Abstract

**Aims:** *Staphylococcus aureus* is a major human pathogen and has to cope with reactive oxygen and chlorine species (ROS, RCS) during infections, which requires efficient protection mechanisms to avoid destruction. Here, we have investigated the changes in the RNA-seq transcriptome by the strong oxidant sodium hypochlorite (NaOCl) in *S. aureus* USA300 to identify novel redox-sensing mechanisms that provide protection under infection conditions.

**Results:** NaOCl stress caused an oxidative stress response in *S. aureus* as indicated by the induction of the PerR, QsrR, HrcA, and SigmaB regulons in the RNA-seq transcriptome. The hypR-merA (USA300HOU_0588-87) operon was most strongly upregulated under NaOCl stress, which encodes for the Rrf2-family regulator HypR and the pyridine nucleotide disulfide reductase MerA. We have characterized HypR as a novel redox-sensitive repressor that controls MerA expression and directly senses and responds to NaOCl and diamide stress via a thiol-based mechanism in *S. aureus*. Mutational analysis identified Cys33 and the conserved Cys99 as essential for NaOCl sensing, while Cys99 is also important for repressor activity of HypR in vivo. The redox-sensing mechanism of HypR involves Cys33-Cys99 intersubunit disulfide formation by NaOCl stress both in vitro and in vivo. Moreover, the HypR-controlled flavin disulfide reductase MerA was shown to protect *S. aureus* against NaOCl stress and increased survival in J774A.1 macrophage infection assays.

**Conclusion and Innovation:** Here, we identified a new member of the widespread Rrf2 family as redox sensor of NaOCl stress in *S. aureus* that uses a thiol/disulfide switch to regulate defense mechanisms against the oxidative burst under infections in *S. aureus*. *Antioxid. Redox Signal.* 29, 615–636.

**Keywords:** *Staphylococcus aureus*, Rrf2, redox-sensing regulator, hypochlorite stress

Introduction

*Staphylococcus aureus* is not only a common commensal bacterium but also a major pathogen that accounts for 40% of nosocomial infections in humans (31). *S. aureus* can cause local wound infections and also severe life-threatening infections, such as septicemia, endocarditis, necrotizing pneumonia, catheter-associated infections, and osteomyelitis (3, 12, 60). A major problem to combat *S. aureus* infections is the increasing prevalence of multiple antibiotic-resistant isolates, such as methicillin-resistant *S. aureus* (MRSA) strains that arise in hospitals as well as in the community (57). *S. aureus* USA300 is a highly virulent community-acquired MRSA strain that encodes many virulence factors,
Staphylococcus aureus is an important human pathogen that has to cope with the oxidative burst of the host innate immune system under infection conditions to ensure survival. Here, we have characterized the novel NaOCl-sensing transcription factor HypR that belongs to the Rrf2-family and controls the flavin disulfide reductase MerA under NaOCl conditions. HypR senses and responds to NaOCl via the formation of Cys33-Cys99 intersubunit disulfides resulting in dissociation from its operator and derepression of hypR-merA transcription.

The hypR-merA operon was required for growth and survival under NaOCl stress and in macrophage infection assays and was identified as an important defense mechanism under hypochlorite stress.

In this study, we applied RNA-seq transcriptomics to identify new redox-sensing regulators in S. aureus USA300 that could specifically sense and respond to NaOCl stress and thereby regulate target genes that confer protection under infection conditions. The hypR-merA operon was most strongly upregulated under NaOCl in the transcriptome that encodes for the novel redox-sensing regulator HypR and the pyridine nucleotide disulfide reductase mercuric ion reductase (MerA). HypR belongs to the widespread Rrf2 family of transcriptional regulators that include iron/sulfur (FeS)-cluster redox sensors for nitric oxide (NsrR) (23, 90, 98), the iron/sulfur status of the cell (IscR) (82, 83), iron metabolism (RirA) (40), or other signals (67). Other Rrf2 family regulators do not contain FeS clusters, such as the cysteine metabolism repressor (CymR) (28, 47, 84) and the redox-sensing SaiR repressor of Bacillus anthracis that controls spxA under disulfide stress conditions and is a close homologue of HypR (68). Using biochemical and genetic approaches, we have characterized HypR as a novel thiol-based redox sensor that senses HOCl stress by intersubunit disulfide formation and protects S. aureus under macrophage infection conditions.

Results

NaOCl stress causes an oxidative stress response in the transcriptome and upregulates most strongly the hypR-merA operon (USA300HOU_0588-87) in S. aureus USA300

We were interested to identify novel redox-sensing regulators in S. aureus that could be specifically involved in the hypochlorite defense under infection conditions. Thus, we studied the changes in the RNA-seq transcriptome in S. aureus USA300 after exposure to sublethal 150 μM NaOCl stress compared to untreated control cells. The NaOCl stress experiments were performed in Belitsky minimal medium (BMM) to avoid quenching by rich medium components as described in our previous studies (46, 58). Three biological replicate experiments were performed and the samples for RNA isolation were taken before (control) and 30 min after NaOCl stress and subjected to RNA-seq analysis as described (44). For significant expression changes, the log2 fold change (m-value) cutoff (log2-fold change NaOCl vs. control) was 10% increased vs. control (M-value) cutoff (log2-fold change NaOCl vs. control)}
of ±1.98 was chosen (95% confidence, $p \leq 0.05$). In total, 89 transcripts were significantly at least fourfold upregulated and 9 were fourfold downregulated under NaOCl stress in the transcriptome data set (Fig. 1 and Supplementary Table S1; Supplementary Data are available online at www.liebertpub.com/ars). In addition, 1442 genes are differentially transcribed below the m-value cutoff under NaOCl stress in the RNA-seq transcriptome of *S. aureus* USA300. All induced and repressed genes above and below the m-value cutoff were sorted into regulons, and a subset of the most interesting upregulated regulons is displayed in Voronoi treemaps (Fig. 2 and Supplementary Table S2).

Previous transcriptome analysis in *Bacillus subtilis* showed that NaOCl stress induces an oxidative stress response as revealed by the induction of the oxidative stress-specific PerR, OhrR, and Spx regulons, as well as by the CtsR regulon indicative of oxidative protein unfolding (20). Furthermore, NaOCl stress resulted in increased expression of the YodB, CatR, HypR, and HxlR regulons in *B. subtilis* that sense reactive electrophilic species (RES), such as quinones and aldehydes (18, 20, 21, 52, 73). In *S. aureus*, we could confirm a similar expression profile under NaOCl stress as revealed by the strong upregulation of the PerR, HrcA, and QsrR (YodB) regulons indicating ROS and RES responses in *S. aureus*. In addition, we noticed the weak two- to threefold induction of few members of the Fur and Zur regulons. Fur and Zur oxidation in their Zn-redox switch motifs was observed under NaOCl stress in our recent OxICAT analyses in *S. aureus* (46) and *Mycobacterium smegmatis* (44). Similarly, the Fur family protein PerR was oxidized to an intramolecular disulfide in *B. subtilis* under NaOCl stress (20). Since Zn-redox switch motifs are known sensors for HOCl stress also in *E. coli* (45, 95), redox regulation of PerR, Fur, and Zur could involve also thiol oxidation in their Zn-motifs.

The SigmaB regulon was also induced under NaOCl stress in *S. aureus* and *B. subtilis*, which might be indicative for an acid stress response and contributes to virulence (20) (Fig. 2). The induction of the SigmaB regulon was previously shown after internalization of *S. aureus* in human epithelial cells and SigmaB was essential for intracellular growth (77). Of note is further induction of GraRS cell wall stress regulon under NaOCl stress in *S. aureus*. GraRS controls resistance to antimicrobial peptides and cell wall-active antibiotics, such as vancomycin, but contributes also to the oxidative stress resistance of *S. aureus* explaining its upregulation under NaOCl stress (29). The oxidative stress response is further indicated by induction of genes that encode for BSH biosynthesis enzymes and bacilliredoxins, such as bshA, bshB, bshC, and brxB in *S. aureus*. The BSH biosynthesis genes are under control of Spx in *B. subtilis* (33) and perhaps also in *S. aureus*.

