37), modulation of immune cell functions by strains ATCC PTA 6475, 100-23, and WU (33, 38–40), and production of antimicrobial compounds (e.g., reuterin and reutericyclin) by ATCC PTA 6475 and many other strains (37, 41, 42). While the genome-wide stress responses of *L. reuteri* to low pH (strain ATCC 23272) (43) and bile salts (strains ATCC PTA 6475 and 23272) (44, 45) have been characterized, little is known about how members of this species respond to ROS, and nothing is known about how *L. reuteri* or any other lactic acid bacterium senses or responds to RCS. The *L. reuteri* genome encodes neither catalase nor superoxide dismutase (46). The oxidative stress repair enzyme methionine sulfoxide reductase (47) is induced by and required for gut colonization by *L. reuteri* strain 100-23 (48, 49), indicating that resistance to oxidative damage is important in vivo. A cysteine-dependent pathway contributing to H$_2$O$_2$ and O$_2$ tolerance has been identified in strain BR11 (50) but did not appear to play a role in the ability of *L. reuteri* BR11 to prevent colitis in mice (51).

In this work, we have taken a transcriptomic approach to characterize genome-wide H$_2$O$_2$- and HOCl-dependent gene regulation in *L. reuteri* ATCC PTA 6475 and to identify genes involved in resistance to killing by these stressors (52), with the goal of finding genes and pathways distinct from those found in the enterobacteria. Our results show that despite not containing close homologs of any of the known RCS-specific transcription factors (22, 53–56), *L. reuteri* is able to mount clearly different stress responses to H$_2$O$_2$ and HOCl and that the presence of O$_2$ has dramatic effects on both gene regulation and survival in response to these stresses. We also identified roles for several genes in surviving H$_2$O$_2$- and HOCl-mediated stress, including those encoding methionine sulfoxide reductase (47), polyphosphate kinase 2 (57, 58), and the lactic acid bacterium-specific small heat shock protein Lo18 (59–61), as well as a role in surviving H$_2$O$_2$ stress for RsIR, previously characterized as an *L. reuteri*-specific regulator of histamine synthesis (35).

### RESULTS AND DISCUSSION

**Growth of *L. reuteri* is inhibited by inflammatory oxidants.** To begin characterizing the response of *L. reuteri* to inflammatory oxidants, we treated anaerobically growing cultures with different concentrations of H$_2$O$_2$ (Fig. 1A) and HOCl (Fig. 1B). *L. reuteri* growth was more sensitive to H$_2$O$_2$ than HOCl, with an increase in culture density of less than 0.1 $A_{600}$ unit 5 h after 0.96 mM H$_2$O$_2$ or 5 mM HOCl treatment. Since
we were interested in characterizing gene regulation during a successful, productive response to bacteriostatic stress, we selected concentrations of 0.12 mM H$_2$O$_2$ and 1.25 mM HOCl for further analysis. These concentrations resulted in reductions in growth rate after stress treatment to 80% of that of untreated cells, followed by complete recovery (Fig. 1A and C). Growth was significantly inhibited by 0.12 mM H$_2$O$_2$ between 1 and 4 h after treatment and by 1.25 mM HOCl between 1.5 and 3 h after treatment (Fig. 1B and D). The growth rates of untreated cells differed between these experiments (0.31 division h$^{-1}$ for the H$_2$O$_2$ treatment experiment and 0.39 division h$^{-1}$ for the HOCl experiment), which we hypothesize may reflect batch-to-batch variations in the MEI-C medium (see Materials and Methods) used. These concentrations of oxidants had no significant effect on cellular NAD$^+$/NADH ratios (Fig. 1E and F and Fig. S1), indicating that the bacteriostatic effects of these H$_2$O$_2$ and HOCl concentrations did not involve major disruptions to the redox state of the bacterial cells.

