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INVESTIGATION

# Multiple Mechanisms Contribute to Lateral Transfer of an Organophosphate Degradation (*opd*) Island in *Sphingobium fuliginis* ATCC 27551

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ABSTRACT The complete sequence of pPDL2 (37,317 bp), an indigenous plasmid of Sphingobium fuliginis ATCC 27551 that encodes genes for organophosphate degradation (opd), revealed the existence of a sitespecific integrase (int) gene with an attachment site attP, typically seen in integrative mobilizable elements (IME). In agreement with this sequence information, site-specific recombination was observed between pPDL2 and an artificial plasmid having a temperature-sensitive replicon and a cloned attB site at the 3' end of the servil tRNA gene of Sphingobium japonicum. The opd gene cluster on pPDL2 was found to be part of an active catabolic transposon with mobile elements y4qE and Tn3 at its flanking ends. Besides the previously reported opd cluster, this transposon contains genes coding for protocatechuate dioxygenase and for two transport proteins from the major facilitator family that are predicted to be involved in transport and metabolism of aromatic compounds. A pPDL2 derivative, pPDL2-K, was horizontally transferred into Escherichia coli and Acinetobacter strains, suggesting that the oriT identified in pPDL2 is functional. A welldefined replicative origin (oriV), repA was identified along with a plasmid addiction module re/B/re/E that would support stable maintenance of pPDL2 in Sphingobium fuliginis ATCC 27551. However, if pPDL2 is laterally transferred into hosts that do not support its replication, the opd cluster appears to integrate into the host chromosome, either through transposition or through site-specific integration. The data presented in this study help to explain the existence of identical opd genes among soil bacteria.

#### **KEYWORDS**

integrative conjugative elements (ICE) genomic islands catabolic transposons phosphotriesterase (PTE) organophosphates

Microbial metabolism of organophosphates (OP) attracted the attention of microbiologists as it contributes to the elimination of toxic OP insecticide residues from agricultural soils (Singh 2009). Microbial enzymes involved in degradation of OP compounds are divided into

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three major groups: organophosphate acid anhydrolases (OAA), phosphotriesteases, and methyl parathion hydrolases (Singh 2009). The OAAs were later shown to be prolidases involved in hydrolysis of the peptide bond of dipeptides with proline at their C terminus (Cheng et al. 1997; DeFrank and Cheng 1991; Vyas et al. 2010). Nerve agents like soman (GD; O-pinacolyl methylphosphonofluoridate), sarin (GB; O-isopropyl methylphosphonofluoridate), GF (O-cyclohexylmethylphosphonofluoridate), and tabun (GA; ethyl N,N-dimethylphosphoramidocyanidate) (Cheng et al. 1999) have been shown to be fortuitous substrates of prolidases (Cheng et al. 1999). Unlike the prolidases, no physiological substrate has yet been found for the other two OP hydrolyzing enzymes. Phosphotriesteases (PTE), encoded by the organophosphate degradation (opd) gene, are membrane-associated metalloenzymes and contain divalent Zn ions at their catalytic site (Benning et al. 2001; Mulbry and Karns 1989). The PTEs catalyze hydrolysis of the ester linkage found in structurally diverse groups of organophosphates, including nerve agents (Benning et al. 1994; Cheng and DeFrank 2000; Cho et al. 2004; Lai et al. 1995). The PTE purified

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Sequence data from this article have been deposited with the EMBL/GenBank Data Libraries under accession no. JX31-2671.

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#### Table 1 Bacterial strains and plasmids

Strain or Plasmid	Genotype or Phenotype	Reference or Source
E. coli DH5α	supE44 $\Delta$ lacU169 (Ø80 lacZ $\Delta$ M15) hsdR17 recA1 endA1 gyrA96 thi1 relA1	Hanahan (1983)
E. coli BL21 DE3	hsdS gal(λ clts857 ind1 sam7 nin5lac UV5 T7 gene 1	Studier and Moffatt (1986)
E. coli EC100D pir <sup>+</sup> 116	F <sup>-</sup> mcrA $\Delta$ (mrr hsdRMS <sup>-</sup> mcrBC) φ80dlacZ $\Delta$ M15 $\Delta$ lacX74 recA1 endA1 araD139 $\Delta$ (ara, leu)7697 galU galK $\lambda^-$ rpsL (StrR) nupG pir-116(DHFR)	Epicentre Biotechnologies, USA
E. coli HB101	$F^{-}$ mcrB mrr hsdS20( $r_{B}^{-}$ m $_{B}^{-}$ ) recA13 leuB6 ara-14 proA2 lacY1 galK2 xyl-5 mtl-1 rpsL20(Sm <sup>R</sup> ) glnV44 $\lambda^{-}$	Boyer and Roulland-Dussoix (1969)
E. coli S17-1	thi pro hsdR hsdM recA RP4 2-Tc::Mu-Kn <sup>r</sup> ::Tn7(Tp <sup>r</sup> Sp <sup>r</sup> Sm <sup>r</sup> )	Simon <i>et al.</i> (1983)
Sphingobium fuliginis ATCC 27551	Wild type strain, Sm <sup>r</sup> , Pm <sup>r</sup> , OPH <sup>+</sup>	Kawahara <i>et al.</i> (2010)
Acinetobacter sp.DS002	Cm <sup>r</sup> , Sm <sup>r</sup> , Ben <sup>+</sup>	Unpublished results from our lab
pRSET-A	Amp <sup>r</sup> , expression vector	Invitrogen, USA
pBluescript KS(II)	Amp <sup>r</sup> , <i>lacZ</i> <sup>+</sup> , cloning vector	Fermentas, USA
pET-23b(+)	Amp <sup>r</sup> , expression vector	Novagen, USA
pMMB206	Cm <sup>r</sup> , a low copy number broad host range expression vector	Morales et al. (1991)
pJQ210	Gm <sup>r</sup> , sacB <sup>+</sup>	Quandt and Hynes (1993)
pKD46	Amp <sup>r</sup> , Red recombinase expression plasmid	Datsenko and Wanner (2000)
pRK2013	Kmr, ColE1-based restricted-host-range helper plasmid	Ditta <i>et al.</i> (1980)
pSM3	Amp <sup>r</sup> , Tc <sup>r</sup> , opd gene replaced with opd::tet in plasmid pSM2	Siddavattam <i>et al.</i> (2003)
pPDL2	opd <sup>+</sup> , 37.3Kb indigenous plasmid	Mulbry et al. (1986)
pPDL2-K	<i>opd</i> <sup>+,</sup> Km <sup>r</sup> , pPDL2 having single minitransposon EZ-Tn5 <r6kγori kan-2=""> insertion</r6kγori>	This study
pPDL2-KT	opd <sup>+,</sup> Km <sup>r</sup> , Tet <sup>r</sup> , pPDL2-K with replacement of opd with opd::tet	This study
pP4I	oriT sequence of pPDL2 cloned in pBluescript KS(II)	This study
pSFT1	repB gene cloned in pRSET-A as BamHI and Xhol fragment	This study
pSFT2	Tc <sup>r</sup> , Mini replicon generated by ligating <i>oriV</i> and <i>repA</i> region of pPDL2 to <i>opd::tet</i> cassette	This study
pSDP7	Amp <sup>r</sup> , temperature sensitive cloning vector	This study
pSDP8	Amp <sup>r</sup> , attB sequence cloned in pSDP7 as HindIII fragment	This study
pSDP9	ligA and ligB genes cloned in pET-23b(+) as NdeI and HindIII fragment	This study

from *Brevundimonas diminuta* hydrolyses parathion, the model OP compound, at a rate close to its diffusion limits and is considered to be an end point in enzyme evolution (Caldwell and Raushel 1991; Scanlan and Reid 1995). The OPs were recently (65 years ago) introduced into agriculture pest management, mainly as replacements for more persistent organochlorides (Davies *et al.* 1985). Therefore, evolution of PTEs in such a short period has become an interesting model in which to study molecular evolution (Elias and Tawfik 2012; Scanlan and Reid 1995). The studies conducted to date have shown quorum-quenching lactonases as progenitors of PTEs (Afriat *et al.* 2006; Elias and Tawfik 2012; Roodveldt and Tawfik 2005).

