OxyR Activation in *Porphyromonas gingivalis* in Response to a Hemin-Limited Environment

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*Porphyromonas gingivalis* is a Gram-negative obligately anaerobic bacterium associated with several forms of periodontal disease, most closely with chronic periodontitis. Previous studies demonstrated that OxyR plays an important role in the aerotolerance of *P. gingivalis* by upregulating the expression of oxidative-stress genes. Increases in oxygen tension and in H$_2$O$_2$, both induce activation of OxyR. It is also known that *P. gingivalis* requires hemin as an iron source for its growth. In this study, we found that a hemin-limited growth environment significantly enhanced OxyR activity in *P. gingivalis*. As a result, expression of *sod, dps*, and *ahpC* was also upregulated. Using a chromatin immunoprecipitation quantitative PCR (qPCR) analysis, DNA binding of activated OxyR to the promoter of the *sod* gene was enhanced in *P. gingivalis* grown under hemin-limited conditions compared to excess-hemin conditions. Cellular tolerance of H$_2$O$_2$ was also enhanced when hemin was limited in the growth medium of *P. gingivalis*. Our work supports a model in which hemin serves as a signal for the regulation of OxyR activity and indicates that *P. gingivalis* coordinately regulates expression of oxidative-stress-related genes by this hemin concentration-dependent pathway.

Many organisms, including microbial cells, require iron for various metabolic processes, such as electron transport, glycolysis, and DNA synthesis (22, 27). Like other organisms, *P. gingivalis*, a well-known periodontal pathogen associated with several forms of periodontitis, requires iron for its growth (5, 19, 32). *In vitro*, *P. gingivalis* can grow well only in media supplemented with hemin, the oxidized form of ferrous iron bound to tetrapyrole. This bacterium possesses a complex system for iron acquisition, and at least 50 proteins of *P. gingivalis* are likely involved directly or indirectly in iron utilization (16, 19). One of the characterized virulence features of *P. gingivalis* is its ability to accumulate hemin on its surface. It appears that its hemin binding ability is higher when it grows under excess-hemin conditions than under hemin-limited conditions (24).

Virulence gene expression in *P. gingivalis* is regulated by several environmental cues, including temperature, the presence of other bacteria, and iron/hemin concentrations (30). We reported earlier that expression of *fimA*, a gene encoding a major subunit protein of long fimbriae, is repressed in *P. gingivalis* grown in hemin-limited media (30). Previous studies, using proteomic and transcriptomic analyses to identify differentially expressed proteins and genes, showed that expression of 70 proteins and 160 genes was significantly altered when *P. gingivalis* was grown under hemin-limited conditions compared to excess-hemin conditions (6). Another recent study indicated that at least 3% of the genes in the *P. gingivalis* genome were modulated in response to a change in hemin concentration (11). These differentially expressed genes or proteins were linked to bacterial invasion, iron transport, and an oxidative-stress response. A recent study showed that inorganic polyphosphates (potential antibacterial agents) inhibited energy-driven uptake of hemin by *P. gingivalis* and repressed the expression of oxidative-stress-induced proteins, including superoxide dismutase (SOD) (15), which provides indirect evidence linking hemin to SOD expression. To better understand the response of *P. gingivalis* to hemin-limited conditions and the association between iron/hemin concentrations and oxidative stress, we investigated the expression and activity of OxyR in *P. gingivalis* under hemin-limited as well as excess-hemin conditions.

OxyR represents the first redox-regulatory protein that was well characterized in *Escherichia coli* (8). OxyR is a tetrameric DNA-binding protein that is activated under oxidative stress by forming a disulfide bond between two Cys residues that do not seem to form covalently cross-linked multimers, suggesting that the Cys199-Cys208 disulfide bond is intramolecular (21, 33). The disulfide bonds are then rereduced by thioredoxin or glutathione to form inactivated OxyR, suggesting that Cys disulfide bonds are crucial for OxyR activation. The oxyR gene is widely distributed in most Gram-negative and some Gram-positive bacteria (14), and OxyR homologs have been identified in more than a dozen other bacterial species, including *P. gingivalis*. *P. gingivalis* OxyR is a 308-amino-acid protein containing 5 cysteines (Cys24, Cys131, Cys171, Cys198, and Cys207). The two Cys residues at the C-terminal end have positions similar to those of the two Cys residues of *E. coli* OxyR but shifted by one amino acid. Previous studies from different laboratories indicated that OxyR is required for activation of the oxidative stress-related genes of *P. gingivalis*, such as sod and ahpC, encoding alkyldydroperoxide reductase (7, 13, 18). In this study, we report that activation of OxyR occurs when the hemin concentration in the growth environment of *P. gingivalis* is limited and that this OxyR activation leads to alteration of expression of some genes in the OxyR regulon. Our results also demonstrated that the activated OxyR induced by hemin-limited conditions has a higher promoter binding activity. Equally impor-