Apart from the stress regulons, the SaeRS regulon was most strongly induced under NaOCl stress in *S. aureus*. The SaeRS two-component system controls many important toxins, such as α-, β-, and γ-hemolysins, leukocidins, superantigens, surface proteins, proteases, nuclease, coagulase, and immune evasion proteins (7, 38, 56) that exhibit the highest fold change between 20- and 160-fold under NaOCl stress (Supplementary Tables S1 and S2, Figs. 1 and 2). Previous studies established important roles of SaeRS in pathogenicity and biofilm formation in a variety of disease models of infections (56). The SaeRS regulon was most strongly upregulated by neutrophil stress in *S. aureus* as shown in previous microarrays (91). The

**FIG. 1.** RNA-seq transcriptomics of *Staphylococcus aureus* USA300 after 30 min of NaOCl stress. For RNA-seq transcriptome profiling, *S. aureus* USA300 was grown in BMM and treated with 150 μM NaOCl stress for 30 min. The gene expression profile under NaOCl stress is shown as ratio/intensity scatter plot (M/A-plot), which is based on the differential gene expression analysis using DeSeq2. Colored symbols indicate significantly induced (red, magenta, yellow) or repressed (green) transcripts (M-value $\geq 1.98$ or $\leq -1.98$; $p$-value $\leq 0.05$). Black symbols indicate differential transcribed genes below the M-value cutoff of 1.98 or $M$ $> -1.98$ ($p \leq 0.05$). Gray symbols denote transcripts with no fold changes after NaOCl stress ($p > 0.05$). The SaeRS, HypR, MgrA, ArgR, CodY, and QsrR regulons are most strongly upregulated under NaOCl stress. The transcriptome analysis was performed from three biological replicates. The RNA-seq expression data of all genes after NaOCl stress and their regulon classifications are listed in Supplementary Tables S1 and S2. BMM, Belitsky minimal medium; NaOCl, sodium hypochlorite.
SaeRS-controlled virulence factors are essential for innate immune evasion since the survival of the saeRS mutant was impaired in neutrophil phagocytosis assays (92). Reduction of ROS production by SaeRS-regulated virulence factors inside neutrophils was recently shown as mechanism to avoid pathogen killing (37). Thus, the strong induction of the SaeRS virulence regulon under NaOCl stress in our transcriptome is reflecting adaptation to infection conditions, such as neutrophil stress.

Among the genes involved in energy metabolism, the expression of the glycolytic cggR-gap-pgk-tpiA-pgm-eno-operon and the pyruvate dehydrogenase pdhABCD operon was two- to
fourfold elevated. Since the glycolytic Gap is the main thiol switch that is redox controlled and inhibited under NaOCl stress by S-bacillithiolation (46), the lack of energy production due to inhibited glycolysis could result in upregulation of the glycolysis enzymes. Furthermore, the two- to threefold induction of the ATP synthase atpCDGHFE operon under NaOCl stress may indicate ATP depletion. Of note is the further enhanced expression of the CodY regulon under NaOCl that includes the ilv-leu and lysC-asd-dapABD operons involved in leucine, isoleucine, and lysine biosynthesis, as well as the riboflavin biosynthesis ribHBAED operon. Increased demand of riboflavin might be caused by the requirement of flavin cofactors for thiol/disulfide reductases. Moreover, the ArgR regulon genes for biosynthesis of arginine and the ArcR-controlled arginine deiminase operon are strongly induced under NaOCl, which could indicate increased ammonium production to compensate for an acid stress response caused by NaOCl stress. Most interesting was the 180-fold induction of the USA300HOU_0588-0587 operon under NaOCl treatment (Figs. 2 and 3). USA300HOU_0588 encodes for an uncharacterized Rrf2 family regulator, termed as HypR, and USA300HOU_0587 encodes for the flavin disulfide reductase (FDR) MerA. The strong expression of the hypR-merA operon was previously observed in S. aureus isolates during phagocytosis with polymorphonuclear leukocytes (PMNs or neutrophils) (91). Thus, our expression profile under NaOCl stress highly resembles that of infection conditions due to the production of ROS and RCS in activated neutrophils. Accordingly, HypR could specify a novel NaOCl-specific transcription factor and defense mechanism of S. aureus to avoid destruction by neutrophils. Thus, we studied the redox-sensing regulatory mechanism of HypR and the function of MerA in more detail under NaOCl stress conditions. For the detailed HypR functional analysis, we have chosen S. aureus COL, which is our model strain for functional genomics, and protocols for genetic manipulation are best established in our laboratory for COL, which has less resistance determinants compared to USA300.

Northern blot analysis revealed the specific induction of the hypR-merA operon under NaOCl and diamide stress and its negative regulation by the Rrf2 family repressor HypR.

We used Northern blot analysis to study the expression of the hypR-merA operon under different thiol-specific stress conditions in S. aureus COL, including 1 mM NaOCl, 2 mM diamide, 10 mM H₂O₂, 0.75 mM formaldehyde, and 0.5 mM methylglyoxal (Fig. 4A). These experiments were performed in RPMI medium, which resembles infection conditions and allows faster growth of S. aureus, but requires higher sublethal doses of 1 mM NaOCl. The strong upregulation of the hypR-merA-specific 1.8 kb mRNA under 1 mM NaOCl and 2 mM diamide stress was confirmed, but no significant increased transcription was found with 10 mM H₂O₂ and aldehyde stress. This indicates that the hypR-merA operon responds specifically to disulfide stress, but not to H₂O₂ and aldehydes in S. aureus.

Next, we studied the role of HypR in transcriptional regulation of the hypR-merA operon in S. aureus COL. Using the temperature-sensitive plasmid pMAD, a clean hypR deletion mutant was constructed lacking 364 bp of the hypR open reading frame. The truncated hypR-merA transcript of 1.5 kb (denoted as TR-merA) was constitutively expressed in the hypR deletion mutant under nonstress conditions, indicating full derepression of hypR-merA transcription (Fig. 4B). These results identify HypR as transcriptional repressor of the hypR-merA operon that is inactivated under disulfide stress conditions in S. aureus.

FIG. 3. The hypR-merA-operon is most strongly upregulated in the RNA-seq transcriptome of S. aureus USA300 under NaOCl stress. The mapped reads for the gene expression profile of the hypR-merA locus under control and NaOCl stress are shown as displayed using the Read-Explorer software. Transcription of the hypR-merA-operon is 180-fold induced under NaOCl stress in S. aureus USA300. The hypR gene encodes for an Rrf2 transcriptional regulator and merA encodes for a pyridine nucleotide disulfide reductase.
Cys99 is required for HypR repressor activity and Cys33 and Cys99 are the NaOCl-sensing Cys residues in vivo.

The HypR repressor belongs to the Rrf2 family that includes FeS cluster containing redox sensors for nitric oxide (NsrR) (23, 90, 98), and also SaiR of B. anthracis as the closest HypR homologue that does not contain an FeS cluster (68) (Fig. 5A). Three conserved Cys residues (Cys93, Cys99, and Cys105) and Asp8 are essential for FeS cluster coordination in NsrR (90), but only Cys99 is conserved in HypR and SaiR. Instead, HypR has two other Cys residues at the nonconserved positions 33 and 142. The crystal structure of the HypR and SaiR homologue YwnA of B. subtilis is available (PDB code 1xd7), which showed 23.5% sequence identity to HypR and shares also the conserved Cys97 (Fig. 5B). Based on the structure of YwnA as template, the HypR and SaiR structures were modeled with SWISS-MODEL (10). The HypR structural model indicates that Cys33 is located in the turn between the z2 and z3 DNA recognition helices, while Cys99 is at the N-terminus of the long z6 helix that forms the dimer interface (Fig. 5B). The position of Cys142 could not be modeled since the C-terminal z8 helix is not present in the YwnA structure that was used as template.

To study the role of the three Cys residues for HypR repressor activity and redox sensing, we analyzed transcription of the hypR-merA operon in hypRC33A, hypRC99A, and hypRC142A mutants in vivo. The successful complementation of the hypR mutant using the pRB473-based expression of hypR, hypRC33A, hypRC99A, and hypRC142A under control, NaOCl and diamide stress was confirmed using Northern blots with a hypR-specific mRNA probe (Supplementary Fig. S1A, B). Additional Western blot analysis confirmed that the complemented strains expressed HypR, Cys33A, Cys99A, and Cys142A mutant proteins at the same protein levels (Supplementary Fig. S6A, B). Transcriptional analysis of hypR-merA expression revealed that the hypRC33A mutation did not affect the DNA binding activity of HypR, but impaired its redox sensing since transcription was abolished under NaOCl and diamide stress (Fig. 4C, D). In contrast, the hypRC99A mutant was impaired in DNA binding as shown by full derepression of the hypR-merA operon under control conditions. The hypRC142A mutant behaved similar as the wild type and was not impaired in DNA binding and redox sensing under NaOCl and diamide stress. In conclusion, our Northern blot results revealed that Cys33 is important for redox sensing of HypR in vivo. For stress experiments, S. aureus cells were grown in RPMI medium and treated with thiol-reactive compounds at an OD500 of 0.5. The arrows point toward the hypR-merA bicistronic mRNA (1.8 kb) in the wild type or the truncated hypR-merA transcript (TR-merA) (1.5 kb) in the hypR mutant. The methylene blue stain is the RNA loading control indicating the 16S and 23S rRNAs. OD500, optical density at 500 nms. H2O2, hydrogen peroxide.

