Research article

Genomic, genetic and structural analysis of pyoverdine-mediated iron acquisition in the plant growth-promoting bacterium *Pseudomonas fluorescens* SBW25

Christina D Moon¹,5, Xue-Xian Zhang¹,2,6, Sandra Matthijs³, Mathias Schäfer⁴, Herbert Budzikiewicz⁴ and Paul B Rainey*¹,2,6

Address: ¹Department of Plant Sciences, University of Oxford, South Parks Rd, Oxford OX1 3RR, UK, ²School of Biological Sciences, University of Auckland, Private Bag 92019, Auckland, New Zealand, ³Laboratory of Microbial Interactions, Department of Molecular and Cellular Interactions, Flanders Interuniversity Institute for Biotechnology, Vrije Universiteit Brussel, Building E, Pleinlaan 2, Brussels, Belgium, ⁴Institut für Organische Chemie der Universität zu Köln, Greinstr. 4, 50939, Köln, Germany, ⁵AgResearch Limited, Grasslands Research Centre, Private Bag 11008, Palmerston North, New Zealand and ⁶Institute of Molecular Biosciences and New Zealand Institute for Advanced Study, Massey University, Private Bag 102904, Auckland, New Zealand

Email: Christina D Moon - christina.moon@agresearch.co.nz; Xue-Xian Zhang - x.x.zhang1@massey.ac.nz; Sandra Matthijs - sm Matthi@vub.ac.be; Mathias Schäfer - aco78@uni-koeln.de; Herbert Budzikiewicz - aco88@uni-koeln.de; Paul B Rainey* - p.b.rainey@massey.ac.nz

* Corresponding author

Abstract

Background: Pyoverdines (PVDs) are high affinity siderophores, for which the molecular mechanisms of biosynthesis, uptake and regulation have been extensively studied in *Pseudomonas aeruginosa* PAO1. However, the extent to which this regulatory model applies to other pseudomonads is unknown. Here, we describe the results of a genomic, genetic and structural analysis of pyoverdine-mediated iron uptake by the plant growth-promoting bacterium *P. fluorescens* SBW25.

Results: In silico analysis of the complete, but un-annotated, SBW25 genome sequence identified 31 genes putatively involved in PVD biosynthesis, transport or regulation, which are distributed across seven different regions of the genome. PVD gene iron-responsiveness was tested using *lacZ* fusions to five PVD loci, representative of structural and regulatory genes. Transcription of all fusions increased in response to iron starvation. In silico analyses suggested that regulation of fpvR (which is predicted to encode a cytoplasmic membrane-spanning anti-sigma factor) may be unique. Transcriptional assays using gene expression constructs showed that fpvR is positively regulated by FpvI (an extracytoplasmic family (ECF) sigma factor), and not directly by the ferric uptake regulator (Fur) as for PAO1. Deletion of *pvdL*, encoding a predicted non-ribosomal peptide synthetase (NRPS) involved in PVD chromophore biosynthesis confirmed the necessity of PvdL for PVD production and for normal growth in iron-limited media. Structural analysis of the SBW25 PVD shows a partly cyclic seven residue peptide backbone, identical to that of *P. fluorescens* ATCC13525. At least 24 putative siderophore receptor genes are present in the SBW25 genome enabling the bacterium to utilize 19 structurally distinct PVDs from 25 different *Pseudomonas* isolates.

Conclusion: The genome of *P. fluorescens* SBW25 contains an extensively dispersed set of PVD genes in comparison to other sequenced *Pseudomonas* strains. The PAO1 PVD regulatory model, which involves a branched Fpv signaling pathway, is generally conserved in SBW25, however there is a significant difference in fpvR regulation. SBW25 produces PVD with a partly cyclic seven amino acid residue backbone, and is able to utilize a wide variety of exogenous PVDs.
Background
Iron is a common co-factor for redox-dependent enzymes and an essential element for almost all living organisms. In nature iron is abundant; however, in aerobic environments and under general physiological conditions, iron typically exists in the insoluble ferric (Fe3+) form, thus rendering acquisition by organisms difficult. Under conditions of iron limitation, many bacteria produce and secrete low molecular weight ferric-specific ligands known as siderophores (for review, see [11]). The ferrisiderophores deliver iron to the cell via specific receptor and transport systems (for review, see [2]).

Collectively, Pseudomonas spp. produce a wide variety of siderophores [3], the most complex and common of which are pyoverdines (PVDs) [4]. PVDs contain a peptide moiety, usually between 6–12 amino acids in length, and a dihydroxyquinoline chromophore moiety [4], which gives PVD its characteristic yellow-green fluorescent appearance. PVD-mediated iron uptake processes have been extensively characterized in Pseudomonas aeruginosa PAO1, where key regulatory and structural proteins have been identified, and regulatory mechanisms have been elucidated [5-9].

In P. aeruginosa PAO1, the primary level of PVD regulation involves the ferric uptake regulator (Fur), which, upon interaction with ferrous iron (Fe2+), binds to a specific DNA sequence (the Fur-box) in the promoter region of certain iron-regulated genes and blocks transcription. Under iron-deplete conditions (no Fe2+ available to interact with Fur) Fur-dependent repression is relieved [6,10]. PAO1 PVD genes that possess Fur boxes, which have been demonstrated to bind Fur in vitro [5,11], are pvdS and fpvL, which encode extracytoplasmic family (ECF) sigma factors, and fpvR, an inner membrane-spanning anti-sigma factor. The alternative sigma factor, PvdS, when associated with core RNA polymerase, binds iron-starvation (IS) box motifs [12] and directs the expression of a suite of genes involved in the synthesis of PVD [7,13]. The activity of PvdS is regulated by a transmembrane signaling system that comprises of FpvR and FpvA, an outer membrane PVD receptor, and is primarily mediated by PVD [5]. More recently, a second ECF sigma factor (FpvI) was found to direct transcription of fpvA [5,14]. Like PvdS, FpvI is also under the direct regulation of FpvR and forms a second divergent branch of the signaling pathway.

Many plant-associated pseudomonads are known to produce PVDs, including the plant pathogen, P. syringae and the saprophytes, P. putida and P. fluorescens. The importance of siderophores for rhizosphere colonization by P. putida has been previously reported [15,16]. However, little is known about the molecular mechanisms of PVD production and regulation in these organisms.

Here we report a genetic characterization of genes for PVD production in a plant growth-promoting bacterium, Pseudomonas fluorescens SBW25. SBW25 was originally isolated from field-grown sugar beet [17] and its biocontrol activity against the soil-borne pathogen, Pythium ultimum, is related to its considerable plant colonization ability [18]. The present investigation was prompted by initial findings that three genes implicated in iron-acquisition processes were up-regulated on plant surfaces, as revealed by in vivo expression technology (IVET) analysis [19,20]. Two genes had significant homology to putative siderophore receptor genes, and one gene appeared to be homologous to PAO1 pvdL, which encodes a non-ribosomal peptide synthetase (NRPS) involved in PVD biosynthesis [21]. These findings suggested a significant role for iron uptake during seedling colonization.

Our work began with the un-annotated whole genome sequence of SBW25 [22]. Based initially on an analysis of this sequence, 31 genes were identified with predicted roles in PVD biosynthesis, transport and regulation, for which the regulation of a subset of these was investigated by genetic analysis using chromosomally-integrated lacZ fusions and gene expression constructs. The results indicate a not previously realized mechanism of fpvR regulation. Moreover, we determine the chemical structure of SBW25 PVD and examine the biological role of PVD via a siderophore-deficient mutant. The ability of SBW25 to utilize a panel of structurally distinct exogenous PVDs was also assessed.

