Distinct H$_2$O$_2$-Scavenging System in Yersinia pseudotuberculosis: KatG and AhpC Act Together to Scavenge Endogenous Hydrogen Peroxide

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To colonize in the digestive tract of animals and humans, Yersinia pseudotuberculosis has to deal with reactive oxygen species (ROS) produced by host cells and microbiota. However, an understanding of the ROS-scavenging systems and their regulation in this bacterium remains largely elusive. In this study, we identified OxyR as the master transcriptional regulator mediating cellular responses to hydrogen peroxide (H$_2$O$_2$) in Y. pseudotuberculosis through genomics and transcriptomics analyses. OxyR activates transcription of diverse genes, especially the core members of its regulon, including those encoding catalases, peroxidases, and thiol reductases. The data also suggest that sulfur species and manganese may play a particular role in the oxidative stress response of Y. pseudotuberculosis. Among the three H$_2$O$_2$-scavenging systems in Y. pseudotuberculosis, catalase/peroxidase KatE functions as the primary scavenger for high levels of H$_2$O$_2$; NADH peroxidase alkyl hydroperoxide reductase (AhpR) and catalase KatG together are responsible for removing low levels of H$_2$O$_2$. The simultaneous loss of both AhpC (the peroxidatic component of AhpR) and KatG results in activation of OxyR. Moreover, we found that AhpC, unlike its well-characterized Escherichia coli counterpart, has little effect on protecting cells against toxicity of organic peroxides. These findings provide not only novel insights into the structural and functional diversity of bacterial H$_2$O$_2$-scavenging systems but also a basic understanding of how Y. pseudotuberculosis copes with oxidative stress.

Keywords: Yersinia, OxyR, AhpC, catalase, oxidative stress response

INTRODUCTION

Oxidative stress caused by reactive oxygen species (ROS), including superoxide (O$_2$\textsuperscript{−}), hydrogen peroxide (H$_2$O$_2$), and hydroxyl radical (OH\textsuperscript{−}), is inevitable to all organisms that respire oxygen (Li et al., 2016). These strong oxidants and/or radicals could damage virtually all biomolecules, such as nucleic acids, proteins, and lipids (Imlay, 2013). Naturally, detoxification of ROS is extremely critical for survival of diverse bacteria, pathogens in particular, because the host cells release ROS as a deadly weapon to defend against bacterial infections (Fang, 2004). Among ROS, H$_2$O$_2$ not only can be formed very rapidly endogenously (for example, 15 µM s\textsuperscript{−1} in Escherichia coli) but also...
OxyR functions as an activator only for major \( \text{H}_2\text{O}_2 \) oneidensis Neisseria and an activator for certain genes) in it can also function in a dual-control manner (both a repressor residues [Cys 199 and Cys 208 within an intramolecular disulfide bond between two conserved cysteine residues] (Zheng et al., 1998). The regulatory mode of OxyR varies AhpR in \( \text{H}_2\text{O}_2 \) (Cys 208) within \( \text{Cys} \). Therefore, in response to \( \text{H}_2\text{O}_2 \) stress, the regulatory mode of OxyR varies between repressors and activators. The role of OxyR in the context of \( \text{H}_2\text{O}_2 \) stress is not yet fully understood.

**Yersinia pseudotuberculosis** is a Gram-negative enteric pathogen that often causes self-limiting gastrointestinal disorders such as enteritis, diarrhea, and mesenteric lymphadenitis to animals and humans (Brady et al., 2020). During infection, environmental stress and host immunity reactions in human guts can introduce an increase in levels of ROS that \( \text{Y. pseudotuberculosis} \) cells encounter, thus requiring a sophisticated regulation of genes to reduce the intracellular \( \text{H}_2\text{O}_2 \) level to a safe line (Knaus et al., 2017). It has been reported that \( \text{Y. pseudotuberculosis} \) is able to combat oxidative stress by an unconventional way: it imports zinc to mitigate ROS by secreting a zincophore via the type VI secretion system (T6SS; Wang et al., 2016, 2020). However, how this bacterium copes with oxidative stress has not been systematically investigated and thus is poorly understood.

In this study, we carried out the transcriptomics analysis of \( \text{Y. pseudotuberculosis} \) in response to exogenous \( \text{H}_2\text{O}_2 \). We identified an OxyR analog in YPIII and found that its regulon members are highly conserved, including those encoding \( \text{H}_2\text{O}_2 \)-scavenging enzymes, iron-sequestering proteins, and thiol-reducing systems. Our data support that OxyR of YPIII functions in an activator-only mode, and its absence causes a plating defect on LB agar, a result of insufficient production of \( \text{H}_2\text{O}_2 \)-scavenging enzymes, including KatE, KatG, and AhpC. We further showed that KatE functions as the primary scavenger for \( \text{H}_2\text{O}_2 \), but both KatG and AhpC together play an essential role in decomposing low levels of \( \text{H}_2\text{O}_2 \). The simultaneous loss of both KatG and AhpC results in the loss of catalase-peroxidase activity, which is reactivated by OxyR using NADPH as reducing equivalent (Perkins et al., 2015). Atypical OxyR systems, in which OxyR is absent, have been found in some bacteria (Bryk et al., 2000; Baker et al., 2001). For instance, in Helicobacter pylori, OxyR is repressed by a thioredoxin/thioredoxin reductase (TrxB) system, and in Mycobacterium tuberculosis, AhpD reduces AhpC with electrons from NADH through dihydrolipoamide dehydrogenase and dihydrolipoamide succinyltransferase (Baker et al., 2001; Jaeger et al., 2004). Because of the essential roles of catalase and OxyR in \( \text{H}_2\text{O}_2 \)-scavenging, bacterial strains lacking both together display a drastically elevated sensitivity to \( \text{H}_2\text{O}_2 \) and carry an apparent aerobic growth defect (Cosgrove et al., 2007; Ezraty et al., 2017).

