Discovery of an Antibiotic-Related Small Protein of Biocontrol Strain Pseudomonas sp. Os17 by a Genome-Mining Strategy

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Many root-colonizing Pseudomonas spp. exhibiting biocontrol activities produce a wide range of secondary metabolites that exert antibiotic effects against other microbes, nematodes, and insects in the rhizosphere. The expression of these secondary metabolites depends on the Gac/Rsm signal transduction pathway. Based on the findings of a previous genomic study on newly isolated biocontrol pseudomonad strains, we herein investigated the novel gene cluster OS3, which consists of four genes (Os1348–Os1351) that are located upstream of putative efflux transporter genes (Os1352–Os1355). Os1348 was predicted to encode an 85-aa small precursor protein, the expression of which was under the control of GacA, and an X-ray structural analysis suggested that the Os1348 protein formed a dimer. The mutational loss of the Os1348 gene decreased the antibiotic activity of Pseudomonas sp. Os17 without changing its growth rate. The Os1349–1351 genes were predicted to be involved in post-translational modifications. Intracellular levels of the Os1348 protein in the deficient mutant of each gene differed from that in wild-type cells. These results suggest that Os1348 is involved in antibiotic activity and that the structure or expression of this protein is under the control of downstream gene products.

Keywords: comparative genomics, root-colonizing pseudomonads, bacteriocin-like gene cluster, post-translational modification, Gac/Rsm signal transduction pathway, X-ray structural analysis

INTRODUCTION

Root-colonizing fluorescent pseudomonads are known to be plant-beneficial strains that suppress the growth of other microbes, including plant pathogens, nematodes, and insects. Strains in this group have the potential to produce various secondary metabolites and enzymes with antibiotic activities, which enable them to compete with other microbes, thereby facilitating niche adaptation in the rhizosphere. Pseudomonas protegens strains CHA0 and Pf-5 have been used as model strains in research on the biosynthesis and regulation of secondary metabolites by these pseudomonads. They produce 2,4-diacetylphloroglucinol (DAPG), pyrrolnitrin (Prn), pyoluteorin (Plt), hydrogen...
cyanide (HCN), and exoenzymes (e.g., AprA protease) as their typical exoproducts (Haas and Keel, 2003; Loper et al., 2012). Moreover, since the complete genomic sequence of strain Pf-5 was elucidated (Paulsen et al., 2005), it has provided novel insights into the molecular mechanisms underlying the pseudomonad-mediated suppression of plant diseases. Genomic-guided approaches have facilitated the identification of novel natural products, such as the cyclic lipopeptide orfamide A (Gross et al., 2007), rhizoxin analogs (Loper et al., 2008), and the insect toxin FitD (Péchy-Tarr et al., 2008). Although the suppressive activities of these metabolites against plant pathogenic fungi and oomycetes have been extensively examined, limited information is currently available on their antibacterial exoproducts.

This wide range of secondary metabolites is considered to be a rich source of valuable compounds for industrial applications. We recently screened 48 DAPG-producing strains among 2,800 fluorescent pseudomonads isolated from fields in Japan to identify novel factors involved in the antibiotic activity of newly isolated strain(s). Of these, we fully sequenced three strains, Cab57, Os17, and St29. Cab57 was identified as P. protogens based on a 16S rRNA gene analysis (with 100% identity) and whole-genome analysis (Takeuchi et al., 2014), whereas the two other strains were the closest to, but different from P. protogens (Takeuchi et al., 2015). We showed that Os17 and St29 exhibited different biocontrol potentials against Pythium damping-off and root rot in the cucumber; Os17 was as effective as Cab57, whereas St29 was less effective. A whole-genome comparison of these related strains revealed that Os17 and St29 are the same species and have several strain-specific genomic regions. Among these regions, the complete rhizoxin analog biosynthesis gene cluster (ca. 79 kb) was found to be specific to the Os17 genome (Takeuchi et al., 2015). In the rxxB gene of polyketide synthase, essential for the production of rhizoxin analogs, mutation, growth inhibitory activity against fungi and oomycetes as well as plant protection efficacy were weaker than those of wild-type Os17. Therefore, although we successfully identified anti-oomycete metabolites, it currently remains unclear whether these strains possess antibacterial exoproducts. DAPG-producing pseudomonads are regarded as a source of bacteriocin, as demonstrated in a previous study on strain Pf-5 in which the novel lectin-like bacteriocin LlpA was discovered (Parret et al., 2005).