Results and Discussion
Genomic analysis of PVD genes in P. fluorescens SBW25
Interrogation of the complete genome sequence of SBW25 using genes known to be involved in PVD-mediated iron acquisition in P. aeruginosa PAO1 [23] revealed 31 homologues with predicted roles in PVD biosynthesis, transport and regulation. The results are summarized in Figure 1 and Additional file 1.

SBW25 homologues were identified for all PAO1 PVD genes except PA2411, which encodes a thioesterase, and pvdI and pvdD, which encode NRPSs involved in biosynthesis of the PVD peptide backbone. In PAO1, PA2411 is not essential for PVD production, and is possibly redundant since pvdG also encodes a thioesterase [23]. Although homologues of pvdI and pvdD were not identified in SBW25, three additional NRPS genes were identified: two downstream of pvdP (Pflu2552 and Pflu2553) and one downstream of pvdI (Pflu2544; Figure 1). Differences in the sequences and organization of these NRPS genes likely reflect differences between the structures of the PVDs produced by SBW25 and PAO1 (discussed below).
The PVD genes of SBW25 are distributed across seven different loci within the genome (Figure 1), which is the most widespread distribution of PVD genes in a genomic comparison among four various sequenced *Pseudomonas* spp. [23]. This distribution is in stark contrast to that of *P. syringae* where PVD genes form one large cluster [23]. In *P. aeruginosa* PAO1 the PVD genes are confined to two loci separated by just 11.5 kb (Figure 1), which appears to be typical of *P. aeruginosa* isolates in general, where almost all PVD genes are located within the same region of the genome [24]. PVD gene homologues in *P. putida* KT2440 and *P. fluorescens* strains Pf0–1 [22] and Pf-5 [25] are distributed across three loci.

Despite the overall differences in the genomic organization of PVD genes between SBW25 and PAO1, genes within each of the seven SBW25 PVD clusters are largely syntenous with those in PAO1. There are, however, two exceptions: *pvdQ* and *fpvR*, which in SBW25 are found in two non-adjacent regions of the genome and separated from all other PVD genes. The isolated position of *fpvR* (especially in relation to *fpvI* – see Figure 1), is particularly unusual: the genes for ECF sigma factors and their cognate anti-sigma factors are typically adjacent, and usually co-regulated by Fur [2,26,27]. The gene arrangement in SBW25 suggests that *fpvR* and *fpvI* might be under separate regulatory control – a notion given credence by the fact that the promoter sequence of *fpvR* in SBW25 does not have a recognizable Fur-box motif. Interestingly, in place of the expected Fur-box motif is an IS-box motif, albeit one in which the last three bases of the consensus sequence are in reverse order (TAAAT-N16-TGC, rather than TAAAT-N16-CGT). The possibility that in SBW25 *fpvR* is opposed to that of *P. aeruginosa* PAO1 [5], the effects of the two constitutively expressed ECF sigma factors (PvdS, FpvI) and the anti-sigma factor (FpvR) on downstream structural genes (*pvdL* and *fpvA*) were assessed. Expression constructs (pTR130.1, pTR130.2 and pTR130.3 for *pvdS*, *fpvI* and *fpvR* respectively) were created.

**Expression of putative PVD genes in response to iron starvation**

To ascertain whether the SBW25 PVD gene homologues identified by *in silico* analyses are responsive to iron, a selection of genes were assessed using chromosomally integrated 'lacZ' transcriptional fusions. Genes were chosen that represented key regulatory and structural components in PVD mediated iron-uptake, and included the ECF sigma factor encoding genes *pvdS* and *fpvI*, the anti-sigma factor encoding gene *fpvR*, and structural genes involved in the biosynthesis and uptake of PVD, *pvdL* and *fpvA*, respectively (fusion strains TR107.1.1, TR107.4.1, TR107.5.1, TR107.2.1, and TR135.1.1 respectively; Table 1). The β-galactosidase activity of each fusion strain was assessed under high iron (450 μM FeSO₄), moderate iron (450 nM FeSO₄), and low iron (100 μM 2,2’-dipyridyl) conditions. All gene fusions showed decreasing levels of activity as iron concentration was increased (Figure 2), which is consistent with genes that are directly or indirectly under the control of the Fur repressor. The most active gene tested was *fpvA*, the activity of which was almost 3-fold greater than the next most active gene, *pvdS*. To determine whether SBW25 possesses a branched Fpv signaling pathway analogous to that in PAO1 [5], the effects of the two constitutively expressed ECF sigma factors (PvdS, FpvI) and the anti-sigma factor (FpvR) on downstream structural genes (*pvdL* and *fpvA*) were assessed. Expression constructs (pTR130.1, pTR130.2 and pTR130.3 for *pvdS*, *fpvI* and *fpvR* respectively) were created.
in the vector pBroadgate which resulted in the cloned genes coming under the control of the constitutive P lac promoter. The expression constructs, and corresponding control vector, pBroadgate-D, were introduced into the pvdL-'lacZ (TR107.2.1) and fpvA-'lacZ (TR135.1.1) fusion strains, and their activities were monitored in β-galactosidase assays under high and low iron regimes in CAA medium. Expression of pvdL and fpvA was significantly increased as a consequence of constitutive expression of pvdS and fpvI, respectively (Figure 3A and 3B), consistent with the notion that PvdS and FpvI are alternative ECF sigma factors that direct regulation of each of these genes. In both cases, an increase in expression was observed even in the presence of iron, consistent with Fur-independent regulation of the cloned regulators. Furthermore, expression of fpvR caused a significant decrease in the level of fpvA expression, consistent with FpvR being an anti-sigma factor in the Fpv signaling pathway model [5]. For pvdL, repression by FpvR was observed under high iron conditions, though not to a statistically significant level. Thus to verify the presence of the Fpv branched signaling pathway in SBW25, PVD induction of the pathway was demonstrated by providing purified SBW25 PVD to the pvdL-'lacZ and fpvA-'lacZ fusion strains. Here, the exogenous PVD stimulated the signaling pathway, resulting in significant up-regulation of both pvdL (1.9-fold) and fpvA (3.0-fold; both results significant at P = 0.05 using a one-tail t-test). Taken together, these data are consistent with SBW25 possessing a branched Fpv signaling pathway that is activated by PVD.

