Conjugative Plasmids and the Degradation of Arylsulfonates in Comamonas testosteroni

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Comamonas testosteroni T-2 degrades p-toluensulfonate (TSA) via p-sulfobenzoate (PSB) and protocatechuate and degrades toluene carboxylate via terephthalate (TER) and protocatechuate. The appropriate genes are expressed in at least five regulatory units, some of which are also found in C. testosteroni PSB-4 (F. Junker, R. Kiewitz, and A. M. Cook, J. Bacteriol. 179:919–927, 1997). C. testosteroni T-2 was found to contain two plasmids, pTSA (85 kbp) and pT2T (50 kbp); a TSA− mutant (strain TER-1) contained only plasmid pT2T. C. testosteroni PSB-4, which does not degrade TSA, contained one plasmid, pPSB (85 kbp). The type strain contained no plasmids. Conjugation experiments showed that plasmid pTSA (possibly in conjunction with pT2T) was conjugative, and the single copy of the TSA operon (tsaMBCD) with its putative regulator gene (tsaR) in strain T-2 was found on plasmid pTSA, which also carried the PSB genes (psbAC) and presumably transport for both substrates. Plasmid pT2T was assigned to the IncP1 group and was found to carry two copies of insertion element IS1071. Plasmid pPSB (of strain PSB-4), which could be maintained in strains with plasmid pTSA or pT2T, was also conjugative and was found to carry the PSB genes as well as to contain two copies of IS1071. In attempted conjugations with the type strain, no plasmid was recovered, but the PSB+ transconjugant carried two copies of IS1071 in the chromosome. We presume the PSB genes to be located in a composite transposon. The genes encoding the putative TER operon and degradation of protocatechuate, with the meta cleavage pathway, were attributed a chromosomal location in strains T-2 and PSB-4.

Arylsulfonates are widespread pollutants, some of which can easily enter drinking water (7, 15; cf. 39). Despite the multi-millions of tons of sulfonates used each year (6, 13), they seldom accumulate in the environment (34), and some are obviously subject to biodegradation (1, 15). The arylsulfonate whose degradation is best understood is obviously subject to biodegradation (1, 15). The arylsulfonate easily enter drinking water (7, 15; cf. 39). Despite the multi-bolic plasmids under environmental conditions may be less relatively high copy numbers (22). Conjugation of large cat- mids encoding antibiotic and heavy metal resistance are trans- ferred easily among bacteria, being small and present in autonomous maintenance in the bacterial host and transfer to a wide range of gram-negative genera. The copy number per chromosome is four to six for Escherichia coli and two to three for Pseudomonas spp. (23). We now report that regulatory units R1 and R3 in C. testosteroni T-2 are encoded on an 85-kbp IncP1 group plasmid, which is conjugative and which is missing in mutant TER-1. We believe that regulatory unit R3 is located in a transposon involving IS1071 insertion elements. C. testosteroni PSB-4 also encodes regulatory unit R3 on a conjugative plasmid, which does not belong to the IncP1 group but which presumably does involve a composite transposon incorporating IS1071 insertion elements.
irregular box are chromosomally encoded. The type strain of \textit{C. testosteroni} PSB-4, which contains no regulatory unit R1, but contains complete units R2, R3 and R4. The type strain of \textit{C. testosteroni} PSB-4, which was chosen to detect IS1071, includes the 3' end of oxygenase tsaM and the 5' end of \textit{korA}, \textit{trfA2-1} and \textit{tsaMB} because it includes the 3' end of the transposase and the adjacent inverted repeat. The PCR product with primer pair \textit{tsaMB} and \textit{tsa-N} was chosen to detect \textit{tsaMB} as probes (blots of PCR products or total DNA). The PCR product with primer pair \textit{tsaMB} and \textit{tsa-N} was chosen to detect \textit{tsaMB} as probes (blots of PCR products or total DNA). The PCR product with primer pair \textit{tsaMB} and \textit{tsa-N} was chosen to detect \textit{tsaMB} as probes (blots of PCR products or total DNA).

**MATERIALS AND METHODS**

**Bacterial strains and plasmids.** The organisms we used are shown in Table 1. \textit{C. testosteroni} T-2 \textit{C. testosteroni} PSB-4 (DSM 11414) were isolated by Thurnheer et al. (36) and thoroughly identified (4).

**PCR analysis.** PCRs (Table 2) were carried out with either Taq polymerase were obtained from Fermentas, Vilnius, Lithuania.

