Links between Anr and Quorum Sensing in *Pseudomonas aeruginosa* Biofilms

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**ABSTRACT**

In *Pseudomonas aeruginosa*, the transcription factor Anr controls the cellular response to low oxygen or anoxia. Anr activity is high in oxygen-limited environments, including biofilms and populations associated with chronic infections, and Anr is necessary for persistence in a model of pulmonary infection. In this study, we characterized the Anr regulon in biofilm-grown cells at 1% oxygen in the laboratory strain PAO1 and in a quorum sensing (QS)-deficient clinical isolate, J215. As expected, transcripts related to denitrification, arginine fermentation, high-affinity cytochrome oxidases, and CupA fimbriae were lower in the Δanr derivatives. In addition, we observed that transcripts associated with quorum sensing regulation, iron acquisition and storage, type VI secretion, and the catabolism of aromatic compounds were also differentially expressed in the Δanr strains. Prior reports have shown that quorum sensing-defective mutants have higher levels of denitrification, and we found that multiple Anr-regulated processes, including denitrification, were strongly inversely proportional to quorum sensing in both transcriptional and protein-based assays. We also found that in LasR-defective strains but not their LasR-intact counterparts, Anr regulated the production of the 4-hydroxy-2-alkylquinolines, which play roles in quorum sensing and interspecies interactions. These data show that Anr was required for the expression of important metabolic pathways in low-oxygen biofilms, and they reveal an expanded and compensatory role for Anr in the regulation of virulence-related genes in quorum sensing mutants, such as those commonly isolated from infections.

**IMPORTANCE**

*Pseudomonas aeruginosa* causes acute ocular, soft tissue, and pulmonary infections, as well as chronic infections in the airways of cystic fibrosis patients. *P. aeruginosa* uses quorum sensing (QS) to regulate virulence, but mutations in the gene encoding the master regulator of QS, lasR, are frequently observed in clinical isolates. We demonstrated that the regulon attributed to Anr, an oxygen-sensitive transcription factor, was more highly expressed in lasR mutants. Furthermore, we show that Anr regulates the production of several different secreted factors in lasR mutants. These data demonstrate the importance of Anr in naturally occurring quorum sensing mutants in the context of chronic infections.

*Pseudomonas aeruginosa*, a notorious pulmonary pathogen, is frequently a causative agent of nosocomial pneumonias (1), is commonly isolated from the lungs of chronic obstructive pulmonary disease (COPD) patients experiencing exacerbation (2), and is a problematic colonizer of the lungs of individuals with cystic fibrosis (CF) (3). By age 20, 80% of CF patients harbor *P. aeruginosa* in their lungs (4), and the presence of *P. aeruginosa* in the airway is correlated with accelerated lung function decline and poor patient prognosis (5, 6). Evidence suggests that in the context of infections, *P. aeruginosa* is often found in a biofilm state, which contributes to its extreme recalcitrance to antibiotic treatments or clearance by surveilling immune cells (7–9).

Multiple lines of evidence show that oxygen concentrations within *P. aeruginosa* biofilms and *Pseudomonas*-infected mucus in CF patient airways are low due to factors such as reduced ventilation, chronic inflammation, and the consumption of oxygen by microbes (10–14). *P. aeruginosa* senses and responds to low levels of environmental oxygen through the activity of the transcription factor Anr, due to the fact that Anr requires an intact, oxygen-labile [4Fe-4S]2+ cluster for dimerization and subsequent DNA binding (15, 16). In addition, Anr activity is stimulated by phosphatidylycerine (PC) catabolic products that are abundant in vivo (17). While required for anoxic growth via denitrification, *P. aeruginosa* Δanr strains are not impaired in growth under hypoxic (low-oxygen) conditions (18, 19). Anr homologs have been identified as regulators of virulence in other Gram-negative microbes (20–22). The high-level expression of transcripts encoding the denitrification and arginine fermentation machinery, as well as certain high-affinity cytochrome oxidases (23), suggests that Anr activity is high in vivo (11, 18, 24).

In both *in vitro* (25) and clinical (3) *P. aeruginosa* biofilms, cells use quorum sensing (QS) cascades to coordinately regulate gene expression (26). *P. aeruginosa* QS is controlled by three hierarchically arranged systems, with the LasRI system being the regulatory circuit in control of downstream pathways involving RhlRI and *Pseudomonas* quinoline signaling systems. QS-controlled viru-
lence factors include pyocyanin, hydrogen cyanide, protease, and lipase (27). Because quorum sensing positively regulates virulence factors, it may seem paradoxical that loss-of-function mutations in lasR are frequently observed in strains isolated from the CF airway (28) and that the presence of lasR mutants in a CF infection is associated with a higher rate of lung function decline (29). In addition, lasR mutants have been observed in acute infections at other body sites (30–32) and can arise spontaneously in laboratory-grown biofilms (33). Taken together, these data imply that under certain conditions, the loss of lasR confers a selective advantage. Previous studies have demonstrated that lasR mutants grow to higher cell densities on specific amino acids found in CF sputum (33), resist cell lysis in high-density cultures (34), show increased resistance to oxidative stress and antibiotic treatment (33, 35), and have higher rates of denitrification (36). The prevalence of QS mutants in infections and their relationship with disease progression illustrates the importance of understanding how pathogenesis is regulated in these strains.

