Has the Bacterial Biphenyl Catabolic Pathway Evolved Primarily To Degrade Biphenyl? The Diphenylmethane Case

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In this work, we have compared the ability of Pandoraea pnomenusa B356 and of Burkholderia xenovorans LB400 to metabolize diphenylmethane and benzophenone, two biphenyl analogs in which the phenyl rings are bonded to a single carbon. Both chemicals are of environmental concern. P. pnomenusa B356 grew well on diphenylmethane. On the basis of growth kinetics analyses, diphenylmethane and biphenyl were shown to induce the same catabolic pathway. The profile of metabolites produced during growth of strain B356 on diphenylmethane was the same as the one produced by isolated enzymes of the biphenyl catabolic pathway acting individually or in coupled reactions. The biphenyl dioxygenase oxidizes diphenylmethane to 3-benzylcyclohexa-3,5-diene-1,2-diol very efficiently, and ultimately this metabolite is transformed to phenylacetic acid, which is further metabolized by a lower pathway. Strain B356 was also able to cometabolize benzophenone through its biphenyl pathway, although in this case, this substrate was unable to induce the biphenyl catabolical pathway and the degradation was incomplete, with accumulation of 2-hydroxy-6,7-dioxo-7-phenylheptanoic acid. Unlike strain B356, B. xenovorans LB400 did not grow on diphenylmethane. Its biphenyl pathway enzymes metabolized diphenylmethane, but they poorly metabolize benzophenone. The fact that the biphenyl catabolic pathway of strain B356 metabolized diphenylmethane and benzophenone more efficiently than that of strain LB400 brings us to postulate that in strain B356, this pathway evolved divergently to serve other functions not related to biphenyl degradation.

Many investigations have shown that the bacterial biphenyl catabolic pathway enzymes, especially biphenyl dioxygenase (BPDO), which initiates the degradation process, are very versatile (1). The biphenyl pathway, also called the upper pathway, comprises four enzymatic steps that transform biphenyl into benzoic acid, which is further metabolized by a lower pathway (Fig. 1).

Aside from its ability to metabolize polychlorinated biphenyls (PCBs) (1), BPDO metabolizes many biphenyl analogs (2–7) to generate hydroxylated aromatics. BPDO is composed of three components (Fig. 1). The catalytic component, which is a Rieske-type dioxygenase (RO) (BphA), is a heteroexamer made up of three α (BphA) and three β (BphE) subunits. The other two components are ferredoxin (BphF) and ferredoxin reductase (BphG), both of which are involved in electron transfer from NADH to BphA. The catalytic center of the enzyme is located on the C-terminal portion of the BphA α subunit, which also carries the major structural determinants for substrate specificity (8). There are three phylogenetically distinct clusters of BphAαs (9–11), and the structure of a representative BphAα (also called BphA1A2) from each of these three clusters has now been elucidated. Thus, the Protein Data Bank (PDB) coordinate file for Burkholderia xeno-

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genes KF707 BPDO and variants derived from it were able to metabolize diphenylmethane. However, the metabolites produced have not been identified, and the steady-state kinetics of these BPDOs toward diphenylmethane were not determined. On the other hand, the ability of BPDO to metabolize benzophenone has never been examined.

While examining the ability of the biphenyl catabolic enzymes of *P. pnomenusa* B356 and of *B. xenovorans* LB400 to metabolize these two chemicals, we unexpectedly found that strain B356 metabolizes both benzophenone and diphenylmethane, both substrates are metabolized by the same catabolic pathway. This led us to postulate that in strain B356, the metabolites produced from benzophenone by biphenyl-induced cells of strain B356 and of *B. xenovorans* LB400 by diphenylmethane and benzophenone was assessed by monitoring the amount of 4-chlorobenzoate produced from 4-chlorobenzil as described previously (14). We also evaluated the ability of wild-type strains B356 and LB400 to grow on diphenylmethane or benzophenone as the sole growth substrate. Cells grown overnight in LB broth were washed twice in MM30 and suspended in MM30 basal medium to an optical density at 600 nm (OD₆₀₀) of 0.5. This cell suspension was used (100 μl) to inoculate 20 ml of MM30 containing 2 mM biphenyl or diphenylmethane. The cultures were incubated with shaking at 28°C. Cell growth was monitored by determining the CFU. We also used the Bioscreen C system (Growth Curves USA, Piscataway, NJ) to compare the growth kinetics of strain B356 according to the substrate diphenylmethane or biphenyl. In order to prepare the inocula for the Bioscreen C experiments, cells were grown on 1 mM diphenylmethane or biphenyl or on 30 mM sodium acetate for 16 to 18 h, they were washed twice in MM30, and the cells were suspended in the same medium to an OD₆₀₀ of 0.08. Each well of the Bioscreen C microplate contained 237.5 μl of MM30 supplemented with 1 mM biphenyl or diphenylmethane added in 30 μl dimethyl sulfoxide (DMSO) or 30 mM sodium acetate added in 30 μl water, and they were inoculated with 12.5 μl of the suspension described above. A series of cultures also contained 3-chlorobenzoic acid (2 mM) or 3-chlorocatechol (0.2 mM), each added in 5 μl DMSO. Control cultures with no substrate and uninoculated cultures were also run in the experiments. The cultures were incubated at 28°C and set at low revolution. Both biphenyl and diphenylmethane are poorly soluble in water, but when added at a concentration of 1 mM, the nonsoluble portion of the substrate did not interfere with the OD readings during the Bioscreen C experiments. Each set of cultures was run in triplicate. Growth was also monitored by determining the CFU at various intervals of time during the Bioscreen C experiments.

