Importance of Different tfd Genes for Degradation of Chloroaromatics by Ralstonia eutropha JMP134†

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The tfdCDEF and tfdCDEF genes modules of plasmid pJP4 of Ralstonia eutropha JMP134 encode complete sets of functional enzymes for the transformation of chlorocatechols into 3-oxoadipate, which are all expressed during growth on 2,4-dichlorophenoxyacetate (2,4-D). However, activity of tfd1-encoded enzymes was usually higher than that of tfdR-encoded enzymes, both in the wild-type strain grown on 2,4-D and in 3-chlorobenzoate-grown derivatives harboring only one tfd gene module. The tfdD1-encoded chloromuconate cycloisomerase exhibited special kinetic properties, with high activity against 3-chloromuconate and poor activity against 2-chloromuconate and unsubstituted muconate, thus explaining the different phenotypic behaviors of R. eutropha strains containing different tfd gene modules. The enzyme catalyzes the formation of an equilibrium between 2-chloromuconate and 5-chloro-2-chloromuconolactone and very inefficiently catalyzes dehalogenation to form trans-dienelactone as the major product, thus differing from all (chloro)muconate cycloisomerases described thus far.

†For a commentary on this article, see page 4049 in this issue.
by transposon mutagenesis resulted in mutants no longer capable of growing on 2,4-D (8), suggesting that the products of these genes play a major role in the metabolism of this substrate. In contrast, interruption of tfdF resulted in only a slight retardation of growth. The critical importance of the tfdF-encoded chloromuconate cycloisomerase is supported by the fact that diverse attempts to purify chloromuconate cycloisomerase from strain JMP134 resulted in purification of only TfdD (17, 21). In the present investigation, the importance of the different tfd-encoded enzymes, chlorocatechol 1,2-dioxygenase, chloromuconate cycloisomerase, dienelactone hydrolase, and maleylacetate reductase, for growth of *R. eutropha* on chloroaromatics was characterized biochemically.

**MATERIALS AND METHODS**

**Bacterial strains.** The 2,4-D-degrading organism *R. eutropha* JMP134 was isolated by Don and Pemberton (7). *R. eutropha* JMP222 is a derivative of strain JMP134, cured of plasmid pJP4. *R. eutropha* JMP222(pBBR1M-I) and *R. eutropha* JMP222(pBBR1M-II) are derivatives of strain JMP222 containing either module I (pBBR1M-I) or module II chlorocatechol genes as described recently (34).

**Culture conditions and preparation of cell extracts.** Growth in liquid culture was performed in the mineral salts medium described by Dorn et al. (9) containing 50 mM phosphate buffer (pH 7.4). The medium was supplemented with a carbon source, usually at 2.5 or 5 mM. Cells were grown in fluted Erlenmeyer flasks, incubated at 30°C on a rotary shaker at 150 rpm. Growth was monitored by absorbance, incubated at 30°C, except for catechols and substituted catechols, which were added at initial concentrations of 0.2 mM. 3-Chloromuconate, as the substrate for chloromuconate dioxygenase, was used.

**Enzyme assays.** Muconolactone isomerase (EC 5.3.3.4) was assayed as described by Prucha et al. (40) in 50 mM sodium phosphate (pH 7.5) with 0.1 mM (45,55/4R,5R)-5-chloro-3-methylmuconolactone as the substrate. Accumulation of 3-methyl-dienelactone was analyzed spectrophotometrically at 270 nm (ε₃₃₀/μmol/lactone = 15,200 M⁻¹ cm⁻¹). Catechol 1,2-dioxygenase (EC 1.13.11.1), chlorocatechol dioxygenase, muconate cycloisomerase (EC 5.5.1.1), chloromuconate cycloisomerases (EC 5.5.1.7), and dienelactone hydrolase (EC 3.1.1.45) were measured as previously described (10, 43, 44) with catechol, 3-chlorocatechol, muconate, 3-chloromuconate, or cis-dienelactone as the substrate. Substrate concentrations in the enzymatic tests were usually 0.05 mM, except for catechols and substituted catechols, which were added at initial concentrations of 0.2 mM. Chloromuconate, as the substrate for chloromuconate cycloisomerase, was prepared in situ from 4-chlorocatechol by using TfdC that was partially purified by anion-exchange chromatography and that was free of any interfering enzyme activity. Activity of partially purified chloromuconate cycloisomerases was determined in the presence of an excess of dienelactone hydrolase. Generally, TfdE, partially purified by hydrophobic interaction chromatography and free of any interfering enzyme activity, was used. Maleylacetate reductase (EC 1.3.1.32) was measured as described by Seibert et al. (45) by using 0.05 mM maleylacetate freshly prepared by alkaline hydrolysis of cis-dienelactone or through transformation by TfdE, partially purified by hydrophobic interaction chromatography. For determination of the kinetic properties of TfdD, the formation of cis-dienelactone was recorded at 305 nm. A reaction coefficient of 5,300 M⁻¹ cm⁻¹ was calculated after complete transformation of 3-chloromuconate (20 to 50 μM) into cis-dienelactone.

Specific activities are expressed as micromoles of substrate converted or product formed per minute per gram of protein at 25°C. Protein concentrations were determined by the Bradford procedure (2).