Anr activity has been profiled in planktonic cultures grown axonically with nitrate, an important alternative electron acceptor for P. aeruginosa. In this study, we used transcriptome sequencing (RNA-Seq) to examine the Anr regulon in colony biofilms grown in low oxygen without exogenous nitrate, using two strains of P. aeruginosa: PAO1 (a laboratory strain) and a QS-deficient CF clinical isolate. We observed Anr regulation of the denitrification machinery under these conditions, as well as a role for Anr in regulation of high-affinity cytochromes, the arginine fermentation genes, and transcripts associated with CupA fimbriae. We also observed Anr regulation of the 4-hydroxy-2-alkylquinoline (HAQ)-dependent quorum sensing pathway, iron acquisition and storage, type VI secretion, the catabolism of aromatic compounds, and many hypothetical proteins. We established that production of CupA fimbriae, known to be important for acute and chronic infections (37, 38), was strictly dependent on Anr in both laboratory and clinical isolates and that enhanced Anr activity increases CupA production. Using both constructed and naturally occurring lasR mutants, we showed that Anr activity increased in the absence of LasRI signaling. Furthermore, we identified a role for Anr in production of HAQs in LasRI signaling-deficient strains but not their QS-competent counterparts. We propose that in the absence of LasRI signaling, Anr is an important regulator of pathogenic processes and that increased expression of the Anr regulon when LasR signaling is off may help explain the basis of selection for lasR mutants in vivo.

MATERIALS AND METHODS

Growth conditions. All strains used in this study are listed in Table S1 in the supplemental material. J215 is a tracheal isolate from an individual with CF at the Dartmouth-Hitchcock Medical Center in Lebanon, NH. P. aeruginosa and Escherichia coli were routinely cultured in lysogeny broth (LB) at 37°C, and the medium was supplemented with gentamicin (60 μg/ml for Pseudomonas and 15 μg/ml for E. coli) and carbenicillin (300 μg/ml and 100 μg/ml) as required. For studies under low-oxygen conditions, strains were grown at 30°C inside a hypoxic cabinet with an O2 controller and CO2 controller (COY Laboratory Products, Grass Lake, MI) at 1% O2 and 5% CO2. Colony biofilms were inoculated with cells from overnight cultures that had been washed and diluted to an optical density of 600 nm (OD600) of 1.0. Five microliters of this suspension was spotted onto the surface of a T-broth (10 g of tryptone and 5 g of NaCl per liter) agar plate, allowed to dry, and then incubated for 12 to 72 h as indicated.

Construction of in-frame deletion mutants and plasmids. Strains and plasmids were built using a Saccharomyces cerevisiae recombination technique described previously (39). Primers used in the construction of plasmids are listed in Table S1 in the supplemental material. Knockout constructs generated in this study were built using the pMQ30 allelic replacement vector. The gcrABC expression plasmid was built using the Pbal, expression vector pMQ70, and the anr expression plasmids were built using the Pmoe expression vector pMQ123.

Cycle sequencing of lasR, lasI, rhlI, and rhlII in J215. Target genes were PCR amplified from J215 genomic DNA and the products were sequenced at the Molecular Biology Core at the Geisel School of Medicine at Dartmouth. The resulting sequences were aligned to the PA01 genomic sequence using the NCBI BLAST program (40).

RNA sequencing analysis. Colony biofilms of wild-type (WT) or Δanr PAO1 and J215 were grown for 12 h, then harvested in 1 ml of phosphate-buffered saline (PBS) applied to the plate, followed by recovery with an angled glass rod. Samples were pelleted by centrifugation and stored at −80°C. RNA was isolated from pelleted cells using the RNeasy minikit (Qiagen), followed by treatment with RNase-free DNase from Promega, both in accordance with the manufacturers’ instructions. RNA quality was assessed using a Bioanalyzer (Agilent Technologies). Two biological replicates (samples from separate single colonies) were analyzed for each strain. One microgram of total RNA was tested for RNAi and tRNA removal using the MICROBExpress bacterial mRNA enrichment kit (Life Technologies) before sequencing. Single-read RNA-Seq was performed on the HiSeq platform at the Helmholtz Center for Infection Research (Braunschweig, Germany). Raw reads were processed and normalized using the CLC Genomics Workbench platform (v7.5.1) using the default parameter setting installed by the manufacturer. All sequences were trimmed and mapped to the PAO1 (GenBank accession number NC_002516) reference genome using the RNA-Seq analysis tool, and mapped reads were quantile normalized to control for any differences in library size. Very-low-abundance transcripts (<10 mapped reads in all samples) were discarded from further analysis, since there is little power to detect expression changes of genes expressed at low levels.

