Steady-State Hydrogen Peroxide Induces Glycolysis in Staphylococcus aureus and Pseudomonas aeruginosa

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Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) from human pathogens Staphylococcus aureus and Pseudomonas aeruginosa can be readily inhibited by reactive oxygen species (ROS)-mediated direct oxidation of their catalytic active cysteines. Because of the rapid degradation of H$_2$O$_2$ by bacterial catalase, only steady-state but not one-dose treatment with H$_2$O$_2$ rapidly induces glycolysis and the pentose phosphate pathway (PPP). We conducted transcriptome sequencing (RNA-seq) analyses to globally profile the bacterial transcriptomes in response to a steady level of H$_2$O$_2$, which revealed profound transcriptional changes, including the induced expression of glycolytic genes in both bacteria. Our results revealed that the inactivation of GAPDH by H$_2$O$_2$ induces metabolic levels of glycolysis and the PPP; the elevated levels of fructose 1,6-biphosphate (FBP) and 2-keto-3-deoxy-6-phosphogluconate (KDPG) lead to dissociation of their corresponding glycolytic repressors (GapR and HexR, respectively) from their cognate promoters, thus resulting in derepression of the glycolytic genes to overcome H$_2$O$_2$-stalled glycolysis in S. aureus and P. aeruginosa, respectively. Both GapR and HexR may directly sense oxidative stresses, such as menadione.

Pathogenic bacteria, such as Pseudomonas aeruginosa and Staphylococcus aureus, need to conquer high concentrations of reactive oxygen species (ROS) that are produced by host phagocytes for sustained virulence (1). To this end, these bacteria use ROS-reactive small molecules, such as glutathione (GSH) and melanin in P. aeruginosa as well as coenzyme A and staphyloxanthin in S. aureus. These pathogens also produce a group of ROS-detoxifying enzymes, such as catalase, superoxide dismutase, hydroperoxide reductase, thioredoxin, and glutaredoxin, whose expression is induced by oxidative stress (2–4).

It has been well documented that P. aeruginosa and S. aureus also mount global transcriptional changes by utilizing a group of thiol-based ROS-active transcription regulators, such as OxSR, SoxR, MgrA, OhrR, SarA, SarZ, MexR, OspR, CymR, AirSR, and AgrA (5–15). Upon oxidative stress, the specific cysteine groups in these regulatory proteins form sulfuric acids or disulfides, thus inducing conformational changes that attenuate their DNA binding affinities. Our previous work showed that a thiol-based, oxidation-sensing mechanism is utilized by these human pathogens to sense the host immune response and regulate a global change of their properties. ROS leads to activation of defense systems to reduce the oxidative threat as well as a major shift in the life forms of the pathogens (6, 11).

ROS can efficiently oxidize the thiol group of active and allosteric cysteines in bacterial proteins, causing changes in their functions. Previously, we employed an isotopic orthogonal proteolytic–activity-based protein profiling (isoTOP-ABPP) technology to identify around 200 oxidation-sensitive cysteines and further determined that several of these proteins perform important redox-active functions in bacteria. The master quorum sensing regulator LasR of P. aeruginosa undergoes an oxidation-responsive transcriptional regulation. Oxidation induces switching of metabolic pathways by modification of active site and/or allosteric cysteine residues in enzymes, such as acetaldehyde dehydrogenase ExaC, arginine deiminase ArcA, and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (16). GAPDH is readily oxidized and inhibited by oxidation. Pathogenic bacteria exhibit a complex, multilayer response to ROS that includes the rapid adaption of metabolic pathways to oxidative stress challenge (17).

Central metabolism has profound influences on bacterial response to ROS. The pentose phosphate pathway (PPP) contributes bacterial tolerance to oxidative stress by generating the redox currency (NADPH), which is the substrate for other reducing agents (18). NADPH is responsible for generating glutathione (GSH) in P. aeruginosa and reduced thioredoxin in S. aureus (19, 20). In addition, PPP is important for producing nucleotide precursors to repair DNA damage under ROS stress in Deinococcus radiodurans (21), suggesting a potentially conserved mechanism in other bacteria. In S. aureus, PPP is linked with glycolysis, which is strictly regulated by the glycolytic repressor GapR (22). In Bacillus subtilis, fructose 1,6-biphosphate (FBP) is the cognate ligand that derepresses CggR, the ortholog of GapR (23), suggesting a similar interaction between FBP and GapR in S. aureus. In P. aeruginosa, PPP is closely associated with the Entner-Doudoroff (ED) pathway, which is controlled by the repressor HexR. HexR specifically senses 2-keto-3-deoxy-6-phosphogluconate (KDPG),
which results in derepression of several ED operons (24). It has been shown in yeast (Saccharomyces cerevisiae) that upon ROS, oxidative inhibition of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) leads to prompt metabolic redirection from glycolysis to PPP, which generates more NADPH (25). Notably, genes related to anaerobic metabolism and cytochrome uptake and storage are upregulated by oxidative stress. Notably, genes associated with DNA repair, virulence, and iron metabolism were upregulated in P. aeruginosa, and genes associated with DNA repair, virulence, and iron metabolism were upregulated in S. aureus. In the case of P. aeruginosa, DNA repair proteins, catalases, intracellular iron transport, and regulation are important for bacterial adaption to oxidative stress. Hydrogen peroxide induced the expression of all F-, R-, and S-type pyocins, leading to self-killing activity via DNA breakage and lipid biosynthesis inhibition (27).

We recently found that both bacteria can rapidly degrade H₂O₂ in several minutes, which leads to the question of whether the in vitro one-dose treatment with H₂O₂ would induce the comprehensive responses that are elicited by host-derived steady-state levels of H₂O₂ in vivo (16). As expected, our transcriptome sequencing (RNA-seq) analyses using a steady level of H₂O₂ stress uncovered significantly more genes that belong to pathways, such as those involved in glycolysis, virulence, translation, and RNA metabolism. The subsequent assays demonstrated that upon ROS stress, the elevated levels of fructose 1,6-biphosphate (FBP) and 2-keto-3-deoxy-6-phosphogluconate (KDPG) lead to dissociation of their corresponding glycolytic repressors from their cognate promoters, thus causing derepression of glycolytic genes to overcome H₂O₂-stalled glycolysis.

