Anaerobiosis-induced Loss of Cytotoxicity is Due to Inactivation of Quorum Sensing in *Pseudomonas aeruginosa*

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Abstract

*Pseudomonas aeruginosa*, an opportunistic pathogen of clinical importance, causes chronic airway infections in patients with cystic fibrosis (CF). Current literature suggests that pockets with reduced oxygen tension exist in the CF airway mucus. However, virulence features of this opportunistic pathogen under such conditions are largely unknown. Cell-free supernatant of the standard laboratory *P. aeruginosa* strain PAO1 obtained from anaerobic culture, but not aerobic culture, failed to kill A549 human airway epithelial cells. Further investigation revealed that this reduced cytotoxicity upon anaerobiosis was due to the suppressed secretion of elastase, a virulence factor controlled by *P. aeruginosa* quorum sensing (QS). Both a *lacZ*-reporter fusion assay and quantitative RT-PCR analysis demonstrated that transcription of the elastase-encoding *lasB* gene was substantially decreased during anaerobic growth compared with aerobic growth. Moreover, transcription of other genes controlled by the LasI/R QS system, such as *rhlR*, *vqsR*, *mvfR*, and *rsaL*, was also repressed under the same anaerobic growth conditions. Importantly, synthesis of 3-oxo-C12 HSL (PAI-1), an autoinducer molecule that mediates induction of the LasI/R QS system, was >22-fold decreased during anaerobic growth while C4-HSL (PAI-2), which mediates RhlI/R QS, was non-detectable under the same growth conditions. Transcription of the *lasB* gene was restored by exogenous supplementation with autoinducers, with PAI-2 more effective than PAI-1 or PQS at restoring transcription of the *lasB* gene. Together, these results suggest that anaerobiosis deprives *P. aeruginosa* of the ability to regulate its virulence via QS and this misregulation attenuates the pathogenic potential of this important pathogen.
Introduction

*Pseudomonas aeruginosa* is a clinically important Gram-negative bacterium that is the causative agent of chronic airway infections in patients suffering from pneumonia and bronchiectasis, including cystic fibrosis (CF) (57). *P. aeruginosa* has developed highly sophisticated virulence mechanisms and secretes a wide range of extracellular virulence factors, such as proteases (50), exotoxin A (53), rhamnolipids (23), pyocyanin (24) and siderophores (11). Production of these virulence factors is regulated to a large extent by a cell density-dependent gene regulatory mechanism termed quorum sensing (QS) (35). The importance of QS in *P. aeruginosa* virulence has been clearly elucidated in studies using a range of infection models (10, 33, 48) and cultured host cells (8, 41).

There are three well-characterized QS systems in *P. aeruginosa*: the *las*, *rhl* and *pqs* systems, each of which plays a distinct role in orchestrating the expression of numerous virulence-associated genes (44). The *las* and *rhl* systems were initially identified to be essential for elastase and rhamnolipid production, respectively (35). Each system is composed of a transcriptional activator protein (LasR or RhlR) and a cognate autoinducer synthase, LasI or RhlI, that produces N-(3-oxododecanoyl)-L-homoserine lactone (3-Oxo-C12-HSL, PAI-1) and N-butyryl-L-homoserine (C4-HSL, PAI-2), respectively. Upon binding to its cognate signal molecule, LasR or RhlR activates the transcription of target genes (35). *P. aeruginosa* QS is also regulated by another system, which involves potentiation of transcriptional activation by MvfR (also known as PqsR) upon binding of a *Pseudomonas* quinolone signal (PQS) (55). The PQS/MvfR complex actively participates in the intertwined *P. aeruginosa* QS network and accumulating evidence now suggests that PQS-mediated QS is absolutely required.
for the uninterrupted production of elastase (29, 36).

In the CF lung, the lack of a functional CFTR channel results in the overproduction of a viscous and stagnant mucus layer (26), to which \textit{P. aeruginosa} becomes established as a microbial community known as a biofilm. This abnormally altered CF airway has been reported to harbor regions with a steep oxygen gradient ranging from aerobic to anaerobic (40, 54). Given the fact that \textit{P. aeruginosa} is able to grow anaerobically in the presence of alternative electron acceptors such as nitrate (NO$_3^-$) or nitrite (NO$_2^-$) that are present in sufficient quantity in a CF mucus layer (21, 32, 59), further research on bacterial responses to an anaerobic environment should be pursued for an integrated understanding of its virulence mechanisms. From this perspective, it is of particular interest that \textit{P. aeruginosa} growing by anaerobic respiration forms a significantly more robust biofilm than that formed during aerobic growth, allowing the establishment of a resistant mode of bacterial proliferation (31, 59). Moreover, bactericidal activity of polymorphonuclear neutrophils (PMN) was significantly decreased under conditions of low oxygen tension due to the impaired production of hydrogen peroxide (30). Together, these results suggest that long-term survival of \textit{P. aeruginosa} can be facilitated by the growth under reduced oxygen tension in the CF airway.