Western blot analysis of CupA1 and OprF. Cells grown as colony biofilms were harvested as described for RNA extraction. Cells were pelleted by centrifugation and boiled in SDS loading buffer for 10 min to generate a whole-cell lysate. Protein concentrations were determined using a bicinchoninic acid (BCA) protein assay reagent (Pierce Biotechnology, Inc.). Proteins were separated on a 15% acrylamide gel via SDS-PAGE for 1 h at 180 V. Proteins were transferred to a polyvinylidene difluoride membrane, washed, and probed with polyclonal serum directed against CupA1 as the primary antibody (41) and a peroxidase-conjugated goat anti-rabbit antibody as the secondary antibody (Sigma-Aldrich). Bound antibodies were visualized by SuperSignal West Pico chemiluminescent substrate (Pierce). Densitometry measurements of CupA1 were conducted using ImageJ (42).

NanoString analysis of P. aeruginosa transcripts. The NanoString nCounter analysis system (NanoString Technologies) was used to analyze the transcript abundance for 75 transcripts and was used with a custom-designed codeset. Each reaction mixture contained 80 ng of RNA in 5 μl of hybridization buffer containing reporter probes, capture probes, and 6 positive and 8 negative controls. Overnight hybridization of RNA with reporter and capture probes was conducted at 65°C and was followed by sample preparation using the NanoString prep station. Finally, targets were counted on the nCounter using 255 fields of view per sample. Data were analyzed using nSolver Analysis software v1.1. Raw counts for all transcripts were normalized to the arithmetic mean of six positive controls and to the geometric mean of three P. aeruginosa housekeeping genes (fpf, ppiD, and rpoD).

Identification of J215 pqsA::TnMar, pqsB::TnMar, and pqsH::TnMar mutants. Overnight cultures of E. coli S17-1 Apir carrying the pBT20 plasmid and J215 recipient strain were subcultured and grown to an OD600 of 1.0, at which point 1 ml of each culture was washed and sus-
pended in 1 ml LB, and the J215 culture was incubated at 42°C for 10 min. Five hundred microliters from each culture was combined, pelleted, and suspended in 40 μl of LB. This mixture was spotted on LB agar and incubated at 30°C for 22 h. The entire colony was collected, suspended in 100 μl of LB, spread on LB agar containing gentamicin (60 μg/ml) and nalidixic acid (40 μg/ml), and incubated at 30°C and 1% O2. After 3 days, colonies that failed to produce an iridescent sheen were identified and analyzed by arbitrarily primed PCR as previously described (43). Returned sequences were mapped using the Pseudomonas Genome Database BLAST function (44). These three mutants were among those identified.

S. aureus inhibition assays. P. aeruginosa strains to be tested were grown overnight in LB at 37°C, then washed, and suspended to an O.D. of 1.0. This suspension (5 μl) was spotted on Whatman paper discs on T-broth agar and incubated for 24 h at the desired oxygen concentration. S. aureus strain 8325-4 was grown with shaking overnight at 37°C in tryptic soy broth, then washed, and suspended to an O.D. of 0.1. S. aureus suspension (100 μl) was spread on tryptic soy agar plates using glass beads. Whatman paper discs with P. aeruginosa biofilms were transferred to the plates, and the zone of inhibition was observed after an additional 16 h of incubation.

Statistical analyses. Fold change values and significance statistics between RNA-Seq samples were calculated using the “Empirical analysis of DGE” algorithm in the CLC Genomics Workbench, which is a reimplementation of the “Exact Test” from the EdgeR Bioconductor package (45, 46) and which was conducted between all pairs, with a total count filter cutoff of 5.0. For the comparison between LasR-regulated transcripts in PAO1 and J215 (see Fig. S1 in the supplemental material), significance was determined using a Wilcoxon rank sum test, with a P value of ≤0.05 considered significant. Differences in expression of LasR and Anr-regulated transcripts in lasR mutants (see Fig. 3) were evaluated with a paired t test, and a P value of ≤0.05 was considered significant. In NanoString experiments (see Fig. 5), significance was determined with ratio paired t test, and a P value of ≤0.05 was considered significant.

RNA sequencing data accession number. The raw and processed RNA-Seq data have been deposited into NCBI Gene Expression Omnibus under accession number GSE68534.

RESULTS

Profiling of the Anr regulon in two P. aeruginosa strains grown as colony biofilms in 1% oxygen. We sought to define the Anr regulon under conditions that relate to those in the mucus plgs that form in CF airways and in clinically relevant biofilms (e.g., low oxygen and high cell density) in two P. aeruginosa strains (PAO1 and a clinical isolate, J215) and their Δanr derivatives. PAO1 is a commonly used laboratory strain with intact quorum sensing. Clinical isolate J215 had colony morphology characteristics of lasR loss-of-function mutants, including the lack of pyocyanin production and the presence of an iridescent colony sheen (33). J215 has a lasR588T allele that encodes LasR E196D, a variant shown previously to lead to decreased LasR activity or LasR loss of function (32). Other synonymous mutations in last, rhlR, and rhlII, as well as nonsynonymous mutations in rhlII, were identified (see Table S2 in the supplemental material).