**Transcriptomic analysis of H$_2$O$_2$ and HOCl response by *L. reuteri*.** We next treated anaerobically growing *L. reuteri* with 0.12 mM H$_2$O$_2$ or 1.25 mM HOCl and used RNA sequencing to characterize the transcriptomes of stressed cells before and 5, 15, and 30 min after stress treatment (Fig. 2 and Table S1). Up to 27% of the *L. reuteri* genome was up- or downregulated (>2-fold; Bonferroni-corrected $P$ value $P_{Bonf} < 0.01$; up to 48% without the 2-fold restriction) after stress treatment (Table S2), and there were clear differences in the responses to H$_2$O$_2$ and HOCl, consistent with...
previous reports that bacterial responses to these oxidants are different (22–24). As shown in Fig. 2 and Table S2, the response to H$_2$O$_2$ involved roughly equal numbers of up- and downregulated genes (2-fold more upregulated at 5 min, 1.4-fold more downregulated at 15 min, and 1.1-fold more downregulated at 30 min), with an increase in the number of genes with significant changes in expression over the 30-min course of stress treatment (from 35 at 5 min to 557 at 30 min, a 16-fold increase). In contrast, HOCl treatment caused 2- to 3-fold more genes to be upregulated than downregulated at all time points, and there was a smaller increase in the number of genes with significant changes in gene expression over time (from 78 at 5 min to 148 at 30 min, a 2-fold increase), consistent with the very high reaction rate of HOCl with biological molecules (22, 27, 62). The differences between the H$_2$O$_2$ and HOCl stress responses were also reflected in principal-component analysis of the transcriptomic data (Fig. S2A), which clearly separated the H$_2$O$_2$- and HOCl-treated samples. The untreated samples from the
two experiments did not cluster as closely together as we expected, since these samples were ostensibly identical. To determine whether this reflected batch effects (possibly due to variations in the growth medium, as mentioned above) or inherent variation in expression levels for particular genes, we selected representative genes that had the same or significantly different levels of expression ($n$ each) in the untreated samples from the two transcriptome sequencing (RNA-seq) data sets (Fig. S2B and Table S1) and used quantitative reverse transcription-PCR (qRT-PCR) to measure their expression in independently prepared unstressed $L$. reuteri cultures. We found that the amounts of variation in expression were similar for all six genes analyzed, with no significant differences between experiments (Fig. S2C). To minimize the effect of the differences between untreated samples from the H$_2$O$_2$ and HOCl RNA-seq experiments, all of the untreated samples from both experiments ($n$ = 6) were considered a single group for the DeSeq2 analyses of RNA-seq data.

We next used qRT-PCR to validate our RNA-seq results (Fig. 3). We examined two genes strongly activated by both H$_2$O$_2$ and HOCl ($ahpF$ and $pcl1$, encoding alkylhydroperoxidase and a predicted iron transporter, respectively) and a gene repressed by both oxidants ($moeB$, encoding a subunit of molybdopterin synthase [63]) (Table S1) and confirmed the expected expression patterns. RT-PCR of the redox-responsive $perR$ and $sigH$ genes, encoding predicted transcription factors, also recapitulated the expres-

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**FIG 3** qRT-PCR validation of RNA-seq results for selected genes. (A) Log$_2$ fold change in gene expression for $ahpF$ (LAR_RS05795), $pcl1$ (LAR_RS08080), $moeB$ (LAR_RS05335), $perR$ (LAR_RS06970), $sigH$ (LAR_RS04695), and $rsiR$ (LAR_RS05165), from RNA-seq experiments (Table S1). Bonferroni-corrected $P$ values comparing each time point to expression at 0 min (untreated samples) are indicated (****, $P < 0.0001$). (B) $L$. reuteri ATCC PTA 6475 was grown anaerobically at 37°C to an $A_{600}$ of 0.3 to 0.4 in MEI-C and then treated with the indicated concentrations of H$_2$O$_2$ or HOCl or not treated ($n$ = 3, means ± SDs). qRT-PCR was used to measure log$_2$ fold changes in expression for each gene at the indicated time points. Two-way ANOVA with Holm-Sidak’s multiple-comparison test was used to determine differences between H$_2$O$_2$- or HOCl-treated samples and untreated samples at each time point (*, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$; ****, $P < 0.0001$).
Finally, we examined expression of rsiR, a known regulator of anti-inflammatory mechanisms in *L. reuteri* ATCC PTA 6475 (34, 35) that has also been reported to regulate some redox response genes (35), and we confirmed that its expression is not regulated by either H2O2 or HOCl.

