Staphylococcus aureus HemX Modulates Glutamyl-tRNA Reductase Abundance To Regulate Heme Biosynthesis

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ABSTRACT Staphylococcus aureus is responsible for a significant amount of devastating disease. Its ability to colonize the host and cause infection is supported by a variety of proteins that are dependent on the cofactor heme. Heme is a porphyrin used broadly across kingdoms and is synthesized de novo from common cellular precursors and iron. While heme is critical to bacterial physiology, it is also toxic in high concentrations, requiring that organisms encode regulatory processes to control heme homeostasis. In this work, we describe a posttranscriptional regulatory strategy in S. aureus heme biosynthesis. The first committed enzyme in the S. aureus heme biosynthetic pathway, glutamyl-tRNA reductase (GtrR), is regulated by heme abundance and the integral membrane protein HemX. GtrR abundance increases dramatically in response to heme deficiency, suggesting a mechanism by which S. aureus responds to the need to increase heme synthesis. Additionally, HemX is required to maintain low levels of GtrR in heme-proficient cells, and inactivation of hemX leads to increased heme synthesis. Excess heme synthesis in a ΔhemX mutant activates the staphylococcal heme stress response, suggesting that regulation of heme synthesis is critical to reduce self-imposed heme toxicity. Analysis of diverse organisms indicates that HemX is widely conserved among heme-synthesizing bacteria, suggesting that HemX is a common factor involved in the regulation of GtrR abundance. Together, this work demonstrates that S. aureus regulates heme synthesis by modulating GtrR abundance in response to heme deficiency and through the activity of the broadly conserved HemX.

IMPORTANCE Staphylococcus aureus is a leading cause of skin and soft tissue infections, endocarditis, bacteremia, and osteomyelitis, making it a critical health care concern. Development of new antimicrobials against S. aureus requires knowledge of the physiology that supports this organism’s pathogenesis. One component of staphylococcal physiology that contributes to growth and virulence is heme. Heme is a widely utilized cofactor that enables diverse chemical reactions across many enzyme families. S. aureus relies on many critical heme-dependent proteins and is sensitive to excess heme toxicity, suggesting S. aureus must maintain proper intracellular heme homeostasis. Because S. aureus provides heme for heme-dependent enzymes via synthesis from common precursors, we hypothesized that regulation of heme synthesis is one mechanism to maintain heme homeostasis. In this study, we identify that S. aureus posttranscriptionally regulates heme synthesis by restraining abundance of the first heme biosynthetic enzyme, GtrR, via heme and the broadly conserved membrane protein HemX.

KEYWORDS Staphylococcus aureus, heme, tetrapyrroles
The tetrapyrrole cofactor heme is critical to the physiology of organisms from humans to bacteria. Heme is composed of a porphyrin ring complexed to iron at its center, making it an excellent redox-active moiety for a variety of enzymes. Across kingdoms, heme is used to shuttle electrons in the respiratory chain and is also required for the function of many critical proteins, including nitric oxide synthase, catalase, and hemoglobin. To satisfy the cellular need for heme, most heme-dependent organisms synthesize heme de novo from simple and abundant precursors.

The versatility of heme as a cofactor is based on its reactivity, which also results in its toxicity at high concentrations. Excess heme can cause damage to cellular macromolecules, and the redox cycling of heme-iron produces reactive oxygen species via Fenton chemistry. Bacteria encode a variety of mechanisms to resist heme toxicity, but the most important of these strategies may be the prevention of self-imposed toxicity by regulating endogenous heme synthesis. A variety of transcriptional and posttranscriptional strategies have evolved to regulate heme synthesis centered around providing sufficient heme to occupy hemoproteins while preventing excess heme synthesis to limit unnecessary consumption of substrates and preclude toxicity.

In this study, we sought to uncover regulatory pathways controlling heme synthesis in the human pathogen *Staphylococcus aureus*. *S. aureus* is a Gram-positive bacterium that causes a variety of devastating diseases, including skin and soft tissue infections, osteomyelitis, endocarditis, and bacteremia. *S. aureus*, as a facultative anaerobe, generates energy through aerobic respiration, anaerobic respiration, or fermentation. The final step in aerobic respiration is reduction of oxygen to water, which *S. aureus* performs with either of the heme-dependent QoxABCD or CydAB terminal oxidases. Although a great deal is known about heme synthesis, heme utilization, and heme toxicity in *S. aureus*, no heme synthesis regulatory pathway has been identified in this organism. *S. aureus* encodes the newly appreciated coproporphyrin-dependent heme synthesis pathway to populate its hemoproteins. These include the terminal oxidases, catalase, and bacterial nitric oxide synthase, all of which contribute to growth, protection from host defenses, and pathogenesis. Under conditions of excess exogenous heme, the heme stress response in *S. aureus* is activated by the heme-sensing two-component system HssRS, which regulates the transcription of a putative efflux pump, HrtAB. This system is critical for growth and survival in toxic concentrations of heme and modulates pathogenesis in a murine model of disease. Sensing or regulatory pathways that connect heme synthesis with heme availability, hemoprotein abundance, or HssRS activation have not been identified.

*S. aureus* synthesizes δ-aminolevulinic acid (ALA), the first dedicated and universal precursor for protoheme synthesis, via the conversion of glutamyl-tRNA to glutamate-1-semialdehyde by glutamyl-tRNA reductase (GtrR) and subsequent production of ALA by glutamate-1-semialdehyde 2,1-aminomutase. Uroporphyrinogen is the precursor to heme, siroheme, and other tetrapyrroles, and the stepwise transformation of ALA to uroporphyrinogen comprises the core of the synthesis pathway. The pathway from uroporphyrinogen to heme was historically considered to be universally conserved for all organisms. However, the field’s understanding of bacterial heme synthesis has undergone a revolution as recent studies uncovered diversity in bacterial strategies to convert uroporphyrinogen to heme (reviewed in reference 14). Gram-positive bacteria proceed through a coproporphyrin-dependent branch that is unique from the classic protoporphyrin-dependent branch in humans and many Gram-negative model organisms.

In this work, we identify GtrR abundance as a critical regulator of *S. aureus* heme synthesis. GtrR is posttranscriptionally maintained at low abundance in heme-proficient cells by the membrane protein HemX, but levels increase when *S. aureus* is deprived of heme. Without HemX regulation, GtrR abundance increases, which results in the concomitant increase in flux through the heme synthesis pathway and accumulation of heme. This excess heme synthesis activates HssRS and disrupts iron homeostasis. Together, this report reveals that *S. aureus* regulates heme synthesis by modulating GtrR abundance via intracellular heme levels and the widely conserved HemX.
GtrR abundance increases specifically in response to heme deficiency. To identify key steps in the regulation of heme synthesis (Fig. 1A and B), we measured the abundance of each biosynthetic enzyme by liquid chromatography-multiple reaction monitoring-tandem mass spectrometry (LC-MRM-MS/MS). This technique allows for quantification with high resolution of even very-low-abundance cellular proteins (16).