For the RNA-Seq analysis, RNA was harvested from P. aeruginosa strains PAO1 and J215 grown in colony biofilms at 30°C in 1% O2 and 5% CO2 on T-broth agar for 12 h. We have shown previously that Anr activity is high under these conditions (17). It is important to note that the culture medium was not amended with nitrate or other compounds that can support energy generation by denitrification. The wild type (WT) and corresponding Δanr derivative in each background grew similarly, with 1.0E8 ± 0.29E8 CFU and 1.1E8 ± 0.27E8 CFU per colony for WT and Δanr PAO1, respectively, and 5.7E7 ± 0.2E7 and 9.0E7 ± 1.7E7 CFU per colony for WT and Δanr J215, respectively. Consistent with the lasRG588T allele encoding a loss-of-function LasR variant, we saw that 67 of 72 LasR-regulated transcripts were expressed at a significantly lower level in J215 than in PAO1 (P < 0.5) (see Fig. S1 in the supplemental material).

Deletion of anr had considerable effects on transcription in both PAO1 and J215. Two hundred fifty-nine genes were significantly different (P < 0.05), more than 2-fold, between the WT and the anr mutant in both backgrounds (see Data set S1 in the supplemental material). A summary of the major genes and pathways transcriptionally affected by the loss of anr is presented in Fig. 1. Below, we first describe Anr regulation of known Anr-regulated transcripts, many of which are involved in metabolism when oxygen is limiting. In addition, we describe the discovery that loss of Anr influences expression of pathways involved in the production of secreted molecules and factors.

Differential expression of known Anr-regulated pathways in colonies grown in the absence of nitrate at 1% oxygen. Many of the genes differentially regulated in both strains are known to be under the control of Anr and encode proteins involved in the metabolic response to a low-oxygen environment. For example, Anr impacted the expression of terminal oxidases involved in aerobic respiration. Anr was required for induction of the cbb-2-oxidase (ccoN2O2Q2P2) and repression of the cyanide-insensitive oxidase (cioAB, PA3928) (47). Levels of ccoN2O2Q2P2 transcript were 12- to 60-fold lower and levels of cioAB were 2- to 8-fold higher in anr mutants from both strains. Transcripts for both the cbb-1 (ccoN1O1P1) and cytochrome bo, (cyoABC) oxidases were higher in the anr mutants in both PAO1 and J215 (2- to 3-fold and 2- to 6-fold, respectively). In contrast, the loss of anr did not affect expression of the aa3 oxidase encoded by ccoAB-coll. Together, these data confirm that Anr participates in the control of the adaptation of respiration under low-oxygen conditions. Anr also controlled expression of genes involved in heme biosynthesis. Both hemF and hemN, are known to be controlled by Anr/Dnr (23, 48) and were expressed between 2- and 5-fold lower in the anr mutants. Because high-affinity cytochromes require heme as a cofactor, Anr may mediate their coordinated expression. The diheme cytochrome c555 peroxidase precursor, encoded by cpxR, is appreciated to be regulated by Anr (23), and it was reduced ~20-fold in both strains.

Transcripts involved in arginine fermentation (arcDABC) were 6- to 23-fold higher in WT strains than in the anr mutants. Our data additionally showed Anr-dependent expression of multiple genes involved in other fermentation pathways, including those for two putative alcohol dehydrogenases that are Anr regulated, adhA (39-fold and 65-fold lower in the Δanr mutants of PAO1 and J215, respectively) and PA2119 (6- and 4-fold lower in the Δanr mutants) (23, 49), as well as a phosphate acetyltransferase pta (5- and 3-fold lower) and an acetate kinase ackA (4- and 3-fold lower) gene. Studies have established a role for P. aeruginosa fermentation pathways and universal stress response proteins in long-term survival within anaerobic biofilms (50, 51). In line with previous work, we found that Anr-dependent expression of the stress response genes uspK, uspL, uspN, and uspO was lower in both Δanr mutants than in their parental strains.

Despite the absence of added nitrate, nitrite or nitric oxide, genes involved in denitrification (dnr, narK1K2GHI, narXL, nirSMCFLGHJEN, nirQ, norCBD, and norR2DFLY, as well as cooperonic hypothetical protein genes) were expressed in the WT
strains and were much lower in the Δnar derivatives (up to ~150-fold reduced). Interestingly, the nar genes responsible for the initial reduction of nitrate to nitrite were more highly expressed in PAO1 than in J215 (Fig. 1).

Anr also regulates the production of OprG and of CupA fimbriae; it is not yet known if these factors influence metabolism when oxygen is limiting. The gene that encodes outer membrane protein OprG was reduced 18- and 35-fold in the Δanr mutants from PAO1 and J215, respectively, consistent with previous reports (52). The chaperone usher pili, including the CupA fimbrial appendages, are expressed on the cell surface and have been implicated in biofilm formation and disease (37, 38, 53, 54). Anr positively regulates the expression of CupA-encoding genes through the activity of a trimeric regulator encoded by the 3-gene operon cgrABC (55, 56). The cupA1-5 transcripts were regulated by Anr in both PAO1 and J215. We also noted that expression of the cupA1-5 genes in J215 was 2- to 29-fold higher than in PAO1 (Fig. 1).