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tantly, we were able to show that the ability of *P. gingivalis* to survive under oxidative stress was greatly enhanced when the bacteria were grown under hemin-limited conditions.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** Bacterial strains used in this study are listed in Table 1. *P. gingivalis* strains were grown from frozen stocks in Trypticase soy broth (TSB) or on TSB blood agar plates supplemented with yeast extract (1 mg/ml), hemin (5 μg/ml), and menadione (1 μg/ml) and incubated at 37°C in an anaerobic chamber (85% N₂, 10% H₂, 5% CO₂). Antibiotics were used when appropriate, at the following concentrations: gentamicin (100 μg/ml) and erythromycin (5 μg/ml). For growth of *P. gingivalis* under hemin-limited conditions, the bacteria were first grown in standard TSB supplemented with hemin to reach an optical density at 600 nm (OD₆₀₀) of 0.8 (mid-exponential phase). *P. gingivalis* cells were then subcultured in TSB without hemin supplement for another three passages; cells collected from the third passage were considered to have been grown under hemin-limited conditions.

**RNA isolation and qPCR.** *P. gingivalis* strains were grown anaerobically in 5 ml of TSB. Bacteria were harvested by centrifugation at 10,000 × g and homogenized in TRIzol reagent (Invitrogen, Carlsbad, CA). The RNA in the supernatant was then purified using an RNeasy mini-spin column (Qiagen, Valencia, CA). RNA samples were digested on the column with RNase-free DNase. The total RNA was tested using an Agilent 2100 Bioanalyzer to ensure the quality of the samples. Real-time reverse transcription-PCR (RT-PCR) analysis was performed by using a QuantiTect SYBR green RT-PCR kit (Qiagen) on an iCycler MyiQ real-time PCR detection system (Bio-Rad Laboratories, Inc., Hercules, CA) according to the manufacturer’s instructions. Primers are listed in Table 2. Amplification reactions consisted of a reverse transcription cycle at 50°C for 30 min, an initial activation at 95°C for 15 min, and 40 cycles of 94°C for 15 s, 58°C for 30 s, and 72°C for 30 s. The melting curve profile was analyzed to verify a single peak for each sample, which indicated primer specificity. The expression levels of the investigated genes for the test sample were determined relative to the untreated calibrator sample by using the comparative cycle threshold (ΔCₜ) method. The ΔCₜ values were calculated by

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<td><em>P. gingivalis</em> strain</td>
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<td>33277</td>
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<td>Pgn0368E</td>
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<td>Pgn1373E</td>
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*Emr*, resistance to erythromycin.

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<th>TABLE 2 Oligonucleotide primers used in this study</th>
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*The sequences corresponding to those of the erythromycin resistance (Emr') gene are underlined.*
subtracting the average $C_T$ value of the test sample from the average $C_T$ value of the calibrator sample and were then used to calculate the ratio between the two by assuming 100% amplification efficiency. Finally, expression levels of testing genes were normalized by expression of a control gene (glk).

**Construction of pgn0373 (trx) mutants.** An insertional trx (pgn0373) mutant was generated by using an overlap extension PCR method (20). A 2.1-kb *ermF-ermAM* cassette was introduced into the trx gene by three steps of PCR to yield a *trx-erm-trx* DNA fragment as described previously (29). Specific primers were used listed in Table 2. The final PCR products were then introduced into *P. gingivalis* 33277 by electroporation. The *trx*-deficient mutant was generated via a double-crossover event that replaced *trx* with the *trx-erm-trx* DNA fragment into the 33277 chromosome. The mutants were selected on TSB plates containing erythromycin (5 μg/ml). The insertional mutation was confirmed by PCR analysis, and the mutants were designated *P. gingivalis* 0373E.