### Table 1: Bacterial strains and plasmids used in this study.

| Strain/plasmid | Relevant genotype and/or other characteristics | Reference |
|----------------|-----------------------------------------------|-----------|
| **P. fluorescens** | | |
| SBW25 | Wild-type | [17] |
| PBR840 | SBW25ΔpvdL or ΔPflu4387 | This study |
| 1SF3 | PBR840 with mini-Tn phoA3 disruption of secondary siderophore biosynthesis genes, GmR | S. Matthijs, unpublished |
| TR107.1.1<sup>a</sup> | SBW25 carrying the pvdS-'lacZ fusion, SmR | This study |
| TR107.2.1<sup>a</sup> | SBW25 carrying the pvdL-'lacZ fusion, SmR | This study |
| TR107.4.1<sup>a</sup> | SBW25 carrying the fpv::lacZ fusion, SmR | This study |
| TR107.5.1<sup>a</sup> | SBW25 carrying the fpvR::lacZ fusion, SmR | This study |
| TR135.1.1<sup>a</sup> | SBW25 carrying the fpvA::lacZ fusion, SmR | This study |
| TR137.2.1 | TR107.2.1 with pTr130.1, GmR | This study |
| TR137.3.1 | TR107.2.1 with pTr130.2, GmR | This study |
| TR137.4.1 | TR107.2.1 with pTr130.3, GmR | This study |
| TR156.1.1 | TR107.2.1 with pBroadgate-D, GmR | This study |
| TR138.2.1 | TR135.1.1 with pTr130.1, GmR | This study |
| TR138.3.1 | TR135.1.1 with pTr130.2, GmR | This study |
| TR138.4.1 | TR135.1.1 with pTr130.3, GmR | This study |
| TR156.3.1 | TR135.1.1 with pBroadgate-D | This study |
| TR168.1.1 | TR107.5.1 with pTr130.1, GmR | This study |
| TR168.2.1 | TR107.5.1 with pTr130.2, GmR | This study |
| TR168.3.1 | TR107.5.1 with pTr130.3, GmR | This study |
| TR168.4.1 | TR107.5.1 with pBroadgate-D | This study |
| **Plasmids** | | |
| pUIC3 | Integration vector with 'lacZ, TcR | [20] |
| pHPS45<sup>Ω</sup> | Source of the omega cassette, SmR | [51] |
| plVET-Sm | pUIC3 with tet gene replaced by Smr/Spr<sup>Ω</sup> fragment from pHPS45<sup>Ω</sup>, SmR | This study |
| pUIC3-40 | pUIC3 containing pvdL deletion fragment | This study |
| pCR2.1 | PCR product cloning vector, KmR | Invitrogen |
| pRK2013<sup>a</sup> | helper plasmid, KmR, Tra<sup>+</sup> | [52] |
| pTR100.1.1 | pIVET-Sm carrying the pvdS-'lacZ fusion | This study |
| pTR100.2.1 | pIVET-Sm carrying the pvdL-'lacZ fusion | This study |
| pTR100.4.1 | pIVET-Sm carrying the fpvL-'lacZ fusion | This study |
| pTR100.5.1 | pIVET-Sm carrying the fpvR-'lacZ fusion | This study |
| pTR128.1.1 | pIVET-Sm carrying the fpvA-'lacZ fusion | This study |
| pBroadgate | Gateway expression vector with P<sub>tra</sub> promoter, GmR | Thwaites and Mansfield |
| pBroadgate-D | pBroadgate with ccdB gene replaced with the linker fragment, GmR<sup>Ω</sup>; | This study |
| pTR130.1 | pBroadgate::pvdS, GmR | This study |
| pTR130.2 | pBroadgate::fpvL, GmR | This study |
| pTR130.3 | pBroadgate::fpvR, GmR | This study |

<sup>a</sup> The 'lacZ fusion reporter was integrated into the PVD locus by insertion-duplication thus the gene function was not affected.
Unique regulation of fpvR in SBW25

Next we tested the hypothesis that fpvR is regulated by PvdS rather than Fur, which is the direct regulator of anti-sigma factors in many other bacterial iron-acquisition systems [27]. The regulatory gene constitutive expression constructs (pTR130.1, pTR130.2 and pTR130.3 for pvdS, fpvI and fpvR respectively) were introduced into the fpvR-'lacZ fusion strain (TR107.5.1). β-galactosidase activities of the resulting strains were determined under high (450 µM FeSO₄) and low (100 µM 2,2'-dipyridyl) iron regimes. Contrary to expectation, the results presented in Figure 3C showed that expression of FpvI, and not PvdS, resulted in the up-regulation of fpvR, even in the presence of iron. It is unknown whether FpvI directly regulates fpvR, or if an intermediate regulatory component is involved. However, a conserved sequence motif, 5’-TAATGAGAA-3’, was identified 56 nt upstream of fpvA and 128 nt upstream of fpvR, which may potentially serve as an FpvI binding site.

Genomic comparisons suggest that the non-adjacent arrangement of fpvI and fpvR in SBW25 may not be unique. Homologues of fpvR were not reported in P. put.

Figure 2
PVD gene activities in response to iron. The activities of pvdL, pvdS, fpvA, fpvl and fpvR in response to iron were determined from lacZ reporter fusion strains, TR107.2.1, TR107.1.1, TR135.1.1, TR107.4.1, and TR107.5.1, respectively. β-galactosidase activity (fmol 4 MU/min/cell) was assayed for cells growing in CAA media supplemented with 100 µM 2,2'-dipyridyl (black bars), 450 nM FeSO₄ (hatched bars), and 450 µM FeSO₄ (white bars). Data are the means ± the standard errors of four independent experiments for each strain. Asterisks indicate that the promoter activity in the given iron treatment (100 µM 2,2'-dipyridyl or 450 nM FeSO₄) was significantly different to the background activity in 450 µM FeSO₄ at the P = 0.05 level of significance using Dunnett’s method.

Figure 3
PVD gene activities in response to constitutively-expressed putative regulators. The activities of pvdL-lacZ (A), fpvA-lacZ (B), and fpvR-lacZ (C) (fusion strains: TR107.2.1, TR135.1.1, and TR107.5.1, respectively), in response to constitutively-expressed regulator genes pvdS, fpvl and fpvR (in expression constructs pTR130.1, pTR130.2 and pTR130.3, respectively). β-galactosidase activity (fmol 4 MU/min/cell) was assayed for cells growing in CAA media supplemented with 100 µM 2,2'-dipyridyl (black bars), or 450 µM FeSO₄ (white bars). Data are the means ± the standard errors of four independent experiments. Asterisks indicate that under the given iron regime, the reporter strain activity with expression construct significantly differs from the background activity of the reporter strain containing the pBroadgate-D (vector) using Dunnett’s method at the P = 0.05 level of significance.
ida KT2440 and *P. fluorescens* Pf0–1, and *fpvR* and *fpvi* were not identified in *P. syringae* pv. tomato DC3000 [23], however, this is likely to reflect a lack of strong sequence homology to the characterized counterparts in PAO1. Indeed, in SBW25, the amino acid sequence identity between FpvR and its PAO1 homologue was the lowest among all genes examined (48.7%; Additional file 1). A search for *fpvR* genes in other sequenced *P. fluorescens* strains revealed putative homologues in Pf0–1 and Pf-5 (48.1% and 48.4% amino acid sequence identity to PAO1 FpvR, respectively), however neither gene was adjacent to its cognate *fpvi* homologue, nor possessed a recognizable Fur-box motif in the upstream sequence. A putative IS-box was observed 44 bp upstream of the Pf0–1 homologue, with 7 out of 8 bases matching the IS-box consensus. However, an IS-box was not observed for the Pf-5 homologue, nor was the potential Fpvi-binding site sequence identified upstream of SBW25 *fpva* and *fpvR*. The presence of key genes encoding components of the Fpv branched signaling pathway (*fpva*, *fpvR*, *pvdS* and *fpvi*) in the Pf0–1 and Pf-5 genomes suggests that this pathway may be conserved in these *P. fluorescens* strains – a hypothesis that can be tested in future work.

