Biodegradation of endocrine disruptor dibutyl phthalate (DBP) by a newly isolated Methylobacillus sp. V29b and the DBP degradation pathway

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Abstract Bacteria of the genus Methylobacillus are methanotrophs, a metabolic feature that is widespread in the phylum Proteobacteria. The study demonstrates the isolation and characterization of a newly isolated Methylobacillus sp. V29b, which grows on methanol, protocatechuate, monobutyl phthalate, dibutyl phthalate, diethyl phthalate, benzyl butyl phthalate, dioctyl phthalate and diisodecyl phthalate. Methylobacillus sp. V29b was characterized with scanning electron microscopy, transmission electron microscopy, Gram staining, antibiotics sensitivity tests and biochemical characterization. It degrades 70 % of the initial DBP in minimal salt medium and 65 % of the initial DBP in samples contaminated with DBP. DBP biodegradation kinetics was explained by the Monod growth inhibition model. Values for maximum specific growth rate (μmax) and half-velocity constant (Ks) are 0.07 h⁻¹ and 998.2 mg/l, respectively. Stoichiometry for DBP degradation was calculated for Methylobacillus sp. V29b. Four metabolic intermediates, dibutyl phthalate (DBP), monobutyl phthalate, phthalic acid and pyrocatechol, were identified. Based on the metabolic intermediates identified, a chemical pathway for DBP degradation was proposed. Six genes for phthalic acid degradation were identified from the genome of Methylobacillus sp. V29b.

Keywords Endocrine disruptor · Degradation kinetics · Stoichiometry · Gene identification · Phthalate ester degradation pathway

Introduction

Phthalic acid esters (PAEs) are a class of compounds widely used as plasticizers to provide mechanical strength and flexibility to the resins (Cartwright et al. 2000; Staples et al. 1997). They are ingredients of paints, adhesives, house-building materials, defoaming agents, PVC pipes, food packing materials, toys, plastics, solubilizers of cosmetic products, medical devices, photography films, textile fabrics, pesticide carriers, lubricating oils and are used in aerospace technology (Gesler 1973; Gross and Colony 1973; Hauser et al. 2007; Ito et al. 2005; Krauskopf 1973; Marcel 1973; Teil et al. 2006; Tepper 1973; Wang et al. 1995b; Wilkinson and Lamb 1999). They have low solubility in water; therefore, they are stable in the environment over a longer period of time (Huang et al. 2008; Vikelsøe et al. 2002; Wang et al. 2008; Yuan et al. 2002). They have been detected in various environments including landfill leachates (Schwarzbauer et al. 2002; Zheng et al. 2007), air (Wensing et al. 2005), soils, sediments, and natural waters (Staples et al. 1997). PAEs have been classified as top priority toxicants by the European Union, China National Environmental Monitoring Center and US Environment Protection Agency (Chen et al. 2007; Council of the European Union 1993). PAEs are responsible for carcinogenicity and endocrine disruption (Colborn et al. 1993; David et al. 1999; Jobling et al. 1995; Piersma et al. 2000). They can elicit cellular estrogenic responses and suppress calcium signaling (Kaun-Yu et al. 2004; Lovekamp-Swan and Davis 2003; Moore 2000). They are responsible for hypospadias, cryptorchidism and malformation of the reproductive tract in mice (Fisher 2004; Gray et al. 2000; Jaeger and Rubin 1970; Li et al. 1998; Zhu et al. 2006). They are known for irritation of eyes, skin, respiratory tract, blurred vision and induce stone formation in the
bladder (Dai et al. 2005a, b; Wang and Gu 2006). Dibutyl phthalate (DBP) is the most widely used plasticizer and has been detected in different environments (Eaton 2001; Feiler 1980; Hashizume et al. 2002; Keith and Telliard 1979; Xu et al. 2005).

The natural processes of degradation such as hydrolysis and photodecomposition are not efficient in the degradation of these pollutants (Lu et al. 2009). Therefore, microbial degradation is the major route for their degradation (Staples et al. 1997; Vamsee-Krishna et al. 2006; Vamsee-Krishna and Phale 2008). Bacteria having potential to degrade PAEs have been isolated from various environments including mangrove sediments, activated sludge and wastewater (Liang et al. 2008). Aerobic degradation of PAEs is much more efficient as compared to anaerobic degradation (Cheung et al. 2007; Fang et al. 2010). Sequential hydrolysis of PAEs has been demonstrated by a few researchers (Engelhardt and Wallnöfer 1978; Jiao et al. 2013; Staples et al. 1997).