We hypothesized that comparing the *S. aureus* wild type (WT) to a strain incapable of synthesizing heme (*pbgS* mutant) (Fig. 1A and B) would allow the identification of specific steps in heme synthesis that respond to cellular heme content, directly or

**RESULTS**

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indirectly. Abundance of GtrR is approximately 30-fold higher in the \textit{pbgS} mutant relative to the WT, while the abundances of all other biosynthetic enzymes are nearly unchanged (Fig. 1C). In WT cells, GtrR abundance is low relative to other heme synthesis enzymes (see Fig. S1 in the supplemental material). The \textit{pbgS} mutant is a heme auxotroph and therefore adopts the respiration-deficient small-colony variant (SCV) phenotype. SCVs arise as the result of inactivation of respiration via inactivation of heme synthesis, the terminal oxidases, or the electron carrier menaquinone, and SCVs have a dramatically different physiology than respiration-proficient cells (17). Therefore, we sought to determine whether the increase in GtrR abundance in the \textit{pbgS} mutant was the result of heme deficiency or a general defect in respiration. To confirm that the menaquinone auxotroph SCV \textit{ΔmenB} strain and the \textit{ΔqoxB cydB} strain lacking both terminal cytochrome oxidases synthesize heme despite being unable to respire, each strain was streaked onto agar and assessed for catalase activity (Fig. 1D). Activity of the heme-dependent catalase KatA leads to the production of oxygen bubbles when hydrogen peroxide is added. The \textit{ΔmenB} and \textit{ΔqoxB cydB} mutants produce bubbles, demonstrating that these SCVs synthesize heme and are not heme auxotrophs. We measured GtrR abundance in a variety of SCV strains by LC-MRM-MS/MS. GtrR abundance increases relative to the WT only in \textit{pbgS} and \textit{ΔchdC} strains (Fig. 1B and E), which are heme auxotroph SCVs (Fig. 1D). When chemically complemented with heme, GtrR abundance returned to WT levels for both strains. GtrR levels do not increase in the \textit{ΔmenB} or \textit{ΔqoxB cydB} strain. Together these data demonstrate that the abundance of GtrR is low in heme-proficient cells but increases specifically in response to heme deficiency.

**HemX controls GtrR abundance in heme-proficient cells to regulate heme synthesis.** Among both Gram-negative and Gram-positive bacteria, regulation of GtrR abundance is a common feature of heme synthesis regulation pathways (18–20). In the model organism \textit{Bacillus subtilis}, which is also a member of the \textit{Firmicutes} phylum, GtrR abundance is impacted by the membrane protein HemX through an unknown mechanism (18, 21). While \textit{S. aureus} is in the same \textit{Bacillales} order as \textit{B. subtilis}, \textit{S. aureus} heme homeostasis is distinct because of its access to host heme and its resistance to heme toxicity mediated by HssRS. Both \textit{B. subtilis} and \textit{S. aureus} carry an operon comprised of gtrR-hemX-hmbS-uroS-pbgS-gsaM (formerly hemAXCBDL) (22, 23). We therefore hypothesized that in \textit{S. aureus}, HemX also impacts GtrR abundance in heme-proficient cells. We created an in-frame unmarked deletion of \textit{hemX} and integrated either pJC1111 \textit{P}_{lg} or \textit{P}_{lg} \textit{hemX} at a neutral site in the chromosome (24). GtrR abundance was measured by LC-MRM-MS/MS and is increased in the \textit{ΔhemX::P}_{lg} strain relative to WT::\textit{P}_{lg} (Fig. 2A). The phenotype can be complemented when \textit{hemX} is provided in cis, showing that it is the result of deletion of \textit{hemX} and not other effects of disrupting the operon. These data are consistent with the hypothesis that HemX regulates GtrR abundance in heme-proficient cells (18, 21).

We next sought to determine whether the increase in GtrR at the protein level in both the \textit{pbgS} and \textit{ΔhemX} strains is the result of an increase in mRNA transcript abundance of \textit{gtrR}. Therefore, the \textit{pbgS} and \textit{ΔhemX} strains were grown to the mid-exponential phase, and RNA was isolated, converted to cDNA, and quantified by quantitative PCR (qPCR) (Fig. 2B). The steady-state mRNA abundance of \textit{gtrR} transcript is unchanged in the \textit{ΔhemX} or \textit{pbgS} strain relative to the WT, suggesting that the increase in GtrR abundance in these strains is not the result of a transcriptional change. Additionally, the increase in GtrR levels in the \textit{pbgS} strain is not affected by the insertion of \textit{ermB} to interrupt the \textit{pbgS} gene, which is upstream of \textit{gsaM} in the operon; there is no change in the transcript abundance of \textit{gsaM} in the \textit{pbgS} strain relative to the WT (see Fig. S2 in the supplemental material).

We hypothesized that the increase in GtrR observed in the \textit{ΔhemX} strain would increase the amount of heme synthesized by increasing abundance of the heme precursors downstream of GtrR. As glutamate-1-semialdehyde is unstable and can convert to δ-aminolevulinic acid (ALA) in the absence of enzyme (25), we measured ALA abundance via a colorimetric method. ALA abundance increases approximately 50% in
the ΔhemX strain relative to the WT (Fig. 2C). We subsequently sought to determine the impact of increased ALA availability on downstream heme intermediates and heme abundance. Total cellular porphyrins were extracted from the WT and ΔhemX strains and analyzed by quantitative exact-mass liquid chromatography-quadrupole time of flight mass spectrometry (LC-qTOF-MS); total extracted ion chromatograms for porphyrins that were observed above the limits of detection are shown in Fig. S3 in the supplemental material, where a dramatic change in porphyrin levels is visible. Based on standard curves for individual porphyrins (including PbgS, UroP, CprP, CopP, and Heme) and enumeration of colony-forming units, absolute quantifications were obtained and