To further characterize the differences and overlaps between the H2O2 and HOCl stress responses of *L. reuteri*, we plotted changes in gene expression under each tested condition against each other condition (Fig. 4 and Table S2). There were 78 genes that were upregulated (at least one time point) by both stressors (287 for H2O2 and 133 for HOCl) and 40 genes that were downregulated (at least one time point) by both stressors (346 for H2O2 and 69 for HOCl). As shown in Fig. 4, there were 12 genes (11 unique) that had reciprocal responses at the same time point (see diagonals of gray-outlined plots in Fig. 4). In general, there were 16 unique genes that were up in at least one time point in H2O2 and down in at least one time point in HOCl (or vice versa, vice versa).
versa). Furthermore, there were 73, 282, and 531 genes at 5, 15 and 30 min, respectively, that were significantly changed ($P_{\text{Bonf}} < 0.01$, 2-fold up or down) with one stressor but not the other, all of which suggests that, despite lacking homologs of known HOCl-sensing transcription factors (22, 53–55, 64, 65), *L. reuteri* has a sophisticated ability to distinguish between H$_2$O$_2$ and HOCl and differentially control transcription. This is consistent with results with *E. coli* and *Bacillus subtilis*, in which H$_2$O$_2$ and HOCl stress responses partially overlap but have substantial oxidant-specific components (22, 27, 56, 62, 66). Cluster analysis of genome-wide expression patterns (Fig. 5)
reinforced this result, and we were able to identify genes whose expression was controlled in very similar ways by the different oxidants as well as groups of genes with very different expression patterns in response to H2O2 and HOCl, including, for example, cgl and cyuABC, which encode a previously characterized cysteine-dependent redox stress response pathway (50). Examples of genes with distinct patterns of regulation are illustrated in the rightmost columns of Fig. 5.

*L. reuteri*s’ response to H2O2 was generally consistent with what has been previously observed with other catalase-negative Gram-positive bacteria (21, 23, 24, 31, 65, 67, 68). Highly upregulated genes included genes encoding alkylhydroperoxidase (*ahpCF*) (69), NADH oxidase (*noxB*) (70), and methionine sulfoxide reductase (*msrB*) (47), DNA repair genes (*uvrABD, xthA, and umuC*) (71), and genes for predicted metal transporters (*pcl1* and *pcl2*) and the peroxide-sensing transcription factor *PerR* (65). The response to HOCl was also, in broad strokes, similar to that of previously characterized bacteria (22), in that upregulated genes included those involved in proteostasis (*groSL, clpE, and hsp20/lo18*), metal stress (*pcl1, pcl2, and copAR*), thioredoxins (*trxABD*), and cysteine and methionine synthesis (*cysK* and *metE*). Genes upregulated by both stressors included not only *msrB, ahpCF, perR*, and the genes for predicted iron transporters (*pcl1* and *pcl2*) but also genes for a variety of predicted sugar and amino acid transporters and metabolic enzymes (*oxc*, encoding oxalyl-CoA decarboxylase (72), for example). These may represent responses to changes in the nutritional environment *L. reuteri* might encounter in the inflamed gut (6, 8).