We also demonstrated that the function of the well-known Gac/Rsm signal transduction pathway (Lapouge et al., 2008; Kidarsa et al., 2013; Valentini and Filloux, 2016) was conserved in Os17, in which the production of rhizoxin analogs was under the control of GacA (Takeuchi et al., 2015). In Pseudomonas spp., the GacS sensor kinase is phosphorylated and initiates this pathway, which enables the activation of the cognate GacA response regulator by phosphotransfer, leading to the transcription of non-coding small RNAs (sRNAs) (Kay et al., 2005, 2006; Lalaouna et al., 2012). GacA positively controls the expression of three sRNAs (RsmX, RsmY, and RsmZ) that have high affinity to the RNA-binding repressor proteins RsmA and RsmE, which induces the expression of mRNAs encoding the biosynthetic gene cluster of secondary metabolites (Heeb et al., 2002; Valverde et al., 2003; Kay et al., 2005; Reimmann et al., 2005; Song et al., 2015). The RsmA and RsmE proteins repress the translation of genes by sequestering a conserved ANGGAN motif in or near to the Shine–Dalgarno sequence and this repression typically occurs during rapid growth (Reimmann et al., 2005; Lapouge et al., 2008). When RsmX/Y/Z sRNAs are induced during restricted growth at a high cell population density, they relieve the translational repression of target genes by sequestering the RsmA and RsmE proteins. Therefore, the biofilm state and secondary metabolism are favored (Lapouge et al., 2008; Valentini and Filloux, 2016). In strain Os17, the gacA mutant exhibited the down-regulated expression of rxxB and rxxI, both of which have the consensus sequence ANGGAN in their upstream region (Takeuchi et al., 2015). In addition to the wide range of genomic repertoires available for the synthesis of the secondary metabolites described above, fine-tuning of the Gac/Rsm regulatory system is considered to contribute to the versatility of P. protogens and related strains for various environmental niches. Previous studies demonstrated that the ability of mutants defective in the Gac/Rsm system to synthesize biocontrol factors, suppress plant diseases, and produce insecticides was reduced (Haas and Défago, 2005; Kupferschmied et al., 2013).

Comparative genomics among closely related strains revealed phenotype-specific genes, thereby enabling us to identify strain-specific biocontrol factor(s). In addition to these strain-specific gene clusters, clusters commonly shared by the genomes of Os17 and St29, but absent from the genomes of P. protogens, are also of interest because they may contribute to establishing the uniqueness of this putative novel species. Comparative genomics among the three strains revealed that in addition to 5,321 core genes, 519 coding sequences (CDSs) were specifically shared by the genomes of Os17 and St29, and a number of these CDSs (i.e., 281 out of 519 CDSs) formed gene clusters (Takeuchi et al., 2015). Some clusters were predicted to be involved in the production of secondary metabolites, the function of which remains unknown. In the present study, we focused on one of these gene clusters, Cluster OS3 (consisting of four successive genes Os1348–1351 in the genome of Os17), which was predicted to be a bacteriocin gene cluster, and we constructed a deficient mutant of each gene. We found that the deficient mutant of Os1348, which encodes an 85-aa small protein, exhibited weaker antibacterial activity than wild-type Os17. We also demonstrated that the downstream genes of Os1348 affected the protein status of Os1348.

MATERIALS AND METHODS

Bacterial Strains and Growth Conditions

The bacterial strains and plasmids used in the present study are listed in Supplementary Table 1. Pseudomonas sp. Os17 has been deposited in the NARO Genebank as MAFF212089 (Takeuchi et al., 2015). Escherichia coli and Pseudomonas strains were routinely grown in NYB [2.5% (w/v) nutrient broth, 0.5% (w/v) yeast extract] and Luria-Bertani (LB) medium with shaking, or on nutrient agar plates [4% (w/v) blood agar base, 0.5% (w/v) yeast extract] amended with the following antibiotics when required: ampicillin, 100 µg/ml; kanamycin, 25 µg/ml; or...
tetracycline, 50 μg/ml (100 μg/ml for selection of Pseudomonas sp.). Incubation temperatures were 28°C for Pseudomonas sp. and 37°C for E. coli.

In antibiotic assays, bacteria were grown in modified glycerol-casamino acid medium (GCM; Maurhofer et al., 1998) without the amendment of minerals.

**DNA Manipulation**
Small-scale plasmid extraction was performed with a QIAprep spin miniprep kit (QIAGEN); large-scale preparations were obtained with a Qiagen plasmid midi kit. Chromosomal DNA from Pseudomonas sp. Os17 was prepared using Qiaagen genomic tips. DNA fragments were purified from agarose gels with a QIAquick gel extraction kit (QIAGEN). The oligonucleotides used are listed in Supplementary Table 2.

**Generation of Mutants of Os1348-Related Genes and the phlD Gene**
The primers used are listed in Supplementary Table 2. An in-frame deletion in chromosomal Os1348-related genes and the phlD gene of Pseudomonas sp. Os17 was created as follows. Fragments of ca. 750–800-bp regions flanking 1348, 1349, 1350, 1351, 1353 and the phlD gene were amplified by PCR with the primer pairs 1348UF/1348UR and 1348DF/1348DR for 1348, 1349UF/1349UR and 1349DF/1349DR for 1349, 1350UF/1350UR and 1350DF/1350DR for 1350, 1351UF/1351UR and 1351DF/1351DR for 1351, 1353UF/1353UR and 1353DF/1353DR for 1353, or PhlDUF/PhlDUR and PhlDDF/PhlDDR for phlD. High-fidelity DNA polymerase KOD Plus™ (Toyobo) and the genomic DNA of Pseudomonas sp. Os17 as a template were used for amplification.

Each of the two corresponding fragments were annealed and amplified as a 1.5- or 1.6-kb fragment using the primer pair 1348UF/1348DR, 1349UF/1349DR, 1350UF/1350DR, 1351UF/1351DR, 1353UF/1353DR, or PhlDUF/PhlDDR. These 1.5- and 1.6-kb fragments were cloned into pCR-Blunt II-TOPO™ (Thermo Fisher Scientific). The inserts obtained were confirmed by sequencing. After sequencing, these 1.5- and 1.6-kb fragments were cloned into pME3087 cut with KpnI and HindIII to give pME3087-1348D, pME3087-1349, pME3087-1350, pME3087-1351, pME3087-1353, and pME3087-phlD, and pME3087 cut with KpnI and HindIII to give pME3087-1349. These plasmids were mobilized from E. coli DH5α to Pseudomonas sp. Os17 by triparental mating with E. coli HB101/pME497. Excision of the vector via second crossing-over was obtained after enrichment for tetracycline-sensitive cells, generating the 1348 complement mutant. Complementation of the 1348 gene into the chromosome was confirmed by PCR with the primers 1349UF and 1349DR (Supplementary Table 2).

