**Phthalate Esters Metabolic Strain** *Gordonia* sp. GZ-YC7, A Potential Soil Degrader for High Concentration Di-(2-ethylhexyl) Phthalate

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**Abstract:** As commonly used chemical plasticizers in plastic products, phthalate esters had become a serious ubiquitous environmental pollutant, such as in soil of plastic film mulch culture. Microbial degradation or transformation was regarded as a suitable strategy to solve the phthalate esters pollution. Thus, a new phthalate esters degrading strain *Gordonia* sp. GZ-YC7 was isolated in this study, which exhibited the highest di-(2-ethylhexyl) phthalate degradation efficiency under 1000 mg/L and the strongest tolerance to 4000 mg/L. The comparative genomic analysis showed that there exist diverse degradation pathways for various phthalate esters such as di-(2-ethylhexyl) phthalate and dibutyl phthalate in *Gordonia* sp. GZ-YC7, which possibly contributes to its broad substrate spectrum, high degrading efficiency and high tolerance to phthalate esters. *Gordonia* sp. GZ-YC7 is potential for bioremediation of phthalate esters in polluted soil environments.

**Keywords:** PAEs; Comparative genomics analysis; DEHP; Bioremediation

**1. Introduction**

Global increasing of plastic production and utilization, while only 9% of various plastic waste being recycled, is leading to the accumulation and persistence of plastic debris in the environment [1]. As commonly used chemical plasticizers in plastic products to improve the flexibility and workability of various plastic products [2], phthalate esters (PAEs) also enriched in many natural environments, such as wastewater treatment plants [3], river water [4], and Chinese plastic film mulch soil [5-8]. Plastic debris can disperse throughout the oceans and their presence and distribution even in inhabited and remote locations on the world [9]. Thus, plastic debris has been recognized as a global health threat such as the marine debris could be harmful to marine organisms and seabirds, and microplastics are detrimental to chlorophyta and Ochrophyta most in growth or photosynthesis [10,11]. Also, PAEs had already been recognized as endocrine-disrupting chemicals (EDCs) which could interfere with the physiological endocrine function of animals and humans [12]. Nowadays study found that exposure to microplastics could enhance the antibiotic resistome of mice gut microbiota which would make traditional antibiotic treatments difficult and costly [13]. Thus, effective methods to eliminate PAEs pollution are essential. Until now, more than sixty kinds of PAEs have been produced and consumed in plastic production [14]. PAEs are differentiated by the various side chains such as the simple-side-chain compound (diethyl phthalate, DEP and butyl benzyl phthalate, BBP), and complex-side-chain compound (di-(2-ethylhexyl) phthalate, DEHP) [14,15]. One of the most widely studied PAE compound is DEHP, which accounts for nearly 50% of the total PAEs pollution production [16]. In addition, DEHP is found to have the highest concentration among other PAEs in water, air, soils and sediments.
The degradation strategies of PAEs includes hydrolytic, photolytic, and microbial degradation [17]. As hydrolytic and photolytic method works slowly and weakly, microbial degradation is regarded as the most promising strategy to solve the PAEs pollution with many advantages such as high efficiency, fast kinetics, mild reaction conditions and no secondary pollution [18]. The microbial degradation pathways of different PAEs were systematically identified. And it has been known that the PAEs degradation efficiency was influenced by pH, temperature, retention time, and microbial communities of the environment [19].

Two main steps are involved in the PAEs biodegradation metabolic pathways according to the intermediates identification: (I) transformation of PAEs to phthalic acid (PA) and (II) complete degradation of PA [20].

Nowadays, more than 80 PAEs-degrading strains isolated from various soil bodies or other environments have been identified, belonging to Gordonia, Sphingomonas [21], Pseudomonas [22], Rhodococcus [23] and Comamonas [24]. Some PAEs-degrading strains can tolerate 500-2000 mg/L PAEs, but their application range was restricted due to their slow growth and reproduction, being unable to tolerance high concentration PAEs, and poor environmental adaptability. Furthermore, PAEs in natural environment usually exist in mixed states including different proportions of dibutyl phthalate (DBP), DEHP, DEP, BBP, di-octyl phthalate (DnOP), and dimethyl phthalate (DMP). DEHP, DEP, DBP, BBP, DnOP, and dimethyl phthalate (DMP) have been ranked as one of the high-priority control pollutants by the US Environmental Protection Agency (USEPA) [25]. But most reported PAEs-degrading bacteria show limited substrates spectrum and degradation efficiency in soil or in aqueous environment. For example, *Gordonia polysaprophytica* G1 could degrade 243 mg/kg DEHP in soil to 50% over 2 days, while *Pseudomonas* sp. YJB6 could completely reduce 200 mg/L DBP in 3 days [22,26]. Therefore, isolating more PAEs-degrading strains with stronger ability, higher efficiency to degrade a broader variety of PAEs is essential to directly apply or construct engineering chassis cells for PAEs bioremediation.

This study aims to (1) isolate novel PAEs degrading strains exhibiting both higher efficiency and broader spectrum than reported strains, from Chinese PAEs polluted soil sample; (2) reveal the potential mechanism and degradation pathways of PAEs metabolism of isolated strains, based on comparative genomic analysis; (3) evaluate application potential of isolated strains for aqueous and soil environment bioremediation and for the construction of PAEs-degrading chassis cell factories.

2. Materials and Methods

2.1. Chemicals and media

Diethyl phthalate (DEP), dipropyl phthalate (DPrP), dibutyl phthalate (DBP), benzyl butyl phthalate (BBP), di-(2-ethylhexyl) phthalate (DEHP), dioctyl phthalate (DnOP), diisononyl phthalate (DiNP) with 99% purity were purchased from Aladdin Chemistry Co., Ltd. (Shanghai, China). Methanol was HPLC grade while all other chemical reagents used in this experiment were analytical grade.

Luria-Bertani (LB) medium consisted of (g/L): NaCl 10, tryptone 10, yeast extract 5. Basic inorganic salt medium (BSM) consisted of (g/L): K2HPO4·3H2O 1.0, NaCl 1.0, (NH4)2SO4 0.5, MgSO4·7H2O 0.4, CaCl2 0.0755, FeCl3 0.0143.

For the preparation of solid media, 20 g/L agar was added to LB and BSM liquid media.

2.2. Isolation and identification of phthalate esters degrading bacteria

Soil sample was collected from the LaoHeiShan landfill, Liupanshui, Guizhou Province, China (39° 52′ 48″ N, 105° 30′ 5″ E), from which 25 g soil sample was added into 20 mL sterile water for mixture completely with glass beads. Then 2 mL supernatant was added to 100 mL BSM medium containing 200 mg/L DEHP and the mixture was cultured under 30 °C and 180 rpm for 4 days. Then 2 mL mixture was transferred to 100 mL fresh
BSM medium containing 200 mg/L DEHP for another round of 4 days culture. After several rounds, the cultures were gradient-diluted and spread onto the solid LB medium, and then incubated at 30 °C for 4 days. The single colonies were selected and inoculated into BSM medium containing different 200 mg/L PAEs, respectively, to test the PAEs degradation abilities.