**Chromatin immunoprecipitation (ChIP) qPCR assay.** ChIP assays were conducted as previously described (17, 28). Briefly, formaldehyde (final concentration, 1%) was added to 20 ml of a *P. gingivalis* 33277 culture. The cross-linking reaction was stopped by the addition of glycine (125 mM, final concentration). Cells were resuspended in 2 ml lysis buffer (20 mM Tris-HCl [pH 8.0], 10 mM EDTA, 0.5 mg/ml TLCK [Nα-p-tosyl-L-lysine chloromethyl ketone], 10 mg/ml lysozyme) for 10 min at room temperature, followed by the addition of an equal volume of 2X immunoprecipitation (IP) buffer (0.1 M Tris-HCl [pH 7.0], 0.3 M NaCl, 2% Triton X-100, 0.2% sodium deoxycholate). Cells were sonicated to fragment chromosomal DNA; these samples were used as the input fraction for the ChIP assay.

Anti-OxyR antibodies (25 μl of the pooled sera) were added to the input fraction, and the mixture was rotated overnight at 4°C, complexes were then incubated for 1 h with a 20-μl bed volume of pre-equilibrated protein A Sepharose CL-4B beads (Sigma, St. Louis, MO). DNA samples were precipitated by ethanol and resuspended in 30 μl distilled H2O.

Real-time PCR was performed by using a Quantitect SYBR green PCR kit (Qiagen) on an iCycler MyIQ real-time PCR detection system (Bio-Rad Laboratories, Inc.), according to the manufacturer's instructions. Primers were designed to amplify the promoter regions of *sod* and *pgn0373* (29). Primers are listed in Table 2. Amplification reactions consisted of an initial activation at 95°C for 15 min and 40 cycles of 94°C for 15 s, 58°C for 30 s, and 72°C for 30 s. Enrichment values (fold) were calculated according to the output/input ratio relative to that of a housekeeping gene, *gkh*, encoding glucokinase.

**Bacterial survival assay.** Bacterial resistance to hydrogen peroxide (H2O2) was determined using a bacterial survival assay as previously described (7). Briefly, 200 μl of an overnight culture of *P. gingivalis* 33277 was inoculated into 10 ml of TSB medium. Hydrogen peroxide (H2O2; 0 , 400 μM) (Sigma) was added to the cultures at 8 h after the initial inoculation. Growth curves over a 48-h period were determined by measuring optical densities (OD600) in 200-μl samples at 8- to 16-h intervals. All experiments were performed in triplicate. The doubling time of bacterial growth was calculated using the formula $T_2 = T_1 \ln 2 / (\ln OD_2 - \ln OD_1)$, where $T_1$ and $T_2$ are starting and ending times and OD1 and OD2 are optical densities at these two times. The length of the lag phase was determined by monitoring the time required for bacteria to reach an OD600 of 0.2.

**Enzyme-linked immunosorbent assay (ELISA).** ELISA was performed according to our previous description, but with some modifications (31). Briefly, 96-well microtiter plates (Thermo Scientific, Rochester, NY) were coated with proteins (25 μg) extracted by sonication of samples of *P. gingivalis* 33277 grown under excess-hemin or limited-hemin conditions for 16 h at 4°C. The unbound proteins were removed by washing with PBS containing 0.1% Tween 20, pH 7.4. The plates were blocked with 3% bovine serum albumin in PBS-Tween 20 for 2 h at 37°C. A series dilutions of anti-Tpx polyclonal antibodies (Thermo Scientific) or anti-OxyR antibodies (28) were then applied to the plates, which were then incubated for 3 h at room temperature. The plates were washed and incubated with horseradish peroxidase-conjugated antibodies against rabbit IgG (1:3,000; Amersham Biosciences, Piscataway, NJ) for 1 h at room temperature. After the wells were washed five times with PBS, peroxide substrate (Sigma) was added to each well. The reaction was stopped by the addition of 100 μl of 1 N H2SO4. The results were read at 450 nm on a Benchmark Plus microplate reader (Bio-Rad). All samples were assayed in triplicate for each sample.

**Statistical analyses.** Student’s t test was used to determine statistical significance of the differences in gene expression profiles and growth rates of *P. gingivalis* strains. A P value of <0.05 was considered significant.

