Oxylipins mediate cell-to-cell communication in *Pseudomonas aeruginosa*

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Oxygenated unsaturated fatty acids, known as oxylipins, are signaling molecules commonly used for cell-to-cell communication in eukaryotes. However, a role for oxylipins in mediating communication in prokaryotes has not previously been described. Bacteria mainly communicate via quorum sensing, which involves the production and detection of diverse small molecules termed autoinducers. Here we show that oleic acid-derived oxylipins produced by *Pseudomonas aeruginosa* function as autoinducers of a novel quorum sensing system. We found that this system controls the cell density-dependent expression of a gene subset independently of the quorum sensing systems thus far described in this bacterium. We identified a LysR-type transcriptional regulator as the primary receptor of the oxylipin signal. The discovery of this oxylipin-dependent quorum sensing system reveals that prokaryote-derived oxylipins also mediate cell-to-cell communication in bacteria.
Bacteria regulate gene expression in response to changes in cell density using a sophisticated cell-to-cell communication process known as quorum sensing. Quorum sensing controls biochemical pathways that are not needed in an isolated individual cell, but become beneficial as part of a population. Diverse quorum sensing systems regulate important biological processes such as bioluminescence, DNA transfer, antibiotic resistance, motility, biofilm formation and virulence. Population density is perceived through the synthesis, release and detection by the bacterial cells of small diffusible molecules referred to as autoinducers. An increase in the bacterial population causes a proportional increase in the extracellular concentration of the autoinducers. Once a threshold concentration is reached, they are detected by quorum sensing signal receptors that trigger a high cell density-specific gene expression program.

In Gram-negative bacteria, N-acyl homoserine lactones (AHLs) are the most common class of autoinducers. However, several bacterial species (Gram-positive and -negative) possess alternative quorum sensing mediators, such as alkyquinolones, α-hydroxyketones, peptides and fatty acid-like molecules. The Gram-negative pathogen Pseudomonas aeruginosa offers one of the best studied models of quorum sensing networks in bacteria. Four interconnected quorum sensing systems have been described in this bacterium thus far: las, rhl, PQS (Pseudomonas quinolone signal) and IQS (Integrating quorum signal). The quorum sensing system las uses 3-oxo-C12-homoserine lactone, while rhl uses N-butyrylhomoserine lactone (both AHLs) as autoinducers. PQS and IQS on the other hand produce and sense the mediators 3,4-dihydroxy-2-heptylquinoline and 2-(2-hydroxyphenyl)-thiazole-4-carbaldehyde, respectively. These quorum sensing systems establish diverse interactions among them, creating a complex regulatory quorum sensing circuitry.

Oxylipins are common cell-to-cell communication mediators in eukaryotes. However, very little is known about the role of prokaryote-derived oxylipins in bacterial physiology. P. aeruginosa possesses a fatty acid dioxygenase that catalyzes the stereospecific oxygenation of exogenous oleic acid. The enzymes responsible for this activity are two fatty acid-dhime cytochrome C peroxidases localized in the periplasm. We recently reported that the oxylipins (10S)-hydroxy-(8E)-octadecenoic acid (10-HOME) and 7S,10S-dihydroxy-(8E)-octadecenoic acid (7,10-DiHOME) generated by the dioxygenase activity are involved in several biological processes, such as motility, biofilm formation and virulence in P. aeruginosa. Compelled by these findings and the role of eukaryote-derived oxylipins in cell-to-cell communication, we investigated whether dioxygenase-derived oxylipins are involved in cell-to-cell communication in P. aeruginosa. We found that P. aeruginosa produces and senses oxylipins in a cell density-dependent manner through a novel quorum sensing system we termed ODS (oxylipin-dependent quorum sensing system). This system operates independently of the hierarchical quorum sensing circuitry of P. aeruginosa. We identified the protein encoded by PA2076 gene of PAO1 as the primary receptor of oxylipins in this bacterium. This protein, which we refer to as OdsR (oxylipin-dependent dioxygenase regulator), is a LysR-type transcriptional regulator (LTTR) that mediates oxylipin-dependent induction of the dioxygenase enzymes. The discovery of ODS reveals for the first time that prokaryote-derived oxylipins are also signaling molecules mediating cell-to-cell communication in bacteria.

**Results**

A positive regulatory loop controls the dioxygenase operon expression. The dioxygenase enzymes of P. aeruginosa are encoded by the PA2077 and PA2078 genes, which together form an operon (Supplementary Fig. 1). Once expressed, these enzymes localize mainly in the periplasm. We found that addition of OA to the culture was required to isolate a periplasmic fraction of P. aeruginosa displaying dioxygenase activity in vitro (Fig. 1a). This observation suggested that expression of the dioxygenase enzymes is dependent on exogenous OA. To confirm this we used P. aeruginosa PAO1 strain containing a genetic fusion between the dioxygenase promoter and the Escherichia coli lacZ reporter gene cloned into plasmid pDSp-lacZ (Supplementary Table 1). The β-galactosidase (β-gal) activity in this strain was dependent on the addition of OA to the medium (Fig. 1b). Surprisingly, when pDSp-lacZ was introduced in a dioxygenase-lacking background strain, the OA failed to fully induce the expression of β-gal activity (Fig. 1b, third column). This result indicated that oxylipins derived from the dioxygenase operon on OA were required to fully induce expression of the dioxygenase operon. We next tested the ability of 10-HOME and 7,10-DiHOME oxylipins, purified from a PAO1 culture supernatant, to induce the β-gal activity in ΔDS (pDSp-lacZ). Each oxylipin, used at 0.1 mg mL\(^{-1}\), induced the β-gal activity in ΔDS (pDSp-lacZ) more than fivefold greater than OA used at the same concentration (Fig. 1c). Altogether, these data indicate that production of oxylipins in P. aeruginosa is regulated by a positive regulatory circuit in which oxylipins induce full expression of their own biosynthetic enzymes.