In many bacteria, OxyR is a primary transcriptional regulator that mediates cellular response to oxidative stress (Imlay, 2013; Fu et al., 2015). As a LysR-family DNA-binding protein, OxyR senses and responds to \( \text{H}_2\text{O}_2 \) stress via formation of an intramolecular disulfide bond between two conserved cysteine residues [Cys 199 and Cys 208 within \( \text{E. coli} \) OxyR (EcOxyR); Zheng et al., 1998]. The regulatory mode of OxyR varies among bacteria. For example, in \( \text{E. coli} \), Salmonella enterica serovar Typhimurium (S. Typh), and many other bacteria, OxyR functions as an activator only for major \( \text{H}_2\text{O}_2 \)-scavenging proteins, such as catalases and AhpR (Imlay, 2008). However, it can also function in a dual-control manner (both a repressor and an activator for certain genes) in Neisseria and Shewanella oneidensis or as a repressor only in Corynebacterium diphtheriae (Iva et al., 2008; Kim and Holmes, 2012; Jiang et al., 2014). OxyR proteins now have been generally regarded as a global regulator implicated in diverse biological processes, but the core members of their regulons are consistently constituted by operons responsible for \( \text{H}_2\text{O}_2 \) degradation such as catalases and peroxidases, iron-sequestering proteins, and thioredoxin and glutathione antioxidant systems (Imlay, 2015).

**Materials and Methods**

**Bacterial Strains, Plasmids, and Culture Conditions**

All bacterial strains and plasmids used in this study are listed in Table 1. All chemicals were obtained from Sigma (Shanghai, China) unless otherwise noted. For genetic manipulation, \( \text{E. coli} \) and \( \text{Y. pseudotuberculosis} \) were grown in LB (containing 1% tryptone, 0.5% yeast extract, and 0.5% NaCl) under aerobic conditions at 37 and 26°C. When needed, the following chemicals were added to the growth medium: 2,6-diaminopimelic acid (DAP), 0.3 mM; ampicillin, 50 \( \mu \text{g/ml} \); kanamycin, 50 \( \mu \text{g/ml} \); gentamycin, 15 \( \mu \text{g/ml} \); and streptomycin, 100 \( \mu \text{g/ml} \).
TABLE 1 | Bacterial strains and plasmids used in this study.

| Strains                      | Description                                      | Source or references |
|------------------------------|--------------------------------------------------|----------------------|
| **Strains**                  |                                                  |                      |
| E. coli                      | Host strain for cloning                          | Lab stock            |
| DH15a                        | ΔdapA, donor strain for conjugation              | W. Metcalf, UIUC     |
| WM3064                       | ΔkatG derived from YPIII                          | Rosqvist et al., 1988|
| S. oneidensis MR-1           | Wild type                                        | Jiang et al., 2014   |
| HG1070                       | ΔkatB derived from MR-1                          | Feng et al., 2020    |
| HG00988-1070                 | ΔkatEΔahpC derived from MR-1                     | This study           |
| Y. pseudotuberculosis YPIII  | Wild type                                        | This study           |
| YPK_RS20585                  | ΔoxyR derived from YPIII                          | This study           |
| YPK_RS14285                  | ΔkatE derived from YPIII                          | This study           |
| YPK_RS17025                  | ΔkatG derived from YPIII                          | This study           |
| YPK_RS16370                  | ΔahpC derived from YPIII                          | This study           |
| YPK17025-16370               | ΔkatGΔahpC derived from YPIII                    | This study           |
| YPK14285-17025               | ΔkatEΔkatG derived from YPIII                    | This study           |
| YPK3CAT                      | ΔkatEΔkatGΔahpC derived from YPIII                | This study           |
| **Plasmids**                 |                                                  |                      |
| pHGM01                       | Ap'/Gm'/Cm'/att-based suicide vector             | Jin et al., 2013     |
| pHG101                       | Promoterless vector for complementation          | Wu et al., 2011      |
| pHGE101                      | Km', integrative lacZ reporter vector            | Fu et al., 2014      |
| pHGE101-katE                 | pHGE101 containing the katE promoter             | This study           |
| pHGE101-katG                 | pHGE101 containing the katG promoter             | This study           |
| pHGE101-ahpC                 | pHGE101 containing the ahpC promoter             | This study           |

Transcriptomics Analysis

For transcriptomics analysis, cell samples were prepared as described previously (Jiang et al., 2014). In brief, cultures of the mid-exponential phase (≈0.4 of OD₆₀₀, the same throughout the study) were subjected to the H₂O₂ treatment. H₂O₂ was added to a final concentration of 0.5 mM, and cells were collected before and 5 min after the addition. Cells were pelleted at 14,000 rpm for 30 s at room temperature and frozen immediately in liquid nitrogen. Three biological replicates under each condition were prepared. Total RNA was extracted with RNeasy Mini Kit (QIAGEN, Alameda, United States), and RNA sequencing (RNA-seq) analysis was conducted as before (Gao et al., 2004; Sun et al., 2020). Data analysis was conducted according to standard procedures used previously (Sun et al., 2020). Genes that passed statistical analysis by analysis of variance (ANOVA; p < 0.05) with Benjamini–Hochberg false discovery rate multiple-testing correction and showed two-fold difference between the H₂O₂-treated and untreated control samples were discussed in the study. NCBI SRA accession number is PRJNA671546 for raw transcriptomics analysis data.

Real-Time Quantitative RT-PCR

Quantitative reverse transcription-PCR (qRT-PCR) was performed to verify the expression of key OxyR regulon members with an ABI7300 96-well qRT-PCR system (Applied Biosystems) as described previously (Jiang et al., 2014). The expression of each gene was determined from four replicas in a single real-time qRT-PCR experiment. The cycle threshold (Cₚ) values for each gene of interest were averaged and normalized against the Cₚ value of the 16s rRNA gene, whose abundance was constant during the exponential phase. Relative abundance (RA) of each gene was standardized to the Cₚ values of the 16s rRNA gene using the equation RA = 2⁻ΔCₚ.

Mutagenesis, Complementation of Mutant Strains

Yersinia pseudotuberculosis in-frame deletion strains were constructed by the att-based Fusion PCR method as described previously (Jin et al., 2013). In brief, two fragments flanking the target gene were generated by PCR with primers containing attB and the gene-specific sequence, which were linked by a linker sequence via second round of PCR. The fusion fragments were integrated into plasmid pHGM01 by site-specific recombination using Gateway BP clonase II enzyme mix (Invitrogen). The resulting vectors were introduced in E. coli WM3064 and transferred to Y. pseudotuberculosis by conjugation. Integration of the mutagenesis constructs into the chromosome was selected by resistance to gentamycin and confirmed by PCR. Verified trans-conjugants were grown in LB broth without NaCl and plated on LB supplemented with 10% sucrose. Gentamycin-sensitive and sucrose-resistant colonies were screened by PCR for deletion of the target gene. To facilitate growth of mutants, catalase (from bovine liver, Sigma) was added onto the plates. All mutations were verified by sequencing the mutated regions.