**Chromatographic separation of enzyme activities.** Cells were harvested during late-exponential growth, and the cell extract (usually containing between 8 and 20 mg of protein per ml) was applied directly to a MonoQ HR5/5 column (Amersham Pharmacia Biotech) or mixed with an equal volume of 2 M (NH₄)₂SO₄ and, after centrifugation, applied to a phenyl-Superose HR5/5 column (Amersham Biosciences, Freiburg, Germany). At least three independent experiments were performed for each type of growth condition to verify the reproducibility of the method. Elution of the respective activities was highly reproducible and varied between independent experiments by only ± 1 fraction, which corresponds to differences of 0.01 M NaCl or 0.02 M (NH₄)₂SO₄. Proteins were eluted at a flow rate of 0.1 ml/min, applied directly to a MonoQ HR5/5 column by a linear gradient of NaCl (0 to 0.5 M) in Tris-HCl (50 mM, pH 7.5, supplemented with 2 mM MnCl₂) in a total volume of 25 ml or by a stepwise gradient of 0 to 0.14 M over 2 ml, 0.14 to 0.29 M over 17 ml, and 0.25 to 0.5 M over 6 ml. Proteins were eluted from the phenyl-Superose HR5/5 column by a linear gradient of (NH₄)₂SO₄ (1 to 0 M) in Tris-HCl (50 mM, pH 7.5, supplemented with 2 mM MnCl₂) over 25 ml or by a stepwise gradient of 1 to 0.8 M over 2 ml, 0.8 to 0.5 M over 8 ml, 0.5 to 0.3 M over 2 ml, 0.3 to 0.2 M over 6 ml, and 0.2 to 0.1 M over 10 ml. For chromatographic separation of maleylacetate reductases, proteins were eluted from the MonoQ HR5/5 column by the gradients described above, with Tris-HCl buffer exchanged for phosphate buffer. The flow rate was always 1 ml/min, and the fraction volume was 0.5 ml. Partial purification of TfdD was achieved by successive separation using hydrophobic interaction and gel filtration. The two fractions from the phenyl-Superose HR5/5 chromatography containing the most TfdD activity were pooled, concentrated by ultracentrifugation to a final volume of 0.5 ml, applied to a Superose 12 column (HR10/30) (Amersham Biosciences), and eluted with 50 mM Tris-HCl, pH 7.5, at a flow rate of 0.3 ml/min (fraction volume, 0.5 ml). Similarly, fractions from MonoQ HR5/5 chromatography containing TfdFI activity were pooled, concentrated, and subjected to gel filtration.

**Electrophoretic methods.** Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed on a Bio-Rad MiniProtein II essentially as described by Laemmli (23). The acrylamide concentrations for the concentrating and separating gels were 5 and 10% (wt/vol), respectively. The proteins were
FIG. 2. Growth of *R. eutropha* JMP222(pBBR1M-I) (left) and *R. eutropha* JMP222(pBBR1M-II) (right) on 3CB. Substrate depletion and product formation were monitored by HPLC.

Stained by Coomassie brilliant blue R250. Molecular mass standards (Bio-Rad, Munich, Germany) were lysozyme (14.4 kDa), trypsin inhibitor (21.5 kDa), carbonic anhydrase (31.0 kDa), albumin (45.0 kDa), bovine serum albumin (66.2 kDa), and phosphorylase b (97.4 kDa). For quantification of TfdDII in partially purified fractions, gels were stained with the fluorescent dye Sypro Ruby (Molecular Probes Inc., MoBiTec GmbH, Göttingen, Germany). Gels were scanned with a Fujifilm LAS-1000 charge-coupled device camera. The relative amount of the TfdDII protein band was determined with the AIDA, version 2.1, software package (Raytest Isopenmessgeräte GmbH, Straubenhardt, Germany).

For two-dimensional gel electrophoresis, a 250-μl aliquot of a TfdF-containing fraction obtained by anion-exchange chromatography was precipitated with 10% trichloroacetic acid, and then the precipitate was washed with ice-cold acetone. The protein was dissolved in 300 μl of reswelling solution containing 7.4 M urea, 2 M thiourea, 4% CHAPS (3-[3-cholamidopropyl]-dimethylammonioyl-1-propanesulfonate), 20 mM Tris base, 30 mM dithiothreitol (DTT), one-half tablet of protease inhibitor (MiniComplete; Roche Applied Biosciences, Mannheim, Germany), 5% IPG buffer, pH 4 to 7 (Amersham Biosciences) and applied to isoelectric focusing (IEF) ReadyStrip (Bio-Rad). After 100 kV·h focusing time in the first dimension (IEF), the gel strip was equilibrated for the second dimension (SDS-PAGE) twice for 15 min by using 5 ml of equilibration solution consisting of 6 M urea, 30% glycerol, 2% SDS, 0.06 mM bromophenol blue, 50 mM Tris base, pH 8.8, 65 mM DTT, and 260 mM iodoacetamide. The strip was applied then to an SDS–10 to 15% PAGE gel (1.5 mm thick) and developed overnight in an IsoDALT chamber (Amersham Biosciences).

N-terminal amino acid sequencing. Proteins were electrotroiled onto a polyvinylidene difluoride membrane, and the membrane was stained with Coomassie brilliant blue R250. N-terminal amino acid sequencing was performed with an Applied Biosystems model 494A Procise HT sequencer.

Analytical methods. High-performance liquid chromatography (HPLC) of low-molecular-weight compounds was performed with a Lichrospher SC 100 RP8 reverse-phase column (125 by 4.6 mm; Bishoff, Leonberg, Germany). Methanol-H2O containing 0.1% (vol/vol) H3PO4 was used as the eluant at a flow rate of 1 ml/min. The column effluent was monitored simultaneously at 210, 260, and 270 nm by a diode array detector (Shimadzu). Typical retention volumes using 25% (vol/vol) methanol were 7.7 (2-chloro-cis,cis-muconate), 2.5 (2-chloromuconolactone), 1.1 (5-chloromuconolactone), 4.1 (cis-dienelactone), and 1.8 ml (trans-dienelactone); those using 40% (vol/vol) methanol were 2.1 (2-chloro-cis,cis-muconate) and 1.4 ml (cis-dienelactone); those using 50% (vol/vol) methanol were 4.4 (CB) and 0.7 ml (2-chloro-cis,cis-muconate). Kinetic measurements were recorded on an UV 2100 spectrophotometer (Shimadzu Corporation, Kyoto, Japan).

