Identification of a novel Rhl/R-PrrH-LasI/PhzC/PhzD signalling cascade and its implication in *P. aeruginosa* virulence

Yang Lu*, Honglin Li*, Jieying Pu*, Qian Xiao*, Chanjing Zhao*, Yimei Liu*, Lina Wang*, Youqiang Li*, Bin Huang*, Jianming Zeng* and Cha Chen*

*Department of Laboratory Medicine, The Second Affiliated Hospital of Guangzhou University of Chinese Medicine, Guangzhou, P. R. People’s Republic of China; †Department of Laboratory Medicine, The First Affiliated Hospital of Sun Yat-sen University, Guangzhou, P. R. People’s Republic of China

**ABSTRACT**

Small regulatory RNAs (sRNAs) act as key regulators in many bacterial signalling cascades. However, in *P. aeruginosa*, the sRNAs involved in quorum sensing (QS) regulation and their function are still largely unknown. Here, we explored how the *prrH* locus sRNA influences *P. aeruginosa* virulence in the context of the QS regulatory network. First, gain- and loss-of-function studies showed that PrrH affects pyocyanin, elastase and rhamnolipid production; biofilm formation; and swimming and swarming motility and impaired the viability of *P. aeruginosa* in human whole blood. Next, our investigation disclosed that LasI and PhzC/D were directly repressed by PrrH. In addition, RhlI, the key member of the *rhl* QS system, diminished the expression of PrrH and enhanced the expression of downstream genes. Bioinformatics analysis found two binding sites of RhlR, the transcription factor of the *rhl* system, on the promoter region of *prrH*. Further β-galactosidase reporter and qPCR assays confirmed that PrrH was transcriptionally repressed by RhlR. Collectively, our data identified a novel Rhl/R-PrrH-LasI/PhzC/PhzD regulatory circuitry that may contribute to *P. aeruginosa* pathogenesis. Our findings indicate that PrrH is a quorum regulatory RNA (Qrr) in *P. aeruginosa* and provide new insight into PrrH’s function.

**ARTICLE HISTORY**

Received 30 July 2019; Revised 13 October 2019; Accepted 28 October 2019

**KEYWORDS** PrrH; sRNA; quorum sensing; *P. aeruginosa*; virulence

**Introduction**

The opportunistic pathogen *Pseudomonas aeruginosa* (*P. aeruginosa*) is the cause of a variety of human infections, including acute infections in patients who are injured or severely burned and chronic lung infections in individuals with cystic fibrosis [1]. *P. aeruginosa* pathogenicity is due to mainly a large amount of cell-associated and extracellular virulence factors, many of which are regulated by two acyl-homoserine lactone (AHL) quorum sensing (QS) systems, LasI-LasR (*las* system) and RhlI-RhlR (*rhl* system) [2]. In the *las* system, the *lasI* gene product directly forms the signal of this system. AHL diffuses more easily outside the cell and acts as an extracellular signalling molecule, which interacts with LasR to activate a number of virulence factors, including the LasA and LasB elastases, exotoxin A, and alkaline protease. In an analogous manner, the *rhlI* and *rhlR* genes encode for a *N*-butanoyl-L-homoserine lactone (C4-HSL) synthase and receptor, respectively. The downstream virulence genes of the *rhl* system include those encoding pyocyanin, rhamnolipids and the type III secretion system [3,4]. In commonly studied *P. aeruginosa* strains, *las* and *rhl* systems are hierarchically connected, and both *rhlI* and *rhlR* are positively regulated by the *las* system [5]. However, recent research has shown that once LasR mutants emerge among populations capable of C4-HSL production, they can rapidly evolve an active LasR-independent *rhl* QS system [2,6]. In addition, these two AHL QS systems interact with a non-AHL signalling system called the *Pseudomonas* quinolone signal (PQS) system, which is regulated by mainly the *pqs* operon [7].

Small regulatory RNAs (sRNAs) are central regulators of gene expression in bacteria and post-translationally control target genes by limited base pairing with their mRNAs, thus negatively or positively affecting the transcript stability and/or the translation rate [8,9]. In recent years, increasing studies have shown that sRNAs regulate a wide variety of genes that encode...
for proteins involved in many cellular processes and pathogenesis, including QS and virulence activities [10–12]. In P. aeruginosa, ReaL is negatively regulated by the las system and enhances the synthesis of PQS by targeting Pqsc [13]. In addition, PhrS and PrrF modulate PQS by regulating PqsR and AntR, respectively [14–17]. However, the number of sRNAs that have been implicated in the regulation of QS, especially the las and rhl system, is limited. An extensive investigation of quorum regulatory RNAs (Qrrs) [18] will help to disclose the elaborate regulatory network of P. aeruginosa virulence.