**Redox-regulated transcription factors in *L. reuteri***. While many bacterial transcription factors that respond to H2O2 and/or HOCl have been described, *L. reuteri* encodes only a few homologs of known H2O2-detecting transcription factors (e.g., *PerR* and *VicK* (24, 68)) and no close homologs of any of the known HOCl-detecting transcription factors (22, 53–55, 64, 65). This suggested that among the 102 predicted transcription factors encoded by the *L. reuteri* genome, there are likely to be novel redox-sensing regulators. To begin to assess this possibility, we performed cluster analysis of the expression of genes encoding transcription factors that showed a significant change with either stressor at any time point (*n* = 73) under both stress conditions (Fig. 6), reasoning that many bacterial transcription factors are autoregulated and that changes in expression of transcription factors are useful signposts for identifying regulatory stress response networks (53, 66, 73). We found genes encoding predicted transcription factors whose expression was activated by both H2O2 and HOCl (e.g., *perR, spxA*, and LAR_RS09770), repressed by both H2O2 and HOCl (e.g., *kgdR* and *fabT*), activated only by H2O2 (e.g., *lexA* and LAR_RS07525), activated only by HOCl (e.g., *ctsR* and *copR*), repressed only by H2O2 (e.g., *sigH* and *rex*), and repressed only by HOCl (e.g., *malR3* and LAR_RS02755), indicating the presence of a complex regulatory response to both oxidants. Some of these regulators have known functions, which give useful insights into the in vivo effects of H2O2 and HOCl on *L. reuteri*. For example, only HOCl activated expression of *ctsR*, a conserved regulator of protein quality control in Gram-positive bacteria (74, 75), consistent with the known ability of HOCl to unfold and aggregate proteins (76, 77) and the activation of the heat shock response in many species of HOCl-stressed bacteria (22). On the other hand, only H2O2 activated expression of the DNA-damage responsive *lexA* regulator (71), consistent with the known ability of H2O2 to damage DNA (23) and suggesting that HOCl does not cause DNA damage at the concentration used in this experiment. However, most of the transcription factors in *L. reuteri* have no known function, and the expression patterns of many of these genes were affected by the redox stress treatments. For example, the only alternative sigma factor (78) encoded in the *L. reuteri* genome (*sigH*) was downregulated strongly by H2O2 but unaffected by HOCl. We do not currently know what genes these uncharacterized regulators regulate, what role(s) they may play in surviving redox stress, or what transcription factor(s) is responsible for HOCl-specific regulation in *L. reuteri*.
Oxygen affects H$_2$O$_2$- and HOCl-dependent gene expression in L. reuteri. We used quantitative RT-PCR to measure the dose responsiveness of changes in expression of selected genes in anaerobically grown L. reuteri 15 min after treatment with concentrations of H$_2$O$_2$ and HOCl at, above, and below the nonbactericidal concentrations used in previous experiments (Fig. 7A). Interestingly, the genes differed in their dose-response patterns, with moeB equally repressed at all H$_2$O$_2$ and HOCl concentrations, ahpF equally upregulated by all three H$_2$O$_2$ concentrations but activated more strongly by increasing doses of HOCl, and pcl1 upregulated more strongly at lower doses of H$_2$O$_2$ and at higher doses of HOCl. Expression of sigH was repressed at higher HOCl concentrations (2.5 mM), indicating that its control is not strictly H$_2$O$_2$ specific.