Recent reports revealed that a considerable proportion of \textit{P. aeruginosa} isolates from CF patients possess mutations in the \textit{lasR} gene (12, 19). Being contradictory to the current view that \textit{lasR}-mediated QS system is essential for \textit{P. aeruginosa} virulence, these findings suggest that (i) QS in the CF airway may not be required for bacterial survival, especially at the chronic stage and (ii) QS regulation may occur differently under conditions with reduced oxygen tension. Although a study of elastase production in response to varying degree of oxygen potential was reported (39), no in-depth
understanding of anaerobiosis-induced modulation of QS has been achieved. We undertook the present study to gain insight into how QS regulation is modulated upon growth under anaerobic conditions and to determine the effect of this modulation on bacterial virulence. Understanding the mode of QS regulation under such conditions will aid the development of evidence-based clinical guidelines for the management of *P. aeruginosa* airway infections.
Materials and Methods

Bacterial strains and growth conditions

*P. aeruginosa* laboratory strains (PAO1, PA14, PAK and FRD1) and pneumonia patient isolates have been previously described (25, 47, 57, 58). PAO1 ∆lasB mutant was purchased from a *P. aeruginosa* transposon mutant library (www.genome.washington.edu/UWGC/pseudomonas) and sequence verified. Unless otherwise indicated, the strains were routinely grown in Luria-Bertani broth (LB, 10 g tryptone, 5 g NaCl, 5 g yeast extract per liter) at 37 °C. *P. aeruginosa* was grown anaerobically in an anaerobic chamber (Coylab Inc. Grass Lake, MI) that was filled with mixed gas (nitrogen 90 %, hydrogen 5 % and carbon dioxide 5 %) and maintained at a temperature of 37°C.

Chamber operation to achieve and maintain anaerobic environments was performed following the manufacturer’s instructions. To enhance anaerobic growth, bacteria were inoculated in a flask with a stirrer bar in it and the flask was placed on top of the stirrer plate to allow homogeneous mixing. Anaerobic growth was supported by the addition of 0.4 % KNO₃ to the culture medium (60).

Cell viability assay

To compare the cytotoxic potential of PAO1 grown aerobically or anaerobically, cell-free culture supernatants were harvested from aerobic and anaerobic cultures that have grown to the similar final cell density. A549 human airway epithelial cells (4) were grown in minimum essential medium (MEM, Gibco/BRL, Rockville, MD, USA) containing 10 % fetal bovine serum and 1 % penicillin/streptomycin (Gibco/BRL). The cells were placed in the wells of 96-well plates at a density of 1×10⁴ cells/well and the plates were incubated overnight under normal culture conditions (37 °C and 5 % CO₂).

After 1 hr adjustment with serum-free media, bacterial culture supernatants...
reconstituted in the same serum-free media were added to the A549 cells. After a 6-hr treatment, A549 cell viability was assessed using an MTT assay kit (Sigma-Aldridge) following the instructions provided. A549 cell viability was also examined by trypan blue viability assay (49).

**Construction of a $P_{\text{lasB}}$:lacZ reporter strain and $\beta$-galactosidase assay**

The $\text{lasB}$ promoter region was PCR-amplified from the *P. aeruginosa* PAO1 chromosome using primers lasB-PF (5'-CATATACTAGTAACCTAGCTGCCACCTGC TT-3’) and lasB-PR (5’-GTAAAGGATCCCTTGTTCAGTTCTCCTGTTTTTTTC-3’), and the $\text{lacZ}$-encoding open reading frame (ORF) was amplified from the pTnKGL3 (61) vector using primers lacZ-F (5’-TATACGGATCCATGACCATGATTACGGATTCACTG-3’) and lacZ-R (5’-TGGTTCTCAGACCTTTAATAGATTATATTACTAATTAATTGGGA-3’). The $\text{lasB}$ promoter region was double-digested with SpeI/BamHI, and $\text{lacZ}$ ORF was digested with both BamHI and XhoI. The sequence-specific chromosomal delivery vector pUC18T mini-Tn7T-Gm (7) was also double-digested with SpeI/XhoI and the two digested PCR products were ligated into the cut vector. The constructed plasmid was then integrated into the *P. aeruginosa* PAO1 genome as described previously (6). $\beta$-galactosidase activity assay was performed as described previously (61).

**Quantitative real-time PCR (qRT-PCR) analysis**

Total RNA was extracted from harvested cells using TRIzol® (Invitrogen) and a RNeasy kit (Qiagen) following the manufacturers’ instructions. RNA quantification was performed using a Nanodrop spectrophotometer (model no. ASP2680, CellTAGen Inc., Seoul Korea). cDNA was synthesized using a Primescript reverse transcriptase kit
(Takara Bio Inc., Shiga, Japan) with random primers (5'-NSNSNSNSNS-3', where N=A,T,C, or G and S=C or G). Real-time PCR reactions were monitored using a StepOne Real-time PCR system (Applied Biosystems, Carlsbad, CA). SYBR premix Ex Taq (Takara) was used for PCR reactions according to the manufacturer’s instructions. Transcript levels of the rpoD gene were similar in cells grown under aerobic or anaerobic conditions and transcript levels of rpoD were thus used to normalize the real-time PCR results. The primers used for qRT-PCR are listed in Table S1.