Oxylipin production depends on cell density. Positive auto-regulatory circuits, in which the autoinducer molecules positively regulate their own synthesis in a cell density-dependent manner, are characteristic of quorum sensing systems. Based on these data we reasoned that oxylipin accumulation in the supernatant of PAO1 could follow a production kinetics similar to that of quorum sensing autoinducers. To test this, we performed a kinetic experiment to follow the accumulation of oxylipins in the supernatant of PAO1 cultured in M63 medium supplemented with 1 mg mL\(^{-1}\) of OA. No detectable amounts of oxylipins were found in the first 3 h of growth (exponential phase), whereas the concentration of oxylipins rapidly increased after 4 h of growth (early stationary phase) (Fig. 2a). In addition, after 6 h, when the OA was mostly consumed, the concentration of 10-HOME rapidly decreased, while the 7,10-DiHOME oxylipin remained longer in the culture. This experiment suggested that production of 10-HOME and 7,10-DiHOME oxylipins is regulated in a cell density-dependent manner, although 7,10-DiHOME had a slower kinetics of consumption compared to 10-HOME (Fig. 2a). To determine the level of the dioxygenase operon expression in relation to the cell density, the kinetics of β-gal activity in PAO1 (pDSp-lacZ) was monitored. The expression of β-gal activity was evident at the early stationary phase of the culture, rapidly increasing (from 4 to 5 h), following a sigmoid curve (Fig. 2b, in dark blue). When the same experiment was done with ΔDS (pDSp-lacZ), there was poor expression of the dioxygenase operon (Fig. 2b, in pale blue). These results suggest that expression of the dioxygenase operon is regulated in a cell density-dependent manner through a positively autoregulated circuit mediated by 10-HOME and 7,10-DiHOME.

PA2076 protein controls the expression of the dioxygenase operon. A positive autoregulatory circuit usually involves a transcriptional regulator that binds the inducer molecule and in turn activates the expression of the inducer biosynthetic enzymes. In many cases, the transcriptional regulator-encoding gene localizes near the genes encoding the autoinducer biosynthetic enzymes (e.g., LasR–LasI and RhlR–RhlI pairs in P. aeruginosa or LuxR–LuxI in Vibrio fischeri). In PAO1, the gene PA2076, encoding a putative
LTTR, localizes downstream of the diol synthase operon (Supplementary Fig. 1A). In order to determine if the product of PA2076 was involved in regulation of the diol synthase operon, a PA2076 deletion mutant was created (ΔPA2076). It was found that ΔPA2076 failed to produce oxylipins in M63 supplemented with OA (Fig. 3a). To rule out any polar effect potentially caused by PA2076 deletion on the contiguous diol synthase operon, PA2076 protein was expressed in trans from plasmid pBB-odsR-His. The ability of the complemented APA2076 strain to produce oxylipin recovered (Fig. 3a). In addition, it was shown that ΔPA2076 (pDSp-lacZ) did not express detectable amounts of β-gal activity, indicating that PA2076 is required for expression of the diol synthase operon (Fig. 3b). The β-gal activity in ΔPA2076 (pDSp-lacZ) was restored by reintroducing PA2076 into the chromosome. These results suggest that PA2076 encodes a positive transcriptional regulator (hereafter termed OdsR) of the operon. An electrophoretic mobility shift assay (EMSA) using a purified His-tag version of OdsR (OdsR-His) showed that OdsR specifically bound to a DNA probe consisting of the 200 bp sequence immediately upstream of the first gene (PA2078) of the diol synthase operon, but not to an unrelated DNA probe used as control (Fig. 3c, Supplementary Fig. 2).

Some transcriptional regulators involved in positive autoregulatory feedback are able to induce their own expression in the presence of their respective autoinducers (e.g., LasR and RhlR). To assess whether OdsR was induced by OA or its derivative oxylipins, a His-tag version of the odsR gene was cloned in the pBBR1MCS vector under control of its native promoter, yielding the plasmid pBB-odsR-His (Supplementary Table 1). A western blot analysis using an anti-poly-histidine antibody indicated that PAO1 (pBB-odsR-His) produced OdsR-His at the same level in the presence or absence of OA. This suggests that OdsR is constitutively expressed in P. aeruginosa (Supplementary Fig. 1B). The EMSA experiment also showed that OdsR binds to the diol synthase operon promoter in the presence or absence of the oxylipins. These results suggest that upon binding to the OdsR-DNA complex, the oxylipins induce a conformational change that allows the RNA polymerase to transcribe the diol synthase operon. This is similar to the mechanism of action of other known LTTRs[20].

LTTRs contain a DNA binding domain and a regulatory domain formed by two subdomains. The inducer molecule usually binds to the region between the two subdomains,
inducing a conformational change of the protein that activates gene expression\textsuperscript{19}. We used the online I-TASSER server to predict the three-dimensional (3D) structure of OdsR. It predicted a large hydrophobic pocket in the inter-subdomain region of the regulatory domain that might serve as the binding site for the oxylipins (Fig. 3d). In agreement with this hypothesis, it was determined that three independent mutations (Y98Q, L143Q and L245T), in which three non-adjacent hydrophobic amino acids in this region were replaced by hydrophilic amino acids, abolished the oxylipin-dependent induction of the diol synthase activity in PAO1 (Supplementary Fig. 1C). As a control, mutagenesis of an amino acid localized outside of the predicted hydrophobic pocket had no effect on the diol synthase activity (Supplementary Fig. 1C). These results suggest that the hydrophobic pocket found in the predicted structure of OdsR is an oxylipin binding site.