Despite research on degradation of PAEs by various researchers, these studies lack in perspectives such as efficient DBP degradation at higher concentrations, elucidation of DBP degradation pathway, kinetics of DBP degradation and identification of genes responsible for PAEs degradation. Extensive research in the above aspects is required to remove these pollutants from the environment. The aim of the study was isolation, characterization and identification of efficient DBP-degrading bacterial strain from municipal solid waste (MSW) leachate and to examine the degradation potential of the isolate toward degradation of DBP in both minimal media and in PAEs-contaminated samples collected from the landfill site.

**Materials and methods**

**Chemicals**

HPLC-grade monobutyl phthalate, diisodecyl phthalate, dioctyl phthalate, protocatechuate, benzyl butyl phthalate, diethyl phthalate and dibutyl phthalate were purchased from Sigma-Aldrich (USA) and used as substrate for growth of the bacteria. HPLC-grade acetonitrile purchased from Sigma-Aldrich (USA) was used as solvent in the analysis of DBP.

**Isolation and characterization of DBP-degrading bacteria**

Municipal solid waste (MSW) leachate samples were collected from a municipal solid waste landfill site at Ghazipur, New Delhi, India. The location co-ordinates of Ghazipur landfill site are 28° 37’ 22.4”N and 77° 19’ 25.7”E. The physical parameters of the site are: pH 8.4, TDS 29,700, COD 31,600, Fe 9.81 and Cl 1174.2. After collection, the samples were stored at 4 °C. The MSW leachate was inoculated in minimal salt medium (MSM) supplemented with DBP [DBP emulsified with 0.1 % (vol/vol) Tween 80] as the sole carbon and energy source. MSM was prepared by dissolving 3.5 g K2HPO4, 1.5 g KH2PO4, 0.27 g MgSO4, 1 g NH4Cl, 0.03 g Fe2(SO4)3, 7H2O and 0.03 g CaCl2 in 1 L distilled water. The pH of MSM was adjusted to 6.8 and sterilized by autoclaving at 121 °C and 15 psi for 20 min. A separate iron sulfate and magnesium sulfate solution was prepared, filter sterilized with 0.22 μm membrane filter and added to MSM to avoid precipitation formation (Vega and Bastide 2003). For isolation of DBP-degrading bacteria, 1 ml MSW leachate was inoculated to 100 ml MSM supplemented with 10 mg/l of DBP and incubated at 30 °C and 180 rpm. The amount of DBP was increased in subsequent transfer cultures. Culture from the flask was streaked on MSM-agar plates to obtain a pure culture. A colony numbered 29 was able to grow in the presence of DBP and it was designated as strain 29D. Size and cell morphology were observed using scanning electron microscopy (SEM). Internal features of the bacteria were observed using transmission electron microscopy (TEM). SEM and TEM of strain 29D were performed at the Advanced Instrumentation Research Facility (AIRF), Jawaharlal Nehru University, New Delhi. For electron microscopy, bacterial cells were fixed in 3 % glutaraldehyde at room temperature for 3 h. After fixation, bacterial cells were washed with 0.1 M phosphate buffer thrice for 10 min. Post-fixation was performed in 1–2 % osmium tetroxide solution followed by dehydration with increasing concentrations of ethanol in water solution (Smith 1955). Characterization of strain 29D was performed using biochemical tests (Vos et al. 2011). Gram staining reaction was performed using Gram-stain kit from Himedia Lifesciences. Determination of catalase activity was performed by assessment of bubble production in 3 % (v/v) H2O2. Determination of oxidase activity was performed using 1 % (w/v) tetramethyl-p-phenylenediamine from Himedia lifesciences. Hydrolysis of starch was determined by growing bacteria on MSM plates containing 0.2 % (w/v) starch. Susceptibility to antibiotics was determined by spreading the bacterial suspension on MSM plates amended with 10–100 μg/ml of antibiotic tested. To assess the capability of strain 29D to cleave the benzene ring, Rothera’s test was performed. For the test, 5 ml of bacterial culture was saturated with solid ammonium sulfate and mixed with a few drops of 2 % sodium nitroprusside solution and liquor ammonia. The mixture was left for 15 min. A bluish-purple ring indicates the presence of the ketone bodies in it (Rothera 1908).
**16S-rRNA gene amplification, sequencing and phylogenetic analysis**