Quantification of PAI-1 and PAI-2 in cell-free culture supernatants

Filter-passed bacterial culture supernatants were sequentially extracted with two equal volumes of ethyl acetate containing 0.01% (final concentration) glacial acetic acid. The ethyl acetate phase was collected and then evaporated to dryness. The dried residues were then dissolved in HPLC-grade ethyl acetate and stored at -20 °C. Quantification of PAI-1 was performed using GC/MS with commercially purchased PAI-1 (Sigma-Aldridge) as a standard. Gas chromatographic analyses of PAI-1 in the solvent extracts was carried out using a Agilent 6890 Plus gas chromatograph equipped with a DB-5 MS capillary column (30 m×0.25 mm i.d., 0.25 μm film thickness, 5 % diphenyl-95 % dimethylsiloxane phase, J&W Scientific, Folsom, CA). Mass spectra were obtained using a quadruple mass spectrometer system with a 5973N mass selective detector (Agilent Technologies Inc., Santa Clara CA). Chromobacterium violaceum CV026 (27) was used to quantify PAI-2 present in the culture supernatants with commercially purchased PAI-2 (Sigma-Aldridge) as a standard. CV026 was inoculated in LB supplemented with PAI-2 of known concentration or the supernatant to be tested and grown for 16–18 hr at room temperature with vigorous shaking. A 1-ml aliquot of each culture was centrifuged to precipitate the insoluble violacein. Then, 1 ml of DMSO was
added to dissolve the pellet. The absorbance of the completely solubilized violacein was measured with a ThermoMax microplate reader (Molecular Devices, Sunnyvale, CA) at a wavelength of 585 nm.

**Western blot analysis and cellular fractionation**

One milliliter aliquots of aerobic or anaerobic cultures of PAO1 grown to a similar final cell density (OD$_{600}$ of ~3.0) were centrifuged at 14,000 rpm for 5 min. Supernatants were passed through a 0.2 µm Acrodisc® Syringe Filter (Pall Life Science Inc., Ann Arbor, MI) and saved as culture supernatants. Cell pellets were resuspended in 100 µl of B-PER® protein extraction reagent (Thermo Fisher Scientific Inc., Rockford, IL) and incubated for 10 min at room temperature. Lysed cells were centrifuged at 14,000 rpm for 10 min and supernatants were recovered (cell extract fractions). Insoluble precipitates were then resuspended with 50 µl of the same B-PER® protein extraction reagent (membrane fractions). An antibody against *P. aeruginosa* elastase was a kind gift from Dr. Efrat Kessler (Tel Aviv University, Israel). Twenty microliters of culture supernatant and 20 µg of cell extract fraction and membrane fraction were loaded onto 12 % polyacrylamide gels. Proteins separated on the gel by electrophoresis were transferred to nitrocellulose membranes (Hybond™-ECL, GE healthcare) and membranes were blocked with 5 % skim mile in TBST buffer. Membranes were then probed with anti-elastase antibody (1:5,000) for 2 hrs and washed six times with TBST for 10 min each time. Membranes were then re-probed with anti-rabbit horseradish peroxidase-conjugated secondary antibody (1:5,000) and washed three more times with TBST. Finally, the membranes were incubated with ECL solution for 2 min and signals were detected on X-ray film (Kodak).
Statistical analysis

Data are expressed as mean ± SD (standard deviation). An unpaired Student’s t-test was used to analyze the data. To compare differences among more than three groups, one-way analysis of variance (ANOVA) was used. A p-value of <0.05 was considered statistically significant. All the experiments were repeated for reproducibility.
Results

Cell-free supernatants of anaerobic cultures failed to kill human airway epithelial cells. To compare the cytotoxicity of virulence factors secreted during aerobic vs. anaerobic growth, we treated A549 human airway epithelial cells with the culture supernatants from bacteria growing aerobically or anaerobically. Because production of virulence factors occurs in a cell density-dependent manner, cell-free supernatants were harvested from cultures that grew to a similar final cell density (i.e., OD$_{600}$ of ~3.75 for aerobic culture vs. ~3.59 for anaerobic culture). To support anaerobic growth, media was supplemented with 0.4 % NO$_3^-$, which acts as an alternative electron acceptor (59); this medium was also used for the aerobic cultures. As shown in Fig. 1A, A549 cells lost their viability upon treatment with aerobic culture supernatants as assessed by the MTT cell viability assay. After a 6-hr treatment, the mean OD$_{570}$ value derived from live cells had decreased to ~13 % of that of the control treatment. In contrast, A549 cells treated with anaerobic culture supernatants remained viable (Fig. 1A). The contrasting cytotoxic activity of these two culture supernatants was further confirmed by trypan blue stain assay (Fig. 1B). We next examined whether the differential cytotoxic effects of the two supernatants were reflected by host cell morphological changes. As shown in Fig. 1C, A549 cells treated with aerobic culture supernatants completely lost their normal cellular morphology, while such changes in cell shape were not observed in cells treated with anaerobic culture supernatants.

