Rhodococcus opacus expresses the \( xsc \) gene to utilize taurine as a carbon source or as a nitrogen source but not as a sulfur source

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The Gram-positive bacteria *Rhodococcus opacus* ISO-5 and *Rhodococcus* sp. RHA1 utilized taurine (2-aminoethanesulphonate) as the sole source of carbon or of nitrogen or of sulfur for growth. Different gene clusters and enzymes were active under these different metabolic situations. Under carbon- or nitrogen-limited conditions three enzymes were induced, though to different levels: taurine–pyruvate aminotransferase (Tpa), alanine dehydrogenase (Ald) and sulfoacetaldehyde acetyltransferase (Xsc). The specific activities of these enzymes in *R. opacus* ISO-5 were sufficient to explain the growth rates under the different conditions. These three enzymes were purified and characterized, and the nature of each reaction was confirmed. Analyses of the genome of *Rhodococcus* sp. RHA1 revealed a gene cluster, \( \tau a u R-\text{ald-tpa} \), putatively encoding regulation and oxidation of taurine, located 20 kbp from the \( xsc \) gene and separate from two candidate phosphotransacetylase (pta) genes, as well as many candidate ABC transporters (\( \tau a u R\-\text{ald-tpa} \)). PCR primers allowed the amplification and sequencing of the \( \tau a u R\-\text{ald-tpa} \) gene cluster and the \( xsc \) gene in *R. opacus* ISO-5. The N-terminal sequences of the three tested proteins matched the derived amino acid sequences of the corresponding genes. The sequences of the four genes found in each *Rhodococcus* strain shared high degrees of identity (>95% identical positions). RT-PCR studies proved transcription of the \( xsc \) gene when taurine was the source of carbon or of nitrogen. Under sulfur-limited conditions no \( xsc \) mRNA was generated and no Xsc was detected. Taurine dioxygenase (TauD), the enzyme catalysing the anticipated desulfonative reaction when taurine sulfur is assimilated, was presumed to be present because oxygen-dependent taurine disappearance was demonstrated with taurine-grown cells only. A putative \( tauD \) gene (with three other candidates) was detected in strain ISO-5.

Regulation of the different forms of metabolism of taurine remains to be elucidated.

INTRODUCTION

Taurine (2-aminoethanesulphonate) is a ‘phylogenetically ancient compound’ (Huxtable, 1992), which has long been known as a sole source of sulfur for growth of aerobic microorganisms (Huxtable, 1992; Uria-Nickelsen et al., 1993). The compound also serves as a source of carbon and energy for many aerobic bacteria (Cook & Denger, 2002), and Chien et al. (1999) considered it to be a source of carbon, energy, nitrogen and sulfur for growth of *Rhodococcus* spp., though the nutritional elements were not all tested individually. Seitz et al. (1993) isolated bacteria which could apparently utilize taurine as a source of carbon and sulfur, but which mutated to lose the ability to utilize it as a source of carbon: this indicates that different enzymes are involved in the physiologically different processes of carbon dissimilation, on the one hand, and assimilation of sulfonate-sulfur on the other (cf. Cook & Denger, 2002; Kertesz, 2000). Kertesz (2000) reviewed the nature of the desulfonative oxygenase (taurine dioxygenase [EC 1.14.11.17], TauD) (Elkins et al., 2002) (Fig. 1B) involved in assimilation of sulfur from taurine in aerobic bacteria, and the global regulation of its expression. Cook & Denger (2002) reviewed the desulphonation of sulfoacetaldehyde involved in the dissimilation of taurine (sulfoacetaldehyde acetyltransferase [EC 2.3.3.15], Xsc) (Fig. 1A), and postulated regulated gene clusters involved in taurine catabolism in different bacteria (see also Brüggemann et al., 2004; Ruff et al., 2003).

*Rhodococcus opacus* ISO-5 was isolated for its ability to desulphonate many aromatic compounds (Schleheck et al., 2003). Following on the work of Chien et al. (1999), the organism was found to utilize taurine as a sole source of

Abbreviations: Ald, alanine dehydrogenase; PLP, pyridoxal 5’-phosphate; Pta, phosphate acetyltransferase; TauD, taurine deoxygenase; TauR, taurine regulator; Tpa, taurine–pyruvate aminotransferase; ThDP, thiamin diphosphate; Xsc, sulfoacetaldehyde acetyltransferase.