**FIG 2** HemX regulates heme synthesis by maintaining low levels of GtrR in heme-proficient cells. (A) The abundance of GtrR was measured by LC-MRM-MS/MS in multiple *S. aureus* strains. The data are the average from a single experiment performed in biological triplicate with standard deviation shown. Statistical significance was determined using a one-way ANOVA with Dunnett's correction for multiple comparisons, comparing GtrR abundance for each strain relative to the ΔhemX::Plgt strain. ***, P < 0.001.** (B) Steady-state transcript abundance of gtrR mRNA isolated from mid-exponential growth of *S. aureus* strains was measured by qRT-PCR and is graphed as fold change relative to the WT. Data are combined from two independent experiments in biological triplicate with standard deviation shown. "ns" indicates no significance by one-way ANOVA with Dunnett's correction for multiple comparisons, comparing fold change of the pbgS and ΔhemX strains to the WT. (C) ALA abundance was measured in *S. aureus* strains by a spectrophotometric quantification. Graphed is the fold change of ALA in the ΔhemX mutant relative to the WT, with data combined from two independent experiments with three biological replicates with standard error of the mean shown. (D) Uroporphyrinogen III (detected as uroporphyrin III), coproporphyrin III, coproheme III, and heme were quantified by LC-qTOF-MS. Graphed is the fold change of metabolite abundance in the ΔhemX mutant relative to the WT, from a single experiment performed in biological triplicate with standard error of the mean shown. For panels C and D, statistical significance was determined with Student's *t* test comparing the ΔhemX mutant to the WT before data transformation to fold change. *, *P* < 0.05; ***, *P* < 0.0001.
referred to by their fold change relative to the WT. As shown in Fig. 2D; the \( \Delta \text{hem}X \) strain exhibits increased abundance of uroporphyrin III, coproporphyrin III, coproheme III, and heme b relative to the WT. Notably, because samples were prepared aerobically, the metabolite uroporphyrinogen III was detected as uroporphyrin III, in which its methylene bridge carbons have spontaneously oxidized in air; likewise, any coproporphyrinogen III that might have been present would be detected as the oxidation product, coproporphyrin III, which is also the product of the enzyme CgoX (Fig. 1). Hydroxymethylbilane spontaneously cyclizes to uroporphyrinogen I, which is decarboxylated by UroD to coproporphyrinogen I. The absence of uroporphyrin or coproporphyrin I isomers indicates that hydroxymethylbilane did not accrue in the \( \Delta \text{hem}X \) mutant. We hypothesize that, in the presence of excess GtrR, the initial step of the pathway may no longer be rate limiting. This may allow other subsequent steps in the pathway to become partly rate limiting, leading to the observed pattern of metabolite accumulation. Finally, the increase in heme abundance in the \( \Delta \text{hem}X \) strain observed by LC-qTOF-MS is complemented when \( \text{hem}X \) is provided in cis from a neutral site in the chromosome, as measured by the pyridine hemochromagen method (see Fig. S4A in the supplemental material). Together, these data demonstrate that inactivation of \( \text{hem}X \) results in increased GtrR abundance, which increases abundances of both early- and late-pathway biosynthetic precursors and cellular heme. Therefore, dysregulation of GtrR alone is sufficient to disrupt heme homeostasis.

**Excess endogenous heme synthesis in the \( \Delta \text{hem}X \) mutant activates the heme stress response.** The unregulated GtrR abundance in the \( \Delta \text{hem}X \) mutant results in greater cellular heme levels (Fig. 2D), which we hypothesized would activate the heme sensor system HssRS in the absence of exogenous heme, leading to transcriptional induction of the hrtAB efflux pump. The WT and \( \Delta \text{hem}X \) mutant strains were transformed with plasmids containing the luminescence-producing operon \( \text{luxABCDE} \) cloned from *Photorhabdus luminescens* without a promoter (pXen-1) or controlled by the HssRS-regulated promoter \( \text{P}_{\text{hrt}} \). \( \text{P}_{\text{hrt}} \) promoter activity, visualized by luminescent imaging, shows that HssRS is activated in the \( \Delta \text{hem}X \) strain in the absence of exogenous heme, whereas HssRS is not activated in the WT (Fig. 3A). The \( \text{P}_{\text{hrt}} \) activity in WT becomes apparent when 20 \( \mu \text{M} \) exogenous heme is added to the agar medium, and luminescence depends on the heme-responsive \( \text{P}_{\text{hrt}} \). To more quantitatively measure \( \text{P}_{\text{hrt}} \) activity as a readout of HssRS activation by endogenous heme, we transformed WT, \( \Delta \text{hem}X \), and \( \Delta \text{hssRS} \) strains with the pOS1 \( \text{P}_{\text{hrt}} \text{XylE} \) plasmid. These strains report \( \text{P}_{\text{hrt}} \) activity with the production of the XylE catechol oxidase enzyme, which can be quantified spectrophotometrically from cell lysate. Data in Fig. 3B demonstrate that in the absence of exogenous heme, \( \text{P}_{\text{hrt}} \) is induced in the \( \Delta \text{hem}X \) strain. \( \text{P}_{\text{hrt}} \) activity does increase in the WT and \( \Delta \text{hem}X \) strains in a dose-dependent manner as exogenous heme is added, but \( \text{P}_{\text{hrt}} \) activity remains higher in the \( \Delta \text{hem}X \) mutant than the WT at all tested heme concentrations. Additionally, XylE activity in this system is dependent on the HssRS two-component system. Taken together, these data suggest that excess endogenous heme synthesized in the \( \Delta \text{hem}X \) strain is sufficient to activate the HssRS two-component system.

We next hypothesized that the intermediate levels of HssRS activation in the \( \Delta \text{hem}X \) mutant, in the absence of exogenous heme (Fig. 3A and B), would be sufficient to preadapt the \( \Delta \text{hem}X \) mutant to heme toxicity. As the HssRS-HrtAB heme stress response provides resistance to heme toxicity, pretreatment with subtoxic concentrations of heme adapts *S. aureus* to subsequent growth in toxic concentrations of heme by activating HssRS and increasing the abundance of HrtAB (10). The WT grown in 10 \( \mu \text{M} \) heme without preadaptation has a severe growth defect evident by a 6-h lag time (Fig. 3C). When preadapted in 2 \( \mu \text{M} \) heme, the WT demonstrates a reduced lag time and greater overall growth, albeit at a lower rate and yield than when grown without heme. In contrast, the \( \Delta \text{hem}X \) mutant grown in 10 \( \mu \text{M} \) heme with or without preadaptation exhibits increased growth compared to the WT. The enhanced growth of the \( \Delta \text{hem}X \) mutant in 10 \( \mu \text{M} \) heme is dependent on the HrtAB efflux system, as the \( \Delta \text{hem}X \)
The ΔhemX strain does not grow in 10 μM heme (Fig. 3C). Additionally, preadaptation of the ΔhemX strain in this assay can be complemented by providing hemX in the chromosome (Fig. S4B). Similarly, the ΔgtrR-hemX pos1 P_{xyIE} strain is resistant to heme toxicity, but becomes sensitive again when hemX is introduced on the plasmid (Fig. S4C). This is further evidence that HemX control of GtrR is not transcriptional, as HemX exerts its effect independent of the native gtrR promoter and ribosome binding site in this assay. Further, the ΔhemX mutant is resistant to the bactericidal effects of acute heme toxicity, compared to a 4-log reduction in viable WT cells after 2 h in the presence of 20 μM heme (Fig. 3D). In sum, these data demonstrate that increased cellular heme in the ΔhemX mutant is sufficient to activate HssRS and cause expression of HrtAB, which leads to resistance to heme toxicity.

**Excess heme synthesis disrupts iron homeostasis.** Considering that every molecule of heme contains an atom of iron, we hypothesized that unregulated heme synthesis disrupts iron homeostasis.