The third prominent organophosphate-degrading enzymes are methyl parathion hydrolases (MPH), encoded by the methyl parathion degradation (mpd) gene. The MPH enzymes known to date have been purified from Pseudomonas strains isolated from OP-polluted Chinese agricultural soils (Liu et al. 2005; Zhang et al. 2006; Zhongli et al. 2001). Despite having functional similarity, MPH enzymes share no homology with PTEs, indicating the existence of structurally independent organophosphate-degrading enzymes among soil bacteria (Dong et al. 2005). Unlike organophosphate hydrolase (OPH), the MPH has been shown to be a structural homolog of  $\beta$ -lactamases, the enzymes that confer resistance to  $\beta$ -lactam-derived antibiotics (Tian et al. 2008). Interestingly these two structurally different enzymes have an identical active site structure (Dong et al. 2005). This type of functional convergence pointed to the existence of independent paths in the evolution of organophosphate-hydrolyzing enzymes. Lateral transfer of mpd genes became evident with the discovery of identical mpd genes among bacterial strains isolated from

OP-polluted Chinese soil samples (Zhang *et al.* 2006), and the *mpd* genes were shown to be part of an active transposon (Wei *et al.* 2009).

Identical *opd* genes have been found in soil bacteria isolated from diverse geographical regions. The dissimilar *opd* plasmids pCMS1 and pPDL2 were isolated from *B. diminuta* (pCMS1) and *Flavobacterium* sp. ATCC 27551 (pPDL2) that were enriched, respectively, from soil samples collected from agricultural fields of Texas, USA, and the International Rice Research Institute (IRRI), Philippines. They were shown to contain identical *opd* genes (Mulbry *et al.* 1986; Pandeeti *et al.* 2011), and a 7 kb region around the *opd* gene apparently constituted the only identity between these two dissimilar plasmids (Mulbry *et al.* 1986; Pandeeti *et al.* 2011; Siddavattam *et al.* 2003). However the sequence of the *opd* gene cluster in the self-transmissible pCMS1 showed no features of a transposable element (Pandeeti *et al.* 2011), and although a transposon-like organization was found in pPDL2, the *opd* cluster of pPDL2 has not been shown to be an active transposon (Siddavattam *et al.* 2003).

In the present study, we report the complete sequence of pPDL2, and we show experimentally that pPDL2 is a mobilizable plasmid having the capability to spread the *opd* cluster laterally by functioning both as an integrative mobilizable element and as an active transposon.

## **MATERIALS AND METHODS**

#### Strains and plasmids

Bacterial strains and plasmids used in the present study are shown in Table 1. Oligonucleotide primers used while performing PCR reactions are listed in supporting information, Table S1. *E. coli* and *Acinetobacter* sp. DS002 cells were grown in LB medium at 37° and 30°,

respectively. The cultures of *Sphingobium fuliginis* ATCC 27551 were grown in modified Wakimoto medium at 30°. When necessary, the antibiotics ampicillin (100  $\mu$ g/ml), tetracyclin (30  $\mu$ g/ml), gentamycin (20  $\mu$ g/ml), and kanamycin (30  $\mu$ g/ml) supplemented the culture medium. The LB sucrose plates were prepared by adding 5% sucrose to LB agar plates. *Acinetobacter* sp. DS002 strains were grown in M9 medium containing filter-sterilized benzoate (5 mM) as a carbon source. Exconjugants of *Acinetobacter* sp. DS002 strains were selected on M9 agar plates supplemented with kanamycin.

# Isolation and rescue cloning of pPDL2 from *Sphingobium fuliginis* ATCC 27551

Large indigenous plasmids of Sphingobium fuliginis ATCC 27551 were isolated by following the Currier Nester protocol with minor modifications (Currier and Nester 1976; Pandeeti et al. 2011). The plasmid preparation made from Sphingobium fuliginis ATCC 27551 was directly used for tagging with minitransposon EZ-Tn5<R6Kyori/KAN-2> using the EZTn5 <R6Kyori/KAN-2> insertion kit (Epicenter Biotechnologies, USA) following the manufacturer's protocols. The isolated plasmid preparation and minitransposon was taken in equimolar concentrations and incubated with transposase for 2 hr at 37° to complete in vitro transposition. After 2 hr, transposase was inactivated by adding 1  $\mu$ l of 1% SDS before incubating the reaction mixture for 10 min at 70°. The reaction mixture was then electroporated into E. coli EC100D pir-116 cells by setting the electroporator (Gene Pulser, Bio-Rad Laboratories, USA) at 2.5 kV, 200 $\Omega$  for 4.5 msec. After electroporation, the cells were plated on LB plates supplemented with kanamycin to select transformants. Colonies having plasmid pPDL2 were identified by performing colony PCR using opd specific primers. The colonies that gave amplification of the opd gene were used to isolate plasmids having the minitransposon Tn5<R6Kyori/KAN-2>. The derivatives of plasmid pPDL2, named pPDL2-K, were then isolated from E. coli pir-116 cells using the BAC isolation protocol (Sambrook et al. 1989). The restriction profile of pPDL2-K was generated by digesting with SmaI, BamHI, PstI, SalI, XhoI, HindIII, and EcoRI, and it was compared with the restriction profile of pPDL2 (Mulbry et al. 1987) to identify the exact point of minitransposon insertion. Further, all fragments of pPDL2-K obtained after digestion with PstI and EcoRI were ligated independently into pBluescript (II) KS (+) digested with similar enzymes.

# Generation of the complete pPDL2 sequence

DNA sequencing was performed by using BigDye Terminator v1.1 Cycle Sequencing Kit (Applied Biosystems, USA), and an ABI PRISM 3100 Genetic Analyzer was used following manufacturer's protocol. Either pPDL2 derivatives or subclones generated by cloning *Eco*RI and *Pst*I fragments were used as templates. Sequence reactions were generated using transposon-specific sequence primers when pPDL2 derivatives were used as templates, whereas universal forward and reverse primers were used to generate sequence from subclones. To fill gaps, a primer-walking strategy was employed. If the cloned fragment size was large, it was digested with different restriction enzymes and subcloned into pBluescript (II) KS(+) and were used for sequencing.

#### Sequence assembly and annotation of pPDL2

All sequences were viewed and edited to remove vector sequences by using Chromas 2.13 software (www.technelysium.com.au/chromas). Sequences were assembled into contigs by using the program Contig Express of VectorNTI software (Invitrogen Technologies, USA). The assembled sequence of plasmid pPDL2 was annotated using Artemis sequence annotation tool (http://www.sanger.ac.uk/resources/software/ artemis/) (Rutherford *et al.* 2000). The open reading frames (ORFs) were identified by using the built-in tool of the Artemis software, and the start codons in the predicted ORFs were fixed with the help of BLAST searches. BLAST searches were made against the nonredundant database of NCBI using BLAST program (www.ncbi.nlm.nih.gov/BLAST). IS elements, transposons, and their repeat elements were identified by doing a pairwise alignment using BlastN program against the ISfinder database (http://www-is.biotoul.fr/).

# Prediction of origin of replication (oriV)

The *oriV* was predicted based on sequence homology to other annotated or predicted *oriVs* and by performing GC skew analysis (Grigoriev 1998). While calculating the GC skew, the ratio of (G-C) to (G +C) was calculated per each window of equal length split over the total sequence of pPDL2. The GC skew was multiplied by w/c to find its dependence on subsequence length, where w and c are lengths of the subsequence and total sequence of pPDL2, respectively. As maximum and minimum value of GC skew is usually associated with termination and origin of replication (Frank and Lobry 1999), a similar approach was followed to predict *oriV* of plasmid pPDL2 origin.