**Oxylipins control a gene subset expression.** Previously it was found that 10-HOME and 7,10-DiHOME oxylipins play a role in several physiological processes in *P. aeruginosa*\textsuperscript{18}. However, the specific metabolic pathways regulated by oxylipins in *P. aeruginosa* remain unknown. To identify proteins specifically induced by oxylipins, a comparative proteomic analysis of PAO1 and ∆odsr, both grown in the presence and absence of OA or each of the oxylipins, was performed. In general, we identified 17 proteins that were induced (excluding PA2077 and PA2078) and 16 proteins that were inhibited more than fivefold by 10-HOME and/or 7,10-DiHOME in the proteomic analysis (Supplementary Table 2). As expected, oxylipin-dependent induction of the diol synthase enzymes (PA2077 and PA2078) occurred in PAO1, but not in ∆odsr. In fact, no other protein, apart from PA2077 and PA2078, seems to be directly regulated by OdsR. This observation suggests that OdsR is restricted to regulating the diol synthase operon and therefore controls the amount of oxylipins produced in a cell density-dependent manner. However, a set of proteins were up- or down-shifted fivefold or more in both PAO1 and ∆odsr when treated with oxylipins. These findings suggest that the oxylipins regulate the expression of these proteins through mediation of other as yet unidentified transcriptional regulators that use 10-HOME or 7,10-DiHOME oxylipins as inducers.

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**Fig. 3** OdsR (oxylipin-dependent diol synthase regulator) is an oxylipin-dependent inducer of the diol synthase operon. **a** Production of oxylipins in the supernatant of PAO1, ∆PA2076 and ∆PA2076 complemented with a copy of odsR gene (∆PA2076/pBB-odsR-His). ∆PA2076 failed to produce oxylipins. **b** β-Galactosidase (β-gal) expression in PAO1 (pDSp-lacZ), ∆PA2076 (pDSp-lacZ) and ∆PA2076 complemented with PA2076 (∆PA2076 (pDSp-lacZ+pBB-odsR-His)). ∆PA2076 (pDSp-lacZ) did not express detectable β-gal activity in the presence of oleic acid (OA; 1 mg mL\textsuperscript{−1}) (t-test, P < 0.0001). **c** Electrophoretic mobility shift assay (EMSA) gel showing specific binding of OdsR protein to the diol synthase promoter probe (Dspr, 200 bp), but not to a control unrelated DNA probe (CTL, 150 bp). **d** I-TASSER predicted model of OdsR three-dimensional (3D) structure. The hydrophobic pocket localized in between the two regulatory subdomains is indicated. The hydrophobic amino acids in the pocket that were replaced by hydrophilic counterparts are represented in red. These changes abolished the ability of OdsR to induce the β-gal activity of the DSp-lacZ genetic fusion. The amino acid L91 (in blue) outside of the pocket had no effect on OdsR function. Data from a, b are the means and s.d. of three independent experiments. ****Significantly different, unpaired two-tailed t-test, P < 0.0001
We also analyzed the transcriptome of ΔodsR in the presence or absence of OA or each of the oxylipins. We identified 29 genes that were induced by oxylipins and 10 genes that were repressed fivefold or more (Supplementary Table 3). As expected, most of the proteins identified as regulated by oxylipins were identified in both the proteomics and the transcriptomic studies (Supplementary Tables 2 and 3). In addition, we identified differences in the proteome and the transcriptome analysis that might reflect the sensitivity of each method and/or the specificities in posttranscriptional regulation. In both studies we found a stronger effect of 10-HOME than that observed for 7,10-DIHOME.

To confirm the validity of the -omics studies, lacZ was transcriptionally fused with the promoter of PA3427 as a representative gene of those strongly induced by the oxylipins. PAO1 strain containing this fusion [PAO1 (PA3427p-lacZ)] expressed β-gal activity only when grown in the presence of OA (Supplementary Fig. 3). However, ΔodsR (PA3427p-lacZ) grown in OA displayed very poor β-gal activity. This observation validated the findings of the proteome and the transcriptome studies by confirming that at least the PA3427 gene is up-regulated by the oxylipins derived from the diol synthase activity. It was also shown that purified 10-HOME and 7,10-DIHOME induced β-gal activity in both strains, PAO1 (PA3427p-lacZ) and ΔodsR (PA3427p-lacZ) (Supplementary Fig. 3), indicating that PA3427 is regulated by oxylipins independently of OdsR and through an, as of yet, unidentified transcriptional regulator.

Oxylipins function as quorum sensing signals. The observation that P. aeruginosa produces 10-HOME and 7,10-DIHOME oxylipins in a cell density-dependent manner suggests that those genes outside of the diol synthase operon and regulated by oxylipins should also follow a cell density-dependent expression pattern. To evaluate this hypothesis, an expression kinetics assay was performed by following the β-gal activity in PAO1 (PA3427p-lacZ). This strain expressed β-gal activity in a cell density-dependent manner, correlating with the level of oxylipins in the culture supernatant (Fig. 4a). This effect was similar to that observed with quorum sensing-regulated genes. For example, PA3427p-lacZ was prematurely induced when free-cell supernatants of stationary cultures of PAO1 (containing high level of oxylipins) were added to fresh cultures at 4 h (Fig. 4b). As expected, this was not observed when free-cell supernatants of ΔDS cultures at stationary phase were added (Fig. 4b). In addition, pure oxylipins, 10-HOME or 7,10-DIHOME, prematurely induced β-gal activity when added to PAO1 (PA3427p-lacZ) cultures at early exponential phase (Fig. 4c). Altogether, these results demonstrate that oxylipins produced by P. aeruginosa are autoinducers of an oxylipin-dependent quorum sensing system that we refer to as ODS.