The morphology characteristics of pure colony were identified, while their 16S rRNA gene were sequenced (Tsingke Biotechnology Co., Ltd) after amplified using universal primers 27F (5′-AGAGTTTGATCMTGGCTCAG-3′) and 1492R (5′-GGTTACCTTGTTACGACTT-3′). The 16S rRNA sequences comparison between the obtained strains and the reported strains in GenBank was conducted by BLAST and phylogenetic tree was then constructed using the neighbor-joining method with MEGAX software.

2.3. Culture conditions and analytic method for phthalate esters

From slant, PAEs degrading bacterial strains were inoculated into LB medium and incubated under 30 °C and 180 rpm until the OD600 of broth reached 0.6-0.8. Cells were harvested by centrifugation at 6000 rpm for 3 min and washed with sterile water for three times. Cells were then suspended for a final seed solution with the OD600 value of 0.8, which was inoculated into 50 mL BSM medium by a 2 % (v/v) inoculation amount. Each group for control or treatment was in triplicate.

The broth was directly used to detect PAEs content. Firstly, PAEs were extracted with an equal volume of dichloromethane, and then the organic phase was transferred to a rotary evaporator for drying. Then the samples were dissolved in 5 mL methanol and filtered through a 0.22 μm membrane filters. PAEs content was detected by HPLC (Agilent 1260) under the follow conditions: the UV wavelength of 235 nm; C18 column; the mobile phase contained 90% (vol) methanol and 10% (vol) water; the flow rate of 1.0 mL/min.

2.4. Phthalate esters degradation ability of isolated strains in BSM medium

2.4.1. Degrading substrates spectrum

To test the substrate spectrum of isolated PAEs-degrading strains, five hundred milligrams of DEP, DPrP, DBP, BBP, DEHP, DnOP, and DiNP were added respectively in 1 L liquid BSM medium as the sole source of carbon and energy. After inoculation, all the cultures were incubated at 180 rpm and 30 °C for 2 days. Then the residue PAEs in the broth was detected by HPLC, while the cell growth OD600 was monitored by a spectrophotometer (UV5200, Shanghai Metash Instruments Co., Ltd, Shanghai).

2.4.2. Tolerance to high concentrations of phthalate esters degrading strains

Five kinds of commonly used PAEs (DBP, BBP, DEHP, DnOP, and DiNP) were selected. The initial PAEs concentrations were set at 4 g/L. Strain was inoculated into liquid BSM medium containing DBP, BBP, DEHP, DnOP, and DiNP, respectively, and the residual concentrations of PAEs were determined on days 1, 3, and 5, respectively.

2.4.3. Mixed phthalate esters degradation

PAEs usually existed in the natural environment as mixed format. Thus, four kinds of PAEs (200 mg/L DEP, DBP, DEHP, and DnOP, respectively) were mixed to afford a final PEAs concentration of 800 mg/L in BSM medium. The cell growth (OD600) and residue concentration of DEHP were measured at each 12-hour intervals for 3 days.

2.4.4. Effects of environmental factors on di-(2-ethylhexyl) phthalate degradation

A pH value range of 5, 6, 7, 8, 9, 10, a temperature range of 15 °C, 25 °C, 30 °C, 37 °C, 42 °C, and a NaCl content range of 2 %, 4 %, 6 %, 8 %, 10 % (v/v), were selected to evaluate the effects of environmental factors on the DEHP degradation. The cell growth (OD600) and residue concentration of DEHP were detected after 2 days incubation at 180 rpm and 30 °C. The group without bacterial inoculation was set as control simultaneously.

2.4.5. Degradation kinetics of di-(2-ethylhexyl) phthalate

Under optimal conditions, the initial DEHP concentrations were set as 200, 500, 1000 and 2000 mg/L. The DEHP residue concentration in cultures was measured at each 12-hour intervals for 3 days. In order to explore the effect of the initial concentration of DEHP on the degradation efficiency, the first-order kinetic equation was used to describe the degradation efficiency. The formula was as the following: \[ \ln C = -Kt + A \], where the C is
2.5. Degradation of di-(2-ethylhexyl) phthalate in soil

The soil was collected from a campus garden of the Zhejiang University of Technology (Hangzhou, China), in which large materials such as leaves, branches, roots, and stones, were removed. The soil sample was then sieved using a 2 mm mesh. The pH value and the water content of the soil were 7.19 and 17.79 % (water to soil ratio at 2.5: 1 w/v) respectively, thus the soil sample was dried naturally at room temperature. Meanwhile, another sterilized part of the soil sample was prepared by autoclaving at 121 °C for 30 min. Then, 4 g/L DEHP acetone solution was added into natural soil and sterile soil respectively to a final concentration of 500 mg/kg. The resultant soil contaminated by DEHP was left overnight until the acetone had completely volatilized. Then, 40 g prepared soil sample was putted in a 50 ml beaker and mixed with sterile water to restore the soil water content to the initial value (2.5: 1 w/v). Two gram soil samples were collected at days 1, 2, 3, 4, 5, respectively, and stored at -80 °C. The freeze-dried soil samples were mixed with 3 mL methanol for 30 min ultrasonic dissolution, and soaked overnight to dissolve fully. The supernatant was collected by centrifugation at 12000 rpm for 5 min and filtered via 0.22 μm membrane filter before HPLC detection for residue DEHP.

2.6. Genome sequence analysis of strain GZ-YC7

The complete genome sequencing of GZ-YC7 was performed using a combined sequencing platform of DNBSEQ and PacBio at the Beijing Genomics Institute by BGI Genomics Co. Ltd., Shenzhen, China. And unavailable PacBio subreads (length < 1 kb) were removed.

The Canu program was selected for self-correction. Then, draft genomic unitigs were assembled by Canu according to the high-quality corrected circular consensus sequence subreads set. And GATK (https://www.broadinstitute.org/gatk/) was taken to make corrections of single-base. Gene prediction of GZ-YC7 was performed using glimmer3 (http://www.cbcb.umd.edu/software/glimmer/) with Hidden Markov models. tRNA, rRNA and sRNAs were predicted according to tRNAscan-SE, RNAmmer, and the Rfam database.

