PhcQ mainly contributes to the regulation of quorum sensing-dependent genes, in which PhcR is partially involved, in *Ralstonia pseudosolanacearum* strain OE1-1

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**Abstract**
The gram-negative plant-pathogenic β-proteobacterium *Ralstonia pseudosolanacearum* strain OE1-1 produces methyl 3-hydroxymyristate as a quorum sensing (QS) signal via the methyltransferase PhcB and senses the chemical through the sensor histidine kinase PhcS. This leads to functionalization of the LysR family transcriptional regulator PhcA, regulating QS-dependent genes responsible for the QS-dependent phenotypes including virulence. The phc operon consists of *phcB*, *phcS*, *phcR*, and *phcQ*, with the latter two encoding regulator proteins with a receiver domain and a histidine kinase domain and with a receiver domain, respectively. To elucidate the function of PhcR and PhcQ in the regulation of QS-dependent genes, we generated *phcR*-deletion and *phcQ*-deletion mutants. Though the QS-dependent phenotypes of the *phcR*-deletion mutant were largely unchanged, deletion of *phcQ* led to a significant change in the QS-dependent phenotypes. Transcriptome analysis coupled with quantitative reverse transcription-PCR and RNA-sequencing revealed that *phcB*, *phcK*, and *phcA* in the *phcR*-deletion and *phcQ*-deletion mutants were expressed at similar levels as in strain OE1-1. Compared with strain OE1-1, expression of 22.9% and 26.4% of positively and negatively QS-dependent genes, respectively, was significantly altered in the *phcR*-deletion mutant. However, expression of 96.8% and 66.9% of positively and negatively QS-dependent genes, respectively, was significantly altered in the *phcQ*-deletion mutant. Furthermore, a strong positive correlation of expression of these QS-dependent genes was observed between the *phcQ*-deletion and *phcA*-deletion mutants. Our results indicate that PhcQ mainly contributes to the regulation of QS-dependent genes, in which PhcR is partially involved.
Quorum sensing (QS) allows bacterial cells to communicate for the cooperative regulation of physiological processes coordinating various bacterial community activities (Ham, 2013; Waters & Bassler, 2005). To recognize their own populations, bacterial cells produce and secrete QS signals, which are small, diffusible molecules. Bacteria monitor QS signals to track changes in their cell numbers and to activate QS for the synchronous control of the expression of genes beneficial for vigorous replication and adaptation to environmental conditions, including virulence, such as the formation of biofilms and the production of virulence factors (Galloway et al., 2011; Rutherford & Bassler, 2012).

The gram-negative plant-pathogenic β-proteobacterium Ralstonia solanacearum species complex (RSSC; Robinson et al., 2010) is globally distributed under diverse environmental conditions; strains infect more than 250 plant species in over 50 families and cause a potentially devastating bacterial wilt disease that seriously affects plant production worldwide (Mansfield et al., 2012). The RSSC is composed of four phylogenotypes (Fegan & Prior, 2006) and assigned to three distinct species: Ralstonia pseudosolanacearum (phylogenotypes I and III), R. solanacearum (phylogenotype II), and Ralstonia syzygii (phylogenotype IV) (Safni et al., 2014).

RSSC strains first invade the intercellular spaces of plant roots (Araud-Razou et al., 1998; Hikichi et al., 2017; Vasse et al., 1995). After invading the roots, the bacterial cells attach to the surface of host cells, where they avoid plant innate immune responses (Genin & Denny, 2012; Hikichi et al., 2017; Kiba et al., 2018, 2020; Nakano et al., 2013) and grow vigorously to activate QS (Hikichi et al., 2017). The bacteria then invade xylem vessels, with the bacteria systemically spreading and multiplying throughout the xylem to activate QS (Genin & Denny, 2012; Hikichi et al., 2017; Vasse et al., 1995). This bacterial proliferation in the xylem leads to wilting symptoms in infected tomato plants. QS-deficient mutants lose their virulence and their ability to invade xylem vessels (Genin & Denny, 2012; Hayashi et al., 2019a; Hikichi et al., 2017; Schell, 2000; Senuma et al., 2020). In addition, QS is thought to be conserved in all RSSC strains and is thus a required network for RSSC virulence (Castillo & Agathos, 2019; Genin & Denny, 2012).

Ralstonia has developed a genus-specific QS system consisting of Phc QS cascade regulatory elements that respond to a unique fatty acid derivative signal (Flavier et al., 1997). Each strain of the RSSC produces either methyl 3-hydroxypalmitate (3-OH PAME) or methyl 3-hydroxymyristate (3-OH MAME) as a QS signal (Flavier et al., 1997; Kai et al., 2015; Schell, 2000; Ujita et al., 2019). RSSC strains synthesize the QS signal by the methyltransferase PhcB and sense the chemical through the sensor histidine kinase PhcS, activating QS (Genin & Denny, 2012; Kai et al., 2015; Schell, 2000; Ujita et al., 2019). The sensor histidine kinase PhcK is required for full expression of phcA, which encodes the LysR family transcriptional regulator PhcA, independently of QS signal production of PhcB (Senuma et al., 2020). In the active state of QS, PhcA activated through QS signal sensing of PhcS regulates QS-dependent genes responsible for QS-dependent phenotypes including virulence (Genin & Denny, 2012). This process leads to the induction of the production of ralfuranones, which are aryl-furanone secondary metabolites, and the major exopolysaccharide EPS I, which is involved in the virulence of RSSC strains (Genin & Denny, 2012; Kai et al., 2014, 2015; Pauly et al., 2013; Schell, 2000; Wackler et al., 2011). Furthermore, these secondary metabolites are also associated with the feedback loop of QS-dependent gene regulation (Hayashi et al., 2019b; Mori et al., 2018).