**Deletion analysis of pvdL**

To investigate the biological roles of the PVD genes predicted from *in silico* analysis, an SBW25 derivative strain containing a deletion of *pvdL* was constructed (PBR840, ∆*pvdL*). In SBW25, *pvdL* (*Pflu4387*) is predicted to encode a 4297 amino acid NRPS involved in PVD chromophore biosynthesis. PBR840 displayed a PVD negative phenotype when grown on LB agar: wild-type SBW25 produced a typical yellow-green siderophore, whereas PBR840 did not (Figure 4A). It was also observed that colonies of PBR840 were notably smaller than those of SBW25 (Figure 4A). It is unknown whether this is solely due to impairment in iron uptake, or whether an iron-independent factor negatively affects PBR840 growth.

The growth kinetics of PBR840 were determined in iron-limited CAA broth and the same medium supplemented with 450 µM FeCl₃ in microtitre plate cultures. Results are shown in Figure 4B. A repeated measures MANOVA (see figure caption for details) showed that the optical density of PBR840 cells was significantly reduced compared to wild-type in iron deplete medium from 14 h [F₁,₈ = 28.02, *P* < 0.0001]. No significant reduction was detected during exponential growth, for example, at 4, 6 and 8 h, *P* values were 0.849, 0.333 and 0.398, respectively. No significant difference in growth was observed between wild-type and PBR840 in iron replete medium [F₁,₈ = 0.03, *P* = 0.862].

Growth analysis of PBR840 took place initially in 200 µl microtitre plate cultures, which although incubated with periodic shaking, could have become oxygen limited thus

**Figure 4**

*Growth phenotype of mutant PBR840 (∆pvdL) in iron-limited media.* **A:** Growth of SBW25 and PBR840 on LB agar for two days, after which the mutant produced no visible PVD and colonies were smaller than wild-type. **B:** Growth in CAA broth (SBW25, clear circle; PBR840, filled circle) and in CAA broth supplemented with 450 µM FeCl₃ (SBW25, clear square; PBR840, filled square). Results are shown for clarity. Data from each medium were analyzed separately using a multivariate model (MANOVA) with repeated measures (2 h intervals): no significant difference was detected between genotypes grown in CAA supplemented with FeCl₃ [F₁,₈ = 0.03, *P* = 0.862], however, a significant genotype effect was evident in iron deficient medium [F₁,₈ = 37.22, *P* < 0.001]. An analysis at each time point revealed a highly significant difference between genotypes from (and including) the 14 h time point [F₁,₈ = 28.02, *P* < 0.0001]. **C:** PBR840 cells were plated on CAA media with EDDHA and overlaid with filter paper disks impregnated with sterile water (left) and 50 nmol purified SBW25 PVD (right). PBR840 was not able to grow on the strongly iron-chelated media, growth was restored when supplemented with PVD.
masking the effect of the \textit{pwdL} mutation during exponential growth. To test this hypothesis, growth of PBR840 and SBW25 was further investigated using strongly aerated cultures grown in CAA broth, where iron is more likely to be present in the ferric form. The growth dynamics (data not shown) were comparable to those observed in the static cultures, with no statistical difference in cell density being detected during exponential growth, however differences in final yield were evident at 24 hours (the mean A450 reading and standard error for PBR840 was 0.9239 ± 0.014 (n = 4), and for SBW25 was 1.3510 ± 0.016 (n = 4)). Furthermore, PBR840 was incapable of growing in CAA medium supplemented with the strong iron-chelator EDDHA (ethylenediaminedihydroxyphenylacetic acid, 0.5 mg/ml), whereas growth was restored upon provision of purified SBW25 PVD (Figure 4C). Taken together, the data confirmed the predicted role of \textit{pwdL} (Pflu4387) in PVD production.

\textbf{Structural analysis of SBW25 PVD}

The chemical structure of SBW25 PVD was determined by mass spectral analysis using standard methods as previously described [28]. Comparison of the mass spectra to published data [29-32] revealed that SBW25 PVD comprises the characteristic PVD quinoline chromophore [33] linked to a partly cyclic seven residue peptide backbone, identical to that of \textit{P. fluorescens} ATCC 13525 [34]: Lys-Gly-\textit{fOHO}rn-\textit{c}(Lys-fOHO}rn-Ser) (fOHO}rn = N\textsuperscript{5}-formyl-N\textsuperscript{3}-hydroxyornithine and underlined residues are in the \textit{D} configuration) (Figure 5). The side chain that is connected amidically to the NH\textsubscript{2}-group of the chromophore varied as succinic acid or ketoglutaric acid. The peptide backbone sequence differs in length and composition to that of PAO1 PVD (Ser-Arg-Ser-fOHO}rn-c(Lys-fOHO}rn-Thr-Thr) [35]). \textit{In silico} analyses [36,37] of the protein sequences of the putative PVD NRPSs: PvdL, Pflu2544, Pflu2552, Pflu2553 and PvdL, were performed to predict their amino acid substrates (results are shown in Additional file 2). Comparison of the substrate predictions with the actual PVD structure suggested that only PvdL and Pflu2544 are involved in the biosynthesis of the SBW25 PVD backbone, with predictions for three of the amino acid residues being correct (Ser, Gly and Ser). Dhb/sal-like residues were consistently predicted for both occurrences of fOHO}rn for PvdL and Pflu2544 (see Additional file 2). It is unclear why the algorithm has made this prediction, as fOHO}rn does not share similar physico-chemical properties with dhb and sal, however, similar analyses of the PAO1 PVD NRPSs have also consistently predicted dhb/sal-like specificities for the fOHO}rn binding modules (Additional file 2). The roles of Pflu2552 and Pflu2553 are presently unknown, however the presence of an IS-box-like motif preceding these genes, and their proximity to other PVD gene homologues (Figure 1) suggests that they have, or have had, a role in PVD-mediated iron-uptake processes. As the SBW25 PVD chromophore structure is identical to that of PAO1 PVD [33], it was not surprising that the \textit{in silico} substrate predictions for SBW25 PvdL were the same as those for PAO1 PvdL (Additional file 2), although only one of the three residues was predicted correctly (glu) in each case.

\textbf{Utilization of exogenous PVDs by \textit{P. fluorescens} SBW25}

The identification of two SBW25 plant-induced genes with homology to siderophore receptor genes [19,20] prompted us to examine the ability of SBW25 to uptake and utilize exogenous siderophores. Initially, a genome-wide search for putative TonB-dependent siderophore receptor genes was undertaken, which revealed 24 putative siderophore receptor genes dispersed throughout the genome (listed in Additional file 3). Nine of these were immediately adjacent to predicted sigma factor/anti-sigma factor gene pairs (Additional file 3), that are presumably involved in their regulation. The number of putative receptor genes is similar to that (26) for \textit{P. fluorescens} Pf0–1 [3], and include \textit{fpvA} (Pflu2545), and Pflu5798 and Pflu6132 which were both previously shown to be induced \textit{in planta} [19]. These results strongly support the notion that SBW25 has the capability to utilize a considerable number of exogenous iron-chelating siderophores produced by other microbes.