The 1348 mutant was also complemented with the plasmid pME6031-OS3, which contained a 4.4-kb fragment carrying the 1348-1351 genes (Cluster OS3). This fragment was amplified with the primers OS3F and OS3R, high-fidelity DNA polymerase KOD Plus™ (Toyobo), and the genomic DNA of Pseudomonas sp. Os17 as a template. The 4.4-kb fragment was cloned into pPCR-Blunt II-TOPO™ (Thermo Fisher Scientific). The insert obtained was confirmed by sequencing. After sequencing, the fragment was cloned into pME6031 cut with HindIII and KpnI. The resultant plasmid, pME6031-OS3, was introduced into the 1348 mutant by electroporation.

**Expression and Protein Purification of the Recombinant Os1348 of Pseudomonas sp. Os17 by E. coli**
The fragment carrying 1348 was amplified from the genomic DNA of Pseudomonas sp. Os17 by PCR with the primers 25F1348 and 25R1348 (Supplementary Table S2). The PCR product was inserted into the Ndel/BamHI sites of the expression vector pET25b (+) (Novagen) with the In-Fusion HD Cloning system (Clontech). The resultant plasmid, pET25-1348, was introduced into E. coli BL21 (DE3) (Merck) and the strain was grown at 37°C. The expression of the recombinant Os1348 protein was induced by the addition of isopropyl-β-D-thiogalactopyranoside at a final concentration of 0.4 mM and growth continued at 16°C for 16 h. Harvested cells were resuspended in PBS (pH 7.4) and lysed by sonication. Debris was removed by centrifugation at 40,000 × g for 30 min. The lysate was applied to a 5-ml HiTrap SP HP column (GE Healthcare). The column was washed with 20 mM sodium acetate (pH 5.2) and eluted with a linear gradient of 0–500 mM NaCl. Os1348 fractions were purified further on HiLoad 26/60 Superdex75 prep grade (GE Healthcare) pre-equilibrated with buffer consisting of 10 mM Tris–HCl (pH 7.5) and 150 mM NaCl. After dialysis at 4°C for 16 h against 20 mM sodium acetate (pH 5.2), the supernatant was purified by a Mono S 10/100 GL column (GE Healthcare) and a 0–300 mM NaCl elution gradient.

**X-ray Structural Elucidation of Os1348**
The Os1348 protein was concentrated to 2.0 mg ml⁻¹ and crystallized by the sitting-drop vapor diffusion method at 293 K using a precipitant solution consisting of 30% (w/v) polyethylene glycol monomethyl ester 2000 and 0.2 M MgCl₂. Plate-type crystals with dimensions of 0.2 × 0.2 × 0.05 mm appeared...
within 2 weeks using 50 µl of the reservoir solution with a drop consisting of 0.5 µl of the protein solution and 0.5 µl of the reservoir solution.

Diffraction experiments for Os1348 crystals were conducted at the beamline BL-5A of Photon Factory (PF), High Energy Accelerator Research Organization, Tsukuba, Japan. Diffraction data were collected using the ADSC Q315 CCD detector (Area Detector Systems Corp., Poway, CA, United States). Crystals were mounted in a quartz glass capillary with a diameter of 0.3 mm (Capillary Tube Supplies Ltd., United Kingdom) and cryocooled in a nitrogen gas stream to 95 K. Crystals diffraeted to 1.8 Å resolution. Data were integrated and scaled using the programs DENGZ and SCALEPACK in the HKL2000 program suite (Otwinowski and Minor, 1997).

The crystal structure was elucidated by a direct method using the program AMPLEx (Bibby et al., 2012) incorporated in the CCP4 program suite (Winn et al., 2011). Manual model building and molecular refinement were performed using Coot (Emsley and Cowtan, 2004) and Refmac5 (Murshudov et al., 2011). Data collection and refinement statistics are shown in Table 4. Model stereochemistry was assessed with the program Molprobity (Chen et al., 2010). Structural illustrations were drawn with the program PyMOL (DeLano, 2002).

Detection of the Os1348 Protein
The presence of the Os1348 protein was examined in cells using western blotting with rabbit polyclonal antibodies raised against the recombinant Os1348 protein. The anti-Os1348 antibody was produced by Biogate (Gifu, Japan) according to their established protocol. Equal amounts of cells corresponding to 20 µl of a suspension with an OD_{600} nm of 4.0 were collected. The pellet was solubilized in buffer using 4× Laemmli Sample Buffer (Bio-Rad) and boiled at 100°C for 5 min in a heating block. Samples were subjected to Tricine-SDS-PAGE (15% polyacrylamide gel) and subsequently analyzed by western blotting with the anti-Os1348 antibody.

β-Galactosidase Assays
β-Galactosidase activity was quantified by the Miller method (Miller, 1972). Pseudomonas strains were grown at 28°C in 50-ml flasks containing 15 ml of NYB amended with 0.05% Triton X-100 with shaking at 180 rpm. Triton X-100 was required to prevent cell aggregation.