Loss of Anr impacts the expression of genes involved in iron acquisition and quorum sensing. A notable signature from our data sets reflected a change in expression levels of iron acquisition and storage pathways upon the loss of Anr. The pchDCBA and pchEFHI pyochelin biosynthesis and transport genes showed lower expression in the Δanr mutant than in the WT for both PAO1 and J215 (between 2- and 14-fold); the pyochelin outer membrane receptor gene fptA had a similar expression pattern. We also saw decreased expression of feoB and bfrA, which encode a ferrous iron transporter and bacterioferritin, respectively. Genes encoding the other siderophore produced by P. aeruginosa, pyoverdine, were not differentially expressed in PAO1 but showed slightly higher expression in the J215 Δanr mutant than in the wild type (pvdA, pvdN, pvdM, and pvdS were induced 2- to 5-fold). However, the transcript encoding ExbB1, involved in pyoverdine uptake, was more than 5-fold lower in the Δanr mutants from both strains, and exbD1, also involved in pyoverdine uptake, was 16-fold lower in J215 Δanr than in the wild type but not differentially expressed in PAO1.

Our analysis also revealed a role for Anr in regulating expression of multiple pathways related to quorum sensing. Among these were hcnABC, involved in production of hydrogen cyanide. The hcn operon is regulated by both LasR and Anr (57), and while transcripts from this operon were much lower in J215 than in PAO1, we observed a reduction of expression in both Δanr mutants (Fig. 1). Our data set also showed a strong role for Anr in regulation of the small RNA PhrS, which is part of the Pseudomonas quinoline system and has been shown to be controlled by Anr previously (58). PhrS levels were reduced 200-fold in PAO1 and 100-fold in J215. The role for Anr in regulating other LasR-controlled transcripts is discussed further below.

In both strains, a very strong feature of the RNA-Seq data set was the upregulation of genes involved in degradation of aromatic compounds in the Δanr mutants. Such transcripts included antABC and catBCA, which were expressed between 50- and 150-fold higher in Δanr strains. AntR, the positive regulator of antABC, was expressed at approximately 10-fold-higher levels. Anr also repressed the gene for the hypothetical protein adjacent to catA, PA2506. The ant and cat gene products degrade anthranilate, which is a precursor to the QS molecule HHQ (59). The list of additional genes that were differentially expressed upon the loss of Anr in both strains includes numerous genes that encode hypothetical proteins (see Data set SI in the supplemental material).
Anr-dependent regulation of CupA is greater in the absence of LasR. Visual inspection of transcripts strongly regulated by Anr (e.g., ccoN2O2Q2P2, arcDABC, and cupA1-A5) suggested a larger difference in expression between the WT and the Δanr mutant for J215 compared to PAO1. To test this, we performed a Western blot analysis of Anr-regulated CupA fimbriae using an anti-J215 compared to PAO1. Levels of CupA1 and OprF, a reference protein, were determined by Western blotting of whole-cell lysates from 48-h colony biofilms grown on T-broth agar. (A) The wild type (WT) and mutant derivatives of strains PAO1 and J215 grown in 1% O2. (B) Comparison of CupA1 production in WT and Δanr J215 after growth at 21% and 1% O2. (C) PAO1 and PA14 wild types compared to their corresponding lasR mutant derivatives, after growth in 1% O2. A PA14 lasR Δanr double mutant is also shown. (D) A lasR-negative clinical isolate (lasR− C.I.; NC-AMT0101-1), a lasR loss-of-function mutant isolated from a CF airway infection, and its lasR+ C.I. parental strain (NC-AMT0101-2) (28) are shown along with their isogenic Δanr derivatives after growth at 1% O2.