Plasmid pHG-101 was used in genetic complementation of mutants as described before (Wu et al., 2011). For complementation of genes next to their promoter, a fragment containing the gene of interest and its native promoter was amplified by PCR and cloned into pHG-101. After sequencing verification, the resulting vectors were transferred into the relevant strains via conjugation.

Growth and Susceptibility to H₂O₂ or t-BHP

The spotting assay was used to evaluate the plating defect on LB plates. Cells of the mid-exponential phase were collected by centrifugation and adjusted to 10⁹ cells/ml, which was set as the undiluted (dilution factor 0). Ten-fold serial dilutions were prepared with fresh LB medium. Five microliter of each dilution was spotted onto LB plates. The plates were incubated for 24 h or longer in the dark before being photographed. All experiments were repeated at least three times.
Disk diffusion assays to test for sensitivity to oxidative stress conditions were performed with *Y. pseudotuberculosis* strains. Two hundred microliter of mid-exponential phase cultures were spread onto LB plates, 6 mm (in diameter) paper disks loaded with 10 µl H$_2$O$_2$ or tert-butyl hydroperoxide (t-BHP) of various concentrations were placed onto the bacterial lawn grown for 6 h, and plates were incubated 26°C for 16 h.

**H$_2$O$_2$ Quantification**

H$_2$O$_2$ at high concentrations (>100 µM) was quantified using the ferrous ion oxidation-xylene orange method (Wolff, 1994). In brief, cells of the mid-exponential phase grown in liquid LB were collected by centrifugation, washed in PBS, and resuspended in the same buffer to an OD$_{600}$ of 0.1. H$_2$O$_2$ was added to final concentrations indicated in the figure legends. Cells were filtered out at different time points, and elutions were assayed immediately for the remaining H$_2$O$_2$.

H$_2$O$_2$ at low concentrations (<50 µM) was quantified using Amplex red fluorescent method (Seaver and Imlay, 2001). In the presence of H$_2$O$_2$, Amplex red can be oxidized by horseradish peroxidase to the fluorescent product resorufin. To measure H$_2$O$_2$, 200 µl of samples from cells grown in MS medium was mixed with 100 µl of stock solutions for Amplex red and horseradish peroxidase prepared the same as described elsewhere (Seaver and Imlay, 2001). Fluorescence was then measured in a Synergy 2 Pro200 Multi-Detection Microplate Reader (Tecan) and converted to H$_2$O$_2$ concentration using a curve obtained from standard samples.

**β-Galactosidase Activity Assay**

β-Galactosidase activity assay was used to determine gene expression. The sequence in question length (~400 bp) upstream of gene of interest was amplified and inserted in front of the full-length *E. coli* lacZ gene in plasmid pHGEI01 (Fu et al., 2014). The resulting plasmid was verified by sequencing, introduced into *E. coli* WM3064, and then conjugated with relevant *Y. pseudotuberculosis* strains. Cultures of the mid-exponential phase were collected by centrifugation, washed with PBS, and treated with lysis buffer (0.25 M Tris/HCl, 0.5% Triton X-100, and pH 7.5). Extracts were collected by centrifugation and applied for enzyme assay by adding o-nitrophenyl-β-D-galactopyranoside (4 mg/ml). Changes in absorption over time were monitored at 420 nM with a Synergy 2 Pro200 Multi-Detection Microplate Reader (Tecan), and the results were presented as Miller units.

**Bioinformatics and Statistical Analyses**

Multiple sequence alignment was carried out with Clustal Omega (Madeira et al., 2019). Sequence logos were generated by using WebLogo (Crooks et al., 2004). Three-dimensional structures of YpAhpC were predicted using Phyre (Kelley et al., 2015). The predicted structures were then visualized by software Pymol (DeLano Scientific LLC). Genome screening for OxyR-binding sites based on established weight matrices from various bacteria was performed using regulatory sequence analysis tools (RSATs; Medina-Rivera et al., 2015). For statistical analysis, Student's t-test was performed for pairwise comparisons of groups, and values are presented as means ± standard deviation (SD).

**RESULTS**

**Genomics Analysis of *Y. pseudotuberculosis* YPIII With Respect to Oxidative Stress Response**

To identify the oxidative stress response regulator(s) in YPIII, a BLASTp search of functional analogs of established oxidative stress-responding regulators, including EcOxyR and SoxRS as well as *Bacillus subtilis* PerR, against the YPIII proteome was performed. While no homolog of *E. coli* SoxRS or of *B. subtilis* PerR was found, YPIII possesses a highly confident homolog of EcOxyR, YPK_RS20585 (BLASTp E-value = 0; identities, 88%; Supplementary Figure 1). Like all 2-Cys OxyRs, YPK_RS20585 contains two conserved cysteine residues (Cys199 and Cys208 within both YPK_RS20585 and EcOxyR) implicated in the activation by disulfide bond formation, suggesting a possible role in the oxidative stress response of YPIII, and therefore we named it OxyR (YpOxyR). However, the sequence similarities between YpOxyR and dual-activity OxyRs, such as that of *S. oneidensis*, are substantially lower (E-value = 7e-47; identities, 33%; Supplementary Figure 2), implying that YpOxyR might function as an activator only.

The current knowledge on bacterial oxidative stress response is most well developed in *E. coli*, whose OxyR regulon is composed of over 20 operons (Imlay, 2015). The most important and conserved EcOxyR regulon members encode proteins involved in detoxification and prevention and/or repair of oxidative damage, such as catalases and peroxidases, iron-sequestering proteins, thioredoxin, and glutathione antioxidant systems. Similarly, the YPIII genome encodes two catalases (HPII KatE and HPI KatG), iron-sequestering protein Dps, and a complete set of thioredoxin and glutathione antioxidant proteins, including TrxA (thioredoxin), TrxB (thioredoxin-disulfide reductase), TrxC (thioredoxin), GrxA (glutaredoxin), YPK_RS20590 (glutathione peroxidase), and GorA (glutathione-disulfide reductase; Table 2).