While the intestine is primarily an anaerobic environment (5), exact oxygen concentrations are difficult to measure in vivo and may vary depending on anatomical position or specific microenvironments in the intestine (79). Recent evidence suggests that inflammation, antibiotic treatment, and infection with enteric pathogens may increase the amount of oxygen available to microbes in the gut (6, 7). We therefore wanted to assess how much of an effect oxygen has on expression of redox-regulated genes in L. reuteri. We repeated our RT-PCR experiment with microaerobic cultures, which were prepared aerobically and grown in full screw-cap tubes without shaking.
low-oxygen conditions under which *L. reuteri*, like other lactic acid bacteria (70, 80), can remove oxygen from liquid media and grow to the same density as under anaerobic conditions (Fig. S3). The results of this experiment (Fig. 7B) revealed that the presence of even the low levels of oxygen expected in these cultures had large effects on redox-responsive gene expression. In contrast to what we observed anaerobically, expression of *ahpF*, *pcl1*, *moeB*, and *sigH* was unaffected by *H*2*O*2 under these conditions, and activation of *perR* was reduced. HOCl activation of *ahpF*, *pcl1*, and *perR* expression was eliminated in the presence of oxygen, and expression of both *moeB* and *rsiR* expression was HOCl repressed. These results showed that oxygen can dramatically affect how bacteria regulate gene expression in response to inflammatory oxidants and that studies of redox responses in the presence of even small amounts of oxygen may not necessarily reflect how bacteria respond in anaerobic environments and vice versa.

**Identifying genes important for surviving oxidative stress in *L. reuteri*.** Finally, we wanted to use the gene expression data generated as described above to begin

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**FIG 7** Dose-responsive control of gene expression by oxidative stress. *L. reuteri* ATCC PTA 6475 was grown anaerobically or microaerobically at 37°C to an *A*600 of 0.3 to 0.4 in MEI-C and then treated for 15 min with the indicated concentrations of *H*2*O*2 or HOCl. Change in expression of the indicated genes relative to untreated control cells was measured by quantitative RT-PCR (*n* = 3, means ± SDs). Asterisks indicate significant differences in expression at different oxidant concentrations under a given growth condition (two-way ANOVA with Holm-Sidak’s multiple-comparison correction) as follows: *, *P* < 0.05, and **, *P* < 0.01.
identifying genes involved in protecting *L. reuteri* against the toxicity of H$_2$O$_2$ and HOCl, based on the simple hypothesis that genes strongly upregulated by a certain stress may be involved in protecting the cell against that stress (81). We were particularly interested in identifying genes encoding factors that protect *L. reuteri* against HOCl, since much less is known about HOCl defense in bacteria in general (22), and no previous studies have examined how lactic acid bacteria survive reactive chlorine stress. We therefore identified bactericidal doses of H$_2$O$_2$ and HOCl for *L. reuteri* (Fig. S4A) and found that 1.5 mM H$_2$O$_2$ was sufficient to cause a 99.9% loss in viable cells of *L. reuteri* over the course of an hour both anaerobically and microaerobically. Subsequent titration of bactericidal HOCl concentrations (Fig. S4B) showed that doses resulting in a rate of viability loss comparable to that seen with 1.5 mM H$_2$O$_2$ in the first 40 min after treatment also resulted in recovery of viable cells after 60 min. We therefore used bactericidal concentrations of HOCl that resulted in a 99.999% loss in viable cells with no recovery during the 1-h course of the experiment (7.5 mM anaerobically and 2.5 mM microaerobiocally).

We constructed several strains containing null mutations of genes that we predicted to be involved in defense against either H$_2$O$_2$ or HOCl, based on known bacterial redox stress response mechanisms (22, 23, 53, 82) and on our transcriptomic data. We obtained mutants lacking *msrB* and *perR*, which we expected to be involved in H$_2$O$_2$ response, as well as four mutants lacking genes we expected to be involved in HOCl response: *ppk1* and *ppk2*, encoding two different kinases able to produce inorganic polyphosphate (polyP), which protects against HOCl-mediated protein damage in *E. coli* (57, 76, 83); *rclA*, encoding a conserved flavoprotein known to protect *E. coli* against HOCl (53); and *hsIO*, encoding the widely conserved HOCl-activated chaperone Hsp33 (77). Finally, we knocked out four genes found in *L. reuteri* but not in enterobacteria with either interesting redox-responsive expression patterns or, in the case of *rsiR*, a known role in probiotic action: *sigH*, *rsiR*, *lo18* (*hsp20*), which encodes a small heat shock protein found only in lactic acid bacteria (59, 60) and whose expression was more strongly activated by HOCl than by H$_2$O$_2$, and LAR_RS09945, encoding a predicted oxidoreductase that was very strongly upregulated by HOCl but not by H$_2$O$_2$. The ability of each of these strains to survive bactericidal oxidative stress was measured by comparison to the viability of the wild-type strain under the same conditions (Fig. 8).