The phc operon consists of phcB and phcS, which encode Phc QS cascade regulatory elements, along with phcR and phcQ (Clough et al., 1997; Salanoubat et al., 2002; Tang et al., 2020). PhcR is an intracellular soluble two-component protein composed of a sensor histidine kinase domain and a receiver domain without a DNA-binding domain. PhcQ is an intracellular soluble regulator containing a receiver domain without a DNA-binding site. Schell (2000) proposed a model for the QS signalling pathway of the phylotype IIA strain AW1 of R. solanacearum, which produces 3-OH PAME as the QS signal (Flavier et al., 1997). At low levels of 3-OH PAME, the regulator protein PhcR may interact with PhcA, leading to PhcA dysfunction. At the threshold concentration of 3-OH PAME, the sensor histidine kinase PhcS senses 3-OH PAME and carries out the phosphorylation by itself. This event may induce the ability of PhcS to phosphorylate the cognate PhcR, resulting in functional PhcA. Furthermore, Tang et al. (2020) demonstrated that PhcQ is involved in the dynamics of activation of PhcA in response to the bacterial density of phylotype I strain GM1000 of R. pseudosolanacearum, which produces 3-OH MAME as the QS signal (Kai et al., 2015). However, the effects of these regulator proteins on the transcription of QS-dependent genes have not been validated experimentally.

In this study, we aimed to elucidate the function of PhcR and PhcQ in the regulation of QS-dependent genes. To achieve this goal, we first created phcR-deletion (ΔphcR) and phcQ-deletion (ΔphcQ) mutants from the phylotype I strain OE1-1 of R. pseudosolanacearum, which produces 3-OH MAME as the QS signal (Kai et al., 2015), and analysed their QS-dependent phenotypes as well as their virulence on tomato plants. We then analysed the transcriptomes of R. pseudosolanacearum strains using quantitative reverse transcription-PCR (RT-qPCR) and RNA-sequencing (RNA-seq).

2 | RESULTS

2.1 | Phylogenetic analysis of the deduced amino acid sequences of PhcR and PhcQ among RSSC strains

To analyse the genetic variation of PhcR and PhcQ among 34 RSSC strains (phylogenotype I, nine strains; phylogenotype IIA, six strains;
phytotype IIB, five strains; phytotype III, three strains; and phytotype IV, 11 strains, including blood disease bacterial strain R229 and *R. syzygii* strain R24; Table S1), the deduced amino acid sequences of PhcR and PhcQ were analysed with ClustalW and phylogenetic trees were constructed with TreeView. The phylogenetic trees regarding PhcR (Figure S1a) as well as PhcQ (Figure S1b) showed that the 34 strains were divided into four clades, consistent with their phylotypes.

### 2.2 | Deletion of phcQ, but not phcR, led to significant changes in QS-dependent phenotypes

We created ΔphcR (Table 1) and ΔphcQ (Table 1) mutants of strain OE1-1 and analysed their QS-dependent phenotypes. We first assayed biofilm formation with crystal violet staining of *R. solanacearum* strains grown in quarter-strength M63 medium. The ΔphcR mutant exhibited slightly less biofilm formation than wild-type strain OE1-1 (Figure 1a). The ΔphcQ mutant produced significantly less biofilm than strain OE1-1, similar to the ΔphcB mutant (Table 1; Kai et al., 2015) and the ΔphcA mutant (Table 1; Mori et al., 2016) (p < .05, t test). The ΔphcR mutant produced less EPS I (Figure 1b) and more ralfuranone A (Figure 1c), which is one of the ralfuranones, than strain OE1-1, whereas the phcQ deletion led to significantly reduced production of EPS I (p < .05, t test; Figure 1b) and ralfuranone A (p < .05, t test; Figure 1c), similar to the effects of phcB and phcA deletions. Compared with strain OE1-1, the ΔphcR mutant, when grown on quarter-strength M63 medium solidified with 0.25% agar, exhibited slightly enhanced swimming motility (Figure 1d). The swimming motility of the ΔphcQ mutant was significantly greater than that of strain OE1-1, similar to the ΔphcB and ΔphcA mutants (p < .05, t test).

The induced expression of two genes—epsB, which is part of the eps operon and required for EPS I biosynthesis (Huang & Schell, 1995), and ralA, which encodes a ralfuranone synthase (Kai et al., 2014; Wackler et al., 2011)—is dependent on QS. In addition, the expression of the flagellar motility-related gene fliC, which encodes flagellin, is suppressed in the active state of QS (Tans-Kersten et al., 2001). To analyse expression levels of these genes in *R. pseudosolanacearum* strains grown in quarter-strength M63 medium until OD_{600} = 0.3, we conducted RT-qPCR assays. Expression levels of ralA and epsB in the ΔphcQ mutant but not in the ΔphcR mutant were significantly lower than those in OE1-1 (p < .05, t test; Figure 2). In contrast, fliC was more highly expressed in the ΔphcQ mutant, but not in the ΔphcR mutant, compared with the OE1-1 strain (p < .05, t test).

| TABLE 1 | Strains and plasmids used in this study |
|----------|----------------------------------------|
| **Plasmids** | **Relevant characteristics** | **Source** |
| pUC18 | Amp′ | Takara Bio |
| pK18mobSacB | Km′, oriT (RP4), sacB, lacZα | Kvitko and Collmer (2011) |
| pUC18-mini-Tn7-T-Gm | Gm′ | Choi et al. (2005) |
| pTNS2 | Helper plasmid carrying T7 transposase gene | Choi et al. (2005) |
| pMD20delta-phcR | pMD20 derivative carrying a 1.4-kb DNA fragment for phcR deletion, Amp′ | This study |
| pdelta-phcR | pK18mobSacB derivative carrying a 1.4-kb DNA fragment for phcR deletion, Km′ | This study |
| pdelta-phcQ | pK18mobSacB derivative carrying a 1.5-kb DNA fragment for phcR deletion, Km′ | This study |
| pUC18-mini-Tn7-T-Gm-phcQ | pUC18-mini-Tn7-T-Gm derivative carrying a 2.8-kb fragment for phcR complementation, Gm′ | This study |
| **Escherichia coli strain** | | |
| DH5α | recA1 endA1 gyrA96 thi-1 hsdR17supE44 Δ(lacU169)lac80lacΔM15 | Takara Bio |
| **R. solanacearum strains** | | |
| OE1-1 | Wild-type strain, phytotype I, race 1, biovar 4 | Kanda et al. (2003) |
| ΔphcB | phcB-deletion mutant of OE1-1 | Kai et al. (2015) |
| ΔphcA | phcA-deletion mutant of OE1-1 | Mori et al. (2016) |
| ΔphcK | phcK-deletion mutant of OE1-1 | Senuma et al. (2020) |
| ΔphcR | phcR-deletion mutant of OE1-1 | This study |
| ΔphcQ | phcQ-deletion mutant of OE1-1 | This study |
| phcQ-comp | A transformant of ΔphcQ with pUC18-mini-Tn7-T-Gm-phcQ containing native phcQ, Gm′ | This study |
| lecM-M | lecM::Tn EZ::TN <KAN-2> mutant, Km′ | Mori et al. (2016) |
2.3 | Transformation of native PhcQ recovered QS-dependent phenotypes of the ΔphcQ mutant