# Prediction of attP sites

For predicting *attP* sites, two independent approaches were used. In the first approach, sequences of all available plasmids having identical integrases were collected from the NCBI database. From bacteria having these plasmids, a dataset of all tRNA sequences along with their upstream and downstream sequences was created. Pairwise alignments were made between the plasmids and the tRNA sequences to identify the *att* sites.

In a second approach, the genome sequences of bacteria having plasmids with integrase genes were collected from NCBI database. These genomic sequences were then used to predict genomic islands (GI) using Islandviewer software (http://www.pathogenomics.sfu.ca/islandviewer/query.php). The predicted GI sequences were then used to make pairwise alignments with sequences of tRNA genes of *Sphingomonas wittichii* and *Sphingobium japonicum*. Short sequence repeats that exactly matched the 3' end of tRNA sequences were taken as putative *att* sites. The predicted *att* sites with a low E-value were considered as potential *att*P and *att*B sites. The *att* sites predicted in this manner were then used to align with plasmid pPDL2 sequence using BlastN to identify *attP* homologs. Alternatively, tRNA gene sequences that perfectly matched the 3' end of tRNA genes.

#### **Bacterial conjugation experiments**

Biparental mating experiments were done using *E. coli pir*-116 (pPDL2-K) as donor and *Acinetobacter* sp. DS002 as recipient. The triparental mating experiments were performed by including *E. coli* strain HB101 (pRK2013) as a helper. Biparental and triparental experiments were performed following standard protocols described elsewhere (Figurski and Helinski 1979). The exconjugants were selected by plating on selection plates having minimal media supplemented with 5 mM benzoate and kanamycin. Parent strains treated in a similar manner were plated on selection plates and served as negative controls.

# Cloning of synthetic *attB* site in a temperaturesensitive vector

A temperature-sensitive plasmid vector was constructed by using vectors pMMB206 (Morales et al. 1991) and pKD46 (Datsenko and

Wanner 2000) as source plasmids. The temperature-sensitive replicative origin along with the bla gene were amplified as an EcoRV/SmaI fragment by using primers DSF005 and DSF006 as forward and reverse primers. The multiple cloning site and lac reporter system were taken from plasmid pMMB206 by digesting it with EcoRV/DraI. These two fragments were then ligated to give plasmid pSDP7. The servl tRNA with an attB sequence was synthesized before cloning it in pSDP7. Equal concentrations of the overlapping primers DSF001 and DSF002 were placed in a PCR tube in boiling water and left until the water reached room temperature. The contents were then brought down by briefly spinning, and then 2.5 µmol dNTP, a unit of pfu, and 10  $\mu$ l 10× pfu buffer were added. The contents were made up to 100  $\mu$ l with MilliQ water before incubating the tube at 72° for 10 min. An aliquot of 10 µl of the reaction mixture was then taken as a template to amplify attB by performing PCR using the primer set DSF003/DSF004 with HindIII recognition sites at the 5' end. The amplicon was then digested with HindIII and cloned into pSDP7 to give pSDP8.

#### In vivo integration assay

Plasmid pSDP8 was transformed into *E. coli pir*-116 (pPDL2-K), and the resultant cells were grown at  $30^{\circ}$  for 12 hr in LB medium with kanamycin and ampicillin. After incubation, cells were collected from 1 ml of culture and thoroughly washed before reinoculating them in fresh LB medium containing only kanamycin. The cultures were then grown for a further period of 12 hr at  $37^{\circ}$  to inhibit the replication of the temperature-sensitive plasmid pSDP8. After incubation, the culture was serially diluted, and appropriate dilutions were plated on LB agar with kanamycin and ampicillin. Plasmids were isolated from the resultant colonies, and the formation of a cointegrate due to sitespecific recombination between *attP* and *attB* was determined by performing restriction analysis. The *attB/P* and *attP/B* sites were identified by DNA sequencing using universal forward and reverse primers.

#### **Transposition assay**

For the transposition assay, we created a pPDL2-K derivative by replacing the *opd* gene with *opd::tet* through homologous recombination. *E. coli pir*-116 (pPDL2-K) was first transformed with the temperature-sensitive plasmid pKD46, which codes for red recombinase, in the presence of arabinose (Datsenko and Wanner 2000). The mid log-phase cultures obtained by growing in LB medium with 1 mM arabinose were made electrocompetent and immediately used to electroporate with insertionally inactivated *opd* (*opd::tet*) taken as a *PstI* fragment from pMS3 (Siddavattam *et al.* 2003). The tetracycline-resistant colonies were then used to perform colony PCR using *opd*-specific primers. The colonies identified as carrying *opd::tet* were grown on them at  $37^{\circ}$  to eliminate temperature-sensitive pKD46. The ampicilin-sensitive, tetracycline-resistant colonies were then used to isolate the pPDL2-K derivative pPDL2-KT and to confirm the presence of *opd::tet* by restriction analysis.

The transposition assay used in the present study is described elsewhere (Siddavattam *et al.* 2003), except that the *sacB* containing pJQ210 (Quandt and Hynes 1993) was used as a reporter plasmid and plasmid pPDL2-KT served as donor of *opd* cluster. *E. coli* (pJQ210) cells were made electrocompetent and plasmid pPDL2-KT was electroporated following standard protocols (Dower *et al.* 1988). The electroporated cells were then grown in LB medium having no antibiotics for 2 hr. The cells were then plated on LB plates having gentamycin, sucrose, and tetracyclin. As plasmid pPDL2-KT requires a *pir* background, the tetracycline-resistant colonies found on sucrose plates were considered to have been generated due a transposition event disrupting the *sacB* gene. These colonies were then used to isolate plasmid and to analyze junction sites by performing restriction profile and PCR amplification. The PCR amplification was done by using *sacB*-specific primers (DSF0015/DSF0016) and primers (DSF0017/DSF0018) designed using pPDL2 sequence found upstream of the predicted Tn3-specific terminal repeats.

#### Interactions between oriT and Mob protein

The predicted repB gene downstream of oriT was amplified using a primer set DSF009/DSF0010 appended with BamHI and XhoI sites. The amplicon was then cloned in expression vector pRSETA as a BamHI/XhoI fragment to generate recombinant plasmid pSFT1. E. coli BL21 (pSFT1) cells were induced to express RepB<sub>C-6His</sub> and affinity purified following procedures described elsewhere (Pandey et al. 2009). A 250 bp DNA fragment containing the oriT region was amplified using a primer pair (DSF0019 and DSF0020) designed by taking the sequence flanking oriT of pPDL2. The PCR amplicon containing oriT was end-labeled with  $[\gamma^{-32}P]$  ATP using T4 polynucleotide kinase following established protocols (Sambrook et al. 1989). While performing mobility shift assay, the pure labeled oriT (2 pmol) was taken in 20 µl of binding buffer [20 mM Tris-HCl (pH 8.0), 1.0 mM EDTA, 6 mM MgCl<sub>2</sub>, 50 mM KCl, 50 µg/ml BSA, 5% (w/v) glycerol, 5 µg/ml of herring sperm DNA] and incubated for 20 min at 25° with various concentrations (0 ng, 100 ng, 250 ng, 500 ng and 1000 ng) of RepB<sub>C-6His</sub>. The reaction mixture was then resolved on a 5% (w/v) native polyacrylamide gel, and the mobility of labeled oriT was captured on X-ray film by performing autoradiography. Control reactions were performed either by omitting RepB<sub>C-6His</sub> or by including increased concentrations (0.2 to 2 pmol) of cold oriT.