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**FIG 3** Excess heme synthesis in the ΔhemX mutant activates the heme stress response. (A) Bioluminescence was imaged on agar medium plates containing vehicle or heme onto which strains were streaked. All four plates were imaged simultaneously, and luminescence was converted to a heat map with the scale shown on the right. (B) XyE catechol oxidase activity was measured in S. aureus strains after growth in vehicle or increasing concentrations of heme. The data are the average from three independent experiments each in biological triplicate with standard deviation shown. Statistical significance was determined using a two-way ANOVA with Tukey’s correction for multiple comparisons, comparing log-transformed data for the WT ΔhemX pOS1 P_{XYIE} strain at each heme concentration to that of each other strain. * P < 0.01; ** P < 0.001; *** P < 0.0001. (C) Growth as measured by OD_{600} was monitored over time for S. aureus strains in medium containing either vehicle or 10 μM heme. Prior to the measured growth, the strains were pregrown to the stationary phase in medium containing vehicle or 2 μM heme. The data are the average of the means from at least three independent experiments each in biological triplicate with standard error of the mean shown. (D) Viable bacteria from S. aureus strains were enumerated after incubation for 2 h in medium containing vehicle or increasing amounts of heme. The data are the average of the means from three independent experiments each in biological triplicate with standard error of the mean shown. The y axis is set to the limit of detection. Statistical significance was determined using a two-way ANOVA with Tukey’s correction for multiple comparisons, comparing log-transformed data for the WT and ΔhrtB strains to the ΔhemX mutant at each heme concentration. * P < 0.01; *** P < 0.0001.
synthesis in the ΔhemX mutant would consume high levels of iron and alter iron homeostasis. To test this hypothesis, growth in minimal medium containing the iron chelator EDDHA [ethylenediamine-N,N′-bis(2-hydroxyphenylacetic acid)] was compared to growth in minimal medium alone. As shown in Fig. 4A, the ΔhemX::P_{lgt} strain demonstrates reduced total yield after growth for 24 h relative to WT::P_{lgt} and the complemented ΔhemX::P_{lgthemX} strain. To corroborate this finding, we assessed promoter activity using a P_{isdA}gfp reporter plasmid. P_{isdA} is controlled by the ferric uptake regulator (Fur) and is derepressed under iron-depleted conditions (26). Data in Fig. 4B show that after growth in rich medium with an alternative iron chelator, 2,2-dipyridyl, the ΔhemX pOS1 P_{lgdA}gfp strain has enhanced P}_{isdA} activity relative to WT pOS1 P_{lgdA}gfp. These data suggest excess heme synthesis depletes the cell of available iron.

**Inactivation of HemX reduces GtrR abundance in heme deficiency.** Based on the observations that HemX and cellular heme both impact GtrR abundance, we hypothesized that measuring GtrR abundance in a strain lacking hemX and unable to synthesize heme would uncover the nature of the relationship between HemX, heme, and GtrR. Surprisingly, GtrR abundance in the ΔhemX pbgS strain is unchanged from that in the ΔhemX strain and lower than that in the pbgS strain (Fig. 5A), and this effect is not...
the result of a change in gtrR transcription in the ΔhemX pbgS strain relative to the pbgS strain (see Fig. S5 in the supplemental material). Similarly, the ΔhemX ΔchdC mutant has lower levels of GtrR than the ΔchdC mutant (Fig. 5A). To corroborate these findings, we measured total cellular porphyrins by LC-qTOF-MS as in Fig. 2; total extracted ion chromatograms are shown in Fig. S6 in the supplemental material. Consistent with the abundance of GtrR, porphyrin intermediates are drastically increased in the ΔhemX mutant relative to the WT. The ΔchdC mutant demonstrates intermediate buildup through coproheme because of elevated GtrR levels but is unable to convert coproheme to heme (Fig. 1A). As expected, based on the reduced GtrR abundance shown in Fig. 5A, the porphyrin intermediates are at lower levels in the ΔhemX ΔchdC mutant relative to the ΔhemX or ΔchdC mutant. These data suggest that heme and HemX do not independently and directly repress GtrR levels, because if so, removal of both would likely have an additive effect on GtrR abundance. Instead the relationship between HemX, heme synthesis, and GtrR levels is still unclear. However, the data are consistent with a model whereby the increase in GtrR levels in heme-deficient strains is dependent on the activity of HemX.

Siroheme synthesis impacts GtrR levels under conditions of nitrite reduction.

The increase in GtrR levels identified in the ΔhemX mutant likely impacts siroheme synthesis, as the cofactor siroheme is synthesized in *S. aureus* from the shared uroporphyrinogen III intermediate (14) (Fig. 1A). We therefore hypothesized that siroheme levels might also affect GtrR abundance. In the experiments presented thus far, in which *S. aureus* is grown aerobically, siroheme has likely not been synthesized. The siroheme synthesis and siroheme-dependent nitrite reductase genes are transcribed primarily under anaerobic conditions (27). To therefore test the role of siroheme, we first identified conditions under which siroheme synthesis via CysG and siroheme-dependent nitrite reduction by the NirD nitrite reductase were important for growth. As demonstrated in Fig. S7 in the supplemental material, when grown anaerobically, the growth of WT is enhanced when the terminal electron acceptor nitrate is provided. Mutants lacking cysG or nirD cannot grow to WT levels when nitrate is provided, suggesting that WT cells synthesize siroheme and utilize it in NirD. It is thought that under these conditions, the anaerobic nitrate reductase will reduce nitrate to nitrite, followed by NirD-dependent reduction of nitrite. Deletion of hemX does not overtly impact nitrite reduction, as the ΔhemX strain grows well in nitrate, the ΔhemX nirD strain phenocopies the nirD strain, and the ΔhemX cysG strain phenocopies the cysG strain. Therefore, GtrR abundance was measured by LC-MRM-MS/MS after growth in tryptic soy broth (TSB) containing nitrate (Fig. 6). The strain lacking cysG, which can
make heme but not siroheme, does not demonstrate elevated GtrR levels. However, the ΔhemX cysG strain has reduced levels compared to the ΔhemX strain, suggesting that siroheme synthesis could impact GtrR regulation.

**HemX co-occurs with capacity for heme biosynthesis, and the corresponding genes often colocalize on the chromosome.** We hypothesized that *B. subtilis* HemX and *S. aureus* HemX might represent only a subset of HemX homologues that exist across bacterial phyla and function to regulate heme synthesis. Diverse genomes from 978 organisms (924 bacterial and 54 archaeal) were analyzed for the presence of *hemX*. Of these, 113 encode HemX; representative members of this analysis are shown in Fig. 7A. These newly identified homologues expand past the *Bacillales* order, of which representative HemX homologues were previously identified and shown to share function (13). HemX appears to represent an ancient protein family, as it is present in some of the evolutionarily oldest taxa, including *Firmicutes*, *Aquificae*, and *Planctomycetes*. The distribution of *hemX* strongly correlates with the capacity for de novo heme synthesis, as *hemX* never occurs in a genome without *gtrR*, and *hemX* never occurs without the capacity for de novo heme synthesis. This correlation holds across the microbial kingdom, where *hemX* never occurs in taxa lacking heme biosynthesis genes (within any representatives with sequenced genomes now available): e.g., *Thermotogae*, *Fusobacteria*, and *Mollicutes*. Additionally, the distribution of *hemX* among the members of the *Firmicutes* phylum supports this correlation; *hemX* is present largely in *Bacillales* but does not occur in *Lactobacillales* and only rarely in *Clostridia* (in 2 out 91 genomes analyzed), which is consistent with the frequent capacity for heme synthesis in *Bacillales* relative to *Lactobacillales* and *Clostridia*. Notably, the genomic co-occurrence of *hemX* and *gtrR* holds true in organisms that synthesize heme via any of the 3 heme biosynthetic pathways identified to date: the coproporphyrin-dependent, siroheme-dependent, or classic protoporphyrin-dependent route (Fig. 7A).