PrrH is characterized as a haem-responsive sRNA encoded by the prrF locus. Transcription of PrrH initiates at the 5’ end of prrF1, proceeds through the prrF1 terminator and prrF1-prrF2 intergenic sequence, and terminates at the 3’ end of the prrF2 gene. The widely known function of this sRNA family is its participation in iron homeostasis, homologous to the RyhB RNAs encoded by Escherichia coli, Shigella flexneri, Shigella dysenteriae, and Vibrio cholerae [19,20]. In this study, we showed that perturbation of PrrH levels affected many processes that are known to be influenced by QS and regulated the virulence of P. aeruginosa. Further evidence demonstrated that the LasI and phzA-G operons were direct targets of PrrH. More importantly, we also disclosed that the virulence inhibition function of PrrH was under the control of the rhl system. Our findings highlight the importance of PrrH in P. aeruginosa pathogenesis and implicate PrrH as a potential therapeutic target for P. aeruginosa.

Results

PrrH influences P. aeruginosa virulence and bloodstream infection ability in vitro.

To assess the function of PrrH, we first constructed prrH gene-deficient and overexpression strains in PAO1 (ΔprrH, WT/PrrH, respectively), as well as a prrH-overexpression strain in the ΔprrH mutation background (ΔprrH/PrrH). The expression levels of the PrrH in all these strains were confirmed by RT-qPCR (Figure S1). Compared to the wild-type strain, the strain with overexpression of prrH had slightly repressed growth, and deficiency in prrH barely affected growth (Figure S2). However, it was notably that overexpression of prrH inhibited the production of pyocyanan (Figure 1A), while deficiency in prrH in the genome resulted in an increase in pyocyanin, which was attenuated by introduction of a prrH-overexpression plasmid (Figure 1B). Furthermore, as azithromycin (AZM) has been shown to potentially inhibit QS signal molecules and attenuate pyocyanin synthesis by P. aeruginosa [21,22], we found that when P. aeruginosa was exposed to 2, 8, and 32 μg/ml AZM (1/64th, 1/16th, and 1/4th of the MIC), pyocyanin synthesis was decreased in a dose-dependent manner (Figure 1C), accompanied with enhanced expression of PrrH, also in a dose-dependent manner (Figure 1D). These data suggest that PrrH negatively regulates pyocyanin synthesis in P. aeruginosa.

To further measure the effects of PrrH on the other QS-linked virulence traits, we next analysed elastase and rhamnolipid production, biofilm formation, and swimming and swarming motility in conditions of prrH overexpression or deletion. As shown in Figure 2A, B and C, deletion of prrH showed no or little promotion of elastase production, rhamnolipid synthesis and biofilm formation, while PrrH overexpression significantly inhibited the above phenotypes. In contrast, swimming and swarming were slightly repressed by deletion of prrH and obviously enhanced by PrrH overexpression (Figure 2D). PrrH appeared to have opposite effects on swimming and swarming compared to those on biofilm formation. This result was consisted with the observation that swimming motility and biofilm formation have an inverse relationship in P. aeruginosa strains [13]. Overall, these results indicated that PrrH plays a negative regulatory role in elastase production, rhamnolipid synthesis and biofilm formation, while its role in swimming and swarming motility is positive.

Since virulence may be a contributing factor for P. aeruginosa bacteremia [23], we then examined the effect of PrrH on the ability of P. aeruginosa to survive in a bloodstream infection by transferring growing bacteria to whole blood to mimic their transition from an extravascular site into the bloodstream [24,25]. As shown in Figure 3A, within 30 min after incubation in whole blood from human volunteers, the count of the prrH-overexpression strain was dramatically reduced, while that of the wild-type or prrH-deficient strain hardly changed. At 60, 90 and 120 min, more than half of both the wild-type and prrH-overexpression PAO1 was killed compared with the initial inoculation quantity. However, it was intriguing that the number of the ΔprrH strain was almost constant throughout the incubation. To further investigate the role of PrrH and in P. aeruginosa bacteremia, we tested the expression level of PrrH after whole blood incubation. Allowing for the great differences in RNA quality between these two cultures from Luria–Bertani (LB) medium and whole blood, we reduced the application number in whole blood to 10% and extended the incubation time. The result showed that the expression of prrH was enhanced at 4 h and faded at 8 h (Figure 3B). The upregulation of PrrH was coincident with the death of PAO1, which might be explained by a disinfection effect of blood in the early stage, while the reduction might be a result of bacterial adaption to blood.
Collectively, these data suggest that PrrH plays a crucial role in impairing the virulence and bacteraemia of PAO1.