**RESULTS**

**Activation of OxyR in *P. gingivalis* in response to hemin limitation.** To test the role of hemin in OxyR activation, *P. gingivalis* wild-type strain 33277 and the *oxyR* mutant were first grown in TSB with hemin (WH, 5 μg/ml) and then grown without supplemental hemin for three passages (HL). Finally, both strains were inoculated into TSB containing hemin for another two passages. Because of its ability to accumulate hemin on its surface, the growth rate of *P. gingivalis* with hemin was not significantly different from that seen in TSB without supplemental hemin during the first two passages. However, the third passage in TSB without hemin showed a slightly lower growth rate. There was also no significant difference in growth rates between the wild-type and the *oxyR* mutant when they were incubated under anaerobic conditions (data not shown). *P. gingivalis* cells were harvested from each passage at mid-log phase (OD600 = 0.8). The expression level of *sod*, a gene that is well known for being positively controlled by OxyR (7, 28), was determined by quantitative RT-PCR (qRT-PCR) and used as an indicator of OxyR activation. As shown in Fig. 1, expression of *sod* increased as much as 8-fold after *P. gingivalis* 33277 (wild type) was cultured for three passages under hemin-limited conditions compared to that in the cells grown with hemin. The expression of *sod* in 33277 after it had been grown for two additional passages in standard TSB declined to the baseline level seen in *P. gingivalis* grown with hemin, indicating a reversible hemin-dependent regulation of OxyR activation. In contrast, this differential expression of *sod* was not observed in the *oxyR* mutants under excess-hemin or hemin-limited conditions, suggesting an involvement of OxyR in hemin-dependent regulation of *sod* expression.

Our previous studies identified a community development and hemin regulator (CdhR) that acted as a transcriptional activator of the *hmu* locus, which encodes an iron acquisition system (4, 29). In those studies, the expression of both *cdhR* and the genes of the *hmu* loci was significantly decreased when *P. gingivalis* cells were grown at a higher cell density. To test if *CdHR* and *HmuY*, encoded by the first gene (*hmuY*) in the *hmu* locus, play a role in OxyR activation, expression of *sod* was measured in *cdhR* and *hmuY* mutants grown in the presence or absence of hemin. As shown in Fig. 2, similar to the *sod* expression pattern in wild-type strain 33277, an elevated expression of *sod* was observed in *cdhR* - and *hmuY*-deficient strains under hemin-limited conditions. These
data suggest that cell density-dependent expression of cdhR and hmuY is not essential for OxyR activation.

Expression of oxidative stress-related genes in *P. gingivalis* under hemin-limited conditions. To test the possibility that transcription of oxyR is elevated in *P. gingivalis* in response to hemin limitation, we compared mRNA levels of oxyR in *P. gingivalis* grown in standard TSB supplemented with hemin to that in hemin-limited TSB. Interestingly, expression of oxyR was not altered despite a change in hemin concentration (Fig. 3). The results demonstrate that OxyR activity but not oxyR expression was activated in *P. gingivalis* in response to hemin-limited conditions. Thioredoxins are known for their fundamental roles in different cellular processes, including OxyR activation (32a). Expression of trx genes is also known to be under the control of OxyR. Based on their DNA sequences, two genes of *P. gingivalis*, pgn0033 and pgn0373, are annotated as encoding thioredoxins (Trx), and pgn1232 is annotated as encoding a thioredoxin reductase (TrxB) (20). As shown in Fig. 3, one of the trx genes, pgn0373, was expressed at 6-fold-higher levels in *P. gingivalis* 33277 grown under hemin-limited conditions than in the same strain under excess-hemin conditions. This alteration of gene expression induced by hemin-limited conditions was not found in the oxyR mutant, indicating that *P. gingivalis* OxyR serves as a transcriptional activator of a particular trx gene (pgn0033). Although the difference is

FIG 1 Differential expression of sod in *P. gingivalis* strains. *P. gingivalis* strains were first grown in TSB medium without hemin for three generations (HL, hemin limitation) and then grown in standard TSB medium for another two generations (WH, with hemin). Expression of the sod gene in both the wild-type strain 33277 and the oxyR mutant was determined using real-time RT-PCR analysis. The change in expression levels was calculated by the \( \Delta \Delta C_T \) method, where \( \Delta C_T = C_T(\text{cells grown under standard conditions}) - C_T(\text{cells grown under hemin limitation}) \), and normalized to that of glk. Standard deviations are indicated (n = 3).