Interestingly, HemX is more commonly found in organisms that encode the ability to synthesize both heme and siroheme (see Fig. S8A in the supplemental material) than in organisms that synthesize heme and not siroheme. This suggests that HemX likely impacts siroheme synthesis as a consequence of affecting GtrR abundance by increasing abundance of uroporphyrinogen III, the final shared biosynthetic intermediate. The co-occurrence of *hemX*, *gtrR*, and *cysG* is also consistent with our findings in Fig. 6 that siroheme synthesis impacts GtrR abundance.

Next we examined the genomic context of *hemX* homologues across 113 organisms carrying *hemX*, as genes associated with the same pathway or area of metabolism tend to colocalize in prokaryotic genomes (28). In 106 (94%) out of these 113 genomes, the
FIG 7  hemX is conserved across bacterial phyla and invariably co-occurs (A) and colocalizes (B) with gtrR. (A) The occurrence of hemX (stars), gtrR (circles), and alaS (squares) homologues (outermost rings) was mapped onto the tree of life (S3). The pathway by which protoheme is synthesized in each of the analyzed organisms is presented in the middle ring as follows (adapted from reference 14): the classic protoporphyrin-dependent pathway (teal), coproporphyrin-dependent path (purple), or siroheme-dependent path (gold). Gray

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hemX and gtrR genes are adjacent and likely cotranscribed, which is a very strong indicator of their functional association. Seven representatives of these organisms are shown in Fig. 7B, highlighting the common genomic context of hemX, gtrR, and other genes involved in uroporphyrinogen synthesis (hmbS, uroS, pbgS, and gsaM) and siroheme synthesis (cysG).

**HemX homologues share predicted membrane topology and residues.** Comparative genome analysis identified several contextual characteristics of HemX homologues. We therefore investigated the sequences of representative HemX homologs. A multiple-sequence alignment revealed relatively low overall identity among HemX sequences (Fig. 7B); however, the alignment presented in Fig. S8B shows that each HemX homologue shares the same predicted eight-transmembrane-domain topology with N- and C-termini predicted to be extracytoplasmic. Additionally, these divergent homologues share four conserved residues, all in predicted transmembrane domains. Taken together with comparative genome analysis, identification of HemX across bacteria uncovered a strong correlation between gtrR, hemX, and de novo heme synthesis, suggesting that HemX control of GtrR to modulate heme synthesis is a common regulatory strategy among bacteria.

**DISCUSSION**

In this report, we identify GtrR abundance as a critical regulator of S. aureus heme biosynthesis. GtrR catalyzes the initial step in the heme biosynthetic pathway and is maintained at low levels in WT cells proficient for heme biosynthesis, but specifically increases in response to heme deficiency (Fig. 1B). In this study, we used heme auxotrophs to stimulate production of GtrR (Fig. 1D), but we would predict that in particular niches during infection, an increase in heme synthesis is required and GtrR abundance increases to accommodate this need. Host-imposed nitrosative stress, oxidative stress, and hypoxia, for example, all cause S. aureus to increase expression of heme-dependent cytochrome oxidases, catalase, and nitric oxide synthase (29–31).

The drastic difference in GtrR enrichment between the heme auxotroph pbgS and ΔchdC mutants (Fig. 1D and 5A), with deficits in genes at the beginning and end of the heme biosynthetic pathway, respectively, suggests that GtrR abundance during heme deficiency could be impacted by mechanisms other than heme availability. This is an interesting observation in light of the comparative genome analysis of hemX, which suggests that HemX could impact siroheme synthesis as well as heme synthesis. It is possible that GtrR abundance is impacted by differences in heme or siroheme abundance as well as abundance in earlier precursor levels, explaining the difference in GtrR levels in the pbgS strain relative to the ΔchdC strain.

In addition to the impact of heme deficiency on GtrR, we also identify HemX as a key regulator of GtrR in heme-proficient cells (Fig. 2A). Our broad genomic analysis has identified HemX homologues across bacterial phyla, suggesting that HemX control of heme synthesis via GtrR is a conserved strategy (Fig. 7; Fig. S8). This finding is consistent with the model set forth by Hederstedt and colleagues (18, 21); in both B. subtilis and now S. aureus, it appears that HemX regulates GtrR abundance posttranscriptionally (Fig. 2B) through an as-yet-undefined mechanism. B. subtilis HemX is sufficient to affect GtrR abundance when both are expressed ectopically in Escherichia coli (18); however, the contribution of heme or conserved E. coli proteins in this system is unclear, making it challenging to conclude if HemX directly interacts with GtrR. Together, our findings are consistent with a model whereby GtrR is regulated by heme abundance and HemX through a multiprotein mechanism. Our finding that GtrR abundance is reduced in the

**FIG 7** Legend (Continued)

rectangles mark the organisms that contain unusual combinations of genes normally involved in different pathways for protoheme synthesis (hybrid paths [14]). The absence of a rectangle in the middle ring indicates the absence of any known route for protoheme synthesis in an organism. Likewise, the absence of a circle (gtrR) or square (alaS) in the outermost ring shows the inability of an organism to produce tetrapyrroles of any kind. Note that hemX does not occur in such organisms. (B) The immediate genomic neighborhood of the hemX gene in seven representative genomes, with ClustalW alignment scores for HemX and GtrR for each organism relative to S. aureus.
\( \Delta \text{hemX} \) \( \text{pbgS} \) strain relative to the \( \text{pbgS} \) strain and not increased supports this model: heme and HemX both do not directly repress GtrR levels (Fig. 5A). In \( \text{Salmonella} \) strains, which do not encode HemX, GtrR is regulated by N-terminal proteolysis by ClpAP and Lon proteases to keep levels low (32). Additionally, \( \text{Salmonella} \) GtrR binds excess heme through a Cys-170 residue (33). Mutagenesis of the N-terminus degradation sequence or heme binding cysteine disrupts regulation, and it has been proposed that heme-bound GtrR but not apo-GtrR is a substrate for the proteases (32, 34). The mechanism by which \( \text{Salmonella} \) GtrR is regulated by proteases via its N terminus and heme binding is not fully understood, and these regulatory amino acids are not conserved in \( \text{S. aureus} \) GtrR. Likewise, further work is needed to dissect the unique regulatory effects of heme and HemX on GtrR levels and the potential involvement of proteolysis in this process in \( \text{S. aureus} \).

In the absence of HemX, \( \text{S. aureus} \) synthesizes excess heme. The increase in heme synthesis disrupts intracellular iron homeostasis (Fig. 4), which could additionally disrupt the expression of the staphyloferrin B siderophore biosynthetic genes, which were recently identified to be under the control of a transcriptional activator that is inactive when bound to heme (35). This altered iron homeostasis would likely reduce the function of many important Fe-S cluster-containing enzymes critical to staphylococcal physiology. Additionally, excess heme synthesized in the \( \Delta \text{hemX} \) mutant activates the heme stress response (Fig. 3). While activation of the HssRS two-component system was first recognized as the result of exogenous heme, our findings add to a growing body of literature that supports a model whereby endogenous heme and exogenous heme both contribute to HssRS activation and heme toxicity. We have previously identified small molecule activators of heme synthesis that increase intracellular heme and activate HssRS (36–38), adding to our genetic evidence presented in this work that endogenous heme activates HssRS. Here, the increase in endogenous heme in the \( \Delta \text{hemX} \) mutant is not toxic because of the HssRS-HrtAB detoxification response. Rather, endogenous heme activation of HssRS provides resistance to heme toxicity through preadaptation and expression of hrtAB (Fig. 3C and D). The impact of inactivation of hemX on the fitness of pathogens that do not encode the HssRS-HrtAB system would offer insight into whether dysregulated heme synthesis is sufficient to induce heme toxicity from within.