**PrrH directly regulates LasI, PhzC and PhzD expression**

Next, we explored the molecular mechanisms responsible for the functions of PrrH that were observed above. Potential target genes of PrrH were first predicted using the publicly available database IntaRNA. LasI, PhzC and PhzD were chosen for further analysis (Figure 4A) because LasI was the key member of a QS system, while phenazine operons, phzA-G, played a vital role in pyocyanin synthesis [26]. To verify whether LasI, PhzC and PhzD are direct targets of PrrH, a green fluorescent protein (GFP) reporter system was first constructed (Figure 4B). Next, analysis showed that co-expression of PrrH significantly inhibited the intensity of GFP that carried a wild-type but not mutant base-pairing site of LasI, PhzC or PhzD (Figure 4C and D), indicating that PrrH may suppress gene expression through its base-pairing sequence at the CDS of a target gene. Further investigation showed that PrrH can also diminish endogenous expression of both PhzC and PhzD (Figure 4E), which may be attributed to degradation caused by sRNA. For LasI, the mRNA level was hardly affected by PrrH, however, the mRNA level of lasA, a downstream virulence gene of LasI, was significantly decreased upon PrrH overexpression, suggesting that PrrH regulates LasI expression at the translational level (Figure 4E).

Taken together, these results implicate that PrrH may negatively regulate lasI, PhzC and PhzD expression by directly targeting the CDS of their mRNAs.

**Regulation of PrrH transcription by RhlI/R**

Considering that the synthesis of virulence factors is primarily under the control of QS [2], we inferred that PrrH may be transcribed by QS-related genes. Since the activation of QS depends on the density of bacteria, we tested the expression levels of lasI, rhlI and prrH in PAO1 during the early exponential phase (t = 2 h), the mid-exponential phase (t = 4 h) and the early stationary phase (t = 6 h) in a corresponding culture model (Figure S3). As shown in Figure 5A, the mRNA of LasI and RhlI gradually rose from 2 h to 4 h and remained stable at 6 h; it’s important that PrrH displayed an inverse trend of expression over time. Moreover, the expression of PrrH was also negatively related to LasI and RhlI in the whole blood culture model (Figure S4). Therefore, we evaluated the role of LasI and RhlI in PrrH transcription. The results showed that deletion or overexpression of lasI, as well as treatment with 3-oxo-C12-HSL barely affected...
PrrH expression (Figure 5B and S5A). However, rhlI deletion caused an increase in endogenous PrrH expression, which was attenuated by restoration of RhlI (Figure 5B) or C4-HSL (Figure S5B). More importantly, similar to overexpression of PrrH, deficiency in rhlI also led to downregulation of LasA, PhzC and PhzD but not of LasI, and restoration of RhlI rescued this phenomenon (Figure 5C).

Since the gene expression-regulated function of RhlI is mediated by the transcription factor (TF) RhlR, we further used the bioinformatics tool PRODORIC, a comprehensive database for gene regulation and expression in prokaryotes, to analyse the DNA motif recognized by RhlR. It seems that genes, such as rhlAB, pa1L and migA, with the specific DNA sequence 5'-CT(N12)AG-3' or 5'-CT(N12)GC-3' in the promoter region may be potential targets of RhlR. Based on this model, within the −100 bp region upstream of PrrH, two RhlR consensus binding sites were found (Figure 5D). Then, a DNA fragment of prrH promoter carrying these predicted sites was cloned upstream of the β-galactosidase gene in the pQF50-promoter reporter. When transformed into BW25113 (an E. coli strain without endogenous β-galactosidase activity), the β-galactosidase activity of pQF50-PrrH was greatly elevated compared to that of the control, while overexpression of RhlR impaired this activity (Figure 5E). Moreover, overexpression of RhlR also resulted in downregulation of PrrH (Figure 5F). These results suggest that the virulence regulatory role of PrrH is under the control of rhlI system.

Figure 2. PrrH influences elastase and rhamnolipid production, biofilm formation, swimming and swarming motility of P. aeruginosa. (A) PrrH overexpression inhibited elastase production. PAO1 carrying pROp200 (WT/Vector), PAO1ΔprrH carrying pROp200 or pROp200-prrH (ΔprrH/Vector, ΔprrH/PrrH) were cultured in LB for 8 h, and the activity of elastase in supernatant was determined. (B) PrrH overexpression inhibited rhamnolipid synthesis. The indicated PAO1 strains were cultured in M9 minimal salts medium for 8 h at 37 °C, and rhamnolipid in the supernatant was measured. (C) PrrH inhibited biofilm formation. The indicated PAO1 strains were cultured in LB in the 12-well plates for 24 h at 37°C. The biofilm was quantified by measuring solubilized crystal violet staining biofilm cells at OD600. (D) PrrH positively regulated swimming and swarming motility. 5μl cultures of the indicated strains were spotted onto the swarming or swimming medium and incubated at 37°C for 16 h. Values are the mean ± SD of at least three independent experiments. ns, not significant; **, *P < 0.01; ***, *P < 0.001.
Taken together, our results elucidate a novel Rhl/I/R-PrrH-LasI/PhzC/PhzD signalling cascade and further identify the importance of PrrH in inhibiting pyocyanin synthesis and virulence (Figure 6), which substantially extend our understanding of the P. aeruginosa QS circuit and the function of PrrH.