To test the ability of SBW25 to utilize exogenous PVDs, PVDs purified from 25 different \textit{Pseudomonas} isolates (Table 2), each with a different siderotype profile (P. Corneils, personal communication), were supplied to 15F3 (a secondary siderophore-deficient mutant strain of PBR840) that had been overlaid as a dilute suspension onto CAA medium chelated with EDDHA. The mutant was unable to grow when water was provided as a negative control. However, growth was restored by provision of PVDs purified from 19 different isolates (Table 2). These include the three types (I, II and III) of PVD from \textit{P. aeru-
ginosa. Thus, it is apparent that SBW25 is able to utilize a range of exogenous PVDs produced by other pseudomonads.

Conclusion

The genomic era has revolutionized the scientific method of comparative biology, where the general applicability of biological traits and systems that have often been evaluated in few model organisms may now be readily assessed across taxa. Although PVDs are common siderophores of Pseudomonas spp., the regulation of the genes involved in the biosynthesis and uptake of PVD has been elucidated almost exclusively in P. aeruginosa PAO1. The availability of the P. fluorescens SBW25 genome sequence has revealed that PVD genes are considerably more widespread throughout the SBW25 genome than in other sequenced pseudomonads, and has enabled a comparative analysis of PVD regulation in SBW25 and PAO1 to be undertaken. The PVD regulatory model established for PAO1 involves a unique branched signaling pathway [5]. Prior to this study it was not known whether this model applies to other Pseudomonas isolates. Using chromosomally-integrated gene reporter fusions and gene expression constructs, we found that the PAO1 branched signaling model is present in SBW25. In addition, we reveal a significant difference between PAO1 and SBW25 in terms of the regulation of fpvR. In SBW25, over-expression of fpvI results in constitutive expression of fpvR, suggesting that fpvR is positively regulated by FpvI, rather than directly by Fur, as is the case of fpvR of PAO1 [5]. The structure of SBW25 PVD was elucidated and it was found that the peptide backbone is identical to that of P. fluorescens ATCC 13525, comprising a partly cyclic peptide of seven residues. The PVD structure implicated the involvement of two NRPSs in the synthesis of the PVD peptide backbone. The SBW25 genome harbors 24 putative TonB-dependent siderophore receptors, two of which are previously shown to be plant-inducible [19]. Exogenous PVD uptake assays demonstrate that SBW25 is able to utilize a variety of structurally distinct siderophores produced by other Pseudomonas isolates. These data lay the foundation for further detailed functional analysis of these putative siderophore receptors in term of regulation and transporter specificity.

Methods

Bacterial strains, plasmids, and growth conditions

Escherichia coli DH5αλpir [38] was used as a recipient strain for gene cloning and then a donor for conjugative transfer into Pseudomonas cells. Pseudomonas strains and plasmids used in this study are listed in Table 1. E. coli cul-

| Strain PVD isolated from | 15F3 growth stimulation | Reference |
|--------------------------|-------------------------|-----------|
| P. aeruginosa PAO1a       | +                       | [40, 54]  |
| P. aeruginosa 7NSK2       | +                       | [55]      |
| P. aeruginosa 59.20b      | +                       | [39]      |
| P. aeruginosa LMG 2112    | +                       | BCCM/LMG  |
| P. aeruginosa LMG 5147    | +                       | BCCM/LMG  |
| P. fluorescens LMG 14562  | +                       | BCCM/LMG  |
| P. fluorescens 17400::qbsF | -                      | [56]      |
| P. fluorescens 5          | +                       | A. Sarniguet, INRA, France |
| P. fluorescens C7R12      | +                       | P. Lemanceau, INRA/Université de Bourgogne, France |
| P. fluorescens SBW25      | +                       | [17]      |
| P. libanensis LMG 21606   | -                       | BCCM/LMG  |
| P. salomonii LMG 22120    | +                       | BCCM/LMG  |
| P. syringae LMG 1247      | +                       | BCCM/LMG  |
| P. vancouverensis LMG 20222 | -                     | BCCM/LMG  |
| Pseudomonas sp. W2Ap9      | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Aug1     | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Aug7     | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Aug36    | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Dec18    | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Dec29    | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Dec33    | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W15Feb31B  | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W2Jun14    | -                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W15Ap2     | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W15Aug1    | +                       | S. Matthijs, VUB, Brussels |
| Pseudomonas sp. W15Oct32   | +                       | S. Matthijs, VUB, Brussels |

*P. aeruginosa* strains PAO1, 7NSK2 and 59.20 produce types I, II and III PVDs, respectively.

*BCCM/LMG: Belgian co-ordinated collections of micro-organisms bacteria collection

Secondary siderophore (thioquinolobactin) production is inactivated in this strain.
tures were grown at 37°C in Luria-Bertani (LB) medium [38] and P. fluorescens cultures were grown at 28°C in LB or casamino acids medium (CAA) [39]. Antibiotics were used at the following final concentrations (µg/ml): ampicillin (Ap) 100, chloramphenicol (Cm) 25, gentamycin (Gm) 20, kanamycin (Km) 50, streptomycin (Sm) 100 and tetracycline (Tc) 10. 5-Bromo-4-chloro-3-indolyl β-D-galactopyranoside (X-gal; Melford Laboratories, Chelsworth, UK) was used at a final concentration of 40 µg/ml.

To monitor PVD gene activities, reporter strains were grown in the presence of 450 nM – 450 µM FeSO₄ for iron supplemented media. To chelate iron, 100 µM 2, 2’-dipyridyl (Sigma, St Louis MO, USA), or 0.5 mg/ml ethylenediaminedi hydroxyphenylacetic acid (EDDHA) was used. Purified PVD was supplied at 50 µM.

Growth curves of bacterial strains were obtained using a VersaMax microtiter plate reader with SOFTmax PRO software (Molecular Devices, Sunnyvale CA, USA). Inoculum was prepared by culturing strains stored at -80°C in LB broth overnight, then sub-culturing into CAA broth with growth overnight. For growth experiments, cells from the CAA culture were washed once with sterile water and 2 µl was used inoculate 200 µl CAA or CAA supplemented with 450 µM FeCl₃ in wells of the 96-well microtiter plate. Plates were incubated at 28°C and absorbance readings at 450 nm were taken every five minutes after brief shaking. Growth curves were also obtained from highly aerated cultures grown in 250 ml flasks containing 20 ml CAA broth, and inoculated to an initial OD450 of ~0.15. Cultures were grown at 28°C with shaking at 200 rpm and absorbance readings at 450 nm were taken over a 24 hour period.

In silico analyses
Putative homologues of PVD-related genes in P. fluorescens SBW25 were identified by interrogating the un-annotated SBW25 genome [22] with P. aeruginosa PAO1 PVD gene sequences [23,40] using nucleotide-nucleotide Basic Local Alignment Search Tool (BLASTN). Complete SBW25 open reading frames (ORFs) were identified from matching sequence hits, and these were used to re-query the PAO1 genome via translated query vs. protein sequence BLAST (tBLASTX) algorithms. Sequences upstream of putative homologues were examined by eye for the PAO1 Fur-box consensus sequence, 5’-GATAAT-GATAAATCATTATC-3’ [11], and iron starvation (IS) boxes, 5’-TAAAT-N₁₆-CGT-3’ [12,41]. The best matches to Fur-box motifs were confirmed using MotifScanner [42], in conjunction with a Fur-box position weight matrix (accession MX000013) from the Prodiric website [43,44]. Genes were mapped to the genome using Artemis software [45], and the percentage identity and similarity at the amino acid sequence level between PAO1 and SBW25 homologues was determined by the Needleman-Wunsch global alignment algorithm implemented in EMBOSS needle [46]. Common DNA sequence motifs were identified by MEME version 3.0 [37]. In silico PVD peptide backbone structural predictions were conducted using the methods of Rausch et al. and Stachelhaus et al.[36,37] implemented in NRPSpredictor [47]. Putative siderophore receptor genes were identified by hits to protein family (PFam) accessions PF00593 (TonB-dependent receptor) and PF07715 (TonB-dependent receptor plug domain) [22], in combination with BLAST homology to putative siderophore receptor genes. Putative sigma factor and anti-sigma factor genes adjacent to receptor genes were identified by hits to PFam accessions PF04542 (Sigma-70 region) and PF04773 (FecR protein), respectively.