Anaerobically, the *msrB* mutant was extremely sensitive to H$_2$O$_2$ treatment, as expected (23, 47), and the *perR* mutant, which is expected to have constitutively high expression of peroxide defense genes (84), was significantly protected. A mutant lacking *rsiR* was significantly more sensitive to H$_2$O$_2$, suggesting that despite the fact that its expression is not controlled by this oxidant (Fig. 3), it is important for surviving H$_2$O$_2$ treatment (35). Surprisingly, only the *perR* mutant was significantly more sensitive than the wild type to HOCl under anaerobic conditions. However, knocking out *lo18* had a significant and unexpected protective effect. This was particularly surprising since *lo18* expression was strongly upregulated in response to HOCl. Under microaerobic conditions, the results of survival assays were considerably different. There were only minor differences in survival of a bactericidal dose of H$_2$O$_2$ in microaerobic cultures for any of the mutants, with *msrB*, *rsiR*, and *rclA* mutants showing very small but statistically significant defects in survival at the 1-h time point. In contrast, there were more significant differences in survival of HOCl stress under microaerobic conditions. The *msrB*, *perR*, *lo18*, and *ppk2* mutants had significant defects in HOCl stress survival under these conditions. The *perR* and LAR_RS09945 mutants were significantly protected at the 20-min time point, but this effect was lost at later time points. There was no difference in HOCl survival between the wild type and *sigH* or *hsIO* mutants. The ability of each mutant to grow in media containing nonbactericidal concentrations of H$_2$O$_2$ and HOCl (Fig. S5) was more variable and the differences between mutants and the wild type of smaller magnitude, but *perR*, *hsIO*, *lo18*, *rsiR*, and LAR_RS09945 mutants had significant, >2-h lags in growth relative to that of the wild type after inoculation into anaerobic media containing either H$_2$O$_2$ or HOCl and a *perR* mutant had a significant growth advantage in microaerobic media containing HOCl 7 to 12 h after inoculation.
Overall, our results show that different redox stress treatment methods can give different results, further emphasize that oxygen concentration has dramatic effects on oxidative stress survival, and indicate that it will be important to quantify what oxygen levels gut bacteria are exposed to in inflamed and noninflamed gut environments (5–7) to understand what genes are likely to play roles in ROS and RCS resistance in vivo.

Screening mutants lacking HOCl-induced genes has successfully identified HOCl resistance factors in other bacterial species (53, 54, 56, 66), but this strategy had limited success in L. reuteri. Neither sigH nor LAR_RS09945 mutations, for example, had any
effect on resistance to the stresses which regulated their expression. In future work, a genome-wide mutant screening approach (e.g., transposon sequencing) (52) may be valuable for identifying additional genes required for H₂O₂ and HOCl stress survival, and complementation and overexpression analysis will be necessary to confirm that the observed phenotypes are specific to the constructed mutations in each gene. Nevertheless, our targeted mutagenesis approach did allow us to identify several important players in oxidative stress resistance. Clearly, methionine sulfoxide reductase is a major contributor to the ability of _L. reuteri_ to resist oxidative stress both anaerobically and microaerobically, consistent with its enzymatic activity (47) and known role in colonization (48, 49). While PerR is relatively unimportant microaerobically, anaerobically it plays a key role in regulating H₂O₂ resistance, as expected (65), although for unknown reasons it appears that the constitutive overexpression of H₂O₂ resistance genes expected in a perR mutant is detrimental in the presence of HOCl.