**Southern blot and dot blot hybridizations.** Southern blotting, dot blotting, and \textit{corA} were used as probes of PCR products or total DNA dot blots under stringent conditions appropriate for 75 to 100% nucleotide sequence similarity. In addition, probes of digoxigenin-labeled PCR products for \textit{IncN(oriT)}, \textit{IncQ(oriV)}, and \textit{IncW(oriP)} plasmids were used under the same stringent conditions. The probes and primer systems used to identify the incompatibility group are not suitable to detect all plasmids belonging to the corresponding incompatibility group (8).

**Filter matings.** Kanamycin (KM)-resistant recipient strains from \textit{C. testosteroni} PSB-4, TER-1, and ATCC 11996 (type strain) were generated by screening for spontaneous mutants on plates containing KM (50 \mu g/ml). Cultures (10 ml) of donor and recipient cells were grown overnight to an optical density at 600 nm of 0.6, centrifuged, washed twice in 0.9% NaCl, and resuspended in 10 ml of 0.9% NaCl. Selection for the transconjugants of the donor-recipient pairs T2-PSB4, T2-TER1, and T2-TYPE was done on plates containing KM (50 \mu g/ml) and that for PSB4-TER1 and PSB-4–ATCC 11996

**TABLE 1. Bacteria used in this study**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relevant characteristic(s)</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{C. testosteroni} T-2</td>
<td>TSA, PSB, TER, PCA</td>
<td>DSM 6577^a</td>
</tr>
<tr>
<td>PSB-4</td>
<td>PSB, TER, PCA</td>
<td>DSM 11414^a</td>
</tr>
<tr>
<td>PSB-4-K^b</td>
<td>PSB, TER, PCA, KM'</td>
<td>This study</td>
</tr>
<tr>
<td>TER-1</td>
<td>TER, PCA</td>
<td>14</td>
</tr>
<tr>
<td>TER-1-K^b</td>
<td>TER, PCA, KM'</td>
<td>This study</td>
</tr>
<tr>
<td>TYPE</td>
<td>PCA</td>
<td>DSM 50244</td>
</tr>
<tr>
<td>TYPE^b</td>
<td>PCA, KM'</td>
<td>This study</td>
</tr>
</tbody>
</table>

**E. coli**

J53R4 Met^-, Pro^-, plasmid K. Smalla
J53R751 Plasmid R751 (TP') K. Smalla

\textit{Pseudomonas putida} mt-2 TOL-plasmid pWW0 DSM 3931

^a Abbreviations indicate ability to degrade the respective compound. KM', TC', AP', and TP' indicates resistance to kanamycin, tetracycline, ampicillin, and trimethoprim, respectively. Met' and Pro' indicate auxotrophy for these amino acids.

^b These KM' organisms were used as recipients in conjugations, but to simplify nomenclature in the text we do not refer to the resistance marker there.

^c See reference 36.

**FIG. 1.** Degradative pathways for TSA and TCA in \textit{C. testosteroni} T-2, their regulation, and the portions of the pathway found in mutant TER-1 and in \textit{C. testosteroni} PSB-4. TSA enters the cell via a physiologically characterized transport system, termed R0, which is coregulated with regulatory unit R1 but is distant from the known R1 operon \textit{tsaMBD} and \textit{tsaR} (13, 18, 29). TSA is oxygenated to \textit{p-sulfinobenzaldehyde} (SYD) to PSB, which is oxygenatively desulphonated to \textit{PCA}. \textit{meta}-cleavage to 4-carboxy-2-hydroxymuconate semialdehyde (CHS) follows. The appropriate catabolic enzymes are encoded in three regulatory units, R1 (the TSA operon), R3 (\textit{psbAC}), and R4 (at least \textit{pcpP}) (13, 29). TCA presumably requires transport into the cell independent of TSA transport (13, 18), and TCA is oxygenated to \textit{carboxyphenyl alcohol} (COL) and further oxidized via \textit{carboxybenzaldehyde} (CYD) to TER, which is oxygenated to the diene-diol (DDC) and decarboxylated to \textit{PCA}. The appropriate catabolic enzymes are encoded in the \textit{C. testosteroni} operon and regulatory unit R2 (the TER operon) (13, 17, 41). The enzymes are represented by abbreviations in boldface. A second organism under study is a mutant of strain T-2, named strain TER-1, which has lost regulatory units R1 and R3, but retains R2 and R4 (14). The third organism is \textit{C. testosteroni} PSB-4, which contains no regulatory unit R1, but contains complete units R2, R3 and R4. The type strain of \textit{C. testosteroni} ATCC 11996, according to the literature (42), encodes the equivalent of regulatory unit R4. The enzymes shown in the regular box are chromosomally encoded.
was done on plates containing sulfobenzoate and KM (50 μg/ml). Controls for donor and recipient cells were plated separately on the same selective plates. Conjugations were carried out with *E. coli* J53, harboring plasmid RP4, as donor, and strain T-2 or transconjugant T2–PSB4 as recipient, and transconjugants were selected on 10 mM succinate minimal medium plates containing TC (50 μg/ml).