**Analysis of the metabolites produced from diphenylmethane and benzophenone by strains B356 and LB400 and by enzymes of their biphenyl catabolic pathway.** The metabolites produced during growth of strain B356 on 2 mM diphenylmethane in 20 ml MM30 were extracted with ethyl acetate at neutral pH and at pH 4 from the supernatant of 22-h-old cultures. They were then treated with butylboronate (nBuB) or N,O-bis(trimethylsilyl) trifluoroacetamide (TMS) for gas chromatography-mass spectrometry (GC-MS) analyses according to previously described protocols (30). A similar protocol was used to examine the metabolites produced from benzophenone by biphenyl-induced cells of strain B356. However, in this case, the cells were grown on biphenyl to support growth of strains B356 and LB400 and to induce their biphenyl catabolic pathway. Induction of the biphenyl catabolic pathway of strains B356 and LB400 by diphenylmethane and benzophenone was assessed by monitoring the amount of 4-chlorobenzoate produced from 4-chlorobenzil as described previously (14).

**Materials and Methods**

**Bacterial strains, plasmids, chemicals, and general protocols.** *Escherichia coli* DH115 (23) and *E. coli* C41(DE3) (24) were used in this study. Wild-type strains *P. pnomenusa* B356 and *B. xenovorans* LB400 were described previously (25, 26). BphAE4 is a mutant of BphAED400, described previously (8), which was obtained by substitution at two residues, Thr335Ala and Phe336Met. This mutant exhibits an expanded substrate range compared to that of the parent enzyme. Most plasmids used in this study were described previously and are listed in Table 1. Plasmids pET14b[B356-bphF] and pET14b[B356-bphG] carry the genes encoding BphF₃56 and BphG₃56, respectively, and they were prepared by subcloning the respective genes from pQE31[B356-bphF] and pQE31[B356-bphG] (27) into pET14b. The culture media used were *Luria-Bertani* (LB) broth (28), basal medium M9 (28), or minimal mineral medium 30 (MM30) (29) amended with various sources of carbon and antibiotics, depending on the experiment. DNA general protocols were done according to Sambrook et al. (28). Diphenylmethane and benzophenone (99% pure) were from Sigma-Aldrich.

**Assays to assess ability of diphenylmethane and benzophenone to support growth of strains B356 and LB400 and to induce their biphenyl catabolic pathway.** Induction of the biphenyl catabolic pathway of strains B356 and LB400 by diphenylmethane and benzophenone was assessed by monitoring the amount of 4-chlorobenzoate produced from 4-chlorobenzil as described previously (14).

**Table 1**

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Protein(s) expressed</th>
<th>Reference or source</th>
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<tr>
<td>pET14b[B400-bphAE]</td>
<td>BphAE₄₄₄₀</td>
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</tr>
<tr>
<td>pET14b[p-p-bphAE]</td>
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<td>8</td>
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<tr>
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<td>BphAE₃₅₆</td>
<td>16</td>
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<td>pQE31[B356-bphAE]</td>
<td>BphAE₃₅₆</td>
<td>16</td>
</tr>
<tr>
<td>pDB31[B400-bphFG]</td>
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</tr>
<tr>
<td>pDH131[B400-bphFGBC]</td>
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<td>48</td>
</tr>
<tr>
<td>pET14b[B356-bphF]</td>
<td>BphF₃₅₆</td>
<td>This study</td>
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<tr>
<td>pET14b[B356-bphG]</td>
<td>BphG₃₅₆</td>
<td>This study</td>
</tr>
<tr>
<td>pET14b[B356-bphB]</td>
<td>BphB₃₅₆</td>
<td>49</td>
</tr>
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reach log phase, and then they were suspended at an OD_{600} of 3.0 in M9 medium and incubated at 28°C and 100 rpm for 60 min in the presence of 0.2 mM benzophenone.

Metabolites from diphenylmethane and benzophenone also were analyzed from suspensions of isopropyl-β-D-thiogalactopyranoside (IPTG)-induced whole cells of *E. coli* [pDB31 B356-bphFGBC] or *E. coli* [pYPH31 LB400-bphFGBC] also harboring pQE31[B356-bphAE] or pQE31[LB400-bphaE] according to a previously published protocol (30). The induced cells were suspended at an OD_{600} of 5.0 in 50 mM sodium phosphate buffer, pH 7.0, and the metabolites generated after 30 min of incubation at 37°C were extracted and analyzed by GC-MS.

GC-MS analyses were performed using a Hewlett Packard HP6890 series gas chromatograph interfaced with an HP5973 mass selective detector (Agilent Technologies). The mass selective detector was operated in electron impact (EI) mode and used a quadrupole mass analyzer. Under these conditions, the instrument resolution is 0.1 atomic mass units, which is sufficient to clearly distinguish between two compounds of atomic masses differing by a single atomic mass unit.