Elastase secretion is reduced under anaerobic growth conditions. Elastase, a major virulence factor (3), is the most abundant protein secreted into the culture media during aerobic growth of P. aeruginosa (Fig. 2A, black arrow). We first monitored the time
profile of elastase secretion during aerobic growth of PAO1. As growth time progressed, increased levels of elastase were detected in the culture supernatant (Fig. 2A). A band corresponding to elastase started to appear after 3 hr, when the culture reached an OD₆₀₀ of ~1.16, and continued to increase steadily as the culture reached stationary phase. As expected, the elastase band was not observed in the culture supernatants of the ΔlasB mutant (Fig. S1). To investigate the potential role of elastase in the observed cytotoxicity of the aerobic culture supernatants, we treated A549 human airway epithelial cells with the culture supernatants of PAO1 harvested at various time points. As shown in Fig. 2B, the viability of A549 cells decreased in proportion to the level of elastase present in the culture supernatants. The relative survival rate for each treatment was normalized using the value obtained from the control treatment, in which A549 cells were treated with the medium, LB+0.4 % NO₃⁻. As expected, the culture supernatant of a mutant with the elastase-encoding lasB gene interrupted by transposon insertion failed to kill A549 cells under the same experimental conditions (Fig. 2C), providing conclusive evidence that elastase was responsible for the cytotoxic activity against A549 cells.

We then examined whether the non-cytotoxic nature of the anaerobic culture supernatant was due to a factor associated with the modulation of elastase secretion. Indeed, the levels of elastase present in anaerobic culture supernatants were significantly lower in our SDS-PAGE analysis than those present in the aerobic supernatants (Fig. 2D). Again, the final cell density (OD₆₀₀ of ~3.75 vs. ~3.59) after 18 hrs of growth was similar between these two cultures, implying that the anaerobiosis-specific repression of elastase secretion was not due to retarded bacterial growth. Together, these results suggest that elastase secretion is highly suppressed during
anaerobic growth and such repression is responsible for the loss of cytotoxicity towards
A549 human airway epithelial cells.

Next, we examined whether the anaerobiosis-induced decrease in elastase secretion
was also observed in other \textit{P. aeruginosa} strains. As in the case of PAO1, PA14 and
three non-mucoid pneumonia patient isolates produced sufficient levels of elastase
during aerobic growth. In contrast, PAK, a highly piliated \textit{P. aeruginosa} strain and
FRD1, a mucoid CF patient isolate produced a negligible or significantly decreased
level of elastase, respectively (Fig. S2A). During anaerobic growth, however, all tested
strains produced very low levels of elastase, thereby further confirming our results in
PAO1 (Fig. S2A). Again, relative cytotoxicity was directly proportional to the amount
of elastase present in the culture supernatants (Fig. S2B).

\textbf{Production, but not secretion, of elastase was decreased under anaerobic}
\textbf{conditions.} Elastase has been reported to be synthesized as a \textasciitilde53 kDa preproenzyme
containing a \textasciitilde2.4 kDa signal sequence and to be translocated to the periplasmic space
after synthesis. The resultant periplasmic protein is then further processed to generate
the mature secretory \textasciitilde33 kDa elastase and \textasciitilde18 kDa propeptide, which acts as an
elastase inhibitor by forming a complex with the processed elastase (22). To determine
whether the decrease in elastase secretion during anaerobic growth was caused by
decreased production or decreased levels of secretion, we analyzed levels of elastase in
each of three different cellular fractions; the culture supernatant (CS), the cell extract
(Ext), and the membrane fraction (Mem) by western blot analysis. As shown in Fig. 3A,
elastase was detected only in the CS fractions and a significantly higher amount of
elastase was present in the aerobic CS than in the anaerobic CS, further validating the
results shown in Fig. 2D. Cell extracts and membrane fractions in both preparations did
not contain either unprocessed or processed elastase. This suggests that the reduced elastase secretion observed during anaerobic growth was due not to incomplete post-translational processes, but to decreased production \textit{per se}.

To examine whether the differential elastase production is mirrored in $\text{lasB}$ transcription, we compared its transcript level in PAO1 grown aerobically versus anaerobically. Since $\text{lasB}$ gene expression occurs in a cell-density dependent manner, bacteria were grown to $\text{OD}_{600}$ of $\sim 3.0$ under both conditions. Fig 3B and C shows $\beta$-galactosidase activity of PAO1 harboring chromosomal copy of $\text{lasB}$ promoter $\text{lacZ}$ fusion and quantitative RT-PCR analysis of $\text{lasB}$ mRNA, respectively. In these two independent assays, mRNA expression of $\text{lasB}$ gene was $\geq 12$-fold and $\geq 40$-fold decreased, respectively in bacteria grown by anaerobic respiration further corroborating that anaerobiosis-induced suppression of elastase secretion is regulated at the transcriptional level.