Gene reporter and expression constructs
DNA manipulation was performed by standard techniques [38]. Promoter reporter fusion strains were generated by cloning the promoter region in the front of a promoterless ‘lacZ carried in the cloning vector pIVET-Sm, a Sm² derivative of pIVETD [19]. Promoter regions were created by PCR amplifying regions upstream of, and including the 5’-portion of PVD genes using primers listed in Table 3. To facilitate directional cloning, BglIII and XhoI sites were incorporated into the 5’-ends of the sense and anti-sense strand primers, respectively. PCR products were digested with BglIII and Xhol, and ligated to BglIII-Xholl digested pIVETD-Sm, creating fusions to promoterless ‘lacZ. Reporter strains were created by transforming SBW25 with the promoter fusion constructs via electroporation or conjugation. Transformants were selected based on streptomycin resistance and PCR screened using a primer upstream of the cloned promoter fragment and a ‘lacZ primer to confirm the construct had integrated into the genome by single-crossover homologous recombination within the cloned PVD promoter fragment region.

Gene expression constructs for pvdS, fpvI and fpvR were created using the Gateway cloning system (Invitrogen). Fragments containing the ribosome-binding site and entire ORF of pvdS, fpvI and fpvR were PCR-amplified using primers in Table 3. Fragments were directionally cloned into pENTR/D-TOPO (Invitrogen) following the manufacturers instructions. The sequences of the cloned fragments were verified, and LR recombination reactions using Gateway LR Clonase enzyme mix (Invitrogen) were performed to transfer the cloned fragment to the broad host range destination expression vector, pBroadgate (gift from R. Thwaites and J. Mansfield). pBroadgate is a derivative of pBBR1MCS-5 [48] that has been modified as a Gateway destination vector by incorporation of an attR cassette, containing a Cm resistance determinant and a
PCR using primers pvdL1 and pvdL4 respectively (Table 3). The two fragments were ligated in a third phase after the sequence of the cloned fragment was verified, the fragment was retrieved by digestion and cloned into the vector pUIC3 to create pUIC3–40.

β-galactosidase assays
β-galactosidase activities of lacZ reporter fusion strains were determined from cultures grown to mid- to late log phase in CAA media, supplemented with FeSO₄ or 2,2'-dipyridyl as required. Assays were based on the hydrolysis of 4-methylumbelliferyl-β-D-galactoside to yield the fluorescent product, 7-hydroxy-4-methylcoumarin (4MU) as described [49]. 4MU was detected at 460 nm after excitation at 365 nm using a Hoefer DyNA Quant 200 fluorometer (Pharmacia Biotech, San Francisco CA, USA), or FLI/Qstar (BMG Labtech, Offenburg, Germany).

Construction of PBR840 (ΔpvdL)
An ORF Pflu4387 (pvdL) knockout mutant was constructed by deleting a 10.9 kb internal portion of the 12.9 kb pvdL gene. This was achieved by a combination of gene splicing by overlap extension (SOE) PCR [50] and a two-step allelic exchange strategy [20]. Briefly, two 1 kb fragments from each of the 5’- and 3’-end of pvdL were PCR-amplified from SBW25 genomic DNA using the primer pairs pvdL1 and pvdL2, and pvdL3 and pvdL4, respectively (Table 3). The two fragments were ligated in a third PCR using primers pvdL1 and pvdL4 via the complementary residues incorporated at the 5’-ends of primers pvdL2 and pvdL3 (Table 3). The resulting 2 kb pvdL deletion fragment was cloned into pCR8/GW/TOPO (Invitrogen). After the sequence of the cloned fragment was verified, the lethal ccdB gene (Invitrogen). P. fluorescens reporter strains were transformed with expression and pBroadgate-D control constructs by electroporation.

PVD structure determination
To determine the structure of SBW25 PVD, mass spectral data were obtained with a MAT 900 ST instrument provided by Finnigan MAT, Bremen, Germany); spray voltage 3.4–3.6 kV, capillary temperature 230°C. Source conditions were set to minimize fragmentation, resolution ca. 5000 (10% valley). Trifluoroacetic acid was added to the sample dissolved in water (100:0.2, v/v). Fragmentation induced by low energy collision activation was effected in the octapole, equipped with an ESI II ion source (Finnigan MAT, Bremen, Germany); spray voltage 3.4–3.6 kV, capillary temperature 230°C. Source conditions were set to minimize fragmentation, resolution ca. 5000 (10% valley). Trifluoroacetic acid was added to the sample dissolved in water (100:0.2, v/v). Fragmentation induced by low energy collision activation was effected in the octapole unit, located in front of the QIT and in the QIT itself (~2·10⁻³ Pa He as bath gas diffusing in the collision octapole).
**PVD purification and exogenous siderophore uptake by SBW25**

PVD was purified from bacterial cultures grown for 48 h in 1 l of CAA at 28°C. For *P. aeruginosa*, the strains were grown at 37°C. The cell-free supernatant was passed on a C-18 column (5 × 2.5 cm) conditioned with methanol and rinsed with sterile H2O. PVD was eluted with 80% methanol. Subsequently the methanol was evaporated and the PVD lyophilized. Purified PVD was quantified spectrophotometrically [39].

To estimate the diversity of exogenous PVDs that may be utilized by SBW25, PVDs were purified from a panel of 25 *Pseudomonas* isolates and strains, representing a high diversity of species that had previously been shown to each produce a unique PVD (each having a unique siderotype) as shown by isoelectric focusing (S. Matthijs, unpublished). These isolates included *Pseudomonas* type strains, and environmental isolates collected from the Woluwe river, Brussels, Belgium (Table 2). CAA agar plates containing 0.5 mg/ml EDDHA were overlaid with 5 × 10⁶ cells of strain 15F3, a siderophore-deficient derivative of PBR840 that contains transposon mini-Tn phosphoA3 disruption of biosynthetic genes for a secondary siderophore (S. Matthijs, unpublished). Filter-paper disks impregnated with 5 µl of 8 mM purified siderophore were placed on the agar [39]. Plates were incubated at 28°C and scored for the presence of detectable growth after one day.

**List of abbreviations used**

CAA, casamino acids medium; EDDHA, ethylenediaminedihydroxyphenylacetic acid; IVET, *in vivo* expression technology; LB, Luria-Bertani; NRPS, non-ribosomal peptide synthetase; PCR, polymerase chain reaction; PVD, pyoverdine; QIT, quadrupole ion trap; SOE, splicing by overlap extension; X-gal, 5-bromo-4-chloro-3-indolyl β-D-galactopyranoside; 4MU, 7-hydroxy-4-methylcoumarin.