Assays to identify diphenylmethane and benzophenone metabolites produced from BphAE_{B356} BphAE_{L4000} and BphAp_{Bip} and to determine their steady-state kinetics. Reconstituted His-tagged BPDO preparations were used in these experiments. His-tagged purified enzyme components were produced and purified by following protocols published previously (31). The enzyme assays were performed in a volume of 200 μl in 50 mM morpholineethanesulfonic (MES) buffer, pH 6.0, at 37°C as described previously (32). For metabolite analyses, the reaction medium was incubated for 10 min and the metabolites were extracted at pH 6.0 with ethyl acetate, and then they were treated with a TMS for GC-MS analyses as described above. The steady-state kinetics were determined by recording the oxygen consumption rates according to a protocol described previously, using a Clarke-type Hansatech model DW1 oxygraph (33). They were determined from three separately prepared purifications of the enzymes.

Purification and NMR analysis of 2,2',3,3'-tetrahydroxybenzophenone. 2,2',3,3'-Tetrahydroxybenzophenone was prepared using a coupled reaction composed of His-tagged purified preparations of B356 BPDO (BphAEFG_{B356}) plus BphBG_{B356}. Each enzyme reaction mixture contained 50 nmol benzophenone, 0.6 nmol of each B356 BPDO component (BphAE_{B356}, BphF_{B356} and BphG_{B356}), 2 nmol BphBG_{B356}, 100 nmol NADH, and 100 nmol NAD in 200 μl (total volume) of 50 mM MES buffer (pH 6.0). The mixture was incubated for 15 min at 37°C and then extracted at pH 6.0 with ethyl acetate. The extract was concentrated 20-fold by evaporation under a stream of nitrogen, and this preparation was dissolved in deuterated acetone at room temperature.

Docking and structure analysis. Dimer AB of BphAE_{L4000} (RCBS Protein Data Bank [PDB] entry 2XXR) and of BphAE_{B356} (PDB entry 3GZX) were used as protein targets, and they were prepared as previously described (16). Ligands were all downloaded as sdf files from PubChem (http://pubchem.ncbi.nlm.nih.gov) and converted into pdb format in Discover Studio Visualizer 2.5. Both proteins and ligands were processed with AutoDockTools to obtain their proper pdbqt format. The searching space for the ligand was centered on mononuclear iron and contained 20 Å in each x, y, and z direction. Autodock 4 (34) with default parameters was used to perform the automatic docking.

**RESULTS**

Growth of *P. promenusa* B356 and *B. xenovorans* LB400 on diphenylmethane and benzophenone. Neither *P. promenusa* B356 nor *B. xenovorans* LB400 grew when benzophenone was used as the sole growth substrate. Using a previously described 4-chlorobiphenyl conversion assay (14), we showed that benzophenone was unable to induce the biphenyl catabolic pathway of both strains. Similarly, diphenylmethane did not serve as the growth substrate for strain LB400 and did not induce its biphenyl catabolic pathway. However, remarkably, strain B356 grew very well in MM30 containing 2 mM diphenylmethane as the sole growth substrate. Under those conditions, CFU values of up to 4 × 10^{9} cells/ml were obtained within 36 h at 28°C. In addition, using the 4-chlorobiphenyl conversion assay (14), we found that a suspension of log-phase cells grown on 2 mM diphenylmethane and adjusted to an OD_{600} of 1.0 produced 329 ± 20 μM 4-chlorobenzoate from 1.25 mM 4-chlorobiphenyl after 2 h of incubation. In comparison, a resting suspension of log-phase cells of strain B356 grown on 2 mM biphenyl and tested under the same conditions produced 330 ± 17 μM 4-chlorobenzoate. This suggested that the biphenyl catabolic pathway of B356 was induced during growth on diphenylmethane. Using the Bioscreen C system, we have compared the growth kinetics of strain B356 according to the substrate used, biphenyl or diphenylmethane. In addition, we have determined the effect of interchanging the growth substrate on growth kinetics (biphenyl or diphenylmethane) to prepare the inocula. Results shown in Fig. 2 clearly demonstrate that strain B356 grows as well on diphenylmethane as on biphenyl, sometimes even better. Furthermore, replacing diphenylmethane with biphenyl to prepare the inoculum for the Bioscreen C experiments did not affect significantly the growth kinetics on diphenylmethane; the lag phases were identical and the growth rates were similar (0.35 ± 0.01 h^{-1} and 0.34 ± 0.02 h^{-1} for diphenylmethane- or biphenyl-induced cells, respectively). Likewise, when biphenyl was used as the growth substrate, the lag phases were very similar whether the inocula were prepared by growing the cells on biphenyl or diphenylmethane, and the growth rates were identical (0.25 ± 0.02 h^{-1}). Conversely, when cells were grown on sodium acetate to prepare the inoculum, the cells grew very poorly whether the Bioscreen C experiments were run using biphenyl or diphenylmethane as the substrate (Fig. 2), exhibiting long lag phases and low growth rates (0.14 h^{-1}), and the CFU count did not exceed 5 × 10^{7} cells/ml at the end of the log phase. We have no data explaining why B356 grew better on diphenylmethane than on biphenyl. This result shows that the combined upper and lower pathways metabolized diphenylmethane more efficiently than biphenyl. However, the facts that interchanging the substrates to prepare the inocula for the Bioscreen C experiments did not affect the growth kinetics and did not prolong the lag phase provide strong evidence that the two substrates were degraded by the same pathway, which they both induced.