**_L. reuteri*-specific defenses against H₂O₂ and HOCl stress.** The H₂O₂ sensitivity of the _rsiR_ mutant was somewhat surprising, since this _L. reuteri*-specific gene has largely been characterized for its role in regulating the expression of the histamine-producing histidine decarboxylase locus of _L. reuteri_, where _rsiR_ is essential for histamine-dependent anti-inflammatory phenotypes (34, 35). However, RsiR is a global regulator, activating and repressing transcription of 195 and 143 genes, respectively, many of which are involved in redox homeostasis (including _ahpC, perR_, and genes involved in cysteine and methionine synthesis) (35). It is currently unclear what signal(s) RsiR responds to, which RsiR-regulated genes contribute to H₂O₂ sensitivity, or what role H₂O₂ resistance plays in RsiR-dependent anti-inflammatory effects _in vivo_, and these are exciting issues for future research exploring the connections between inflammatory oxidants and anti-inflammatory probiotic mechanisms.

The small heat shock protein Hsp33 and the flavoprotein RclA are RCS-specific defense factors in _E. coli_ (53, 77), so we were also surprised to find that mutations of these genes had no apparent effect on HOCl resistance in _L. reuteri_, despite the fact that _rclA_ expression was induced more strongly by HOCl treatment than by H₂O₂ (Fig. 8A and Table S1). This could be due to the redundant nature of RCS resistance mechanisms (22) or could reflect fundamental differences in RCS response between _L. reuteri_ and _E. coli_. Supporting the second hypothesis is the fact that mutations in _lo18_ and _ppk2_, genes not found in _E. coli_, had very strong effects on HOCl resistance. Lo18 is a chaperone found only in a subset of _Lactobacillus_ and _Oenococcus_ species that stabilizes proteins and membranes under heat and ethanol stress conditions (59, 60). While this could easily explain how Lo18 protects _L. reuteri_ against the protein-unfolding activity of HOCl, as we saw under microaerobic conditions, it is much less intuitive why the loss of Lo18 protected _L. reuteri_ against HOCl anaerobically, and more work will be needed to understand the mechanism underlying this effect. PolyP plays a role in stress resistance and probiotic phenotypes in several different _Lactobacillus_ species (85–90). In _E. coli_, the polyP kinase PPK (homologous to _L. reuteri_ PPK1) is required for HOCl resistance (76), but deletion of _ppk1_ had only a modest, nonstatistically significant effect on HOCl resistance in _L. reuteri_. In contrast, deletion of _ppk2_, which encodes an unrelated polyP kinase (PPK2) whose primary physiological role is generally thought to be in generating NTPs from NDPs or NMPs and polyP (57, 58), led to a highly significant defect in HOCl resistance, albeit only in the presence of oxygen. Whether polyP production in response to HOCl stress is driven by PPK1 or PPK2 in _L. reuteri_ remains to be determined, as does the relative importance of PPK2’s polyP- and NTP-synthesizing activities. PPK2 is not present in enterobacteria but is found in many species of commensal bacteria (including lactobacilli, _Bacteroidetes_, and _Clostridiaceae_) (58, 91, 92).

Our results clearly demonstrate that HOCl resistance in _L. reuteri_ depends on factors different than in _E. coli_ or _B. subtilis_. These differences may represent possible targets for differentially sensitizing gut bacteria to oxidative stress. Interestingly, the frontline IBD drug mesalamine has recently been shown to be an inhibitor of PPK1 (93), and it is tempting to speculate that mesalamine may therefore have a larger impact on the
ability of enterobacteria to survive in the inflamed gut than on PPK2-encoding commensals, although more data will be needed to test this hypothesis.

Conclusions. Manipulating the microbiome is likely to be a key element in future treatments for inflammatory diseases of the gut. Development of such treatments will require a sophisticated understanding of how gut bacteria respond to changes in their environment. The differences we have now begun to uncover in oxidative stress response between anti-inflammatory, health-associated bacteria and proinflammatory, disease-associated species may present opportunities for new therapies. We hope that our results will ultimately make it possible to sensitize enterobacteria to inflammatory oxidants while simultaneously protecting the healthy gut community.