**PFGE.** A contour-clamped homogeneous electric field system (model CHEF-DRII; Bio-Rad Laboratories, Richmond, Calif.) was used to separate plasmid DNA on 0.9% agarose gels. Preparation of total DNA was done essentially as described previously (32). All experiments were done in TBE buffer (100 mM Tris-HCl, 100 mM boric acid, 0.2 mM EDTA [pH 8.3]) at 4°C. The pulsed-field gel electrophoresis (PFGE) conditions used were 200 V; running time, 22 h; and pulse time, 20 to 120 s. The chromosome of *Saccharomyces cerevisiae* YNN295 (Bio-Rad) was used as a linear DNA size marker. The plasmids pWW0, RP4, and R751 were used as supercoiled DNA standard markers.

**Plasmid purification.** Purifications of plasmids pTSA, pWW0, and R751 were performed according to the procedure described by Ish-Horowicz and Burke (10). The protocol was modified to include a phenol-chloroform extraction (8), and the isolated plasmid was further purified with a Geneclean Plasmid Spin-Kit (BIO 101).

**Mercury and antibiotic resistance.** Ampicillin (AM), KM, streptomycin, TC, trimethoprim (TP), or Hg resistance (at 10- and 50-μg/ml concentration) was tested in liquid medium for strains T-2, PSB-4, and TER-1.

**Nucleotide sequence accession number.** The accession number for the fragment of *trfA2* we sequenced is GenBank U73743.

## RESULTS

**Plasmid pTSA of *C. testosteroni* T-2.** PFGE showed that strain T-2 contains at least two megaplasmids (Fig. 2, lane 1). The larger plasmid was designated pTSA (from the TSA operon) (13), and the smaller plasmid was designated pT2T (see below). In contrast, mutant TER-1 (TSA–, PSB–, PCA+) contained only the smaller plasmid, pT2T (Fig. 2, lane 4; Table 3). The type strain (TYPE) of *C. testosteroni* (TSA–, PSB–, PCA+)

### TABLE 2. Primer systems

<table>
<thead>
<tr>
<th>Target</th>
<th>Primer pair (5’–3’)</th>
<th>Size of PCR product (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>tsaMB</td>
<td><em>tsaM</em>-C, AAAATCTTGAGCCAGGT;</td>
<td>386</td>
</tr>
<tr>
<td></td>
<td><em>tsaB</em>-N, TTGAGCTTCTGGAATCT</td>
<td></td>
</tr>
<tr>
<td>IS1071</td>
<td><em>tpA</em>, GGGAAATCTTGAGCCAGATG;</td>
<td>338</td>
</tr>
<tr>
<td></td>
<td><em>IR</em>, CTTTGTAGTCGAAATG</td>
<td></td>
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<tr>
<td>IncP1</td>
<td><em>orT</em>-1, CACCGCTCGCAGGAGCAGATG;</td>
<td>110</td>
</tr>
<tr>
<td></td>
<td><em>orT</em>-2, CACCGCGGCCAGGATAGGTAAGT</td>
<td></td>
</tr>
<tr>
<td>trfA2</td>
<td>*CAGAAATTCATGTGGTGAAGGAAATG;</td>
<td>240</td>
</tr>
<tr>
<td></td>
<td><em>CAGTTGCAATGCACAGGTC</em></td>
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</tr>
<tr>
<td>korA1</td>
<td><em>ATGAAAGAAACGCGTACCGA</em></td>
<td>294</td>
</tr>
<tr>
<td></td>
<td><em>TTCTCGTGTCTTGGCGTC</em></td>
<td></td>
</tr>
</tbody>
</table>

*a* Primers based on published sequence data were used to detect IncP1 plasmids, insertion element IS1071 and the genes *tsaMB*, *tsaMB* toluenesulfonate methyl-monoxygenase (13); IS1071, class II insertion sequence element (20); IncP1, incompatibility group P-1 plasmid; *orT*, origin of replication; *trfA2*, trans-acting replication function; *korA*, host-killing override factor (8).