Transformation of 2-chloromuconate by chloromuconate cycloisomerase. Transformation of 2-chloromuconate was usually performed in 50 mM Tris-HCl, pH 8. The reaction mixtures contained 500 μM 2-chloromuconate and 20 to 100 μM (as determined with 50 μM 3-chloromuconate as the substrate) of chloromuconate cycloisomerase per ml. Substrate transformation was monitored by HPLC.

Chemicals. Chemicals were purchased from Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany; Baker Chemikalien, Griesheim, Germany; and Merck AG, Darmstadt, Germany. 3-Chlorocatechol was obtained from Promochem, Wessel, Germany; 2-Chloro-cis,cis-muconate and (4R,5S)-5-chloro-3-methylmuconolactone were prepared as described by Pieper et al. (36, 38), (4R,5S)-5-Chloromuconolactone and (4S)-2-chloromuconolactone were prepared as described by Prucha et al. (40). cis-Dienelactone was a generous gift from Stefan Kaschabek and Walter Reinecke (Bergische Universität-Gesamthochschule, Wuppertal, Germany).

RESULTS

Growth of *R. eutropha* JMP222 derivatives containing pBBR1M-I or pBBR1M-II on 3CB and expression of Tfd gene products. Differences in the growth of *R. eutropha* JMP222 derivatives harboring *tfd* module I or II on 3CB have been reported previously (34). Accordingly, *R. eutropha* JMP222(pBBR1M-I) exhibited significantly higher growth rates (0.19/h) and higher growth yields (0.22 units of optical density at 546 nm (OD546/mM 3CB)) during growth on 3CB than did *R. eutropha* JMP222(pBBR1M-II) (0.06/h and 0.13 OD546 units/mM 3CB), as indicated by the turbidity of the culture obtained after complete transformation of the 3CB growth substrate (Fig. 2). The observed differences in growth yield seem to be due to the significantly different amounts of...
intermediates produced during growth of the two strains. Thus, whereas _R. eutropha_ JMP222(pBBR1M-I) accumulated 15% ± 5% of added 3CB as 2-chloromuconate, more than 50% of added 3CB appeared as 2-chloromuconate in the supernatant during growth of _R. eutropha_ JMP222(pBBR1M-II).

Whereas 2-chloromuconate was the only intermediate excreted during the growth of _R. eutropha_ JMP222(pBBR1M-I), _R. eutropha_ JMP222(pBBR1M-II) also excreted minute amounts of cis-dienelactone, corresponding to 2% of the substrate transformed. The accumulation of large amounts of 2-chloromuconate in strain JMP222 expressing module II, compared to the amounts accumulated in strain JMP222 expressing module I, can be explained by the recently reported very low activity of TfdDII chloromuconate cycloisomerase with this substrate (34). Accumulation of cis-dienelactone can be assumed to have been due to the observed low activity of TfdEII dienelactone hydrolase in _R. eutropha_ JMP222(pBBR1M-II) (Table 1).

Maleylacetate, produced by the activity of dienelactone hydrolase on cis- or trans-dienelactone, was never observed in the culture supernatants, indicating that maleylacetate reductase did not constitute a pathway bottleneck in _R. eutropha_ JMP222(pBBR1M-II) or in _R. eutropha_ JMP222(pBBR1M-I), despite the fact that only poor maleylacetate reductase activities have been recently reported to be induced during growth of strain JMP222 expressing _tfd_ module I (34). This discrepancy can be explained by the fact that the former enzyme activity measurements were performed with maleylacetate produced in situ by cell extracts. However, by using maleylacetate freshly prepared from cis-dienelactone by alkaline hydrolysis or purified dienelactone hydrolase and by measuring the enzymatic activity immediately after the preparation of a cell extract containing at least 5 mg of protein per ml, it was possible to reproducibly detect significant maleylacetate reductase activities in _R. eutropha_ JMP222(pBBR1M-I) (Table 1). All other enzyme activities were as previously reported (Table 1) (34).

**Levels of Tfd activity during growth of _R. eutropha_ JMP134 on chloroaromatics.** Due to the fact that all four enzymes involved in the chlorocatechol degradation of _tfd_ modules I and II were obviously active and based on detection of the respective mRNAs, which showed that both gene modules were expressed during growth on 2,4-D (22, 24), the importance of each gene module for the degradation of chloroaromatics in wild-type strain _R. eutropha_ JMP134(pJP4) was investigated. Chromatographic methods were applied to partially purify and separate the respective isoenzymes from cell extracts without significant loss of activity. Strains _R. eutropha_ JMP222(pBBR1M-I) and _R. eutropha_ JMP222(pBBR1M-II) were used for the establishment of an optimized purification scheme.