# Expression of the lig operon

The *lig* operon comprising the *ligB* and *ligA* genes was amplified from plasmid pPDL2 using primer set DSF0013/DSF0014 appended with NdeI-HindIII, respectively. The NdeI site in the forward primer corresponds to the 5' region of ligB and overlaps with its start codon. Similarly, the reverse primer designed by taking the 3' end of ligA contains a HindIII site created by changing the stop codon. The amplicon containing the ligBA genes was then cloned in pET23b as an NdeI/HindIII fragment to generate plasmid pSDP9. The cloning strategy followed to generate pSDP9 places both ligB and ligA under the transcriptional and translational signals of the expression vector. As the stop codon of ligA was eliminated to facilitate fusion of ligA with the vector-coded histidine tag, the LigA alone was expressed with C-terminal histidine tag. As active protocatechuate dioxygenease contained both LigB and LigA subunits, affinity purification using Niaffinity column was expected to yield LigBA complex. Expression of LigBA was monitored both by measuring dioxygenease activity (Ono et al. 1970) and by detecting LigA<sub>C-6His</sub> from Western blots using antihis antibody (Pandey et al. 2009).

#### **Enzyme assays**

Protocatechuate-4,5-dioxygenase assay was performed using purified LigBA complex. The reaction mixture contained protocatechate (20  $\mu$ M) and 5  $\mu$ g of purified LigBA complex in 100 mM Tris acetate buffer (pH 7.2). Formation of the ring clevage product, 2-hydroxy-4-carboxymuconic semialdehyde, was measured spectroscopically following established procedures (Ono *et al.* 1970). Similarly, formation of 2-hydroxymuconic semialdehyde was monitored at 375 nm when

catechol was used as a substrate (Nozaki *et al.* 1970). The organophosphate hydrolase (OPH) assay was performed using methyl parathion as substrate. The OPH activity was measured by estimating the concentration of *p*-nitrophenol (Pakala *et al.* 2007). Specific activity of enzymes was expressed as micromoles of product formed per minute per milligram of protein.

#### RESULTS

Flavobacterium sp. ATCC 27551 was one of the first organophosphate-degrading bacteria isolated from the agricultural soils of the International Rice Research Institute, Manila, Philippines (Sethunathan and Yoshida 1973). Recently it was reclassified as Sphingobium fuliginis ATCC 27551 (Kawahara et al. 2010). The OP degradation property has been shown to be encoded on one of its four indigenous plasmids, pPDL2 (Mulbry et al. 1986), the other three not being involved in this activity. Lateral gene transfer of opd is evident due to the existence of identical opd genes in bacterial strains belonging to different taxonomic groups. However, no systematic studies are available to explain the mechanism of lateral transfer of the opd gene cluster. Because pPDL2 contains a transposon-like opd cluster, we have taken it as a model system to study horizontal gene transfer (HGT) of the opd cluster. Initially pPDL2 was isolated from the other indigenous plasmids in S. fuliginis ATCC 27551 by performing rescue cloning. After in vitro transposition using the minitransposon Tn5 <R6Kyori-Kan2>, a pPDL2 derivative (pPDL2-K) having a single minitransposon insertion was identified by colony PCR using opd gene-specific primers.

The complete nucleotide sequence of 37,317 bp plasmid pPDL2 is deposited in GenBank (accession no. JX31-2671). Its GC content (62.37%) is very similar to the GC content of indigenous plasmids pCHQ1 (63%), pUT1 (63.7%), and pUT2 (61.0%) found in S. japonicum (Nagata et al. 2010, 2011). The GC content, GC skew, and the length of the plasmid indicates multiple changes in the sequence of pPDL2 either due to acquisition of foreign DNA or rearrangement of its own sequences (Figure 1). A total of 48 protein-coding regions were annotated, which represents a coding density of 91.3% (Table S2). Out of the 48 orfs, 18 of them were annotated as ORFs coding for hypothetical proteins, and 7 of them code for either transposases or associated resolvases (Table S2). Based on comparison with database entries, the ORFs coded by plasmid pPDL2 can be divided into a number of functional modules. Prominent among them are modules for replication and partition, mobilization, integration, and OP degradation, as well as mobile genetic elements.

#### **Replication initiation proteins**

In pPDL2, two ORFs, orf22 and orf45, show high homology to replication initiator proteins. The protein coded by orf22 (367aa) shows absolute identity to the replication initiation protein RepA of plasmid pUT1 of S. japonicum pUT26S (Nagata et al. 2010). It contains a sequence that shows similarity with the well-defined RPA and Rep-3 domains. The proteins having these conserved domains have been shown to play a key role in replication initiation of plasmids (Wickner et al. 1991). As in plasmid pUT1, in the upstream of repA, an oriV sequence was identified with characteristic iterons, an AT-rich sequence, and a dnaA box (Figure 2). Iterons play a key role both in initiation and regulation of replication in theta-replicating plasmids (del Solar et al. 1997; Nordstrom 1990). There is a significant similarity among the oriV sequences found in plasmids pPDL2, pUT1, and pPS10 (Figure 2). If the structural conservation of both oriV and RepA is taken into consideration, they qualify to have all the potential to replicate plasmid pPDL2 (Figure 2). To gain further evidence on their role in pPDL2 replication, we generated a mini-replicon by ligating

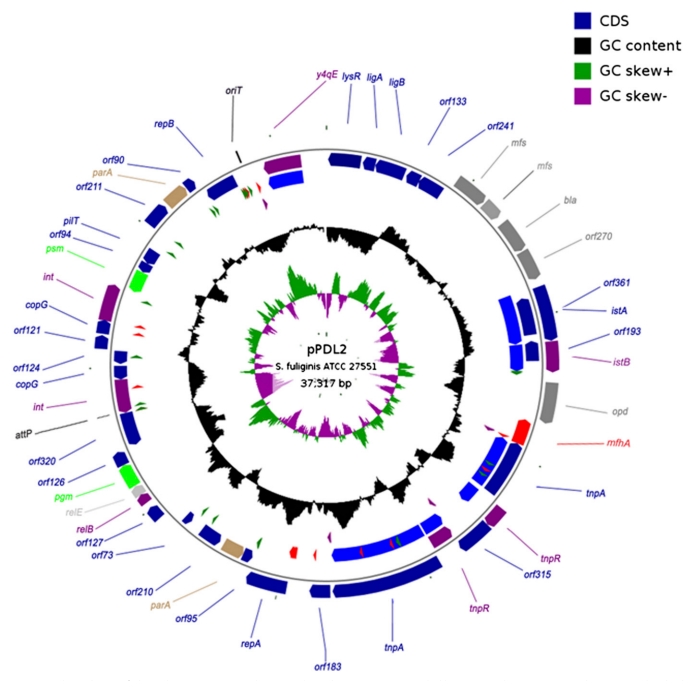
the repA oriV region (18,815-20,914) to a tetracycline-resistant gene. When electroporated into Sphingobium fuliginis ATCC 27551, the resultant mini-replicon (pSFT2) gave tetracycline-resistant colonies, indicating the predicted oriV and repA contribution for the replication of pPDL2. Further, the generated mini-replicon remained in the cells even after growing them in the absence of antibiotic for several generations. Existence of the mini-replicon was also evident in plasmid preparations made from the transformed cells (data not shown). The RepA sequences of theta-replicating plasmids have been showed to follow a characteristic phylogenetic pattern and, to some extent, serve as signatures when assigning incompatibility groups to the plasmids (del Solar et al. 1997; Suzuki et al. 2010). Therefore, the RepA sequence of plasmid pPDL2 was aligned with the available RepA sequences in the GenBank (Suzuki et al. 2010). RepA of pPDL2 has more than 50% identity covering the complete sequence of RepA sequences of the IncP group of plasmids, pADP-1, pEST4011, and pUB1 (Suzuki et al. 2010). Such similarity is not seen with RepA sequences of other incompatibility groups of plasmids, suggesting that pPDL2 could be an IncP group of plasmid.