Identification of strain 29D was performed by 16S-rRNA gene identification. Genomic DNA of the strain 29D was isolated using Fast DNA® SPIN Kit for soil from MP Bio. The 16S-rRNA gene amplification was performed with bacterial universal primers 27F and 1492R (Weisburg et al. 1991). For PCR, a 50 µl reaction was used containing 25 µl PCR master mix (Thermo Scientific), 2 µl forward primer, 2 µl reverse primer, 23 µl sterile water and 1 µl genomic DNA. Time programming used for the thermo cycler (Applied Biosystems GeneAmp PCR system 9700) was 10 min at 95 °C, 35 cycles, 60 s at 95 °C, 90 s at 54 °C, and 60 s at 72 °C and 5 min at 72 °C. The amplified PCR product was gel purified with HiYield™ PCR DNA Mini Kit from Real Genomics™ (Ref catalog no. YDF100). Sequencing of the purified product was performed at the DNA sequencing facility UDSC, University of Delhi, New Delhi, India. Sequencing was performed using dideoxy termination method with bacterial universal primers: 27F (5'-AGAGTTTGATCCTGGCTCAG-3') and 1492R (5'-GGCTACCTTGTTACGACTT-3'). The obtained sequences were combined with Biodeit program. The resultant 16S-rRNA sequence was submitted to NCBI. Phylogenetic analyses were carried out with maximum composite likelihood method (Felsenstein 1981) using MEGA version 5 (Tamura et al. 2011).

**Substrate utilization**

Substrate utilization of strain 29D was performed to examine its ability to grow on different substrates. Six substrates were chosen including methanol, diethyl phthalate (DEP), dioctyl phthalate (DOP), monobutyl phthalate (MBP), diisodecyl phthalate (DIDP), benzyl butyl phthalate (BBP) and protocatechuic acid (PC). Substrate concentration was 2000 mg/l. Growth of strain 29D was measured with Perkin Elmer Lambda 25 UV/vis spectrophotometer at 600 nm.

**Kinetics and stoichiometry of DBP degradation by strain 29D**

For degradation studies, strain 29D was inoculated in MSM and PAEs-contaminated samples and flasks were incubated at 30 °C and 180 revolution per minute in an incubator. Samples for residual DBP analysis and metabolic intermediates identification were collected every 24 h. Collected samples were extracted with ethyl acetate in 1:1 ratio and residues were dissolved in methanol. The obtained extract was filtered through a 0.22 µm membrane filter and the filtrate was transferred to an autosampler vials for HPLC and gas chromatography analysis (Jin et al. 2010). For HPLC analysis of the extracts, 20 µl of the filtrate was injected to the Shimadzu HPLC system. Analysis of the samples was performed using Ascentis® C 18 column, 5 µm, 25 cm × 4.6 cm from Sigma-Aldrich. A gradient program having two mobile phases, a water/acetonitrile (15:85) v/v and B 100 % acetonitrile, was used. Time programing was: 0–3 min a 100 % A, 6.5–19.5 min 100 % B. A total flow rate of 0.6 ml/min was maintained. Run time of the samples was 45 min. DBP was detected using a UV detector at 225 nm (Thuren 1986). Residual DBP in the samples was quantified by preparing a standard curve for DBP (Park et al. 2016). Metabolic intermediates for DBP degradation were identified by the GC–MS system at Advanced Instrumentation Research Facility, Jawaharlal Nehru University, New Delhi, India, with column temperature of 100 °C, injection temperature 250 °C and total flow 16.3 ml/min. For calculation of biomass in terms of dry weight, bacterial cells were harvested and filtered with 0.45 µm membrane filter and dried in an oven at 100 °C. The dried biomass was measured with a weighing balance (Bratbak and Dundas 1984).