**Anion-exchange chromatography of cell extracts of 3CB-grown** _R. eutropha_ JMP222(pBBR1M-I) or JMP(pBBR1M-II) resulted in high recoveries of TfdCII (60% ± 10% of applied activity), TfdCII (80% ± 10%), TfdDII (70% ± 10%), TfdEII (60% ± 10%), TfdEI (90% ± 10%), and TfdEII (90% ± 10%) activity (Fig. 3A and B), although only low activities of maleylacetate reductase (activity of TfdFI eluting at 0.21 ± 0.02 M NaCl was usually lower than 5% of the applied activity, and activity of TfdFII eluting at 0.16 ± 0.01 M NaCl was usually lower than 10%) could be recovered. However, TfdCII (eluting at 0.29 M NaCl) showed a distinctly different retention behavior than did TfdCII (eluting at 0.19 M NaCl). TfdDII and TfdDII also exhibited different properties (eluting at 0.43 and 0.17 M NaCl, respectively). TfdEII and TfdEII could not be separated by such a chromatographic method (eluting at 0.16 M NaCl). Optimization of the applied NaCl gradient did not result in a significant enhancement of separation.

<table>
<thead>
<tr>
<th>Enzyme activity</th>
<th>Assay substrate</th>
<th>Sp act (U/g of protein) for indicated strain/substrate</th>
</tr>
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<tbody>
<tr>
<td>(Chloro) catechol 1,2-dioxygenase</td>
<td>Catechol</td>
<td>430 ± 90/840 ± 120/550 ± 90/540 ± 90/290 ± 70/260 ± 50</td>
</tr>
<tr>
<td>3-Chlorocatechol</td>
<td>Catechol</td>
<td>420 ± 90/380 ± 80/540 ± 90/290 ± 70/260 ± 50</td>
</tr>
<tr>
<td>4-Chlorocatechol</td>
<td>Catechol</td>
<td>410 ± 80/370 ± 80/520 ± 90/260 ± 50</td>
</tr>
<tr>
<td>(Chloro) muconate cycloisomerase</td>
<td>Muconate</td>
<td>&lt;5/120 ± 20/&lt;5/&lt;5/90 ± 30</td>
</tr>
<tr>
<td>2-Chloromuconate</td>
<td>Muconate</td>
<td>25 ± 5/15 ± 5/35 ± 10/140 ± 50</td>
</tr>
<tr>
<td>3-Chloromuconate</td>
<td>Muconate</td>
<td>270 ± 40/180 ± 40/360 ± 40/140 ± 50</td>
</tr>
<tr>
<td>Diene lactone hydrolase</td>
<td>cis-Diene lactone</td>
<td>1,050 ± 120/980 ± 110/1,200 ± 400/55 ± 15</td>
</tr>
<tr>
<td>Maleylacetate reductase</td>
<td>Maleylacetate</td>
<td>750 ± 170/510 ± 150/470 ± 150/1,520 ± 300</td>
</tr>
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</table>

*Values are averages of at least three independent experiments.*
been observed in 3CB-grown cells of *R. eutropha* JMP134 (36), whereas those enzymes were not induced during growth of this strain on 2,4-D (21). However, neither catechol 1,2-dioxygenase nor muconate cycloisomerase was induced during growth of strain JMP222(pBBR1M-I) with 3CB. The observed dienelactone hydrolase activity in strain JMP222(pBBR1M-II) was not identical to that of chromosomally encoded dienelactone hydrolase of strain JMP222 observed during growth on

FIG. 3. Separation of proteins from cell extracts of *R. eutropha* JMP134, JMP222, and derivatives by means of a MonoQ HR5/5 anion-exchange (A to C) or a phenyl-Superose HR5/5 hydrophobic-interaction column (D to F). Cell extracts were either directly applied (A to C) or applied after addition of ammonium sulfate (final concentration of 1 M) (D to E), and proteins were eluted with a linear gradient of NaCl (0 to 0.5 M) or (NH₄)₂SO₄ (1 to 0 M). The eluted fractions (0.5 ml) were analyzed for activities against the respective assay substrates (usually applied at a concentration of 50 µM, except for catechol and 3-chlorocatechol, which were added at a concentration of 200 µM). Enzymes eluting with a yield of ≥40% are shown. Arrows, enzymes eluting with a low yield (<20%; TfdF₁ and TfdF₄). Elution was performed in Tris-HCl buffer, except for panel C, where phosphate buffer was used.

(A) Separation of a cell extract of 3CB-grown cells of JMP222(pBBR1M-I) (6.1 mg of protein). (Inset) Activity of chromosomally encoded maleylacetate reductase after separation of a cell extract of 4-fluorobenzoate (4FB)-grown cells of JMP222 (7.9 mg of protein). (B) Separation of a cell extract of 3CB-grown cells of JMP222(pBBR1M-II) (11.6 mg of protein). (Inset) Detailed view of the activities observed in fractions 30 to 41. (C) Separation of a cell extract of 3CB-grown cells of JMP222(pBBR1M-I) (5.6 mg of protein). (Inset) Activity of chromosomally encoded maleylacetate reductase after separation of a cell extract of 4FB-grown cells of JMP222 (6.5 mg of protein). Muconate cycloisomerase activities could not be recovered after elution with phosphate buffer, as phosphate completely inhibited those enzymes. (D) Separation of a cell extract of 3CB-grown cells of JMP222(pBBR1M-I) (4.7 mg of protein). (Inset) Activity of chromosomally encoded maleylacetate reductase after separation of a cell extract of 4FB-grown cells of JMP222 (8.1 mg of protein). (E) Separation of a cell extract of 3CB-grown cells of JMP222(pBBR1M-II) (4.5 mg of protein). (F) Separation of a cell extract of 2,4-D-grown cells of JMP222(pBBR1M-I) (3.6 mg of protein).
4-fluorobenzoate (43), as the enzyme induced in strain JMP222(pBBR1M-II) exhibited similar activities against both cis- and trans-dienelactone (activity with trans-dienelactone was 60% of that with cis-dienelactone at substrate concentrations of 50 to 100 μM). The chromosomally encoded enzyme, in contrast, has been shown to exhibit activity only against trans-dienelactone (43).