The protein coded by orf24 of pPDL2 has 100% sequence coverage and identity with the partitioning protein ParA of plasmid pUT1 of S. japonicum UT26S. ParA, in association with ParB, plays a major role in plasmid segregation. However, no parB homolog is found in pPDL2. In the absence of parB, pPDL2 may depend on a ParB protein encoded either by the chromosome or by other indigenous plasmids present in S. fuliginis ATCC 27551. Interestingly, a well-defined plasmid addiction module was identified in pPDL2. The ORFs orf29 and orf30 code for proteins showing more than 95% identity to the RelB/ RelE toxin/antitoxin system found in a variety of Gram-negative bacteria and archaea (Gotfredsen and Gerdes 1998; Gronlund and Gerdes 1999). The RelE and RelB pair of E. coli is well characterized. The 11kD RelE protein acts as toxin to bacterial cells, and a similarly sized relB product counteracts the toxic effect of RelE (Gotfredsen and Gerdes 1998; Gronlund and Gerdes 1999). Presence of partition protein ParA, along with a post-segregational killing mechanism (RelE/ RelB), suggests faithful partitioning and maintenance of pPLD2 in daughter cells of Sphingobium fuliginis ATCC 27551.

## Plasmid pPDL2 is a mobilizable plasmid

Since oriT is an essential element for mediating horizontal gene transfer we searched for a similar sequence in pPDL2. A number of functionally validated oriT sequences have been shown to contain typical secondary structures in a characteristic GC rich region (Lee and Grossman 2007) and such an oriT-like sequence (34823-35011) was identified in pPDL2. The predicted oriT has very high similarity to an oriT sequence of a self-transmissible plasmid pLB1 of *S. japonicum* UT26S (Miyazaki *et al.* 2006). If the location of oriT is taken into consideration, along with the presence of four tandem repeats immediately downstream, the predicted sequence has all features of a functional oriT (Figure 3A). Interestingly the putative oriT sequence is found upstream of an ORF (orf45) that codes for a protein showing high similarity (98%) to the plasmid-borne RepB sequences of *Sphingobiaceae* members (Bertalan *et al.* 2009).

The predicted RepB sequence includes a Rep-3 domain, involved in typical nicking-closing like activity (Dyda and Hickman 2003; Moscoso *et al.* 1995). Proteins having a Rep-3 domain also perform strand-transfer reactions generally seen during rolling circle (RC) replication (Moscoso *et al.* 1995; Ruiz-Maso *et al.* 2007). RC replication is also seen during horizontal transfer of indigenous plasmids. However, it is initiated from a well-defined *oriT* sequence (Pansegrau and Lanka

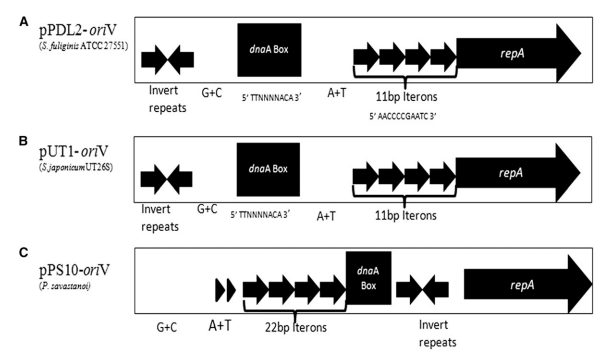


**Figure 1** Physical map of plasmid pPDL2. Outer and inner circles indicate proteins encoded by sense and anti-sense strands, respectively. Third circle indicates mobile elements and repeat sequences. Direct (DR) and Inverted (IR) repeats are shown with filled red and green triangles, respectively. Tn3-specific repeats appear with filled purple triangles in the fourth circle. The fifth circle shows GC content across the plasmid sequence. The sixth and seventh circles represent GC-skew in sense and anti-sense strands.

1991). Since we have identified both *oriT* and RepB in the sequence of pPDL2 we have attempted to validate their involvement in the lateral transfer of pPDL2. We initially assessed whether pPDL2 is a mobilizable plasmid by performing triparental mating experiments, involving *E. coli pir*-116 (pPDL2-K) as donor, with *Acinetobacter* sp. DS002 and *E. coli* (pRK2013) as recipient and helper strains respectively. Kanamycin-resistent exconjugants of *Acinetobacter* sp. DS002 have appeared on a selection plate at a frequency of  $3.4x10^{-5}$  per recipient. As expected, all exconjugants carried the *opd* gene and showed OPH activity, properties not found in *Acinetobacter* sp. DS002 (Figure 3D).

Mobility of plasmid pPDL2 was also observed when *E. coli* BL21 was used as recipient. Though pPDL2 is stably replicated in *Acinetobacter* sp. DS002, it failed to replicate in an *E. coli pir* negative background. Although kanamycin resistant *E. coli* colonies were found, they subsequently lost the plasmid. In certain colonies that retained the ability to grow on kanamycin we could not detect plasmid from these cells suggesting that either complete pPDL2-K or the region containing kanamycin gene has probably integrated into the genome.

To understand RepB and *oriT* interactions better we performed mobility shift assays by incubating the PCR amplicon carrying *oriT* 



**Figure 2** Replication module. Comparison of replicative origin (*oriV*) of plasmid pPDL2 (panel A) with *oriV* sequences of plasmids pUT1 (panel B) and pPS10 (panel C). Solid inverted arrows indicate inverted repeat sequences. The *dnaA* Box and iterons are shown with solid box and tandem arrows, respectively. The transcription orientation of *repA* is shown with a solid arrow. Filled triangles indicate repeats found in the A+T-rich region of pPS10.

with purified RepB<sub>C-6His</sub> (Figure 3B). There was a RepB<sub>C-6His</sub> dependent shift in the electrophoretic mobility of oriT (Figure 3C) and no shift in presence of a BSA control, indicating specific interactions between RepB<sub>C-6His</sub> and oriT. Supporting this observation the intensity of shifted oriT band got proportionately reduced when increased concentrations of cold oriT was added to the reaction mixture (data not shown). To assess if RepB might function as a relaxase (Mob) the sequence of RepB was aligned with other relaxase sequences (Achtman 2011; Francia et al. 2004; Garcillan-Barcia et al. 2009; Guglielmini et al. 2011) and a weak homology was observed with relaxase of the MOB<sub>F</sub> plasmid. The role of RepB in lateral transfer of pPDL2, therefore needs to be further investigated. Despite not detecting tra genes in the sequence of pPDL2, experiments were also performed to assess whether pPDL2 is self-transmissible but no exconjugants were observed from a biparental mating using E. coli pir-116 (pPDL2-K) as donor and Acinetobacter sp. DS002 as recipient. Hence, in summary, our data clearly indicate that pPDL2 is a mobilizable but not a selftransmissible plasmid.