FIG. 4. Northern blot analysis of hypR-merA transcription in S. aureus COL under NaOCl, diamide, H2O2, and aldehyde stress and in hypR Cys-Ala mutants. (A) Northern blot analysis was performed using RNA isolated from S. aureus COL wild type before (co) and 15 and 30 min after exposure to 1 mM NaOCl, 2 mM diamide, 10 mM H2O2, 0.75 mM formaldehyde, and 0.5 mM methylglyoxal stress. (B) Transcription of the hypR-merA operon was analyzed in the COL wild-type and in the ΔhypR mutant under 1 mM NaOCl stress indicating strong derepression of hypR-merA transcription under control conditions in the absence of HypR. (C, D) Northern blot analysis of hypR-merA operon transcription in the ΔhypR deletion mutant and in ΔhypR mutants complemented with hypR, hypRC33A, hypRC99A, and hypRC142A before and 15 and 30 min after exposure to 1 mM NaOCl (C) or 2 mM diamide stress (D). The results indicate that Cys33 is required for redox sensing of HypR in vivo. For stress experiments, S. aureus cells were grown in RPMI medium and treated with thiol-reactive compounds at an OD500 of 0.5. The arrows point toward the hypR-merA bicistronic mRNA (1.8 kb) in the wild type or the truncated hypR-merA transcript (TR-merA) (1.5 kb) in the hypR mutant. The methylene blue stain is the RNA loading control indicating the 16S and 23S rRNAs. OD500, optical density at 500 nms. H2O2, hydrogen peroxide.
HypR binds specifically to the **hypR-merA** operator and DNA binding is reversibly inhibited under NaOCl and diamide stress in vitro

The results above suggested that HypR binds directly to the **hypR-merA** promoter region and that Cys33 is the redox-sensing Cys residue that is oxidized under NaOCl and diamide stress. Rrf2 regulators are known to bind as homodimers to operator sequences with dyad symmetry (84). Based on the RNA-seq data, we used the MEME software to identify the putative promoter sequence and a conserved inverted repeat in the upstream region of the **hypR-merA** operon (Fig. 6A). The 12-3-12 bp inverted repeat sequence TAATTGTAACTA-N3-CAGTTACAATTA was detected in the **hypR-merA** upstream region as possible HypR binding site that overlapped with the putative -10 region. Similar inverted repeats are characteristic as full-length binding sites for other Rrf2 regulators, such as SaiR, CymR, IscR, and RsrR (67, 68, 82, 84). We searched for the conservation of the putative HypR operator sequence upstream of homologous **hypR-merA** operons in the genomes of other **Staphylococcus** species. The multiple-sequence alignment showed that the 12-3-12 bp inverted repeat sequence is highly conserved in the **hypR-merA** upstream promoter regions across **Staphylococci** (Fig. 6B).

To investigate whether HypR binds directly and specifically to the HypR operator sequence under reduced conditions with dithiothreitol (DTT), we used electrophoretic mobility shift assays (EMSAs). The gel shift results revealed that purified reduced HypR protein binds with high affinity to the **hypR-merA** upstream region in vitro (Fig. 7A). To analyze the specific binding of HypR to the 12-3-12 inverted repeat sequence, we exchanged two nucleotides in each half of the inverted repeat by G-T and A-C (IR-m1 probe) or T-G and C-A (IR-m2 probe) and analyzed the DNA-binding activity of HypR to these mutated DNA probes (Supplementary Fig. S2). HypR was unable to bind to the mutated inverted repeats IR-m1 and IR-m2 in vitro supporting the specific binding of HypR to the identified operator sequence. Furthermore, HypR was also

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unable to bind to nonspecific DNA probes (trxA and RNAIII genes) (Supplementary Fig. S2B).

Next, the effects of Cys-Ala mutations in Cys33, Cys99, and Cys142 on the DNA-binding activity of HypR were investigated using gel shift experiments. The reduced HypR-C33A and HypRC142A mutant proteins showed similar DNA-binding affinities compared with the HypR wild-type protein (Fig. 7A). However, the reduced HypRC99A and HypRC99S mutant proteins were unable to bind to the hypR-merA promoter, which confirms our in vivo results. Next, we analyzed the effect of thiol oxidation of the HypR, HypRC33A, and HypRC142A proteins on DNA-binding activity in vitro. Treatment of HypR, HypRC33A, and HypRC142A mutant proteins with 1–20 μM NaOCl or 10–20 μM diamide resulted in their fast dissociation from the hypR operator DNA (Fig. 7B, C). This indicates that HypR as well as the HypRC33A and HypRC142A mutant proteins respond similar to NaOCl and diamide stress in vitro. The reduction of the oxidized HypR proteins with DTT restored the DNA-binding ability indicating that HypR responds to NaOCl stress by reversible thiol oxidation. The responsiveness of the HypRC33A mutant to NaOCl stress in vitro is in contrast to lack of NaOCl response and hypR-merA transcription in the hypR-C33A mutant in vivo. Cys33 was identified as redox-sensing Cys in vivo, but the C33A mutant was not impaired in redox sensing under NaOCl and diamide in vitro. It might be possible that the purified C33A mutant protein is oxidized to the “wrong” disulfides under disulfide stress in vitro, which are not stabilized inside S. aureus. In summary, our results show that HypR is a redox-sensing regulator that controls expression of the hypR-merA operon and is inactivated due to reversible thiol oxidation under NaOCl and diamide stress leading to release from the promoter DNA.

HypR responds to NaOCl and diamide stress by intermolecular disulfide formation that involves the redox-sensing Cys33 and Cys99

The EMSA results and Northern blots revealed that the DNA-binding activity of HypR is redox regulated by reversible thiol oxidation under NaOCl and diamide stress. Thus, we were interested to elucidate the thiol switch mechanism leading to release of HypR from the target DNA. HypR can be classified as 2-Cys-type redox regulator, such as OhrR from Xanthomonas campestris and YodB from B. subtilis that sense different thiol reactive compounds by intermolecular disulfide formation between the opposing subunits in the homodimer (2, 27, 43, 74). Thus, HypR most likely also senses and responds...
to NaOCl via intersubunit disulfide bonds. Indeed, when we exposed purified HypR to NaOCl or diamide stress, it was quickly oxidized to the disulfide-linked dimer that migrates at the size of 35 kDa in the nonreducing SDS-PAGE (Fig. 8A and Supplementary Fig. S4A). The oxidation of HypR to the intersubunit disulfide was reversible with DTT as shown in the reducing SDS-PAGE (Supplementary Figs. S3A and S4C). Of note, two bands migrated closely together at the size of the HypR disulfide-linked dimer, labeled as 1 and 2 (Fig. 8A). We hypothesized that these two dimers harbor either one or two disulfide bonds between the opposing HypR subunits. The reversible oxidation to the intermolecular disulfide was also observed for the HypRC33A and HypRC142A mutant proteins under NaOCl and diamide stress, but not for the HypRC99A mutant (Fig. 8A, B, Supplementary Figs. S3A and S4C). These results are in agreement with the in vitro DNA binding assays where the HypRC33A and HypRC142A mutant proteins still responded to NaOCl and diamide stress leading to the relief of repression, while the HypRC99A mutant was impaired in DNA binding. However, the Cys33A mutant was less oxidized to intermolecular disulfides under diamide stress compared to NaOCl, supporting that Cys33 is required for disulfide formation. In conclusion, our results show that HypR is oxidized to intermolecular disulfides under NaOCl and diamide stress, most likely involving Cys33 and Cys99 as the redox-sensing Cys residues in vitro.

We were interested to identify the Cys residues oxidized to the intermolecular disulfide in wild-type HypR under disulfide stress. The bands 1 and 2 of both oxidized HypR dimers (Fig. 8A) were tryptic digested and subjected to matrix-assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF-MS) (Fig. 9). In the overview MS1 scan, a peptide was identified with the size of 3522.74 Da corresponding to the size of the Cys33-SS-Cys99 intermolecular disulfide peptide (Fig. 9A). During fragmentation of this Cys33-SS-Cys99 peptide, the parent ion immediately disappeared and the disulfide bond fragmented into both single Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B). Moreover, fragment ions of both Cys peptides were observed without the sulfur and with the second sulfur atom that was engaged in the disulfide linkage in the parent ion. Thus, the fragmentation spectrum of this 3522.74 Da peptide is characteristic for the disulfide bond between the Cys33 and Cys99 peptides with the sizes of 1534.72 and 1992.90 Da (Fig. 9B).
In addition, we aimed to identify the disulfide crosslinked Cys residues in the Cys33A mutant that formed under NaOCl stress in vitro (Fig. 8A). MALDI-TOF-MS analysis revealed that Cys99 is fully reduced in the oxidized Cys33A mutant protein as shown by the iodoacetamide (IAM)-alkylated Cys99 peptide that was not detected in the oxidized HypR protein (Supplementary Fig. S5C, D). However, Cys142 could not be detected in the Cys33A mutant protein sample. These results suggest that HypR could be oxidized to the wrong Cys142-Cys142 disulfide that may crosslink two HypR dimers in vitro. The large peptide size of this putative Cys142-Cys142 disulfide peptide excludes detection by MS.