2.7. Gene annotation and protein classification

The Blast alignment tool was chosen for function annotation. Seven databases such as KEGG (Kyoto Encyclopedia of Genes and Genomes), COG (Clusters of Orthologous Groups), NR (Non-Redundant Protein Database data bases), Swiss-Prot, and GO (Gene Ontology), TrEMBL and EggNOG were used for general function annotation. Virulence factors and resistance gene were identified based on the core dataset in VFDB (Virulence Factors of Pathogenic Bacteria), ARDB (Antibiotic Resistance Genes Database) database, and CAZy (Carbohydrate-Active Enzymes Database). All genome sequences used in this research were downloaded from NCBI database.

3. Results

3.1. Isolation and characterization of strains

Among all isolated PAEs-degrading strains, strain GZ-YC7 exhibited the broadest substrate spectrum and was then identified. Its colonies on LB solid plates appear orange color and its cell is round shape and Gram positive (Fig. 1a, 1b, 1c). GZ-YC7 could utilize glucose, fructose, xylose, mannose and starch. GZ-YC7 showed positive results both on catalase and hydrogen sulfide experiments, but it showed negative results on urease, V-P, methyl red, and casein hydrolysis tests.

The length of GZ-YC7 16S rRNA was 1,417 bp and the sequence information has been uploaded to GenBank with the accession number of OM049462. It was found that GZ-YC7 shared 99 % similarity to Gordonia alkanivorans strain 1960BRRJ (MK182084.1), and the corresponding phylogenetic tree was displayed in Fig. 1d. Based on the phylogenetic analysis of 16S rRNA sequences, morphological, biochemical, and physiological
characteristics, the strain GZ-YC7 was finally identified as a species belonging to Genus Gordonia and named as *Gordonia* sp. GZ-YC7. *Gordonia* sp. GZ-YC7 has been deposited in the China Center for Type Culture Collection (CCTCC), Wuhan University, China, with the number of CCTCC M2022045.

### 3.2. Phthalate esters degradation ability of isolated strains in BSM medium

#### 3.2.1. Degrading substrate spectrum

Most PAEs-degrading strains always showed preference to finite substrates which might limit their applicability. For example, only DBP (1200 mg/L) was reported to be mineralized (approximately 90%) in 48 h by strain HD-1 which was enriched from activated sludge (He et al., 2013). Thus, the degrading substrate spectrum was an important evaluation criterion for the applied potential of PAEs-degrading strains.

In this study, all seven kinds of PAEs could be efficiently degraded by *Gordonia* sp. GZ-YC7 within 48 h (Fig. 2a). GZ-YC7 exhibited the best degrading efficiency on BBP and DEHP that were fully degraded, while its degradation ratio of DnOP and DiNP also reached 91.22% and 97.57%, respectively. However, the degradation ratio of short-chain PAEs (DBP, DPrP, and DEP) was only 88.02%, 66.77% and 59.72%.

It is generally assumed that PAEs containing long chains (DiNP, DnOP, DEHP) and benzene ring side chains (BBP) are hardly degraded because of steric hindrance [27]. For examples, the degradation rate of DBP was significantly higher than DEHP and DnOP by *Rhodococcus* sp. strain WJ4 [28]. Arthrobacter sp. ZJUTW can efficiently degrade DBP, while it cannot degrade DEHP [29]. However, some PAEs degrading strains showed versatile ability for PAEs with different side chains. For instances, *Rhodococcus* sp. 2G could degrade 7 types of PAEs, but the degradation efficiency increased gradually with an increase in the side chain length [24]. *Gordonia alkanivorans* strain YC-RL2 was able to degrade all kinds of PAEs with short-chain length, long-chain length even with the benzene ring side chains, the degradation efficiency of DEHP with the concentration of 100 mg/L reached higher than 99 % in 7 days [30].

*Gordonia* sp. GZ-YC7 also showed degrading ability of structurally diverse phthalate esters, especially long-side chains PAEs (DiNP, DnOP, and DEHP) and benzene ring side-chains (BBP), suggesting it has great potential for application in the future environmental bioremediation.

#### 3.2.2. Tolerance to high concentrations of phthalate esters degrading strains

The ability of GZ-YC7 to degrade and tolerate DBP, BBP, DEHP, DnOP and DiNP was measured respectively, under a high concentration of 4 g/L which was the highest test concentration among the known test conditions in the previous publications. The results (Fig. 2b) showed that GZ-YC7 could degrade 70.71% of high concentration DEHP in 5 days, which was much higher than the DEHP degradation efficiency of *Gordonia alkanivorans* YC-RL2 (< 10 % of 4 g/L DEHP degraded in 7 days) [30]. As for DnOP and DiNP with long chains, the degradation efficiency of GZ-YC7 was 47.82% and 54.54%, respectively. In conclusion, GZ-YC7 could keep high degradation efficiency for high concentration PAEs, especially for DEHP, which indicated its higher tolerance than other strains and potential degradation to various PAEs.

#### 3.2.3. Mixed phthalate esters degradation

PAEs usually present in coexisting forms in the natural environment, thus the degradation ability of mixed PAEs is essential for a strain to be practically utilized. Four types of commonly used PAEs including DEP, DBP, DEHP, and DnOP were mixed equally for the degradation experiment (Fig. 3a). It was found that DEP and DBP with short chains were degraded first while the longest-chain DnOP was degraded after 24 h
cultivation. In addition, GZ-YC7 could degrade DEHP and DBP with the efficiency over 95% at 72 h, while the degradation efficiency of DEP and DnOP was 77.56% and 79.97%, respectively. Moreover, the degradation rate of DnOP in the mixed PAEs was significantly lower than that of only DnOP, suggesting the substrate preference of GZ-YC7 in PAEs mixed environment. Thus, the result illustrated that the PAEs degradation ability might different in various environment even of the same strain that could provide new lights for polluted environmental remediation.

3.2.4. Effects of environmental factors on di-(2-ethylhexyl) phthalate degradation

The effects of initial pH (Fig. 2c), temperature (Fig. 2d), and NaCl content (Fig. 2e) on growth and DEHP degradation efficiency of GZ-YC7 were investigated in this study. Fig. 2c showed that GZ-YC7 could degrade DEHP in the pH range of 5 to 10, and 500 mg/L DEHP was completely removed after 48 h incubation under pH 7.0 to 10.0. Meanwhile, the optimum pH for GZ-YC7 growth was 8.0. DEHP could be completely degraded by GZ-YC7 at temperature of 25 °C, 30 °C and 37 °C, meanwhile, the optimum temperature for GZ-YC7 growth was 30°C (Fig. 2d). The concentration of NaCl showed a significant effect on DEHP degradation. GZ-YC7 could completely degrade DEHP at the concentration of 2% and 4% while the strain growth and DEHP degradation ability were significantly inhibited when the NaCl concentration increased above 6% (Fig. 2e).