As further evidence that growth on diphenylmethane proceeded through the biphenyl catabolic pathway, we examined the effect of adding 3-chlorocatechol and 3-chlorobenzoate to the growth medium. In the Bioscreen C experiments, 0.2 mM 3-chlorocatechol completely inhibited growth of strain B356 on both biphenyl and diphenylmethane (data not shown). In a previous
to metabolize 3-chlorobenzoate into 3-chlorocatechol is induced during growth on diphenylmethane, and 3-chlorocatechol strongly impairs the metabolism of the upper pathway. The fact that both the diphenylmethane and biphenyl degradation pathways responded similarly to the presence of 3-chlorocatechol and 3-chlorobenzoate provides additional evidence that both substrates are metabolized by the same pathway.

Because phenylacetate would be the expected end product if diphenylmethane was metabolized by the enzymes of the upper biphenyl pathway, we have determined whether it could serve as the growth substrate for strains B356 and LB400. In both cases, cells grew very well, reaching CFU values exceeding 10^9 cells/ml within 18 h at 28°C.

**Metabolism of diphenylmethane and benzophenone by *P. phomomenus* B356.** When diphenylmethane-grown cells of strain B356 were inoculated in MM30 containing 2 mM diphenylmethane, the substrate (approximately 40 μmol) was almost completely metabolized after 22 h. The mass spectral features of the major metabolite in the acidic ethyl acetate extract were identical to those of an authentic standard of phenylactic acid (Fig. 3A; spectra are not shown). Based on the area under the GC-MS peak, approximately 15 nmol was present in the 22-h-old cultures. Among the minor metabolites, one peak eluting at 14.76 min (Fig. 3A) exhibited mass spectral features that correspond to those of a *meta*-fission metabolite resulting from the catalytic cleavage of a catechol derivative of diphenylmethane. This metabolite was tentatively identified as 2-hydroxy-6-oxo-7-phenylhepta-2,4-dienoic acid (7-phenyl HODA) \( m/z 376 (M^+) \), \( m/z 361 (M^+ - CH_3) \), \( m/z 333 (M^+ - CH_3 - CO) \), and \( m/z 259 (M^+ - COO(CH_3)_2Si) \) (see Fig. S1 in the supplemental material for the spectrum). This is the metabolite that would be expected if the hydroxylation reaction had occurred on the *ortho-meta* carbons of one benzene ring of diphenylmethane to generate 3-benzylcyclohexa-3,5-diene-1,2-diol. The identity of the acidic metabolite as 7-phenyl HODA is consistent with the fact that a metabolite exhibiting the same mass spectral features was produced from diphenylmethane by resting cells of *E. coli* pQE31[B356-bphAE]+pYH31[LB400-bphFGBC] (Fig. 3A). The *E. coli* resting cell assay also shows that BphB_{B356} and BphC_{LB400} can further metabolize 3-benzylcyclohexa-3,5-diene-1,2-diol generated by BphAE_{B356}.

The acidic extracts of cells of strain B356 grown on diphenylmethane also contained a metabolite whose TMS derivative exhibited a molecular mass at \( m/z 256 \) corresponding to a mono-hydroxy-diphenylmethane (Fig. 3A; spectrum not shown). This metabolite was presumably generated from the corresponding dihydrodiol metabolite during the extraction procedure at pH 4.

In the culture extracts prepared at neutral pH, 3-benzylcyclohexa-3,5-diene-1,2-diol and 2,3-dihydroxydiphenylmethane were detected only in trace amounts (not shown). The fact that only small amounts of acidic and neutral metabolites were detected in the culture medium of 22-h-old cultures shows diphenylmethane metabolism is very efficient, and this is consistent with the rapid growth on this substrate.

Strain B356 was unable to grow on benzophenone. However, biphenyl- or diphenylmethane-grown resting cell suspensions metabolized benzophenone and produced a yellow *meta*-fission metabolite which was detected within minutes after substrate addition. Two metabolites (Fig. 3B) with similar spectral features were detected by GC-MS analysis of their TMS derivatives \( m/z 390 (M^+) \), \( m/z 375 (M^+ - CH_3) \), \( m/z 347 (M^+ - CH_3 - CO) \), \( m/z 331 (M^+ - CH_2CO) \), and \( m/z 319 (M^+ - CH_2CO - CH_3) \) (Fig. 3B).
C-1 carboxylic functions may promote the isomerization of the double bonds at C-2 and C-4.