\textbf{Production of 3-oxo-C12-HSL and C4-HSL was suppressed under anaerobic growth conditions.} Although the LasI/R QS system plays a dominant role in the complex QS hierarchy in \textit{P. aeruginosa} (34), three QS systems, namely LasI/R, RhlI/R, and PQS, all participate in a complex signaling network to regulate $\text{lasB}$ gene expression (29). Recently, it was reported that production of PQS is completely abrogated during anaerobic growth of \textit{P. aeruginosa} (43, 51). Given the fact that synthesis of the PQS signal molecule is coordinately regulated by the LasI/R and RhlI/R components of the QS system (28), this finding suggests that these two QS systems might also be differentially modulated during anaerobic vs. aerobic growth. To gain a better understanding of the molecular basis of the anaerobiosis-induced suppression of $\text{lasB}$ transcription, we measured the levels of 3-oxo-C12-HSL (PAI-1) and C4-HSL.
(PAI-2) in cell-free culture supernatants of PAO1 grown either aerobically or anaerobically. PAI-1 and PAI-2 were quantified using GC/MS and a CV026 reporter-based bioassay, respectively. In our GC/MS analysis using commercially purchased purified PAI-1 as a standard, we found that aerobic culture supernatant contained ~9.72 µM PAI-1. In contrast, only ~0.44 µM was detected in the cell-free supernatant of anaerobic cultures (Fig. 4A). We then measured the level of PAI-2 in cell-free culture supernatants of PAO1 using C. violaceum CV026, a reporter strain that produces violacein in response to exogenously added PAI-2 (27). The level of PAI-2 in aerobic cultures grown to an OD_{600} of ~3.0 was determined to be 1.23±0.05 µM, a value ~7.9 times lower than PAI-1 (Fig. 4A). In contrast, the level of PAI-2 in the cell-free supernatant of an equally dense anaerobic culture was below the detection limits. Together, these results indicate that during anaerobic growth, production of PAI-1 and PAI-2 is highly suppressed, likely rendering P. aeruginosa QS incompetent under such conditions. We then examined whether the substantially decreased production of autoinducers is associated with altered expression of the lasI and rhlI genes. Figure 4B shows that lasI gene expression was similar in bacteria irrespective of the growth conditions, while the expression of rhlI was rather increased in PAO1 grown under anaerobic conditions. This suggests that the dramatically reduced levels of PAI-1 and PAI-2 observed under anaerobic growth conditions are not attributable to transcriptional regulation of the genes involved in their synthesis.

Transcription of most virulence-associated genes was suppressed during anaerobic growth. Next, we sought to elucidate the effect of suppressed synthesis of PAI-1 and PAI-2 on the expression of downstream virulence genes. We tested eight genes reported to be directly regulated by LasR (17). Figure 5A shows the relative expression of these
selected genes as measured by quantitative RT-PCR analysis. Similar to the case of lasB, transcript levels of the mvfR, rsaL, vqsR, and rhlR genes, which encode major regulators in the P. aeruginosa QS system were ~10-fold lower in bacteria grown by anaerobic respiration compared to their aerobically grown counterparts. Transcript levels of rhlA, which encodes rhamnosyltransferase (34) and PA3904, a hypothetical gene with unknown function (45) were ~20% and ~50% of the transcript levels observed under aerobic growth, respectively. In contrast, mRNA levels of PA4677 (45) and xcpP (5) were similar in cells grown under either condition. These results suggest that anaerobiosis downregulated the expression of several, but not all, LasR-regulated genes.

Next, we examined the downstream effects of anaerobiosis-induced suppression of mvfR, a transcriptional regulator of the synthesis of the PQS signal molecule (14). To address this question, we analyzed transcript levels of two genes known to be regulated by MvfR, namely pqsA and pqsC, which encodes enzymes involved in PQS synthesis (56). As shown in Fig. 5B, expression levels of pqsA and pqsC during anaerobic growth were only ~6.7% and ~2% of those during aerobic growth, respectively. This result provides further basis for the significant suppression of PQS synthesis during anaerobic growth.

Addition of autoinducers restored lasB transcription during anaerobic growth. To further verify that anaerobiosis-induced abrogation of lasB transcription was due to an insufficient level of autoinducers, we examined whether lasB transcription was activated by the addition of exogenous autoinducers. Higher levels of lasB transcription, expressed as β-galactosidase activity, were observed after the addition of 10 µM PAI-2 (Fig. 6, first three bars) than PAI-1 or PQS, suggesting that PAI-2 plays a more important role in inducing lasB transcription under anaerobic conditions than the other.
two autoinducers. When pairs of autoinducers were added, the pair of PAI-2 and PQS was more effective at activating lasB transcription than the other two pairs (i.e., PAI-1+PQS and PAI-1+PAI-2), which failed to induce additive β-gal activity (Fig. 6, fourth to sixth bars from left). The highest level of lasB transcription was achieved when all three autoinducers were added; ~74% of the level observed for aerobic growth (Fig. 3B). Again, no significant lasB transcription occurred in the control treatment (bacteria treated with the same concentration of MeOH used to dissolve the signal molecules) (Fig. 6, first bar from right). When the same experiment was repeated under aerobic condition, during which autoinducers are normally produced, no significant changes in β-galactosidase activity were observed in response to the added extraneous autoinducers (Fig. 6B). This result further proves that the anaerobiosis-induced suppression of lasB transcription is due to a lack of autoinducers and also suggests that PAI-1, PAI-2, and PQS are all required for maximal lasB transcription.
Discussion

LasR, the most upstream QS regulon in the *P. aeruginosa* QS hierarchy, regulates the expression of more than 300 virulence-associated genes (17). However, recent genetic studies using diverse *P. aeruginosa* clinical isolates reported that adaptive mutations in the lasR gene occur spontaneously in the course of chronic airway infection in CF (9, 12, 19, 42). Phenotype changes conferred on *P. aeruginosa* by these mutations include (i) facilitated growth on amino acids present in relatively large quantities in CF airways (1, 12), (ii) an efficient shift to an anaerobic mode of growth using nitrate over oxygen (20), and (iii) elevated antibiotic resistance (12, 20). Therefore, frequent identification of lasR mutants suggests that *P. aeruginosa* may acquire the mutation to increase its survival fitness in harsh host environment at the expense of its ability to regulate QS-mediated virulence properties. Furthermore, this notion also supports the idea that QS machinery may be dispensable once chronic infection is successfully established in the patient airway.