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**FIG. 2.** PFGE of DNA from different strains of *C. testosteroni* and growth characteristics of these organisms. +, growth; −, no growth with substrate as sole source of carbon and energy. Lane 1, strain T-2; lane 2, plasmid pTSA; lane 3, transconjugant J53RP4–T2; lane 4, strain TER-1; lane 5, supercoiled plasmids pWW0 and R751; lane 6, yeast chromosomal DNA standard; lane 7, strain PSB-4; lane 8, transconjugant PSB4–TER1; lane 9, transconjugant T2–PSB4; lane 10, transconjugant J53RP4–(T2–PSB4). Sizes indicated at left for lanes 1 to 5 and at right for lanes 6 to 10.
The ability to utilize toluenesulfonate and sulfobenzoate (but not TER) could be transferred via conjugation from donor strain T-2 to the type strain by transferring plasmid pTSA, whose presence (with pT2T) in T2-TYPE transconjugants was detected after PFGE (Table 3). Similarly, T2-TER1 transconjugants were TSA\(^+\) and PSB\(^-\) (and TER\(^-\)) and contained both plasmids pTSA and pT2T (Table 3), and T2-PSB4 transconjugants were also TSA\(^+\) but lacked plasmid pT2T (Table 3). Each of these transconjugants (T2-TYPE, T2-TER2, and T2-PSB4) was shown to contain the genes \(tsaMB\) by PCR (data not shown) and by Southern blot analysis (Fig. 3A and B, lanes 7 and 10; see also reference 14). These genes were first located on the pTSA plasmid in strain T-2 (Fig. 3A and B, lanes 1 and 3) but not in strain PSB-4, TER-1, or ATCC 11996 (Fig. 3A and B, lanes 4 to 6). Association of the \(tsaMBCD\) genes (R1) with plasmid pTSA, which was shown to be conjugative (at least in the presence of plasmid pT2T), was thus confirmed. Plasmid pT2T is further detailed below.

The first attempt to obtain an organism which harbors only plasmid pTSA, transconjugant T2-TYPE, contained both plasmids (see above), although the selection was only for degradation of toluenesulfonate; no other attempts succeeded (see below). However, plasmid preparations from strain T-2, separated by PFGE, showed that plasmid pTSA was largely recovered, with only traces of plasmid pT2T and chromosomal DNA (Fig. 2, lane 2). Restriction enzyme analysis of \(Pst\) I and \(Bgl\) II-SalI digests of plasmid pTSA and Southern blot hybridization with a probe against the monoxygenase gene \(tsaMB\) revealed no signals of the sizes (1.8 kbp for \(Pst\) I [Fig. 3A and B, lane 1] and 4 kbp for \(Bgl\) II-SalI) predicted by the restriction map and sequence of the TSA operon (13). Southern blot hybridizations of purified plasmid pTSA separated on PFGE gave signals with probes against \(tsaMB\) (data not shown). The size of plasmid pTSA, 85 ± 10 kbp, was first estimated by summing the sizes of restriction fragments obtained from a \(Pst\) I (Fig. 3A, lane 1) or from a \(Bgl\) II-SalI digest. Supercopied plasmid pTSA was located between plasmids pWW0 (117 kbp) and R751 (53 kbp), which confirmed the estimate of 85 kbp.

Catabolic genes are often associated with IncP1 plasmids (3, 21, 24, 31, 33, 43), so we probed by PCR for the presence of three conserved genes of IncP1 plasmids (Table 2) in total DNA or in plasmid preparations from strains T-2, TER-1, and PSB-4. Products of the expected sizes (Table 2) were obtained from strain T-2 only (data not shown); no specific product was obtained from strains TER-1 and PSB-4. This presence of an IncP1 gene (e.g., \(traB2\)) in strain T-2, but not in strain TER-1 or PSB-4, was confirmed by Southern blot analysis (Fig. 3, lanes 3 to 5), and the gene was located on plasmid pTSA (Fig. 3, lane 1). We confirmed pTSA as being an IncP1 plasmid by mating strain T-2 (recipient) with \(E. coli\) J53RP4, which harbors the IncP1\(\alpha\) plasmid RP4. Selection for \(C. testosteroni\) and the IncP1\(\alpha\) plasmid RP4 gave the transconjugant J53RP4-T2 (TSA\(^-\), PSB\(^-\), TER\(^-\)), which had lost plasmid pTSA but contained plasmid RP4 instead, as well as plasmid pT2T (Fig. 2, lanes 1 and 3; Table 3). We were able to assign pTSA to IncP1\(\beta\) by first labeling the PCR products (\(oriT, traB2,\) and \(korA\)) of the IncP1\(\beta\) reference plasmid R751 with digoxigenin and then using these as probes in Southern blots of the PCR products and dot blots of total DNA from strain T-2 (data not shown) (8). Signals were obtained from the PCR products of strain T-2 and from the positive control, plasmid R751, but not for the IncP1\(\alpha\) control plasmid RP4.