The GenBank accession numbers for the sequences reported in this paper are: \( xsc_{\text{ISO-5}} \) AY498611; \( tauR\-\text{ald-tpa}_{\text{ISO-5}} \) AY498612.
carbon (Ruff et al., 2003) via an Xsc encoded by a gene in xsc subgroup 1 (Cook & Denger, 2002), or as a sole source of sulfur (Schleheck et al., 2003). It was also observed that *Rhodococcus* sp. strain RHA1 contained potential *xsc* and *tauD* genes (W. W. Mohn, unpublished data). Further analysis of the genome sequence showed the presence of candidates for genes encoding a regulator (TauR) (Brüggemann et al., 2004), taurine–pyruvate aminotransferase (Tpa) [EC 2.6.1.77] and alanine dehydrogenase (Ald) [EC 1.4.1.1], which suggested that the dissimilatory pathway in Fig. 1(A) was operative in *Rhodococcus* spp. This dissimilatory pathway is widespread (Cook & Denger, 2002), but analyses of the first genome sequences, which were found to contain putative *xsc* genes, reintroduced the alternative pathway via taurine dehydrogenase [EC 1.4.99.2, probably EC 1.4.2.], TauXY (Brüggemann et al., 2004; Kondo et al., 1973; Ruff et al., 2003).

We now report that *R. opacus* ISO-5 and *Rhodococcus* sp. strain RHA1, which were isolated on different continents for reasons independent of taurine metabolism, supplied two independent, but almost identical, pictures of the utilization of the individual nutritional macroelements of taurine, with the advantage that the genome of one of these organisms is being sequenced. Taurine was dissimilated via Tpa, which had not been observed previously in a sequenced organism. The dissimilatory genes, and those encoding assimilative desulfonation, seem not to be clustered in either organism. Further, the high level of identity between corresponding genes in the two organisms was not matched in the DNA sequences surrounding those genes. The use of two organisms thus allowed us to confirm the generality of the phenomena under study.

**METHODS**

**Material.** Commercial chemicals were of the highest purity available, and they were purchased from Fluka, Roth, Merck, Serva or Sigma. The preparation of sulfoacetaldehyde, as the bisulfite addition complex, was described elsewhere (Denger et al., 2001).

**Organisms, growth, harvesting of cells and preparation of cell-free extracts.** *Rhodococcus* opacus ISO-5 was isolated in this laboratory (Schleheck et al., 2003) and deposited with the German Culture Collection DSMZ, Braunschweig, Germany, under the accession number DSM 44600. *Rhodococcus* sp. strain RHA1 was a kind gift of W. W. Mohn, University of British Columbia, Vancouver, Canada.

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**Fig. 1.** Pathway of dissimilation of taurine in *R. opacus* ISO-5 with putative homeostatic mechanisms to detoxify sulfite and maintain the osmolarity in the cell (A), and dioxygenolytic desulfonation of taurine to aminoacetaldehyde for the assimilation of taurine sulfur (B). Different transport systems are inferred for the high fluxes involved in supplying carbon and nitrogen from taurine for growth (A), compared with the low flux and high affinity required to supply sulfur from taurine (B). Abbreviations not used in the Introduction: PLP, pyridoxal 5'-phosphate; ThDP, thiamin diphosphate.
Cells under carbon-limited conditions were grown aerobically at 30 °C in a phosphate-buffered mineral-salts medium (Thurnheer et al., 1986) with 10–20 mM taurine or acetate, or 3 mM benzoate, p-aminobenzenesulfonate, 4-sulfobenzoate, 4-phenolsulfonate, 4-toluene sulfonate, benzenesulfonate, salicylate, 3-phenylpropionate, 4-phenylbutyrate or phenylacetate as the sole added source of carbon. Nitrogen-limited cells were grown aerobically at 30 °C in the same medium but with the omission of ammonium chloride and the addition of 5 mM glucose or 20 mM acetate as carbon source; taurine or ammonium ion (2 mM) served as sole added source of nitrogen. Sulfur-limited Tris-buffered salts medium (Laue et al., 1996) with glucose or acetate as carbon source was used for sulfur-limited conditions. In this medium 100 µM taurine or sodium sulfate, or 30 µM 4-toluene-, benzene-, anthracene-2-, p-nitrobenzene-, p-aminobenzenesulfonic-, p-chlorobenzenesulfonic-, 1,2-ethanedi-, or phenylacetate, or 4-phenylbutyrate or phenylacetate as the sole added source of sulfur.

Precultures and cultures for the determination of the substrate range were grown in 30 ml screw-cap tubes in a roller. Growth experiments were done on the 100 ml scale in 1 litre Erlenmeyer flasks on a shaker. Samples were taken at intervals to measure optical density for chromatography. This generated a precipitate, which was removed by centrifugation (10 000 g, 3 min, room temperature). The clear supernatant fluid was loaded on to a Mono Q anion-exchange column and protein was eluted as described elsewhere (Ruff et al., 2003). Representative fractions were assayed for Tpa, Ald and Xsc.

Pooled fractions with Xsc activity were concentrated, desalted with equilibrated PD 10 columns (Pharmacia) according to the manufacturer’s instructions and subjected to a second separation on Mono Q equilibrated with 20 mM potassium phosphate buffer of higher pH (pH 6-9).