**MATERIALS AND METHODS**

**Strains, plasmids, and primers.** Strains, plasmids, and primers are listed in Table S1 in the supplemental material. The Pseudomonas aeruginosa MPA01 strain was maintained in LB medium. The Staphylococcus aureus Newman strain was cultured in Trypticase soy broth (TSB) medium. For plasmid maintenance in P. aeruginosa and Escherichia coli, the medium was supplemented with 50 µg/ml carbenicillin and 100 µg/ml ampicillin, respectively.

**Metabolite preparation and quantification using LC-MS.** For isolation of water-soluble metabolites, an ethanol-water protocol was used as previously described (28). Briefly, wild-type (WT) P. aeruginosa MPA01 or S. aureus Newman strains were grown aerobically in LB or TSB medium for overnight at 37°C, diluted 100-fold in 20 ml of fresh medium, and incubated at 37°C with shaking at 250 rpm for 3 h (optical density at 600 nm [OD₆₀₀] of ~0.6). To examine the effect of oxidative stress on the metabolites, 20 ml of mid-log-phase cultures was placed in a dialysis bag (10 kDa) with shaking in 1 liter of LB or TSB containing 10 mM H₂O₂ for 10 min. The control sample was also dialyzed against medium without H₂O₂ for 10 min. Bacteria were collected by centrifugation for 5 min at 5,000 × g at 4°C and then washed once by prechilled 0.6% NaCl solution at 4°C. One milliliter of prechilled 60% ethanol solution was used to resuspend the pellet, which was then snap-frozen in liquid nitrogen. The bacterial pellets were subjected to two rounds of bead disruption (Fast Prep EP120 instrument; Qiagen) at 4°C. After centrifugation, the supernatant was stored at −80°C or injected directly into the mass spectrometer. Detailed liquid chromatography-mass spectrometry (LC-MS) procedures have been described previously (29). Four biological repeats were included for each sample.

**RNA-seq, data analyses, and qRT-PCR verifications.** To examine the effect of oxidative stress on the transcriptome, 10 ml of mid-log-phase bacterial cultures (P. aeruginosa and S. aureus) were placed in a dialysis bag (10 kDa) with shaking in 1 liter of LB or TSB containing 10 mM H₂O₂ for 10 min. The control sample was also dialyzed against medium without H₂O₂ for 10 min. An RNAeasy minikit (Qiagen) was used for subsequent RNA purification with DNase I treatment. After removing RNA by using the MICROBExpress kit (Ambion), mRNA was used to generate the cDNA library according to the TruSeq RNA sample prep kit protocol (Illumina), which was then sequenced using the HiSeq 2000 system (Illumina). Bacterial RNA-seq reads were mapped to the P. aeruginosa and S. aureus genomes by using TopHat (version 2.0.0), with two mismatches allowed (30). Only the uniquely mapped reads were kept for the subsequent analyses. The gene differential expression analysis was performed using Cuffdiff software (version 2.0.0) (31). Quantitative reverse transcription-PCR (qRT-PCR) was performed to verify the transcriptional changes for several glycolytic genes (gapA, gapR, and fbp of S. aureus, as well as gapA, zwf, fic, fdB, fkd, algU, algG, and mucA from P. aeruginosa [primers shown in Table S4 in the supplemental material]). GO enrichment analyses were conducted on all differentially transcribed genes using DAVID (32) before data sets were imported into Cytoscape with an Enrichment Map plugin (33).

**Statistical analysis.** RNA-seq analyses were repeated twice. All other experiments were repeated at least three times. Two-tailed Student’s t-tests were performed using Microsoft Office Excel 2011.

**MIC measurements.** MICs of H₂O₂ were measured by using a microdilution technique according to NCCLS guidelines (34) in Mueller-Hinton broth. TSB medium was used to grow S. aureus in a 96-well plate. The MIC value was recorded as the lowest concentration at which there was no visible growth of S. aureus.

**Protein purification for GapR, HexR, and Eda.** For the expression of GapR, HexR, or Eda, we used the ligation-independent cloning (35) method (36). The respective coding regions were PCR amplified from either S. aureus (GapR) or P. aeruginosa (HexR and Eda) genomic DNA with the primers listed in Table S1 in the supplemental material (GapR-EXP/F for GapR, HexR-EXP/F for HexR, and Eda-EXP/F for Eda). The PCR products were treated with T4 DNA polymerase in the presence of dCTP for 30 min at room temperature. Target vector pMC5G19 (36) was digested with SspI, gel purified, and then treated with T4 DNA polymerase in the presence of dCTP for 15 min at 16°C. The T4 DNA polymerase-treated plasmid vector and PCR product were gel purified, mixed, incubated for 5 min at room temperature, and then transformed into E. coli strain DH5. The resulting plasmid was transformed into BL21 Star(DE3) containing a plasmid (pRK1037) expressing tobacco vein mottling virus (TVMV) protease (Science Reagents, Inc.), and the transformants were selected on LB agar plates with 100 µg/ml ampicillin and 50 µg/ml kanamycin. The BL21 Star(DE3) strain carrying the plasmid was grown in LB to an optical density at 600 nm (OD₆₀₀) of 0.6, and then 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) was added. After overnight induction at 16°C, the cells were harvested and frozen at −80°C. The expressed protein was purified from the frozen cells with a HisTrap column (GE Healthcare, Inc.) by following the column manufacturer’s recommendations. The purified protein was supplemented with 20% glycerol and stored at −80°C.