Next, we used nonreducing Western blot analysis with polyclonal HypR antibodies to confirm the intermolecular disulfides between Cys33 and Cys99 in HypR under NaOCl stress in vivo. Protein extracts were prepared from the S. aureus COL wild type, the ΔhypR mutant, and the hypR, hypRC33A, hypRC99A, and hypRC142A complemented strains under control and NaOCl stress, and the redox state of HypR was analyzed using nonreducing Western blots (Fig. 8C, D). Unfortunately, the HypR polyclonal antibodies showed cross-reactivity with other proteins of S. aureus which could not be eliminated after preincubation of the antiserum with Western blots of the ΔhypR mutant. However, the HypR specific bands could be clearly distinguished by using protein extracts from hypR immunoprecipitates shown in Supplementary Figure S7. MALDI-TOF-MS, matrix-assisted laser desorption ionization-time of flight mass spectrometry.

FIG. 8. HypR senses NaOCl stress by intermolecular disulfides, which requires Cys33 and Cys99 in vitro and in vivo. (A, B) The purified HypR wild-type and Cys mutant proteins were treated with increasing NaOCl concentrations in vitro and subjected to nonreducing SDS-PAGE analysis. The reduction of the HypR disulfides after DTT treatment is shown in the reducing SDS-PAGE analysis in Supplementary Figure S3. The HypR intermolecular disulfides (bands 1 and 2) were cut, tryptic digested, and subjected to MALDI-TOF MS/MS analysis as shown in Figure 9 to verify the Cys33-Cys99 disulfide. The bands of the C33A mutant protein that were used for tryptic digestion and MS are boxed and labeled with 3 and 4. The MALDI-TOF results of the C33A mutant tryptic peptides are shown in Supplementary Figure S5B, C. (C, D) For the analysis of HypR disulfides in vivo, we used S. aureus COL with plasmid pRB473-hypR, the ΔhypR deletion mutant and ΔhypR mutant strains complemented with hypR, hypRC33A, hypRC99A, and hypRC142A. S. aureus strains were exposed to NaOCl stress, alkylated with NEM, and protein extracts were subjected to nonreducing Western blot analysis using polyclonal rabbit anti-HypR antibodies. The reducing Western blot analysis of the HypR disulfides and loading controls is shown in Supplementary Figure S6. (E) Nonreducing/reducing diagonal SDS-PAGE and HypR-specific Western blot analysis of alkylated protein extracts were performed to verify the intersubunit disulfides for HypR and the HypR-C142A mutant protein under NaOCl stress in vivo, but not in the HypRC33A and HypRC99A mutants. Additional diagonal assays using the HypR immunoprecipitates are shown in Supplementary Figure S7.
Fig. S6A, B). However, both Cys33A and Cys99A mutants did not form intermolecular disulfides under NaNCl exposure and failed to respond to NaNCl stress in vivo (Fig. 8D). These results are in agreement with the transcriptional data of hypR-merA expression and indicate that HypR responds to NaNCl stress by the formation of intersubunit disulfides between Cys33 and Cys99 in vivo and in vitro. To confirm the formation of the intermolecular disulfide in HypR and the involvement of Cys33 and Cys99 in the disulfide crosslink in vivo, diagonal nonreducing/reducing SDS-PAGE analyses were performed using the crude protein extracts and immunoprecipitated HypR. The diagonal assays clearly showed that HypR and the HypRC142A mutant protein migrate at the right side of the diagonal in NaNCl-treated cells indicating intermolecular disulfide formation in vivo (Fig. 8E). In contrast, no HypR intermolecular disulfide was detected in the HypRC33A and HypRC99A mutant proteins in the diagonal assays (Fig. 8E and Supplementary Fig. S7). These data reveal that Cys33 and Cys99 are the redox-sensing Cys residues of HypR, required for disulfide bond formation both in vitro and in vivo.

Analysis of the oligomerization states by size exclusion multiangle light scattering and circular dichroism spectroscopy of HypR and HypR Cys mutant proteins

Cys99 is located at the end of the long α6 helix that forms the dimer interface in the HypR structure (Fig. 5B). The differences in the DNA-binding properties between HypR and the HypRC99A mutant protein raised the question whether the Cys99A mutation affects dimerization of HypR. Thus, we analyzed the oligomerization states of reduced HypR and HypR Cys-Ala mutant proteins using size exclusion chromatography/multiangle light scattering (SEC-MALS) (Supplementary Fig. S8). The results showed that all reduced proteins eluted mostly as dimers with a molecular weight of ~32–34 kDa. Due to the redox sensitivity of Cys33 and Cys99, we observed higher molecular weight oligomeric species with molecular masses ranging from 67 to 78 kDa. The oligomer amounts differed between each Cys mutant (15–17%) and might be related to disulfide crosslinks of two HypR dimers due to incomplete reduction of the proteins. Overall, the SEC-MALS results indicate that HypR and the three Cys-Ala mutant proteins are mostly present as dimers in solution.

Next, we used circular dichroism (CD) spectroscopy to compare the secondary structures between reduced HypR and the three Cys-Ala mutants to see whether the Cys99A mutation causes a conformational change. The far UV-CD spectra of HypR and the three Cys-Ala mutants revealed a significant α-helical content in all proteins (Supplementary Fig. S9). The calculation of secondary structure elements using the program DichroWeb (http://dichroweb.cryst.bbk.ac.uk) (86, 94) showed that reduced HypR and all three Cys-Ala mutants contain similar contents of α-helices (34–37%), β-sheets (11–13%), and β-turns (18–19%), confirming our
structural model (Fig. 5B). Thus, under reduced conditions, the dimerization and secondary structure elements of HypR are retained in all three HypR Cys-Ala mutant proteins.

HypR and MerA play important roles in the defense of S. aureus against NaOCl stress

Next, we analyzed the role of MerA in protection under NaOCl stress in S. aureus COL. MerA is annotated as nicotinamide adenine dinucleotide phosphate (NADPH)-dependent FDR, which contains an active site Cys43 in a highly conserved C_43XXXXC_48 motif (Supplementary Fig. S10) (4). The active site Cys43 of MerA was also predicted in the THIOREDOXOME database (30). We hypothesized that MerA could be involved in reduction of cellular disulfides formed under oxidative stress and infection conditions. Thus, we analyzed the growth and survival of the S. aureus merA and hypR mutants and the complemented strains under NaOCl stress. The growth of the merA mutant was significantly impaired under sublethal 1.5 mM NaOCl treatment in RPMI medium indicating that the merA mutant is more sensitive compared with the wild type (Fig. 10A). The NaOCl-sensitive growth defect of the merA mutant could be restored back to wild-type level after complementation with merA expressed from plasmid pRB473 (Fig. 10B), but not when the active site Cys43 of MerA was replaced by a serine. This indicates that MerA confers resistance to NaOCl and likely functions as defense mechanism under infection conditions in S. aureus. However, the hypR mutant was also slightly more sensitive to 1.75 mM NaOCl stress compared with the wild type suggesting that constitutive MerA expression does not confer a resistance phenotype (Fig. 10C, D).

We further studied the survival of the merA and hypR mutants and their complemented strains under NaOCl stress. Both merA and hypR mutants were impaired in their survival after exposure to lethal concentrations of 3.5 mM NaOCl (Fig. 11). The survival defect of the merA and hypR mutants could be restored back to wild-type level after complementation with merA and hypR, respectively. In agreement with the growth curves, Cys43 of MerA was important for the function of MerA in the NaOCl stress defense in the survival assay. Moreover, the role of Cys33 for redox sensing of HypR was confirmed also in our survival phenotype assays.

MerA and HypR are important for the survival of S. aureus in macrophage infection assays

Since the hypR-merA operon was most strongly induced in infection assays (91), we determined the survival of merA and hypR mutants inside murine macrophages in phagocytosis assays using the cell line J-774A.1. The macrophages were infected with S. aureus wild-type and mutant strains and the uptake of bacteria was stopped after 1 h with gentamycin to kill extracellular bacteria. The colony forming units (CFUs) of intracellular S. aureus were determined after 2, 4, and 24 h postinfection by plating serial dilutions of the host cell lysates. After 4 h of infection, the CFUs of intracellular S. aureus cells were determined as approximately 10^6 cells for the wild-type strain (Fig. 11A). The results are from four biological replicate experiments. MerA, mercuric ion reductase; OD_{500}, optical density at 500 nm.
type, hypR, and merA mutants. However, 24 h after phagocytosis, the viable counts of intracellular S. aureus cells were decreased to 22.8% for the wild-type mutant and to 11.2% and 13.4% for the hypR and merA mutants, respectively (Fig. 12A). This indicates that merA and hypR mutants showed 40% and 50% decreased survival inside macrophages compared with the wild type after 24 h postinfection (Fig. 12C). This survival phenotype could be restored to 75–86% of wild-type level in both mutants after complementation with plasmid-encoded hypR and merA, respectively (Fig. 12B). These observations strongly indicate that MerA is required for the protection of S. aureus against the host innate immune defense under infection conditions. However, our results also show that constitutive expression of MerA in the hypR mutant does not increase intramacrophage survival. In conclusion, our in vitro growth and survival assays under NaOCl stress and the macrophage infection assays support that MerA is a major NaOCl defense mechanism enhancing survival of S. aureus in infection assays.