3.2.5. Degradation kinetics of di-(2-ethylhexyl) phthalate by strain GZ-YC7

In some previous publications, it was found that PAEs degradation was inhibited by the increasing initial concentration of PAEs [31,32]. The initial DEHP concentration also influenced the growth of a DEHP-degrading *Pseudoxanthomonas* sp. N4. However, the strain N4 growth was obviously limited when the initial DEHP concentration decreased below 500 mg/L, while its growth stopped when the initial DEHP concentration increased above 500 mg/L [33].

Thus, the degradation ability of GZ-YC7 with the initial DEHP concentration of 200, 500, 1000, and 2000 mg/L was also measured respectively. The results (Fig. 2f) showed that the degradation rate really decreased when the initial DEHP concentration increased from 200 to 2000 mg/L, meanwhile the half-life increased gradually (Table B.1). DEHP was completely degraded by GZ-YC7 within 24 h under the DEHP initial concentration of 500 mg/L, while GZ-YC7 growth remained until 36 h (Fig. 2f). Only 87.11% DEHP was degraded in 3 days under the DEHP initial concentration of 2000 mg/L. However, GZ-YC7 exhibited a higher degradation efficiency and tolerant ability to high concentration of DEHP when compared to reported strains (Table 1). For examples, the DEHP degradation efficiency of GZ-YC7 was significantly higher than that of *Agromyces* sp. MT-O (35.0% of 1000 mg/L DEHP degraded within 7 days), *Microbacterium* sp. CQ0110Y (half-life of 2000 mg/L DEHP was 2.36 day), and *Rhodococcus pyridinivorans* XB (87.5 % of 800 mg/L DEHP degraded within 3 days) [26,34,35]. It was convinced that high PAEs initial concentration could inhibit the DEHP degradation by GZ-YC7, but the half-life kept at a level from 0.45 to 1.49 days when the DEHP concentration increased from 200 mg/L to 2000 mg/L.

It was also found that most strains only grew and kept degrading in the concentration of DEHP lower than 1200 mg/L, but the DEHP degradation efficiency of *Gordonia* sp. GZ-YC7 was higher than any other DEHP-degrading strains (Table 1). When the DEHP concentration was below 500 mg/L, GZ-YC7 could completely remove the DEHP in one day while *Bacillus mojavensis* B1811 need four days cultivation in 500 mg/L DEHP. When DEHP concentration reached 1000 mg/L, *Mycolicibacterium phocaicum* RL-HY01 had the highest DEHP degradation efficiency among the reported strains that DEHP was completely removed in 3 days, while GZ-YC7 only need 2.5 days. Overall, *Gordonia* sp. GZ-YC7 had the highest degradation efficiency under the DEHP concentration lower than 1000 mg/L. Most DEHP concentration in natural environments were at low level.
(<1000 mg/L), the new DEHP-degrading strain *Gordonia* sp. GZ-YC7 with the highest DEHP degradation efficiency indicated greater potential in environmental governance.

In addition, natural environments were diverse and dynamic, where sometimes the concentration of DEHP might be much higher than conventional condition. Thus, strains with high tolerance ability of DEHP concentration had wider applications and could provide new ideas for the remediation of extreme polluted environments. The DEHP concentration of 4000 mg/L degraded by *Gordonia* sp. GZ-YC7 was known to be the highest DEHP concentration. The results showed that GZ-YC7 could still grow in 4000 mg/L DEHP and DEHP degradation reached 70.71 % after 5 days. It indicated that *Gordonia* sp. GZ-YC7 had the extreme DEHP stress tolerance with the highest degradation efficiency under low DEHP concentration (<1000 mg/L) and the strongest tolerance in 4000 mg/L DEHP. Thus GZ-YC7 was a suitable chose for DEHP biodegradation.

### 3.3. Degradation of di-(2-ethylhexyl) phthalate in soil

DEHP was more easily enriched in soil than in water and leaded to the seriously soil contamination [26]. Thus, the DEHP degradation efficiency of strain GZ-YC7 in soil was necessary to evaluate. The time course of DEHP concentration changes in soil samples was showed in Fig. 3b. No significant change of DEHP concentration was detected in both group NSS and group SS within 5 days. After the inoculation of strain GZ-YC7, 45.22% and 47.33% DEHP with the initial concentration of 500 mg/kg was degraded in group NSS7 and SS7 respectively. It could be found that there was no significant difference of DEHP degradation by GZ-YC7 with or without soil sterilization, suggesting that there was no synergy of GZ-YC7 with native microorganisms in the soil and only GZ-YC7 played the role of DEHP degrader in soil samples [34,36]. In the previous publications, it was found that the soil microbial activity was inhibited when DEHP concentration reached 100 mg/kg [27,35,37,38]. However, GZ-YC7 could still grow and efficiently degrade 500 mg/kg DEHP in soil. GZ-YC7 was potential for application in high concentration DEHP polluted environments.

### 3.4. Genome sequencing and analysis

To date, the complete genomic information of only six *Gordonia* sp. strains including WA4-43, KTR9, JH63, 135, PDNC005 and YC-JH1, had been uploaded to NCBI (2022/1/11). The whole genome circle map of GZ-YC7 was completed by bioinformatics analysis after the quality control (Fig. A1). The sequencing data statistics of DNBSEQ and PacBio are displayed in Table B2.

The complete genome of strain GZ-YC7 was identified to be 5,027,874 bp with the GC content of 67.93 % and there was no plasmid detected in GZ-YC7. This genome revealed 4,654 predicted genes including 72 none-coding RNA (ncRNA) (Table B3). A total of 9 sRNA candidates were predicted, which may regulate some specific biological functions such as biofilm formation, iron metabolism and so on [39]. All genes were annotated against twelve databases including VFDB, ARDB, TREMBL, CAZY, IPR, Swiss-Prot, COG, CARD, GO, KEGG, NR, and T3SS. The details of all above genome data were displayed in Table B3.

### 3.5. Comparison of esterase enzymes

In the process of catalysis of PAEs by esterase (Fig. A2a), esterase could be divided into three types. As shown in Table 2, seven putative esterase genes were predicted in GZ-YC7 genome based on reported esterase genes in previous publications (Table B4). The gene GZ-YC7GL001189 of GZ-YC7 shared 98.94 % similarity to the reported gene *mehpH* in *Gordonia* sp. P8219 [40]. The similarity between GZ-YC7GL004260 and the gene *estS1* of *Sulfolobus acidophilus* DSM10332 was only 31.62 % (Zhang et al., 2014). The phylogenetic tree of the 7 putative esterase genes and other reported esterase genes was
constructed as in Fig. A 2b. It was confirmed that three types of esterase genes simultaneously exist in GZ-YC7 genome. The diversity of esterase enzymes existing in GZ-YC7 might play role on the degradation ability of various PAEs compounds.