No 2,3-dioxo-3-phenylpropanoic acid, which presumably is the metabolite produced from 7-phenyl DODA by the phenyl HODA hydrolase (BphD), was detected, even after 1 h of incubation in the presence of the substrate. Therefore, we were unable to obtain evidence that benzenophene metabolism goes beyond the meta-fission reaction. On the other hand, two metabolites (Fig. 3B) that exhibited spectral features that could correspond to pyranol isomers [m/z 462 (M⁺), m/z 447 (M⁺ – CO), m/z 419 (M⁺ – CH₃ – CO), m/z 345 (M⁺ – COO(CH₃)₃Si)] (Fig. 3B; also see Fig. S3 in the supplemental material for the spectrum) were detected. These pyranol isomers may have been generated from a cyclization reaction through intramolecular rearrangements. At this time, there is no documented evidence of spontaneous automerization or cyclization of phenyl HODAs. The fact that the same metabolites were produced by an IPTG-induced resting cell suspension of a 1:1 mixture of recombinant E. coli pQE31[B356-bphAE] + pYH31[LB400-bphFGBC] plus E. coli pQE31[B356-bphC] (not shown) suggests they were produced spontaneously from 7-phenyl DODA in the cell suspensions or during the TMS reaction. However, we cannot exclude other mechanisms of formation involving unspecific enzymatic reactions occurring in both strain B356 and E. coli. Therefore, their production remains unexplained at this time.

**Metabolism of diphenylmethane by purified BphAE<sub>B356</sub>, BphAE<sub>LB400</sub> and BphAE<sub>plasmid</sub>.** To confirm that the biphenyl catalytic enzymes of strain B356 metabolize diphenylmethane efficiently, we have examined the catalytic properties of BphAE<sub>B356</sub> toward diphenylmethane, and we have compared them to those of BphAE<sub>LB400</sub> and BphAE<sub>plasmid</sub>. GC-MS analysis of the nBuB-treated diphenylmethane metabolites revealed two metabolites (Fig. 4A). Since nBuB reacts only with vicinal hydroxyl groups, the mass spectral features (see Fig. S4 in the supplemental material) of the major metabolite (m/z 268 [M⁺], m/z 211 [M⁺ – nBu], m/z 184 [M⁺ – nBuBO], m/z 177 [M⁺ – C₇H₇], m/z 168 [M⁺ – nBuBO₂], m/z 156 [M⁺ – nBuBO – CO]) must correspond to a dihydrodiol. Docking experiments (see below) suggested the oxygenation occurred on the ortho-meta carbons. Therefore, on the basis of its mass spectral features, the major metabolite was tentatively identified as 3-benzylcyclohexa-3,5-diene-1,2-diol.

The minor metabolite contains two pairs of vicinal hydroxyl groups (Fig. 4A). This metabolite presumably was generated from the catalytic ortho-meta hydroxylation of the nonhydroxylated ring of 3-benzylcyclohexa-3,5-diene-1,2-diol. It was tentatively identified as 3-[(5,6-dihydroxycyclohexa-1,3-dien-1-yl)methyl] cyclohexa-3,5-diene-1,2-diol, showing ions at m/z 368 (M⁺), 284 (M⁺ – nBuBO), 200 (M⁺ – nBuBO – nBuBO₂), 177 (M⁺ – nBuBO₂ – C₇H₇), 168 (M⁺ – nBuBO₂ – nBuBO₂), and 156 (M⁺ – nBuBO₂ – nBuBO – CO) (see Fig. S5 in the supplemental material). The fragmentation ion at m/z 177 provides evidence that both rings were oxidized.

The production of 3-benzylcyclohexa-3,5-diene-1,2-diol and 3-[(5,6-dihydroxycyclohexa-1,3-dien-1-yl)methyl]cyclohexa-3,5-diene-1,2-diol as sole products of the B356-BPDO reaction was confirmed by the fact that a coupled reaction composed of purified preparations of BphAE<sub>B356</sub> Plus BphB<sub>B356</sub> generated two metabolites. On the basis of the mass spectra of their TMS derivatives, the major one was identified as 2,3-dihydroxydiphenylmethane (3-benzylbenzene-1,2-diol) and the minor one as 2,2',3,3'-tetra-
hydroxydiphenylmethane \(\{3-[(2,3\text{-dihydroxyphenyl})\text{methyl}\text{]benzene-1,2-diol}\}\) (Fig. 4B). The diagnostically important ions for the TMS-derived 2,3-dihydroxydiphenylmethane (see Fig. S6 in the supplemental material) comprise the molecular ion at \(m/z\) 344 and fragmentation ions at \(m/z\) 329 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{CH}_3)\), 271 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\), 255 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\), 225 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si}/\text{H}1_{1002}\text{CH}_3)\). For the TMS-derived 2,2',3,3'-tetrahydroxydiphenylmethane (see Fig. S7), they are \(m/z\) 520 \((\text{M/H}1_{1001})\), 505 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{CH}_3)\), 447 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\), 431 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\), 342 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\), 253 \((\text{M/H}1_{1001}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si}/\text{H}1_{1002}\text{O(CH}_3)_3\text{Si})\). The vicinity of the two hydroxyl groups in these metabolites was confirmed by the mass spectra of the nBuB derivatives (not shown).

When BphAE\(_{B356}\) was incubated for 10 min with the substrate, it produced approximately the same amount of 3-benzylcyclohexa-3,5-diene-1,2-diol from diphenylmethane as BphAE\(_{B356}\) but no \(3-[(5,6\text{-dihydroxy}cyclohexa-1,3\text{-dien-1-yl})\text{methyl}\text{]cyclohexa-3,5-diene-1,2-diol}\) was produced (Fig. 4A). The metabolic pattern of BphAE\(_{p4}\) toward diphenylmethane was very similar to that of the parental enzyme BphAE\(_{LB400}\) (not shown). Therefore, BphAE\(_{B356}\) was the only one of the three enzymes to further oxidize the dihydroxylated metabolite.