Hydrophobic interaction chromatography was observed to be superior to anion-exchange chromatography, allowing separation of the different chlorocatechol dioxygenases and chloromuconate cycloisomerases, as well as the dienelactone hydrolases (Fig. 3D and E). However, the yields (50% ± 10% for TfdDI and TfdDII; 60% ± 10% for TfdEII, TfdEIII, and TfdCIII; and 70% ± 10% for TfdCII) were usually slightly lower than those obtained by anion-exchange chromatography, and hydrophobic-interaction chromatography also failed to give highly active preparations of maleylacetate reductase. Although a reasonable maleylacetate reductase activity, corresponding to approximately 5% of the observed activity against 3-chlorocatechol was observed in the initially eluting fractions during separation of the JMP222(pBBR1M-II)-derived extract. Furthermore, no precipitation of dienelactone hydrolase activities. However, no maleylacetate reductase under these conditions has been reported (43). Therefore, the possibility that the maleylacetate reductase activities observed during growth of strain JMP222(pBBR1M-I) or JMP222(pBBR1M-II) on 3CB (Table 1) are, at least partially, due to recruitment of this chromosomally encoded maleylacetate reductase cannot be excluded. Partial purification of this enzyme activity, which was usually induced at levels of 300 ± 50 U/g of protein in 4-fluorobenzoate-grown cells of strain JMP222, showed that this enzyme, in contrast to the activities induced in strains JMP222(pBBR1M-I) and JMP222(pBBR1M-II), exhibits high stability. Approximately 50% of the applied activity could be recovered by hydrophobic-interaction chromatography; the activity eluted at 0.38 M ammonium sulfate (Fig. 3D, inset). As no significant activity was observed in the respective fractions after separation of JMP222(pBBR1M-I)- or JMP222(pBBR1M-II)-derived cell extracts, it can be concluded that the chromosomally encoded maleylacetate reductase activity was not recruited in these strains for growth on 3CB. Similarly, as no significant activity was observed in the respective fractions after separation of a cell extract of 2,4-D-grown cells of strain JMP134 (Fig. 3F), it seems that the chromosomally encoded maleylacetate reductase does not play a major role in the degradation of this substrate in the wild-type strain.

Approximately 90% of the applied maleylacetate reductase activity of 4-fluorobenzoate-grown cells of R. eutropha JMP222 was recovered by anion-exchange chromatography; the activity eluted at 0.2 M NaCl (Fig. 3A, inset), a condition similar to those under which the activities of JMP222(pBBR1M-I) and JMP222(pBBR1M-II) eluted. However, the yields of the latter maleylacetate preparations were <5 and 10% ± 5%, respectively, again indicating that the chromosomally encoded maleylacetate reductase plays no significant role during 3CB degradation by strain JMP222(pBBR1M-I) or JMP222(pBBR1M-II).

As neither hydrophobic-interaction chromatography nor anion-exchange chromatography resulted in sufficient resolution of the three maleylacetate reductase activities or in reasonable yields of TfdEII, or, particularly, TfdF, anion-exchange chromatography was applied and Tris-HCl buffer in the eluant was replaced by phosphate buffer. This simple change had a marked effect on the retention behavior of the chromosomally encoded maleylacetate reductase, as well as on the TfdEII enzyme, as their interactions with the column material were nearly abolished (Fig. 3C). Moreover, the use of phosphate buffer dramatically increased the yield of TfdF. In a typical separation run, after application of a cell extract of strain JMP222(pBBR1M-I) containing 15 mg of protein, the two most active fractions contained, together, 7,050 mU of TfdF activity and 0.48 mg of protein (corresponding to a specific activity of 14,700 U/g of protein) in a complete volume of 1 ml. Further purification by gel filtration resulted in a significant loss of activity. Less than 10% of the applied activity was recovered, with specific activities of up to only 800 U/g of protein. To further demonstrate that the maleylacetate reductase induced in 3CB-grown strain JMP222(pBBR1M-I) is identical to TfdF, an aliquot of a highly active fraction obtained by anion-exchange chromatography, containing 240 μg of protein, was subjected to two-dimensional gel electrophoresis and blotted onto a polyvinylidene difluoride membrane and a major protein spot corresponding to the estimated molecular mass of

R. eutropha JMP134 was grown on 2,4-D, and, subsequently, the soluble proteins were separated by means of hydrophobic-interaction chromatography (Fig. 3F). The activities of tfdF-encoded enzymes TfdCII, TfdDII, and TfdEII were approximately 40 to 70% of those induced during growth of strain JMP222(pBBR1M-I) on 3CB, and those of tfdF-encoded enzymes TfdCII, TfdDII, and TfdEII were approximately 25 to 50% of those induced during growth of strain JMP222(pBBR1M-II) on 3CB. As the yields of the respective isoenzymes after chromatographic separation were always similar, the comparison of total activities in the respective fractions indicates the actual importance of the respective enzymes for the degradation process. As shown in Fig. 3F, in 2,4-D-grown cells of R. eutropha JMP134, the TfdF-encoded enzyme activities comprised 20% ± 5% of the observed total activity against 3-chlorocatechol and 3-chloromuconate but only approximately 5% of the observed activity against cis-dienelactone.