**Identification of PAEs-degrading genes**

Genes responsible for phthalate esters degradation from the strain 29D were identified by PCR. Primers known for PAEs degradation were synthesized and amplified (Table 2). The programing used for the PCR thermocycler was: 5 min at 95 °C, 30 cycles of 30 s at 95 °C, 30 s at Tm of corresponding primers, 90 s at 72 °C and final extension for 7 min at 72 °C (Han 2008). Gel-purified amplification products were sequenced at the DNA sequencing facility UDSC, University of Delhi, New Delhi, India. Sequencing of the amplicons was performed by specific primers for each gene and amplicons were cloned in M13 vector.

**Statistical analysis**

Statistical analysis of DBP degradation was performed by F test and one-way ANOVA with three replicates data using data analysis tool pack in Microsoft excel. For analysis, the hypothesis was made at the 5 % level of significance to calculate P and F values.

**Results and discussions**

**Isolation, characterization and identification of the bacteria**

Cells of the strain 29D were rod shaped, Gram negative, aerobic and without flagella, forming round and creamy
Colonies on agar plates. The cells grow on MSM with 1% methanol. Strain 29D was catalase and oxidase positive which is the characteristic of bacteria form of the genus *Methylobacillus* (Doronina et al. 2004; Urakami and Komagata 1986; Anthony 1982; Lidstrom 2006). Negative results were obtained for nitrate production, urease activity, H2S production, ammonia production, methyl red, Voges–Proskauer and starch hydrolysis. The strain was positive for indole production, lysine utilization and β-galactosidase activity. It was resistant to streptomycin (100 μg ml⁻¹) and susceptible to ampicillin (10 μg ml⁻¹), penicillin (10 μg ml⁻¹), kanamycin (10 μg ml⁻¹), tetracycline (10 μg ml⁻¹), and chloramphenicol (10 μg ml⁻¹). SEM at 20 KX revealed that strain 29D was rod shaped, smooth, without flagella with length ~2 μm and width ~0.2 μm. TEM of strain 29D at 20 KX revealed the presence of the outer membrane, peptidoglycan layer and plasma membrane (Fig. 1).

On comparison with the 16S-rRNA gene sequence of strain 29D, it found maximum identity with *Methylobacillus* arboreus clone SY117 (Accession No. KM041246.1). The strain 29D was designated as *Methylobacillus* sp. V29b and the obtained sequence was submitted to NCBI accession No. KM219114. The phylogenetic relationship of *Methylobacillus* sp. V29b is presented in Fig. 2.

**DBP biodegradation by Methylobacillus sp. V29b in MSM and contaminated samples**

Biodegradation experiments were conducted by growing *Methylobacillus* sp. V29b in MSM supplemented with DBP at 2000 mg/l. To quantify the residual DBP in the bacterial culture, samples were collected every 24 h and quantified using HPLC. Figure 3a presents the relationship between the growth of *Methylobacillus* sp. V29b and DBP degradation. Figure 3b presents the degradation of DBP in the PAEs-contaminated sample from a landfill site at Ghazipur, New Delhi, India. The amount of DBP quantified in contaminated samples was 441 mg/l. From Fig. 3a, it was observed that from the initial DBP concentration of 1997 mg/l, *Methylobacillus* sp. V29b degraded half DBP in 120 h and only 590.40 mg/l DBP was left after 192 h. Therefore, 70.5% of DBP was degraded in 192 h. From Fig. 3b, it may be observed that the *Methylobacillus* sp. V29b degraded half of the initial amount DBP in 96 h and 64.5% the initial amount of DBP in 144 h.

**Biodegradation kinetics and DBP degradation stoichiometry**

Researchers have reported degradation kinetics using various models. Degradation kinetics of organic pollutants was explained by first-order equations (Lu et al. 2009; Xu et al. 2005; Zeng et al. 2004). A second-order equation was also reported for degradation of phthalate esters by algae *Chlorella pyrenoidosa* (Yan et al. 1995). A modified Gompertz model was used to describe the effect of initial DBP concentration on DBP biodegradation by *Gordonia* sp. QH-11 (Jin et al. 2012). Haldane substrate inhibition model was used to explain DBP degradation in a continuous culture system (Wang et al. 1998). DBP biodegradation kinetics by *Methylobacillus* sp. V29b was explained by drawing a plot of specific growth rate (μ) and DBP concentration (Sav) (Fig. 4). It was observed that as the concentration of DBP was increased, there was increase in specific growth rate, but when DBP concentration reached 1900 mg/l, there was a decline in the specific growth rate. This behavior was explained by the growth inhibition model. Equation (1) represents the Monod model for growth inhibition. The calculated maximum specific growth (μmax) and half-velocity constant (Ks) were: 0.07 h⁻¹ and 998.2 mg/l, respectively.

\[
\mu = \frac{\mu_{\text{max}} S}{(K_s + S)},
\]

(1)
where $\mu$ is the specific growth rate of the microorganism, $\mu_{\text{max}}$ the maximum specific growth rate of the microorganism, $S$ the concentration of the limiting substrate for growth and $K_s$ the half-velocity constant.