MATERIALS AND METHODS

Bacterial strains and growth conditions. All strains and plasmids used in this study are listed in Table 1. All L. reuteri strains were derivatives of strain ATCC PTA 6475 (Biogaia) (94). Strain 6475rsiR-Stop (35) was a gift from James Versalovic (Baylor College of Medicine), and plasmid pJP042 (recT+ /erm+) (94) was a gift from Jan-Peter van Pijkeren (University of Wisconsin—Madison). L. reuteri was grown at 37°C in MEI broth (86) without added cysteine (MEI-C) or on solid De Man, Rogosa, and Sharpe (MRS) agar (Difco). Anaerobic cultures were incubated in an anaerobic chamber (Coy Laboratory Products) in an atmosphere of 90% nitrogen, 5% CO2, and 5% H2 or in Hungate tubes prepared, inoculated, and sealed in that chamber. Liquid media were made anaerobic before use by equilibration for at least 24 h in the anaerobic chamber. MRS plates for CFU plate counts were incubated in sealed containers made anaerobic using GasPak EZ sachets (Becton, Dickinson). Microaerobic cultures were incubated aerobically without shaking in 16- by 125-mm screw-cap test tubes containing 15 ml of MEI-C. Methylene blue (2 mg liter−1) was added when indicated (95). Aerobic cultures (5 ml in a 16-mm diameter test tube) were incubated with shaking (200 rpm). Details of H2O2 and HOCl stress treatments, transcript quantification, and phenotype analysis are described in the supplemental material.

Molecular methods. Oligonucleotide-directed recombineering was used to construct null mutations in the chromosome of L. reuteri using the pJP042-encoded RecT recombinase as previously described (94). Null mutations were designed to incorporate in-frame stop codons near the 5′ end of each gene. Mutagenic primers used are listed in Table S3. Primers used for quantitative RT-PCR were designed with Primer Quest (Integrated DNA Technologies; parameter set “qPCR 2 primers intercalating dyes” for qRT-PCR primer design) and are listed in Table S4. Additional primers for PCR amplification, screening, and sequencing were designed using WebPrimer (www.candidagenome.org/cgi-bin/compute/web-primer). All chromosomal mutations were confirmed by Sanger sequencing (UAB Heflin Center for Genomic Sciences).

Data availability. All strains generated in the course of this work are available from the authors upon request. RNA sequencing data have been deposited in NCBI’s Gene Expression Omnibus (96) and are accessible through GEO Series accession number GSE127961.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mSystems.00453-19.

TEXT S1, DOCX file, 0.03 MB.

FIG S1, EPS file, 1.1 MB.

FIG S2, EPS file, 1.6 MB.

FIG S3, EPS file, 2.5 MB.

### TABLE 1 Strains and plasmids used in this study

| Strain or plasmid | Relevant genotype | Source and/or reference |
|-------------------|-------------------|-------------------------|
| L. reuteri strains |                   |                         |
| ATCC PTA 6475     | Wild type; human breast milk isolate | Biogaia (94) |
| 6475rsiR-Stop     | rsiR (LAR_RS05165) | 35                      |
| MJG0562           | ppp1 (LAR_RS01770) |                         |
| MJG0569           | ppp2 (LAR_RS00075) |                         |
| MJG0570           | rclA (LAR_RS00915) |                         |
| MJG0977           | msrB (LAR_RS00975) |                         |
| MJG0979           | hslO (LAR_RS01385) |                         |
| MJG1017           | iol18 (LAR_RS07000) |                         |
| MJG1056           | LAR_RS09945       |                         |
| MJG1278           | sigH (LAR_RS04695) |                         |
| MJG1573           | perR (LAR_RS06970) |                         |
| Plasmid pJP042    | recT+ /erm+       | 94                      |

*Unless otherwise indicated, all strains were generated in the course of this work.*
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