Perhaps one of the most striking observations is that YPIII seemingly lacks the counterpart of *E. coli* AhpF, the cognate reductase for peroxidase component AhpC of AhpR (Tartaglia et al., 1990; Poole and Ellis, 1996). A BLASTp search using *E. coli* AhpCF against the YPIII proteome revealed a single putative homolog for AhpC and AhpF, YPK_RS16370 (peroxiredoxin C; E-value, 3e-38) and TrxB (E-value, 3e-42), which are not in proximity on the chromosome. Combining the fact that AhpCF is encoded by a single operon in other bacteria and TrxB is clearly the counterpart of *EcTrxB* (E-value, 0) but not *EcaHpf*, these data strongly suggest that YPIII lacks a conventional AhpF. Furthermore, the YPIII genome also lacks a gene for a homolog of Ohr, the primary enzyme that decomposes OPs. Given the involvement of AhpR in scavenging OPs, thus it is particularly
TABLE 2 | Highly regulated genes involved in the response to H$_2$O$_2$ and genes encoding analogs of OxyR regulon members.

| Locus tag | Gene | OxyR regulon$^a$ | YpOxyR motif W$^b$ | Fold Change$^c$ | COG type$^d$ | Description |
|-----------|------|-----------------|----------------|----------------|----------------|-------------|
| **Top 20 up-regulated genes** | | | | | | |
| YPK_RS20590 | Y | 689.5 | O | Glutathione peroxidase |
| YPK_RS20595 | Y | 89.3 | O | Dihydrolipoyl dehydrogenase |
| YPK_RS16780 | Y | 75.4 | J | Thioredoxin TrxC |
| YPK_RS08095 | Y | 15.4 | P | Iron-sequestringer protein Dps |
| YPK_RS17250 | Y | 38.5 | P | Adenylyl-sulfate kinase |
| YPK_RS22535 | Y | 34.0 | P | Hypothetical protein |
| YPK_RS17260 | Y | 32.0 | P | Sulfate adenylytransferase subunit CysD |
| YPK_RS17255 | Y | 28.3 | P | Sulfate adenylytransferase subunit CysN |
| YPK_RS14285 | Y | 22.7 | P | Catalase, OxyR regulon |
| YPK_RS13675 | Y | 12.6 | S | GnuA family glutaredoxin |
| YPK_RS17265 | Y | 18.0 | C | Uroporphyrinogen-III C-methyltransferase |
| YPK_RS00550 | Y | 16.2 | C | Isopenicillin N synthase family oxygene |
| YPK_RS13735 | Y | 16.2 | C | Glutathione-disulfide reductase |
| YPK_RS13730 | Y | 15.4 | P | ABC transporter substrate-binding protein |
| YPK_RS03760 | Y | 13.2 | P | TonB-dependent receptor |
| YPK_RS07100 | Y | 13.1 | P | CMD domain-containing protein |
| YPK_RS16785 | Y | 13.0 | P | Sulfate/thiosulfate ABC transporter ATP-binding protein CysA |
| YPK_RS07190 | Y | 12.7 | P | DTW domain-containing protein |
| YPK_RS10680 | Y | 12.3 | G | Glucose-6-phosphate dehydrogenase |
| YPK_RS07190 | Y | 11.9 | G | Cysteine synthase A |
| **Other OxyR regulon members** | | | | | | |
| YPK_RS17025 | Y | 23.1 | O | Catalase/peroxidase HPI |
| YPK_RS20585 | Y | 7.2 | P | DNA-binding transcriptional regulator OxyR |
| YPK_RS13445 | Y | 4.5 | O | Thioredoxin-disulfide reductase |
| YPK_RS16370 | Y | 4.5 | O | Peroxiredoxin C |
| YPK_RS09285 | Y | 3.5 | O | Fe-S cluster assembly protein |

$^a$YpOxyR regulon members in at least three bacterial genera (E. coli, S. Typhi, and Neisseria meningitidis).
$^b$Weight values of predicted YpOxyR regulon members by using regulatory sequence analysis tool (RSAT).
$^c$Fold changes in transcripts (the ratio of the H$_2$O$_2$-treated sample of the wild type to the control) revealed in transcriptomics analysis.
$^d$COG: O: posttranslational modification; J: translation, ribosomal and biogenesis; P: metabolism; S: Function unknown; C: energy production; and G: carbohydrate transport and metabolism.

important and interesting to understand the physiological role of AhpC in YPIII.

**Transcriptomics Analysis of YPIII in Response to H$_2$O$_2$ Stress**

In order to gain a comprehensive understanding of the cellular response of YPIII to H$_2$O$_2$, we performed an RNA-seq analysis to obtain gene expression changes resulting from the H$_2$O$_2$ treatment. Our early stress response studies suggest that the best concentration of H$_2$O$_2$ for transcriptomics analyses would be the dosage at which the agent arrests growth of cells but does show an evident killing effect on them (Gao et al., 2004; Jiang et al., 2014). To determine the concentration, the minimum inhibitory concentration (MIC) of H$_2$O$_2$ against the wild-type strain of YPIII was assessed. The results revealed MIC to be 4 mM, which is four times lower than that of E. coli (Figure 1A), indicating that YPIII is significantly more sensitive to H$_2$O$_2$ than E. coli. Then, impacts of H$_2$O$_2$ over a range of concentrations (under MIC) on growth and viability of actively growing cells (≈0.4 of OD$_{600}$) were examined. Upon addition of H$_2$O$_2$ at 0.2 mM or higher, growth paused immediately and resumed after lag periods that increase with the H$_2$O$_2$ concentrations (Figure 1B). Additionally, viability assays revealed that H$_2$O$_2$ at 0.5 mM or lower did not show a significant killing effect (Supplementary Figure 3). Based on these observations, for RNA-seq transcriptomics analysis, we collected cells before and 5 min after the addition of...
0.5 mM H$_2$O$_2$ as the untreated control and the treated samples, respectively.