The quorum sensing signal molecules are “social public goods” that can be exploited by signal-producing deficient mutants. For example, a lasR mutant of P. aeruginosa unable to produce 3O-C12-HSL responds to the signal produced by a wild-type strain in co-culture21. This process is relevant as a signal-deficient strain can act as a “social cheater” in an environment where quorum sensing is required to grow23. Thus, we explored the ability of oxylipin-deficient strains (ΔDS or ΔodsR) to exploit the oxylipin signal produced by an oxylipin-proficient strain. For this purpose we co-cultured ΔDS (PA3427p-lacZ) with PAO1 and ΔodsR (PA3427p-lacZ) with PAO1 to measure the ability of the oxylipins produced by PAO1 bacterial cells to induce the β-gal activity in the neighbor cells of the oxylipin-deficient strains. As expected, both ΔDS (PA3427p-lacZ) and ΔodsR (PA3427p-lacZ) were able to sense the oxylipins produced by PAO1 and expressed a level of β-gal activity similar to that observed for PAO1 (Fig. 5a, b, respectively). These observations are in agreement with our previous results reporting that ΔDS profit from the extracellular oxylipins produced by PAO1 to complement its biofilm deficiency when they are co-cultured16.

ODS is a self-regulated quorum sensing system. P. aeruginosa is one of the more intensively studied model of quorum sensing systems in bacteria. In addition to the ODS system described here, this bacterium contains at least two AHL-dependent quorum sensing systems, las and rhl, and a 4-quinolone-dependent quorum sensing system, PQS11. These systems are interconnected forming a complex quorum sensing circuitry. In order to determine if ODS is a self-regulated system or if it is dependent of one of the above-mentioned quorum sensing systems we measured the kinetic of oxylipin production in transposon insertion mutants of the genes encoding LasR, RhlR and MvfR, which are the autoinducer receptors of las, rhl and the PQS systems, respectively. While the production of oxylipins by rhlR::tn showed a normal kinetic compared to PAO1, lasR::tn and mvfR::tn showed a slower kinetics of 10-HOME or 7,10-DIHOME production (Fig. 6a, b). However, although slower, the production of 10-HOME and 7,10-DIHOME in lasR::tn and mvfR::tn was still regulated in a cell density-dependent manner, demonstrating that ODS is, to a great extent, a self-regulated system.

Discussion
Herein, we report a novel quorum sensing system in P. aeruginosa that uses oxylipins as autoinducers and regulates the
expression of a gene subset in a cell density-dependent manner. The working model of ODS function is shown in Fig. 7. Based on these results, we propose a model in which the OdsR receptor is expressed constitutively. When P. aeruginosa encounters an environment where OA is present, this molecule directly or indirectly induces a basal expression of the diol synthase operon at low cell density. This basal induction causes a basal production of oxylipins. When the cell density increases, a certain oxylipin concentration threshold is achieved at which oxylipins bind OdsR and further activate the diol synthase operon. This positive feedback of the diol synthase operon expression causes a sudden increase in the amount of

Fig. 5 Oxylipin-deficient P. aeruginosa profit from oxylipins produced by PAO1 wild-type strain. a Graph showing the β-galactosidase (β-gal) activity of ΔDS (PA3427p-lacZ) grown alone or in co-culture with PAO1. b Graph of the same experiment, but using ΔodsR (PA3427p-lacZ)

Fig. 6 Oxylipin-dependent quorum sensing system (ODS) is self-regulated. a Time course of 10-HOME and b 7,10-DiHOME production in the supernatants of PAO1 and the transposon mutants lasR::tn, rhlR::tn and mvfR::tn. The mutants rhlR::tn and mvfR::tn showed a slower kinetics of 10-HOME and 7,10-DiHOME production. Means and s.d. are from three independent experiments. 10-HOME (10$^S$)-hydroxy-(8$^E$)-octadecenoic acid, 7,10-DiHOME 7$^S$,10$^S$-dihydroxy-(8$^E$)-octadecenoic acid

Fig. 7 Model of the oxylipin-dependent quorum sensing system (ODS). a The OdsR (oxylipin-dependent diol synthase regulator) receptor (represented as a blue square) is expressed constitutively. At low cell density in the presence of oleic acid (OA), this molecule (or an unknown derivative molecule) binds OdsR inducing a basal expression of the ODS operon, shown with a single plus symbol above the DS promoter (DSp). P. aeruginosa grown under these conditions produces a small amount of oxylipins. b When the cell density increases, the oxylipins reach a threshold concentration at which they bind to OdsR. OdsR in turn induces the expression of the DS enzymes at a higher rate than that induced by OA (shown with a triple plus symbol), creating a positive regulatory feedback that further increases the extracellular concentration of oxylipins. Subsequently, the oxylipins are sensed by other as yet unidentified secondary receptor(s) (shown as a green circle) that ultimately regulate the expression of the ODS effector genes
extracellular oxylipins, which in turn activates the expression of the ODS effector genes.