Characterization of maleylacetate reductases induced during growth of R. eutropha JMP134 and derivatives on chloroaromatics. R. eutropha JMP222 is capable of growth on 4-fluorobenzoate, and induction of a chromosomally encoded maleylacetate reductase under these conditions has been reported (43). Therefore, the possibility that the maleylacetate reductase activities observed during growth of strain JMP222(pBBR1M-I) or JMP222(pBBR1M-II) on 3CB (Table 1) are, at least partially, due to recruitment of this chromosomally encoded maleylacetate reductase cannot be excluded. Partial purification of this enzyme activity, which was usually induced at levels of 300 ± 50 U/g of protein in 4-fluorobenzoate-grown cells of strain JMP222, showed that this enzyme, in contrast to the activities induced in strains JMP222(pBBR1M-I) and JMP222(pBBR1M-II), exhibits high stability. Approximately 50% of the applied activity could be recovered by hydrophobic-interaction chromatography; the activity eluted at 0.38 M ammonium sulfate (Fig. 3D, inset). As no significant activity was observed in the respective fractions after separation of JMP222(pBBR1M-I)- or JMP222(pBBR1M-II)-derived cell extracts, it can be concluded that the chromosomally encoded maleylacetate reductase activity was not recruited in these strains for growth on 3CB. Similarly, as no significant activity was observed in the respective fractions after separation of a cell extract of 2,4-D-grown cells of strain JMP134 (Fig. 3F), it seems that the chromosomally encoded maleylacetate reductase does not play a major role during 3CB degradation by strain JMP222(pBBR1M-I) or JMP222(pBBR1M-II).

As neither hydrophobic-interaction chromatography nor anion-exchange chromatography resulted in sufficient resolution of the three maleylacetate reductase activities or in reasonable yields of TfdFII, or, particularly, TfdF, anion-exchange chromatography was applied and Tris-HCl buffer in the eluant was replaced by phosphate buffer. This simple change had a marked effect on the retention behavior of the chromosomally encoded maleylacetate reductase, as well as on the TfdEII enzyme, as their interactions with the column material were nearly abolished (Fig. 3C). Moreover, the use of phosphate buffer dramatically increased the yield of TfdF. In a typical separation run, after application of a cell extract of strain JMP222(pBBR1M-I) containing 15 mg of protein, the two most active fractions contained, together, 7,050 mU of TfdF activity and 0.48 mg of protein (corresponding to a specific activity of 14,700 U/g of protein) in a complete volume of 1 ml. Further purification by gel filtration resulted in a significant loss of activity. Less than 10% of the applied activity was recovered, with specific activities of up to only 800 U/g of protein. To further demonstrate that the maleylacetate reductase induced in 3CB-grown strain JMP222(pBBR1M-I) is identical to TfdF, an aliquot of a highly active fraction obtained by anion-exchange chromatography, containing 240 μg of protein, was subjected to two-dimensional gel electrophoresis and blotted onto a polyvinylidene difluoride membrane and a major protein spot corresponding to the estimated molecular mass of
the TfdF subunit (37.9 kDa) was analyzed by N-terminal sequencing. The amino-terminal sequence was identified as MKKFTLDYLSPR; this sequence was identical to the sequence deduced from the *tfdF* gene. Therefore, it can be assumed that the maleylacetate reductase activity present in 3CB-grown cells of strain JMP222(pBBR1M-I) is, in fact, due to expression of *tfdF*.

Thus, the use of anion-exchange chromatography with phosphate buffer has enabled the characterization of the expression of TfdFII, as well as of the different dienelactone hydrolases and catechol 1,2-dioxygenases, during growth of *R. eutropha* JMP134 and JMP222 derivatives. 2,4-D-grown cells of strain JMP134 induce significant levels of TfdFII, comprising roughly half of the total maleylacetate reductase activity. Calculations of the levels of TfdCII versus TfdEII in those cells confirmed the results obtained above by hydrophobic interaction chromatography, with 20% of the total chlorocatechol 1,2-dioxygenase activity due to TfdCII and 5% of the total dienelactone hydrolase activity due to TfdEII.

**Characterization of chloromuconate cycloisomerase TfdDII**

Compared to the previously described chloromuconate cycloisomerases from gram-negative microorganisms (21, 51) the chloromuconate cycloisomerase activity encoded by *tfd*DII exhibited very poor activity against 2-chloromuconate, possibly resembling chloromuconate cycloisomerase of *Rhodococcus opacus* 1CP (46). Therefore, this enzyme activity was further characterized. TfdDII was partially purified from 3CB-grown cells of strain JMP222(pBBR1M-II) by anion-exchange chromatography and gel filtration. A total of 10 U (calculated for 3CB-grown cells of strain JMP222(pBBR1M-II)) was subjected to gel filtration. A total of 3.8 U was recovered in four fractions, with specific activities as high as 3,800 U/g. Kinetic experiments were performed by quantification of cis-dienelactone formation at 305 nm, and a $K_m$ value of 190 ± 15 μM was calculated. For quantification of TfdDII in active fractions, aliquots were separated by PAGE (40 to 300 ng of protein per fraction) and stained with Sypro Ruby. A major band with a molecular mass of 41 ± 1 kDa, which contained 30 to 60% of the total protein in the respective fractions, was observed. The N-terminal sequence of this protein (MLTEKAIADSPN) was identical to that expected for TfdDII. Thus, maximal transformation rates calculated during kinetics experiments could be related to the amount of TfdDII present. Taking into account the predicted subunit molecular mass of 36.9 kDa, a $k_{cat}$ value of 1,950 ± 150 min$^{-1}$ was calculated.

By the photometric test (depletion in the presence of an excess of dienelactone hydrolase as measured at $\lambda = 260$ nm), TfdDII exhibited significant activity only against 3-chloromuconate and poor activity against 2-chloro-substituted muconate, as well as against unsubstituted muconate. At a substrate concentration of 0.1 mM, the activity with muconate was only 0.4% ± 0.1% of that with 3-chloromuconate and the activity with 2-chloromuconate was 0.8% ± 0.2%.