In total, 364 genes displayed significant differences in transcription levels between the H$_2$O$_2$-treated and untreated control samples (Supplementary Table 1). Among these genes, 186 genes were up-regulated, while 178 genes were down-regulated (Figure 1C). Genes displaying significant differences in expression levels due to H$_2$O$_2$-induced oxidative stress were observed in almost every functional category, indicating that YPIII transcriptionally responds to H$_2$O$_2$ in a global scale (Supplementary Table 1). The high quality of the expression data was validated with a statistical analysis as previously described (Gao et al., 2004) and by real-time qRT-PCR. Eight genes were selected for analysis with the same RNA samples used in the RNA-seq based on the level and reproducibility of changes observed in the RNA-seq experiments. A high level of concordance ($R^2 = 0.96$) was observed between RNA-seq and real-time qRT-PCR data despite quantitative differences in the level of change (Supplementary Figure 4), suggesting that the RNA-seq results are an accurate reflection of the gene expression profile. The highly induced included many genes encoding proteins combating oxidative stress. In contrast, genes encoding metabolic enzymes were down-regulated in general (Figure 1D), an observation that reflects a paused/reduced growth rate rather than a specific response to the stress has been commonly found in other stress response studies (Gao et al., 2004; Jiang et al., 2014). Intriguingly, many ABC transporters and sulfur metabolic enzymes are among differentially expressed genes, implicating that sulfur species and certain small molecules may have a particular significance in the oxidative stress response of YPIII.

To stay focused, here we only discussed genes whose transcription was significantly induced upon the addition of H$_2$O$_2$, especially those specific to H$_2$O$_2$-induced oxidative stress (Table 2). With respect to OxyR regulon members, our transcriptomics data revealed that YPK_RS20590 (glutathione peroxidase), trxC, dps, katE, grxA, and gorA were among the top 20 most up-regulated genes in H$_2$O$_2$-treated cells (Table 2). Other putative OxyR regulon members that were induced but not among the top 10 included katG, trxB, ahpC, and sufABC (Fe-S cluster assembly proteins; Table 2). Intriguingly, the entire thioredoxin and glutathione antioxidant systems, which play a supporting role in combating oxidative stress in many bacteria (Feng et al., 2019), were found to be highly induced by H$_2$O$_2$ (for instance, YPK_RS20590 was induced 689-fold), implying that these systems may have more significant contribution in protecting YPIII cells from oxidative damage. It is also worth mentioning that the oxyR gene was transcribed at substantially elevated levels (4.5-fold) upon H$_2$O$_2$ stress (Table 2), suggesting that the regulator per se is also subjected to quantity control, in addition to activity transformation upon oxidation.
Among the remaining top 20 most significantly induced genes, those for sulfur species transport and metabolism (cysC, cysD, cysN, cysA, and cysK) drew special attention. In many organisms, sulfur species are critically involved in cell protection against oxidative damage (Wu et al., 2015; Peng et al., 2017; Korshunov et al., 2020). Along with highly induced transcription of cysI, cysW, cysT; and cysJ in H₂O₂-challenged cells, these data suggest that sulfur species may be critical for oxidative stress response in YPIII. In addition, we also noticed that genes encoding iron/manganese ABC transport system (yfeA, yfeB, yfeC, and yfeD) were transcribed at significantly elevated levels upon the treatment. This may not be surprising given that the intracellular Mn/Fe ratios are correlated with bacterial resistance to oxidative stress (Chandrangsu et al., 2017). To maintain the activity of vulnerable mononuclear iron proteins in the presence of H₂O₂, many bacteria replace the iron atom with manganese as the active center (Anjem et al., 2009; Imlay et al., 2019). These data concur well with the result from the Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analysis in which sulfur metabolic pathways are among the most enriched (Figure 1D). Together with the missing of AhpF, these results suggest that the mechanism underlying the response of YPIII to H₂O₂-induced oxidative stress carries novel characteristics that are different from the classical mode established in E. coli.

Characterization of YPIII oxyR Mutant
In many bacteria, OxyR-binding motifs have been determined. Given sequence and function conservation, it is not surprising that these motifs are highly similar (Wan et al., 2018). To predict the operons under the direct control of YpOxyR, we constructed a matrix for the OxyR-binding motif with verified OxyR-binding sequences from closely related bacterial species and used it to screen the entire genome of YPIII by RSAT (Medina-Rivera et al., 2015). In total, 38 putative YpOxyR-binding sites were identified with weight values greater than 7, an arbitrary cutoff implemented in RSAT (Supplementary Table 2). On the top of the list are well-established OxyR regulon members, including katG, katE, ahpC, trxB, grxA, dps, and trxC, supporting that YpOxyR recognizes a similar DNA motif (Figure 2A). For the remaining genes on the list, we supposed that many of them may not belong to the YpOxyR regulon because weight values of
their binding motifs are relatively low (<10). But this requires further verification.

To determine the impacts of OxyR on oxidative stress response, an oxyR in-frame deletion strain (ΔoxyR) was constructed and characterized in YPIII. In liquid LB, ΔoxyR grew indistinguishably from the wild type (Supplementary Figure 5), indicating that YpOxyR is not required for normal growth. We then assessed the role of YpOxyR in combating oxidative stress. Disk diffusion assays demonstrated that ΔoxyR was more sensitive to H₂O₂, generating inhibition zones that were substantially larger than those of the wild type (Figure 2B). The phenotype can be confidently attributed to the loss of OxyR given successful genetic complementation. To provide further support to this, we compared the mutant and the wild type with respect to H₂O₂ consumption. Cells at the mid-exponential phase were collected, adjusted to identical densities, and incubated with 0.2 mM H₂O₂. As shown in Figure 2C, the ΔoxyR strain degraded H₂O₂ at a rate much lower than the wild type did; 14% and 49% of H₂O₂ remained 2 min after the reaction started for the wild type and the mutant, respectively. These results indicate that the oxyR deletion greatly impaired the ability of YPIII to remove exogenous H₂O₂. Furthermore, we also confirmed that the ΔoxyR strain carried a plating defect on LB agar plates (Figure 2D), a common phenotype of the oxyR mutants in bacteria such as E. coli and S. oneidensis because of compromised H₂O₂-scavenging capacity (Jiang et al., 2014). As exogenous catalase completely suppressed the plate defect of the ΔoxyR strain, it is reasonable to propose that the same mechanism is responsible for the YPIII oxyR mutant (Jiang et al., 2014; Shi et al., 2015).

In many bacteria lacking Ohr, such as E. coli and S. Typhi, OxyR also mediates cellular response to OPs (Imlay, 2015). Clearly, this is also true in the case of YPIII because the oxyR mutant exhibited significantly increased sensitivity to t-BHP, a widely used representative OP (Figure 2E). All together, these data conclude that YPIII employs transcriptional regulator OxyR to mediate an oxidative stress response.