We next transformed the ΔphcQ mutant with pUC18-mini-Tn7T-Gm-phcQ (Table 1) harbouring the native phcQ gene fused with the promoter of the phcBSRQ operon (Kai et al., 2015) to generate the complemented ΔphcQ mutant strain phcQ-comp (Table 1). Transformation of the ΔphcQ mutant with the pUC18-mini-Tn7T-Gm-phcQ construct led to enhanced biofilm formation (Figure 1a) and production of EPS I (Figure 1b) and ralfuranone A (Figure 1c) and reduced swimming motility (Figure 1d).

2.4 | Deletion of phcQ, but not phcR, led to a loss of bacterial virulence

To investigate the effects of phcR and phcQ on the virulence of strain OE1-1, we inoculated 5-week-old tomato plants with R. pseudosolanacearum strains by the root-dip method and then assayed the population dynamics and behaviour of these strains in the tomato plants as well as disease development. The population of the ΔphcQ mutant at 3 days after inoculation (DAI) was significantly smaller than that of OE1-1 and the ΔphcR mutant, similar to ΔphcB and ΔphcA mutants (p < .05, t test; Figure 3a). In a plate-printing
assay, we detected OE1-1 and the ΔphcR mutant in the inoculated roots and stems of tomato plants, whereas no ΔphcQ was observed beyond the inoculated roots, similar to ΔphcB and ΔphcA mutants (Figure 3b). The tomato plants inoculated with the OE1-1 strain exhibited wilt symptoms at 5 DAI and died by 10 DAI (Figure 3c). The ΔphcQ mutant was not virulent on tomato plants 10 DAI, similar to the ΔphcB and ΔphcA mutants, whereas the virulence of the ΔphcR mutant resembled that of the OE1-1 strain.

2.5 | Deletion of phcR or phcQ did not influence the regulation of QS-related genes

The ΔphcQ mutant exhibited significantly changed QS-dependent phenotypes including the virulence compared to strain OE1-1, though the QS-dependent phenotypes of the ΔphcR mutant were largely unchanged (Figures 2 and 3). To analyse the effects of deletion of phcR and phcQ on the gene expression of phcB, phcK, and phcA (QS-related genes), we monitored the expression levels of QS-related genes in R. pseudosolanacearum strains grown in quarter-strength M63 medium until OD_{600} = 0.3 by RT-qPCR. No significant differences in expression levels of phcB, phcK, and phcA were detected among the ΔphcQ mutant, the ΔphcR mutant, and OE1-1 (p < .05, t test; Figure 2). Furthermore, expression levels of phcQ and phcR in the ΔphcR mutant and ΔphcQ mutant, respectively, were similar to those in OE1-1 (p < .05, t test).

2.6 | RNA-seq transcriptome analysis of R. pseudosolanacearum strains

The deletion of phcR or phcQ did not influence the regulation of QS-related genes (Figure 2). To analyse the effects of phcR and phcQ deletion on the regulation of QS-dependent genes, we performed an RNA-seq transcriptome analysis of R. pseudosolanacearum strains grown in quarter-strength M63 medium until OD_{600} = 0.3. Mapping of RNA-seq reads of the OE1-1 strain to the GMI1000 genome (Salanoubat et al., 2002) resulted in the identification of 4,437 protein-coding transcripts (Table S2). To extract genes with significant expression changes, the following thresholds were applied: q < .05 and |log(fold change [FC])| ≥ 2. Compared with their expression levels in OE1-1, 371 and 174 genes in the ΔphcB mutant were significantly down-regulated and up-regulated, respectively (Figure S2). Among them, 345 and 163 genes in the ΔphcA mutant were significantly down-regulated and up-regulated, respectively, and were thus inferred to be positively and negatively QS-dependent genes, respectively. The transcriptome analysis with RNA-seq showed that 97 genes in the ΔphcR mutant were significantly down-regulated compared with their expression levels in OE1-1, and were thus inferred to be positively PhcR-regulated genes (Figure 4a; Table S3), while 67 genes (negatively PhcR-regulated genes) in the ΔphcR mutant were significantly up-regulated (Figure 4b; Table S3). Among the positively PhcR-regulated genes, 79 genes, including norB, lecM, and xpsR, were positively QS-dependent genes (Figure 4a; Table S3). Among the
negatively PhcR-regulated genes, 43 genes, including some flagellin biosynthesis-related genes and chemotaxis-related genes, were negatively QS-dependent genes (Figure 4b; Table S3). The log(FC) values of these PhcR-regulated QS-dependent genes between strain OE1-1 and the ΔphcR mutant were moderately correlated with those between OE1-1 and the ΔphcB mutant (y, log(FC) of ΔphcB; x, log(FC) of ΔphcR; y = 1.5886x − 0.371, r² = .8727; Figure 4c) or the ΔphcA mutant (y, log(FC) of ΔphcA; x, log(FC) of ΔphcR; y = 1.6716x − 0.4498, r² = .8761; Figure 4c). It is thus thought that PhcR is partially involved in the regulation of 79 (22.9%) and 43 (26.4%) of the positively and negatively QS-dependent genes, respectively.