The PCR product from the \(traB2\) primers with plasmid pTSA was cloned into pBluescript KS and sequenced. The deduced amino acids (from 240 nucleotides) were aligned with the \(traB2\) gene products from plasmids R751 and RK2 (Fig. 4). Identities of 95 and 87% between the PCR product and TrfA2 from plasmids R751 (IncP1\(\beta\)) and RK2 (IncP1\(\alpha\)), respectively, were found. The higher identity between the \(traB2\) gene of plasmid pTSA and R751, compared with that between plasmids pTSA and RK2, further supports the identification of pTSA as an IncP1\(\beta\) plasmid.

We conclude that the conjugative, 85-kbp megaplasmid pTSA of strain T-2 belongs to the IncP1\(\beta\) group. Plasmid pTSA carries at least the operon \(tsaMBCD\) (R1) (Fig. 1), and the \(psbA(C)\) gene(s) of regulatory unit R3 and presumably transport R0.

**Plasmid pT2T of strain T-2 and its mutant TER-1.** The smaller megaplasmid, pT2T, of strains T-2 and TER-1 (Fig. 2, lanes 1 and 4), was found to have the same electrophoretic properties (by PFGE) as supercoiled IncP1\(\beta\) plasmid R751 and was estimated to be about 50 kbp in length. Plasmid preparations of plasmid pT2T (from strain T-2 or TER-1) gave

<table>
<thead>
<tr>
<th>Strain or transconjugants</th>
<th>Plasmid(^{a,b})</th>
<th>IS1071 copy no.</th>
<th>Growth substrate(^{c})</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>pTSA</td>
<td>pPSB</td>
<td>pT2T</td>
</tr>
<tr>
<td>T-2</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>PSB-4</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>TER-1</td>
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<td>TYPE</td>
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</table>

<table>
<thead>
<tr>
<th>Transconjugants (donor-recipient)</th>
<th>Plasmid(^{a,b})</th>
<th>IS1071 copy no.</th>
<th>Growth substrate(^{c})</th>
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<tr>
<td>T2-PSB4</td>
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<tr>
<td>T2-TER1</td>
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<td>+</td>
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<td>PSB4-TER1</td>
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<tr>
<td>J53RP4-T2</td>
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<td>+</td>
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<td>+</td>
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<tr>
<td>PSB4-TYPE</td>
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- \(^{a}\) +, present (plasmid) or utilized (substrate); -, absent (plasmid) or not utilized (substrate).
- \(^{b}\) Plasmid, IncP1\(\beta\) plasmid; RP4, IncP1\(\alpha\). Incompatibility group for pPSB or pT2T is unknown.
- \(^{c}\) ND, not determined.
low yields, and no catabolic function was found. However, plasmid pT2T was transferred in a mating experiment between strains T-2 (donor) and TYPE (Table 3) but not between strains T-2 (donor) and PSB-4 (Table 3). Its presence or absence did not lead to a change in the utilization of the substrates we studied, and it was not lost under nonselective conditions (growth on acetate or succinate). It does not belong to the IncP1 group, because it coexists with IncP1 plasmids pTSA (strain T-2 and transconjugant T2-TYPE) and RP4 (transconjugant J53RP4-T2) and no signals were obtained with specific IncP1 primers in PCR and Southern blots (Fig. 3, lane 5). Southern blots with IncW, IncN, and IncQ probes did not give a signal, but an affiliation with one of these groups cannot be excluded as the probes do not detect all members of the corresponding incompatibility group.

Plasmid pPSB of strain PSB-4. Only one plasmid was detected in C. testosteroni PSB-4, and it was termed pPSB (Fig. 2, lane 7). Mating experiments with strains PSB-4 (donor) and TER-1 yielded a PSB + transconjugant, PSB4-TER1 (Fig. 2, lane 8; Table 3), which contained plasmids pPSB and pT2T. Plasmid pPSB is thus conjugative and carries the genes pshAC for the oxygenation of sulfobenzoate (Fig. 1).