**EMSA.** The electrophoretic mobility shift assay (EMSA) was performed as follows. DNA probes containing promoter regions of gapR (S. aureus), zwf (P. aeruginosa), and gapA (P. aeruginosa) were PCR amplified using primers gapR-GSF/R, zwf-GSF/R, and gapA-GSF/R, respectively (listed in Table S1 in the supplemental material). The PCR products were radiolabeled with T4 polynucleotide kinase (NEB) and [γ-32P]ATP (PefinKin-Elmer). The radioactive probe (2 ng) was mixed with various amounts of the GapR or HexR protein in 20 µl of gel shift loading buffer (10 mM Tris-HCl, pH 7.4, 50 mM KCl, 5 mM MgCl₂, 10% glycerol, 3 µg/ml sheared salmon sperm DNA). After being incubated at room temperature for 20 min, the samples were analyzed by 8% (for S. aureus gapR)
or 6% (for *P. aeruginosa zwf* or gapA) polyacrylamide gel electrophoresis (100 V for prerun and 85 V for 45 min for sample separation). The gels were dried and subjected to autoradiography on a phosphor screen (BAS-IP; Fuji). The assay was repeated at least for three times with similar results.

**Dye primer-based DNase I footprint assay.** The DNase I footprint procedures were modified according to reference 37. The promoter regions from gapR or zwf were generated by PCR with the primers gapR-FF-6FAM and gapR-GSR as well as zwf-FF-6FAM and zwf-GSR (see Table S1 in the supplemental material). About 50 ng of 6-carboxyfluorescein FP-6FAM and gapR-GSR as well as zwf-FP-6FAM and zwf-GSR (see Table S1 in the supplemental material). The primers PA3001DGapF (with an XbaI site) and PA3001DGapR (with an XbaI site) were used for amplification of 2 kb of PA3001 downstream region. The two respective PCR products were digested with HindIII-XbaI and XbaI-EcoRI, respectively, and then cloned into HindIII/EcoRI-digested gene replacement vector pEX18Ap via a three-piece ligation, yielding pEX18Ap-PA3001UD. A 0.8-kb gentamicin resistance cassette was cut from pPS858 with XbaI and then cloned into pEX18Ap-PA3001UD, yielding pEX18Ap-PA3001UGD. The resultant plasmid was electroporated into MPAO1 with selection for gentamicin resistance. Colonies were screened for gentamicin sensitivity and loss of sucrose (5%) sensitivity, which typically indicates a double-crossover event and is indicative of gene replacement. The resulting ΔPA3001 strain was further confirmed by PCR analysis.

**NADPH assay.** The intracellular concentration of NADPH was measured using the EnzyChrom NADP+/NADPH assay kit from BioAssay Systems. Bacteria of the wild-type *P. aeruginosa* MPAO1 or *S. aureus* Newman strain were grown in LB or TSB medium overnight at 37°C, diluted 100-fold in 20 ml of fresh medium, and incubated at 37°C with shaking at 250 rpm for 3 h (OD600 of ~0.6). To examine the effect of oxidative stress on NADPH, 20 ml of mid-log-phase culture was placed in a dialysis bag (10 kDa) with shaking in 1 liter of LB or TSB containing 10 mM H2O2 for 10 min. Bacteria were collected by centrifugation for 5 min at 5,000 × g at 4°C and then washed once with prechilled 0.6% NaCl solution at 4°C before following the manufacturer’s instructions.

**H2O2 assay.** The concentrations of H2O2 inside the dialysis bag during the 10-min dialysis (starting with 10 mM H2O2) were measured using the Amplex Red hydrogen peroxide/peroxidase assay kit (LifeTechnologies). The medium was taken from the dialysis bag at certain time points before following the manufacturer’s instructions.

**Microarray data accession number.** The RNA-Seq data files have been deposited in NCBI’s Gene Expression Omnibus (GEO) and can be accessed through GEO series accession no. GSE55258.

## RESULTS

Steady-state levels, but not one-dose treatments with H2O2, induce PPP following inhibition of GAPDH. Like many other organisms, GAPDH from both *P. aeruginosa* and *S. aureus* can be readily inhibited by ROS-mediated direct oxidation of their catalytically active cysteines (16). We speculated that inhibition of GAPDH leads to an elevated pentose phosphate pathway (PPP), which has been observed in yeast (*Saccharomyces cerevisiae*) (25). This hypothesis also implies that bacteria generate more NADPH to protect from ROS; NADPH is the main cellular reducing agent used by bacterial protective enzymes in reducing H2O2, and related oxidants (44). In order to verify this speculation, we cultured *P. aeruginosa* and *S. aureus* until mid-log phase and treated the bacteria with one dose of H2O2 (10 mM) for 20 min as commonly used previously. After extracting intracellular metabolites from the bacteria, we utilized LC-tandem MS (LC-MS/MS) to quantify the intracellular concentrations of a group of glycolytic and PPP metabolites in H2O2-treated versus untreated controls. There were no significant changes observed in metabolite levels post-H2O2 treatment (see Fig. S1A in the supplemental material).
Given that both bacteria are able to completely degrade 10 mM H\textsubscript{2}O\textsubscript{2} within minutes (16), we changed our approach and continuously treated bacteria (10 ml of mid-log-phase culture) with 1 liter medium containing 10 mM H\textsubscript{2}O\textsubscript{2} for 10 min in a dialysis bag (10-kDa cutoff). We measured the actual H\textsubscript{2}O\textsubscript{2} concentrations inside the dialysis bag at different time points (0, 1, 2, 5, and 10 min) during the 10-min dialysis. As shown in Fig. S1B in the supplemental material, the concentration ranges from 3 to 7 mM, indicating the steady state of H\textsubscript{2}O\textsubscript{2} exposed to bacteria via this method. Under the new conditions, the levels of glucose-6-phosphate/fructose-6-phosphate (G6P/F6P), glucose or fructose bisphosphate (G2P/F2P), and ribose-5-phosphate (R5P) were significantly increased after H\textsubscript{2}O\textsubscript{2} exposure in both pathogens (Fig. 1A), indicating that the inactivation of GAPDH likely elevates PPP metabolites. Finally, we also detected a higher concentration of the reduced glutathione (GSH) in the H\textsubscript{2}O\textsubscript{2}-treated P. aeruginosa, which strongly suggests that the increased NADPH is used to generate more GSH (via glutathione reductase) to counter oxidative stress in P. aeruginosa (Fig. 1A). The different effects between one-dose and steady-dose H\textsubscript{2}O\textsubscript{2} led us to test if one-dose H\textsubscript{2}O\textsubscript{2} would also elicit a similar metabolic change at early time points. As expected, we were able to observe induced levels of glycolytic and PPP metabolites at 2 min but not after 10 min after one dose of 10 mM H\textsubscript{2}O\textsubscript{2} (see Fig. S1C).