**KatE Is the Primary Catalase in YPIII Positively Regulated by OxyR**

In general, multiple catalases are encoded in a bacterial genome, but only one exhibits predominant H₂O₂-scavenging activity in vivo (Mishra and Imlay, 2012). To test whether this also holds true in YPIII, we assessed the contribution of catalases to decomposition of H₂O₂. Strains lacking katE and katG were generated and characterized with the disk diffusion assay and the H₂O₂-decomposing assay. As shown in Figure 3A, the katE mutant (ΔkatE) was hypersensitive to H₂O₂, generating inhibition zones that were approximately two and a half times larger than those of the wild type. On the contrary, the difference in zone sizes between the wild type and ΔkatG was insignificant. Consistent with the results in disk diffusion assay, the katE deletion resulted in severely impaired ability to decompose H₂O₂, with nearly 60% of H₂O₂ remaining 2 min after the reaction started (Figure 3B). In the case of ΔkatG, the assay revealed that this mutation slightly but still significantly compromised the ability of YPIII to decompose H₂O₂ when compared with that of ΔkatE, validating that KatG is also functioning in YPIII.

Impacts of YpOxyR on the expression of some of its regulon members predicted by the in silico analysis, including four high-confident (weight value > 12; trxB, dps, katE, and grxA) and two low-confident genes (weight value ≈9; acnA and zwf; Table 2 and Supplementary Table 2), were assessed by qRT-PCR. In cells of ΔoxyR prepared the same as for transcriptomics analysis, we found that none of the four high-confident genes were responsive to the H₂O₂ treatment, whereas two control genes, yfeC and tauA, which lack a predicted OxyR-binding motif, were upregulated upon exposure to H₂O₂ as in the wild type (Figure 3C). In the case of acnA and zwf, however, we found that the loss of OxyR negatively affected but not completely abolished their transcription in response to H₂O₂ (Figure 3C). Apparently, these data support that the predicted DNA motifs largely determine OxyR-dependent transcription.

Given the central role of catalases in protecting cells from H₂O₂ damage, we further compared the expression of katE and katG genes in the wild type and ΔoxyR strains by measuring activities of katE and katG promoters with an integrative lacZ reporter system (Figure 3D). In line with the transcriptomics data, the expression of katE and katG increased nearly six-fold in the wild-type cells when challenged by H₂O₂. While in the ΔoxyR strain, both genes were no longer responsive to H₂O₂, being expressed at levels even lower than those observed in the untreated wild-type cells. In summary, these data validate that expression of both KatE and KatG is under the positive control of OxyR, and KatE functions as the primary catalase to cope with H₂O₂ stress.

**AhpC Possesses Scavenging Activity Against H₂O₂ but Not Organic Peroxides**

The genomics analysis as presented above indicates that YPIII possesses an atypical AhpR, whose AhpF is missing. Like katE and katG, the ahpC gene is positively regulated by OxyR (Figure 3C), and therefore, the protein is likely produced when OxyR is activated. Given the importance of AhpR in scavenging both H₂O₂ and OPs, we reasoned that AhpC may be still functional in YPIII because it can also be reduced by TrxB and/or glutathione (GSH) reductase (Gor), albeit low in efficiency (Feng et al., 2020). To test this, we knocked out the ahpC gene from mutants lacking KatE, KatG, or both. The removal of AhpC alone had no detectable impact on H₂O₂ consumption (Figure 4A), an expected result because of the dominance of catalase in H₂O₂ degradation (Mishra and Imlay, 2012). In the absence of KatE, the effect of the AhpC loss became significant, indicating that AhpC does have some capacity of H₂O₂ degradation. Compared to ΔkatEΔahpC, the loss of both catalases (ΔkatEΔkatG) significantly impaired the ability of YPIII to decompose H₂O₂ (Figure 4A). Furthermore, additional removal of AhpC from ΔkatEΔkatG nearly completely abolished the H₂O₂-decomposing capacity of YPIII. Therefore, these data conclude that all of KatE, KatG, and AhpC are capable of decomposing H₂O₂ in YPIII. Intriguingly, we found that
the strain lacking both KatG and AhpC had stronger H$_2$O$_2$-decomposing capacity than the wild type. All of these data were verified by genetic complementation, in which one of the missing genes was expressed in trans in deletion mutants (Supplementary Figure 6).

To provide additional evidence, we then examined if the removal of catalases and/or AhpC would cause YPIII a plating defect, as this is also the case with S. oneidensis (Shi et al., 2015). Similar to an S. oneidensis katB (encoding dictating catalase) mutant, the ΔkatE strain of YPIII exhibited a plating defect, albeit much less severe (Figure 4B). In contrast, the removal of either katG or ahpc did not introduce a significant defect, supporting that KatE alone is sufficient to protect cells from the killing of H$_2$O$_2$ generated spontaneously on the plates (Shi et al., 2015). Compared to the ΔkatE strain, the additional removal of either katG or ahpc only marginally deteriorated the defect (Figure 4B). However, the triple mutant ΔkatE ΔkatG Δahpc exhibited significantly further lowered viability, resembling the scenario observed with the S. oneidensis ΔkatB Δahpc strain. Given that S. oneidensis AhpC alone is responsible for cleaning endogenous H$_2$O$_2$, these data suggest that YPIII would require both KatG and AhpC to be a functional equivalence.
AhpC has been identified as a primary OP scavenger in many bacteria lacking Ohr, such as *E. coli*, *B. subtilis*, and *S. Typhi* (Niihara et al., 1995; Antelmann et al., 1996). We therefore predicted that AhpC is likely involved in combating against OPs in YPIII. To test this, we compared the abilities of the wild type and strains lacking catalase and/or AhpC to scavenge t-BHP. Results revealed that the removal of any of the genes under test (*katE*, *katG*, and *ahpC*) or all together did not introduce a significant difference in YPIII susceptibility to t-BHP (Supplementary Figure 7A). Quantification of the remaining t-BHP in the reaction also showed that the t-BHP-scavenging capacity of YPIII was not affected by any of these mutations or combined (Supplementary Figure 7B). In summary, these data suggest that AhpC in YPIII is able to decompose H$_2$O$_2$ but not OPs.