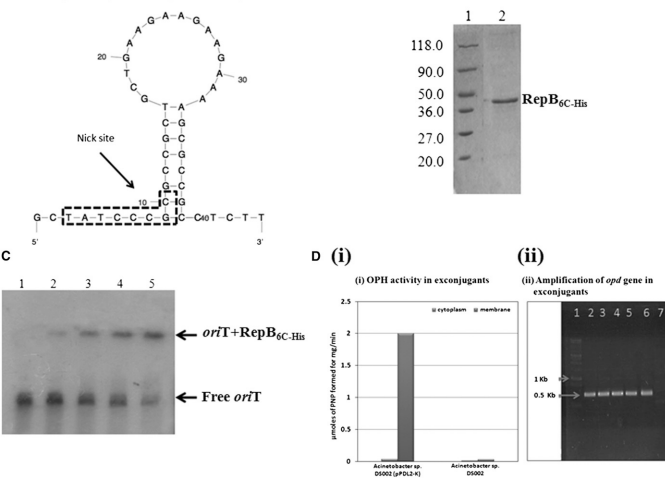
#### Integration module

An interesting observation in the sequence of pPDL2 is the existence of an uniquely organized integrase module (nucleotide position 26697 to 30249). The module contains two units, CIP-I and CIP-II, each consisting of *copG*, *int* and *pgm* genes and having divergent transcription orientation (Figure 4A). The integrase of CIP-I is a 328 amino acid protein with strong identity (86%) to a plasmid pUT1-borne phage integrase of *S. japonicum* UT26 and 70% identity to a sitespecific recombinase of *Pseudomonas syringae* DC 3000. The CopG/ MetJ/Arc family regulatory protein (subsequently referred to as CopG) coded by pPDL2 shows highest similarity to the CopG protein coded by plasmid pUT1 of *S. japonicum* UT26S. The 131 residue protein shows homology to a number of other proteins belonging to CopG/MetJ/Arc family of transcriptional regulators that can act both as transcriptional repressors and activators (del Solar *et al.* 1995; del Solar and Espinosa 1992; Gomis-Ruth *et al.* 1998; He *et al.* 2002). In a well-characterized Streptococcal plasmid, pMV185, CopG has been shown to act as a repressor of *repB* by binding to a pseudosymmetirc region overlapping the -35 sequence of the  $\sigma^{70}$ dependent promoter, thereby preventing transcription of the *copG*, *repB* operon in plasmid pMV158 (Costa *et al.* 2001).

In pPDL2, the *copG* and *int* genes appear to be organized as one operon in both CIP-I and CIP-II units. A potential  $\sigma^{70}$ -dependent promoter was only seen upstream of each of the *copG* sequences, and the stop codon of *copG* overlaps with the start codon of the *int* genes suggesting that these two genes are cotranscriptional. Due to the existence of opposite transcriptional orientation, an overlap of 5bp is seen between the putative copG1 and copG2 promoters (Figure 4A). As CopG1 proteins are known to act as transcription repressors, the intergenic region found between the copG1, int1 and copG2, int2 operons was analyzed to identify *cis*-elements that can serve as target sites for CopG proteins (Figure 4A). In studies conducted by del Solar and his associates, CopG is shown to bind to a pseudosymmetric ciselement present overlapping -35 region (del Solar et al. 2002). Inverted repeats are also shown to be putative binding sites of CopG in Sulfolobus neozealandicus (Greve et al. 2005). Interestingly, in both CIP-I and CIP-II units, typical CopG binding sites were predicted upstream of the start codon of copG and putative promoter sequence (Figure 4A). Transcriptional organization of copG and *int* genes and the existence of such CopG binding sites, clearly indicates involvement of CopG in regulation of expression of the integrase in S. fuliginisATCC 27551.

In addition to these two transcriptionally coupled integrase and copG genes an additional ORF that codes for a protein showing identity (98%) to phosphoglycerate mutase of *S. japonicum* UT26S is

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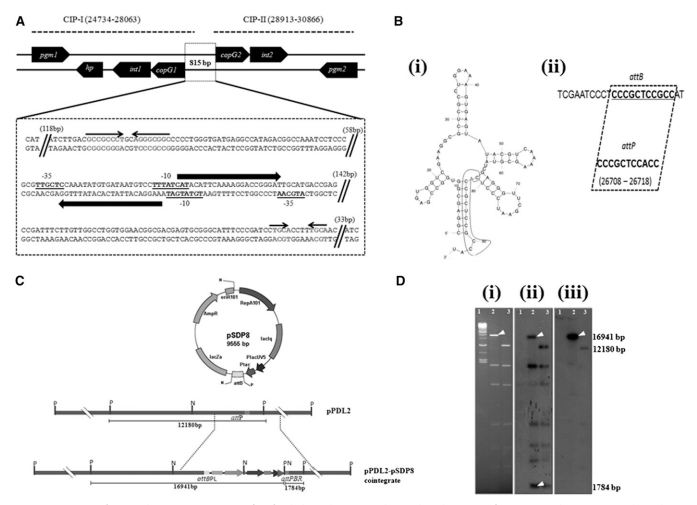


**Figure 3** Mobilization of pPDL2. Panel A represents secondary structure of predicted *ori*T sequence. The nick site that matches perfectly with the nick site of plasmid pLB1 of *Sphingobium japonicum* is shown in a dotted box. Affinity purification of  $\text{RepB}_{C-6His}$  is shown in panel B. Panel C indicates the mobility shift assay done using radiolabeled *ori*T fragment and purified  $\text{RepB}_{C-6His}$ . Lane 1 represents <sup>32</sup>P labeled *ori*T without  $\text{RepB}_{C-6His}$ . Lanes 2–5 represent labeled *ori*T incubated with increased concentrations [0.1 µg (2), 0.25 µg (3), 0.5 µg (4), and 1 µg (5)] of  $\text{RepB}_{C-6His}$ . Arrows indicate either free *ori*T or *ori*T-RepB<sub>C-6His</sub> complex. OPH activity (i) and amplification of *opd* gene (ii) in exconjugants are shown in panel D.

present at the 3' end of the *int* gene in each of the two units of the integrase modules (Figure 4A). Though, the phosphoglycerate mutase gene in CIP-I module is found 1.1 Kb away from the stop codon of integrase, in CIP-II, it is found immediately downstream of transcription terminator sequence of int2 gene. This 206 residue protein shows high identity (98%) to phosphoglycerate mutase encoded on plasmid pUT1 of S. japonicum UT26S. Phosphoglycerate mutases (PGMs) are very well characterized group of enzymes found both in prokaryotes and eukaryotes. PGMs are basically transferases involved in transfer of phosphate group (Parkinson and Kofoid 1992). Conversion of glyceraldehyde 3-phosphate to glyceraldehyde 2-phosphate, by transfer of phosphates from the 3<sup>rd</sup> position to the 2<sup>nd</sup> position, is the classical biochemical reaction catalyzed by this group of enzymes. However, PGMs containing an HP-PGM-like domain are also known to be involved in signal transduction process (Matsubara and Mizuno 2000). The histidine present in the catalytic site of this group of enzymes undergoes phosphorylation during the signal relay process (Parkinson and Kofoid 1992). The strong link between PGM, CopG and integrase suggests a related function of these proteins, and a similar organization is also seen in S. japonicum UT26S plasmids pCHQ1 (NCBI Reference no: NC014007) and pUT1 (NCBI Reference no:

NC014005), and *Sphingomonas wittichi* RW1 (pSWIT01) adding strength to the proposed hypothesis.

To assess whether the CIP-I and CIP-II units were generated by gene duplication we compared the sequences of these two units. If they are products of gene duplication there should have been absolute identity between the proteins coded by these two units of the integrase module. However there is only 67% identity between these two proteins. The N-terminal region of CopG2 encoded by CIP-II unit was found to be much longer. In addition to the size difference, there was no sequence conservation at the C-terminus of the protein. Nevertheless, the central regions of CopG1 and CopG2 are almost identical. Such diversity in the primary sequence suggests a divergent origin of CopG1 and CopG2. A similar situation was seen when int1 and pgm1 were aligned with int2 and pgm2. Between int1 and int2 only 80% identity was observed although this identity continued throughout the sequence. The percent identity between PGM1 and PGM2 is high (91%), the difference only being seen at the C-terminus and N-terminus. Considering the diversity in the primary sequence of the proteins coded by CIP-I and CIP-II of the integrase module, the plasmid pPDL2 is proposed to have acquired the CIP-I and CIP-II integrase modules from independent sources, possibly through a unique recombination process.