If the metabolic change is dependent on GAPDH, its mutant would display a similar metabolic change as elicited by the steady dose of hydrogen peroxide supplement. There are two GAPDH enzymes in S. aureus, GapA and GapB, which are predicted to catalyze the glycolytic oxidation of GAP and the reverse gluconeogenic reaction, respectively (22). In order to verify this hypothesis, we obtained an S. aureus gapA (SAOUHSC_00795 [encodes GAPDH]) deletion mutant from J. A. Morrissey at the University of Leicester, and constructed a P. aeruginosa PA3001 (encodes GAPDH) deletion mutant by exchanging a gentamicin resistance gene with the PA3001 gene in the chromosome. Intracellular metabolites were extracted from these mutants and their parent wild-type (WT) strains and quantified by LC-MS. As shown in Fig. S1E in the supplemental material, these mutants indeed displayed a decrease in NADH and elevated levels of PPP metabolites. Different from the WT, these mutants displayed no significant metabolic changes in the absence or presence of a steady dose of H\textsubscript{2}O\textsubscript{2} treatment (see Fig. S1F), which further confirms the essential role of GAPDH in hydrogen peroxide-induced metabolic change.

We next tested if inactivation of GAPDH could result in an increased anti-ROS response in bacteria. The mutation in gapA affected growth and therefore was not used (data not shown). We then supplemented the culture medium with iodoacetic acid (IAA [400 \mu M]), which is a known inhibitor of GAPDH (45), and then measured the MIC to H\textsubscript{2}O\textsubscript{2} of IAA-treated and untreated bacteria. As shown in Fig. 1B, IAA-treated S. aureus displayed a 4-fold higher MIC to H\textsubscript{2}O\textsubscript{2} than the untreated control, indicating that the GAPDH blockage indeed enables bacterial resistance to ROS. Our metabolic quantification and subsequent MIC measurements indicate that oxidative inactivation of GAPDH functions as a metabolic switch that maintains NADPH/NADP\textsuperscript{+} equilibrium during oxidative stress in bacterial pathogen.

Recently, a combination of specific glycolytic metabolites (glucose, mannitol, fructose, or pyruvate) and aminoglycosides have been presented as the first strategy capable of eradicating bacterial persisters, a notorious subpopulation of dormant bacteria that can tolerate antibiotic treatment (38). Catabolism of these metabolites generates NADH via glycolysis. After being oxidized by the electron transport chain, NADH contributes to proton motive force (PMF) that promotes aminoglycoside uptake and the killing of persisters. This observation points to the importance of the bacterial metabolites in antibiotic mechanism of action. Integrating these results with H\textsubscript{2}O\textsubscript{2}-mediated inhibition of GAPDH led us to speculate that ROS inactivation of GAPDH could decrease NADH and PMF, thus disabling persister eradication. To test this hypothesis, we cultured S. aureus until the mid-log phase and treated bacteria with 1 mM H\textsubscript{2}O\textsubscript{2} (in a dialysis bag) or 400 \mu M IAA for 1 h. After extracting intracellular metabolites from the bacteria, we utilized LC-MS/MS to quantify the intracellular concentration of NADH and compared it to that of the untreated controls. The NADH level in S. aureus was significantly decreased after exposure to H\textsubscript{2}O\textsubscript{2} or IAA (Fig. 1C). We measured the S. aureus persister killing by both glucose and kanamycin in the presence of 1 mM H\textsubscript{2}O\textsubscript{2} or 400 \mu M IAA. As shown in Fig. 1D, a 100-fold increase of persister survival was observed in the presence of 1 mM H\textsubscript{2}O\textsubscript{2} or 400 \mu M IAA compared to the untreated control. In the absence of H\textsubscript{2}O\textsubscript{2}, the persisters derived from the gapA deletion displayed higher resistance to kanamycin than those from the WT (Fig. 1D). Taken together, our results showed that the activity of GAPDH plays an important role in the metabolite-based persister eradication (Fig. 1E).