The transcriptome analysis with RNA-seq showed that in the ΔphcQ mutant, 435 positively PhcQ-regulated genes were significantly down-regulated (Figure 4a; Table S4) and 197 negatively PhcQ-regulated genes were significantly up-regulated relative to strain OE1-1 (Figure 4b; Table S4). Among the positively PhcQ-regulated genes, 334 genes were positively QS-dependent genes (Figure 4a; Table S4). Among the negatively PhcQ-regulated genes, 109 genes were negatively QS-dependent genes (Figure 4b; Table S4). A strong positive correlation of the expression of these QS-dependent genes among PhcQ-regulated genes was observed between the ΔphcQ mutant and the ΔphcB mutant (y, log(FC) of ΔphcB; x, log(FC) of
ΔphcQ; $y = 0.9308x + 0.1356$, $r^2 = 0.9432$; Figure 4d) and between the ΔphcQ mutant and the ΔphcA mutant ($y$, log(FC) of ΔphcA; $x$, log(FC) of ΔphcQ; $y = 0.9721x + 0.0186$, $r^2 = 0.9402$; Figure 4d).

Therefore, PhcQ is mainly involved in the regulation of 334 (96.8%) and 109 (66.9%) of the positively and negatively QS-dependent genes, respectively.

We then carried out hierarchical clustering of the QS-related gene-deletion mutants and the ΔphcR and ΔphcQ mutants based on their relative expression levels normalized against those of QS-dependent genes. In the resulting dendrogram, the ΔphcR mutant clustered with strain OE1-1, whereas the ΔphcQ mutant grouped with the ΔphcB mutant, and ΔphcA clustered separately with the phcK-deletion (ΔphcK) mutant (Figure 4e).

### 2.7 Deletion of phcQ led to a significant reduction in 3-OH MAME content

Tang et al. (2020) demonstrated that PhcQ contributes to the synthesis of 3-OH MAME by strain GM1000. In the active state of QS, expression of lecM, which encodes the lectin LecM, is induced and
LecM affects the activation of QS through regulating the stability of secreted 3-OH MAME (Hayashi et al., 2019a). Deletion of phcR or phcQ did not influence the regulation of QS-related genes including phcB (Figure 2). We assayed 3-OH MAME content purified from R. pseudosolanacearum strains. The lecM-Δ mutant exhibited a significantly lower 3-OH MAME content than strain OE1-1, similar to the ΔphcQ mutant (p < .05, Figure 5). Though the ΔphcR mutant exhibited significantly lower 3-OH MAME content compared to strain OE1-1, 3-OH MAME content in the ΔphcR mutant was higher than that in other mutants, including the ΔphcA mutant.

To analyse the expression levels of lecM in R. pseudosolanacearum strains grown in quarter-strength M63 medium until OD<sub>600</sub> = 0.3, we conducted RT-qPCR assays. The expression levels of lecM in the ΔphcQ and ΔphcA mutants was significantly lower than in strain OE1-1 (p < .05, t-test; Figure 2). Though the expression level of lecM in the ΔphcR mutant was significantly lower than in strain OE1-1, the lecM expression level in the ΔphcR mutant was higher than in the ΔphcA mutant.

2.8 | Deletion of phcR led to a slight change in expression levels of QS-dependent genes at lower bacterial density

In the inactive state of QS, PhcR reportedly inhibits PhcA function in strain AW1 (Schell, 2000). To analyse the influence of phcR deletion on expression levels of QS-dependent genes in the inactive state of QS, using RNA isolated from the ΔphcR mutant and strain OE1-1 grown in quarter-strength M63 medium until OD<sub>600</sub> = 0.01, we conducted RT-qPCR assays to assess relative expression levels of lecM, ralA, epsB, and fliC in the ΔphcR mutant compared to strain OE1-1. The expression levels of the positively QS-dependent genes lecM, ralA, and epsB were slightly but significantly lower in the ΔphcR mutant than in strain OE1-1 (p < .05, t-test; Figure 6). In addition, the expression level of the negatively QS-dependent fliC was significantly higher in the ΔphcR mutant than in strain OE1-1 (p < .05, t-test).

2.9 | Exogenous 3-OH MAME application did not lead to a change in the QS-dependent phenotypes of ΔphcR and ΔphcQ mutants

QS activity in strain OE1-1 is dependent on the exogenous levels of 3-OH MAME (Hayashi et al., 2019b; Kai et al., 2015). We examined the influence of exogenous 3-OH MAME application on QS-dependent phenotypes, biofilm formation, EPS I production, and swimming motility of these mutants. Exogenous application of 3-OH MAME at a concentration of 0.1 μM enhanced biofilm formation and EPS I production and reduced swimming motility of strain OE1-1 and the ΔphcB mutant (p < .05, t-test; Figure 7a). However, exogenous 3-OH MAME application had no effect on these QS-dependent phenotypes of ΔphcR and ΔphcQ mutants.

To analyse the influence of exogenous 3-OH MAME application on the expression levels of the QS-dependent genes lecM, ralA, epsB, and fliC, we conducted RT-qPCR assays using R. pseudosolanacearum strains grown in quarter-strength M63 medium containing 3-OH MAME at a concentration of 0.1 μM until OD<sub>600</sub> = 0.3. Exogenous 3-OH MAME application significantly enhanced the expression levels of lecM, ralA, and epsB and significantly reduced the expression levels of fliC in strain OE1-1 and the ΔphcB mutant but not in the ΔphcQ mutant (p < .05, t-test; Figure 7b). In contrast, in the ΔphcR mutant, exogenous 3-OH MAME application slightly but significantly enhanced the expression levels of lecM and epsB, but not ralA, and slightly but significantly reduced the expression levels of fliC (p < .05, t-test).