The loss of LasR signaling causes an Anr-dependent increase in CupA fimbriae. The observation that CupA1 levels were higher in J215 than in PAO1 suggested a connection between the loss of LasR/N-3-oxo-dodecanoyl-homoserine lactone (3OC12HSL) quorum sensing and increased Anr activity. In order to test this model further, we measured production of CupA fimbriae in strain PAO1, strain PA14 (another laboratory strain), and their respective ΔlasR derivatives. In both instances the ΔlasR mutant had higher levels of CupA fimbriae than the wild-type parental strain (10-fold higher in PAO1 ΔlasR than in the PAO1 wild type and 4-fold higher in PA14 ΔlasR than in the PA14 wild type) (Fig. 2C). We also measured CupA production in a pair of genetically related CF clinical strains isolated from the same subject (28). Previous genomic analyses revealed that one isolate, NC-AMT0101-2, has a functional allele of lasR, while NC-AMT0101-1 had acquired a lasR mutation. The lasR-defective clinical isolate produced 7.5-fold more CupA fimbriae than the parental strain (Fig. 2D), and in both NC-AMT0101-1 and NC-AMT0101-2, deletion of anr abolished CupA1 production (Fig. 2D). Taken together, these results suggest that lasR mutants have higher levels of Anr activity and that this leads to higher production of CupA fimbriae.
Anr is repressed by LasR/3OC12HSL. To further test the hypothesis that QS signaling is inversely correlated with Anr activity, we used a multiplex method for the simultaneous analysis of multiple transcripts with the nCounter NanoString technology. NanoString mRNA quantitation uses fluorescent probes to capture and count specific mRNA targets (60), and we developed a NanoString codeset that included a number of QS and Anr-regulated transcripts. The genes directly controlled by Anr in our codeset were arcA, arcD, adhA, conN2, ccoP2, dnr, narG, nirS, norC, cgrA, and PA1673. The QS-controlled genes were lasI, lasB, rhlI, rhlA, pqsA, pqsE, pqsH, phzA2, phzC, phzM, and phzG. We analyzed mRNA levels of these transcripts in PAO1 and PA14 and their lasR mutant derivatives, the paired clinical isolates NC-AMT0101-2 and NC-AMT0101-1 (described above), and another set of genetically related clinical isolates with and without functional LasR (AMT0047-2 and AMT0047-3, respectively) (28). We observed that nearly all Anr-regulated genes were more highly expressed in the lasR mutants than in their cognate lasR-intact strains (Fig. 3). For example, expression of Anr-regulated ccoN2 was 4- to 12-fold higher in the lasR mutants and PA1673 expression was 5- to 25-fold higher in the lasR mutants. As expected, QS-regulated transcripts were uniformly lower in the absence of functional LasR. In addition to comparing wild-type and LasR-defective strains, we examined Anr activity in a ΔlasI mutant, which lacks the 3OC12HSL synthase, in the absence and presence of 3OC12HSL (61). We found that complementation of the ΔlasI strain with exogenous 3OC12HSL was sufficient to rescue the expression of QS-controlled genes and led to decreased expression of Anr-regulated genes (Fig. 3). Across all five pairs of samples, 7 of 11 Anr-regulated transcripts and 8 of 11 QS-regulated transcripts were significantly different (P < 0.05, paired t-test). We did observe that in both AMT0047-3 and NC-AMT0101-1 (carrying natural lasR defective alleles) but not the other lasR or lasI mutants, narG expression was reduced. There is a potential LasR binding motif (CTTACTGTTTAAAAAG) 150 bp upstream of narKI translational start (the first gene in the operon containing narG), and our data may indicate a positive role for LasR in regulating expression of this operon in these strain backgrounds. This is consistent with reduced expression of the narKI-PA3871 operon in J215 compared with that in PAO1 (Fig. 1). Expression of norC was also decreased in AMT0047-3 compared to its parental strain, AMT0047-2. In contrast, four flagellar transcripts (fliG, fliC, and flgK, and flgD) did not vary between the pairs (Fig. 3). The repression of Anr activity upon addition of 3OC12HSL to a ΔlasI strain indicated that Anr was directly responsive to LasR signaling and that increased Anr activity was not due to a secondary effect common in the absence of LasR. The complete NanoString data set from these experiments is provided in Data set S2 in the supplemental material. LasR regulation of Anr is likely indirect, as there is no evidence that LasR binds the anr promoter (62), and anr transcript levels are not altered in transcriptional profiling analyses of the LasR regulon (63, 64) (see Data set S2). LasR positively regulates 3,4-dihydroxy-2-heptylquinolone (PQS) production (65), and PQS has been shown to inhibit denitrification (66). To determine if increased Anr activity is due to a decrease in PQS, we tested whether CupA1 production was greater in three mutants in the PQS pathway in J215. There were no detectable differences (see Fig. S2E in the supplemental material), indicating that HAQ production does not likely effect Anr activity in J215.

Anr partially compensates for the loss of LasR signaling in the regulation of HAQs. We showed above that mutants or strains with lower levels of LasR activity have increased Anr activity. Our studies also showed that in LasR-defective backgrounds, Anr plays roles that are not evident in lasR- laboratory strains. When active, LasR regulates production of 4-hydroxy-2-alkyquinolines (HAQs), including PQS and its direct precursor 4-hydroxy-2-heptylquinolone (HHQ). The iridescent sheen that is characteristic of lasR mutants is caused by an accumulation of HHQ, which is due to an inability to properly induce the pqsH product, the enzyme that converts HHQ to PQS, in the absence of LasR (33). In contrast to the J215 WT, J215 Δanr did not make HHQ sheen when grown in 1% oxygen (Fig. 4A). These visual phenotypes were supported by the RNA-Seq data, which showed that three of the five transcripts in the HHQ biosynthetic operon (pqsA, pqsC, and pqsE) were 2- to 8-fold lower in J215 Δanr than in the J215 WT. In contrast, the loss of anr had no effect on colony morphology in PAO1 (see Fig. S3A in the supplemental material), and pqs transcripts were not different between the PAO1 WT and
PAO1 Δanr. Interestingly, anr was not required for HHQ production when cells were grown at 21% oxygen, a condition correlated with lower Anr activity (see Fig. S3B). Consistent with the RNA-Seq analysis of HAQ-related transcripts and phenotypic data, a targeted analysis of pqsA and pqsE in J215 was performed. In the absence of Anr, pqsA and pqsE were both reduced in the J215 background (Fig. 5A; see also Data set S3 in the supplemental material). The cgrA, cupA1, and cupA3 transcripts, as expected based on the data shown above, also followed this pattern. Expression of these genes was restored upon complementation with anr at the native locus. Anr-dependent regulation of the pqs and cupA genes was also observed in the PA14 ΔlasR and PA14 ΔlasR Δanr pair (Fig. 5B; see also Data set S3). Together, these data suggest that Anr activity impacts HAQ production in strains with low or absent LasR activity.