**Figure 4** Site-specific recombination. Integration of artificial plasmid pSDP8 at the predicted *attP* site of pPDL2. Panel A represents physical map of integration module. The underlined sequences indicate putative promoter elements. The CopG binding sites are shown with inverted arrows. The transcription orientation of integration modules I and II are shown with solid arrows. The synthetic *attB* site created using sequence of *Sphingobium japonicum* UT26S and the similarities between predicted pPDL2 *attP* and *attB* sequences are shown in (i) and (ii) of panel B, respectively. The physical map of pSDP8 and site of integration of pSDP8 in pPDL2 is shown in panel C. The restriction profile of the cointegrate (i) and the corresponding southern blot developed using either labeled pPDL2 (ii) or pSDP8 (iii) are shown in panel D. The increase in size of 4.7 kb *Pstl* fragment due to integration of a 6.4 Kb plasmid pSD8 is shown with an arrow.

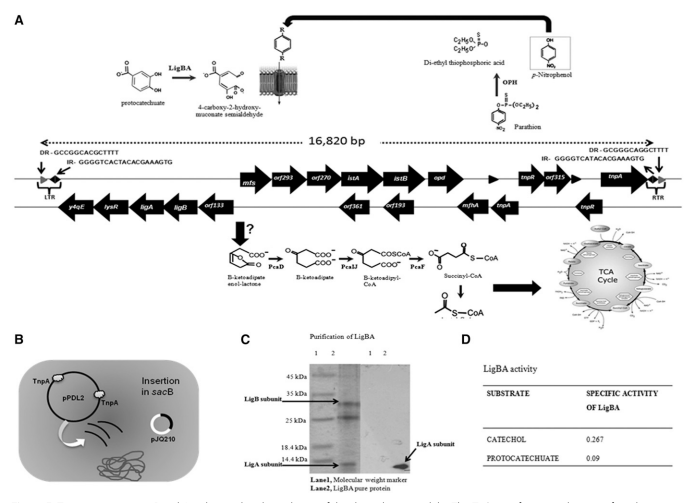
#### Plasmid pPDL2 is an integrative mobilizable element

Having identified an integrase module, we performed experiments to determine whether site-specific integration of pPDL2 takes place at an artificially created attB site cloned in temperature-sensitive plasmid pDSP8. Initially we performed bioinformatic studies to identify attP sites on the plasmid pPDL2. While perfoming these predictions, we took tRNA genes from the genome sequence of S. japonicum UT26S, whose total genome sequence is known (Nagata et al. 2010). The reasons for taking tRNA genes from S. japonicum are that parts of pPDL2 show considerable homology with its indigenous plasmid pUT1 and that it encodes an integrase module very similar to the one present in pPDL2. These tRNA gene sequences were then blasted with the total sequence of pPDL2 to obtain a library of identical sequences with a minimum length of 10 bp. Among these sequences, the ones found in intergenic regions were predicted to be potential attP sites. The servl tRNA gene of S. japonicum has sequence identity with the predicted attP at its 3' end and served as an attB sequence [Figure 4B, part (i)]. E. coli pir-116 strains carrying pPDL2-K and pSDP8 were grown independently for 12 hr at 30° and 37°, and then

plated on kanamycin and ampicillin plates. A lawn of cells were seen on plates seeded with the  $30^{\circ}$ -grown culture, whereas only a few colonies appeared in plates seeded with the  $37^{\circ}$ -grown culture. In the absence of *attB*, no colonies appeared, suggesting that the generated colonies were due to occurrence of site-specific recombination between *attB* and *attP* sites [Figure 4B, part (ii)]. The restriction profile of plamids isolated from randomly picked colonies showed integration of pSDP8 in a 12.0 kb *PstI* fragment of pPDL2 (Figure 4, C and D), and sequence analysis revealed *attPBL* and *attBPR* sequences, suggesting that the *attP* site predicted at the 3' end of *int1* served as an *attP* site in pPDL2.

#### Degradation module

In our previous studies, we analyzed the 7 kb DNA region of pPDL2 that shows sequence identity with the pCMS1-borne *opd* cluster from *B. diminuta*, and we identified a transposon-like organization of the *opd* gene cluster in *S. fuliginis* ATCC 27551 (Siddavattam *et al.* 2003). Such structural organization and the existence of identical *opd* clusters in bacterial strains representing diverse taxonomic groups suggested



**Figure 5** Transposition assay. Panel A indicates the physical map of the degradative module. The Tn3-specific terminal repeats found upstream and downstream of transposable element Tn3 and *y4qE* are shown with arrows. The reactions catalyzed by OPH, LigBA are shown. The transposon assay performed to assess transposition of the degradative module is shown in panel B. Panel C indicates affinity purification of LigBA and the corresponding Western blot using anti-His antibody. The dioxygenase activity of purified dioxygenase (LigBA) with various aromatic compounds is shown in panel D.

horizontal transfer of the *opd* cluster through transposition (Figure S1). However, in our previous studies, we were unable to detect any transposition events, and we concluded that it is a defective transposon due to the existence of a truncated *tnpA* gene in the Tn3 element identified at the flanking end of the *opd* cluster.

The full sequence of pPDL2 shows the existence of second copy of Tn3 immediately downstream of the defective Tn3 element, and so we have revisited the question of whether the opd cluster of pPDL2 functions as an active transposon. Bioinformatic analysis identified Tn3-specific terminal repeats (Siguier et al. 2006) in pPDL2: two at the terminal ends of Tn3 and a third at the upstream region of transposon *y4qE* (Figure 5A). The genetic map of the DNA region between y4qE and Tn3 includes the previously reported opd gene clusters (Siddavattam et al. 2003), transposons Tn3 and y4qE, and the ligBA operon coding for the protocatechuate-4,5-dioxygenase alpha and beta subunits, along with a gene (orf1) coding for a LysR-type transcription regulator. The ligBA operon codes for protocatechuate-4,5-dioxygenase alpha and beta subunits showing 83% identity to the similar proteins of Xanthomonas campestris pv. campestris str. ATCC 33913 (NCBI reference no. NP\_636196.1). While evaluating the functional status of lig operon, an expression plasmid (pSDP9) was constructed by placing it under the control of vector-specific transcriptional and translational signals. The resting cells prepared from induced cultures of E. coli BL21 (pDSP9) successfully generated ring-cleavage products from proctocatechuate, catechol, and benzenetriol indicating the operon codes for a functional dioxygenase. As the cloning strategy translationally fuses ligA with vector sequences coding for a His-tag, the E., coli BL21 (pDSP9) cells encode a LigA<sub>C-6His</sub> protein. Affinity purification yielded both LigA<sub>C-6His</sub> and LigB in approximately equimolar concentration, revealing that these two functionally related proteins form a complex (Figure 5A). Supporting the resting cell assay, the affinity-purified dioxygenase also generated ring-cleavage products from both protocatechuate and catechols (Figure 5D). In addition to LigB and LigA<sub>C-6His</sub> proteins, an additional band with a size of 25 kD was enriched during affinity purification (Figure 5C). It is not clear whether this 25 KDa protein is copurified due to specific interactions with Lig proteins. Protocatechuate-4,5-dioxygenase is the key enzyme in the benzoate (hydroxylation pathway) and 2,4-dichlorobenzoate degradation pathways (Adriaens et al. 1989). In addition to these two compounds, many aromatic compounds such as vanillate, isovanillate, phthalates, and benzoate derivatives are shown to be channelled

through the protocatechuate degradation pathway (Providenti *et al.* 2006). When OP insecticides, like methyl parathion and parathion, are hydrolyzed by OPH, aromatic compounds such as *p*-nitroarophenols are generated (Sethunathan and Yoshida 1973). These compounds are found to be more toxic to the microflora than the parent compounds (US Environmental Protection Agency 2007). The existence of a ringcleavage dioxygenase as part of the *opd* gene cluster allows channeling of these aromatic byproducts of OP insecticides into the TCA cycle, contributing to their complete mineralization.