**Authors’ contributions**

CDM conducted the bioinformatic and genetic work, and drafted the manuscript. XXZ created the ΔpvdL mutant, assessed growth characteristics, and contributed to the manuscript. SM performed the assay for the heterologous usage of PVDs. MS and HB determined the structure of SBW25 PVD. PBR assisted with experimental design and critically reviewed the manuscript. All authors have read and approved the final manuscript.

**Additional material**

**Additional file 1**

List of *P. fluorescens* SBW25 genes with predicted function in PVD biosynthesis and uptake in comparison with homologues in *P. aeruginosa* PA01. The data list all putative PVD genes identified in SBW25, and show predicted amino acid identities to their PA01 homologues.

Click here for file [http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S1.doc](http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S1.doc)

**Additional file 2**

NRPS substrate in silico predictions. The data show the substrates of the PVD NRPS modules predicted by specificity conferring code and transduction support vector machine methods.

Click here for file [http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S2.doc](http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S2.doc)

**Additional file 3**

List of putative siderophore receptors in *P. fluorescens* SBW25. The data list all putative siderophore receptors in the SBW25 genome, and whether these are adjacent to predicted sigma factor/anti-sigma factor gene pairs.

Click here for file [http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S3.doc](http://www.biomedcentral.com/content supplementary/1471-2180-8-7-S3.doc)

**Acknowledgements**

We are grateful to I.L. Lamont and P. Cornelis for helpful discussions. Many thanks also to R.W. Jackson for statistical analyses and helpful discussion, Z.A. Park-Ng for statistical analyses, S.R. Giddens and C.G. Knight for bioinformatics support, R. Thwaites and J. Mansfield for the kind gift of pBroadgate, and J. Bull for technical support. This research was supported by a grant from the Biotechnology and Biological Sciences Research Council, U.K. to P.B.R.

**References**

1. Neillands JB: Siderophores: structure and function of microbial iron transport compounds. *J Biol Chem* 1995, 270(45):26723-26726.
2. Crosa JH: Signal transduction and transcriptional and post-transcriptional control of iron-regulated genes in bacteria. *Microbiol Mol Biol Rev* 1997, 61(3):319-336.
3. Cornelis P, Matthijs S: Diversity of siderophore-mediated iron uptake systems in fluorescent pseudomonads: not only pyoverdines. *Environ Microbiol* 2002, 4(12):787-798.
4. Meyer JH: Pyoverdines: pigments, siderophores and potential taxonomic markers of fluorescent Pseudomonas species. *Arch Microbiol* 2000, 174(1):135-142.
5. Beare PA, For RJ, Martin LW, Lamont IL: Siderophore-mediated cell signalling in *Pseudomonas aeruginosa*: divergent pathways regulate virulence factor production and siderophore receptor synthesis. *Mol Microbiol* 2003, 47(1):195-207.
6. Vasil ML, Ochsner UA: The response of *Pseudomonas aeruginosa* to iron: genetics, biochemistry and virulence. *Mol Microbiol* 1999, 34(3):399-413.
7. Lamont IL, Beare PA, Ochsner U, Vasil AI, Vasil ML: Siderophore-mediated signaling regulates virulence factor production in *Pseudomonas aeruginosa*. *Proc Natl Acad Sci USA* 2002, 99(10):7072-7077.
8. Redly GA, Poole K: FpvIR control of fpvA ferric pyoverdine receptor gene expression in Pseudomonas aeruginosa: demon-
stration of an interaction between FpvL and FpvR and identifica-
tion of mutants in each compromising this inter-
action. J Bacteriol 2005, 187(16):5648-5657.
9. Lamont IL, Martin LW, Sims T, Scott A, Wallace M: Characteriza-
tion of a gene encoding an acetylase required for pyoverdine synthesis in Pseudomonas aeruginosa. J Bacteriol 2006, 188(8):3149-3152.
10. Prince RW, Cox CD, Vasil ML: Coordinate regulation of siderophore and exotoxin A production: molecular cloning and sequencing of the Pseudomonas aeruginosa fur gene. J Bacteriol 1993, 175(2):258-259.
11. Ochsner UA, Vasil ML: Gene repression by the ferric uptake regulator in Pseudomonas aeruginosa: cycle selection of iron-regulated genes. Proc Natl Acad Sci USA 1996, 93(9):4409-4414.
12. Cunliffe HE, Merriman TR, Lamont IL: Analysis of promoters rec-
ognized by PvdS, an extracytoplasmic-function sigma factor protein from Pseudomonas aeruginosa. J Bacteriol 2001, 183(6):2151-2155.
13. Cunliffe HE, Merriman TR, Lamont IL: Cloning and characteriza-
tion of a PvdS gene required for pyoverdine synthesis in Pseudomonas aeruginosa: PvdS is probably an alternative sigma factor. J Bacteriol 1995, 177(10):2744-2750.
14. Redly GA, Poole K: Pyoverdine-mediated regulation of FpvA synthesis in Pseudomonas aeruginosa: involvement of a preglobular extracytoplasmic-function sigma factor, FpvL. J Bacteriol 2003, 185(4):1261-1265.
15. Loper JE, Henkel MD: Utilization of heterologous siderophores enhances levels of iron available to Pse-
domonas putida in the rhizosphere. Appl Environ Microbiol 1999, 65(12):3537-3541.
16. Molina MA, Godoy P, Ramos-Gonzalez MI, Munoz N, Ramos JL, Espinosa-Urgel M: Role of iron and the TonB system in coloni-
zation of corn seeds and roots by Pseudomonas putida KT2440. Environ Microbiol 2005, 7(3):443-449.
17. Bailey MJ, Ulley AK, Thompson IP, Rainey PB, Ellis RJ: Site-directed chromosomal marking of a fluorescent pseudomonad iso-
lated from the phytosphere of sugar beet; stability and potential for marker gene transfer. Mol Ecol 1995, 4(6):579-583.
18. Ellis RJ, Timms-Wilson TM, Bailey MJ: Identification of conserved traits in fluorescent pseudomonads with antifungal activity. Environ Microbiol 2000, 2(3):274-284.
19. Gal M, Preston GM, Massey RC, Spiers AJ, Rainey PB: Genes encod-
ing a cellulosic polyester contribute toward the ecological success of Pseudomonas fluorescens SBW25 on plant sur-
faces. Mol Ecol 2003, 12(11):3109-3121.
20. Rainey PB: Adaptation of Pseudomonas fluorescens to the plant rhizosphere. Environ Microbiol 1999, 1(3):243-257.
21. Mossialos D, Ochiaver LJ, Yavrova C, Chablan P, Fournier J, Koedam N, Budzikiewicz H, Fernandez DU, Schafer M, Ravel J, Cornelis P: Identifi-
cation of new, conserved, non-ribosomal peptide synthesis-
tases from fluorescent pseudomonads involved in the biosynthesis of the siderophore pyoverdine. Mol Microbiol 2002, 44(6):1673-1685.
22. http://www.sanger.ac.uk/Projects/P.fluorescens/ [http://www.sanger.ac.uk/Projects/P_fluorescens/]
23. Ravel J, Cornelis P: Genomics of pyoverdine-mediated iron uptake in pseudomonads. Trends Microbiol 2003, 11(5):195-200.
24. Smith EE, Sima EP, Spencer DC, Kaur R, Olson MV: Evidence for diversifying selection at the pyoverdine locus of Pse-
domonas aeruginosa. J Bacteriol 2005, 187(6):2138-2147.
25. Paulsen IT, Press CM, Ravel J, Kobayashi DT, Myers GSA, Mavrilo-
 di DV, DeBoy RT, Seshadri R, Ren Q, Maduup R, Dodson RJ, Durkin AS, Brinkak LM, Daugherty SC, Sullivan SA, Rosovitz MJ, Gwinn ML, Zhou L, Schneider DJ, Cattinhow SW, Nelson WC, Weidman J, Watkins K, Tran K, Khouri H, Pierson EA, Pierson LS, Thomashow LS, Loper JE: Complete genome sequence of the plant commensal Pseudomonas fluorescens PS-5. Nature Biotechnology 2005, 23(7):873-878.
26. Koster M, van Klompenburg W, Bitter W, Leong J, Weisbeck P: Role for the outer membrane ferric siderophore receptor PupB in signal transduction across the bacterial cell envelope. EMBO J 1994, 13(12):2805-2813.
27. Visca P, Leoni L, Wilson MJ, Lamont IL: Iron transport and regu-
lation, cell signalling and genomics: lessons from Escherichia coli and Pseudomonas. Mol Microbiol 2002, 45(5):1771-190.
28. Fuchs R, Budzikiewicz H: Structural studies of pyoverdines by mass spectrometry. Curr Org Chem 2001, 5:265-288.
29. Budzikiewicz H, Schafer M, Fernandez DU, Mathijs S, Cornelis P: Characterization of the chromophores of pyoverdines and related siderophores by electrospray tandem mass spec-
26trometry. BioMetals 2007, 20(2):135-144.
30. Budzikiewicz H, Schafer M, Meyer JM: Siderotyping of fluorescent pseudomonads - problems in the determination of molecular masses by mass spectrometry. Mini-Rev Org Chem 2007, 4(6):579-583.
31. Fuchs R, Budzikiewicz H: Structural studies of pyoverdines with cyclopeticidic substrucutures by electrospray ionization and collision induced fragmentation. Spectroscopy 2000, 14:229-246.
32. Hohneicher U, Hartmann R, Taraz K, Budzikiewicz H: Pyoverdine, ferrioxamine, azothobactin - a wide trade of siderophores from Pseudomonas chlororaphis ATCC 9446 and its relation to Pseudomonas fluorescens ATCC 13525. Z Naturforsch 1995, 50:337-344.
33. Budzikiewicz H: Siderophores of the Pseudomonadaceae sensu stricto (fluorescent and non-fluorescent Pseudomonas spp.). J Bacteriol 2004, 187:317-3140.
34. Linget C, Azadi P, MacLeod JK, Dell A, Abdalla M: Bacterial siderophores: The structures of the pyoverdins of Pse-
domonas fluorescens ATCC 13525. Tetrahedron Lett 1992, 33(1):73-76.
35. Abdalla MA: Pyoverdines and pseudobactins. In CRC handbook of microbial iron chelates Edited by: Winkelmans B. Boca Raton, Florida , CRC Press.; 1991:139-152.
36. Rausch C, Weber T, Kohlbacher O, Wohlleben W, Hsuan DH: Spe-
cificity prediction of adenyl ation domains in nonribosomal peptide synthetases (NRPS) using transductive support vec-
tor machines (TSVMs). Nucleic Acids Res 2005, 33(18):3599-3608.
37. Schachteleus T, Mooz HD, Mariash ML: The specificity-confer-
ing code of adenylation domains in nonribosomal peptide synthetases. Chem Biol 1999, 6(8):493-505.
38. Sambrook J, Fritsch EF, Maniatis T: Molecular Cloning: A Labora-
toary Manual. 2nd edition. Plainview, , Cold Spring Harbor Labora-
tory Press.; 1989.
39. Meyer JM, Stintzi A, De Vos D, Cornelis P, Tappe R, Taraz K, Budzik-
iewicz H: Use of siderophores to type pseudomonads: the three Pseudomonas aeruginosa pyoverdine systems. Microbi-
ology 1997, 143:353-35.
40. Stover CK, Pham XQ, Erwin AL, Mizoguchi SD, Warrender P, Hickey M, Brinkman FS, Hufnagle WO, Kowalik D, Lagrou M, Garber RL, Golyt L, Tolentino E, Westbrook-Wadman S, Yuen Y, Brody LL, Coulter SN, Folger KR, Kas A, Larbig K, Lim R, Smith K, Spencer D, Dong WG, Wu Z, Paulsen IT, Reizer J, Saier PH, Hancock RE, Lory S, Olson MV: Complete genome sequence of Pseudomonas aeruginosa PA01, an opportunistic pathogen. Nature 2000, 406(6799):959-964.
41. Lamont IL, Martin LW: Identification and characterization of novel pyoverdine synthesis genes in Pseudomonas aerugi-