This substrate spectrum significantly resembles that reported for chloromuconate cycloisomerase of *Rhodococcus opacus* (46). This cycloisomerase differs from those of previously described gram-negative organisms in that it lacks the ability to convert 2-chloromuconolactone and to form trans-dienelactone from 2-chloromuconate. Thus, the kinetic properties of the TfdDII chloromuconate cycloisomerase were assessed by using the partially purified enzyme fractions described above. These fractions were free of any muconate cycloisomerase (capable of transforming 2-chloromuconolactone), muconolactone isomerase (capable of dehalogenating intermediate 5-chloromuconolactone), or dienelactone hydrolase activities. As shown in Fig. 4, the predominant products formed from 2-chloromuconate were identified as 2- and 5-chloromuconolactone. The ratio between 2-chloromuconate, 5-chloromuconolactone, and 2-chloromuconolactone, after a short incubation period, was about 20:4:3, as previously reported for the equilibrium between those compounds after addition of muconate cycloisomerases (50). Further incubation resulted in the formation of mainly trans-dienelactone and minor amounts of cis-dienelactone. A control incubation using partially purified muconate cycloisomerase resulted in the formation of a similar equilibrium between 2-chloromuconate and 5- and 2-chloromuconolactone. However, the rate of trans-dienelactone formation was negligible in those assays (less than 2 μM h$^{-1}$ in the presence of 100 mM of muconate cyc-
loisomerase per ml, compared to 12 μM h⁻¹ in the presence of 100 mU of TfdI per ml).

Incubation of TfdDII with 2-chloro muconolactone resulted in formation of the equilibrium between 2-chloro muconate and 2- and 5-chloro muconolactone described above, demonstrating that TfdDII, in contrast to the Rhodococcus enzyme, can transform 2-chloro muconolactone. Considerable levels of trans-dienelactone formation were visible only after significant accumulation of 5-chloromuconolactone.

DISCUSSION

Chloroaromatic degradation by R. eutropha JMP134 has been the subject of investigations for decades. The recently discovered tfdDI genes have elucidated a new layer of complexity for chloroaromatic degradation by this strain. It has been shown that tfdDI genes are transcribed in R. eutropha JMP134 upon exposure to 2,4-D (22, 24). Moreover, R. eutropha JMP222 strains containing tfdII genes were capable of growing on 3CB (34), and expression in Escherichia coli showed that tfdII genes encode functional enzymes (22). We have demonstrated here that, besides tfdFII (45), at least tfdDII, tfdCII, and tfdEII are translated into functional enzymes in R. eutropha JMP134. We have also shown that tfdFII, which was previously thought to be nonfunctional or poorly functional (34, 45), is transcribed and translated into a functional enzyme. Thus, tfdII, as well as tfdI genes, encode a complete set of functional and active enzymes for the transformation of chlorocatechols into 3-oxoadipate. However, whereas significant activities of TfdDII, TfdCII, and TfdFII were observed during the growth of strain JMP134 on 2,4-D and also of strain JMP222 containing tfdII genes, activities of TfdDII were negligible compared with those of TfdEII. The basis for this poor activity remains to be elucidated. A second obvious difference between tfdI and tfdII-encoded proteins lies in the TfdD proteins. On the first view, both TfdD (21, 51) and TfdDII are characterized by their poor activity with 2-chloro muconate and muconate, compared to that with 3-chloro muconate and thus differ only slightly in substrate specificity. However, whereas the relative TfdD activity with 2-chloro muconate at substrate concentrations between 10 and 100 μM is 6 to 9% of that with 3-chloro muconate (51), the relative activity of TfdDII is less than 1%. This poor activity with 2-chloro muconate is reflected in the higher accumulation of this intermediate and lower growth yield of strain JMP222 containing tfdII genes, compared to those of strain JMP222 containing tfdI genes, when the strain is grown on 3CB.

However, the most striking difference between TfdDII and TfdDII is the poor capability of TfdDII to dechlorinate during cycloisomerization of 2-chloro muconate. In contrast to what was found for other gram-negative bacteria chloro muconate cycloisomerases, where only small amounts of 2- and 5-chloro muconolactone accumulated during 2-chloro muconate turnover (52), dehalogenation to form trans-dienelactone by TfdDII is slow compared to cycloisomerization. Thus, TfdDII seems to be specifically dedicated to the transformation of 3-chloro-substituted muconates, a feature recently described for chloro muconate cycloisomerase of Rhodococcus opacus 1CP (46). However, whereas the Rhodococcus enzyme neither transforms 2-chloro muconolactone nor dehalogenates 2-chloro muconate or 5-chloro muconolactone during cycloisomerization, TfdDII exhibits both capabilities. TfdDII thus appears to be only distantly related, by sequence homology (12) as well as by kinetic properties, to all chloro muconate cycloisomerases described to date (Fig. 5). With respect to biochemical properties, TfdDII is intermediate to muconate and chloro muconate cycloisomerases of gram-negative bacteria, making TfdDII an interesting subject for evolutionary study. Regarding 3-chloro muconate turnover, no differences between TfdDII and other chloro muconate cycloisomerases described so far (e.g., a kcat/Km value of 10 μM⁻¹ min⁻¹ compared to 7 to 38 μM⁻¹ min⁻¹ for the Pseudomonas sp. strain P51- and Pseudomonas sp. strain B13-derived and TfdDII chloro muconate cycloisomerases) were evident.