**Loss of AhpC and KatG Together Activates OxyR**

The data presented thus far have concluded that in YPIII, KatE is the primary catalase in combating H$_2$O$_2$-induced oxidative stress, whereas both KatG and AhpC contribute. It has been established that catalase functions to scavenge high concentrations of H$_2$O$_2$, whereas AhpC is a more kinetically efficient scavenger of trace H$_2$O$_2$ (Seaver and Imlay, 2001). To determine whether the YPIII H$_2$O$_2$-scavenging enzymes act in a similar manner, we measured the rates at which cells decomposed low (2 μM) and high (150 μM) concentrations of H$_2$O$_2$. The result showed that the *katE* mutant scavenge 2 μM H$_2$O$_2$ significantly more rapidly than ΔkatG and ΔahpC strains did (Figure 5A), indicating that in wild type, both of KatG and AhpC carry out the decomposition of low-dose H$_2$O$_2$ mostly. Interestingly, a ΔkatGΔahpC strain exhibited a faster rate in reducing 2 μM H$_2$O$_2$ than strains without either of these two proteins. Given that the additional removal of KatE totally disabled the ΔkatGΔahpC strain, this observation implies that the loss of KatG and AhpC induces the expression of KatE. On the contrary, with 150 μM H$_2$O$_2$, the *katE* mutant worked very poorly (Figure 5B), an expected result from a strain without the primary catalase. While both ΔkatG and ΔahpC strains showed normal scavenging activity, the ΔkatGΔahpC strain decomposed 150 μM H$_2$O$_2$ nearly two times faster than did the wild type, supporting an increased production of KatE.

In *E. coli*, the removal of AhpC results in H$_2$O$_2$ accumulation, leading to activation of OxyR and thereby increased catalase production (Seaver and Imlay, 2001). Although this is apparently not the case in YPIII, the data presented above imply that the loss of both KatG and AhpC is likely able to stimulate the production of KatE. To test this, we assessed the effects of depletion of KatG, AhpC, and both on KatE expression. As shown in Figure 5C, the *katE* gene was expressed undistinguishably in the wild-type, ΔahpC, and ΔkatG strains, and the same scenario was observed in H$_2$O$_2$-treated cells. These observations indicated that the loss of either AhpC or KatG alone does not significantly influence KatE production, eliminating the possibility that neither its absence is sufficient to activate OxyR in YPIII. Conversely, in the ΔkatGΔahpC cells grown under normal aerobic conditions, we found that the *katE* promoter activity increased 3.7 times (Figure 5C). Despite this, activity of the *katE* promoter can be elevated further under the treatment of H$_2$O$_2$ (Figure 5C), implying that the oxidative stress risen from the loss of KatG and AhpC probably activates only a portion of OxyR molecules, which is consistent with our earlier proposal that the ratio between oxidized and reduced OxyR is in a dynamic equilibrium (Wan et al., 2018, Wan et al., 2019).

**DISCUSSION**

Resistance to oxidative stress resulting from ubiquitous ROS generated both endogenously and exogenously belongs to key virulent factors of bacterial pathogens. To deal with H$_2$O$_2$-induced oxidative stress, Gram-negative bacteria widely employ OxyR to sense the oxidant and mediate transcription of their regulon in a concerted manner (Imlay, 2013). Although regarded as a pleiotropic regulator, OxyR has evolved to consistently possess operons that are involved in H$_2$O$_2$ decomposition and damage control as the core member of regulons (Imlay, 2015). This is seemingly the case in *Y. pseudotuberculosis* YPIII. Our result in Supplementary Table 2 predicted that all H$_2$O$_2$-scavenging enzymes, iron-sequestering proteins, and thioredoxin and glutathione antioxidant systems are encoded by operons whose promoter regions contain most conserved OxyR-binding motifs with the highest similarities. OxyR of YPIII, highly similar to its *E. coli* counterpart in terms of sequence identity (88%) and the almost identical binding motifs, exhibits stimulating activity only (Zheng et al., 1998). This gains support from our transcriptomics data, which reveal elevated transcription of all predicted OxyR regulon members in the H$_2$O$_2$-treated cells. In addition to OxyR regulon members, genes for sulfur species transport and metabolism and iron–manganese transport system are among the top up-regulated in the H$_2$O$_2$-treated cells. Sulfur species, especially hydrogen sulfide (H$_2$S) and cysteine, are well recognized as important factors in bacterial oxidative stress response. These reductants have been shown to play a critical role in the detoxification of H$_2$O$_2$ in the periplasm (Ohtsu et al., 2010; Ezraty et al., 2017). Intracellular sulfur homeostasis should be carefully maintained because H$_2$S and cysteine in vast excess promote oxidative damages by inhibiting catalases (Park and Imlay, 2003; Wu et al., 2015). Consistently, all components of thiol-based antioxidant systems of YPIII are most highly induced upon exposure to H$_2$O$_2$, conceivably requiring an active metabolism and fast shuttling of sulfur species (Feng et al., 2019). In parallel, genes for iron/manganese transport system are highly induced by exogenous H$_2$O$_2$. In bacteria, many enzymes using iron as a cofactor may become inactive due to the loss of the metal upon oxidative stress (Anjem et al., 2009; Anjem and Imlay, 2012). A common way to overcome this is to replace iron with manganese, and hence more manganese is imported when cells are challenged by H$_2$O$_2$ (Imlay et al., 2019). Although further investigations are needed, all of these observations suggest that sulfur species and manganese are important for YPIII to combat oxidative stress.
Simultaneous loss of AhpC and KatG increases H$_2$O$_2$ resistance. Efficiencies of AhpC, KatE, and KatG at different H$_2$O$_2$ concentrations. H$_2$O$_2$ was added at a final concentration of 2 µM (A) and 150 µM (B) to cultures of YPIII strains indicated. 2 min after addition of H$_2$O$_2$, the H$_2$O$_2$ concentration was measured. (C) KatE expression in strains indicated before and after the H$_2$O$_2$ treatment. In all panels, asterisks indicate statistically significant differences of the values compared (n = 4; ns, not significant; *p < 0.05; **p < 0.01; and ***p < 0.001).