In addition to impacting pqs transcription and HHQ production, deletion of Anr also affected production of the HHQ-derived exoproduct 4-hydroxy-2-heptylquinoline N-oxide (HQNO), which can slow the growth of *Staphylococcus aureus* and other Gram-positive organisms by inhibiting electron transport (67, 68). We tested the role for anr in this interaction by exposing a lawn of *S. aureus* to J215 biofilms and observing the zone of growth inhibition. We saw that biofilms formed by J215 WT inhibited *S. aureus* in 1% oxygen and that this ability required anr and pqsA (Fig. 4A). To further study Anr regulation of HAQs in *lasR* mutant backgrounds, we examined the role for Anr in PA14 ΔlasR. (Strain PAO1 ΔlasR was not used because it does not over-produce HHQ, a fact that has been previously published [33].) Deletion of anr in a PA14 *lasR* mutant led to a marked decrease in HHQ production and an inability to inhibit *S. aureus* at 1% oxygen, while deletion of anr in a *lasR* intact background had no effect on either colony morphology or *S. aureus* inhibition (Fig. 4B).

![FIG 4](http://jb.asm.org/)

**FIG 4** Anr is required for HHQ-dependent colony sheen and HQNO-dependent *S. aureus* inhibition at 1% oxygen in *lasR* mutants. (A) J215; (B) PA14. Top row, colony biofilms at 5 days on T-broth agar. Bottom row, Whatman paper discs with 24-h biofilms grown on T-broth agar were transferred to tryptic soy agar plates spread with a lawn of *S. aureus* and grown for an additional 16 h.

![FIG 5](http://jb.asm.org/)

**FIG 5** Anr is required for expression of virulence-associated pathways in *lasR* mutants. NanoString data from colony biofilms grown in 1% O₂ for 12 h are shown. (A) Data represent the average number of transcript copies from 3 biological replicates. Bars represent standard deviations. Significance was determined by ratio paired t test, *, *P* < 0.05; **, *P* < 0.01. (B) Heat map representation of one of the experiments included in panel A, as well as a separate experiment measuring PA14 Δ*lasR* and PA14 Δ*lasR Δanr*. Z-scoring was done by row.

With J215, anr was not required for HHQ production in PA14 Δ*lasR* at 21% oxygen (see Fig. S3B in the supplemental material). It is important to note that Anr activity was not sufficient to maintain the same level of HAQ production as was seen in *lasR*-intact strains; transcriptional data showed that pqs transcripts were less abundant in J215 than in PAO1 (Fig. 1), and the zone of *S. aureus* inhibition is larger in PA14 than in PA14 Δ*lasR* (Fig. 4B). However, these data indicate that Anr supports biologically active levels of HAQ production in low-oxygen biofilms in the absence of LasR.

We tested a number of potential mechanisms by which Anr could impact HAQ production in *lasR* mutants. RhlR has been shown to activate HAQ production in *lasR* mutants under certain conditions (69, 70), but as anr is required for the production of HAQs in both the PA14 Δ*lasR* mutant and the PA14 Δ*lasR* Δ*rhlR* double mutant, we conclude that Anr-dependent regulation of HAQ production is not through RhlR (see Fig. S4A in the supplemental material). Sonnleitner et al. described a connection between Anr and HAQs through PhrS, a small ncRNA that activates translation of *pqsR* and is induced by Anr (58). In our RNA-Seq experiment, both PAO1 and J215 anr mutants showed a strong reduction in phrS expression (200-fold and 100-fold, respec-
tively). However, a J215 ΔphrS mutant did not show a decrease in the HHQ-dependent colony phenotype that is lost upon mutation of anr (see Fig. S4B). The antABC, catBCA, and xylXYZL gene products are involved in degradation of aromatic compounds, including the HHQ/HQNO precursor anthranilate. These genes were induced up to 140-fold in both Δanr mutants, and we reasoned that overactivity of these pathways could deplete intracellular anthranilate and lead to an inability to synthesize HAQs. However, deletion of antA in J215 Δanr did not restore HHQ production, as colony biofilms from this strain remained smooth (see Fig. S4B). Because Anr is necessary for expression of arcD, which encodes an arginine-ornithine antiporter, and because arginine has been linked with HHQ-mediated modulation of P. aeruginosa swarming motility (71), we hypothesized that the anr defect may be linked to an inability to acquire or synthesize arginine. However, growth on media supplemented with 0.4% arginine did not restore colony sheen in J215 Δanr (see Fig. S4B). Finally, Dnr, a major downstream regulator under the control of Anr, does not participate in Anr regulation of HAQs, as a Δdnr strain retains HHQ production, while the Anr mutant does not (see Fig. S4B). Taken together, these data indicate that the Anr effect on HAQ production is likely due to a confluence of factors or through an unrecognized pathway.