**Discussion**

In this study, we disclose a novel QS/PrrH-PhzC/PhzD regulatory circuitry, that is, Rhl/I/R represses the expression of PrrH, whereas the expression of PrrH reduces the levels of LasI, Rhl/I and PhzC/PhzD, thereby inhibiting QS signalling and abrogates QS-promoted virulence in P. aeruginosa. Our results elucidate the new regulation mechanism and function of PrrH.

One of the most fascinating aspects of the prrF locus sRNAs in P. aeruginosa is the tandem arrangement of the prrF1 and prrF2 genes, allowing for expression of a third sRNA, PrrH [27]. Most recent publications about prrF locus sRNAs have focused on the role of PrrF in iron homeostasis or PQS regulation [17,19,27,28]. Few reports suggest that there is an extra haem-responsive function for PrrH, as a result of the unique sequence derived from the prrF1-prrF2 intergenic sequence (PrrHIG) [20]. In this study, we showed that deletion or overexpression of the whole sequence of prrH resulted in a change in virulence and bloodstream infection ability. However, our genetic manipulation not only changed the PrrH level but also caused gain or loss of PrrF expression. In addition, the recognition site of PrrH on LasI, PhzC and PhzD was not in PrrHIG but shared with PrrF. Thus, the assessed function of PrrH in our work relies on that of the prrH locus sRNAs and not on PrrH only.

In P. aeruginosa PA01, pyocyanin is synthesized from chorismate in a complex series of intermediates by enzymes encoded by the homologous phzA1B1C1-D1E1F1G1 and phzA2B2C2D2E2F2G2 operons. The two phz operons are 98.3% identical at the DNA level [26,29]. As a result, the recognition site of PrrH in PhzC1/D1 is the same as that in PhzC2/D2. We confirmed that both PhzC1/D1 and PhzC2/D2 were targets of PrrH. More interestingly, the pairing bases among these four target mRNAs were conserved, meaning that the PrrH family used the same base-pairing site to recognize a protein family. In the perspective of evolution, this conservation means that these base-pairing events are likely to be physiologically relevant.

It has been shown that the transcription of quite a few sRNAs is regulated by TFs and that many sRNAs regulate TF-encoding genes. This strategy implies that TFs, sRNAs and target genes may form a complex regulatory circuit [8,9,30]. For example, in Vibrio harveyi and Vibrio cholerae, LuxO activates the expression of the Qrr sRNAs, which in turn repress the expression of the QS master regulators LuxR and LuxO [30–33]; thus, sRNAs may function at the centre of QS circuits. P. aeruginosa is a model for studies of QS. However, the homologous Qrr sRNAs in P. aeruginosa have not yet been characterized. Here, we showed that Rhl/I/R directly repressed PrrH and promoted PhzC/D expression, while PhzC/D was negatively regulated by PrrH. This finding suggests that Rhl/I/R, PrrH and PhzC/D combine in a feed-forward loop to regulate pyocyanin production. On the other hand, PrrH also repressed Rhl/I/R expression (Figure S6) through targeting LasI, forming a negative-negative feedback loop to fine-tune QS. Our data suggest that prrH locus sRNAs may be the Qrr sRNAs of P. aeruginosa.

A few publications have reported several genes involved in the link between iron and QS systems in P. aeruginosa. MvfR, for example, was transcriptionally activated under low-iron concentrations or by the cell-cell signalling molecules 4-hydroxy-2-heptylquinoline (HHQ) and 3,4-dihydroxy-2-heptylquinoline (PQS), subsequently controlling transcription of iron-related genes and many virulence-related factors [34,35]. Previous works have disclosed that PrrF was transcriptionally repressed by Fur [19]. Its target mRNA involved in iron homeostasis was PhuS, and AntR and VreR were targets involved in virulence [17,27]. In our work, LasI and PhzC/D were identified as new virulence-associated targets for this locus sRNA. More importantly, we also revealed that the expression of PrrH was under the control of Rhl/I/R. To our knowledge, this study is the first time that a sRNA has been shown to connect AHL QS to iron homeostasis in P. aeruginosa.