Like *E. coli*, YPIII contains two catalases: HPII KatE and HPI KatG (Mishra and Imlay, 2012). Although both KatE and KatG are highly induced upon H$_2$O$_2$ treatment, we first anticipate that YPIII uses KatG as the predominant H$_2$O$_2$-scavenging enzyme because *E. coli* does so (Seaver and Imlay, 2001). Unexpectedly, it is KatE that confers the H$_2$O$_2$-decomposing ability at high concentrations in YPIII. Instead, KatG exhibits a much minor but still significant contribution in scavenging high-dose H$_2$O$_2$. More importantly, KatG shows a strong scavenging activity toward low concentrations of H$_2$O$_2$, a role that is exclusively played by AhpC in *E. coli*. Despite this, given that *Yp*OxyR becomes activated in the absence of both AhpC and KatG, but not of either one, it is clear that both KatG and AhpC, while neither is sufficient, is able to scavenge low concentrations of H$_2$O$_2$ in YPIII. It is well established that AhpC allows *E. coli* cells to degrade low concentrations of H$_2$O$_2$ because of its high kinetic efficiency ($k_{cat}/K_M$, ∼$10^{-8}$ M$^{-1}$ s$^{-1}$; Mishra and Imlay, 2012). We believe that this is also likely true in the case of YPIII KatG. But the result of impaired H$_2$O$_2$-scavenging ability of $\Delta$katG and $\Delta$ahpC at low levels in YPIII implies that YPIII KatG possesses a kinetic efficiency similar to, or at least not significantly lower than, that of AhpC, the catalytic activity that is substantially greater than those of KatE and KatG in *E. coli* whose activities are similar to each other (∼$10^{-6}$ M$^{-1}$ s$^{-1}$; Obinger et al., 1997; Hillar et al., 2000). We are working to test this notion.
Bacterial AhpC is regarded as the founding member of the peroxiredoxin family, and its homologs appear to be more widely distributed than catalase/peroxidase (Chae et al., 1994). The AhpR complex is highly conserved among diverse bacteria, with respect to not only amino acid sequence but also gene organization (Feng et al., 2020). To date, all bacteria that employ OxyR as the master H<sub>2</sub>O<sub>2</sub>-responding regulator are equipped with an ahp operon encoding both AhpC and its cognate reductase AhpF, including <i>M. tuberculosis</i> in which AhpF is replaced by AhpD to reduce AhpC (Baker et al., 2001; Jaeger et al., 2004). Apart from that, AhpC without a co-transcribed cognate reductase is found in <i>H. pylori</i>, a gastric pathogen that represents an extreme example in terms of oxidative stress response machinery because it lacks homologs of the oxidative stress regulators present in other bacteria, including OxyR, SoxR, SoxS, and PerR (Baker et al., 2001; Wang et al., 2006). A sequence alignment reveals that AhpC of <i>Y. pseudotuberculosis</i> is more closely related to typical AhpCs than to the <i>H. pylori</i> AhpC: there are significantly more identical residues in AhpCs of <i>E. coli</i>, <i>S. oneidensis</i>, and <i[Y. pseudotuberculosis</i> (<i>Supplementary Figure 8</i>). Despite this, we also observed that the linear arrangement of AhpC gene is highly conserved in <i>Yersinia</i> species (<i>Supplementary Figure 9</i>), a phenomenon that is quite rare in other bacteria. Therefore, it would be interesting to demystify how evolution has honed this particular phenomenon.

Although the reductase for YPIII AhpC remains to be determined, we may get clues from <i>H. pylori</i> AhpC in terms of their differences in sequence and structure from the typical ones (with AhpF as the cognate reductase). Reduction of <i>H. pylori</i> AhpC is carried out by thioredoxin TrxA, which is reduced by thioredoxin reductase TrxB (also called TrxR) using NADPH as the electron donor (Wang et al., 2006). A common feature that <i>Yp</i>AhpC and <iHp</i>AhpC share is that they have an additional eight-residue segment in the C-terminal end, which is missing in the typical AhpC proteins (<i>Figure 6A</i> and <i>Supplementary Figure 8</i>). In fact, the last few residues in the C terminus of AhpC have been shown to be particularly critical in maintaining an enzymatically active AhpR complex in <i>vitro</i> (Dip et al., 2014a). This C-terminal tail of AhpC is crucial in complexation of AhpC with AhpF, TrxR, or glutathione-disulfide reductase GorA (Dip et al., 2014a; Feng et al., 2020). Coincidently, earlier studies of certain <i>E. coli</i> and <i>S. oneidensis</i> AhpC mutants have established that certain mutations, especially in the C-terminal region, can transform AhpC to a surprisingly malleable enzyme, reducible not only by Trx system but also by Grx system (Faulkner et al., 2008; Feng et al., 2020). Therefore, we speculate that the extra eight residues at the C-terminal end of YPIII AhpC may affect the specificity of the protein for its reducing partner.

The difference in the C-terminal region may also provide an explanation for the finding that YPIII AhpC does not have the ability to reduce OPs. It has been shown that catalytic efficiency of AhpC with small hydroperoxides, especially H<sub>2</sub>O<sub>2</sub>, is substantially higher (~100-fold) than that with bulky, tertiary hydroperoxides, such as t-BHP (Parsonage et al., 2008). For both types of oxidants, the catalytic reaction adopts a binding and releasing mechanism that enables the assembly of AhpC and the reducing partner to undergo efficient catalytic cycles of transferring electrons without compromising the catalytic turnover rate (Parsonage et al., 2015). A prerequisite for such a mechanism is the correct fitting of the C-terminal tail of AhpC into the “arch-like” groove on the binding surface of the reducing partner, resulting in formation of an active-site pocket, to which small peroxides are more accessible (Dip et al., 2014b). The additional C-terminal tail of YPIII AhpC may completely prevent large OPs from entering the active-site pocket (<i>Figure 6B</i>). Compared to the <i>E. coli</i> AhpC structure (PDB accession number 4o5r; Dip et al., 2014b), <i>Y. pseudotuberculosis</i> AhpC has an α-helix that is formed by the extra residues. Unfortunately, how this α-helix folds remains unknown because the available <i>H. pylori</i> AhpC structure is truncated without this tail (PDB accession number 1zo; Papinutto et al., 2005; <i>Supplementary Figure 10</i>). But one may imagine that the extra C-terminus α-helix may hinder the reduction of OPs. Given that YpAhpC, unlike typical AhpCs, is not sufficient to scavenge endogenous H<sub>2</sub>O<sub>2</sub>, we speculate that the α-helix also compromises the efficiency of H<sub>2</sub>O<sub>2</sub> reduction. If this holds, the difference in the kinetic efficiencies for H<sub>2</sub>O<sub>2</sub> between AhpC and catalases of YPIII is likely significantly smaller than that between the <i>E. coli</i> counterparts.

### DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

### AUTHOR CONTRIBUTIONS

FW and HG designed and supported the research. FW, XF, and JY performed the research and analyzed the data. FW, XF, and HG wrote the manuscript. All authors contributed to the article and approved the submitted version.

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### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.626874/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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