Discussion

During infections, S. aureus has to cope with the oxidative burst of activated macrophages and neutrophils, requiring a complex regulatory network of virulence and antibiotic regulators, for example, of the SarA/MarR family that senses ROS, RES, and RCS (8, 17, 43). However, the redox-sensing mechanisms of S. aureus in response to strong bactericidal oxidants, such as hypochloric acid, are largely unknown. It is of utmost importance to understand the mechanisms of intracellular survival of S. aureus inside macrophages and neutrophils and the escape from destruction by oxidants to discover new drug targets to combat emerging drug-resistant S. aureus infections.

In this work, we have characterized the novel NaOCl-sensing transcription regulator HypR that belongs to the widely distributed Rrf2 family and controls the NADPH-dependent FDR MerA in S. aureus. The hypR-merA operon was previously highly induced during phagocytosis with neutrophils (91). In this work, we have used a global RNA-seq analysis and identified the hypR-merA operon as most strongly upregulated under NaOCl stress. Detailed Northern blot analysis revealed that NaOCl and diamide stress specifically upregulates the hypR-merA operon, whereas sublethal concentrations of H$_2$O$_2$ and aldehydes do not lead to significant induction. Thus, HypR responds strongly to disulfide stress, but not to H$_2$O$_2$ and RES via intermolecular disulfide formation between Cys33 and Cys99. The sensitivity of S. aureus HypR to disulfide stress resembles the redox-sensing mechanisms of the MarR family regulators HypR and OhrR characterized in B. subtilis (20, 32, 73) and of the TetR family regulator NemR from Escherichia coli (35, 36). These thiol-based regulators sense HOCI, diamide, organic hydroperoxides, or electrophiles, such as N-ethylmaleimide (NEM) and methylglyoxal, and provide protection under HOCI stress. Since diamide is a very unspecific oxidant (49) that is not physiological relevant for S. aureus, we regard the Rrf2 regulator HypR as the most specific hypochlorite defense mechanism for S. aureus.

Some Rrf2 family transcriptional regulators contain FeS clusters, such as NsrR (23, 90, 98) and IcsR (82, 83), while others do not, such as CymR (28, 47, 84) and the HypR homolog SaiR of B. anthracis (68). NsrR and IcsR coordinate [4Fe-4S] clusters with three conserved Cys residues (Cys93, Cys99, and Cys106 in Sc-NsrR) present in the C-terminal domain (Fig. 5A, B) (90). However, only Cys99 in this C-terminal domain is conserved in HypR, while two other Cys residues are located in nonconserved positions at Cys33 and Cys142 (Fig. 5A). Our results support a model that HypR senses and responds to NaOCl stress by intersubunit disulfide bond formation between Cys33 and Cys99 of the opposing subunits of the HypR dimer in vivo (Fig. 13). HypR oxidation leads to inactivation of its repressor function and upregulation of the FDR MerA. Thus, the S. aureus HypR repressor

![Figure 11](image-url)
can be classified as typical two-Cys-type regulator since its mechanism of repressor inactivation resembles that of the two-Cys-type MarR/DUF24 family repressors HypR and YodB of B. subtilis that are redox sensors of NaOCl, diamide, and quinones and inactivated via intersubunit disulfide formation (18, 73). The NemR repressor of E. coli was shown to sense HOCl and electrophiles, such as NEM and methylglyoxal, by intermolecular disulfide as well as sulfenamide formation (34, 36, 50, 72). Thiol oxidation of these repressors leads to conformational changes and dissociation of the repressors from their operator sites leading to derepression of their target genes.

The structural changes on oxidation have been first studied for the 2-Cys-type MarR/OhrR repressor of Xanthomonas campestris that senses organic hydroperoxides by Cys22-Cys127 intersubunit disulfide formation (51). OhrR oxidation breaks a conserved Cys22 hydrogen-bonding network and causes large structural rearrangements in the dimer interface and rigid body rotation of the DNA-binding domains causing dissociation from the DNA (51). Using mutational analysis, we showed that Cys33 and Cys99 of the S. aureus HypR repressor are important for redox sensing of NaOCl and diamide stress in vivo, while Cys99 is required for repressor activity of HypR. Both Cys33 and Cys99 mutants did not form intermolecular disulfides in vivo as shown by nonreducing SDS-PAGE analysis. These results clearly support the two-Cys-type oxidation model of HypR for Cys33-Cys99 intermolecular disulfide bond formation. Thus, both Cys33 and Cys99 are required for redox sensing of NaOCl and are oxidized to the intersubunit disulfide in vitro and in vivo.

In comparison to other Rrf2 transcription factors, HypR shows the highest sequence identity to SaiR of B. anthracis (20.4%) and YwnA of B. subtilis (23.5%) (Fig. 5A). Moreover, HypR and SaiR show common responses to disulfide stress (diamide, NaOCl) and in infection assays inside macrophages in B. anthracis and S. aureus (9, 68). YwnA also responds to diamide stress (71) and probably controls the adjacent ywnB gene encoding for an NADPH-binding reductase. SaiR of B. anthracis may play a related role like HypR of S. aureus in the protection against oxidative burst of activated macrophages and neutrophils under infection.

FIG. 12. MerA and HypR are required for survival of S. aureus COL in murine macrophages. The survival of S. aureus strains was analyzed 2, 4, and 24 h postinfection (p.i.) of the murine macrophage cell line J-774A.1 and the CFUs were determined. (A, B) The percentages in survival of the ΔhypR and ΔmerA mutants and complemented strains were calculated in 5–6 biological replicate experiments and the survival at the 2-h time point was set to 100%. (C) The average percentage in survival was calculated for each mutant and complemented strain in relation to the wild type (WT), which was set to 100%. Results of 5–6 biological replicates are presented as scatter dots in (A, B) and mean values of percentage survival in comparison to the wild type (C). Error bars represent the SEM and the statistics was calculated using one-way ANOVA and Tukey’s multiple comparisons post hoc test using the GraphPad Prism software. The p-values were determined as follows for the scatter dots (A, B): p = 0.0078 for WT/ΔhypR; p = 0.0303 for WT/ΔmerA; p = 0.0461 for WTpRB473/ΔhypRhypR; and p = 0.1234 for WTpRB473/ΔmerAmerA. For the percentage survival (C), the p-values were determined as p < 0.0001 for WT/ΔhypR, p = 0.0012 for WT/ΔmerA, p = 0.0511 for WTpRB473/ΔhypRhypR, p = 0.4684 for WTpRB473/ΔmerAmerA, p = 0.0459 for ΔhypR/ΔhypRhypR, and p = 0.0725 for ΔmerA/ΔmerAmerA. Symbols are defined as follows: *p > 0.05; *p ≤ 0.05; **p ≤ 0.01; and ****p ≤ 0.0001. CFU, colony-forming unit.
respectively, which is required for repressor activity and pathogens via confer protection under oxidative stress conditions in these (68). Thus, SaiR of the SpxA1 was shown to be required for diamide resistance and merase to activate transcription of a large disulfide stress a responds to NaOCl and diamide stress by thiol oxidation and a member of the ArsC (arsenate reductase) family that re-
spxA2 conditions. SaiR controls a five-gene operon that includes spxA2, saiR, and 3 unknown function genes (68). SpxA2 is a member of the ArsC (arsenate reductase) family that responds to NaOCl and diamide stress by thiol oxidation and interacts with the α-C-terminal domain of the RNA polymerase to activate transcription of a large disulfide stress regulon (69, 70, 99). SpxA2 together with its paralogue SpxA1 was shown to be required for diamide resistance and the saiR mutant conferred a peroxide-resistant phenotype (6, 68). Thus, SaiR of B. anthracis and HypR of S. aureus both confer protection under oxidative stress conditions in these pathogens via control of SpxA2 and MerA, respectively.