In summary, we disclose a novel QS-PrrH regulatory circuit and demonstrate that PrrH is capable of repressing P. aeruginosa virulence and bloodstream
infection ability in vitro. Our findings advance the fundamental understanding of QS circuits in P. aeruginosa and indicate that PrrH might be a potential target for therapeutic development.

**Experimental procedures**

**Bacterial strains, plasmids and growth conditions**

The bacterial strains and plasmids used in this study are listed in Table S1. Bacteria were grown in LB medium or on LB plates containing 1.5% agar, unless otherwise indicated. When necessary, antibiotics were used at the following concentrations: 100 μg/ml ampicillin (AMP), 25 μg/ml gentamicin (GM), 16 μg/ml chloramphenicol (CM), and 2, 8, and 32 μg/ml AZM.

**Plasmid construction**

The whole sequence of PrrH or the coding sequences of LasI, RhlI and RhlR were cloned into the EcoRI site of pROp200 with a Ready-to-Use Seamless Cloning Kit (BBI, USA, Code NO. B632219) to generate expression vectors named pROp200-prrH, pROp200-lasl, pROp200-rhII and pROp200-rhIR, respectively. For another PrrH expression vector, pSTV28-prrH, the prrH gene was cloned into the PstI/Hind III sites of pSTV28.

To construct the target-gfp translational fusion vectors pUCP30T-lasl-gfp, pUCP30T-phzC-gfp and pUCP30T-phzD-gfp, the coding sequence of GFP was first cloned into XbaI/HindIII sites downstream of the P<sub>lac</sub> promoter in pUCP30T to generate the
reporter vector pUCP30T-gfp, and then a wild-type fragment of LasI, PhzC or PhzD mRNA that contained putative binding sites for PrrH was PCR amplified and inserted into the XbaI/NcoI sites upstream of the first codon of GFP in the pUCP30T-gfp vector. pUCP30T-mRNA-mut-gfp, which carried a mutated sequence in the complementary site for PrrH was generated using fusion PCR based on wild-type pUCP30T-mRNA-gfp.

The plasmid pQF50 containing a promoter-less lacZ reporter was used for prrH promoter analysis. A DNA fragment of the prrH promoter spanning −92 to +74 was PCR-amplified and inserted into BamHI/Hind III sites upstream of the lacZ reporter in pQF50 to generate pQF50-PprrH.

All constructs were confirmed by direct sequencing. Primers used in this study are listed in Table S2.

**Construction of PAO1 prrH-deficient mutants**

The sacB-based suicide vector system was employed for the knockout of prrH in PAO1 as described by Zeng et al in our previous work [21]. Briefly, the upstream and downstream recombinant fragment of prrH was cloned into XbaI/Sacl sites in pGSM to generate the recombinant plasmid pGSM-ΔprrH, and then it was transformed into PAO1 to generate a prrH mutant strain (PAO1ΔprrH).

**Real-time PCR**

Total RNA was extracted using RNAiso Plus reagent (Takara, Dalian, Liaoning, China). Reverse transcription (1 μg of total RNA) was performed with the PrimeScript RT Reagent Kit (Takara, Dalian, Liaoning, China; code No. RR047A). The cDNA was subjected to qPCR on a ViiATM 7 Dx system (Applied Biosystems, Foster, CA, USA) using SYBR Green qPCR Master Mix (Takara, Dalian, Liaoning, China). The expression levels of the target genes were normalized to the expression of an internal control gene (rpoD) using the 2−ΔΔCt method. The sequences of the primers are listed in Table S2.

**Fluorescence intensity measurement**

E. coli DH5α carrying the reporter pUCP30T-mRNA-gfp or pUCP30T-mRNA-mut-gfp combined with pSTV28 or pSTV28-ΔprrH was grown overnight (8~10 h) at 37°C and was re-incubated to 0.5 McFarland
standard (MCF), and then 100 µl of cultures were added into 3 ml of LB to grow for 6 h. Subsequently, cultures were collected by centrifugation and washed twice and resuspended in 0.9% NaCl, and 200 µl aliquots were transferred to black polystyrene 96-well microplates with a clear, flat bottom. The absorbance (Abs595) and fluorescence intensity (F485/535) were measured in a Tecan Infinity M1000Pro Reader. GFP activity was expressed in arbitrary units (AU) as F485/535/Abs595. All of the tests were carried out independently and at least in triplicate.

β-galactosidase assays
β-galactosidase assays were carried out by the Miller method when cells were grown to mid-log phase at 37°C [36]. All of the tests were carried out independently and at least in triplicate.

Pyocyanin production assay
Pyocyanin in 5 ml of P. aeruginosa culture supernatant was extracted with 3 ml of chloroform and 1 ml of 0.2 N HCl, and then the absorbance of the extract was measured at 520 and 600 nm. The concentration of pyocyanin was determined using the following formula: (A520/A600×17.072) = µg/ml [37]. All of the tests were performed independently and at least in triplicate.