2.10 | PhcA was involved in the negative control of siderophore-mediated iron acquisition activity, independently of PhcR and PhcQ

RSSC strains produce micacidin and staphyloferrin B as siderophores to acquire iron from extracellular environments (Bhatt & Denny, 2004; Kreutzer et al., 2011). The RSPO424 gene (ssd),

![Figure 6](image-url)

**Figure 6**: Expression of the positively QS-dependent genes lecM, ralA, and epsB, and the negatively QS-dependent gene fliC in Ralstonia pseudosolanacearum strains OE1-1 and ΔphcR grown in quarter-strength M63 medium until OD<sub>600</sub> = 0.01, as determined by quantitative reverse transcription-PCR. Two replicate experiments conducted using independent samples with eight technical replicates per experiment produced similar results. Results of a single representative sample are provided. Bars indicate standard errors. Asterisks indicate values significantly different from those of OE1-1 (p < .05, t-test)
siderophore-mediated iron acquisition activity, we measured the influence of PhcR and PhcQ on siderophore-mediated iron acquisition activity of R. pseudosolanacearum strains OE1-1, ΔphcB, ΔphcR, and ΔphcQ. (b) Expression levels of the QS-dependent genes epsB, ralA, and fliC in the same strains. (a) For the biofilm formation and EPS I production assays, R. pseudosolanacearum strains were grown in quarter-strength M63 or quarter-strength M63 medium containing 0.1 µM 3-OH MAME. For the swimming motility assay, 5-µl aliquots of cell suspensions at 5 × 10^8 cfu/ml were added to the centre of plates containing quarter-strength M63 medium solidified with 0.25% agar, and the swimming-area diameters were measured at 24 hr postinoculation. Three replicate experiments conducted using independent samples with seven technical replicates per biofilm formation experiment and five technical replicates per EPS I production or swimming motility experiment produced similar results. The results of a representative experiment are shown. Bars indicate standard errors. Asterisks indicate values significantly different from those of OE1-1 (p < .05, t test). (b) Total RNA was extracted from bacterial cells grown until OD_600 = 0.01. Bars indicate standard errors. Asterisks indicate values significantly different from those of OE1-1 (p < .05, t test)

Balancing selection is a type of positive selection that favours the maintenance of a high genetic diversity within a given population and functions through spatial/temporal heterogeneity as well as overdominant selection and frequency-dependent selection (Charlesworth, 2006; Hedrick, 2012; Stoeckel et al., 2012). Under balancing selection, the phcBSRQ operon plays a significant role in virulence of RSSC strains. (Castillo & Agathos, 2019). PhcS is subject to strong selection from the plant host (Guidot et al., 2014). Based on the deduced PhcB and PhcS amino acid sequences, RSSC strains are divided into two groups according to their QS signal types (Kai et al., 2015). In contrast, the phylogenetic analysis based on the deduced amino acid sequences of PhcR and PhcQ indicates that RSSC strains can be divided according to their phylotypes, similar to that of PhcA (Senuma et al., 2020). Overall, it is thought that extracellular communications through the production and sensing of QS signals reveal the unique evolution among RSSC strains independently of their phylotypes. The following intracellular signalling is under balancing...
Selection according to their phylotypes and is involved in bacterial environmental fitness including virulence.

PhcQ is reportedly involved in the dynamics of activation of PhcA in the active state of QS (Tang et al., 2020). PhcQ is required for systemic infectivity of strain OE1-1 in tomato plants and its virulence on tomato plants, similar to the other QS cascade regulatory elements. The present transcriptome analysis results of the ΔphcQ mutant imply that PhcQ is involved in the regulation of 96.8% and 66.9% of positively and negatively QS-dependent genes, respectively. Furthermore, the expression levels of these PhcQ-regulated genes in the ΔphcQ mutant were strongly correlated with those in ΔphcB or ΔphcA. We previously demonstrated that the putative sensor histidine kinase PhcK is required for full phcA expression independently of 3-OH MAME content (Senuma et al., 2020). However, the deletion of phcQ had no effect on the expression level of phcA. In the hierarchical cluster dendrograms of PhcA-regulated genes in QS-related gene-deletion mutants, the ΔphcQ mutant clustered with the ΔphcB mutant, and the ΔphcA mutant grouped with the ΔphcK mutant. Furthermore, the exogenous application of 3-OH MAME did not lead to a change in the expression levels of QS-dependent genes and QS-dependent phenotypes of the ΔphcQ mutant. It is thus thought that PhcQ may mainly contribute to the regulation of 96.8% and 66.9% of positively and negatively QS-dependent genes, respectively, by PhcA, dependent on 3-OH MAME content.

The supernatant of the phcQ mutant generated from strain GMI1000 does not induce the expression of the positively QS-dependent xpsR, demonstrating involvement of PhcQ in 3-OH MAME production (Tang et al., 2020). However, the influence of extracellular 3-OH MAME application on the expression levels of QS-related genes and other QS-dependent genes in the ΔphcQ mutant is not experimentally validated. In the present study, PhcQ was involved in the induced expression of lecM but not phcB, thereby influencing the stability of secreted 3-OH MAME and thus PhcA activation by QS. Furthermore, phcQ deletion significantly reduced QS-inducible production of secondary metabolites, EPS I and ralfuranones, which are associated with the feedback loop of QS-dependent gene regulation (Hayashi et al., 2019b; Mori et al., 2018). It is thus thought that phcQ deletion leads to a change in the regulation of QS-dependent genes through extracellular 3-OH MAME content and the feedback loop through EPS I and ralfuranones as well as PhcA activation.