Eight putative esterase enzymes were also predicted in *Gordonia alkanivorans* YC-RL2. However, only four of the eight enzymes showed more than 30 % identity: (1) WP_006868835.1, 32 % identity to Est1 of *Sulfobacillus acidophilus* DSM10332; (2) WP_005200181.1, 38 % similarity with EstG of *Sphingobium* sp. SM42; (3) WP_006358366.1 and (4) WP_006358508.1, 35 % and 37 % identity to CarEW of *Bacillus* sp. K91, respectively [30].

3.6. Genetic peculiarities and pathway of phthalate esters degradation by GZ-YC7

It has been reported that the *pht* cluster, *pca* cluster, ben cluster, and *cat* clusters participated in the process of PAEs biodegradation [24,30,41]. After PAEs were catalyzed to phthalic acid (PA) by esterase, the metabolism from PA to TCA cycle was catalyzed by gene clusters *pht*, *pca*, ben, and cat (Fig. 4a). However, no *pht* gene cluster was detected in *Rhodococcus* sp. 2G which contains whole *pca* gene cluster, ben gene cluster, and cat gene cluster annotated in the genome. Similarly, no *pht* gene cluster was detected in *Gordonia alkanivorans* YC-RL2 which contains only ben gene cluster and cat gene cluster annotated in the genome. But what really distinguished *Gordonia* sp. GZ-YC7 is that all gene clusters of *pht*, *pca*, ben, and *cat* were detected in the genome. Two metabolism pathways of PAEs predicted previously were displayed in Fig. 4b [24]. It could be seen genes participated in these two pathways were all existed in GZ-YC7 genome which suggested that two metabolism pathways might coexisted in GZ-YC7. In a word, these two genetic peculiarities of GZ-YC7 may contribute to its high degradation efficiency of PAEs.

The gene distribution comparison of the above gene clusters between *Gordonia* sp. GZ-YC7 and other strains (Fig. 5) showed that the gene distribution of *pht*Aa, *pht*Ab, *pht*Ac, and *pht*Ad belonging to *pht* cluster in *Gordonia* sp. GZ-YC7 were the same to that in *Gordonia* sp. HS-NH1 and *Gordonia* sp. YC-JH1. The *pca* cluster of *Gordonia* sp. GZ-YC7 shared the highest similarity with *Gordonia* sp. YC-JH1. However, more transport genes (*benM, benE, benK*) involved in ben cluster were annotated in GZ-YC7, while *benM* exist only within the *cat* gene cluster in GZ-YC7 genome. These transport genes could influence the benzoate metabolic process. All the above genetic peculiarities and coexisting various esters, both contribute to the substrate diversity and high degrading efficiency of GZ-YC7. Thus, GZ-YC7 could provide abundant raw materials for the construction of totipotent PAEs-degrading strains.

PAEs metabolites gene clusters analysis showed that *Gordonia* sp. GZ-YC7 possessed completely metabolic pathways to degrade various PAEs, which also contributes to the wide substrate spectrum of PAEs degradation. In order to have deeper insight into the genetic background of the new PAEs-degrading strain *Gordonia* sp. GZ-YC7, all the related pathways were predicted and analyzed. According to the annotation of genes associated with PAEs biodegradation processes, the putative pathways were proposed as Fig. 4b. Although genes encoded decarboxylases which could convert PA (Phthalate) to benzoate were not discovered in GZ-YC7, the pathway catalyzed by *pca* gene cluster was completely [42,43]. PA would be converted to phthalate 3,4-cis-dihydrodiol and finally to Acetyl-CoA which was then entered the TCA cycle.

3.2. Figures, Tables and Schemes
Figure 1. GZ-YC7 morphology of (a) colonies on the LB solid plate, (b) cell under TEM (60,000 ×), (c) gram staining result mixed with E. coli (red cells are GZ-YC7), and (d) phylogenetic tree analysis based on 16S rRNA sequences, Neighbor-Joining method with a bootstrap value of 1000.
Figure 2. PAEs degradation and the growth of strain GZ-YC7 under 500 mg/L concentration of PAEs for 2 days (a); different PAEs degradation ability of GZ-YC7 under a high concentration of 4,000 mg/L (b). DEHP degradation ratio and the growth of strain GZ-YC7 under different pH (c), temperature (d), NaCl concentration (e), and DEHP initial concentration with growth curves of GZ-YC7 under 500 mg/L DEHP (f).
Figure 3. (a) PAEs Degradation and growth ability of GZ-YC7 in the BSM containing 800 mg/L mixed four PAEs (200 mg/L of DEP, DBP, DEHP, and DnOP, respectively); (b) DEHP degradation by GZ-YC7 in soil. NSS: Non-sterilized soil without GZ-YC7; NSS7: Non-sterilized soil with GZ-YC7; SS: Sterilized soil without GZ-YC7; SS7: Sterilized soil with GZ-YC7.
Figure 4. Gene clusters (a) the putative pathways (b) of PAEs degradation and in Gordonia sp. GZ-YC7. Module 1 in blue is catalyzed by esters; Module 2 in yellow is catalyzed by the pht gene cluster; Module 3 in green is catalyzed by the ben and cat gene clusters; Module 4 in red is catalyzed by the pca gene cluster.
Figure 5. The comparative genomics analysis of the gene clusters pht (a), pca (b), ben (c), and cat (c).
Table 1. Comparison between the strain GZ-YC7 and reported PAEs-degrading strains