Values of the coefficients:

$\mu_{\text{max}} = 0.07 \text{ h}^{-1},$

$K_s = 998.2 \text{ mg/l},$

Yield $= 0.43.$

Stoichiometry for DBP utilization and biomass formation (Shuler and Kargi 2002) is presented in Eq. (2):

\[
50C_{16}H_{22}O_4 + 893O_2 + 20NH_3 \rightarrow 20C_5H_7O_2N + 546H_2O + 700CO_2.
\]

(2) DBP Biomass

**Identification of metabolic intermediates**

Determination of the metabolic intermediates for DBP degradation was performed by analysis of the GC–MS results. Four metabolic intermediates dibutyl phthalate (DBP), monobutyl phthalate (MBP), phthalic acid (PA), and pyrocatechol (PC) were detected during DBP degradation by comparing the mass spectrum at a particular retention time with published mass spectra from the National Institute of Standards and Technology (NIST) database. Figure 5 presents the HPLC peaks and structure of the identified metabolic intermediates from GC–MS. It was observed that with time the length of the DBP peak was decreased, while the peak lengths of the MBP and PA was increased and on day 8 the peak length was highest for pyrocatechol (PC). The identified intermediates suggest that DBP was converted to MBP, which was converted to PA. The final product of the reaction was PC. The positive result for Rothera’s reaction confirms the benzene ring cleavage.

Few studies reported that PAEs degradation was mediated by a pathway where they are first converted to its monoester by esterase and then to phthalic acid (PA). PA is then transformed to carbon dioxide and water via an intermediate known as protocatechuate (Benckiser and Ottow 1982; Eaton and Ribbons 1982; Engelhardt and Wallnöfer 1978; Wang et al. 1995a, 2003a; Xu et al. 2005, 2007, 2006). Sometimes, two ester linkages in PAEs are cleaved by two different bacteria (Cartwright et al. 2000; Li et al. 2005; Li and Gu 2007). Some studies reported the formation of phthalic acid from DBP mediated by intermediate, monobutyl phthalate (MBP) (Benckiser and Ottow 1982; Gu et al. 2004; Wang et al. 2004; Xu et al. 2005). Based on the reported studies, the identified metabolic intermediates and Rothera’s test, a pathway for DBP degradation by *Methylobacillus* sp. V29b was proposed in Fig. 6. A similar pathway for dibutyl phthalate degradation in landfill bioreactor was reported for *Enterobacter* sp. T5 isolated from municipal solid waste (Fang et al. 2010). The study described the appearance of two major transient metabolites including phthalic acid (PA) and monobutyl phthalate (MBP). Pathway described for DBP degradation by *Methylobacillus* sp. V29b is the extension of the
pathway proposed for DBP degradation by Enterobacter sp. T5. In Methylobacillus sp. V29b DBP, degradation of one more metabolic intermediate pyrocatechol was identified as the final aromatic intermediate which fills the gap between PA to carbon dioxide conversion.

**Identification of phthalates-degrading genes**

*Methylobacillus* sp. V29b was able to grow on substrates protocatechuate, monobutyl phthalate, diethyl phthalate, benzyl butyl phthalate, dioctyl phthalate and dodecyl.
phthalate (Table 1). The growth of *Methylobacillus* sp. V29b decreases as the length and complexity of hydrocarbon attached to the phthalate ring increases (Wang et al. 2003b). Therefore, good growth was observed in PC, MBP and DEP. To examine the possibility for degradation of different phthalate esters by *Methylobacillus* sp. V29b and explore the PAEs degradation pathway, PAEs-degrading genes were amplified. Table 2 presents the list of primers
Table 1 Growth of *Methylobacillus* sp. V29b in different substrates

| Strain name | Methylobacillus sp. V29b |
|-------------|--------------------------|
| Methanol    | +                        |
| PC          | +++                      |
| MBP         | +++                      |
| DEP         | ++                       |
| BBP         | +                        |
| DOP         | +                        |
| DIDP        | +                        |