It has been speculated recently (22) that the main reason for maintenance of the tfdI gene cluster, other than as a mechanism for supplying auxiliary functions, such as the facilitated uptake of 2,4-D mediated by TfdK (26), is the supply of a functional transcriptional activator (TfdR) (25). Furthermore, it has been assumed that TfdFII is the major maleylacetate reductase complementing a nonfunctional or poorly functional TfdFII gene product. However, we can demonstrate now that TfdFII is, in fact, functional and induced at a high level during growth on 2,4-D.

However, besides these observations, the tfdI genes definitively increase the dosage of chlorocatechol genes and of active gene products in strain JMP134 when the strain is grown on chloroaromatics. As has been recently shown, a duplication of the tfd gene modules results in an increased growth rate on 3CB (5). It can be proposed that the number of tfd genes present in the wild-type strain is limiting for growth on 3CB. As TfdDII, TfdCII, and TfdFII enzymes constitute more than 20% of the total activity during growth on chloroaromatics, it is evident that maintenance of the tfdI genes would be advantageous for the strain. The fact that an elevated number of chlorocatechol genes is necessary to achieve growth on chloroaromatics has been reported in various cases. The transfer of single copies of chlorocatechol genes (tcbR and tcbCDEF) originating from Pseudomonas sp. strain P51 (48) into Pseudomonas putida KT2442 did not result in the expected 3CB-degrading derivatives; multiple chromosomal copies (at least two) were needed to achieve this phenotype (20). Similarly, the transfer of single copies of tfdi or tfdI genes into the chromosome of R. eutropha JMP222 did not result in a 3CB-degrading phenotype, in contrast to the situation where those genes were introduced on a medium-copy-number vector (34). P. putida F1 transconjugants containing two copies of the Pseudomonas sp. strain B13-derived clc element were unable to grow on chlorobenzene, and characterization of chlorobenzene-degrading transconjugants revealed that three to eight copies of the clc element were required for growth, with a larger number of clc elements being associated with increasingly vigorous growth (41). Moreover, the currently available genetic evidence suggests that gene amplification plays an important role in the adaptation of bacteria to chloroaromatic degradation in contaminated environments (31, 32, 49).

High levels of catechol 1,2-dioxygenase and muconate cycloisomerase were observed in 3CB-grown cells of strain JMP222 harboring tfdII genes and of strain JMP134 but not in 3CB-grown cells of strain JMP222 harboring tfdI genes or 2,4-
D-grown cells of strain JMP134 (Table 1). Expression of catechol and chlorocatechol operons usually requires LysR-type transcriptional activators and inducer muconate (catechol operons) or 2-chloromuconate (chlorocatechol operons) (28). In strain JMP134, the \( \text{tfdR} \) gene product is responsible for expression of the \( \text{tfd} \) operons (25) and it is activated by chloromuconates (14). It has been proposed that, even in the absence of \( \text{tfdR} \), low-level expression of \( \text{tfdCDEF} \) occurs, implying a cross-activation by chromosomally encoded regulatory elements (14). In \( \text{P. putida} \) it has been shown that CatR, the regulator of the \( \text{clcABD} \) operon (for chlorocatechol degradation), interacts with the \( \text{clcABD} \) promoter region and, likewise, \( \text{clcR} \), the regulator of the \( \text{clcABD} \) operon (for chlorocatechol degradation), was shown to interact with the \( \text{catBC} \) promoter region. CatR could even complement a \( \text{ClcR}/\text{H11002} \) mutant \( \text{P. putida} \) strain harboring the \( \text{clcABD} \) operon for growth on 3CB (29, 33). Moreover, it was recently demonstrated that the LysR-type regulator of the \( \text{cbn} \) operon of \( \text{R. eutropha} \) NH9 is activated by both muconate and 2-chloromuconate (30), implying the presence of significant cross talk among the homologous transcriptional activators. Consequently, it seems likely that the accumulated 2-chloromuconate may drive expression of the catechol pathway genes by acting on a putative \( \text{catR} \)-like element in the chromosomes of strains JMP134 and JMP222. However, induction was observed only in a subset of growth conditions or genetic backgrounds, specifically those that resulted in accumulation of a high level of 2-chloromuconate during growth. Thus, evidently, highly elevated levels of intracellular 2-chloromuconate are necessary to achieve expression of catechol catabolic genes. This agrees with observations of Cosper et al. (6), who proposed that versions of \( \text{Acinetobacter} \) sp. strain ADP1 containing variants of the muconate cycloisomerase with reduced catalytic properties have increased intracellular levels of muconate available, which are responsible for the activation of LysR-type regulator CatM, which is, in turn, capable of activating the expression of genes encoding enzymes for benzoate degradation.

On one hand, the activation of the catechol pathway genes in \( \text{R. eutropha} \) can be regarded as a burden for the strain. However, an induction of this type can be proposed to have advantageous effects, as well, specifically for strains such as JMP222(pBBR1M-II). Catechol 1,2-dioxygenase and muconate cycloisomerase have some activity on chlorinated substrate analogues (21, 37) and thus to a certain extent can be responsible for transformation of pathway intermediates. 2-Chloro- and 3-chloromuconate will, thereby, be transformed into chloromuconolactones (50) and protoanemonin (1), respectively. Protoanemonin accumulation, when this compound is produced at low rates, can be prevented by dienelactone hydrolase (3), and muconolactone isomerase can support the dehalogenation of any 5-chloromuconolactone (39) formed during cycloisomerization of 2-chloromuconate by muconate cycloisomerase or \( \text{TfdD}_{II} \). Thus, it can be assumed that part of 3CB in strain JMP222(pBBR1M-II) is mineralized by a com-

![Diagram](http://jb.asm.org)
plex metabolic interplay between chromosome- and plasmid-encoded enzymes.

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