| Strain                     | Source                     | Degrading substrates spectrum | DEHP degrading efficiency | References |
|---------------------------|----------------------------|-------------------------------|---------------------------|------------|
| Rhodococcus sp. 2G        | activated sludge           | DEHP, DMP, DEP, DBP, BBP, DnOP, DiNP | 200 mg/L, 5d, >95%        | [24]       |
| Rhodococcus ruber YC-YT1  | marine plastic debris      | DEHP, DDP, DNP, DOP, DCHP, BBP, DHPP, DHP, | 100 mg/L, 3d, >95%        | [12]       |
| Rhodococcus pyridinivorans XB | activated sludge          | DEHP, DMP, DEP, DBP          | 400 mg/L, 3d, 100%        | [35]       |
| Agromyces sp. MT-O        | landfill soil              | DEHP, DMP, DEP, DBP, DnOP    | 1000 mg/L, 7d, 65%        | [34]       |
| Bacillus mojavensis B1811 | soil                       | DEHP, DEP, DMP, DBP, BBP, DnOP, DPP | 500 mg/L, 4d, 100%        | [44]       |
| Pseudarthrobacter defluvi E5 | agricultural soil          | DEHP, DMP, DEP, DBP, DHXP    | 1200 mg/L, 2d, >50%       | [42]       |
| Mycolicibacterium phocaicum RL-HY01 | wastewater                | DEHP, DMP, DEP, DBP          | 1000 mg/L, 3d, 100%       | [45]       |
| Achromobacter sp. RX      | activated sludge           | DEHP                          | 300 mg/L, 4d, 96%         | [14]       |
| Pseudoxanthomonas sp. N4  | Denitrification biofilter reactor | DEHP                        | 1250 mg/L, 5d, 30%        | [33]       |
| Gordonia sp. Lff          | river sludge               | DEHP, DMP, DEP, DBP, DOP     | 2000 mg/L, 3d, 91.4%      | [37]       |
| Gordonia alkanivorans YC-RL2 | soil                      | DEHP, DCP, DEP, DMP, DBP     | 1000 mg/L, 7d, 68.3%      | [30]       |
| Gordonia sp. GZ-YC7       | landfill soil              | DEHP, DEP, DPPrP, DBP, BBP, DnOP, DnNP | 4000 mg/L, 5d, 70.71%; This study |
|                           |                            |                               | 2000 mg/L, 3d, 87.11%;   |            |
|                           |                            |                               | 1000 mg/L, 2.5d, 100%;   |            |
|                           |                            |                               | 500 mg/L, 1d, 100%;      |            |
|                           |                            |                               | 200 mg/L, 1d, 100%       |            |

1Red represents PAEs with long side chains; green represents PAEs with short side chains; blue represents PAEs with cyclic side chains.
**Table 2.** Alignment results of esters in *Gordonia* sp. GZ-YC7

| Type | Ester | Strain | Gene accession number in GZ-YC7 | Similarity (%) | References |
|------|-------|--------|---------------------------------|----------------|------------|
| Ⅰ    | EstS1 | *Sulfobacillus acidophilus* DSM10332 | GZ-YC7GL004260 | 31.62 | [2] |
| Ⅰ    | EstSP1 | *Sphingomonas glacialis* PAMC 26605 | GZ-YC7GL000423 | 46.62 | [46] |
| Ⅱ    | mehP | *Gordonia* sp. P8219 | GZ-YC7GL001189 | 98.94 | [40] |
| Ⅱ    | patE | *Rhodococcus jostii* RHA1 | GZ-YC7GL000190 | 38.92 | [47] |
| Ⅲ    | EstG | *Sphingobium* sp. SM42 | GZ-YC7GL000793 | 40.37 | [48] |
| Ⅲ    | CarEW | *Bacillus* sp. K91 | GZ-YC7GL000873 | 37.16 | [49] |
| Ⅲ    | pehA | *Arthrobacter* sp. ZJUTW | GZ-YC7GL001562 | 34.07 | [29] |
4. Discussion

Gordonia sp. GZ-YC7 exhibited the highest degradation efficiency among all reported strains under the DEHP concentration lower than 1000 mg/L. Furthermore, GZ-YC7 could keep high PAEs degradation efficiency even in a high PAEs concentration of 4 g/L under optimal conditions (pH 7.0, 30 °C, and 6 % NaCl), suggesting the highest tolerance to PAEs among all reported strains. The diversity of esterase and metabolic pathway predicted in Gordonia sp. GZ-YC7 genome, also support its great potential for application in the remediation of contaminated soil.

Supplementary Materials: The following supporting information can be downloaded at: www.mdpi.com/xxx/s1, Supplementary A: figures; Supplementary B: tables.

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References

1. Li, Q., Zeng, A., Jiang, X., Gu, X. Are microplastics correlated to phthalates in facility agriculture soil? J Hazard Mater 2021, 412, 125164. doi:10.1016/j.jhazmat.2021.125164
2. Zhang, X. Y., Fan, X., Qiu, Y. J., Li, C. Y., Xing, S., Zheng, Y. T., Xu, J. H. Newly identified thermostable esterase from Sulfobacillus acidophilus: properties and performance in phthalate ester degradation. Appl Environ Microbiol 2014, 80(22), 6870-6878. doi: 10.1128/AEM.02072-14
3. Huang, M.Z., Ma, Y.W., Wang, Y., Wan, J.Q., Zhang, H.P. The fate of di-n-butyl phthalate in a laboratory-scale anaerobic/anoxic/oxic wastewater treatment process. Bioreour. Technol 2010, 101, 7767–7772.
4. Chi, J. Phthalate acid esters in Potamogeton crispus L. from Haihe River, China. Chemosphere 77, 48–52. doi: 10.1016/j.chemosphere.2009.03.050
5. Sun, J., Pan, L., Tsang, D. C. W., Zhan, Y., Zhu, L., Li, X. Organic contamination and remediation in the agricultural soils of China: A critical review. Science of The Total Environment 2018, 615, 724–740. doi:10.1016/j.scitotenv.2018.10.057
6. Zeng, L. J., Huang, Y. H., Chen, X. T., Chen, X. H., Mo, C. H., Feng, Y. X., Lv, H. X., Xiang, L., Li, Y. W., Li, H., Cai, Q., Y., Wong, M. H. Prevalent phthalates in air-soil-vegetable systems of plastic greenhouses in a subtropical city and health risk assessments. Science of The Total Environment 2020, 743, 140755. doi:10.1016/j.scitotenv.2020.140755
7. Hou, H., Min, Y., Liu, X., Wang, P., Zhou, Z., Liu, D., 2021. Occurrence and migration of phthalates in adhesive materials to fruits and vegetables. J Hazard Mater 418, 126277. doi: 10.1016/j.jhazmat.2021.126277
8. Zhou, B., Zhao, L., Sun, Y., Li, X., Weng, L., Li, Y. Contamination and human health risks of phthalate esters in vegetable and crop soils from the Huang-Huai-Hai region of China. Sci Total Environ 2021, 778, 146281. doi:10.1016/j.scitotenv.2021.146281
9. João Pinto da Costa. Micro- and nanoplastics in the environment: Research and policymaking. Curr Opin Environ Sci Health 2018, 1, 12-16. doi: 0.1016/j.coesh.2017.11.002
10. Cheang, C. C., Ma, Y., Fok, L. Occurrence and Composition of Microplastics in the Seabed Sediments of the Coral Communities in Proximity of a Metropolitan Area. Int J Environ Res Public Health 2018, 15(10): 2270. doi: 10.3390/ijerph15102270.
11. Nam, S., Lee, J., An, Y. Towards understanding the impact of plastics on freshwater and marine microalgae: A review of the mechanisms
12. Yang, T., Ren, L., Jia, Y., Fan, S., Wang, J., Wang, J., Yan, Y. Biodegradation of Di-(2-ethylhexyl) Phthalate by Rhodococcus ruber YC-YT1 in Contaminated Water and Soil. Int J Environ Res Public Health 2018, 15(5), 964. doi:10.3390/ijerph15050964