Detection of Antibiotic Activity
The antibiotic activities of Pseudomonas Os17 and the mutant strains were assessed with Bacillus subtilis M168 (DSM 402, NBRC 111470) as the reporter. We also tested other Pseudomonas strains (Pseudomonas syringae pv. pisi 730032 and tabaci 301612, Pseudomonas chlororaphis St508, and P. protegens Cab57) as the reporter to test susceptibility against Pseudomonas Os17. P. syringae, oomycete and fungal strains were obtained from the NARO Genebank in Tsukuba. In the assays with B. subtilis and other bacterial reporter strains, cultures of Os17 and the mutant strains were adjusted to OD_{600} nm = 1.0 and 4-µl samples were spotted onto the modified GCM plate. After an overnight incubation at 28°C, cells were killed by UV irradiation on a transilluminator for 5 min. A soft-agar overlay for plates was prepared with 4-ml half-strength NA [2% (v/v) blood agar base, 0.25% (w/v) yeast extract] inoculated with 400-µl cultures of the reporter strains grown in NYB until the stationary phase. The overlay of B. subtilis revealed antibiotic production by growth inhibition zones. Halo sizes were measured from the edge of the bacterial spot to the end of the clear zone, and the data obtained are shown as the average of 3–6 replicates. In the assay with Pythium ultimum or Fusarium oxysporum, cultures of Pseudomonas strains were adjusted to OD_{600} nm = 5.0 and 20-µl samples were streaked around the edge of the PDA plate, and, after the overnight incubation at 28°C, an inoculum of P. ultimum or F. oxysporum was transferred to the center of the plate. The plate was incubated at room temperature until P. ultimum or F. oxysporum reached its edge.

In the evaluation of DAPG production, strains were grown in modified GCM for 22 h after the inoculation (scaling up from an overnight culture to a fresh culture, 50 times, and grown in Erlenmeyer flasks at 180 rpm, 28°C), corresponding to an OD_{600} nm of ca. 0.5. The supernatants of strains were extracted with an equal volume of ethyl acetate and analyzed by HPLC for DAPG as described previously (Schnider-Keel et al., 2000).

RESULTS
Genomic Structure of the Putative Bacteriocin Gene Cluster in Pseudomonas sp. Strains Os17 and St29
We previously identified 33 gene clusters that are shared by the genomes of Os17 and St29 and absent from the P. protegens Cab57 genome (Clusters OS1 to OS33 in Supplementary Table 4 of Takeuchi et al., 2015). Among these clusters, we focused on Cluster OS3, which was predicted to encode the bacteriocin gene cluster according to the antiSMASH program (Blin et al., 2013). In this program, the downstream genes (gene IDs were given as Os1352 and Os1353, predicted as transporter-related genes) were also included in this region as “bacteriocin.” The genomic structure of the region surrounding Cluster OS3 in Pseudomonas sp. Os17 (gene IDs were given as Os1348–Os1351) is shown in Figure 1 and the annotation of these ORFs is summarized in Table 1. The Os1348 protein was annotated as an alpha chain of nitrile hydratase. Although the actual function of the Os1348 protein as a nitrile hydratase has not yet been clarified, recent bioinformatic research that examined thiazole/oxazole-modified microcin (low-molecular-weight bacteriocins referred to as TOMM) precursors based on comparative genomics revealed a new class of post-translationally modified peptides with strong similarities to the alpha subunit of the nitrile hydratase (Haft et al., 2010). This finding, together with the annotation data of the downstream genes described below, prompted us to speculate that Os1348 encodes the precursor and downstream genes are involved in post-translational modifications; however, conserved motifs among TOMM precursors, such as nitrile hydratase leader peptides, were not found in Os1348. Gene clusters encoding the predicted TOMM-type microcin have been identified in some
Pseudomonad genomes as homologs of the mcb operon that produces the DNA gyrase inhibitor microcin B17 (MccB17) in E. coli (Ghequire and De Mot, 2014). We also predicted the bacteriocin-encoding gene through the genomes of Os17 and St29 using the BAGEL 4 program (van Heel et al., 2018); however, Cluster OS3 was not detected, suggesting that it is not a typical bacteriocin. Only one gene in each strain (POS17_1251 for Os17 and PST29_1284 for St29) was predicted to be a putidacin by this program. We previously predicted that these two genes were the bacteriocin LlpA2 through annotation (Takeuchi et al., 2015). This gene was classified into 5,321 core genes that were shared with P. protegens Cab57.

The downstream genes Os1349–1351 were predicted to encode acetyltransferase, an uncharacterized protein (peptidase), and SagB-type dehydrogenase domain-containing protein, respectively, when predictions were performed using the Microbial Genome Annotation Pipeline (Takeuchi et al., 2015). A recent study on streptolysin, a bacteriocin from Streptococcus pyogenes, revealed that the 53-aa precursor peptide SagA was modified by the downstream gene products of sagBCD (homolog of the mcb operon), which encodes the dehydrogenase SagB, cyclodehydratase SagC, and docking protein SagD, respectively, jointly functioning for the introduction of thiazole and oxazole heterocycles onto SagA to be an active cytolysin (Lee et al., 2008).

In the genome of Os17, the homologs of sagA, sagC, and sagD were not found in our search.

When we performed searches with protein BLAST, the N- and C-terminal regions of Os1350 showed similarities to the peptidase MA superfamily (Gluzincin, pfam13485) and HEXXH motif domain-containing protein (TIGR04267), respectively (Table 1). The Gluzincin family includes several zinc-dependent metallopeptidases and contains HEXXH and EXXXD motifs as part of their active site (NCBI; Hooper, 1994).