Our study was initiated by the observation that cell-free supernatants obtained from anaerobic cultures of PAO1 failed to kill A549 airway epithelial cells, while aerobic culture supernatants were cytotoxic. Because expression of virulence traits is dependent on QS, this result led us to further characterize QS-controlled virulence regulation of *P. aeruginosa* under anaerobic growth conditions. Our subsequent analysis demonstrated that (i) the observed cytotoxicity is mediated by the secretion of elastase (Fig. 2) and (ii) the anaerobiosis-induced loss of cytotoxicity is due to the suppressed level of lasB gene transcription (Fig. 3), which was later found to be mediated by the substantially reduced autoinducer synthesis (Fig. 4) and subsequent decrease in transcription of QS regulators (Fig. 5).
The data presented in Fig. 4A indicate that the concentrations of PAI-1 and PAI-2 in the culture medium during aerobic growth were 9.72 µM and 1.23 µM, respectively, yielding a value of PAI-2/(PAI-1+PAI-2) of ~0.11, consistent with the previous findings of Singh and colleagues (46). In contrast, our data suggested that PAO1 produced significantly suppressed level of PAI-1 and non-detectable PAI-2 during anaerobic growth, respectively. Together with previous reports revealing that the production of PQS is also highly inhibited during anaerobic growth (43, 51), these findings suggest that (i) the three major autoinducers mediating *P. aeruginosa* QS are either not produced or produced at significantly lower levels during anaerobic growth and (ii) QS may therefore not be functional in PAO1 growing under anaerobic conditions. It is of particular interest that the levels of PAI-1 and PAI-2 detected in sputum samples isolated from CF patients colonized with up to $10^8$ CFU/ml of *P. aeruginosa* were significantly lower than those produced during laboratory aerobic cultures (46). Likewise, Erikson and colleagues also reported that autoinducers detected in an independent set of CF sputa were present at very low levels, with concentrations of PAI-2 lower than those of PAI-1 (15). Because anaerobiosis suppresses the production of autoinducers, as was demonstrated in this study, these findings further corroborate that anaerobic respiration is likely a major mode of bacterial growth in the CF mucus (59). Schertzer and colleagues recently elucidated the molecular basis behind the abrogated synthesis of PQS during growth without oxygen (43). Because oxygen and NADH are required as cofactors for the enzymatic synthesis of PQS from its precursor, 2-heptyl-4-quinolone (HHQ), a lack of oxygen prevents the enzymatic conversion of HHQ to PQS from taking place. Thus, the suppressed synthesis of PQS during anaerobic growth is not due to the anaerobiosis-induced altered expression of genes.
whose products are involved in PQS synthesis, but to the absence of molecular oxygen that is physically required for terminal hydroxylation of HHQ (Fig. 7). It appears, however, that molecular oxygen is not directly involved in the synthesis of HSL-based autoinducers. Synthesis of 3-oxo-C12-HSL and C4-HSL by their cognate autoinducer synthase (i.e., LasI and RhlI) requires S-adenosylmethionine (SAM) and 3-oxo-C12-acyl carrier protein or the N-butyrylacyl carrier protein, respectively (18, 38). Our qRT-PCR analysis results shown in Fig. 4B demonstrated that transcript levels of lasI and rhlI were either similar or increased during anaerobic growth compared with those under aerobic growth, suggesting that the anaerobiosis-specific inhibition of autoinducer synthesis is not likely caused by downregulation of associated genes. Interestingly, two independent genome-wide microarray analyses revealed that gene expression of QS-regulated acyl carrier proteins (PA0999, PA1869, PA3333 and PA3334) was invariably decreased during anaerobic growth (16, 52). Although more analysis at the protein level is necessary to allow more robust conclusions, it is likely that suppressed synthesis of PAI-1 and PAI-2 may be due to a limitation of acyl carrier proteins (Fig. 7, red downward arrows).

Our autoinducer “add-back” experiments demonstrated that lasB transcription was restored to intermediate levels by the addition of each of three autoinducer molecules (Fig. 6). These results indicated that (i) virtually none of the autoinducers were produced to sufficient levels during anaerobic growth and (ii) each autoinducer has a distinct role in inducing lasB expression. Consistent with this notion, lasB transcription was restored to its highest level in the presence of all three autoinducers, suggesting that maximal activation of lasB gene transcription is due to the combined effects of the three autoinducers. It is also noteworthy that PAO1 growing anaerobically responded better to
PAI-2 either alone or with PQS to activate lasB transcription (Fig. 6), suggesting that during anaerobic growth, PAI-2-mediated QS may play a more important role than QS induced by the action of other signaling molecules. It was recently reported that PAI-2 production, albeit delayed, still occurs in a lasR mutant strain and that the RhlI/R QS system can override, at least in part, the effects of lasR mutations (13). In addition, comprehensive chronological genetic analysis using a large number of CF isolates revealed that isolates that lost the ability to produce PAI-1 appeared earlier than strains that were unable to produce both PAI-1 and PAI-2 (2). This finding indicates that bacteria may lose the LasI/R QS system more readily than the RhlI/R system during the course of chronic airway infection. Together, these findings suggest that *P. aeruginosa* may have evolved a mechanism by which it can express virulence factors in response to PAI-2 when the LasI/R system is not available.