FIG 2 Comparison of sod expression in *P. gingivalis* 33277 and its mutant strains. The oxyR mutant, the cdhR mutant, the hmuY mutant, and the parent strain 33277 were grown with or without hemin for three passages. Expression of sod was measured using real-time RT-PCR. Each bar represents the increase of sod expression in *P. gingivalis* grown without hemin compared to expression in strains grown with hemin, which was normalized to the change in glk expression. Error bars represent standard deviations. Asterisks indicate a statistically significant difference in sod expression levels in *P. gingivalis* grown with and without hemin (t test; P < 0.05).
Gene expression in *P. gingivalis* in response to hemin limitation. Total RNA was extracted from *P. gingivalis* 33277 and its oxyR mutant grown with or without hemin. Expression of *pgln0033* (thioredoxin), *pgln0373* (putative thioredoxin), *pgln1232* (thioredoxin reductase), and *oxyR* was measured by real-time RT-PCR. Each bar represents the change in *P. gingivalis* strains grown without hemin compared to that in strains grown with hemin, which was normalized to the change in *glk* expression. Error bars represent standard deviations. Asterisks indicate a significant difference in gene expression in *P. gingivalis* strains grown under hemin-limited versus excess-hemin conditions (t test; *P* < 0.05).

To determine if OxyR and Trx production are modified in *P. gingivalis* grown under hemin-limited conditions, we compared protein levels of OxyR and Trx in bacteria grown in different hemin concentrations using ELISA. As shown in Fig. 4, synthesis of OxyR, consistent with its mRNA level, was not significantly different between *P. gingivalis* cultures grown under excess-hemin and hemin-limited conditions. The results confirm that OxyR activity but not *oxyR* expression was activated in *P. gingivalis* in response to hemin-limited conditions. In contrast, thioredoxin (TRX) levels in *P. gingivalis* grown under hemin-limited conditions were 58% higher than those in bacteria grown under excess-hemin conditions (Fig. 4).

**DNA binding affinity of OxyR regulated by hemin concentrations.** We previously showed that expression of *sod* by OxyR involves direct interaction of OxyR and the promoter region of the *sod* gene (28). The fact that OxyR is activated in *P. gingivalis* grown under hemin-limited conditions suggests that a modified form of OxyR may exist under this conditions, one that has a higher affinity for the promoter of the *sod* gene. To test this hypothesis, we carried out a chromatin immunoprecipitation (ChIP) qPCR assay.

![FIG 3](http://iai.asm.org/)  
**FIG 3** Gene expression in *P. gingivalis* in response to hemin limitation. Total RNA was extracted from *P. gingivalis* 33277 and its oxyR mutant grown with or without hemin. Expression of *glk*, *pgln0033* (thioredoxin), *pgln0373* (putative thioredoxin), *pgln1232* (thioredoxin reductase), and *oxyR* was measured by real-time RT-PCR. Each bar represents the change in *P. gingivalis* strains grown without hemin compared to that in strains grown with hemin, which was normalized to the change in *glk* expression. Error bars represent standard deviations. Asterisks indicate a significant difference in gene expression in *P. gingivalis* strains grown under hemin-limited versus excess-hemin conditions (t test; *P* < 0.05).

![FIG 4](http://iai.asm.org/)  
**FIG 4** Differential expression of OxyR and Trx (thioredoxin) proteins. *P. gingivalis* 33277 cells were grown under excess-hemin (WH) or hemin-limited (HL) conditions. OxyR and Trx levels were determined by ELISA and reported as the mean absorbance from three independent experiments. Error bars represent standard deviations. Asterisks indicate a statistically significant difference in protein expressions in *P. gingivalis* grown under different hemin concentrations (t test; *P* < 0.001).
to assess the interaction of OxyR and the sod promoter in vivo. P. gingivalis 33277 cells were grown under hemin-limited or excess conditions. OxyR-DNA complexes were immune-precipitated with anti-OxyR antibodies and measured using a qPCR analysis with specific primers corresponding to the promoter region of sod. As shown in Fig. 5, enrichment of the promoter region of sod in P. gingivalis grown under the hemin-limited conditions was significantly greater than that in P. gingivalis grown under excess-hemin conditions. Interaction between OxyR and the promoter of pgn2075, a gene encoding excinuclease, was tested as a control. There was no significant difference in immune precipitation of the pgn2075 promoter by OxyR antibodies. Moreover, the differential enrichment of the sod promoter was not observed in parallel samples using preimmune serum from the same rabbit (data not shown).