SaiR and HypR both share the conserved Cys96 and Cys99, respectively, which is required for repressor activity in vitro and in vivo as well as for redox sensing in vitro (68). To explain, why Cys99 is required for DNA binding, the HypR structure was modeled based on the template of B. subtilis YwaA (1xd7) (Fig. 5A, B). The structural model suggests that Cys33 is located in the turn between the DNA recognition α2 and α3 helices, while the conserved Cys99 is at the N-terminus of the long α6 helix that forms the highly hydrophobic dimer interface required for the elongated fold of Rrf2 family regulators (Fig. 5B). Recently, the structure of Streptomyces coelicolor NsrR was resolved with its [4Fe-4S] cluster in comparison to the three-Cys-Ala (3CA) mutant (90). Interestingly, the FeS cluster ligand Cys99 is in close proximity to Gly37 of the α3 DNA recognition helix of the opposing subunit. In holo-NsrR, Gly37 forms a hydrogen bond with Asn97, which is interrupted in the 3CA mutant resulting in a shift of the α3 DNA binding helix (90). Gly37 of NsrR is the counterpart of Cys33 in HypR, suggesting that the Cys99A mutant also could break hydrogen bonds of Cys33 leading to structural changes in the DNA-binding helices. This is supported by our mutational results since the C99A and Cys99S mutants are unable to bind DNA both in vitro and in vivo. The possible proximity of Cys33 and Cys99 in the structural model (based on the YwaA template) (Fig. 5B) further explains that both are involved in redox sensing of NaOCl and well positioned to form intersubunit disulfides as we confirmed by MS.

Our CD results further showed that HypR contains the predicted high α-helical content. However, we could not show major structural changes in the secondary structure elements between HypR and the three Cys-Ala mutants. It will be interesting to further investigate the crystal structures of reduced and oxidized HypR proteins and the structural changes of the DNA-binding helices on oxidation.

Our gel shift assays and nonreducing SDS-PAGE and Western blot results further revealed that the Cys33 mutant was impaired in redox sensing in vivo, but still responsive to NaOCl and diamide stress in vitro. However, compared to the wild type HypR protein, the Cys33A mutant showed a lower oxidation under diamide stress in the nonreducing SDS-PAGE in vitro. Since we could only detect reduced (alkylated) Cys99 and not the Cys142 peptide in the mass spectrum of the intersubunit disulfide of the Cys33A mutant, the Cys33 mutant protein should be oxidized to the ‘‘wrong’’ ‘‘Cys142-Cys142’’ intersubunit disulfides, crosslinking two HypR dimers in vitro that also lead to HypR inactivation. However, this alternative oxidation was not observed in vivo, and thus, Cys33-Cys99 intersubunit disulfide formation is the physiologically relevant HOCl-sensing mechanism of HypR. Exposure of the purified Cys33 mutant protein to low doses of NaOCl stress might have much stronger effects in vitro and this wrong disulfide might be not stabilized in vivo.

Another unanswered question is why merA and hypR mutants display both sensitivity to NaOCl stress and macrophage infections, significant phenotypes that could be restored by complementation with MerA and HypR. Our unpublished RNA-seq data of the wild-type and the hypR mutant identified only the hypR-merA operon as upregulated in the hypR mutant. Thus, the deficiency of MerA impaired the growth and survival under infections, and also its constitutive overproduction did not confer resistance. It might be possible that overproduction of MerA causes deficiency in the NADPH level, causing in turn a redox poise. The transcriptional results revealed a very low basal level of hypR-merA expression indicating that MerA expression is tightly regulated and its constitutive overexpression perhaps toxic for S. aureus explaining the NaOCl-sensitive phenotype of the hypR mutant.

However, our results clearly identify the NADPH-dependent disulfide reductase MerA as the major HOCl-defense mechanism of S. aureus. MerA also enhances the survival of S. aureus in J774A.1 murine macrophages during infection assays. The J774A.1 cell line was shown to produce NADPH oxidase and MPO during infections and on activation by drug treatment in vitro (55, 65). Increased ROS levels significantly contributed to the antimicrobial activity and killing of intracellular pathogens by the J774A.1 macrophage cell line (54). Thus, MerA could be involved in the defense of S. aureus against ROS and HOCl also during macrophage infections. Of note, there is a striking 49.2% sequence identity between MerA of S. aureus and the FDR RclA of E. coli that is controlled by the RCS-specific transcriptional activator RclR (75) (Supplementary Fig. S10). These conserved flavoenzymes may function as major defense mechanisms under hypochlorite stress in bacteria and are controlled by different redox regulators, the Rrf2 repressor HypR in S. aureus and the AraC activator RclR in E. coli (75).
MerA is annotated as mercuric reductase, but it lacks the auxiliary C-terminal Cys residues that are important for Hg(II) reduction to elemental Hg\(^{0}\) in mercuroid ion reductases (4, 63, 64). Instead, MerA belongs to group-I FDR that uses a redox-active disulfide in the highly conserved C\(_{43}\)XXX\(_{48}\) motif to transfer electrons from NADPH via an FAD cofactor to an unknown disulfide substrate (4). Our results have shown that the MerA-active site Cys43 is important for NaOCl survival since the merAC43S mutant was unable to complement the NaOCl-sensitive phenotype of the merA mutant. The group-I FDR enzymes with known structures include glutathione disulfide reductase (Gor), thioredoxin reductase (TrxR), and mycothiol disulfide reductase (Mtr). We speculate that MerA could be involved in reduction of cellular LMW thiol/disulfides, such as oxidized bacillithiol disulfide (BSSB) or oxidized coenzymeA disulfide (CoAS2), that are increased under NaOCl stress. Our previous results showed an increased BSH redox potential using the genetically encoded Bacilliredoxin-redox-sensitive green fluorescent protein biosensor (58) and increased protein thiol oxidation under NaOCl stress in S. aureus indicating an increased BSSB level (46). Thus, it could be possible that MerA helps to reduce BSSB and CoAS2 to restore the redox balance. On the contrary, expression of MerA and HypR is coregulated, suggesting that both could constitute a redox couple and MerA could be involved in HypR reduction to regenerate the DNA-binding activity of HypR during recovery from HOCl stress. Our future studies are directed to identify the physiological disulfide substrate of MerA to understand the important role of this flavoenzyme in the defense against oxidative burst under macrophage and neutrophil infections.

**Experimental procedures**

**Bacterial strains, growth, and survival assays.** Bacterial strains, plasmids, and primers are listed in Supplementary Tables S3–S5. For cloning and genetic manipulation, *E. coli* was cultivated in Luria Bertani (LB) medium. For stress experiments, *S. aureus* USA300 and COL strains were cultivated in LB, RPMI, or BMM as described previously (58). Depending on the growth medium, different concentrations of NaOCl were applied due to the quenching effect of rich medium components, which affected the effective sublethal NaOCl doses. Thus, we applied sublethal 150 \(\mu M\) NaOCl in BMM as described before (58) for RNA-seq analysis and 1–1.75 \(mM\) sublethal NaOCl in rich RPMI medium to analyze the expression and growth phenotype. For survival phenotype assays, *S. aureus* was grown in RPMI medium until an optical density at 500 nm (OD\(_{500}\)) of 0.5, exposed to a lethal dose of 3.5 \(mM\) NaOCl for 2–4 h, and 10 \(\mu L\) of serial dilutions were spotted onto LB agar plates for 24 h. Complemented mutants that carried the pRB473 plasmid were grown in the presence of 1% xylose and 10 \(\mu g/mL\) chloramphenicol. Sodium hypochlorite, diamide, DTT, Tris(2-carboxyethyl)phosphine (TCEP), \(H_2O_2\) (35% w/v), formaldehyde, and methylglyoxal were purchased from Sigma Aldrich.

**RNA isolation, library preparation, and next-generation cDNA sequencing.** *S. aureus* USA300 was cultivated in BMM and treated with 150 \(\mu M\) NaOCl stress as described previously (58). *S. aureus* cells were harvested before (as untreated control) and 30 min after exposure to 150 \(\mu M\) NaOCl and disrupted in 3 \(mM\) ethylenediaminetetraacetic acid (EDTA)/200 \(mM\) NaCl lysis buffer with a Precellys24 ribolyzer. RNA isolation was performed using the acid phenol extraction protocol as described (19). The RNA quality was checked by Trinean Xpose (Gentbrugge, Belgium) and the Agilent RNA Nano 6000 kit using an Agilent 2100 Bioanalyzer (Agilent Technologies, Böblingen, Germany). Ribo-Zero rRNA Removal Kit (Bacteria) from Illumina (San Diego, CA) was used to remove the rRNA. TruSeq Stranded mRNA Library Prep Kit from Illumina was applied to prepare the cDNA libraries. The cDNAs were sequenced paired end on an Illumina HiSeq 1500 and MiSeq system (San Diego, CA) using 50 and 70 bp read length. The transcriptome sequencing raw data files are available in the ArrayExpress database under accession number: E-MTAB-5666.

**Bioinformatics data analysis, read mapping, data visualization, and analysis of differential gene expression.** Trimmed reads (26 nt) were mapped to the *S. aureus* USA300_TCH1516 genome sequence (accession number CP000730) (41) using SARUMAN (11), allowing one error per read. The forward and reverse reads, if both present and with a maximum distance of 1 kb, were combined to one read containing the reference sequence as insert. Paired mappings with a distance >1 kb were discarded, and paired reads with either only the forward or the reverse read mapping were retained as single mapping reads. For the visualization and counting of short read alignments, ReadXplorer v2.2 (42) was used.