This study found that regulation of GtrR abundance is sufficient to regulate total heme synthesis (Fig. 2), consistent with multiple reports that ALA formation is a critical rate-limiting step in heme synthesis (14). Indeed, regulation of ALA synthesis via control of either GtrR or ALAS has emerged as a theme across kingdoms. In metazoans, two ALAS isoforms exist and are impacted by heme (reviewed in reference 39). In the plant model organism \( \text{Arabidopsis thaliana} \), ALA synthesis is regulated by degradation of GluTR via Clp proteolysis from the N terminus as well as stabilization and activation of a regulatory binding protein (40, 41). In Gram-negative model organisms, as mentioned above, GtrR abundance is regulated by heme and proteolysis (32). Our findings extend this paradigm further into the \( \text{Firmicutes} \) phylum of bacteria.

The specific mechanism by which HemX impacts GtrR abundance remains undefined. HemX is annotated as a member of the cytochrome c assembly protein family (Pfam accession no. PF01578), suggesting that it may be involved in heme binding and trafficking at the membrane. However, \( \text{S. aureus} \) does not encode c-type cytochromes. The capacity of HemX to bind heme has not been experimentally validated, but excess heme does accumulate in the membrane (42), which is suggestive of a potential role for membrane-localized heme reservoirs or chaperones. The limited regions of HemX predicted to be cytoplasmic suggest that protein-protein interactions likely occur between other membrane proteins, but no HemX-interacting partners have been identified to date. Additionally, GtrR residues that impact regulation by either heme or HemX are unknown, but would offer information as to the regulatory steps between heme, HemX, and GtrR, which appear to be complex. Although heme-dependent inhibition of \( \text{S. aureus} \) heme synthesis was first proposed in 1962 (43), the impact of HemX and heme on GtrR abundance continues to warrant further investigation.
TABLE 1  S. aureus strains used in this study

| Strain          | Genotype | Description                                                                 | Source or reference |
|-----------------|----------|------------------------------------------------------------------------------|---------------------|
| Newman          | WT       | Wild-type, methicillin-sensitive clinical isolate                            | 57                  |
| Newman          | pbgS     | pbgS (NWMN_1562) gene interrupted with erythromycin resistance gene ermB by homologous recombination, transduced into Newman | 3                   |
| Newman          | ΔchdC    | In-frame unmarked deletion of chdC (NWMN_0550) generated by allelic exchange in ΔhemX strain | 58                  |
| Newman          | ΔhemX ΔchdC | In-frame unmarked deletion of chdC (NWMN_0550) generated by allelic exchange | This work          |
| Newman          | ΔmenB    | In-frame unmarked deletion of menB generated by allelic exchange             | 59                  |
| Newman          | ΔgtxB cylD | In-frame unmarked deletion of qoxB and cylD::Tn                            | 3                   |
| Newman          | katA     | katA::Tn (NE1366)                                                          | BEI (46)           |
| Newman          | katA     | katA::Tn (NE1366), transduced into Newman                                    | This work          |
| Newman          | ΔhemX    | In-frame unmarked deletion of hemX (NWMN_1565)                              | This work          |
| NE1279          | ΔgtrR    | gtrR-hemX integrated into chromosome                                          | This work          |
| NE1931          | ΔchdC    | gtrR-hemX integrated into chromosome                                          | This work          |
| NE1366          | ΔmenB    | menB interrupted with hemX                                                   | This work          |
| NE1931          | ΔhemX ΔmenB | In-frame unmarked deletion of menB generated by allelic exchange             | This work          |
| RN9011          |          | RN4220 carrying pRN7023 integrase plasmid                                    | 24                  |
| RN9011          | attC::P<sub>gr</sub>hemX | pJC1111 P<sub>gr</sub>hemX integrated into chromosome at attC locus | This work          |
| Newman          | ΔhemX attC::P<sub>gr</sub> | pJC1111 P<sub>gr</sub> integrated into chromosome at attC locus | This work          |
| Newman          | ΔhemX attC::P<sub>gr</sub>hemX | pJC1111 P<sub>gr</sub>hemX integrated into chromosome at attC locus | This work          |
| Newman          | Δpsg          | pJC1111 P<sub>gr</sub> integrated into chromosome at attC locus             | This work          |
| Newman          | ΔhemX pbgS | In-frame unmarked deletion of hemX (NWMN_1565) in pbgS strain               | This work          |
| Newman          | ΔhssRS     | In-frame unmarked deletion of hssRS generated by allelic exchange           | 60                  |
| Newman          | ΔhrtB      | In-frame unmarked deletion of hrtB generated by allelic exchange            | 61                  |
| Newman          | hrtB       | hrtB::Tn(PhNE01762)                                                        | 10                  |
| Newman          | ΔhemX hrtB | hrtB::Tn(PhNE01762) allele transduced to ΔhemX strain                       | This work          |
| Newman          | ΔgtrR-hemX | In-frame unmarked deletion of gtrR and hemX (NWMN_1565-1566) generated by single allelic exchange | This work          |
| JE2             | cysG      | cysG::Tn (NE1931; SAUSA300_2533::Tn)                                        | BEI (46)           |
| Newman          | cysG      | cysG::Tn (NE1931) transduced into Newman                                     | This work          |
| Newman          | ΔhemX cysG | cysG::Tn (NE1931) transduced into Newman ΔhemX                                | This work          |
| Newman          | nirD      | nirD::Tn (NE1279)                                                          | BEI (46)           |
| Newman          | ΔhemX nirD | nirD::Tn (NE1279) transduced into Newman ΔhemX                               | This work          |
| Newman          | ΔgtrR-hemX | In-frame unmarked deletion of gtrR and hemX (NWMN_1565-1566) generated by single allelic exchange | This work          |
| RN4220          |            | Restriction-deficient cloning intermediate strain                             | 62                  |

MATERIALS AND METHODS

Bacterial strains and reagents. Bacterial strains (Table 1), plasmids (see Table S1 in the supplemental material), and primers (Table S1) are listed in the specified table. S. aureus strains were grown routinely on tryptic soy agar (TSA) or broth (TSB) supplemented with 10 μg/ml chloramphenicol or 10 μg/ml erythromycin when necessary. When used, heme (hemin chloride) was used at the concentrations noted. Heme was prepared fresh at 10 mM in 0.1 M NaOH; for experiments in which heme was used, an equal volume of 0.1 M NaOH was used for all conditions. E. coli strains were grown on lysogeny broth (LB) or LB agar (LBA), supplemented with 50 μg/ml carbenicillin when necessary. For growth in liquid medium, an Innova44 incubator shaking at 180 rpm was used. For standard cultures of 4 to 5 ml, 15-ml round-bottomed polypropylene tubes with aeration lids were used, at a 45° angle in the incubator. For cloning and mutagenesis in plasmids, all constructs were confirmed by sequencing (GeneWiz). Unless noted otherwise, all chemicals are from Sigma. All molecular biology reagents were from New England Biolabs (NEB) and used according to the manufacturer's instructions, unless otherwise noted. Phusion 2X Hi-fidelity master mix was used for all PCRs for cloning. As necessary, plasmids were transformed by electroporation from E. coli into the S. aureus cloning intermediate strain RN4220 before isolation and subsequent electroporation into the final S. aureus strains.