Role of hemin in survival ability of P. gingivalis under oxidative stresses. P. gingivalis OxyR plays a central role in the bacterial aerotolerance by activating transcription of oxidation-related genes under anaerobic conditions, including sod, dps (encoding a DNA-binding protein), and ahpC (encoding alkyl hydroperoxide reductase) (7, 28). A previous study suggested that SOD was responsible for tolerance to atmospheric oxygen but did not appear to protect P. gingivalis against hydrogen peroxide (H₂O₂) (12). On the other hand, P. gingivalis Dps was found to be essential for protection of the organism from H₂O₂ (26). Therefore, the expression of dps (pgn2037) and ahpC (pgn0060) in P. gingivalis 33277 grown with or without hemin was determined using qRT-PCR. As expected, expression of dps and ahpC was enhanced approximately 2.5- and 3.5-fold, respectively, in response to hemin limitation (Fig. 6). Similar to the change in sod expression (Fig. 1), expression of dps and ahpC was also reduced to baseline levels when P. gingivalis cells grown under hemin-limited conditions for three passages were cultured in a standard TSB medium.

Based on these observations, we speculate that P. gingivalis grown under hemin-limited conditions may have a higher tolerance to H₂O₂ due to OxyR activation. To determine the hemin concentration-dependent tolerance to H₂O₂, we compared growth curves of P. gingivalis 33277 in the absence or presence of hemin and exogenous H₂O₂. P. gingivalis was grown in TSB with or without supplemental hemin for 48 h. There was no significant difference (t test) in the lengths of the lag phase (9.25 h versus 9 h) or doubling times (4.6 h versus 4.7 h) between P. gingivalis cultures grown with and without hemin (Fig. 7). This indicates that accumulation of hemin on the surface of P. gingivalis is sufficient to support growth over a 48-h period; this is true even though consumption of the resident hemin on the surface affected gene expression of some proteins. Addition of 200 and 400 µM H₂O₂ at 8 h after initial inoculations had a significant impact on the growth rates of P. gingivalis. Thus, in the presence of H₂O₂, P. gingivalis required a significantly longer time to reach its log phase, which is in agreement with a previous report (7). More interestingly, there was a significantly increased ability to resist H₂O₂ in P. gingivalis grown under hemin-limited conditions compared to bacteria grown under excess-hemin conditions. As shown in Fig. 7, P. gingivalis grown under hemin-limited conditions reached its log phase in 10 h in the presence of 200 µM and 26 h in the presence of 400 µM H₂O₂, while P. gingivalis grown under excess-hemin conditions required 12 h in the presence of 200 µM H₂O₂ and 36 h in the presence of 400 µM H₂O₂ to reach log phase. Similar patterns were observed in the bacterial doubling times. These data demonstrate that resistance to H₂O₂ was enhanced in P. gingivalis in response to he-
min-limited conditions, which was likely due to activation of OxyR and the proteins of the OxyR regulon.

To establish a role for Trx (PGN0373) in P. gingivalis OxyR activation, sensitivities to H$_2$O$_2$ of the wild-type strain 33277, the oxyR mutant, and the trx (pgn0373) mutant were determined and compared. There was no significant difference in growth rates when all three strains were grown in standard TSB (Fig. 8). However, significant differences in sensitivities to H$_2$O$_2$ (400 μM) were found among these three strains. The oxyR mutant was not able to recover from the H$_2$O$_2$ attack at all, while 33277 recovered grad-

FIG 6 Differential expression of dps and ahpC in P. gingivalis 33277 in response to hemin concentrations. Total RNAs were extracted from P. gingivalis grown under hemin-limited (HL) conditions or those grown under hemin-limited conditions first and then cultured in standard TSB with hemin (BH). Expression of dps and ahpC in P. gingivalis grown without hemin was measured using qRT-PCR and compared to that in P. gingivalis with hemin after normalized with expression level of glk. Error bars represent standard deviations. An asterisk indicates a statistically significant difference between expression levels of genes in P. gingivalis grown without hemin and those in P. gingivalis grown in TSB with hemin (t test; P < 0.05).

FIG 7 Comparison of growth curves and tolerance to hydrogen peroxide of P. gingivalis under different growth conditions. P. gingivalis 33277 cells were grown under excess-hemin (WH) or hemin-limited (HL) conditions for 48 h. Hydrogen peroxide (HP) was added to the medium 8 h after the initial inoculation. Data are the mean optical densities (OD$_{600}$) of bacterial cultures from four experiments. Error bars represent standard deviations.
ually. The highest recovery rate was found in the trx mutant, likely due to increased OxyR activity in this mutant.