Differential gene expression analysis, including normalization, was performed using Bioconductor package DESeq2 (59) included in the ReadXplorer v2.2 software (42). The signal intensity value (a-value) was calculated by log2 mean of normalized read counts and the signal intensity ratio (m-value) by log2 fold change. The evaluation of the differential RNA-seq data was performed using an adjusted p-value cutoff of \(p \leq 0.05\) and a signal intensity ratio (m-value) cutoff of \(\geq 1.98\) or \(-1.98\). The latter was determined by applying a significance level of 5% to the experiment with the assumption that the majority of genes are not differentially transcribed. Thus, 95% of all m-values should fall in this range. Therefore, the standard deviation (STDEV) for all m-values was calculated and the cutoff was set to \(m = 1.96 \times STDEV\). Genes with an m-value outside this range and \(p \leq 0.05\) were considered as differentially transcribed.

**Construction of Voronoi treemaps.** The software Paver (DECODON GmbH, Greifswald, Germany) was used to generate the regulon treemap showing log2 fold changes (m-values) of selective genes sorted into operons and regulons that are upregulated under NaOCl stress compared to the untreated control. The treemap construction is based on a model of competing particle swarms for the layout optimization as described (62). The hierarchical structure of the treemap is defined by the regulons (on the first level—bold white labels), genes, and operons (second level—small contrast optimized labels). Cell sizes are defined as absolute log2 fold changes of expression levels after NaOCl stress divided by the untreated control. Colors are defined by a symmetric divergent color gradient (negative values—blue and positive values orange—red) of NaOCl versus control log2 fold ratios.
Cloning, expression, and purification of His-tagged HypR and HypR Cys mutant proteins in *E. coli*. The hypR gene (SACOL0641) was amplified from chromosomal DNA of *S. aureus* COL by PCR using primers 0641-pET-for-Nhel and 0641-pET-rev-BamHI (Supplementary Table S5), digested with Nhel and BamHI and inserted into plasmid pET11b (Novagen) that was digested using the same restriction enzymes to generate plasmid pET11b-hypR.

For construction of HypR Cys-to-Ala and Cys-to-Ser variants, the Cys residues of HypR were replaced by alanine or serine, respectively, using PCR mutagenesis. For the hypR-C33A mutant, two first-round PCRs were performed using primers 0641-pET-for-Nhel and 0641-pET-C33A-Rev and primers 0641-pET-for-C33A and 0641-pET-rev-BamHI (Supplementary Table S5). For the hypR-C99A mutant, two first-round PCRs were performed using primers 0641-pET-for-Nhel and 0641-pET-C99A-rev and primers 0641-pET-C99A-for and 0641-pET-rev-BamHI. For the hypR-C99S mutant, two first-round PCRs were performed using primers 0641-pET-for-Nhel and 0641-pET-C99S-rev and primers 0641-pET-C99S-for and 0641-pET-rev-BamHI. The two PCR products of each first-round PCR were hybridized and subsequently amplified by a second round of PCRs using primers 0641-pET-for-Nhel and 0641-pET-rev-BamHI. The two PCR products from the second-round PCRs were digested with Nhel and BamHI and inserted into plasmid pET11b digested with the same enzymes to generate plasmids pET11b-hypR-C33A, pET11b-hypR-C99A, and pET11b-hypR-C99S. The hypR-C142A gene was amplified from chromosomal DNA of *S. aureus* COL by PCR using primers 0641-pET-for-Nhel and 0641-pET-C142A-rev-BamHI, digested with Nhel and BamHI and inserted into plasmid pET11b that was digested using the same restriction enzymes to generate plasmid pET11b-hypR-C142A. The correct sequences of all inserts were confirmed by PCR and DNA sequencing.

For expression and purification of His6-tagged HypR wild-type and Cys mutant proteins, *E. coli* BL21(DE3) plysS was used that contains the plasmids pET11b-hypR, pET11b-hypR-C33A, pET11b-hypR-C99A, pET11b-hypR-C99S, and pET11b-hypR-C142A. These *E. coli* expression strains were cultivated in 1 L LB medium until the exponential growth phase at OD600 of 0.8 followed by addition of 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) at 37°C. E. coli strains expressing recombinant His6-tagged HypR and the HypR Cys mutant proteins were disrupted by sonication in binding buffer (20 mM NaH₂PO₄, 500 mM NaCl, 20 mM imidazole, pH 7.4). Lysates were cleared from cell debris by repeated centrifugation, and purification of the His-tagged proteins was performed using His Trap™ HP Ni-NTA columns (5 mL; GE Healthcare, Chalfont St Giles, United Kingdom) and the AKTA purifier liquid chromatography system (GE Healthcare) according to the instructions of the manufacturer. Supplementary Figure S11 shows the HypR purification profile in the Akta chromatography. A gradient from 0 to 500 mM imidazole was applied in elution buffer (20 mM NaH₂PO₄, 500 mM NaCl, 500 mM imidazole, pH 7.4) and HypR eluted at ~240 mM imidazole. The purity of eluted and pooled fractions was analyzed by reducing SDS-PAGE (Supplementary Fig. S11, inset). The purified proteins were extensively dialyzed against 10 mM Tris-HCl (pH 8.0), 100 mM NaCl, and 30% glycerol and stored at ~80°C. The HypR protein concentrations were determined by UV absorbance using the Thermo Nanodrop 2000c spectrophotometer (Thermo Scientific, Germany). The final HypR protein concentration after dialysis was 2.8 mg/mL (158.7 μM) with a threefold enrichment after dialysis. Before further usage, HypR proteins were freshly reduced with 10 mM DTT or 10 mM TCEP or as otherwise stated in the methods and figure legends.

**Construction of S. aureus COL hypR and merA deletion mutants and complemented Cys-Ala mutant strains.** *S. aureus* COL mutants with clean deletions of hypR (SACOL0641) and merA (SACOL0640) were constructed by allelic replacement via the temperature-sensitive shuttle vector pMAD as described previously (5). Briefly, for construction of the plasmids, pMAD-hypR and pMAD-ΔmerA, the flanking gene regions, including 500 bp upstream and downstream of hypR and merA of *S. aureus* COL were amplified using the primers SACOL0641-pMAD and SACOL0640-pMAD (Supplementary Table S5). The upstream and downstream flanking regions of hypR and merA were each fused by overlap extension PCR and ligated into the BglII and SalI sites of plasmid pMAD. The pMAD constructs were electroporated into the restriction-negative and methylation-positive intermediate *S. aureus* RN4220 strain and further transferred to *S. aureus* COL by phage transduction using phage 80 (81). Transductants were streaked out on LB agar with 10 μg/mL erythromycin and 40 μg/mL 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal) at 30°C. Blue transductants with pMAD integrations were used for plasmid excision by a heat shock and screened for erythromycin-sensitive white colonies on X-gal plates as described (66). The clean deletions of internal gene regions of hypR or merA were confirmed by PCR and sequencing.

The complemented hypR and merA deletion mutants and Cys mutant strains were constructed using the pRB473-XylR plasmid as described previously (58). Briefly, hypR, hypR-C33A, hypR-C99A, hypR-C142A, merA, and merAC43S sequences were amplified from the *E. coli* expression plasmids pET11b-hypR, pET11b-hypR-C33A, pET11b-hypR-C99A, pET11b-hypR-C142A, pET11b-merA, and pET11b-merAC43S using the following primers. Primers 0641-pRB-for-BamHI and 0641-pRB-REV-KpnI were used to construct plasmids pRB473-XylR-hypR, pRB473-XylR-hypR-C33A, and pRB473-XylR-hypR-C99A. Primers 0641-pRB-REV-KpnI and 0641-pRB-C2-3A were used to construct pRB473-XylR-hypR-C142A. Primers 0640-pRB-for-BamHI and 0640-pRB-REV-KpnI were used to construct plasmids pRB473-XylR-merA and pRB473-XylR-merAC43S. Each forward primer also includes the shine-dalgarno sequence of hypR or merA, respectively. The PCR product was digested with BamHI and KpnI and inserted into the pRB473-XylR shuttle vector that was digested using the same enzymes. The recombinant plasmids were introduced into the ΔhypR or ΔmerA deletion mutants via phage transduction as described previously (58).

**Northern blot experiments.** Northern blot analyses were performed as described previously (93) using RNA isolated from the *S. aureus* COL strains that were exposed to 1 mM NaCl, 2 mM diamide, 10 mM H₂O₂, 0.75 mM formaldehyde, and 0.5 mM methylglyoxal (MG) for 15 and 30 min as indicated. Hybridizations were performed with the digoxigenin-labeled hypR- and merA-specific antisense RNA.
probes synthesized in vitro using T7 RNA polymerase as described previously (89). The primer pairs SACOL0641-for and SACOL0641-rev or SACOL0640-for and SACOL0640-rev were each used for generation of the digoxigenin-labeled hypR and merA antisense RNA probes, respectively.