The present transcriptome analysis results showed that PhcR is involved in the regulation of 22.9% and 26.4% of positively and negatively QS-dependent genes, respectively, in strain OE1-1. However, the expression levels of these genes in the ΔphcR mutant were moderately correlated with those in ΔphcB or ΔphcA. The exogenous application of 3-OH MAME did not lead to a significant change in the QS-dependent phenotypes of ΔphcR, but led to a slight but significant change in the expression levels of the QS-dependent
genes lecM, epsB, and fliC, but not ralA, in the ΔphcR mutant. Our transcriptome analysis showed that phcR deletion led to a slight but significant change in the expression levels of lecM, epsB, and fliC, but not ralA. As PhcQ mainly contributes to the regulation of these QS-dependent genes, dependent on 3-OH MAME content, the change in expression levels of lecM, epsB, and fliC in the ΔphcR mutant upon 3-OH MAME treatment may result from enhanced function of PhcQ in the regulation of QS-dependent genes. It is thus thought that PhcR may partially contribute to the regulation of 22.9% and 26.4% of positively and negatively QS-dependent genes, respectively, by PhcA, dependently on 3-OH MAME content. This leads to partial influences on QS-dependent phenotypes of the ΔphcR mutant but not its systemic infectivity and virulence on tomato plants.

The QS model of strain AW1 proposed by Schell (2000) suggests that in the inactive state of QS, the expression of positively and negatively QS-dependent genes should be higher and lower, respectively, in the ΔphcR mutant than in the wild type. However, regardless of bacterial density, the ΔphcR mutant exhibited slightly lower and higher expression levels of positively and negatively QS-dependent genes, respectively, compared to the wild-type strain OE1-1 (Figures 2 and 6), suggesting PhcR partially contributes to the regulation of QS-dependent genes by PhcA. Clough et al. (1997) demonstrated that the deletion of phcR has no obvious effect on the production of EPS I by strain AW1 incubated in EG medium under shaking. The phylogenetic analysis using the deduced amino acid sequences of PhcR indicated that PhcR-mediated intracellular signalling pathways involved in bacterial environmental fitness may be under balancing selection according to phylotypes of RSSC strains. Incubation under different conditions may thus lead to differences in PhcR function in QS-dependent phenotypes between strains OE1-1 (phylotype I) and AW1 (phylotype IIA), though further experimental validations are required.

Strain AW1 produces staphyloferrin B as a siderophore, and ssd, which is negatively regulated by PhcA, is required for staphyloferrin B production and siderophore-mediated iron acquisition activity (Bhatt & Denny, 2004). The strains used in the present study showed siderophore-mediated iron acquisition activity in the order ΔphcA = ΔphcK < ΔphcB < ΔphcQ < OE1-1 < ΔphcR. PhcA regulated not only ssd but also Rsc1271, Rsc2918, Rsc2919, Rsc2920, Rsp0419, and Rsc0421, independently of PhcR and PhcQ. In the hierarchical cluster, the ΔphcA mutant grouped with the ΔphcK mutant, and ΔphcQ clustered separately with the ΔphcB mutant. Furthermore, though the ΔphcR mutant also clustered with strain OE1-1 instead of with the other QS-deficient mutants in the hierarchical cluster, the expression patterns of Rsc1805, Rsc1806, and Rsp0100 in the ΔphcR mutant were different from those in strain OE1-1. PhcR, PhcQ, and PhcA were thus all found to be independently involved in the control of siderophore-mediated iron acquisition activity. Furthermore, the regulation of 3.2% of positively QS-dependent genes and 33.1% of negatively QS-dependent genes was independent of both PhcR and PhcQ.

Strain OE1-1 produces and secretes 3-OH MAME and senses the chemical through PhcS, activating QS (Kai et al., 2015; Ujita et al., 2019). PhcR (Clough et al., 1997) and PhcQ (Tang et al., 2020) are regulator proteins with a receiver domain without a DNA-binding site. Single-receiver-domain proteins play roles as phospho-relays or phospho-sinks (Feldheim et al., 2018). Considering the results of the present study, we propose the following scenario to explain the Phc QS signalling cascade of strain OE1-1. PhcQ may mainly contribute to the activation of PhcA required for the regulation of 96.8% and 66.9% of positively and negatively QS-dependent genes, respectively, dependent on extracellular 3-OH MAME content. PhcR may be partially involved in the activation of PhcA required for the regulation of 22.9% and 26.4% of positively and negatively QS-dependent genes, respectively, along with PhcQ, dependent on extracellular 3-OH MAME content. In addition, an unknown regulator may be involved in the activation of PhcA required for the regulation of 3.2% and 33.1% of positively and negatively QS-dependent genes, respectively. In vitro experiments with purified proteins and phosphorylation assays could be required to demonstrate that PhcQ and PhcR can be phosphorylated by PhcS. The LysR-type transcriptional regulator PhcA has a conserved structure with an N-terminal DNA-binding helix-turn-helix motif and a C-terminal co-inducer-binding domain (Maddock & Oyston, 2008). However, little information on the co-inducer of PhcA is available. The roles of PhcR and PhcQ elucidated in this study in the regulation of QS-dependent genes shed light on the activation mechanisms of PhcA function in the active state of QS.

4 | EXPERIMENTAL PROCEDURES

4.1 | Bacterial strains, plasmids, and growth conditions

We used the following R. pseudosolanacearum strains in this study: OE1-1 (Kanda et al., 2003), the phcA-deletion mutant ΔphcA (Mori et al., 2016), the phcB-deletion mutant ΔphcB (Kai et al., 2015), and the phcK-deletion mutant ΔphcK (Senuma et al., 2020). All R. pseudosolanacearum strains were routinely grown in quarter-strength M63 medium at 30°C. Escherichia coli strains were grown in Luria–Bertani medium (Hanahan, 1983) at 37°C. Gentamycin (50 μg/ml) was used in selective media.