Elastase assay
Elastase activity in P. aeruginosa cultures supernatant was determined by elastin-Congo red (ECR) (Sigma Chemical, America) assay [38]. Mid-exponential phase bacteria were inoculated in 3 ml of LB at an OD600 of 0.05 and grown for 8 h at 37°C. Then, 200 µl culture supernatants were added to 800 µl ECR solution containing 0.1 M Tris (pH 7.2), 1 mM CaCl2 and 3 mg/ml ECR. Reaction tube was incubated 4 h at 37°C with shaking and then 100 µl of 0.12 M EDTA was added to stop reaction. Insoluble ECR was removed by centrifugation, and the OD495 was measured. All results were analyzed of three independent experiments.

Rhamnolipid assay
The detection of rhamnolipid was based on the experimental method used by Neissa M. Pinzon with some modifications [39]. Mid-exponential phase bacteria were resuspended in 3 ml of M9 minimal salts medium that contained 0.4% glucose, 2 mM MgSO4 and 100 µM CaCl2 at OD600 of 0.05 and incubated at 37°C for 8 h. 1ml of culture supernatant was acidified (pH 2.5 ± 0.2) using 1 N HCl, rhamnolipid was extracted with 4 ml chloroform. Then 3 ml of chloroform extract was allowed to react with 100 µl of 1 g/l of methylene blue and 2 ml of distilled water, the OD638 was measured. All results were analyzed of three independent experiments.

Biofilm formation assay
A biofilm formation assay was performed as described previously by Sara Carloni [13]. Mid-exponential phase bacteria were inoculated in 2 ml of LB at an OD600 of 0.05 in 12-well plates and grown for 24 h at 37°C. Biofilm cells attached to plates were stained with 1% crystal violet. Then, 2 ml of 95% ethanol was used to solubilize crystal violet-stained biofilm cells and measured at OD600. All analyses included three independent experiments.

Motility assays
P. aeruginosa strains were grown in LB medium to the mid-exponential phase, and 5 µl of the culture was spotted onto swarming medium (LB containing 0.5% agar and 5 g/l glucose) and swimming medium (tryptone broth containing 0.3% agar), respectively [40]. Swarming and swimming motility were observed after 16 h of incubation at 37°C. At least three replicates were measured for each sample.

Whole blood killing assay
Whole blood was partitioned into wells of a 96-well round bottom plate, and 1 × 10^6 CFU of the indicated PAO1 strain in 0.9% NaCl was added to whole blood to produce a final concentration of 90% whole blood in a 200 µl volume. Samples were taken at time 0, 30, 60, 90, and 120 min, serially diluted, and plated on LB agar plates for CFU enumeration [41]. Experiments were performed at least in triplicate.

Ethics approval and consent to participate
For the whole blood killing assay, informed consent was obtained from each volunteer, and the study was approved by the Ethics Committee of Guangdong Provincial Hospital of Traditional Chinese Medicine (approval notice number: BE-2018-166-01).

Statistical analysis
Data of the results from multiple independent experiments are expressed as the mean ± standard deviation (SD). The differences between groups were analysed using Student’s t-test when two groups were compared or one-way ANOVA when more than two groups were compared. All analyses were performed using GraphPad Prism, version 5 (GraphPad Software, Inc., San Diego, CA, USA). Differences with a P-value (P) of 0.01 < P < 0.05 are represented by *, P < 0.01 are represented by **, and P < 0.001 are represented by ***.

Acknowledgements
We are very grateful to Prof. Shiyun Chen (Wuhan Institute of Virology, Chinese Academy of Sciences, Wuhan, China) for providing the plasmid pROp200, Guangdong Magigene Biotechnology Co., Ltd for the help of bioinformatics analysis.
Disclosure statement

No potential conflict of interest was reported by the authors.

Funding

This work was supported by the National Natural Science Foundation of China (grant number 81702063, 81871703); the Guangzhou Science Technology and Innovation Commission (grant number 201707010296); the Natural Science Foundation of Guangdong Province (grant number 2015A030310439); Guangdong Provincial Hospital of Traditional Chinese Medicine (grant number YN2018QJ01).