EMSAs of HypR. For EMSAs, we amplified a DNA fragment containing the hypR-merA upstream region that covered the region from -128 to +70 relative to the transcriptional start site using PCR with the primer set emsa0641-for and emsa0641-rev (Supplementary Table S5). Approximately 0.75 ng of purified PCR products was incubated with different amounts of purified His-HypR and His-HypR Cys mutant proteins for 45 min at room temperature in EMSA binding buffer (10 mM Tris-HCl [pH 7.5], 100 mM KCl, 5% glycerol, 50 µg/mL bovine serum albumin (BSA), and 5 µg/mL salmon sperm DNA) in the presence of 10 mM DTT. To study the effect of NaOCl and diamide on the DNA-binding activity in vitro, the purified proteins were reduced with 10 mM EDTA for 20 min, desalted with Micro-Bio spin columns (Biorad), and diluted to 0.6 µM in 10 mM Tris-HCl, pH 7.5. Reduced HypR and Cys mutant proteins were oxidized with 1.25–20 µM NaOCl or 2.5–20 µM diamide for 10 min at room temperature and incubated with the DNA fragment (0.75 ng) containing the hypr-merA promoter. DNA-binding reactions were separated by 4% native polyacrylamide gel electrophoresis in 10 mM Tris, 1 mM EDTA buffer, pH 8, containing 2.5% glycerol at room temperature and constant voltage (180 V) for 30 min. Gels were stained with SYBR green (Thermo Fisher Scientific) for 30 min in the dark and fluorescence was visualized using a Typhoon scanner (Typhoon FLA 9500; GE Healthcare Life Sciences).

CD spectroscopy. CD spectra of DTT-reduced HypR and HypR Cys mutant proteins were obtained using a Jasco J-810 spectropolarimeter with a HAAKE WKL recirculating chiller (D-76227, Karlsruhe). The DTT-reduced proteins were measured at 10 µM in 20 mM potassium phosphate buffer, pH 7.5, with 1 mM DTT. The quartz cuvettes (2 mm path length, Suprasil Hellma) were set at a constant temperature of 25°C with a Jasco PTC-423S Peltier-type thermostate. Secondary structure elements were calculated using the program DichroWeb (http://dichroweb.cryst.bbk.ac.uk) (86, 94).

SEC-MALS analysis. HypR proteins were diluted in SEC buffer (10 mM HEPES, 500 mM NaCl, pH 7.4, 1 mM TCEP) and reduced with 10 mM TCEP at 23°C for 1 h. The molecular weights and oligomerization states of HypR and the HypR Cys mutants were determined using SEC–MALS analysis at 20°C running in SEC buffer. The reduced HypR protein samples (170 µM each) were passed over a Superdex 75 10/300 size-exclusion column (GE Healthcare) coupled to a miniDAWN TREOS three-angle light scattering detector (Wyatt Technology) and a RefractoMax520 refractive index detector (ERC). Detectors were aligned, corrected for band broadening, and the photodiodes of the miniDAWN TREOS were normalized with BSA as a reference. Data were analyzed with ASTRA 6.1.4.25 (Wyatt Technology). For calculation of the molecular masses, protein concentrations were determined from the differential refractive index with a specific refractive index increment (dn/dc) of 0.185 mL/g.

Nonreducing/reducing diagonal SDS-PAGE analysis. S. aureus COL strains were cultivated in BMM, treated with 150 µM NaOCl for 30 min as described (58), harvested, and washed in TE-buffer (10 mM Tris-HCl, pH 8; 1 mM EDTA) with 50 mM NEM to alkylate reduced thiols. Cells were disrupted using the ribolyzer, and the protein extracts were obtained after repeated centrifugation. Immunoprecipitation of HypR proteins was performed using HypR-polyclonal antibodies and S. aureus protein extracts with Dynabeads protein A (Invitrogen) according to the instructions of the manufacturer. The precipitated HypR proteins were eluted in nonreducing SDS sample buffer (2% SDS, 62.5 mM Tris-HCl [pH 8.0]). The protein extracts and immunoprecipitated HypR proteins were separated by 15% nonreducing/reducing SDS-PAGE as described previously (53). In the first SDS-PAGE, proteins were separated in nonreducing SDS-PAGE sample buffer. The lanes were cut and incubated in reducing SDS sample buffer (with 50 mM DTT) followed by thiol alkylation with 50 mM iodoacetamide (IAM) for each 15 min. The bands were positioned horizontally on an SDS-PAGE gel, separated using reducing SDS-PAGE, and subjected to HypR-specific Western blot analysis as described previously (22). HypR forms intermolecular disulfides between two subunits that run at the right side of the diagonal.

Western blot analysis. S. aureus COL cells were harvested, washed in TE buffer (pH 8.0) with 50 mM NEM, disrupted using the ribolyzer, and the protein extract was cleared from cell debris by repeated centrifugation as described above. Protein amounts of 25 µg were diluted in nonreducing or reducing SDS sample buffer (as above), separated using 15% SDS–PAGE, and the Western blot analysis was performed as described previously (22). Anti-HypR polyclonal rabbit antiserum was generated using purified His-HypR protein and used at a dilution of 1:500 for Western blot analyses.

MALDI-TOF-MS of in vitro oxidized HypR protein. The purified HypR and HypR-C33A proteins were oxidized with NaOCl and all reduced thiols were blocked with 50 mM IAM resulting in the mass increase of 57 Da at Cys residues. The oxidized HypR and HypR-C33A intermolecular disulfides were separated by nonreducing SDS–PAGE, followed by tryptic digestion of the HypR and HypR-C33A disulfide bands as described (21). The peptide mass spectra were measured using a MALDI-TOF/MS using an Ultraflex-II TOF/TOF instrument (Bruker Daltonics, Bremen, Germany) equipped with a 200 Hz solid-state Smart beam™ laser. The mass spectrometer was operated in the positive reflector mode. Mass spectra were acquired over an m/z range of 600–4000. MS/MS spectra of selected peptides were acquired in the LIFT mode as described previously (87).

Infection assays with murine macrophage cell line J-774A.1. Murine macrophage cell line J-774A.1 (80) was cultivated in Iscove’s modified Dulbecco’s medium (Biochrom) with 10% heat-inactivated fetal bovine serum (FBS) and seeded in cell culture dishes for 2 days under 5% CO2 at 37°C as described (79). Before the infection assay, S. aureus COL strains were plated overnight on LB agar and re-suspended in eukaryotic growth medium. Macrophages were infected with S. aureus cells at a multiplicity of infection (MOI) of 1:25. One hour after infection, the cell culture medium was replaced and 150 µg/mL gentamycin was added.
for 1 h to kill extracellular bacteria and to stop the uptake of \( S. aureus \). The intracellular survival was determined at 2, 4, and 24 h after phagocytosis. Infected macrophages were lysed with 0.1% Triton X-100 and the supernatant with internalized intracellular bacteria was plated on agar plates for CFUs.

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Author Disclosure Statement

No competing financial interests exist.

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**Abbreviations Used**

- **BMM** = belitsky minimal medium
- **BSA** = bovin serum albumin
- **BSH** = bacillithiol
- **BSSB** = oxidized bacillithiol disulfide
- **CD** = circular dichroism
- **CFU** = colony-forming unit
- **Chp** = chemotaxis-inhibitory protein
- **CoAS2** = oxidized coenzyme A disulfide
- **CoASH** = coenzyme A
- **CynR** = cysteine metabolism repressor
- **DTT** = dithiothreitol
- **EDTA** = ethylenediaminetetraacetic acid
- **EMSA** = electrophoretic mobility shift assay
- **FDR** = flavin disulfide reductase
- **FeS** = iron/sulfur
- **H2O2** = hydrogen peroxide
- **HOCI** = hypochloric acid
- **IAM** = iodoacetamide
- **IsR** = iron/sulfur status of the cell
- **LB** = Luria Bertani
- **LMW** = thiol low-molecular-weight thiol
- **MALDI-TOF-MS** = matrix-assisted laser desorption ionization-time of flight mass spectrometry
- **MerA** = mercuric ion reductase
- **MPO** = myeloperoxidase
- **MRSA** = methicillin-resistant *Staphylococcus aureus*
- **M-value** = log2 fold change
- **NADPH** = nicotinamide adenine dinucleotide phosphate
- **NaOCl** = sodium hypochlorite
- **NEM** = N-ethylmaleimide
- **NsrR** = redox sensors for nitric oxide
- **OD500** = optical density at 500 nm
- **RCS** = reactive chlorine species
- **RES** = reactive electrophilic species
- **ROS** = reactive oxygen species
- **SEC-MALS** = size-exclusion chromatography/multangle light scattering
- **STDEV** = standard deviation
- **TCEP** = tris(2-carboxyethyl)phosphine
- **X-gal** = bromo-4-chloro-3-indolyl-β-D-galactopyranoside