**DISCUSSION**

**Ferrous iron, hemin, and OxyR activation.** Iron is essential to most organisms, including bacteria. Iron metabolism in bacteria is associated with their ability to overcome oxidative stress, since iron serves as a cofactor of enzymes such as SOD. *P. gingivalis* cells also contain cytosolic SOD that requires both Fe and Mn for activity (12). Iron is also known as a main determinant of the cellular response to oxidative stress. Excessive free iron can generate the highly reactive and extremely damaging hydroxyl radical (2). In *E. coli*, an increased sensitivity to redox stress agents was observed when cells were grown under iron-rich conditions (1). In addition, deregulation of iron metabolism inactivated the fur gene in *E. coli* and led to an increased sensitivity of the organism to redox stress, an effect that could be reversed by either iron chelation or a tonB mutation that blocked iron uptake (25).

OxyR activity in many Gram-negative bacteria is known to be regulated under oxidative stresses, such as H$_2$O$_2$. Previous studies from independent laboratories have shown that the activity of OxyR in *P. gingivalis* was enhanced when the organism was exposed to H$_2$O$_2$ and atmospheric oxygen (7, 28). In this study, we revealed that expression of several oxidative-stress-related genes, including sod, trx, dps, and ahpC, fluctuated in *P. gingivalis* in response to a hemin-limited environment. Apparently, this hemin-dependent alteration of the expression of oxidative-stress-related genes is controlled by OxyR, since the phenomenon was not observed in the oxyR mutant (Fig. 1). Only baseline expression of the sod gene was found in the oxyR mutant grown with either excess or limited hemin. A recent study showed that iron significantly reduced expression of PerR, which, like OxyR, is a transcriptional regulator controlling transcription of oxidative-stress-related genes in *Campylobacter jejuni* (10). Most interestingly, the effect of iron on PerR transcription was unique, and other metal ions, including copper, cobalt, manganese, and zinc, were ineffective. In contrast, our data demonstrate that expression of the oxyR gene in *P. gingivalis* was not altered as a function of fluctuating hemin concentrations in the growth media and rather that OxyR activity was enhanced under hemin-limited conditions. It is possible that ferrous iron and hemin play different roles in regulating anti-oxidative stress in *P. gingivalis*. According to previous reports, ferrous iron acts as a cofactor for SOD and Dps and is required for the activity of these anti-oxidative stress proteins (12, 26). We demonstrate here that a low hemin concentration acts as a signal for the positive regulation of OxyR activity, which in turn enhances the expression of anti-oxidative stress genes.

*P. gingivalis* utilizes hemin through hemin-binding proteins. The best described are HmuY and HmuY, encoded by hmuY and pg1552. We were the first to identify a transcriptional activator of the hmu gene, CdhR (PGN1373) (29). Thus, we hypothesized in this study that expression of the sod gene would be increased in the cdhR and the hmuY mutants due to an impaired ability to take up hemin, resulting in a decreased intracellular hemin concentration, and that the fold change of sod expression observed in *P. gingivalis* 33277 in response to a limited hemin concentration would not be detected in the mutants. Unexpectedly, the hmuY and the cdhR mutants showed a pattern of enhancing sod expression similar to that in 33277 (Fig. 2). There could be two explanations for this phenomenon. The first one is that other surface proteins are involved in uptake of hemin. At least two other proteins in *P. gingivalis*, hemin uptake system protein A (HusA) and hemin-binding
protein 35 (HBP35), appear to possess hemin binding activity (9, 23). These proteins may compensate for the function of HmuY and thus fulfill the hemin requirements of *P. gingivalis*. The other explanation is that *P. gingivalis* senses a lower hemin concentration in its growth environment, but not a lower intracellular hemin concentration, and then adjusts its cellular response accordingly, in this case to enhance OxyR activity. An investigation of the mechanism by which *P. gingivalis* senses hemin-limited conditions is under way in our laboratory.

It is noteworthy that in a previous study, a rapid, OxyR-dependent response to H$_2$O$_2$ was observed in *E. coli*, such that several OxyR-regulated genes (e.g., *dps*, *ahpC*, and *trx*) were induced more than 20-fold (35). In contrast, another study used real-time PCR analysis and found that the expression of the OxyR-dependent genes, including *sod*, *dps*, *ahpC*, and *trx*, was either repressed or unchanged in *P. gingivalis* treated with H$_2$O$_2$ (10). Those authors suggested that H$_2$O$_2$ is not a strong inducer of OxyR activation in these cells and that OxyR constitutively activated transcription of oxidative-stress-related genes in *P. gingivalis*. The observations reported here, however, indicate that activation of OxyR is inducible in *P. gingivalis* by a low hemin concentration in the growth environment.