13. Liu, J., Lv, M., Sun, A., Ding, J., Wang, Y., Chang, X., Chen, L. Exposure to microplastics reduces the bioaccumulation of sulfamethoxazole but enhances its effects on gut microbiota and the antibiotic resistome of mice. Chemosphere 2022, 294, 133810. doi:10.1016/j.chemosphere.2022.133810

14. Wang, P., Gao, J., Zhao, Y., Zhang, M., Zhou, S. Biodegradability of di-(2-ethylhexyl) phthalate by a newly isolated bacterium Achromobacter sp. RX. Sci Total Environ 2021, 755(Pt 1), 142476. doi:10.1016/j.scitotenv.2020.142476

15. Ren, L., Lin, Z., Liu, H., Hu, H. Bacteria-mediated phthalic acid esters degradation and related molecular mechanisms. Appl Microbiol Biotechno 2018, 102(3), 1085-1096. doi:10.1007/s00253-017-8687-5

16. Ma, T., Wei, Z., Chen, L. K., Wu, L., Luo, Y. Toxicity effects of di-(2-ethylhexyl) phthalate to Eisenia fetida at enzyme, cellular and genetic levels. Plos One 2017, 12(3), e0173957. doi: 10.1371/journal.pone.0173957.

17. Natale, C. D., Onesto, V., Lagreca, E., Vecchione, R., Netti, P. A. Tunable Release of Curcumin with an In Silico-Supported Approach from Mixtures of Highly Porous PLGA Microparticles. Materials 2020, 13(8), 1807. doi: 10.3390/ma13081807

18. Huang, H., Zhang, X. Y., Chen, T. L., Zhao, Y. L., Xu, D. S., Bai, Y. P. Biodegradation of Structurally Diverse Phthalate Esters by a Newly Identified Esterase with Catalytic Activity toward Di(2-ethylhexyl) Phthalate. J Agric Food Chem 2019, 67(31), 8548-8558. doi: 10.1021/acs.jafc.9b02655.

19. Tran, H. T., Lin, C., Bui, X., Nguyen, M. K., Cao, N. D. T., Mukhtar, H., Hoang, H. G., Varjani, S., Ngo, H. H., Nghiem, L. D. Phthalates in the Environment: Characteristics, Fate and Transport, and Advanced Wastewater Treatment Technologies. Bioresour Technol 2022, 344, 126249. doi:10.1016/j.biortech.2021.126249

20. Ding, J., Zhou, Y., Wang, C., Peng, Z., Huang, Z. Development of a whole-cell biocatalyst for diisobutyl phthalate degradation by functional display of a carboxylesterase on the surface of Escherichia coli. Microb Cell Fact 2020, 19(1), 114. doi:10.1186/s12934-020-01373-6.

21. Jin, D., Bai, Z., Chang, D., Hoefel, D., Jin, B., Wang, P., Wei, D., Zhuang, G. Biodegradation of di-n-butyl phthalate by an isolated Gordonia sp strain QH-11: Genetic identification and degradation kinetics. J Hazard Mater 2012, 221-222, 80-85. doi:10.1016/j.jhazmat.2012.04.010.

22. Feng, N. X., Feng, Y. X., Liang, Q. F., Chen, X., Xiang, L., Zhao, H. M., Liu, L. B., Cao, G., Li, Y. W., Li, H., Cai, Q. Y., Mo, C. H. Complete biodegradation of di-n-butyl phthalate (DBP) by a novel Pseudomonas sp. YJB6. Sci Total Environ 2021, 761, 143208. doi: 10.1016/j.scitotenv.

23. He, Z., Niu, C., Lu, Z. Individual or synchronous biodegradation of di-n-butyl phthalate and phenol by Rhodococcus ruber strain DP-2. J Hazard Mater 2014, 273, 104-109. doi:10.1016/j.jhazmat.2014.03.033.

24. Zhao, H. M., Hu, R. W., Du, H., Xin, X. P., Li, Y. W., Li, H., Cai, Q. Y., Mo, C. H., Liu, J. S., Zhou, D. M., Wong, M. H., He, Z. L. Functional genomic analysis of phthalate acid ester (PAE) catabolism genes in the versatile PAE-mineralising bacterium Rhodococcus sp. 2G. Sci Total Environment 2018a, 640-641, 646-652. doi:10.1016/j.scitotenv.2018.05.337.

25. Li, X., Zhou, Y., Zhang, J. Status and associated human health risk of zinc accumulation in agricultural soils across China. Process Saf Environ Prot 2021, 146, 867-876. doi:10.1016/j.psep.2020.12.017

26. Chao, W. L., Cheng, C. Y. Effect of introduced phthalate-degrading bacteria on the diversity of indigenous bacterial communities during di-(2-ethylhexyl) phthalate (DEHP) degradation in a soil microcosm. Chemosphere 2007, 67(3), 482-488. doi:10.1016/j.chemosphere.2006.09.048

27. Ren, L., Jia, Y., Ruth, N., Qiao, C., Wang, J., Zhao, B., Yan, Y. Biodegradation of phthalic acid esters by a newly isolated Mycobacterium sp. YC-RL4 and the bioprocess with environmental samples. Environ Sci Pollut Res Int 2016, 23(16), 16609-16619. doi:10.1007/s11356-016-6829-4

28. Wang, J., Zhang, M.Y., Chen, T., Zhi, Y., Thai, Y., Luo, Y. M., Christie, P. Isolation and Identification of a Di-(2-Ethylhexyl) Phthalate-
Degrading Bacterium and Its Role in the Bioremediation of a Contaminated Soil. Pedosphere 2015, 25(2), 202-211. doi:10.1016/s1002-0160(15)60005-4

29. Liu, T., Li, J., Qiu, L., Zhang, F., Linhardt, R. J., Zhong, W. Combined genomic and transcriptomic analysis of the dibutyl phthalate metabolic pathway in Arthrobacter sp. ZJUTW. Biotechnol Bioeng 2020, 117(12), 3712-3726. doi:10.1002/bit.27524

30. Nahurira, R., Ren, L., Song, J., Jia, Y., Wang, J., Fan, S., Wang, H., Yan, Y. Degradation of Di(2-Ethylhexyl) Phthalate by a Novel Gordonia alkanivorans Strain YC-RL2. Curr Microbiol 2017, 74(3), 309-319. doi:10.1007/s00284-016-1159-9

31. Li, J., Zhang, J., Yadav, M. P., Li, X. Biodegradability and biodegradation pathway of di-(2-ethylhexyl) phthalate by Burkholderia pyrrocinia B1213. Chemosphere 2019, 225, 443-450. doi:10.1016/j.chemosphere.2019.02.194

32. Wen, Z. D., Gao, D. W., Wu, W. M. Biodegradation and kinetic analysis of phthalates by an Arthrobacter strain isolated from constructed wetland soil. Appl Microbiol Biotechnol 2014, 98(10), 4683-4690. doi:10.1007/s00253-014-5568-z

33. Meng, X., Niu, G., Yang, W., Cao, X. Di-(2-ethylhexyl) phthalate biodegradation and denitrification by a Pseudoxanthomonas sp. Strain. Bioresour. Technol 2015, 180, 356-359.