RNA-seq analyses revealed that steady-state levels of H\textsubscript{2}O\textsubscript{2} elicited profound transcriptional changes, including induced glycolysis in pathogenic bacteria. It is estimated that a steady-state level of superoxide is produced by NADPH oxidase in a phagosome (1, 46). The observation of induced glycolysis under only steady levels of ROS stress strongly suggests that a steady-state stress of H\textsubscript{2}O\textsubscript{2} might induce broader and greater transcriptional fluctuation than the one-dose H\textsubscript{2}O\textsubscript{2} that had been used in previous microarray analyses (26, 47). With this in mind, we decided to employ RNA-seq to examine the transcriptomes of S. aureus Newman and P. aeruginosa MPAOI in response to continuous treatment with 10 mM H\textsubscript{2}O\textsubscript{2} (in a dialysis bag for 10 min), a concentration commonly used by other studies since both pathogens can tolerate millimolar levels of H\textsubscript{2}O\textsubscript{2} well (26, 48, 49). The same practice with a dialysis bag has been used in previous studies to expose bacteria to a steady-state level of H\textsubscript{2}O\textsubscript{2} (48, 50). We measured bacterial numbers of both pathogens in the presence and absence of a steady dose of 10 mM H\textsubscript{2}O\textsubscript{2} for 10 min with no difference observed (see Fig. S1D in the supplemental material). Therefore, we added sufficient but not deleterious H\textsubscript{2}O\textsubscript{2} in order to ensure that most steady H\textsubscript{2}O\textsubscript{2}-responsive changes could be revealed in our transcriptomic experiments. Using replicate experiments, we identified a total of 458 S. aureus genes (17.1% of the genome; 208 upregulated and 250 downregulated) and 1,722 P. aeruginosa genes (31.2% of the genome; 1,113 upregulated and 709 downregulated), whose mRNA levels were altered in response to a steady-state treatment with H\textsubscript{2}O\textsubscript{2} compared to the control without H\textsubscript{2}O\textsubscript{2} treatment (Fig. 2A; see Tables S2 and S3 in the supplemental material). As expected, these numbers were significantly greater than those in previous microarray analyses based on one-dose H\textsubscript{2}O\textsubscript{2} treatment (12.7% of the S. aureus genome and 9.3% of the P. aeruginosa genome), suggesting profound global transcriptional changes with steady-state stress of H\textsubscript{2}O\textsubscript{2}. Our new finding represents transcripts of genes that may be activated or
FIG 1. Steady-state treatment with \( \text{H}_2\text{O}_2 \) induces PPP following inhibition of GAPDH. (A) Changes in metabolite levels in \( P. \text{aeruginosa} \) or \( S. \text{aureus} \) treated with and without a steady dose of 10 mM \( \text{H}_2\text{O}_2 \) for 10 min. Bacterial lysates were prepared, and metabolites were quantified by LC-MS/MS. The absolute metabolite concentrations were normalized and are presented as fold changes compared to the untreated control (CK). The asterisks denote that the differences between \( \text{H}_2\text{O}_2 \)-treated and untreated samples are statistically significant \((P < 0.05)\). (B) MIC of \( \text{H}_2\text{O}_2 \) for \( S. \text{aureus} \) RN4220 strains in Mueller-Hinton broth. SA, wild-type strain RN4220; SA + IAA, wild-type RN4220 supplemented with 400 \( \mu \text{M} \) iodoacetic acid. The asterisks denote that the differences from the untreated wild type are statistically significant \((P < 0.05)\). (C) Changes in intracellular NADH levels in \( S. \text{aureus} \) treated with and without 1 mM \( \text{H}_2\text{O}_2 \) or 400 \( \mu \text{M} \) IAA for 10 min. Bacterial lysates were prepared, and metabolites were quantified by LC-MS/MS. The absolute metabolite concentrations were normalized and are presented as changes in percentage compared to the untreated control. The asterisks denote that the differences between \( \text{H}_2\text{O}_2 \)-treated and untreated samples are statistically significant \((P < 0.05)\). (D) Percentage of survival of \( S. \text{aureus} \) persisters after treatment with 30 \( \mu \text{g/ml} \) kanamycin and 1 mM glucose. Addition of 1 mM \( \text{H}_2\text{O}_2 \) or 400 \( \mu \text{M} \) IAA promoted survival of the persisters. Persisters derived from the \( \text{gapA} \) deletion are around 150-fold more resistant to kanamycin than the WT. The asterisk denotes that the difference is statistically significant \((P < 0.05)\). (E) ROS inactivation of GAPDH leads to a metabolic reconfiguration from glycolysis to the pentose phosphate pathway, thus increasing the intracellular levels of NADPH and GSH that are involved in counteracting oxidative stress. ROS-induced inhibition of GAPDH leads to the reduced production of NADH and PMF, thus repressing the eradication of metabolite-based persisters.
repressed during interaction with host macrophages that consistently produce a steady dose of peroxide (1).

Based on SP-PIR (Protein Information Resource) keyword designations, these differentially expressed genes were classified into multiple functional categories, including many metabolic pathways and stress responses (see Fig. S2A and S2B, respectively, in the supplemental material for \textit{S. aureus}, as well as Fig. S2C and S2D, respectively, for \textit{P. aeruginosa}). In \textit{S. aureus}, Gene Ontology (GO) enrichment analysis showed significant enrichment for genes related to glycolysis, the tricarboxylic acid (TCA) cycle, DNA repair, redox homeostasis, transporters, pyrimidine, and fatty acid and amino acid (β-Ala, Trp, and Lys) biosynthesis among the genes upregulated by the steady-state treatment with H2O2 (\(P < 0.005\)) and purine, tRNA, nitrogen, two-component systems, and amino acid (Ala, Asp, and Glu) metabolism among downregulated genes (\(P < 0.005\)) (Fig. 2B and C). Among these categories, genes involved in glycolysis, the TCA cycle, and several amino acid biosynthesis pathways were not identified in previous

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**FIG 2** Steady-state H2O2 stress can elicit profound transcriptional changes in bacteria. (A) Comparison of numbers of changed genes between previous microarray analyses based on one-dose treatment with H2O2 and the current RNA-seq analyses using steady-state treatment with H2O2. (B and C) GO enrichment analysis of all genes in \textit{S. aureus} upregulated or downregulated, respectively, by a steady dose of H2O2. (D and E) GO enrichment analysis of all genes in \textit{P. aeruginosa} upregulated or downregulated, respectively, by a steady dose of H2O2. Pathways that are not reported in previous microarray analyses are highlighted in red. All \(P\) values of the nodes are \(<0.005\).
one-dose-based microarray analyses (highlighted in red in Fig. 2B and C), indicating that these pathways are affected in response to a steady level of H$_2$O$_2$.