The effector genes identified in this work using -omics approaches include those encoding several putative oxidoreductases, dehydrogenases, a protein kinase (PA0588), a putative aminoglycoside phosphotransferase (PA1829) and a nitric oxide reductase (PA0523 and PA0524), among others (Fig. 8, Supplementary Tables 2 and 3). The roles of each of these proteins in the physiology of P. aeruginosa will require further extensive analysis, but their putative functions suggest that they might be involved in adaptation of P. aeruginosa to the host environment, resistance to oxidative stress, nitric oxide toxicity and antibiotics (Fig. 8). We have demonstrated that production and sensing of oxylipins regulates bacterial motility and thus biofilm formation, as well as virulence. However, we did not identify any gene evidently related to biofilm or motility in the -omic studies. This indicates that either these processes are regulated post-transcriptionally by the oxylipins (e.g., the induction of small regulatory RNAs is a possibility) or that some of the identified hypothetical proteins might be directly involved in the regulation of motility.

The ODS system differs from other quorum sensing systems in that it uses a transcriptional regulator (OdsR) to regulate the level of oxylipin production and at least one other receptor to sense the level of oxylipins, and subsequently induces the expression of the transcriptional regulator that serves as receptor of the auto-inducer signal is the same that regulates the expression of the effector genes.

ODS operates as an environment-specific quorum sensing system since it requires exogenous OA as a precursor for the biosynthesis of the autoinducers. This is an interesting strategy through which P. aeruginosa avoids responding indiscriminately to high cell density by regulating the appropriate subset of genes required for a particular niche. OA is a fatty acid present in specific niches, such as decaying biomass and within living organisms. This suggests that ODS plays an important role in vivo during bacterial infections. This is in agreement with our previous results indicating that P. aeruginosa scavenges OA from the tissues of infected Drosophila melanogaster to produce oxylipins, which in turn promotes bacterial virulence.

The finding that P. aeruginosa can regulate cell density-dependent expression of the diol synthase operon even in LasR-, RhlR- and MvfR-deficient strains demonstrates that ODS is to a great extent a self-regulated quorum sensing system, and therefore abrogation of the other quorum sensing systems of P. aeruginosa do not annul ODS functionality. However, this does not preclude the existence of regulatory interactions of ODS with other quorum sensing systems of P. aeruginosa. Indeed, we found a delay in the kinetics of oxylipin production in the lasR and mvfR mutants (Fig. 5), which suggests that ODS interacts with the las and mvf quorum sensing systems at a genetic or posttranscriptional level. Due to the high complexity of the quorum sensing circuitry of P. aeruginosa, the determination of ODS-specific interactome with the quorum sensing network of this bacterium requires extensive investigation which is the target for future research directions.

Fig. 8 The oxylipin-dependent quorum sensing system (ODS) regulon. Categorization of genes regulated by the oleic acid (OA)-derived oxylipins in P. aeruginosa according to their hypothetical assigned functions in the Pseudomonas Genome Database (http://www.pseudomonas.com). Positive regulation is indicated using red arrows and negative regulation by green flat bars.
This study provides the first evidence of a quorum sensing pathway regulated by oxylipins in bacteria. The fact that ODS has not been previously identified in the widely studied model of quorum sensing, P. aeruginosa highlights the importance of approaching the study of quorum sensing under conditions that better reproduce the bacterial natural environments. We believe that oxylipins could mediate cell-to-cell communication in other bacterial species, since we have identified genes homologous to PA2077 and PA2078 in other bacterial species, not only inside the genus Pseudomonas, but also in species from diverse genera, such as Acinetobacter baumannii, Methylocapsa palsarum, Massilia eurypsychrophila and Paracoccus sanguinis (see phylogenetic trees in Supplementary Fig. 4 and 5). Other genes encoding potential oxylipin synthases, such as lipoxigenases, di-heme cytochrome C peroxidases and cytochromes P450, are also widely distributed in bacteria24–26. Interestingly, it was recently reported that the bacterial plant pathogen Xylella fastidiosa produces 10-HOME and 7,10-DiHOME in vivo when inoculated in the plant model Nicotiana tabacum27. These observations suggest that production of oxylipins in bacteria is not confined to P. aeruginosa and thus a more general role of oxylipins in bacterial cell-to-cell communication is plausible. Yet, the discovery of ODS represents another layer of complexity in the already intricate quorum sensing circuitry of P. aeruginosa and a new potential target for the development of antimicrobial drugs aimed to interfere in the infection process of this important multi-drug-resistant pathogen.

Methods

Strains, plasmds and oligonucleotides. Strains, plasmids and oligonucleotides used in this study are described in Supplementary Table 1.

Culture conditions. The strains were routinely grown in lysogeny broth (LB) medium at 37 °C, to which agar was added when solid medium was required. LB agar without NaCl plus 15% sucrose was used to segregate suicide plasmids from merodiploids during construction of ΔodsR and ΔlasR mutants by allelic exchange (see below). When required, P. aeruginosa was grown in M63 media supplemented with 0.2% glucose, 0.1% casamino acids and MgSO4, 1 mM (M63 complete). Antibiotics were added at the following concentrations: Ampicillin (Amp), 100 μg mL−1; Carbenicillin (Cb), 300 μg mL−1; P. aeruginosa; Kanamycin, 25 μg mL−1; for E. coli, and 200 μg mL−1 for P. aeruginosa; Kanamycin, 25 μg mL−1; OA 99% (Sigma) was added to cultures for oxylipin production and purification. M63-complete media were supplemented with OA 99% (Sigma) or purified oxylipins at the specified concentrations.