QS has been studied extensively as an obvious target to alleviate bacterial virulence (37). Such approaches have been considered to be advantageous because targeting QS may not impose selective pressure for the development of resistance as we have witnessed with antibiotics. Our results, however, indicate that QS per se is not actively occurring in *P. aeruginosa* growing in an anaerobic environment. Given the fact that local regions with reduced oxygen tension exist in the CF mucus airway, these results clearly suggest that we need to change the way we understand *P. aeruginosa* pathogenic mechanisms and thus deal with *P. aeruginosa* infections. We anticipate that the data provided in this study will prompt further investigations with the ultimate goal of eradicating this persistent colonizer from anaerobic mucus layers.
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Figure legends

Figure 1. Cytotoxic activity of *P. aeruginosa* culture supernatants towards A549 cells. (A) Relative viability of A549 human epithelial cells treated with cell-free culture supernatants (CS) of PAO1 grown in LB containing 0.4% NO$_3^-$ either aerobically (21% O$_2$) or anaerobically (0% O$_2$). After a 6-hr treatment, 30 µl of MTT reagent (5 mg/ml) was added to the cells and the plate was read at 570 nm. The relative viability is shown as a ratio of OD$_{570}$ values from control media treatment vs. CS treatment. The values shown are the means ± SD from three independent experiments. *p<0.01 vs. treatment with anaerobic CS. (B) Relative viability of A549 cells as assessed by trypan blue staining assay. Experimental conditions for A549 cell growth and treatment were identical to those described for Fig. 1A. Non-viable (i.e., blue-stained) cells were counted and divided by the total number of counted cells. *p<0.01 vs. treatment with anaerobic CS. (C) Morphological changes of A549 cells in response to the treatment with LBN media, aerobic PAO1 CS or anaerobic PAO1 CS. A549 cells were treated for 6 hrs before photos were taken. The images were acquired using a Zeiss Axiovert 200 inverted microscope at a 100× magnification.

Figure 2. Elastase is responsible for A549 cell death and secretion of elastase is significantly repressed during anaerobic growth. (A) Time-dependent accumulation of elastase in the culture supernatant (CS) of aerobic PAO1 culture. Bacteria grown overnight in LB at 37 °C were inoculated at 1:100 in LB + 0.4 % NO$_3^-$, and growth was monitored by measuring OD$_{600}$. Aliquots of the culture were harvested every hour and protein contents present in each CS were analyzed by SDS-PAGE. The protein band that corresponds to the mature elastase was shown with an arrow. (B) Effect of increasing
levels of elastase on A549 cell viability. The viability of A549 cells (1 × 10^4 cells) treated with the same set of CSs for 6 hrs was monitored using an MTT assay as described in Fig. 1. The differences in the mean values among the treatment groups are statistically different (p<0.01, ANOVA). (C) A549 cell viability in response to the CS of a lasB deficient mutant. The mutant was grown aerobically for 8 hrs in LB plus 0.4% (w/v) NO_3^- . Assay conditions were identical to those described in the legend for Fig. 2B.

*p<0.01 vs. treatment with the aerobic CS of ΔlasB mutant. (D) The level of elastase secreted into the culture medium during anaerobic growth. PAO1 were grown in LB plus 0.4% (w/v) NO_3^- inside an anaerobic chamber. The cell density (OD_600) after an 18-hr cultivation in a flask stirred with a magnetic bar to ensure homogeneous mixing was 3.59. The level of elastase was analyzed by SDS-PAGE.

**Figure 3. Anaerobiosis-induced suppression of elastase secretion is controlled at the transcriptional level.** (A) Western blot analysis of elastase in each of three cellular fractions; the culture supernatant (CS), the cell extract (Ext), and the membrane fraction (Mem) of PAO1 grown aerobically (21% O_2) vs. anaerobically (0% O_2). Because the level of elastase secretion is dependent on cell density, bacteria were grown to a similar cell density before harvest (OD_600 of ~3.0) under both conditions. The arrowhead shown to the right indicates elastase. (B) β-galactosidase activity of PAO1 harboring a chromosomal copy of a lasB promoter-lacZ fusion gene. Reporter cells were inoculated in LB plus 0.4% (w/v) NO_3^- and grown to an OD_600 of ~3.0 under both conditions.

*p<0.01 vs. β-galactosidase activity of PAO1 grown aerobically. (C) Quantitative RT-PCR analysis of the expression of the lasB gene encoding elastase in PAO1. qRT-PCR was conducted using cDNA synthesized from 2 µg total RNA extracted from PAO1
grown to an OD$_{600}$ of ~3.0 under both conditions. Transcript levels of the lasB gene were normalized to those of the rpoD gene transcript. *$p<0.01$ vs. lasB transcript level in PAO1 grown aerobically.