The GO enrichment analysis of all changed genes in _P. aeruginosa_ generated more pathways with complicated patterns (Fig. 2D and E). Notably, in the presence of a steady-state level of H$_2$O$_2$, expression of genes coding for proteins involved in glycolysis, the pentose phosphate pathway, flagella, and alginic acid was induced, whereas expression of genes involved in translation, protein disassembly, and thiamine was repressed. Many of these genes were not responsive to one-dose treatment with H$_2$O$_2$ (highlighted in red in Fig. 2D and E). RT-PCR was performed for genes that are involved in flagellum and alginate pathways. As shown in Fig. S2E in the supplemental material, all tested genes (filC, filD, fliE, algU, algG, and mucA) were induced 1.5- to 2-fold by a steady-state level of H$_2$O$_2$ for 10 min. However, these expression changes did not translate into a phenotypic level (by either the swelling or biofilm test [data not shown]). We also took advantage of a recent annotation of _P. aeruginosa_ small RNAs (sRNAs) to identify 18 sRNAs that were differentially transcribed under the steady-state stress of H$_2$O$_2$, such as transfer-messenger RNA (tmRNA), _rapB_, and _crcZ_ (51) (see Table S4 in the supplemental material).

Steady-state H$_2$O$_2$ treatment leads to derepression of glycolytic genes through GapR in _S. aureus_. Among the pathways that were highlighted in the GO enrichment analysis, genes encoding glycolytic enzymes were induced in both tested bacterial strains by steady-state levels of H$_2$O$_2$, suggesting that a common underlying mechanism is involved. In _S. aureus_, a total of seven genes associated with glycolysis (gapR, gapA, pgk, tpiA, pgm, fbp, and pgi) or PPP (zwf) are significantly upregulated by a steady-state treatment with H$_2$O$_2$ (2- to 4-fold) (Fig. 3A). First, to assess the reliability of RNA-seq data in representing the relative levels of individual transcripts, the identical RNA samples were subjected to real-time quantitative reverse transcription-PCR (qRT-PCR) in order to verify the mRNA levels of two glycolytic genes (gapR and gapA) and one gluconeogenic gene (fbp) with both control and steady-state treatments with H$_2$O$_2$. The qRT-PCR results matched with the corresponding RNA-seq data, ensuring its reliability in determining the transcriptional changes (Fig. 3B). We next aimed to test if the upregulated glycolytic genes would eventually lead to elevated levels of glycolytic metabolites. We utilized LC-MS/MS to quantify the intracellular concentrations of a group of glycolytic, PPP, and TCA metabolites in steady-state H$_2$O$_2$–treated versus untreated controls. The levels of phosphoenolpyruvate (PEP), glyceraldehyde-3-phosphate (GAP), 3-phosphoglycerate/2-phosphoglycerate (3PG/2PG), succinate, and citrate were significantly increased after H$_2$O$_2$ exposure in _S. aureus_ (Fig. 3C), indicating that the steady-state challenge with H$_2$O$_2$ not only induces glycolytic genes but also elevates glycolytic and early TCA cycle metabolites.

As aforementioned, we showed that steady-state levels of H$_2$O$_2$ can cause an induction of both glycolysis and the pentose phosphate pathway, following inhibition of GAPDH. In light of this, the new observation of induced glycolytic genes by the steady-state treatment with H$_2$O$_2$ suggests a potential correlation between these two events. We propose that metabolite-mediated transcriptional regulation is involved for the glycolytic genes in _S. aureus_. Although the regulatory mechanism of staphylococcal glycolytic genes is not well documented, _S. aureus_ has a glycolytic operon that is highly homologous to the well-studied counterpart in _B. subtilis_ (23). In _B. subtilis_, the glycolytic operon contains six genes, beginning with _cggR_ encoding the glycolytic repressor CggR, which tunes the transcription of itself and all other genes in the operon (gapA, pgk, tpiA, pgm, and eno) (Fig. 3D). CggR directly binds to a CggR motif consisting of two direct repeats (GGGACN$_6$TGTC-N$_4$CGGGACN$_6$TGTC) in its own promoter (23). It has also been demonstrated that fructose-1,6-biphosphate (FBP) specifically interacts with CggR and functions as its derepressor by reducing its DNA-binding activity. Like _cggR_, _gapR_ in _S. aureus_ has been shown to negatively regulate its own glycolytic operon (Fig. 3E) (22). In the present study, glucose or fructose biphosphatase (G2P/F2P), including FBP, exhibited a 4-fold accumulation upon steady-state treatment with H$_2$O$_2$ (Fig. 1A), suggesting that elevated FBP directly binds to GapR and induces the dissociation of GapR from its target DNA. We used an enzyme-based assay (see Materials and Methods) to measure the intracellular concentrations of FBP in _S. aureus_, which are 2.5 ± 0.5 mM (without H$_2$O$_2$ treatment) and 10 ± 3 mM (with a steady-state level of H$_2$O$_2$). These concentrations are comparable to the published concentration in _E. coli_ (15 mM) (52).

We expressed and purified a His$_6$-tagged full-length _S. aureus_ GapR protein from _E. coli_ grown in Luria broth. As expected, the electrophoretic mobility shift assay (EMSA) showed that GapR binds to its own promoter efficiently and specifically (Fig. 4A; see Fig. S3A in the supplemental material). The dissociation constant ($K_d$) of the interaction between GapR and its own promoter was around 0.2 μM (see Fig. S3A). Importantly, there is a noticeable change in the binding of GapR to its own promoter DNA in the presence of 2.5 mM or 10 mM FBP. The GapR-DNA complex is not sensitive to 10 mM H$_2$O$_2$, or the other inorganic phosphate (3PG) (Fig. 4A). In order to further confirm the binding site of GapR on its own promoter (also the promoter of the glycolytic operon), we performed a DNase I footprint assay by using dye primer sequencing on the Applied Biosystems 3730 DNA analyzer. We were able to uncover a specific GapR-protected region (~100 to ~56 away from ATG) on the GapR promoter (Fig. 4C). Interestingly, a putative GapR box (GAGGTTN$_6$TGTCN$_5$CGGGACN$_6$AGGC, from −94 to −58) was located in this protected region, which is very similar to the CggR motif (CGGGACN$_6$TGTCN$_6$TGTC) found in _B. subtilis_. The predicted putative GapR box is located downstream of the −55 and −10 consensus sequences (highlighted in blue in Fig. 4G), which is characteristic of negative regulation by bacterial transcription factors (53). Indeed, _gapA_ has been shown to be overexpressed in a _gapR_ mutant, proving the direct negative regulation (22).