Genetic constructions. Plasmid pBB-odsR-His was constructed by inserting PA2076 gene and its promoter region into the pBB1R1MCS plasmid28. The PA2076 DNA fragment was amplified using primers PA2076-Stul and PA2076-KanStul (Supplementary Table 1) and PA01 chromosomal DNA as a template. The primers inserted Stul and HindIII restriction sites at the extremes of the amplified fragment, which were used to insert the fragment into pBB1R1MCS digested with HindIII and Smal. Primer PA2076-HindIII also introduced six histidine codons (coding a His-tag) at the 3′ end of the gene. Site-directed mutagenesis of odsR was performed by reverse PCR. Whole plasmid pBB-odsR-His sequence was amplified with oligonucleotides that anneal adjacent and divergently from each other at the sequence inside PA2076 to be mutagenized. Primer pairs used for the mutagenesis are shown in Supplementary Table 1 (primers from 191791-F through 2Y01Q-R). One of the oligonucleotide pairs contained the desired base change(s) (capitalized bases, Supplementary Table 1) to change the targeted amino acid. The amplified fragment was then self-ligated resulting in a plasmid with identical sequence to the original except for the desired mutation inside odsR. For deletion of PA2076 from PA01 chromosome, this gene was first amplified using primers PA2076-F-EcoRI and PA2076-HindIII (Supplementary Table 1), as well as PA01 genomic DNA as a template (Forward primer for PCR amplification using primers PA2076-Stul and PA2076-HindIII digested and inserted between same sites in pEX100Tslink vector29 was used to obtain pEX-PA2076. DNA of this plasmid was used as a template for reverse PCR amplification using primers PA2076-BamHI and PA2076-BamHI. Amplified fragment was BamHI digested and self-ligated, resulting in plasmid pEX-PA2076, which was used with an internal in-frame deletion. This suicide plasmid was used to delete PA2076 from PA01 chromosome by allelic replacement as previously described28.

Plasmid pET22b-odsR was ordered from GenScript. It contains an E. coli codon optimized odsR gene with a C-terminal His-Tag, which was synthesized by the vendor. For deletion of lasR from PA01 chromosome, this gene was amplified using primers lasR-HindIII and lasR-EcoRI (Supplementary Table 1), as well as PA01 genomic DNA as a template. Amplified fragment was EcoRI–HindIII digested and inserted between same sites in pEX100Tslink suicide vector to obtain pEX-lasR. DNA of this plasmid was digested with PstI, which cuts two times inside lasR, and then self-ligates, which results in plasmid pEX-lasR, which contains lasR with an internal in-frame deletion. This suicide plasmid was used to delete lasR from PA01 chromosome by allelic replacement as previously described28. Plasmid pBB-P3437γ-lacZ was fused to the promoter of gene PA3427. To construct this plasmid the lasZ gene from plasmid pIT2 (Manoil lab, University of Washington) was PCR-amplified using primers lacZ-5′-Xhol and lacZ-3′-BamHI (Supplementary Table 1) that introduced Xhol and BamHI restriction sites at respective ends of amplified DNA fragment, which was inserted into the same sites of pBB1R1MCS plasmid to obtain pBB-lacZ. The PA3427 promoter was PCR-amplified from PA01 chromosome using oligonucleotides PA3427-BamHI and PA3427-Xhol and the resulting 333 bp DNA fragment was inserted between BamHI/Xhol-digested pBB-lacZ, which positioned PA3427 promoter correctly oriented in front of lacZ.

Purification of 10-HOME and 7,10-DiHOME oxylipins. PA01 was plated in LB agar and incubated overnight at 30 °C. The bacterial biomass was scraped from the plate and used to inoculate 200 mL of M63 complete supplemented with 1% oleic acid. Production of oxylipins was followed by taking 0.5 mL aliquots of the culture every 30 min, and extracting the lipids with ethyl acetate from hydrochloric acid acidified (pH = 2) supernatant aliquots (aqueous were removed by centrifugation). Production of each aliquot was then analyzed to obtain mass chromatograms using a Quattro Premier XLS mass spectrometer (TCL) until most of the OA was converted into the oxylipins. The culture was then used to purify the oxylipins. First, the culture was centrifuged at 8000 × g for 15 min to remove bacterial cells. The supernatant was recovered and acidified (pH = 2) with hydrochloric acid. Then, a vol/vol organic extraction with ethyl acetate was carried out and the organic phase was evaporated. The dried mixture obtained was dissolved in 3 mL of ethyl acetate and used for purification of the oxylipins on an Isco Teledyne Combiflash RF 200 with four channels with 300CF ELSD (evaporative light scattering detector). Universal RediSep solid sample loading pre-packed cartridges (30 g silica) were used to absorb the crude product and eluted with 24 g silica RediSep RF Gold Silica (20–μm silica) columns using an increasing gradient of ethyl acetate (solvent B) over hexane (solvent A). Fractions collected for each detected peak were combined and evaporated, then dissolved in ethanol.

The purity of the oxylipins was analyzed by high-performance liquid chromatography and mass spectrometry. Briefly, the purified 7,10-DiHOME and 10-HOME oxylipins were dissolved at 1 mg mL−1 in methanol (stock solution). From these solutions were prepared by diluting in ddH2O 0.1% formic acid. A 20 μL aliquot of each sample was loaded onto a Synergi Hydro-RP 80 Å 250 × 2 mm C18 column (Phenomenex) using a Shimadzu Prominence liquid chromatography system. The mobile phase composition was as follows: 60% A (acetonitrile in deuterated water), 40% B (acetonitrile in ddH2O) at 600 °C, respectively. The gradient proceeded from 10–80% B over 11 min, then to 100% B at 14 min, then re-equilibrated back at initial conditions for 6 min for a total of 20 min per evaluation using the SCIEX 4000Q TRAP Quadrupole Mass Spectrometer (Concord, Ontario, Canada). The mass chromatogram (ESI-negative mode) of deuterated OA (20–μm silica) in the electrospray ionization (ESI)-negative ion mode. Nitrogen was used as a nebulizer and curtain gas (CUR = 20). The collision gas, collision energy and temperature were set at 10 (30 eV for 10-HOME, −34 eV for 7,10-DiHOME) and 600 °C, respectively. GS1 and GS2 were set at 40 and 60 respectively. Analyst 1.6.2 software controlled the liquid chromatography-tandem mass spectrometry system.