**Figure 4. Quantification of PAI-1 and PAI-2 in the culture supernatants of PAO1 growth either aerobically or anaerobically.** (A) Concentrations (in $\mu$M) of two homoserine lactone-based autoinducers (PAI-1 and PAI-2) in aerobic (21% $O_2$) and anaerobic (0% $O_2$) CSs. Bacterial cells were grown to similar final density (i.e., OD$_{600}$ of ~3.0) under both conditions. Autoinducers were extracted using acidified ethyl acetate and PAI-1 and PAI-2 were analyzed with GC-MS/MS and a CV026 reporter cell assay, respectively. Quantification was performed using CSs obtained from three independent cultures and results are displayed as mean±SD. ND stands for “Not Detected”. *$p<0.01$ vs. autoinducer level produced aerobically. (B) mRNA transcript levels of lasI and rhlI in PAO1 cells grown aerobically (black bars) and anaerobically (gray bars) as assessed by qRT-PCR analysis. qRT-PCR was conducted using cDNA synthesized from 2 $\mu$g total RNA extracted from PAO1 grown either aerobically (21% $O_2$) or anaerobically (0% $O_2$). Transcript levels of tested genes were normalized to those of the rpoD gene transcript. Three independent experiments were performed and mean±SD values are displayed in each bar. *$p<0.01$ vs. rhlI transcript in PAO1 grown aerobically.

**Figure 5. Anaerobiosis-induced transcriptional modulation of genes involved in QS regulation.** (A) Quantitative RT-PCR analysis of genes that were previously determined to be directly regulated by LasR. Assay conditions were identical to those described in...
Fig. 4B. Three independent experiments were performed and mean±SD values are displayed in each bar. *p<0.01 vs. transcript levels in PAO1 grown aerobically. 

Downstream effects of suppressed transcription of mvfR on the expression of pqsA and pqsC as assessed by qRT-PCR. *p<0.01 vs. transcript levels in PAO1 grown aerobically.

Figure 6. Exogenously-supplemented autoinducers restore lasB transcription during anaerobic growth. PAI-1, PAI-2, and PQS, either alone or in combinations, were added to the anaerobic (A) or aerobic (B) culture of the P_lasc::lacZ reporter strain at a final concentration of 10 µM. β-galactosidase activity was assessed using cells grown for 18 hrs under both conditions. Three independent experiments were performed and mean±SD values are displayed in each bar. *p<0.01 vs. the other two treatments with PAI-1 (first bar) or PQS (third bar), **p<0.01 vs. the other two treatments with PAI-1/PAI-2 (fourth bar) or PAI-1/PQS (sixth bar), ***p<0.01 vs. all other treatments. Methanol was used as a vehicle control (last bar). p = NS in all comparisons (NS; not significant, ANOVA).

Figure 7. A simplified model of P. aeruginosa QS regulation under anaerobic conditions. During anaerobic growth, production of three major QS signal molecules, PAI-1, PAI-2 and PQS is highly suppressed, rendering P. aeruginosa incapable of QS. Gene names are shown in lower-case italic, while protein names are shown in upper-case roman inside gray square boxes. Downregulated gene transcripts and proteins are shown in squares with dashed lines. Anaerobiosis-induced suppression of lasB transcription (shown in the center) is mediated by inactivation of PAI-1- (top), PAI-2- (left), and PQS-mediated (right) QS. Abbreviations and symbols: SAM, S-Adenosyl-
Methionine; ACP, Acyl Carrier Protein; MTA, 5’-Methylthioadenosine; HHQ, 2-Heptyl-4-Quinolone; ANT, Anthranilate; →, activation or production; suppression, suppressed contribution. The lack of oxygen is denoted by a cross. Dashed black arrows indicate suppressed association of QS signals to their cognate regulator proteins.
Fig. 1

A
Relative viability

C
LBN media 
PAO1 CS, 
21% O 
2
PAO1 CS, 
0% O 
2
B
MTT Relative viability 
Trypan blue 
2
1
%
 
O 
2
0
%
 
O 
2
2
1
%
 
O 
2
0
%
 
O 
2
* 
*
Fig. 2

A

\[ \text{O}_2: \ 21\% \ O_2 \]

\[ \text{Media:} \ \text{LB} + 0.4\% \ NO_3^- \]

\[ \text{OD}_{600}: \ 0.03, 0.07, 0.35, 1.16, 2.54, 3.34, 3.69, 3.86, 3.96 \]

\[ \text{Time (hr):} \ 0, 1, 2, 3, 4, 5, 6, 7, 8 \]

B

Relative viability

\[ \rho<0.001, \ ANOVA \]

C

Relative viability

D

\[ \text{O}_2: \ 21\% \ 0\% \]

\[ \text{Media:} \ \text{LB} + 0.4\% \ NO_3^- \]

\[ \text{OD}_{600}: \ 3.75, 3.59 \]

\[ \text{Time (hr):} \ 18, 18 \]

\[ \text{Elastase:} \ \text{control PAO1 ΔlasB} \]
Fig. 3

β-gal activity
PlasB::lacZ
Relative expression
B C
qRT-PCR
*
21% O2
0% O2
*

Elastase
A

B

C

qRT-PCR

Fig. 3
**Fig. 4**

A B lasI rhlI

Relative expression

ND

PAI-1 PAI-2

Concentrations (in μM) of PAI-1 and PAI-2

21% O

2

0% O

2

21% O

2

0% O

2

*
Fig. 5

**A**

Relative expression

**B**

Figure 5
Fig. 6

**A**

β-galactosidase activity (Miller Unit)

<table>
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<th>PAI-2 (10 μM)</th>
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<tr>
<td>0% O₂</td>
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**B**

β-galactosidase activity (Miller Unit)

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Fig. 7