No PCR product was obtained with primers for conserved IncP1 genes (Table 2), so plasmid pPSB does not belong to the

FIG. 3. PstI digests of plasmid pTSA and total DNA and Southern blot hybridizations with three different probes. (A) Southern blot with 0.75% agarose gel; (B) methyl-monoxygenase tsaM probe; (C) IS1071-tnpA probe; (D) IncP1-trfA2 probe. Lane 1, plasmid pTSA; lane 2, DNA ladder (kbp); lane 3, strain T-2; lane 4, strain PSB-4; lane 5, strain TER-1; lane 6, strain TYPE; lane 7, transconjugant T2-PSB4; lane 8, transconjugant J53RP4–T2; lane 9, transconjugant J53RP4-(T2-PSB4); lane 10, transconjugant T2-TYPE; lane 11, transconjugant PSB4-TYPE; lane 12, λ HindIII DNA marker.

FIG. 4. Amino acid sequence alignment of the sequenced trfA2 PCR product (240 bp; AC5U73743) from plasmid pTSA with TrfA2 proteins from plasmids R751 (AC5U07618) and RK2 (AC5U05774). The underlined amino acids were encoded by the trfA2 PCR primers. *, amino acid from TrfA2 from plasmid R751 which is different from that of plasmid pTSA; #, amino acid from TrfA2 from plasmid RK2 which is different from that of plasmid pTSA. The numbering is based on TrfA2 from plasmid R751.
IncP1 group. This was confirmed in mating experiments with strains T-2 (donor) and PSB-4 which yielded a TSA\(^{-}\)-PSB\(^{-}\) transconjugant (T2-PSB4) which was found to contain plasmids pPSB and pTSA, which comigrate, but not pT2T (Fig. 2, lane 9). Plasmid pTSA of strain T2-PSB4 could be cured with plasmid IncP1\(\alpha\)-RP4 (Table 3), yielding an organism that was TSA\(^{-}\} and PSB\(^{-}\} and which contained plasmids pPSB and RP4 [J53RP4\((\{\{\text{T2-PSB4}\}\})\) (Fig. 2, lane 10). Plasmids pPSB and pT2T coexist in strain PSB4-TER1 so they belong to different incompatibility groups. As with plasmid pT2T, plasmid pPSB gave no signal in Southern blots with probes for IncW, IncN, and IncQ plasmids. Attempts to purify plasmid pPSB gave very poor yields. Comigration of plasmids pTSA and pPSB indicates that the latter is about 85 kbp.

Transconjugant PSB4-TYPE (Table 3) did not contain the plasmid but did gain the ability to grow on sulfobenzoate. The recombination of \(psbAC\) with the chromosome and the possible involvement of the insertion element IS1071 is discussed below.

**TER degradation.** TerZoa\(\beta\)R was found to be identical in the independently isolated \(C.\) testosteroni strains PSB-4 and T-2 and in mutant TER-1 (14). The plasmid-free strain TYPE (TER\(^{-}\), PCA\(^{-}\)\)) (Fig. 1) was used as a recipient in mating experiments. Transconjugant T2-TYPE (TSA\(^{+}\}, TCA\(^{-}\}, PSB\(^{-}\}, TER\(^{-}\}, PCA\(^{-}\})\) contained plasmids pTSA and pT2T (Table 3), and transconjugant PSB4-TYPE (PSB\(^{+}\}, TER\(^{-}\}, PCA\(^{-}\})\) (Table 3) did not gain the ability to utilize TER (or toluenecarboxylate). We conclude that regulatory unit R2 (Fig. 1) is missing in these transconjugants. Plasmids pTSA, pPSB, and pT2T do not carry the genes terZoa\(\beta\)R and terY (R2), which are apparently located on the chromosome.

**Insertion element IS1071.** Strain T-2 was found to contain insertion element IS1071 (43a). A Southern blot probe for the transposase of IS1071 (tnpA) (Table 2) revealed that two major copies of IS1071 were found in plasmid pTSA (4- and 1.7-kbp \(Pst\) I fragments) (Fig. 3A and C, lane 1); the trace (below the 4-kbp band) was attributed to chromosomal contamination. In contrast, three copies of IS1071 were found in digests of total DNA from strain T-2; of these, two copies corresponded with those on plasmid pTSA, and the third copy, located at 3.8 kbp, was presumed to be chromosomal (Fig. 3A and C, lane 1). Correspondingly, only this latter copy was detected in strain TER-1 and transconjugant J53RP4-T2 (Fig. 3A and C, lanes 5 and 8), which lack plasmid pTSA, and no such copy was found in strain TYPE (seen here in transconjugant T2-TYPE [Fig. 3A and C, lane 10]). The intensity of the 3.8-kbp signal from T-2 total DNA was about 30 to 50% of that of the two signals of plasmid origin. On this basis, the copy number of plasmid pTSA lay in the range of two to three, which is typical for IncP1 plasmids (23). Plasmid pT2T cannot carry any copy of IS1071, because transconjugant T2-TYPE (Fig. 3C, lane 10), which harbors plasmids pT2T and pTSA, has only the two copies of IS1071, which were shown to originate in plasmid pTSA (Fig. 3C, lane 1).