FBP is also a coactivator for CcpA, the carbon catabolite protein in many Gram-positive bacteria (54). In order to test if CcpA is involved in the FBP-mediated induction of the _gap operon_, we repeated experiments using a _gapR_ mutant (from J. A. Morrisey at University of Leicester). Our RT-PCR assay revealed no significant induction of _gapA_ by the steady-state level of H$_2$O$_2$ in the _gapR_ mutant (see Fig. S3B in the supplemental material), which indicates that _gapR_ is responsible for the induction of the _gap operon_ in the WT.

Steady-state treatments with H$_2$O$_2$ can elevate the level of 2-keto-3-deoxy-6-phosphogluconate (KDPG), which interacts with HexR and derepresses glycolytic genes in _P. aeruginosa_. Like _S. aureus_, a group of glycolytic genes in _P. aeruginosa_ were induced with steady-state treatment with H$_2$O$_2$, such as _gapA_, _pgm_, _edd_, _pgk_, _zwf_, and _pgl_, which were verified by a subsequent...
Our metabolomic analysis also demonstrated that the levels of glycolytic and early TCA metabolites, such as PEP, 3PG/2PG, 6PG/6PF, succinate, citrate, and ATP, were significantly increased after H\textsubscript{2}O\textsubscript{2} exposure in \textit{P. aeruginosa} (Fig. 5C).

In \textit{Pseudomonas} species, glycolysis is linked with the Entner-Doudoroff (ED) pathway, which is extensively studied in \textit{Pseudomonas putida} and contains two operons (\textit{zwf-pgl-eda} and \textit{edd-glk-gltR2-gltS}) (55). The transcription of these operons is controlled by the repressor HexR, which directly binds to an inverted repeat (TTGT\textsubscript{7–8}ACAA) in these two promoters, which is released by the specific binding of the ED pathway intermediate 2-keto-3-deoxy-6-phosphogluconate (KDPG) to HexR (24). The orthologs of these glycolysis ED genes are organized in a similar way in \textit{P. aeruginosa}, which have not been well characterized (Fig. 5D). We propose that the steady-state levels of H\textsubscript{2}O\textsubscript{2} elevate the intracellular level of KDPG, which subsequently binds to HexR en route to derepression of glycolytic genes. The asterisks denote that the differences between H\textsubscript{2}O\textsubscript{2}-treated and untreated samples are statistically significant (\(P < 0.05\)).

qRT-PCR assay (Fig. 5A and B). Our metabolomic analysis also demonstrated that the levels of glycolytic and early TCA metabolites, such as PEP, 3PG/2PG, 6PG/6PF, succinate, citrate, and ATP, were significantly increased after H\textsubscript{2}O\textsubscript{2} exposure in \textit{P. aeruginosa} (Fig. 5C).

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Fructose 1,6-biphosphate (FBP) releases the repression of glycolytic genes through GapR in S. aureus. (A) EMSA shows that GapR directly binds to its own promoter. There is a noticeable dissociation of the GapR-DNA complex in the presence of 10 mM FBP (but not in the presence of 2.5 mM FBP), which is the calculated in vivo concentration after a steady dose of 10 mM H$_2$O$_2$. The GapR-DNA complex is not sensitive to either 10 mM H$_2$O$_2$ or 3PG but is sensitive to 1 mM menadione. (B) GapR directly binds to a CggR box-like motif in its own promoter according to a dye primer-based DNase I footprint assay. Electropherograms show the protection patterns of the gapR promoter after digestion with DNase I following incubation in the absence (upper panel) or presence (lower panel) of 1 μM GapR. ROI, region of interest. (C) gapR promoter sequence (−160 from ATG) with a summary of the DNase I footprint assay results. The −35 and −10 promoter regions are highlighted in blue. The GapR-protected region is underlined, and the two putative repeats in the GapR motif are highlighted in red.

FIG 4 Fructose 1,6-biphosphate (FBP) releases the repression of glycolytic genes through GapR in S. aureus. (A) EMSA shows that GapR directly binds to its own promoter. There is a noticeable dissociation of the GapR-DNA complex in the presence of 10 mM FBP (but not in the presence of 2.5 mM FBP), which is the calculated in vivo concentration after a steady dose of 10 mM H$_2$O$_2$. The GapR-DNA complex is not sensitive to either 10 mM H$_2$O$_2$ or 3PG but is sensitive to 1 mM menadione. (B) GapR directly binds to a CggR box-like motif in its own promoter according to a dye primer-based DNase I footprint assay. Electropherograms show the protection patterns of the gapR promoter after digestion with DNase I following incubation in the absence (upper panel) or presence (lower panel) of 1 μM GapR. ROI, region of interest. (C) gapR promoter sequence (−160 from ATG) with a summary of the DNase I footprint assay results. The −35 and −10 promoter regions are highlighted in blue. The GapR-protected region is underlined, and the two putative repeats in the GapR motif are highlighted in red.
tion of KDPG (from 45 ± 15 to 150 ± 28 μM) (Fig. 5E), which suggests the importance of KDPG in response to ROS. Indeed, we found that an edd (which encodes phosphogluconate dehydratase that produces KDPG) mutant was 4-fold more sensitive to H2O2 (Fig. 5F), demonstrating that KDPG is a critical metabolite involved in resistance to ROS in P. aeruginosa.