4.2 | Phylogenetic analysis

Phylogenetic trees were constructed using deduced amino acid sequences of PhcB (464–468 amino acids), PhcA (347 amino acids), PhcR (368–374 amino acids), and PhcQ (336–337 amino acids) of RSSC strains. We constructed phylogenetic trees using the ClustalW program according to the neighbour-joining method (Saitou & Nei, 1987), with genetic distances computed with Kimura’s two-parameter model (Kimura, 1980). The phylogenetic trees were drawn with TreeView.
4.3 | Generation of the phcR-deletion and phcQ-deletion mutants

The oligonucleotide primers used for construction of recombinant plasmids are listed in Table S6. The fragments delta-R-1 and delta-R-2 for generation of the \( \Delta \text{phcR} \) mutant and the fragments delta-Q-1 and delta-Q-2 for generation of the \( \Delta \text{phcQ} \) mutant were amplified by PCR from the genomic DNA of strain OE1-1. The fragments delta-R and delta-Q amplified using delta-R-1 and delta-R-2 were ligated into a pK18mobsacB vector (Kvitko & Collmer, 2011) to produce recombinant plasmids pdelta-phiCR and pdelta-phiCQ, respectively. Each plasmid was electroporated into OE1-1 competent cells, which were prepared as previously described by Mori et al. (2016). Kanamycin-sensitive, sucrose-resistant recombinants \( \Delta \text{phcR} \) and \( \Delta \text{phcQ} \) were then selected.

4.4 | Generation of a phcQ-deletion mutant transformed with native phcQ

Using the oligonucleotide primers listed in Table S5, the fragments souho-Q-1 and souho-Q-2 were amplified by PCR from the genomic DNA of strain OE1-1. The fragment souho-Q amplified using souho-Q-1 and souho-Q-2 were ligated into a pUC18-mini-Tn7T-Gm vector (Choi et al., 2005) to produce psouhoQ. This plasmid was electroporated into OE1-1 competent cells with a T7 transposase expression vector pTNS2 (Choi et al., 2005). Finally, a gentamycin-resistant transformant, phcQ-comp, was selected.

4.5 | Transcriptome analysis based on RNA-seq

Total RNA was extracted from \( R. \text{pseudosolanacearum} \) strains grown in quarter-strength M63 until OD\(_{600} \) = 0.3 with a High Pure RNA Isolation kit (Roche Diagnostics). Ribosomal RNA was eliminated from the extracted total RNA using a Ribo-Zero RNA Removal kit (gram-negative bacteria; Illumina) as previously described (Hayashi et al., 2019a). Oriented, paired-end RNA-seq (2 \( \times \) 100 bp) was performed on an Illumina HiSeq 2000 system. The generated reads were trimmed with Cutadapt v. 1.1 (http://code.google.com/p/cutadapt/) and Trimmomatic v. 0.32 (http://www.usadellab.org/cms/?page=trimmomatic) and then mapped with the TopHat program v. 2.0.10 (http://tophat.cbcb.umd.edu/). Three independent biological replicates were carried out per strain.

4.6 | Differential gene expression analysis

Statistical analysis of the RNA-seq data was performed in the R environment. Genes with zero counts in at least one OE1-1 sample were excluded. RNA-seq read counts of the remaining genes were normalized using the function calcNormFactors (trimmed mean of M value normalization) in the package edgeR (Robinson et al., 2010).

To extract genes with significant expression changes, the following thresholds were applied: \( q < 0.05 \) and \(|\log(FC)| \geq 2\). The false discovery rate \( (q \text{ value}) \) was calculated in edgeR from Benjamini–Hochberg-corrected \( p \) values. Hierarchical clustering of all normalized mean expression values based on their relative expression (counts per million) was performed using Cluster v. 3.0 software (de Hoon et al., 2004). The average value of three replicates per strain was used. Heatmaps were created in TreeView (Eisen et al., 1998).

4.7 | RT-qPCR

Total RNA was extracted from \( R. \text{pseudosolanacearum} \) strains grown in quarter-strength M63 until OD\(_{600} \) = 0.01 or OD\(_{600} \) = 0.3 with a High Pure RNA Isolation kit. To determine the influence of exogenous 3-OH MAME application on the regulation of QS-dependent genes, total RNA was extracted from \( R. \text{pseudosolanacearum} \) strains grown in quarter-strength M63 containing 0.1 \( \mu \text{M} \) 3-OH MAME until OD\(_{600} \) = 0.3 with a High Pure RNA Isolation kit. An RT-qPCR assay with gene-specific primers (Table S7) was carried out using the SYBR GreenER qPCR Reagent system (Invitrogen) on a 7300 Real-Time PCR system (Applied Biosystems) as previously described (Hayashi et al., 2019a). All values were normalized against the expression level of rpoD, which was used as an internal standard for each cDNA sample. No significant differences in rpoD expression levels were observed among \( R. \text{pseudosolanacearum} \) strains. Two replicate experiments conducted using independent samples with eight technical replicates per experiment produced similar results. Results of a single representative sample are provided.

4.8 | QS-dependent phenotypes

We examined the in vitro biofilm formation of \( R. \text{pseudosolanacearum} \) cells grown in quarter-strength M63 solidified with 1.5% agar with a High Pure RNA Isolation kit. To determine the influence of exogenous 3-OH MAME application on biofilm formation, \( R. \text{pseudosolanacearum} \) strains were grown without shaking in quarter-strength M63 containing 0.1 \( \mu \text{M} \) 3-OH MAME as previously described (Senuma et al., 2020). The biofilm formation was quantified on the basis of the absorbance at 550 nm (\( A_{550} \)). The resulting value was normalized according to the number of cells (OD\(_{600} \)). Three replicate experiments conducted using independent samples with seven technical replicates per experiment produced similar results. The results of a representative experiment are shown.

EPS I production by \( R. \text{pseudosolanacearum} \) cells grown on quarter-strength M63 solidified with 1.5% agar was quantitatively analysed in an enzyme-linked immunosorbent assay (Agdia) as previously described (Mori et al., 2016). To determine the influence of exogenous 3-OH MAME application on EPS I production, \( R. \text{pseudosolanacearum} \) strains were grown on quarter-strength M63 solidified with 1.5% agar containing 0.1 \( \mu \text{M} \) 3-OH MAME. EPS I production
was quantified according to the absorbance at 650 nm ($A_{650}$). Three replicate experiments conducted using independent samples with five technical replicates per experiment produced similar results. The results of a representative experiment are shown.