**Thioredoxins and OxyR.** Activation of bacterial transcription factor OxyR is best described in *E. coli*. A series of studies suggested the presence of a thiold-based-on-off switch in *E. coli* (3, 33, 34). They revealed that active OxyR contains an intramolecular disulfide bond between two cysteine residues, whereas the inactive form loses this disulfide bond through disulfide reductases such as thioredoxin (Trx). OxyR activation in *P. gingivalis* appears to also be controlled by this thiold-based-on-off switch. We demonstrated here that a trx mutant has an increased tolerance to H$_2$O$_2$ (Fig. 8), which probably results from an accumulation of the active form of OxyR due to a lack of Trx. Interestingly, the *trx* genes are also members of the oxyR regulon in some bacteria, including *P. gingivalis* (7). Expression of the *trx* genes is elevated by OxyR, likely via a feedback mechanism crucial for regulation of OxyR activity. We tested the *trx* regulation of *P. gingivalis* in response to different hemin concentrations. An elevated *trx* expression at both the transcriptional and translational levels was found when *P. gingivalis* was grown under hemin-limited conditions, which is presumably due to activation of OxyR. Our observation of an enhanced expression of *sod*, *trx*, and *ahpC* as a consequence of OxyR activation further underscores the essential role of OxyR in *P. gingivalis* against oxidative stresses.

**Role of OxyR regulation in pathogenicity.** The ability to regulate activation of OxyR appears to be an important feature for the survival of *P. gingivalis* in the oral cavity. Our *in vitro* studies indicated that *P. gingivalis* cells grown under hemin-limited conditions showed much higher tolerance to H$_2$O$_2$ than cells grown under excess-hemin conditions (Fig. 7). This was likely due to an activation of OxyR which upregulated expression of oxidative stress defense genes. In fact, expression of *sod*, *dps*, and *ahpC* genes was enhanced severalfold in *P. gingivalis* grown under hemin-limited conditions (Fig. 1 and 6). Previously, Smalley et al. observed that *P. gingivalis* binds to hemin in a μ-oxo dimeric form, as monitored by Mössbauer spectroscopy (24). Presumably, this dimeric form on the surface of *P. gingivalis* ties up free oxygen species and protects the organism from reactive oxidants. In this study, we provided direct evidence and support for a critical role of hemin in OxyR activation in *P. gingivalis* and in the organism’s survival under oxidative stress, which indicates that there is a sophisticated regulation system in *P. gingivalis* to protect the bacteria against harsh environments.

When *P. gingivalis* cells enter the oral cavity, the bacteria are suspended in saliva and to oral surfaces, where the oxygen tension is higher and the hemin concentration is relatively lower than in periodontal pockets. Therefore, *P. gingivalis* may adjust expression of some oxidative-stress-related genes in response to this low hemin signal, which likely greatly increases its ability to survive in this environment. In contrast, at a late stage of infection, *P. gingivalis* lives with other bacteria in more complex mixed microbial communities in deep periodontal pockets, an environment bathed by gingival crevicular fluid, which exudes from vessels of the microcirculation or is present in blood from periodontal inflammation. Proteins in gingival crevicular fluid or blood, such as hemoglobin, are an important source of hemin for *P. gingivalis*. Therefore, hemin concentrations in the periodontal pocket are much higher than those in the oral cavity of a periodontally healthy individual. In the meantime, the oxygen tension is much lower in a mature microbial community and in a deep periodontal pocket. *P. gingivalis*, therefore, represses expression of most oxidative-stress-related genes in response to a higher hemin concentration signal and a lower oxygen tension.

In conclusion, as an anaerobic bacterium, *P. gingivalis* possesses an OxyR that controls expression of some well-known anti-oxidative stress genes, such as *sod*, *trx*, *dps*, and *ahpC*. We revealed that activity of OxyR is inducible in response to a hemin-limited environment, which is likely crucial for *P. gingivalis*’ survival and pathogenicity. The findings presented here provide a foundation for further investigation into a novel mechanism of OxyR activation induced by hemin. Future studies may establish a strategy to block this pathway of OxyR activation, thus diminishing *P. gingivalis*’ ability to survive in the oral cavity.

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**REFERENCES**