The genes Os1348–1351 were not as commonly conserved in other strains as the downstream genes (Os1352-1355) based on the number of top hits of each gene in the Blastp analysis (Table 1). None of the three P. protegens strains tested (Cab57, Pf-5, and CHA0) possessed the genes Os1348–1351, whereas they possessed the homologs of Os1352-1355 (Figure 1). The Os1352-1354 gene cluster was predicted to encode the EmbABC efflux transporter, and Os1355 was annotated as their transcriptional regulator EmbR. The homolog of this region has been characterized in the DAPG-producing biocontrol strain Pseudomonas fluorescens 2P24, in which the deletion of each gene of embABC increased the extracellular accumulation of DAPG, whereas the embR-deficient mutant decreased the biosynthesis of DAPG (Tian et al., 2010).
### TABLE 1 | Similarities in *Pseudomonas* sp. Os17 ORFs to proteins in databases.

| Gene ID of Os17* | Position | Products size (aa) | NCBI Ref. of the closest protein, blastP E-value | Predicted function | Number of Blast top hits** |
|------------------|----------|--------------------|------------------------------------------------|-------------------|---------------------------|
| **Cluster OS3 (1348-1351)** | | | | | |
| 1348 (RS06725) | 1495345..1495602 | 85 | PMI18_04289 \*Pseudomonas sp. GM102 1.0E-29 \Plam14407 5.2E-04 | Nitrile hydratase, alpha chain | 2 |
| | | | Pfam14407 | | Frankia peptide, ribosomally synthesized peptide |
| 1349 (RS06730) | 1495717..1496166 | 149 | PMI18_04288 \*Pseudomonas sp. GM102 2.0E-50 \Plam00683 7.8E-10 | Acetyltransferase | 10 |
| | | | Pfam14407 | | Acetyltransferase (GNAT) family |
| 1350 (RS06735) | 1496187..1497347 | 386 | PMI18_04287 \*Pseudomonas sp. GM102 1.0E-133 \Plam13485 1.5E-04 \TIGR04267 9.3E-08 | Uncharacterized protein | 4 |
| | | | Pfam14407 | | Peptidase MA superfamily (Gluzincin) HEXXH motif domain |
| 1351 (RS06740) | 1497363..1499815 | 660 | PMI30_00628 \*Pseudomonas sp. GM50 0.0 \cd02142 7.6E-41 | SagB-type dehydrogenase domain-containing protein | 6 |
| | | | YP_347010.1 \*Pseudomonas fluorescens Pf0-1 0.0 | mcbC-like oxidoreductase |
| **Putative transporter-related genes (1352-1354)** | | | | | |
| 1352 (RS06745) | 1499376..1500836 (complement) | 486 | YP_258460.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | Acriflavine resistance protein A | 11 |
| | | | YP_004352462.1 \*Pseudomonas brassicacearum subsp. brassicacearum NFM421 0.0 | | > 100 |
| 1353 (RS06750) | 1500833..1503976 (complement) | 1047 | YP_258460.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | Acriflavine resistance protein A | 11 |
| | | | YP_258461.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | | > 100 |
| 1354 (RS06755) | 1503980..1505137 (complement) | 385 | YP_004352462.1 \*Pseudomonas brassicacearum subsp. brassicacearum NFM421 0.0 | Acriflavine resistance protein A | 11 |
| | | | YP_258461.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | | > 100 |
| 1355 (RS06760) | 1505403..1506053 | 216 | YP_258462.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | Transcriptional regulator TglR | 11 |
| | | | YP_258462.1, PFL_1332 \*Pseudomonas protegens Pf-5 0.0 | | Transcriptional regulator EmhR |

*Gene ID given in the *Pseudomonas* genomic database (www.pseudomonas.com) are indicated in brackets. **The sequence identity cut-off was set to 90%. A Blast search was performed on July 23, 2018.

**Construction of Os1348 and Downstream Gene Mutants and Their Effects on Antibiotic Activity**

Annotation data prompted us to predict that this region contributes to the antibiotic activity of the strains that possess it. We investigated whether the deletion of each gene affected the antibiotic activity of strain Os17 in a biotest using *B. subtilis*, which has been used as an indicator strain to test susceptibility against pseudomonads (Takeuchi et al., 2009, 2012, 2014, 2015), on modified GCM plates. The rationale for using this medium was to prevent the excessive halo size observed in strain Os17. Growth inhibitory activity against *B. subtilis* was weaker in the Os1348 mutant than in the wild-type (Table 2 and Figure 2).
Based on the well-known function of bacteriocins, in which they are only deleterious to members of a certain bacterial species or a subset of phylogenetically close relatives of the producer, we also tested other *Pseudomonas* strains (*P. syringae* pvs. *pisi* 730032 and *tabaci* 301612, *P. chlororaphis* St508 and *P. protegens* Cab57) as the reporter of close relatives (Supplementary Figure 2). *P. syringae* and *P. chlororaphis* strains were sensitive to the Os17 wild-type showing a dim halo; however, the Os1348 mutant abolished the clear halo, whereas the single mutant of *phlD* in the *additional mutation in the* gene, which encodes polyketide synthase, a key enzyme in DAPG biosynthesis (Table 3). The deletion in the *Os1353* gene did not affect the production level of DAPG under the conditions tested. The additional mutation in the *phlD* gene on the *Os1348* and *Os1353* mutants also abolished the clear halo, whereas the single mutant of *phlD* did not (Figure 2), suggesting that the antibiotic activity observed in this experiment mainly accounts for the DAPG production level of each strain, and that *Os1348* and *Os1353* are involved