Thin layer chromatography. TLCs were run on 60 Å silica gel plates of 20 × 10 cm and 200 μm thickness (Whatman®). The mobile phase solvent was a mix of hexane, ether and acetic acid (80:20:5). TLC plates were revealed with 10% phosphomolybdic acid in ethanol and dried with a hair dryer.

Proteomics analysis. P. aeruginosa PA01 and ΔodSR strains were grown in quadruple in M63 complete medium (3 mL) up to an OD600 = 1 at 30 °C and shaking (240 rpm). Each strain was then treated with 0.1 mg mL−1 of OA, 10-HOME or 7,10-DiHOME oxylipin or kept untreated as baseline control and incubated for additional 2 h at 30 °C and shaking (240 rpm). Bacterial cells were collected by centrifugation (10,000 rpm, 5 min) from 1 mL aliquots of each culture and freeze dried. Bacterial pellets were then thawed and lysed with 20 μL of NAP-20 column buffer, reduced, denatured and separated by one-dimensional polyacrylamide gel electrophoresis (1D PAGE) as previously referenced30. This step was carried out on an SDS Bis-Tris gel (10%, Invitrogen) as per the manufacturer’s instructions. The gels were stained, the entire lane for each sample was excision into 3 MW fractions and each gel plug was equilibrated in 100 mM ammonium bicarbonate (AmBc). Each gel plug was then digested with Trypsin Gold (Promega) following the manufacturer’s instruction, and peptide
extracts were reconstituted in 0.1% Formic Acid/diH2O at −0.1 µg µL−1. Mass spectrometry runs were carried out, and the data processed, searched, filtered, grouped and quantitated as previously reported (Ludwig et al.32; under section 2.5 NLC-ESI-MS2 and Protein IDs for GelCG31).

RNA sequencing and transcriptomics analysis. P. aeruginosa PA01 strain was grown in quadruplicates in M63 complete medium (3 mL) up to an OD600 = 1 at 30 °C and shaking (240 rpm). The cultures were then treated with 0.1 mg mL−1 of OA, 10-HOME or 7,10-DiHOME or treated with the vehicle only (ethanol) as baseline control and were harvested for additional 3 h at 30 °C and shaking (240 rpm). This was done in duplicates. Bacterial cells were recovered and used to isolate total RNA using a QIAGEN RNeasy Plus Mini Kit (cat. no. 74134). Concentration and purity of RNA samples were determined using the absorbance (A260/A280) using a NanoDrop 2000 spectrophotometer (ThermoFisher Scientific). Total RNA samples were then processed for the depletion of ribosomal RNA (rRNA) following the manufacturer’s user manual for Ribo-Zero RNA removal kit for bacteria from Illumina (Document no. 15066012 V02). RNA libraries were prepared following the Lexogen’s user manual of Sense Total RNA-seq library preparation kit (cat. nos. 009.24, 020.24, 022.24). Briefly, 10 ng of rRNA-depleted RNA was used as the starting material for the generation of libraries. The library generation started with the random hybridization of the starter/stopper heterodimer mix to the rRNA-depleted RNA. These starter/stopper heterodimers contained Illumina-compatible linker sequences. A single-tube reverse transcription and ligation reaction was then performed to expand the starter to the next hybridized heterodimer, where the newly synthesized complementary DNA insert was ligated to the stopper. The second strand synthesis was then performed to hydrolyze the RNA and then the library was converted to double-stranded DNA, which was purified using magnetic beads to adjust the library length and to ensure the complete removal of second strand synthesis reaction components. The library amplification was then performed to add the complete adaptor sequences required for cluster generation and to produce sufficient material for quality control and sequencing; 17 indices were added during this step in order to uniquely multiplex the samples for the sequencing run. The resulting RNA libraries were then purified using magnetic beads to get rid of PCR components, which could interfere with quantification and sequencing on an Illumina MiSeq instrument. Bioinformatics Analysis of RNA-seq was performed using FASTQ files from the MiSeq sequencer were uploaded to Qiagen’s GeneGlobe Data Analysis Center, where the raw sequence reads were aligned to the reference genome and transcript abundances were estimated. These transcript abundances were normalized using their Average Reference Genes (Molecular Tags option and differential expression was calculated from these normalized values. For generating heatmaps and principal component analysis plots, ClustVis was used32.