Three copies of IS1071 were detected in digests of total DNA from strain PSB-4 (25-, 4-, and 3.8-kbp, \(Pst\)I fragments) (Fig. 3A and C, lane 4). The 4- and 3.8-kbp fragments were common to both strains T-2 and PSB-4. By analogy to the data from strain T-2, we presume a chromosomal location for the IS1071 element in the 3.8-kbp \(Pst\) I fragment and a plasmid (pPSB) location for the other copies. Correspondingly, transconjugant T2-PSB4, which harbors the plasmids pTSA and pPSB, showed three strong (plasmid-derived) signals and one weak (chromosomal) signal for the IS1071 probe. Of these three strong bands, the 4-kbp band is derived from two sources, plasmids pTSA and pPSB.

### Table 4. Plasmids found in *C. testosteroni* T-2, PSB-4, and TER-1

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Strain(s)</th>
<th>Type</th>
<th>Size (kbp)</th>
<th>Catabolic genes</th>
<th>Regulatory unit(s)</th>
<th>IS1071 copy nos.</th>
</tr>
</thead>
<tbody>
<tr>
<td>pTSA</td>
<td>T-2</td>
<td>IncP1(\beta)</td>
<td>85</td>
<td>(tsaMBCD) ((tsaR),*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pPSB</td>
<td>PSB-4</td>
<td>Unknown</td>
<td>85</td>
<td>(psbA)</td>
<td>R1, R2</td>
<td>2</td>
</tr>
<tr>
<td>pT2T</td>
<td>T-2, TER-1</td>
<td>Unknown</td>
<td>50</td>
<td>(psbAC)</td>
<td>R2</td>
<td>2</td>
</tr>
</tbody>
</table>

* nd, not detected.

Transconjugant PSB4-TYPE grew with sulfobenzoate but did not contain a plasmid. Two copies of IS1071 were detected in \(Pst\)I digests (4 kbp and 3.8 kbp), but the 25-kbp fragment was missing. The 4-kbp signal was stronger than the 3.8-kbp signal and may be identical with the 4-kbp signal from plasmids pTSA and pPSB. We cannot explain the weak, 3.8-kbp signal, which may resemble the 3.8-kbp chromosomal fragment of strains T-2, PSB-4 and TER-1, and which was not found in strain TYPE (see above), so given other nonspecific signals (see below), we cannot exclude the possibility that this 3.8-kbp \(Pst\)I signal is nonspecific.

Data and interpretations based on \(Pst\)I digests were confirmed by independent \(Bgl\)II-\(Sal\)I digests (data not shown). However, in addition to the signals just discussed, Southern blots of \(Bgl\)II-\(Sal\)I digests showed an additional weak signal which is presumably nonspecific.

Strains T-2, PSB-4, and TER-1 were AM resistant. Streptomycin resistance occurred after a longer lag phase (2 to 3 days) but no other resistance (e.g., mercury) was found. Most *C. testosteroni* strains are AM resistant (42), so it seems likely that this resistance is chromosomally encoded.

### DISCUSSION

Whereas the type strain of *C. testosteroni* contains no detectable plasmids, the more recent isolates from the environment (strains T-2 and PSB-4) contain conjugative megaplasmids from at least three incompatibility groups (Table 4), and these plasmids can maintain themselves in different *C. testosteroni* hosts. We are as yet uncertain whether all three plasmids are conjugal; plasmid pPSB from strain PSB-4 could be transferred among strains of *C. testosteroni*, whereas transfer and maintenance of plasmid pTSA were always in the presence of plasmid pT2T (strain T-2 and transconjugant T2-TYPE) or pPSB (transconjugant T2-PSB4).

Plasmid pTSA, at 85 kbp, is sufficiently large to carry all the genes required for conjugation, which together with the replicatory functions, require some 40 kbp (see Introduction). The catabolic genes [\(tsaMBCD\) with regulator \(tsaR\) and \(psbA\) (C)] (Fig. 1) and presumably those for a transport function for at least toluenesulfonate (13) can, thus, be easily accommodated.

IncP1 plasmids have an average GC content of 62% (23), which we have confirmed for one gene on plasmid pTSA (62.5% for \(trfA\)2 PCR product) (Fig. 4). The GC content of the \(tsaMBCD\) genes, 70%, indicates a different origin of these genes. Chromosomal DNA of *C. testosteroni* also has an average GC content of 62% (42), which is valid for strain T-2 (4), so the origin of the \(tsaMBCD\) genes remains unclear. A GC content of 70% tends to indicate *Pseudomonas*-like organisms.