The steady-state kinetic parameters of purified preparations of BphAE\(_{B356}\), BphAE\(_{p4}\), and BphAE\(_{LB400}\) toward diphenylmethane were calculated from the initial oxygen consumption, and they were consistent with the GC-MS analysis. Diphenylmethane was as good a substrate as biphenyl for all three enzymes. Thus, the \(k_{cat}\) and \(k_{cat}/K_m\) values for BphAE\(_{B356}\) (Table 2) were in the range reported for biphenyl (4.3 s\(^{-1}\) and 63 \(\times\) 10\(^3\) M\(^{-1}\) s\(^{-1}\), respectively) (16) when this enzyme was used under the same reaction conditions. Similarly, the \(k_{cat}\) and \(k_{cat}/K_m\) values for BphAE\(_{LB400}\) and for BphAE\(_{p4}\) (Table 2) were very close to those obtained when biphenyl was the substrate and the reactions were run under identical conditions (the reported values were 0.9 s\(^{-1}\) and 41 \(\times\) 10\(^3\) M\(^{-1}\) s\(^{-1}\) for BphAE\(_{LB400}\) respectively, and 1.0 s\(^{-1}\) and 31 \(\times\) 10\(^3\) M\(^{-1}\) s\(^{-1}\) for BphAE\(_{p4}\) respectively) (31).

**Benzophenone metabolism by purified BphAE\(_{B356}\), BphAE\(_{LB400}\), and BphAE\(_{p4}\).** Unlike diphenylmethane, benzophe-
none is metabolized differently by BphAE_{B356} and BphAE_{LB400}. A purified preparation of BphAE_{B356} completely oxidized 50 nmol benzophenone in 10 min to generate a single metabolite (Fig. 4C), tentatively identified as 3-benzoylcyclohexa-3,5-diene-1,2-diol from the GC-MS spectrum of its nBuBO_2 derivative (see Fig. S8 in the supplemental material) (ions at m/z 282 [M^+] and at 225 [M^+ - nBu], 198 [M^+ - nBuBO], 182 [M^+ - nBuBO_2], and 170 [M^+ - nBuBO - CO]). Under the same conditions, BphAE_{LB400} and BphAE_{p4} oxidized only a fraction of the added 50 nmol benzophenone, but both enzymes produced the same metabolite as BphAE_{B356} (Fig. 4C). This is consistent with the steady-state kinetics shown in Table 2, where the turnover rate of the reaction of BphAE_{B356} toward benzophenone was 12 times higher than that for BphAE_{p4}, and the turnover rate of the reaction of BphAE_{LB400} was too low to obtain reliable values.

When BphB_{B356} was used to oxidize 3-benzoylcyclohexa-3,5-diene-1,2-diol, 2,3-dihydroxybenzophenone was produced (not shown) and was identified from the GC-MS spectral features of its TMS derivative (ions at m/z 358 [M^+] and m/z 343 [M^+ - CH_2], 270 [M^+ - (CH_3)_2Si], and 212 [M^+ - (CH_3)_2Si - (CH_3)_2Si]) (see Fig. S9 in the supplemental material). Remarkably, when benzophenone at a concentration of 50 nmol was the substrate for the coupled reaction composed of BphAE_{B356} plus BphB_{B356} it was almost completely metabolized to 2,2',3,3'-tetrahydroxybenzophenone (Fig. 4D), which was identified from its GC-MS and NMR spectra. The other two metabolites, 2,3-dihydroxybenzophenone and 2,2',3- or 2,3,3'-trihydroxybenzophenone, represented less than 1% of total metabolites produced by this reaction. The spectral features of the TMS-treated major metabolite (see Fig. S10 in the supplemental material) was comprised of a molecular ion at m/z 354 plus fragmentation ions at m/z 319 (M^+ - CH_3), 446 [M^+ - (CH_3)_2Si], and 358 [M^+ - (CH_3)_2Si - (CH_3)_2Si]. The position of the hydroxyl groups was confirmed by NMR analysis. The spectrum showed only three signals for six protons between 6.80 and 7.10 ppm. This indicates that there are two identical aromatic rings, and each one contains 3 protons. The chemical shifts for the protons labeled H_6, H_5, and H_4 in Fig. 5 were recorded at 6.824 ppm (triplet) (H_6), at 7.096 (doublet of doublets), and 6.969 (doublet of doublets) (H_4 and H_5). The coupling constant between the three protons (Fig. 5) and the shape of the peaks (1 triplet and 2 doublet of doublets) revealed that the 3 protons of each aromatic ring were vicinal to each other. With this in mind, the only possibility for the structure of the metabolite was 2,2',3,3'-tetrahydroxybenzophenone.