34. Zhao, H. M., Du, H., Lin, J., Chen, X. B., Li, Y. W., Li, H., Cai, Q. Y., Mo, C. H., Qin, H. M., Wong, M. H. Complete degradation of the endocrine disruptor di-(2-ethylhexyl) phthalate by a novel Agromyces sp. MT-O strain and its application to bioremediation of contaminated soil. Sci Total Environ 2016, 562, 170-178. doi:10.1016/j.scitotenv.2016.03.171

35. Zhao, H. M., Hu, R. W., Chen, X. X., Chen, X. B., Lu, H., Li, Y. W., Wong, M. H. Biodegradation pathway of di-(2-ethylhexyl) phthalate by a novel Rhodococcus pyridinivorans XB and its bioaugmentation for remediation of DEHP contaminated soil. Sci Total Environ 2018b, 640-641, 1121-1131. doi:10.1016/j.scitotenv.2018.05.334

36. Chen, S., Geng, P., Xiao, Y., Hu, M. Bioremediation of beta-cypermethrin and 3-phenoxybenzaldehyde contaminated soils using Streptomyces aureus HP-S-01. Appl Microbiol Biotechnol 2012, 94(2), 505-515. doi:10.1007/s00253-011-3640-5

37. Wang, Y., Zhan, W., Ren, Q., Cheng, S., Wang, J., Ma, X., Zhang, C., Wang, Y., 2019. Biodegradation of di-(2-ethylhexyl) phthalate by a newly isolated Gordonia sp. and its application in the remediation of contaminated soils. Sci Total Environ 689, 645-651. doi:10.1016/j.scitotenv.2019.06.459

38. Zhang, H., Lin, Z., Liu, B., Wang, G., Weng, L., Zhou, J., Hu, H., He, H., Huang, Y., Chen, J., Ruth, N., Li, C., Ren, L., 2020. Bioremediation of di-(2-ethylhexyl) phthalate contaminated red soil by Gordonia terrae RL-JC02: Characterization, metabolic pathway and kinetics. Sci Total Environ 733, 139138. doi:10.1016/j.scitotenv.2020.139138

39. Livny, J., Brencic, A., Lory, S., Waldor, M. K. Identification of 17 Pseudomonas aeruginosa sRNAs and prediction of sRNA-encoding genes in 10 diverse pathogens using the bioinformatic tool sRNAPredict2. Nucleic Acids Res 2006, 34(12), 3484-93. doi: 10.1093/nar/gkl453.

40. Nishioka, T., Iwata, M., Imaoka, T., Mutoh, M., Egashira, Y., Nishiyama, T., Shin, T., Fujii, T. A Mono-2-Ethylhexyl Phthalate Hydrolase from a Gordonia sp. that is Able To Dissimilate Di-2-Ethylhexyl Phthalate. Appl Environ Microbiol 2006, 72(4), 2394-2399. doi:10.1128/AEM.72.4.2394-2399.2006.

41. Fan, S., Wang, J., Li, K., Yang, T., Jia, Y., Zhao, B., Yan, Y. Complete genome sequence of Gordonia sp. YC-JH1, a bacterium efficiently degrading a wide range of phthalic acid esters. J Biotechnol 2018, 279, 55-60. doi: 10.1016/j.jbiotec.2018.05.009.

42. Chen, F., Chen, Y., Chen, C., Feng, L., Dong, Y., Chen, J., Lan, J., Hou, H., 2021. High-efficiency degradation of phthalic acid esters (PAEs) by Pseudarthrobacter defluvii E5: Performance, degradative pathway, and key genes. Sci Total Environ, 794, 148719. doi:10.1016/j.scitotenv.2021.148719

43. Nahurira, R., Wang, J., Yan, Y., Jia, Y., Fan, S., Khokhar, I., Eltoukhy, A. In silico genome analysis reveals the metabolic versatility and biotechnology potential of a halotolerant phthalic acid esters degrading Gordonia alkanivorans strain YC-RL2. AMB Express 2019, 9(1), 21. doi:10.1186/s13568-019-0733-5

44. Zhang, J., Zhang, C., Zhu, Y., Li, J., Li, X. Biodegradation of seven phthalate esters by Bacillus mojavensis B1811. International Biodeterioration & Biodegradation 2018, 132, 200-207. doi:10.1016/j.ibiod.2018.04.006
45. Ren, L., Wang, G., Huang, Y., Guo, J., Li, C., Jia, Y., Chen, S., Zhou, J., Hu, H. Phthalic acid esters degradation by a novel marine bacterial strain *Mycolicibacterium phocaicum* RL-HY01: Characterization, metabolic pathway and bioaugmentation. Sci Total Environ 2021, 791, 148303. doi:10.1016/j.scitotenv.2021.148303

46. Hong, D. K., Jang, S. H., Lee, C. Gene cloning and characterization of a psychrophilic phthalate esterase with organic solvent tolerance from an Arctic bacterium *Sphingomonas glacialis* PAMC 26605. Journal of Molecular Catalysis B: Enzymatic 2016, 133, S337-S345. doi:10.1016/j.molcatb.2017.02.004

47. Hara, H., Stewart, G. R., Mohn, W. W. Involvement of a novel ABC transporter and monoalkyl phthalate ester hydrolase in phthalate ester catabolism by *Rhodococcus jostii* RHA1. Appl Environ Microbiol 2010, 76(5), 1516-1523. doi:10.1128/AEM.02621-09

48. Whangsuk, W., Sungkeeree, P., Nakasiri, M., Thiengmag, S., Mongkolsuk, S., Loprasert, S. Two endocrine disrupting dibutyl phthalate degrading esterases and their compensatory gene expression in *Sphingobium* sp. SM42. International Biodeterioration & Biodegradation 2015, 99, 45-54. doi:10.1016/j.ibiod.2014.12.006

49. Lei, B., Ding, J., Wang, C., Xie, Z., Li, J., Yang, Y., Mu, Y., Tang X., Xu, B., Zhou, J., Huang, Z. Properties of a Newly Identified Esterase from *Bacillus* sp. K91 and Its Novel Function in Diisobutyl Phthalate Degradation. Plos One 2015, 10(3). e0119216. doi:10.1371/journal.pone.0119216