(i) Deletion of genes by allelic exchange. Deletion of hemX and chdC was performed by allelic exchange as described in reference 44 with some modifications. The pKOR1 plasmids containing ~1-kb homologous regions flanking upstream and downstream of the gene to be deleted were prepared using NEB Hi-Fi assembly according to manufacturer's suggestions. The pKOR1 backbone was amplified by PCR using JC291/292, which produces a linear product not including the attB recombination sites. The ~1-kb flanking regions were amplified from the S. aureus Newman genomic DNA. Deletions were confirmed by PCR using isolated genomic DNA and complemented by providing the gene in cis or trans. Additional details are found in Text S1 in the supplemental material.

(ii) hemX chromosomal integration. Chromosomal complementation was performed by cloning P<sub>gr</sub> or P<sub>gr</sub>hemX into pJC1111. P<sub>gr</sub> was PCR amplified from pOS1 P<sub>gr</sub> using JC158/229 and subsequently cloned into the multiple cloning site of pJC1111 after restriction digestion with SalI and BamHI. hemX was cloned into pOS1 P<sub>gr</sub> by amplifying hemX flanked by Ndel and BamHI sites from S. aureus Newman genomic DNA using primers JC157/155 and ligated (T4 ligase) into the multiple cloning site of pOS1 P<sub>gr</sub> after restriction digestion with Ndel and BamHI. P<sub>gr</sub>hemX was amplified from pOS1 P<sub>gr</sub>hemX using...
JC158/155 and subsequently cloned into the SalI and BamHI sites of pJC111 after restriction digestion with SalI and BamHI. pJC111 P_hemX and pJC111 P_gyrA were integrated into the chromosome of strain RN9011 as described previously (24) and then transduced into the S. aureus Newman WT or ΔhemX mutant as noted. Transductions of pJC111 loci were performed with φ85 as described in reference 45, with some modifications: after incubation of donor phage with recipient strains and washing with sodium citrate, cells were allowed to recover for 4 h in TSB with 40 mM sodium citrate at 37°C with shaking and plated to TSA containing 0.15 mM cadmium chloride.

(iii) Transduction of transposon library alleles. For transduction of transposon library alleles, the katA::Tn (NE1366), cysG::Tn (NE1931), nirD::Tn (NE1279), and hrtB::Tn (PhiNE01762) transposon alleles were transduced to the S. aureus Newman and ΔhemX strains as listed in Table 1 as described previously (45) using bacteriophage φ85; alleles were confirmed by an inverse-PCR method and Sanger sequencing (46).

Catalase activity. To assess catalase activity, strains were grown for 16 h in TSB, and then 50 μl of each culture was spotted onto a TSA plate and streaked for isolation. After 24 h of growth at 37°C, 50 μl of 30% H2O2, was added to each strain and immediately imaged.

LC-MRM-MS/MS. Strains were streaked onto TSA and grown for 24 h at 37°C. Cultures were started from single colonies in 5 ml of RPMI plus 1% Casamino Acids and grown at 37°C for 15 h. Overnight cultures were subcultured 1:100 into RPMI plus 1% Casamino Acids and grown until mid-exponential phase. For small-colony variants without chemical complementation, overnight cultures were subcultured 1:25. For conditions under which heme was added, 2 μM was used; for menaquinone, 12.5 μM menaquinone-vitamin K2 was used.

For anaerobic experiments, a Coy (Grass Lake, MI) anaerobic chamber was used, filled with a mixture of 90% nitrogen, 5% carbon dioxide, and 5% hydrogen gases, and hydrogen levels were monitored to ensure a minimum 2% hydrogen concentration. Palladium catalysts (Coy) were used to remove any residual oxygen. A Coy static incubator was maintained at 37°C. Solutions and plasticware were allowed to equilibrate for >24 h inside the glove box before use. For anaerobic samples, strains were streaked onto TSA and grown aerobically for 24 h at 37°C. Cultures were started from single colonies in 5 ml of anaerobic TSB and grown at 37°C for 15 h. Overnight cultures were subcultured 1:100 into anaerobic TSB containing 40 mM sodium nitrate and grown until mid-exponential phase. Protein was collected, tryptically digested, and subjected to LC-MRM-MS/MS as described in Text S1 in the supplemental material.

Anaerobic growth curves. The S. aureus Newman WT, ΔhemX, cysG, ΔhemX cysG, nirD, and ΔhemX nirD strains were streaked onto TSA and grown aerobically for 24 h at 37°C. Cultures were started from single colonies in 5 ml of anaerobic TSB and grown at 37°C for 15 h. Overnight cultures were subcultured 1:100 in round-bottomed 96-well plates with 200 μl of anaerobic TSB containing 40 mM sodium nitrate or an equal volume of sterile water and covered with Breathe-Easy gas-permeable seal (Sigma). Growth was monitored by optical density (OD) over time in a BioTek Synergy H1.

Quantitative reverse transcriptase PCR. Strains were streaked onto TSA and grown at 24 h at 37°C. Cultures were started from single colonies in 5 ml of RPMI plus 1% Casamino Acids and grown at 37°C for 15 h. Overnight cultures were subcultured 1:100 (WT and ΔhemX mutant) or 1:25 (pbgS ΔhemX pbgS strain) into RPMI plus 1% Casamino acids and grown until the mid-exponential phase. An equal volume of ice-cold acetone-ethanol was added, and the mixture was stored at −80°C. RNA was isolated using Tri reagent and chloroform and precipitated with isopropanol. Inverted RNAs were treated with DNase I of ice-cold acetone-ethanol was added, and the mixture was stored at

PYRIDIHEMCHROMAGEN QUANTIFICATION. Strains were streaked onto TSA and grown for 18 h at 37°C. Single colonies were used to start 5-ml cultures of TSB and grown at 37°C for 10 h. Sixty microliters of each culture was added to 6 ml of TSB and grown for 16 h at 37°C. Cells were collected by
centrifugation, and the cell wall was removed by incubation in 20 mM potassium phosphate buffer (pH 7.4) containing 20 µg of lysostaphin for 45 min at 37°C. Samples were lysed by sonication, and unbroken cells were collected by centrifugation. Four hundred fifty microliters of the soluble supernatant was added to 450 µl of 0.2 M NaOH containing 40% pyridine and 500 µM potassium ferricyanide. Absorbance was measured in a Cary 50 Bio UV-Vis spectrophotometer from 540 to 590 nm. Ten microliters of 0.5 M sodium dithionite prepared in 0.5 M NaOH was added to samples, the mixture was incubated for 5 min, and absorbance (A) was measured again from 540 to 590 nm. Heme quantity is calculated using the equation: 

\[ \Delta A = (A_{575 \text{ reduced}}) - (A_{575 \text{ oxidized}}) \]

and an extinction coefficient of 32.4 mM\(^{-1}\) cm\(^{-1}\).