| Test number and bacterial strain | Halo size against *Bacillus subtilis* (mm) |
|---------------------------------|------------------------------------------|
| 1 Os17 (wild-type)              | 7.0 ± 1.3                                |
| Δ1348                           | 5.0 ± 0.6*                               |
| 2 Os17                          | 8.0 ± 0.0                                |
| Δ1349                           | 8.8 ± 1.2                                |
| 3 Os17                          | 8.7 ± 0.9                                |
| Δ1350                           | 8.3 ± 1.1                                |
| 4 Os17                          | 6.8 ± 0.7                                |
| Δ1351                           | 3.5 ± 0.4*                               |
| 5 Os17                          | 6.5 ± 0.4                                |
| Δ1353                           | 10.0 ± 0.0*                              |
| 6 Os17                          | 6.8 ± 0.3                                |
| Δ1348 + 1348                    | 6.9 ± 0.2                                |
| 7 Os17                          | 8.1 ± 0.6                                |
| ΔgacA                           | ND                                       |

*Data are shown as the averages of 3–6 replicates ± standard deviation. Halo sizes were measured from the edge of the bacterial spot to the end of the clear zone. *Indicates a significant difference from that in Os17 at *P* < 0.05 (t-test).*

**FIGURE 2** Effects of the *phlD* mutation on antibiotic activity. The antibiotic activities of *Pseudomonas* strains grown on modified GCM plates were evaluated by the size of the growth inhibition zone of *B. subtilis*. (A) The antibiotic activities of the *Pseudomonas* sp. Os17 wild-type (WT), 1348 mutant (Δ1348), and 1353 mutant (Δ1353) were compared. (B) The effects of the *phlD* mutation on each strain in panel (A) were compared as follows: *phlD* mutant (Δ*phlD*), 1348/*phlD* mutant (Δ1348Δ*phlD*), and 1353/*phlD* mutant (Δ1353Δ*phlD*) were compared.
TABLE 3 | DAPG production levels by Pseudomonas sp. Os17 and its mutants grown in modified GCM medium.

| Bacterial strain tested | [DAPG] (nmol OD\textsubscript{600} \textsuperscript{-1}) \textsuperscript{a} |
|-------------------------|----------------------------------------------------------|
| Os17 (wild-type)        | 1.07 ± 0.18                                              |
| Δ1348                   | 1.87 ± 0.91                                              |
| Δ1353                   | 49.86 ± 15.31\textsuperscript{*}                         |

\textsuperscript{a}Data are shown as the averages of three replicates ± standard deviation. DAPG concentrations were indicated per optical density at 600 nm (OD\textsubscript{600}) in strains grown in modified GCM to OD\textsubscript{600} = 0.5 (one OD\textsubscript{600} unit corresponds to approximately 1 × 10\textsuperscript{9} cells/ml). *Indicates a significant difference from that in Os17 at P < 0.01 (t-test).

in the expression of antibiotic activity other than that caused by DAPG. The hazy halo observed in the 1348/phlD double mutants (and also in 1353/phlD) may have been due to the production of remaining secondary metabolites. This will be addressed in the section “Discussion.”

Expression of the Recombinant Os1348 Protein in E. coli

To obtain a more detailed understanding of the characteristics and structure of Os1348, we generated a full-length recombinant Os1348 protein without the use of tags by expression in E. coli.

The antibiotic activity of the recombinant Os1348 protein toward B. subtilis was tested, but not detected (data not shown), suggesting that the Os1348 protein needs to be modified post-translationally in order to exert its antibiotic effects or that the protein itself does not act as an antimicrobial protein that directly inhibits the growth of B. subtilis.

Downstream Genes of Os1348 Affect the Affinity of the Os1348 Protein Toward the Antibody

We hypothesized that the Os1348 protein was modified post-translationally. Therefore, we assessed the Os1348 protein status in Pseudomonas sp. Os17. By utilizing rabbit polyclonal antibodies raised against the Os1348 recombinant protein, we detected the Os1348 protein (ca. 8.9 kDa) with western blotting in wild-type cells, but not in Os1348 mutant cells. Intracellular levels of the Os1348 protein differed in Os1350- or Os1351-disrupted cells at higher levels of detection and in Os1349-disrupted cells at lower levels of detection from that in wild-type cells, suggesting that the Os1348 protein levels in these mutants are quantitatively different from that in the wild-type or that the expression level of the Os1348 protein is under the control of downstream genes (Figure 3). The increased level of the Os1348 protein in the 1350 mutant is consistent with the predicted function of the Os1350 protein as a peptidase; however, peptidase activity has not yet been demonstrated. The mobility shift in molecular mass was not observed in the 1351 mutant, suggesting that it has little, if any, responsibility for modifications. Assuming that the Os1351...
protein functions as a dehydrogenase, an increase in multiples of 2 Da will be observed in the deficient mutant, which may have been difficult to detect using the present method.

We also subjected the supernatant of the culture medium of the wild-type to western blotting to investigate the status of protein secretion into the medium. However, it was below the detection limit (data not shown).