To assay ralfuranone A production, $R$. pseudosolanacearum strains were incubated in B medium at 30 °C for 4–6 hr and then diluted to an OD$_{600}$ of 0.05 in new MGRL medium containing 3% sucrose (Ujita et al., 2019). The cell suspensions (2 ml) were incubated in 15-ml test tubes at 30 °C with shaking for 2 days. Following growth, ralfuranone A was extracted from the culture supernatants with MonoSpin C18 (GL Science) according to the manufacturer’s instructions. The methanol eluates from the MonoSpin C18 extraction were analysed for ralfuranones by liquid chromatography–mass spectrometry (LC-MS) using an InertSustain C18 column (150 mm × 2.1 mm, 3 μm) under the following conditions: eluent = 20%–95% MeCN in 0.1% aqueous formic acid (0–24 min) and 95% MeCN (24–30 min); column temperature, 40 °C; flow rate, 200 μl/min; injection volume, 5 μl. The experiment was conducted three times using independently prepared samples.

For the swimming motility assay, overnight cultures of $R$. pseudosolanacearum strains were washed with distilled water and then diluted to a cell density of $5 \times 10^5$ cfu/ml, and 5-μl aliquots of cell suspensions were added to the centre of plates containing quarter-strength M63 medium solidified with 0.25% agar. The swimming-area diameters of $R$. pseudosolanacearum strains were measured at 48 hr postinoculation (Mori et al., 2018). To determine the influence of exogenous 3-OH MAME application on swimming motility, 5-μl aliquots of cell suspensions without or with 3-OH MAME at a concentration of 0.1 μM were added to the centre of plates containing quarter-strength M63 medium solidified with 0.25% agar. The swimming-area diameters of $R$. pseudosolanacearum strains were measured at 24 hr postinoculation. Three replicate experiments conducted using independent samples with five technical replicates per experiment produced similar results. The results of a representative experiment are shown.

### 4.9 | 3-OH MAME contents produced by $R$. pseudosolanacearum strains

Strains of $R$. pseudosolanacearum grown in B medium at 30 °C for 4–6 hr were diluted to an OD$_{600}$ of 1.0 with new medium. The cell suspension at a volume of 50 μl was pipetted onto a BG agar plate (90 mm, 25 ml; Kai et al., 2015), and the plate was incubated for 24 hr at 30 °C. The BG agar was cut into small pieces and soaked in ethyl acetate at a volume of 50 ml for 2 hr twice. The combined extracts were dried over Na$_2$SO$_4$ and concentrated. The residue was dissolved in acetonitrile (100 ml) and subjected to LC-MS analysis. LC-MS data were recorded with an LCMS-2020 (Shimadzu) and an InertSustain C18 column (150 mm × 2.1 mm, 3 μm particle size; GL Sciences). The conditions used were as follows: injection, 5 μl; solvent A, 0.1% formic acid; solvent B, acetonitrile containing 0.1% formic acid; gradient, 20%–95% B (0–24 min), 95% B (24–30 min); flow rate, 200 ml/min; detection, positive SIM mode; data reflect four replicates each.

### 4.10 | Siderophore-mediated iron acquisition activity

The siderophore-mediated iron acquisition activity of $R$. pseudosolanacearum strains was analysed using a method modified from Wali et al. (2015). $R$. pseudosolanacearum strains were incubated in PY medium (5 g/L polypeptone and 2 g/L yeast extract) for 18 hr at 30 °C and adjusted to a concentration of $2 \times 10^5$ cfu/ml with 0.1 M PIPES buffer (pH 6.5). After 6 hr of incubation, each culture was filtered through a 0.2-μm pore filter. Next, 100 μl of a culture, or PIPES buffer alone as a reference, was added to 100 μl of chromazurol S (CAS) solution (2.4 mM hexadecyl-trimethyl ammonium bromide, 0.06 mM FeCl$_3$, 0.6 mM HCl, 0.6 mM CAS in PIPES buffer). Absorbance at 630 nm ($A_{630}$) was measured after incubation for 30 min at 30 °C. The activity of the siderophore was calculated by subtracting the absorbance value of the reference from the total absorbance. Three replicate experiments conducted using independent samples with eight technical replicates per experiment produced similar results. Results of a single representative sample are provided.

### 4.11 | Virulence assays

Eight-week-old tomato plants (Solanum lycopersicum ‘Ohgata-Fukuju’) were inoculated with $R$. pseudosolanacearum strains ($10^8$ cfu/ml) using a root-dip inoculation procedure as previously described (Hayashi et al., 2019a).

We assessed populations of $R$. pseudosolanacearum strains in inoculated tomato roots according to their observed growth on Hara–Ono medium as described by Hayashi et al. (2019a). For each bacterial strain, three replicate experiments conducted using independent samples with five technical replicates per experiment produced similar results. Results of a single representative sample are provided.

The behaviour of $R$. pseudosolanacearum strains in tomato plants inoculated by the root-dip method was assessed as described by Hayashi et al. (2019a). A sample from each cut site (Figure 3b) was pressed onto Hara–Ono medium. Twelve plants were analysed per trial, and each assay comprised five successive trials.

Plants were monitored daily for wilting symptoms, which were rated according to the following disease index scale: 0, no wilting; 1, 1%–25% wilting; 2, 26%–50% wilting; 3, 51%–75% wilting; 4, 76%–99%; and 5, dead. For each bacterial strain, three replicate experiments conducted using independent samples with 12 technical replicates per experiment produced similar results. Results of a single representative sample are provided.
4.12 | Statistical analysis

The means of all assays were analysed for significant differences between *R. pseudosolanacearum* strains by Student’s *t* test in Microsoft Excel.

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CONFLICT OF INTEREST
The authors declare that they have no conflicts of interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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SUPPORTING INFORMATION
Additional Supporting Information may be found online in the Supporting Information section.

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