**Bioluminescent reporter assay.** The *S. aureus* WT and ΔhemX strain with pXen-1 or pXenLuxABCDE were streaked onto TSA-chloramphenicol prepared with 0 or 20 µM heme. After 18 h, the plates were imaged using a Xenogen IVIS 2000.

**Xyle reporter assay.** XylE abundance in cellular lysate was assessed spectrophotometrically by measuring formation of 2-hydroxymuconic acid from catechol after growth in TSB containing chloramphenicol and 0 to 2 µM heme, as described previously (10).

**Heme killing assay.** *S. aureus* WT, ΔhemX, and ΔhrtB strains were streaked onto TSA and grown for 24 h at 37°C. Single colonies were used to start 5-ml cultures of TSB and grown at 37°C for 14 h. Two microliters of each culture was added to 148 µl of TSB containing different concentrations of heme in a 96-well round-bottomed plate and incubated at 37°C for 2 h. Samples were serially diluted in phosphate-buffered saline (PBS) and plated to TSA for CFU enumeration after 24 h of growth at 37°C.

**Heme toxicity growth curves.** Strains were streaked onto TSA and grown for 24 h at 37°C. Single colonies were used to start 5-ml cultures of TSB and grown for 16 h at 37°C containing 0 or 2 µM heme as noted. One microliter of each culture was used to inoculate 199 µl of medium containing 0 or 10 µM heme, and growth was monitored over time at 37°C by measuring the optical density at 600 nm (OD\(_{600}\)) in a BioTek Synergy2 spectrophotometer and analyzed with BioTek Gen5 software.

**Growth in minimal medium.** Chemically defined media (CDM) supplemented with 5 mg/ml glucose was prepared as previously described (48), with the exception that iron was not added. Strains were streaked onto TSA and grown for 24 h at 37°C. Single colonies were used to start 5-ml cultures in TSB and grown for 14 h at 37°C. Cells were collected by centrifugation, washed in PBS twice, and then resuspended in 5 ml of PBS. One microliter was added to 199 µl of CDM containing 1 µM ethylenediamine-N,N′,N″-tris(2-hydroxyphenylacetic acid) (EDDHA; LGC Standards) or an equal volume of 0.1 M NaOH (vehicle) in a 96-well round-bottomed plate. Growth was monitored for 24 h with shaking at 37°C in a BioTek EPOCH spectrophotometer and analyzed with BioTek Gen5 software.

**pOS1 P\(_{\text{iscAgfp}}\) reporter assay.** The *S. aureus* WT pOS1 P\(_{\text{iscAgfp}}\) and ΔhemX pOS1 P\(_{\text{iscAgfp}}\) strains were streaked onto TSA-chloramphenicol and grown for 14 h at 37°C. Single colonies were used to inoculate 5-ml cultures of TSB-chloramphenicol and grown for 37°C for 8.5 h. One microliter of each culture was used to inoculate 199 µl of TSB-chloramphenicol containing 1 mM 2,2-dipyridyl or an equal volume of ethanol (vehicle). Growth was monitored over the course of 16 h by measuring OD\(_{600}\) as well as relative fluorescence at 485 nm (excitation) and 510 nm (emission) in a BioTek Cytation5 spectrophotometer and analyzed with BioTek Gen5 software.

**Comparative genome analysis.** With over 100,000 prokaryotic genomes currently available in public databases and many more in the pipelines (http://www.genomesonline.org), it is not practical or possible to perform meaningful comparative analysis on all of them simultaneously. Thus, a set of diverse representative prokaryotic genomes have been developed in the SEED database as follows. The algorithm for computing molecular operational taxonomic units (OTU) based on DNA barcode data (49, 50) was used to group ~12,600 prokaryotic genomes available in the SEED database in October 2013 into about 1,000 taxon groups. One or two representative genomes (rarely three) for each OTU were selected based on the largest amount of published experimental data and the highest level of research interest within the scientific community. The resultant collection of 982 diverse genomes (928 eubacterial and 54 archaean) creates a manageable set that accurately represents the immense diversity of the over 12,000 prokaryotic organisms with sequenced genomes. Importantly, it is not skewed by an overabundance of genera for a few microbial genera (medically or industrially important), such as Enterobacteriaceae, streptococci, mycobacteria, etc.

The HemX protein family was exhaustively annotated for this set of 982 representative microbial genomes in the SEED database (51). Contextual associations for this family were predicted based on the patterns of co-occurrence and/or colocalization of its members with other protein families using the set of tools for comparative genome analysis available in SEED (52) within the functional and genomic contexts provided by the subsystem “Heme Biosynthesis: protoporphyrin-, coproporphyrin- and siroheme-dependent pathways” (http://pubseed.theseed.org//SubsysEditor.cgi?page=ShowSubsystem&subsystem=Heme_Biosynthesis%3A_protoporphyrin-%2C_coproporphyrin-%26_siroheme-dependent_pathways). Phylogenetic distribution of the HemX protein family was mapped onto the tree of life (53), and protoheme biosynthetic pathway analysis was adapted from reference 14.

**HemX multiple sequence alignment and topology prediction.** The HemX multiple sequence alignment was KEGG ClustalW (http://www.genome.jp (accessed March 2017)) (54) using Staphylococcus aureus strain Newman, Staphylococcus epidermidis strain ATCC 12228, Bacillus anthracis strain Sterne, Chlorobium tepidum strain TLS, Aquifex aeolicus strain VF5, Desulfovibrio vulgaris strain DP4, and Geobacter sulfurreducentis strain PCA. The transmembrane domains depicted were predicted by MEMSAT3 (http://bioinf.cs.uc.ac.uk; accessed March 2017) as described previously (55). All models were confirmed using TMHMM 2.0 (http://www.cbs.dtu.dk (accessed March 2017)) (56), and predictions were matched...
across prediction servers, with the exception of Chlorobium tepidum, which TMHMM2.0 predicts to have seven rather than eight transmembrane domains.

**Statistical analysis.** All data analysis and statistical tests were performed using GraphPad Prism 6 software. Replicate numbers and statistical tests for each experiment are listed in the figure legends.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02287-17.

**TEXT S1,** DOCX file, 0.1 MB.

**FIG S1,** TIF file, 0.9 MB.

**FIG S2,** TIF file, 0.3 MB.

**FIG S3,** TIF file, 2.7 MB.

**FIG S4,** TIF file, 6.8 MB.

**FIG S5,** TIF file, 0.7 MB.

**FIG S6,** TIF file, 3.1 MB.

**FIG S7,** TIF file, 4.1 MB.

**FIG S8,** TIF file, 45.3 MB.

**TABLE S1,** DOCX file, 0.1 MB.

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