References

[1] Valentini M, Gonzalez D, Mavridou DA, et al. Lifestyle transitions and adaptive pathogenesis of Pseudomonas aeruginosa. Curr Opin Microbiol. 2018;41:15–20.
[2] Kostylev M, Kim DY, Smalley NE, et al. Evolution of the Pseudomonas aeruginosa quorum-sensing hierarchy. Proc Natl Acad Sci U S A. 2019;116:7027–7032.
[3] Lu Y, Zeng J, Wu B ES, et al. Quorum sensing N-acyl homoserine lactones-SdhA suppresses Escherichia coli-Pseudomonas aeruginosa Conjugation through inhibiting tral expression. Front Cell Infect Microbiol. 2017;7(7):1–7.
[4] Fuqua C, Parsek MR, Greenberg EP. Regulation of gene expression by cell-to-cell communication: acyl-homoserine lactone quorum sensing. Annu Rev Genet. 2001;35:439–468.
[5] Lee J, Zhang L. The hierarchy quorum sensing network in Pseudomonas aeruginosa. Protein Cell. 2015;6:26–41.
[6] Chen R, Déziel E, Groleau M-C, et al. Social cheating in a Pseudomonas aeruginosa quorum-sensing variant. Proc Natl Acad Sci U S A. 2019;116:7021–7026.
[7] Déziel E, Lépine F, Milot S, et al. Analysis of Pseudomonas aeruginosa 4-hydroxy-2-alkylquinolines (HAQs) reveals a role for 4-hydroxy-2-heptylquinoline in cell-to-cell communication. Proc Natl Acad Sci U S A. 2004;101:1339–1344.
[8] Mandin P, Guillier M. Expanding control in bacteria: Interplay between small RNAs and transcriptional regulators to control gene expression. Curr Opin Microbiol. 2013;16:125–132.
[9] Nitzan M, Rehani R, Margalit H. Integration of bacterial small RNAs in regulatory networks. Annu Rev Biophys. 2017;46:131–148.
[10] Sonnleitner E, Romeo A, Bläsi U. Small regulatory RNAs in Pseudomonas aeruginosa. RNA Biol. 2012;9:364–371.
[11] Wagner EGH, Romby P. Small RNAs in bacteria and archaea: who they are, what they do, and how they do it. Adv Genet. 2015;90:133–208.
[12] Felden B, Cattoira V. Bacterial adaptation to antibiotics through regulatory RNAs. Antimicrob Agents Chemother. 2018;62:1–11.
[13] Carloni S, Macchi R, Sattin S, et al. The small RNA Real1: a novel regulatory element embedded in the Pseudomonas aeruginosa quorum sensing networks. Environ Microbiol. 2017;19:4220–4237.
[14] Sonnleitner E, Gonzalez N, Sorger-Domenigg T, et al. The small RNA Phr5 stimulates synthesis of the Pseudomonas aeruginosa quinolone signal. Mol Microbiol. 2011;80:868–885.
[15] Oglesby AG, Farrow JM, Lee J-H, et al. The influence of iron on Pseudomonas aeruginosa physiology: a regulatory link between iron and quorum sensing. J Biol Chem. 2008;283:15558–15567.
[16] Ferrara S, Carloni S, Fulco R, et al. Post-transcriptional regulation of the virulence-associated enzyme AlgC by the r22-dependent small RNA ErsA of Pseudomonas aeruginosa. Environ Microbiol. 2015;17:199–214.
[17] Djapgne L, Panja S, Brewer LK, et al. The Pseudomonas aeruginosa PrrF1 and PrrF2 Small regulatory RNAs Promote 2-Alkyl-4-quinoline production through redundant regulation of the antR mRNA. J Bacteriol. 2018;2001:1–17.
[18] Tu KC, Long T, Svenningsen SL, et al. Negative feedback loops involving small regulatory RNAs precisely control the Vibrio harveyi quorum-sensing response. Mol Cell. 2010;37:567–579.
[19] Wilderman PJ, Sowa NA, FitzGerald DJ, et al. Identification of tandem duplicate regulatory small RNAs in Pseudomonas aeruginosa involved in iron homeostasis. Proc Natl Acad Sci. 2004;101:9792–9797.
[20] Oglesby-Sherrouse AG, Vasil ML. Characterization of a heme-regulated non-coding RNA encoded by the prr locus of Pseudomonas aeruginosa. PLoS One. 2010;5:e9930, 1–11.
[21] Zeng J, Zhang N, Huang B, et al. Mechanism of azithromycin inhibition of HSL synthesis in Pseudomonas aeruginosa. Sci Rep. 2016;6:1–10.
[22] Pérez-Martínez I, Haas D. Azoithromycin inhibits expression of the GacA-dependent small RNAs RsmY and RsmZ in Pseudomonas aeruginosa. Antimicrob Agents Chemother. 2011;55:3399–3405.
[23] Qadir SM, Donkor DA, Bhakta V, et al. Phosphatidylserine externalization and procoagulant activation of erythrocytes induced by Pseudomonas aeruginosa virulence factor pyocyanin. J Cell Mol Med. 2016;20:710–720.
[24] Graham MR, Virtaneva K, Porcella SF, et al. Group A Streptococcus transcriptome dynamics during growth in human blood reveals bacterial adaptive and survival strategies. Am J Pathol. 2005;166:455–466.
[25] Colmer-Hamood JA, Dzvova N, Krucezk C, et al. In vitro analysis of Pseudomonas aeruginosa virulence using conditions that mimic the environment at specific infection sites. Prog Mol Biol Transl Sci. 2016;142:151–191.
[26] Mavrodi DV, Bonsall RF, Delaney SM, et al. Functional analysis of genes for biosynthesis of pyocyanin and phenazine-1-carboxamide from Pseudomonas aeruginosa PA01. J Bacteriol. 2001;183:6454–6465.
[27] Reinhart AA, Powell DA, Nguyen AT, et al. The prrF-encoded small regulatory RNAs are required for iron homeostasis and virulence of Pseudomonas aeruginosa. Infect Immun. 2015;83:863–875.
[28] Reinhart AA, Nguyen AT, Brewer LK, et al. The Pseudomonas aeruginosa PrrF Small RNAs regulate iron homeostasis during acute murine lung infection. Infect Immun. 2017;85:1–15.
[29] Sui SJH, Lo R, Fernandes AR, et al. The heme-regulated non-coding RNA encoded by the hsa RNA promotes the Vibrio harveyi quorum-sensing response. Mol Microbiol. 2010;78:1448–1466.
[30] Mavrodi DV, Bonsall RF, Delaney SM, et al. Functional analysis of genes for biosynthesis of pyocyanin and phenazine-1-carboxamide from Pseudomonas aeruginosa PA01. J Bacteriol. 2001;183:6454–6465.
[31] Reinhart AA, Powell DA, Nguyen AT, et al. The prrF-encoded small regulatory RNAs are required for iron homeostasis and virulence of Pseudomonas aeruginosa. Infect Immun. 2015;83:863–875.
[32] Reinhart AA, Nguyen AT, Brewer LK, et al. The Pseudomonas aeruginosa PrrF Small RNAs regulate iron homeostasis during acute murine lung infection. Infect Immun. 2017;85:1–15.
[33] Sui SJH, Lo R, Fernandes AR, et al. The heme-regulated non-coding RNA encoded by the hsa RNA promotes the Vibrio harveyi quorum-sensing response. Mol Microbiol. 2010;78:1448–1466.
[34] Beisel CL, Storz G. Base pairing small regulatory RNAs and their roles in global regulatory networks. Mol Microbiol. 2010;78:1448–1466.
their roles in target regulation. EMBO J. 2013;32:2158–2171.