nosa. Microbiology 2003, 149:831-842.
42. Thijs G, Lesot C, Marchal K, Rombouts S, De Moor B, Rouzé P, Moreau Y: A higher order background model improves the detection of regulatory elements by Gibbs Sampling. Bioinfor-
matics 2001, 17(12):113-122.
43. http://prodoric.tu-bs.de [http://prodoric.tu-bs.de]
44. Munch R, Hiller K, Barge H, Heldt D, Lenz S, Wingender E, Jahn D: PRODORIC: prokaryotic database of gene regulation. Nucleic Acids Res 2003, 31(1):266-269.
45. http://www.ab.informatic.uni-tuebingen.de/software [http://www.ab.informatic.uni-tuebingen.de/software]
range cloning vector pBRR1MCS, carrying different antibiotic-resistance cassettes. Gene 1995, 166(1):175-176.

49. Pharmacia B: Hoefer DyNA Quant 200 Fluorometer User Manual.

50. Horton RM, Hunt HD, Ho SN, Pullen JK, Pease LR: Engineering hybrid genes without the use of restriction enzymes: gene splicing by overlap extension. Gene 1989, 77:61-68.

51. Prendki P, Krisch HM: In vitro insertional mutagenesis with a selectable DNA fragment. Gene 1984, 29(3):303-313.

52. Figurski DH, Helinski DR: Replication of an origin-containing derivative of plasmid RK2 dependent on a plasmid function provided in trans. Proc Natl Acad Sci USA 1979, 76(4):1648-1652.

53. Giddens SR, Jackson RW, Moon CD, Jacobs MA, Zhang XX, Gehring SM, Rainey PB: Mutational activation of niche-specific genes provides insight into regulatory networks and bacterial function in a complex environment. Proc Natl Acad Sci USA 2007, 104(46):18247-18252.

54. Holloway BW: Genetic recombination in Pseudomonas aeruginosa. J Gen Microbiol 1955, 13(3):572-581.

55. Hofte M, Mergeay M, Verstraete W: Marking the rhizopseudomonas strain 7NSK2 with a Mu d(lac) element for ecological studies. Appl Environ Microbiol 1990, 56(4):1046-1052.

56. Matthijs S, Bayse C, Koedam N, Tehrani KA, Verheyden L, Budzikiewicz H, Schaefer M, Hoorelbeke B, Meyer JM, De Greve H, Cornelis P: The Pseudomonas siderophore quinolobactin is synthesized from xanthurenic acid, an intermediate of the kynurenine pathway. Mol Microbiol 2004, 52(2):371-384.