Docking experiments with diphenylmethane and benzophenone. We docked diphenylmethane and benzophenone in the substrate-bound form of BphAE_{B356} and BphAE_{LB400} after removing biphenyl. For both substrates, the conformation of the top-ranked docked molecules in BphAE_{B356} exhibited an orientation that would enable oxygenation of the ortho-meta carbons of the phenyl ring. In both cases, the ortho-meta carbons closely aligned with carbons 2 and 3 of the oxidized ring of biphenyl in the complexed form (Fig. 6A and B). This suggests BphAE_{B356} produces 3-benzoylcyclohexa-3,5-diene-1,2-diol from diphenylmethane and 3-benzoylcyclohexa-3,5-diene-1,2-diol from benzophenone, and it is consistent with the NMR spectral data that confirm that 2,2',3,3'-tetrahydroxybenzophenone was produced from benzophenone by the combined reaction of BphAE_{B356} plus BphB_{B356}.

As shown in Fig. 6A and B, the conformation of benzophenone inside the catalytic pocket of BphAE_{B356} differs from that of diphenylmethane. Therefore, the carbonyl group strongly influences the interaction between the substrate and the amino acid residues that line the catalytic pocket. However, the ortho-meta carbons of the docked benzophenone aligned very well with the reactive carbons of biphenyl in the biphenyl-bound form of BphAE_{B356} (Fig. 6A). In the case of BphAE_{LB400} the top-ranked conformations for benzophenone were at a distance from the catalytic iron that may have allowed catalytic hydroxylation (Fig. 6C). However, the reactive ring did not align as well for BphAE_{LB400} with the reactive ring of biphenyl, and this may explain why BphAE_{LB400} catalyzed the oxygenation of benzophenone poorly. On the other hand, we cannot exclude the possibility that other types of interactions between protein structures and the substrate, which could not be reproduced in the docking experiments, prevent its productive binding to the BphAE_{LB400} catalytic pocket. Since the conformation of benzophenone in the docked BphAE_{B356} structure appears to be more favorable for the catalytic reaction, we have superposed the benzophenone docked structure of BphAE_{B356} on the structure of the biphenyl-bound BphAE_{LB400} after biphenyl removal (Fig. 6D). As shown in the figure, the benzophenone carbonyl oxygen was very close to both Gly321 and Phe336 of BphAE_{LB400}. Therefore, in this conformation of the substrate, the proximity of these two residues relative to the carbonyl oxygen may hinder proper binding. This is consistent with previous observations (8, 16) where Gly321 and Phe336, two resi-
...)
DISCUSSION

Unlike *B. xenovorans* LB400, *P. pnomenusa* B356 grew remarkably well on diphenylmethane, and many observations made in this work are supportive of the idea that it was metabolized by the enzymes of the biphenyl catabolic pathway to produce phenylacetate, which was further metabolized by a lower pathway. Cells of strain B356 produced the same metabolites from diphenylmethane during their growth on this substrate as those isolated from enzymes of the biphenyl catabolic pathway (Fig. 7). Furthermore, the kinetic parameters of BphAEB356 toward diphenylmethane were in the same range as those for biphenyl, suggesting this enzyme metabolizes diphenylmethane during growth on this substrate. It is noteworthy that when purified preparations of B356 BPDO or B356 BPDO plus BphBB356 were used to metabolize diphenylmethane, significant amounts of tetrahydroxylated metabolites were detected in the reaction medium. However, the tetrahydroxylated metabolites were present in very small amounts in cultures of B356 growing on diphenylmethane (not shown). This shows that during growth of strain B356 on diphenylmethane, 3-benzylcyclohexa-3,5-diene-1,2-diol and subsequent metabolites were oxidized very efficiently by the downstream enzymes of the catabolic pathway, thus preventing further BPDO oxidation of the dihydrodiol and the catechol metabolites.

Judging by its steady-state kinetics toward diphenylmethane, BphAELB400 metabolized this substrate as well as biphenyl. In addition, we found that BphCLB400 catalyzed the ring cleavage of 2,3-dihydroxydiphenylmethane, and the strain grew well on phenylacetate. However, unlike strain B356, cells of strain LB400 grown cometically on diphenylmethane plus sodium acetate were unable to metabolize 4-chlorobiphenyl, showing that diphenylmethane did not induce the biphenyl catabolic pathway of strain LB400. Therefore, the inability of diphenylmethane to induce the biphenyl catabolic pathway in strain LB400 most likely is the reason why this strain is unable to grow on this substrate.
Growth kinetics data provided strong evidence that both diphenylmethane and biphenyl induced the same catabolic enzymes during growth of strain B356. Uninduced cells grew poorly on diphenylmethane, which was the case when the inocula were prepared by growing the cells on sodium acetate. However, interchanging the growth substrates to prepare the inocula did not affect the growth kinetics on diphenylmethane or on biphenyl, which strongly suggests that both substrates induced the same catabolic pathway.