*PC* pyrocatechol, *MBP* monobutyl phthalate, *DEP* diethyl phthalate, *BBP* benzyl butyl phthalate, *DOP* dioctyl phthalate, *DIDP* diisodecyl phthalate

selected and synthesized for gene amplification. Table 3 presents the genes amplified from *Methylobacillus* sp. V29b genome. Gene sequences for primers FEH, HFDH, FOXG, FOXGS and FDK were obtained from Arthrobacter sp. 68b growing on phthalic acid as the sole carbon source (Stanislauskiené et al. 2011). Gene sequence for the primer transporter ATPase was obtained from Arthrobacter sp. 68b growing on phthalic acid as the sole carbon source (Stanislauskiené et al. 2011). Gene sequences for the primers oph-A1, -A2, -B, -C, -D, -H, -R, Tph-A2, -A3, and -B were obtained from the bacteria-degrading phthalate, isophthalate and terephthalate (Han 2008).

Biodegradation of phthalate esters is initiated by their transport inside the cell by phthalate permease (oph-D) which induces phthalate 4,5 dioxygenase. They belong to major facilitator superfamily with 12 hydrophobic membrane-spanning helices. Phthalate permeases are reported as transport enzymes (Eaton 2001; Keyser et al. 1976). Permeases from *P. putida* NMH102-2 and *B. cepacia* ATCC 17616 found similarity with anion-cation symporter family (Chang and Zylstra 1999). Permeases have reported multiple genes and have catalytic operons (Chang and Zylstra 1999; Eaton 2001; Sasoh et al. 2006; Wang et al. 1995b). Phthalate dioxygenase (oph-A1) catalyzes the incorporation of two hydroxyl groups on the phthalate ring to yield phthalate dihydrodiols (Eaton 2001; Keyser et al. 1976). Primer oph-A1 was used to amplify the enzyme called phthalate dioxygenase reductase (PDR). PDR is an iron sulfur flavoprotein which utilizes flavin mononucleotide (FMN) to accomplish electron transfer from reduced nicotinamide adenine nucleotide (NADH) to the one-electron acceptor, [2Fe-2S] (Correll et al. 1992).

Table 2. Primers used for identification of PAEs-degrading genes

| Primer name | Gene name                                      | References                      |
|-------------|------------------------------------------------|---------------------------------|
| Oph-A1      | 3,4-Dioxygenase oxygenase component large subunit | (Han 2008)                     |
| Oph-A2      | 3,4-Dioxygenase oxygenase component small      | (Han 2008)                     |
| Oph-B       | Phthalate dihydrodiol dehydrogenase            | (Han 2008)                     |
| Oph-C       | 3,4-Dihydroxyphthalate decarboxylase           | (Han 2008)                     |
| Oph-D       | d-Galactonate transporter                      | (Han 2008)                     |
| Oph-H       | Hemerythrin-like metal-binding protein         | (Han 2008)                     |
| Oph-R       | Transcriptional regulator, MarR family         | (Han 2008)                     |
| FEH         | Phthalic ester hydrolase                       | (Stanislauskiené et al. 2011)  |
| HFDH        | 3,4-Dihydroxy-3,4-dihydrophthalate dehydrogenase | (Stanislauskiené et al. 2011)  |
| FOXG        | Phthalate dioxygenase large and small subunits | (Stanislauskiené et al. 2011)  |
| FOXGS       | Ferredoxin and reductase subunits              | (Stanislauskiené et al. 2011)  |
| FDK         | 3,4-Dihydroxyphthalate-2-decarboxylase         | (Stanislauskiené et al. 2011)  |
| Prt A       | Transporter ATPase                             | (Choi et al. 2005)             |
| Tph-A2      | Terephthalate 1,2-dioxygenase oxygenase component large subunit | (Han 2008)                     |
| Tph-A3      | Terephthalate 1,2-dioxygenase oxygenase component small subunit | (Han 2008)                     |
| Tph-B       | Terephthalate dihydrodiol dehydrogenase       | (Han 2008)                     |
They occur in invertebrates Sipunculida (peanut worms), Brachiopoda (lamp shells) Priapulida (priapulid worms) and some Annelida (segmented worms, including leeches and polychaete worms) (Klippenstein et al. 1968; Loehr et al. 1978; Long et al. 1992). Hemerythrin-like proteins were reported in prokaryotes, specifically Methylococcus capsulatus and Desulfovibrio Vulgaris, and it was proposed that the oxygen-binding domain acted as an oxygen sensor (Xiong et al. 2000). Genes for hemerythrin-like metal-binding proteins were amplified using oph-H primer from the bacteria-degrading phthalate, isophthalate and terephthalate. They have been found in cluster with genes responsible for phthalate degradation, but their specific function in PAEs degradation is uncertain (Han 2008). Terephthalate metabolism is initiated by double hydroxylation at the position 1 and 2 of the ring by terephthalate 1,2 dioxygenase (Tph-B) to produce 2-hydro-1,2-dihydroxy terephthalic acid (Choi et al. 2005; Schlafli et al. 1994). 2-Hydro-1,2-dihydroxy terephthalic acid is further metabolized to 3,4-dihydroxybenzoate (Vamsee-Krishna et al. 2006; Wang et al. 1995b). Figure 7 presents the amplified gene products for phthalate ester-degrading genes.