Effects of Complementation of the 1348 Mutant

Complementation of the 1348 gene into the chromosome at its original position restored antibiotic activity (Table 2) as well as the expression level of the Os1348 protein (Figure 3) to wild-type levels, indicating that the phenotype was due to inactivation of the 1348 gene and not to polar effects on neighboring genes. Regarding the complementation of the 1348 mutant by a plasmid, we constructed a plasmid harboring 1348-1351 genes (full region of OS3) based on the potential for post-translational modifications by the products of the downstream genes of 1348. Complementation of the 1348 mutant by this plasmid led to higher expression levels of Os1348 than that of the wild-type, presumably because a multicopy plasmid was used (Figure 3). However, complementation by plasmids did not restore antibiotic activity (data not shown), suggesting that the gene needs to be located in the original position to exert its function on antibiotic activity, i.e., the OS3 cluster needs to be located upstream of the putative transporter-related genes (Os1352-1354) in order for the Os1348 protein to be exported. In a number of bacteriocins, the genetic organization of bacteriocin gene cluster have been found to include transporter genes in the downstream region (Haft et al., 2010; Ghequire and De Mot, 2014), suggesting that the positioning of transporter genes is relevant.

X-ray Crystal Structure of Os1348

The full-length Os1348 protein was overexpressed in E. coli, purified, and crystallized. The Os1348 structure was elucidated by the molecular replacement method using the ideal helices mode at a resolution of 1.9 Å (Figure 4 and Table 4). The final model included four Os1348 molecules consisting of 2–79...
residues in the asymmetric unit. The six C-terminal residues were not observed in the electron density map and appeared to be highly flexible.

The Os1348 monomer was composed of six α helices (α1-α6) and formed a Magatama (comma)-shaped α-structure (Figures 4A,B). Two Os1348 monomers joined and formed a dimeric structure. The Os1348 dimer had a semicircular shape (Figure 4C). The dimer interface was composed of many hydrophobic amino acids from the α1, α4, α5, and α6 helices. Hydrophilic interactions, namely, hydrogen bonds, and salt bridges, also appeared to contribute to the stability and solubility of the Os1348 dimer. Based on its three-dimensional structure, the free energy values ΔГint and ΔГdist of the Os1348 dimer were calculated by the PISA website as −20.4 and 17.0 kcal/mol, respectively, suggesting that the Os1348 protein formed a stable homodimer in solution.

The Os1348 protein showed no structural similarities to other bacteriocins or proteins registered in the PDB. Although the Os1348 protein was annotated as an alpha chain of nitrile hydratase, no structural similarities to the nitrile hydratases registered in the PDB were observed. Therefore, the function of the Os1348 protein and mechanisms underlying its antimicrobial activity remain unclear. Based on the results of biological investigations, the full-length Os1348 protein may be a bacteriocin precursor and appears to be post-translationally modified into an active form by the products of downstream genes.

The gacA Mutant Exhibits Decreased Expression Levels of Os1348-lacZ

In P. protegens and other fluorescent pseudomonads, the GacS/GacA system positively controls secondary metabolism. We previously reported that the gacA mutant of strain Os17 lost its antibiotic activity against B. subtilis, similar to other biocontrol strains of pseudomonads. Furthermore, the production of rhizoxin analogs, which were found to be major biocontrol factors of this strain, was also under the control of GacA (Takeuchi et al., 2015). To establish whether the production of the Os1348 protein in strain Os17 was also under the control of the GacS/GacA system, we examined production levels using western blotting. In the gacA mutant, the accumulation of the Os1348 protein was below the detection limit (Figure 3).

In Cluster OS3 of the Os17 genome, the Os1348 gene has a potential RsmA/E binding site (UAAGGAGA) in its upstream region, which may expose the conserved AAGGAG hexa-loop on a U-A base pair (underlined). Therefore, we constructed translational Os1348-lacZ fusion, and compared expression levels in the wild-type and gacA mutant (Figure 5). In the gacA mutant, a decreased level of expression was observed at high cell densities, suggesting that the GacA-dependent regulation of the gene may account for the decreased level of the Os1348 protein in the gacA mutant.

**DISCUSSION**

The present results demonstrated that comparative genomics among closely related root-colonizing pseudomonad strains, including model and non-model strains, revealed a novel gene cluster whose expression contributes to the antibacterial activity of the non-model strain. Since the genomic sequence of model strain Pf-5 was completely elucidated (Pausen et al., 2005), it has provided novel insights into genome-mining strategies to identify antimicrobial secondary metabolites from agriculturally important bacteria (Gross et al., 2007). These strategies have also identified small proteinaceous and peptidic structures, such as bacteriocins, which are ribosomally synthesized and post-translationally modified (Ghequire and De Mot, 2014). The relatively large genome sizes of P. protegens (ca. 6.7–7.1 Mbp) and related strains are attributed to the variety of genomic repertoires for the synthesis of competitive weapons against other organisms in the rhizosphere. The genome sizes of our recently isolated strains Os17 and St29, which were revealed to be same species, were 6.89 and 6.83 Mbp, respectively (Takeuchi et al., 2015). Some clusters conserved in both strains were predicted to be involved in the production of secondary metabolites, and one, named Cluster OS3, was examined in the present study.

In the genome of strain Os17, there are several gene clusters other than phl (for the production of DAPG) that are involved in the biosynthesis of known secondary metabolites with antibiotic activities. The representative clusters are rzx and hcn (for the production of rhizoxin analogs and HCN, respectively).
The aprA gene cluster, for the major extracellular protease AprA, is also conserved in this strain (Takeuchi et al., 2015). Under the experimental conditions tested in the present study, DAPG is the major factor for inhibitory activity of B. subtilis, and the minor is related to the cluster OS3. The hazy halo observed in the 1348/phlD double mutant (and also in 1353/phlD) in Figure 2 may due to the production of remaining secondary metabolites. It is important to note that reciprocal regulation between antibiotics has been reported in P. protegens strains CHA0 and Pf-5, while that of pyoluteorin and the phloroglucinol derivatives has been investigated in the most detail (Schnider-Keel et al., 2000; Brodhagen et al., 2004; Baehler et al., 2005; Kidarsa et al., 2011). In the present study, the mutational loss of the phlD gene may have affected the production of antibiotics other than pyoluteorin, which is absent from Os17, and, thus, resulted in the hazy halo.