Remarkably, although further work will be needed to confirm the conclusion, data suggest that a benzoate pathway is induced when cells are grown on diphenylmethane. This was supported by the fact that 3-chlorobenzoate strongly inhibited growth on diphenylmethane. Based on a previous report (36), in order to inhibit growth on biphenyl, 3-chlorobenzoate must be converted to 3-chlorocatechol, which is a very potent inhibitor of BphC. In this study, we have not elucidated the enzymatic steps involved in phenylacetate catabolism during growth of strain B356 on diphenylmethane. However, since phenylacetate is a key metabolite in the metabolism of many aromatic chemicals, including phenylalanine, most bacteria have the ability to metabolize it through a pathway that proceeds via the formation of a coenzyme A (CoA)-thioester derivative (39). Strain LB400 can metabolize 3- and 4-hydroxyphenylacetate through the homogentisate pathway (40), but the ability of this strain to metabolize phenylacetate through this pathway has not been demonstrated. Focht and Alexander (22, 41) have described a 

Hydrogenomonas isolate that metabolizes diphenylmethane through a pathway that presumably involves the metacleavage of a catechol metabolite, which is further metabolized to generate phenylacetate. Since cells grown on diphenylmethane readily metabolized homogentisate, we postulated that phenylacetate was metabolized via a homogentisic pathway in that 

Hydrogenomonas strain. However, neither the phenylacetyl-coenzyme A nor the homogentisate pathway involves production of a 2,3-catechol derivative, as is the case for the metabolism of benzoic acid in strain B356. Therefore, the growth inhibition caused by 3-chlorobenzoate suggests that diphenylmethane or one of its metabolites induces a pathway involving a benzoate 2,3-dioxygenase, namely, the biphenyl-associated benzoate pathway. Indeed, strain B356 carries two benzoate pathway enzymes, namely the metacleavage of a catechol metabolite, which is further metabolized to generate phenylacetate. Since cells grown on diphenylmethane readily metabolized homogentisate, we postulated that phenylacetate was metabolized via a homogentisic pathway in that 

Hydrogenomonas strain. However, neither the phenylacetyl-coenzyme A nor the homogentisate pathway involves production of a 2,3-catechol derivative, as is the case for the metabolism of benzoic acid in strain B356. Therefore, the growth inhibition caused by 3-chlorobenzoate suggests that diphenylmethane or one of its metabolites induces a pathway involving a benzoate 2,3-dioxygenase, namely, the biphenyl-associated benzoate pathway. Indeed, strain B356 carries two benzoate pathways, only one of which, the one induced when cells are grown on biphenyl, is able to produce 3-chlorocatechol from 3-chlorobenzoate (36).

Few investigations have assessed the ability of the bacterial biphenyl catabolic enzymes to metabolize diphenylmethane or one of its derivatives, such as DDT or 1,1-dichloro-2,2-(4-chlorophenyl)ethane (DDD) (16, 19, 42–44), but no studies have examined the ability of biphenyl-degrading bacteria to grow on diphenylmethane. Therefore, strain B356 might not be the only one with the ability to use the biphenyl catabolic pathway to grow on this substrate.

Neither strain B356 nor LB400 could use benzoephene as a growth substrate; however, BphAE_{B356} metabolized this diphenylmethane analog significantly more efficiently than BphAE_{LB400}. Structural analysis of the docked substrate showed that residues Phe336 and Gly321 were partly responsible for preventing productive binding of this substrate with BphAE_{LB400}. Structural analyses of substrate-bound crystals and docking experiments have shown that the combined effect of these two substitutions increases the space required in the catalytic pocket to accommodate bulky substrates (8, 16). The replacement of Thr335 of BphAE_{LB400} by Gly333 in BphAE_{B356} produces a similar effect (16). However, other structural features that our docking experiments could not identify also may have an influence on the ability of the enzyme to metabolize this substrate, since, on the basis of their steady-state kinetics, BphAE_{B356} metabolized benzophene more efficiently than BphAE_{LB400}, which is a doubly substituted Thr335Ala/Phe336Met mutant of BphAE_{LB400}.

Generally, the investigations related to the biphenyl-degrading bacteria during the last 5 decades were initiated with the objective of designing a biological process to degrade PCBs and other chlorinated aromatics, such as chlorodibenzofurans (45, 46). These bacteria were obtained by enrichments on biphenyl, and traditionally it was believed that the four-enzymatic-step biphenyl pathway evolved in bacteria primarily to transform biphenyl into benzoic acid. However, data presented in the current and previous works (14) bring us to question this belief. Biphenyl is a naturally occurring chemical, but it is not universally distributed in nature. It is noteworthy that bacteria carrying the biphenyl catabolic pathway enzymes were obtained from pristine soils not exposed to biphenyl or chlorobiphenyls (10).

On the basis of their primary amino acid sequences, BphAE_{B356} and BphAE_{LB400} belong to distinct phylogenetic clusters of BphAE (9–11). Recent data suggested that each has acquired a distinct PCB-degrading pattern (9, 15), as well as distinct abilities to metabolize simple flavonoids (14), DDT (16), or DDD (43). In this work, we found that BphAE_{B356} and BphAE_{LB400} metabolized diphenylmethane similarly but benzophenone and 2,3-dihydroxybenzophene very differently. Furthermore, the fact that the biphenyl catabolic pathway of strain B356 is inducible by diphenylmethane allows the strain to use it as a growth substrate. As a result, under natural conditions, strain B356 has the potential to metabolize or cometabolize many chemicals that strain LB400 metabolizes poorly.

Although the enzymes of the peripheral pathways, such as those of the biphenyl catabolic pathway, have evolved to metabolize a broad range of substrates, on the basis of the observations made in this work and a previous one (14), we postulate that the biphenyl catabolic pathway enzymes have evolved divergently in bacteria, in such a way that each phylogenetic branch has specialized to play distinct ecophysiological functions with regard to chemicals naturally found in nature.

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