This study is the first to report the isolation and characterization of a Gram-negative bacterium form of the genus Methylobacillus-degrading DBP. Bacteria from the genus Methylobacillus are methylotrophs and have the ribulose monophosphate (RuMP) pathway for formaldehyde assimilation (Bratina et al. 1992; Doronina et al. 2004; Urakami and Komagata 1986). The isolated Methylobacillus sp. V29b not only grows on methanol, but is able to grow on other substrates, such as protocatechuate, monobutyl phthalate, diethyl phthalate, benzyl butyl phthalate, dioctyl phthalate and diisodecyl phthalate. DBP degradation was reported by Sphingomonas sp. DK4 (5 mg/l) (Chang et al. 2004), Pseudomonas fluorescens B-1(2.5 & 10) (Xu et al. 2005), Acinetobacter Iwoffi (1000 mg/l) (Hashizume et al. 2002), Corynbacterium nitilofius G11 (100 mg/l), R. rhodochrous G2, G7 (100 mg/l) (Chao et al. 2006) and R. Coprophilus G2, G7 (100 mg/l) (Chao et al. 2006). Methylobacillus sp. V29b was able to degrade 1997 mg/l DBP, which is very high reported till so far. Degradation kinetics of organic pollutants were explained by first-order equations (Lu et al. 2009; Xu et al. 2005; Zeng et al. 2004), second-order equation (Yan et al. 1995), modified Gompertz model (Jin et al. 2012) and Haldane model (Wang et al. 1998). Methylobacillus sp. V29b demonstrated DBP degradation by the Monod model for growth inhibition, which is a new perspective to present the degradation of pollutants. While a majority of the studies are concentrated on the degradation of PAEs in minimal media (Chao et al. 2006; Fang et al. 2010; Wu et al. 2010a; Wu et al. 2010b), this study focuses on the degradation of DBP in contaminated samples collected from landfill sites. Landfill sites are composed of wastes from domestic, medical, pharmaceutical and industrial sources containing a variety of plastic items and PAEs. Therefore, it was selected for DBP degradation studies. It was an efficient degrader of DBP in PAEs-contaminated sample; therefore, it can be considered as a potential candidate for bioremediation of the sites contaminated with pollutants.

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| Amplicons | Primer name | Name of the gene | Amplicon size |
|-----------|-------------|------------------|---------------|
| 84        | Tph-B-F2,R2 | Terephthalate dehydroxygenase | 500 kb |
| 85        | Tph-B-F1,R1 | Terephthalate dehydroxygenase | 800 kb |
| 86        | Oph-A1      | Phthalate dioxygenase | 800 kb |
| 87        | Oph-D       | Phthalate permease | 1 kb |
| 89        | Oph-C       | Phthalate decarboxylase | 1 kb |
| 93        | Tph-B-F2,R3 | Terephthalate dehydroxygenase | 500 bp |
| 95        | Oph-H       | Hemerythin-like metal-binding protein | 200 bp |
| 99        | Oph-B       | Phthalate dehydrogenase | 500 bp |
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