IS1071 is found on plasmid pUO1, in *Moraxella* sp. strain B, flanking a dehalogenase (\(dehH2\)) (43). Transposon \(Tn5271\) (17 kbp) of *Alcaligenes* sp. strain BR60 is a composite transposon...
flanked by IS1071 (20). This class I element contains three
genes (cbaABC, which encode 3-chlorobenzoate-3,4 dioxygen-
nase, the corresponding reductase, and a dehydrogen-
ase, respectively) and is located on the 88-kbp IncP1 plasmid,
pBRC60. Tn5271 can integrate into the chromosome when
conjugated with the incompatible plasmid pGS65 or R68
and can be remobilized by plasmid pBRC40 or R68, re-
spectively. It has also been reported that Tn5271 transposes into C.
testosteroni ATCC 11996 (43). However, IS1071 (3.2 kbp
in total with 110-bp inverted repeats on either side) itself belongs
to the class II transposons, and in spite of a missing resolvase,
it has been shown to be a functional transposon (43). CbaAB is a
class IA multicompartment mononuclear iron oxygenase
system, whose oxygenase component has 45% identity with the
N-terminal region of PsbA (class IA) over the available 35-
amino-acid protein microsequence (19, 21), which could indi-
cate a higher than average similarity of the two enzymes in the
class IA.

Both 85-kbp plasmids pTSA and pPSB contain two copies of
insertion element IS1071, which suggests the presence of a
composite transposon possibly analogous to Tn5271. We con-
clude that the transconjugant PSB4-TYPE (PSB4) is in part
generated by a transposition event involving IS1071, so we
assume that genes psbAC are included in the transposition
element, and we are examining this element. Apart from pos-
sible similarities to Tn5271, the oxygenase component PsbA of
dioxigenase PsbAC is identical in plasmid pTSA from strain
T-2 and in plasmid pPSB from strain PSB-4 (14), whereas the
reductase has been lost from, or inactivated in, strain T-2,
where the regulation has also suffered alteration (29). We hope
to establish what led to these changes.

The strains T-2 and PSB-4 have not only the plasmid-borne
IS1071 sequences but also presumably a chromosomal copy of
this insertion element. Wyndham et al. (43) suggested a mech-
nism whereby this could occur during generation of a com-
posite transposon.

We find that the TER genes (R2) (Fig. 1) are chromosom-
ally located, and we presume that the PCA genes (R4), with all
meta cleavage pathway genes, are also chromosomal because
the plasmid-free type strain utilizes PCA (Table 3). In contrast,
the toluenesulfonate genes (R1) and the sulfobenzoate genes
(R3) are located on a plasmid. So the degradation of both
tolueneconcarboxylate via TER and that of toluenesulfonate via
sulfobenzoate in strain T-2 are examples of complementation
between chromosomally and plasmid-encoded catabolic path-
ways which differ in detail. Plasmid pTSA carries all genes (R1
and R3) necessary to degrade toluenesulfonate to PCA; there-
after, the chromosomal genes are required. Tolueneconcar-
boxylate, via TsaMBCD, is taken only to TER before the chromo-
some supports the hypothesis that the evolution of degra-
dative plasmids is based on the assembly of previously inde-
pendent pathways mediated by flexible genetic elements, and
more recent work has indicated the modular nature of these
events (38).

Surprisingly, this would appear to be the first report of
catabolic genes on a defined plasmid in C. testosteroni,
despite the very large number of degradative abilities reported in this
species and its widespread prevalence in the environment (42).
These data confirm preliminary results indicated by Hooper
(9) and suggest that the roles of, e.g., the broad-host-range
IncP1β plasmids and transposable elements in the genetic flex-
ibility of Comamonas spp. have been underestimated in the
past.

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REFERENCES

nology, Zurich, Switzerland.

2. Ausubel, F. M., R. Brent, R. E. Kingston, R. E. Moore, J. G. Seidman, J. A.
New York, N.Y.

Comparative genetic organization of incompatibility P group degradative

xenobiotic-degrading isolates from the beta subclass of the Proteobacteria by
a polyphasic approach including 16S rRNA partial sequencing. Int. J. Syst.

5. Cain, R. B. 1981. Microbial degradation of surfactants and “builder” com-
(ed.), Microbial degradation of xenobiotics and recalcitrant compounds.


31. Wyndham, R. C. Personal communication.