[32] Lenz DH, Mok KC, Lilley BN, et al. The small RNA chaperone Hfq and multiple small RNAs control quorum sensing in *Vibrio harveyi* and *Vibrio cholerae*. Cell. 2004;118:69–82.

[33] Feng L, Rutherford ST, Papenfort K, et al. A qrr non-coding RNA deploys four different regulatory mechanisms to optimize quorum-sensing dynamics. Cell. 2015;160:228–240.

[34] Balasubramanian D, Schneper L, Kumari H, et al. A dynamic and intricate regulatory network determines *Pseudomonas aeruginosa* virulence. Nucleic Acids Res. 2013;41:1–20.

[35] Hazan R, He J, Xiao G, et al. Homeostatic interplay between bacterial cell-cell signaling and iron in virulence. PLoS Pathog. 2010;6:e1000810.

[36] Giacomini A, Corich V, Ollero FJ, et al. Experimental conditions may affect reproducibility of the beta-galactosidase assay. FEBS Microbiol Lett. 1992;100:87–90.

[37] Essar DW, Eberly L, Hadero A, et al. Identification and characterization of genes for a second anthranilate synthase in *Pseudomonas aeruginosa*: interchangeability of the two anthranilate synthases and evolutionary implications. J Bacteriol. 1990;172:884–900.

[38] Pearson JP, Pesci EC, Iglewski BH. Roles of *Pseudomonas aeruginosa* las and rhl quorum-sensing systems in control of elastase and rhamnolipid biosynthesis genes. J Bacteriol. 1997;179:5756–5767.

[39] Pinzon NM, Ju L-K. Analysis of rhamnolipid biosurfactants by methylene blue complexation. Appl Microbiol Biotechnol. 2009;82:975–981.

[40] Rashid MH, Kornberg A. Inorganic polyphosphate is needed for swimming, swarming, and twitching motilities of *Pseudomonas aeruginosa*. Proc Natl Acad Sci U S A. 2000;97:4885–4890.

[41] Munguia J, LaRock DL, Tsumemoto H, et al. The Mla pathway is critical for *Pseudomonas aeruginosa* resistance to outer membrane permeabilization and host innate immune clearance. J Mol Med. 2017;95:1127–1136.