OdsR protein purification. Plasmid pET22b-odsR was transformed into E.coli BL-21 DE3 cells. A single colony was inoculated in LB medium overnight at 37 °C. A total of 5 mL of the culture was added into 500 mL of autocluded medium. The cultures were grown at 37 °C for 4 h, followed by overnight induction at 18 °C. The cells were harvested and cell pellets were resuspended in lysis buffer (20 mM Tris, 50 mM NaCl, 20 mM imidazole, 0.5 mM TCEP, pH 8.0) with 1 mg mL−1 lysozyme and 1 tablet of SIGMAFAST protease inhibitor (Sigma-Aldrich). The cell suspension was sonicated on ice and centrifuged at 15,000 rpm for 1 h. The supernatant was mixed with 1 mL Ni-NTA resin (GE Healthcare Life Sciences) and shaken for 2 h. The Ni-NTA resin was spun down and loaded onto an empty column, which was washed successively with lysis buffer and lysis buffer containing 40 mM imidazole and 60 mM NaCl. OdsR-His protein was eluted in 400 mM imidazole prepared in lysis buffer and further purified by size-exclusion chromatography (SEC) using a Superdex 200 column (16/600, GE Healthcare Life Sciences) in the SEC buffer (20 mM Tris, 150 mM NaCl, 1 mM TCEP, pH 8.0). The fractions containing Ods-His protein were pooled and concentrated to about 8 mg mL−1. The protein solution was aliquot and stored at −80 °C freezer.

Western blotting. Protein expression of PA2078-His (OdsR-His) or its derived single amino acid mutants was studied by western blotting using 1 µg µL−1 of Anti-6X His tag® antibody (HRP) conjugate (Abcam) as previously described34 with some modifications. Briefly, proteins were electrophoresed in 4–12% Bis-Tris Plus gels as per the manufacturer’s instructions (Invitrogen) and the resolved proteins were transferred to a polyvinylene difluoride membrane by electroblotting using an iBlot2 apparatus (Invitrogen). The membrane was transferred to an iBind apparatus (Invitrogen), where all remaining immunoblotting steps were carried out. After last wash, the membrane was developed using Supersignal West Pico (Thermo Scientific). Protein bands were imaged using ImageQuant LAS4000 imaging system (GE Healthcare Life Sciences).

β-Galactosidase activity assay. P. aeruginosa strains to be assayed were grown overnight in LB agar plates, then bacterial suspensions were prepared in fresh M63 complete to OD600 = 0.5 with or without oxylipins or OA (0.1 mg mL−1). Cultures were incubated at 30 °C for the duration of the experiment, then bacterial cells were centrifuged and the supernatant discarded (to remove OA and oxylipins that interfere with the OD600). Bacteria were suspended in M63 medium at an OD600 = 1, then 250 µL of each suspension was mixed with 250 µL of Z buffer [NaH2PO4, Na2HPO4, (0.06 M), KCl (0.04 M), MgSO4 (0.001 M), b-mercaptoethanol (0.05 M), pH 7.0], 5 µL of 0.1% SDS and 100 µL of chloroform and the mix vortexed for 20 s. The tubes were incubated at 30 °C for 5 min and the reaction started by adding 100 µL of o-nitrophenyl-β-D-galactoside (ONPG, 4 mg mL−1) and briefly vortex mixing. Reactions were incubated at 30 °C for 1 h and stopped by adding 250 µL of 1 M Na2CO3. The reaction was centrifuged for 1 min at maximum speed and the supernatant was used to measure the OD at 420 nm and at 550 nm. Finally, β-gal activity was calculated using the equation: Miller Units = 1000 × [(OD420 − 1.75 × OD550)]/(T × V × OD0), where OD420 and OD550 are the final reads from the reaction mixture, OD0 is the initial cell density of the cultures, T is the time of the reaction in min and V the volume of culture used in the assay in mL.

Electrophoretic mobility shift assay. Binding of OdsR to the diol synthase operon (DS promoter) was assessed by EMSA using the purified OdsR-His protein. A DNA fragment comprising 200 bp immediately upstream from the PA2078 start codon was used as a DS promoter probe. Another unrelated 150 bp DNA fragment was used as negative control. The DS promoter (15 ng) or control DNA (20 ng) probes were mixed with 0, 12, 36, 48, 90 pmol of OdsR-His protein in binding buffer 1× (5% composition: 100 mM Hepes, pH 7.6, 5 mM EDTA, 50 mM (NH4)2SO4, 5 mM DTT, Tween 20, 1 % (w/v), 150 mM KCl) and incubated 30 min at room temperature. Each DNA-protein mix was applied in a 6% DNA retardation gel (Invitrogen Cat. No. ECG356BOX) and electrophoresed at 120 volts until the loading dye reached 1 cm from the gel edge. The gel was then stained with SYBR gold (Thermofisher), visualized and photographed.

Co-culture experiments. Strains ΔDS (PA3427p-lacZ) and Δodsr (PA3427p-lacZ:Supplementary Table 1) were each co-cultured with PA01 in 3 mL of M63 complete medium supplemented with OA (1 mg mL−1). Each pair of strains were inoculated from fresh overnight grown pre-cultures at an initial OD600 = 0.1 and incubated for 6 h at 30 °C. The β-gal activity was measured using 250 µL of each co-culture as described above. Strains ΔDS (PA3427p-lacZ) or Δodsr (PA3427p-lacZ) alone cultured under the same conditions were used as negative controls.

Phylogenetic tree construction. The phylogenetic trees were generated using BLAST pairwise alignments by the Neighbor Joining method. The maximum sequence difference (the maximum allowed fraction of mismatched bases in the aligned region between any pair of sequences) was set to 0.7.

Statistical analysis. Unpaired t-test (two-tailed) was used to determine differences between means of varying conditions after it was determined that the variance was similar between groups. All statistical analyses were performed using GraphPad Prism 7 software.

Reporting summary. Further information on experimental design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The authors declare that the data supporting the findings of this study as well as all materials and data generated, such as plasmids and strains, are available within the article and its supplementary information files, or from the corresponding author upon request. The transcriptomic data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus35 and are accessible through GEO Series accession number GSE123356. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE36 partner repository with the dataset identifier PXD012256.

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