We next purified a His6-tagged full-length HexR protein that indeed strongly and specifically binds to the promoters of zwf and gapA in EMSA (Fig. 6A; see Fig. S3C in the supplemental material). Similar to the relationship between GapR and FBP in S. aureus, addition of 150 μM KDPG dissociates significantly more DNA from the HexR-DNA complex than does 50 μM KDPG (Fig. 6A). As a control, the HexR-DNA complexes are not sensitive to 10 mM H2O2. The subsequent DNase I footprint assay further confirmed two specific HexR-protected regions (−80 to −4 away from ATG) on the zwf promoter (Fig. 6B and C). Two putative HexR boxes (TTGn5ACTA, from −78 to −64, and TTGn5CAA, from −18 to −4, respectively) (Fig. 6C) were located in these protected regions, which are very similar to the HexR motif (TTGn5ACTA) in P. putida. These two predicted putative
HexR boxes are located downstream of the −35 and −10 consensus sequences (highlighted in blue in Fig. 6C).

Since the steady-state levels of H$_2$O$_2$ induced both the gene expression and intracellular levels of metabolites in the glycolysis ED pathway, we reason that continuous treatment with H$_2$O$_2$ might boost the glucose uptake in *P. aeruginosa* as well. In order to test this hypothesis, the glucose uptake rates were calculated by measuring the concentration of the remaining glucose in the growth medium posttreatment with IAA for 1 h. Since H$_2$O$_2$ interfered with the measurement of glucose, IAA was used to mimic the inhibitory effect of GAPDH. Indeed, IAA significantly induced the glucose uptake rate, while inhibiting bacterial growth (Fig. 6D and E). On the other hand, GAPDH inhibition did not induce glucose uptake in *S. aureus* (data not shown). We noted that, unlike in *P. aeruginosa*, genes involved in glucose uptake in *S. aureus* (glk, glcU, and glcA) were not significantly induced under the steady-state level of H$_2$O$_2$, which could explain the discrepancy between the two bacteria (see Table S2 in the supplemental material).

**DISCUSSION**

A key measure for bacteria to counteract oxidative damage is to maintain the intracellular redox state, which is mostly governed by ratios of NADH to NAD$^+$ and NADPH to NADP$^+$. NADH and NADPH fuel the antioxidant activities of alkyl hydroperoxidase and glutathione and thioredoxin reductases. NADH acts as a pro-oxidant that feeds reducing equivalents to flavoproteins (56). PPP is the major pathway for NADPH production, which raises the bacterial electrochemical potential that is involved in antioxidant tolerance. A similar role of GAPDH as a metabolic
switch has been previously characterized in yeast, illustrating evolutionary conservation of this strategy in both prokaryotes and eukaryotes (25, 57). The altered levels of metabolites might also function as antioxidant signals. The steady-state level of H$_2$O$_2$ significantly induces a group of metabolic pathways, including fatty acid, tryptophan, and pyrimidine metabolism (Fig. 2), which could serve as a “sink” to dispose PPP intermediates and produce excess NADPH. In addition, H$_2$O$_2$ helps to consume NADPH that is responsible for GSH production. Moreover, we show that the inactivation of GAPDH decreases the intracellular NADH level that contributes to the proton motive force (PMF) and promotes the survival of bacterial persisters that are treated by aminoglycosides and glycolytic metabolites. Although this strategy is limited to only one category of bacteria, it provides a general framework for understanding how prokaryotes can counteract oxidative stress to ensure survival.
of antibiotics, our findings imply that oxidative stress would be beneficial for persister to escape killing.

Previously, we found that *P. aeruginosa* and *S. aureus* were able to largely degrade millimolar levels of *H₂O₂* in several minutes in the mid-log-phase culture *in vitro* (16). However, inside the host, *H₂O₂* can be produced constitutively by the immune response with steady doses (1), which suggests that *H₂O₂* should be continuously supplemented *in vitro* in order to best mimic the physiological conditions under the host immune response. The rationale was validated by the observation of induced glycolysis inside these bacteria when challenged with a steady-state level of *H₂O₂* but not the one-dose *H₂O₂* treatment (Fig. 1A; see Fig. S1A in the supplemental material) commonly used in studying ROS sensing and response in bacteria (26). With this in mind, previous microarray analyses that are based on one-dose *H₂O₂* treatment may not comprehensively represent the *in vivo* transcriptomic changes elicited by host-derived ROS (26, 27, 47). Therefore, we used a dialysis bag to continuously treat bacteria with 10 mM *H₂O₂* before employing RNA-seq to globally profile transcriptional changes.

Interestingly, our present RNA-seq analyses uncovered that glycolytic genes were upregulated and significantly enriched for both tested bacteria. The subsequent biochemical and genetic characterizations demonstrated that following *H₂O₂* inactivation of GAPDH, which induces glycolysis and PPP, the elevated levels of FBP and KDPG lead to dissociation of their corresponding glycolytic repressors (GapR and HexR, respectively) from the cognate promoters, thus resulting in derepression of the glycolytic genes to overcome *H₂O₂*-stalled glycolysis in *S. aureus* and *P. aeruginosa*, respectively (Fig. 7A and B). There is increasing evidence that metabolites are important in modulating glycolytic flux (58). The metabolite quantification further confirmed that the glycolysis is activated by the steady-state treatments with *H₂O₂* for both pathogenic bacteria (Fig. 3C and 5C). This is reminiscent of the observation that zwf was found to be induced by oxidative stress in *Pseudomonas putida* (59). The similar ROS-induced glycolysis has been well documented in eukaryotic systems, including yeast and cancer cells (60, 61). The most intriguing example is the one-dose *H₂O₂* treatment (Fig. 1A; see Fig. S1A in the supplemental material) commonly used in studying ROS sensing and response in bacteria (26). Within this in mind, previous microarray analyses that are based on one-dose *H₂O₂* treatment may not comprehensively represent the *in vivo* transcriptomic changes elicited by host-derived ROS (26, 47). Therefore, we used a dialysis bag to continuously treat bacteria with 10 mM *H₂O₂* before employing RNA-seq to globally profile transcriptional changes.

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We declare that we have no conflicts of interest.

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Steady-State H$_2$O$_2$ Induces Glycolysis in Bacteria

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