In addition to the *lig* operon, *lysR* and two ORFs showing homology to transporter proteins belonging to the major facilitator superfamily were identified in the upstream and downstream region of the *lig* operon (Figure 5A). Transporters belonging to the major facilitator superfamily proteins are known to be involved in transport of aromatic proteins (Pao *et al.* 1998). Hence, in summary, if the sequence found between the Tn3-specific terminal repeats is a transposable unit, it appears to have all necessary information for mineralization of OP insecticides like parathion and methyl parathion.

#### The opd cluster of pPDL2 is an active transposon

The transposition assay that we described in our previous study was repeated by replacing pMS3 with the pPDL2-K derivative pPDL2-KT in which opd is replaced with opd::tet and hence the event of transposition can be monitored by the tetracycline-resistant phenotype. Plasmid pPDL2-KT requires pir protein for its replication and hence replicates only in E. coli pir-116 cells. When pPDL2-KT was electroporated into E. coli (pJQ210) cells, we found colonies resistant to sucrose and tetacycline, indicating disruption of the sacB gene of pJQ210 due to transposition of opd::tet. Further analysis of plasmids isolated from the sucrose/gentamycin/tetracyclin resistant colonies gave a typical restriction profile suggesting existence of opd cluster in the sacB region of pJQ210. The junctions of the opd cluster were then amplified using sacB-specific primers and primers designed for the region upstream of the predicted terminal repeats. The sequence of the PCR amplicons confirmed the existence of Tn3-specific terminal repeats, indicating that the opd cluster of pPDL2 is an active transposon (Figure 5B).

#### DISCUSSION

Horizontal gene transfer in bacteria is an important driving force that plays a critical role in generating genetic variation among bacteria and contributes to shaping of their genomes (Hacker and Carniel 2001; Jain et al. 1999, 2003; Preston et al. 1998; Skippington and Ragan 2011). DNA acquisition by bacteria through various mobile genetic elements, such as plasmids, bacteriophages, IS elements, transposons, integrons, and conjugative transposons, is an established phenomenon (Dobrindt et al. 2004; Garcillan-Barcia et al. 2011; Wiedenbeck and Cohan 2011). Sequences associated with mobile genetic elements differ from the host genome with respect to their GC composition and GC skew, serving as unique makers to assess lateral acquisition. Such delineation represents an island-like existence for mobile genetic elements in the genome of the host (Juhas et al. 2009). Depending on the nature of the genetic information present on these mobile genetic elements, they are designated as pathogenic islands (Dobrindt et al. 2004; Schmidt and Hensel 2004), resistant islands (Larbig et al. 2002), or catabolic islands (Gaillard et al. 2006). Involvement of plasmids and catabolic transposons in lateral transfer of degradative genes is a recognized phenomenon (Topp et al. 2000; Tsuda et al. 1999; van der Meer et al. 2001). Recently genes required for 3-chlorobenzoate degradation have been shown to be part of an integrative conjugative element (ICE). This ICE element, found downstream of a glycyl tRNA gene, contains all the necessary genes for mineralization of 3-chlorobenzoate along

with an *int* gene coding for a site-specific integrase and attachment sites (Gaillard *et al.* 2006).

Plasmid pPDL2 is unique in number of ways. It contains a welldefined replication module with an oriV sequence and a repA gene to support its replication. A mini-replicon generated by ligating the oriV, *repA* region of pPDL2 with an insertionally inactivated *opd* (*opd::tet*) gene successfully replicated in Acinetobacter sp. DS002, suggesting its involvement in replication of pPDL2. The pPDL2 also has an integrase module, consisting of copG, int, and pgm genes (Figure 4A). The organization of the integrase module is similar to the one present in plasmid pUT1 of S. japonicum UT26S, although pUT1 has a single integrase module, whereas pPDL2 has two modules. The data presented in the present study clearly show that pPDL2 can successfully integrate if a suitable attB site is available in the host chromosomal DNA. Plasmid pPDL2 contains an oriT sequence that is identical to the oriT sequence of pLB1 of S. japonicum UT26. In support of this observation, the pPDL2 derivative pPDL2-K was mobilized into Acinetobacter and E. coli cells in the presence of a helper strain E. coli HB101 (pRK2013). The data obtained in the present study clearly show formation of cointegrates between pPDL2-K and pSDP8 due to site-specific recombination. If a suitable attB site is present, such an event is possible between pPDL2-K and the E. coli chromosome. If this property of pPDL2 is taken together with its ability to mediate its own mobilization, designating pPDL2 as an IME appears to be logical. In general, lateral mobility of Sphingobium plasmids is only seen among Sphingomonadaceae members (Basta et al. 2004). However, the self-transmissible Sphingobium plasmid pLB1 derivative pLB1-K has been successfully transmitted from S. japonicum to Sinorhizobium meliloti and Mesorhizobium loti (Miyazaki et al. 2006). Like pLB1, plasmid pPDL2 derivative pPDL2-K can be successfully mobilized from E. coli to Acinetobacter that is not a member of the Sphingomonadaceae. This property of pPDL2 facilitates lateral transfer of the opd cluster among a wide range of bacterial strains. When such lateral transfer takes place into a host where pPDL2 replication is not supported, it can integrate into the genome of the recipient cells due to the existence of an integrase module. This additional feature appears to ensure retention of genetic information evolved for degradation of organophosphates, even in a host that does not permit replication of pPDL2.

Furthermore, pPDL2 contains a catabolic transposon that contains information for hydrolyzing a structurally diverse group of organophosphates and then for degrading aromatic compounds generated due to OPH-mediated hydrolytic cleavage of OP insecticides. As shown in Figure 5A, the entire degradative module appears to be organized as a complex transposon having transposons y4qE and Tn3 at the flanking ends of degradative modules. Tn3-specific terminal repeats were found upstream and downstream of y4qE and Tn3, and consistent with this sequence information, we demonstrated a transposition event that inserted the entire degradative module into the sacB gene of a reporter plasmid. This event suggests the possibility of spreading opd as part of a large catabolic transposon. Surprisingly, most of the genetic information found on 37,317 kb pPDL2 appears to be related either to lateral gene transfer or degradation of organophosphates. It appears that the plasmid is tailor-made through a number of recombination and transposition events for spreading opd information among soil bacteria. The presence of identical opd regions on plasmids pCMS1 and pPDL2 and the existence of an active transposable element with opdA is in itself evidence for acquisition of opd through lateral gene transfer (Mulbry et al. 1987; Horne et al. 2002, 2003; Pandeeti et al. 2011). In the Sphingobiaceae family, only two members, S. fuliginis ATCC 27551 and Sphingomonas sp. JK1, have *opd* genes (NCBI reference no. ACD85809). The existence organophosphates in the soil might have favored the *opd*-containing bacteria to acquire DNA that encodes transporters and aromatic degradation genes as it allows the complete utilization of organophosphates.

Considering the experimental data presented in this study, we have searched genome sequences of Sphingobiaceae members to determine whether sequences that match pPDL2 exist in other genomes. Synteny maps were drawn to show the extent of similarity between pPDL2 and genome sequences of Sphingobiaceae members (S. japonicum and S. wittichii). Plasmid pUT1 of S. japonicum has organizational similarity with pPDL2 in the region containing replication and integrase genes (see Figure S2). If synteny maps are taken into consideration along with GC content and GC skew, it clearly suggests that plasmid pPDL2 evolved by a number of transposition and rare recombination events to integrate DNA from different sources. Although pPDL2 has all the necessary features for spreading opd information, due to the lack of tra genes, it depends on helper plasmids for achieving lateral transfer. However, as Sphingobium fuliginis ATCC 27551 has three more plasmids, complementation of such a function by these plasmids cannot be ruled out.

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