Analysis of links between LasR and Anr in the P. aeruginosa genome. In the RNA-Seq data, Anr-dependent expression of genes in the H2-type VI secretion system was more pronounced in J215 than in PAO1 (Fig. 1). The H2-type VI secretion system delivers a phospholipase with activity against bacterial membranes, PldA, directly into target cells (72) and has been shown to contribute to P. aeruginosa virulence in eukaryotic models of infection (73, 74). Expression of the H2-type VI secretion locus is controlled by LasR/3OC12HSL (73, 74), and a putative Anr-binding site has been identified at bp −174 relative to the start of transcription of the operon (23). Further analysis confirmed that expression of hsiB2, a gene within the H2-type VI secretion system operon, was dependent on Anr in the absence of functional LasR (Fig. 5B).

In order to identify additional genes potentially regulated by both QS and Anr, we cross-referenced data sets from a LasR-ChIP experiment (62) and a microarray experiment comparing a LasI mutant biofilms. In the presence and absence of 3OC12HSL is low. Additionally, our data showing that Anr is necessary for induction of pathways that promote the generation of energy under low-oxygen conditions, including genes encoding high-affinity cytochrome oxidases, the machinery necessary for denitrification, and arginine fermentation enzymes (18, 23). Additionally, our conditions revealed a previously unobserved role for Anr in the control of expression of pathways related to iron acquisition and storage, HAQ production, the catabolism of aromatic compounds, and H2-type VI secretion (Fig. 6).

Because isolates with loss-of-function mutations in lasR are common in CF, it is interesting to speculate that increased Anr activity contributes to the fitness of these strains. We have shown a role for anr in a model of pulmonary infection, and there is evidence to suggest that Anr-regulated pathways are an important part of the long-term adaptation of P. aeruginosa to the CF airway (17, 18, 50, 51, 75). Additionally, our data showing that Anr activity can be reduced in a ΔlasI strain by addition of exogenous 3OC12HSL raise the possibility that Anr has an important role in the regulation of virulence factors in low-cell-density environments, when the concentration of 3OC12HSL is low.

The activities of Anr and also the E. coli Anr homolog Fnr are redox sensitive, due to the requirement of an assembled [4Fe-4S]2+ cluster for dimerization and DNA binding (76, 77). Anr is also likely affected by iron availability, as has been shown to be the
case for Fnr (78). It is possible that either of these factors is altered by LasR activity. Future studies will determine specifically how LasR affects Anr activity in P. aeruginosa.

The role for Anr in regulation of HAQ production may be both direct and indirect. The antABC and catBCA aromatic compound degradation pathways are strongly activated by intracellular anthrinate, suggesting that the increased expression of these pathways in anr mutants could reflect accumulated anthranilate due to inactivity of the pqsABCD operon. Additionally, AntR, when bound to anthranilate, has been shown to inhibit the activity of PqsR, and PqsR represses antA (59). Although deletion of antA was not sufficient to restore HHQ production in J215 Δanr, it is possible that the Δanr phenotype is due to repressive effects of AntR. Another intriguing possibility is that HAQ production is reduced in the J215 Δanr mutant as a result of either an increase in or an inability to appropriately respond to oxidative stress. Multiple genes involved in the oxidative stress response (katB, which encodes a catalase, as well as ahpF and ahpCF, which encode hydroperoxide reductases) were expressed at a level >2-fold higher in both the PAO1 and J215 Δanr mutants. Quinolones have been shown to sensitize P. aeruginosa to oxidative stress (79), and pyochelin can promote oxidative stress (80) and is regulated in response to oxidative stress (81).

The requirement for Anr in HHQ production in LasR-defective strains is interesting in light of data which showed that HHQ, rather than its derivative PQS, is required for infection in a murine model (82). P. aeruginosa cannot produce PQS anaerobically, due to the fact that PqsH (the enzyme that oxidizes HHQ to PQS) requires oxygen as a cofactor (83), and HHQ is readily detected in CF airway secretions (84). We believe that the relationship between Anr and HHQ in quorum sensing mutants may functionally compensate for effects of losing LasR, and this could help explain the ability of P. aeruginosa lasR mutants to thrive in infections. A recent study demonstrated equal infectivity between WT PAO1 and a lasRI rhlRI quadruple mutant in a mouse lung model, suggesting that homoserine lactone signaling in general may be dispensable for infection in this context (85). Future studies will be aimed at measuring the role for Anr regulation of HHQ production in infections.

Thus, LasR loss-of-function mutants show increased expression of metabolic pathways that are valuable in low oxygen, increased production of CupA fimbriae, and functionally active levels of HHQ and HQNO (Fig. 6), all of which are dependent on Anr. We propose that Anr-regulated pathways may contribute significantly to virulence and fitness in lasR mutant isolates, and future studies will be aimed at measuring the specific role for Anr-regulated pathways in lasR mutants in infections.

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REFERENCES


37. Skurnik D, Roux D, Aschard H, Cattoir V, Yoder-Himes D, Lory S, Pier GB. 2013. A comprehensive analysis of *in vitro* and *in vivo* genetic fitness of *Pseudomonas aeruginosa* using high-throughput sequencing of trans-

38. Winsland C, Langille MG, Fothergill JL, Kucikova-Ibrulu I, Paradis-


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