Two gene clusters, plt and prn, which are involved in the production of pyoluteorin (PLT) and pyrrolnitrin (PRN), respectively, were found to be missing in the genomes of strains Os17 and St29. They are two out of four typical gene clusters of P. protegens other than phl and hcn. Genomic variations may contribute to the uniqueness of the species of these strains. The size of strains Os17 and St29 were equivalent to that of P. protegens strains; this may be due to the presence of unknown gene clusters that encode novel secondary metabolites. For example, in the genome of strain Os17, two neighboring clusters were partly found to be homologs of the biosynthetic clusters for the antibiotic mitomycin from Streptomyces lavendulae (Takeuchi et al., 2015). Although the function of these clusters has not yet been elucidated, they will contribute to the identification of novel secondary metabolites.

In the present study, a functional analysis of the gene cluster Os1348–Os1351 was performed, and the Os1349–Os1351 genes were predicted to be involved in post-translational modifications to the small precursor protein Os1348. The deletion of Os1350
resulted in an increased Os1348 protein level (Figure 3). This result is consistent with the predicted function of the Os1350 protein as a peptidase, in which the N-terminal region of Os1350 showed similarities to the peptidase MA superfamily (Gluzincin); however, peptidase activity has not yet been demonstrated. Peptidases and proteases need to have strict selectivity in order to avoid the deleterious and random degradation of functional proteins. A recent study on a metallopeptidase-harbor ing gluzin cin motif revealed that it recognized O-linked glycan modifications on protein substrates for its catalytic machinery (Noach et al., 2017). If the Os1350 protein functions as a peptidase and targets the Os1348 protein, this raises a number of questions about selectivity, i.e., whether post-translationally modified proteins are recognized. There is another speculation on the increased level of Os1348 observed in the 1350 mutant, concerning the inability of the cell to export the unmodified peptide.

Regarding Os1351, which was predicted to encode a SagB-type dehydrogenase domain-containing protein, an important finding was previously reported on streptolysin, a bacteriocin from S. pyogenes, as described above. Besides streptolysin, a similar mechanism of modification is utilized in the biosynthetic pathways for microcin B17 (Ghequire and De Mot, 2014), the patellamides (Schmidt et al., 2005), and the thiopetides (Wieland Brown et al., 2009). The products of these clusters have been collectively classified as TOMMs due to the conservation of their genetic and chemical structures (Haft et al., 2010). Although the relevance of Sag-type modifications remains unknown in Os17, a more detailed understanding of the roles and mechanisms underlying protein modifications using gene products of the OS3 cluster represents a future challenge in bioengineering.

The Os1348 protein in the present study cannot be regarded as a bacteriocin from a rigorous definition point of view because we were unable to demonstrate that it was secreted into the medium due to the detection limit. The mutational loss of the putative efflux transporter gene Os1353 increased the protein accumulation of Os1348 in the cell, which suggests the presence of a transport system. Considering the spectrum of the strains that were tested, Os1348 may not be categorized as a typical bacteriocin, which is defined as toxins against those that are closely related to bacteriocin-producing strains.

In addition to direct antibiotic activity, peptides secreted through a bacteriocin export system have been shown to possess signaling functions in a manner of quorum sensing and bacterial crosstalk (Michiels et al., 2001). The Os1348 protein also showed homology to the Type VI secretion protein ImpH (27% identical at 41 aa without gaps, Supplementary Figure 3), which was originally identified as one of the gene products of the nodulation impairment locus imp in Rhizobium leguminosarum (Roest et al., 1997; Bladergroen et al., 2003). Since the Type VI secretion system (T6SS) has been shown to play various roles in the social behavior of Gram-negative bacteria, besides the primary role assigned to T6SS as a potent weapon during interspecies competition, investigations on the possible functions of the Os1348 protein as a signaling molecule will be an interesting aspect of future studies. Omics approaches, such as RNA-seq analyses based on comparisons of wild-type and mutant strains, may provide a more accurate view of the possible function of this protein in a signaling role.

In the present study, we did not detect the antibiotic activity of the full-length recombinant Os1348 protein, the crystal structure of which showed no structural similarities to other bacteriocins or proteins registered in the PDB. The native Os1348 protein is suggested to be post-translationally modified by the products of downstream genes, and this maturation process may be indispensable for this protein to be an active form. Further studies are needed to investigate whether the native Os1348 protein forms a dimer, and if so, whether the dimerization of the Os1348 protein affects the biological activity of this protein. Taken together with the relevance of post-translational modifications in this protein, a structural analysis of the native Os1348 mature protein combined with that of each mutant of the downstream genes will provide an in-depth quality assessment of the protein.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://doi.org/10.2210/pdb6LK7/pdb.

AUTHOR CONTRIBUTIONS

KT and WT conceived and designed research. KT, WT, ZF, and KY conducted the experiments. NS contributed the strains. KT, WT, ZF, and TY analyzed the data. KT, WT, and ZF wrote the manuscript. All authors read and approved the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020.605705/full#supplementary-material
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