Analytical methods. Absorbance was measured in a Uvikon 922 spectrophotometer (Kontron). Sulfate was quantified as turbidity in the presence of BaCl₂ (Sörbo, 1987). Ammonium ion was assayed colorimetrically by the Berthelot reaction (Gesellschaft-Deutscher-Chemiker, 1996). Acetyl phosphate was determined chemically as iron(III) acetyl hydroxamate (Racker, 1962; Stadtman, 1957). Reversed-phase HPLC was used to quantify taurine (Laue et al., 1997) or sulfoacetaldehyde (Cunningham et al., 1998) after derivatization. Acetate was quantified by GC (Laue et al., 1997). Protein was assayed by protein-dye binding (Bradford, 1976) or with whole cells by a Lowry-type method (Cook & Hütter, 1981). SDS-PAGE and staining were done by standard methods (Laemmli, 1970) to monitor protein purification and to estimate molecular masses under protein-denaturing conditions. The sequence of the N-terminal amino acids was determined by Edman degradation under contract by the sequencing facility of the Fachhochschule Bingen, Germany (http://spa.ihb-bingen.de/). Values of apparent Kₘ were derived by hyperbolic curve-fitting as cited elsewhere (Ruff et al., 2003); the relevant ranges of concentration of substrates were 0–50 mM taurine with Tpa, 0–25 mM alanine for Ald and 0–20 mM sulfoacetaldehyde for Xsc.

Amplification, nucleotide sequencing and sequence analysis of genes. These were done as described elsewhere (Brüggemann et al., 2004); PCR primers are given in Table 1. The protein sequences initially used in BLAST searches of the genome of Rhodococcus sp. strain RHA1 were TauR, TauABC, TauXY, Xsc and Pta from Burkholderia sp. strain LB400 (Brüggemann et al., 2004; Ruff et al., 2003), and Tpa (AF269146) and Ald (AF269148) from Bilephila wadsworthia.

RT-PCR. Cells for the preparation of total RNA were grown in the required selective medium and harvested in the mid-exponential growth phase (100–140 mg protein 1⁻¹); the bacterial culture was stabilized by addition of RNAProtect Bacteria Reagent (Qiagen) prior to harvesting. Total RNA from these cells (0·1 g wet weight) was prepared with the RNaseasy Mini Kit and the RNase-free DNase set (Qiagen) following the protocol of the manufacturer, but with the following modifications: (i) resuspended cells in lysis buffer (manufacturer’s instructions) were disrupted in a bead-mill homogenizer (Fastprep FP120; Savant Instruments) at a setting of 6·5 ms⁻¹ for 45 s, and the lysate was centrifuged (12 000 g, 5 min, 4 °C) to remove particulate matter; (ii) ethanol was added to the supernatant fluid (manufacturer’s instructions), and this solution was vortexed (5 s) to yield clumps of precipitated (chromosomal)
DNA, which were discarded prior to loading this solution on to the RNeasy-column (manufacturer’s instructions). The First Strand cDNA Synthesis Kit (MBI Fermentas) was used for reverse transcription of RNA (10^3 µg) according to the manufacturer’s instructions, and with the use of primer Xsc RT-r (Table 1). RNA was quantified photometrically (A_260). cDNA from these reverse transcriptions was used as template (2 µl of reaction mixture) for PCR reactions (20 µl final volume) to amplify an 860 bp fragment of xsc (primer pair Xsc RT-f and Xsc RT-r; Table 1) under standard PCR conditions as described previously (Tralau et al., 2003). Controls without reverse transcriptase were used to detect any contamination of undigested DNA in the RNA preparations.

### RESULTS

**Carbon-limited growth with taurine initiated by transamination and desulfonation**

*R. opacus* ISO-5 utilized taurine (or acetate) as sole source of carbon and energy for aerobic growth with a molar growth yield that showed mass balance for carbon (Table 2) and a specific growth rate (μ = 0.12 h⁻¹) which allow the specific utilization rate for taurine to be calculated as 3·1 mkat (kg protein)⁻¹. The sulfonate-sulfur was

### Table 1. PCR primers and anticipated product lengths with selected primer pairs

Analyses with the BLAST algorithm allowed a provisional annotation of genes in the genome of *Rhodococcus* sp. strain RHA1. Gene-specific PCR primers were selected or derived as appropriate. The PCR primers for the different xsc subgroups were derived from sequence alignments.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Fragment length with given primer</th>
<th>Source of information</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ald1</td>
<td>CCGGCAGCGAGTGAGGTA</td>
<td>729 with Tpa1</td>
<td>RHA1 genome sequence</td>
</tr>
<tr>
<td>Tpa1</td>
<td>GGGATGTCTSTYGGCGGCTT</td>
<td>1289 with Xsc_{sg1}-r</td>
<td>This paper</td>
</tr>
<tr>
<td>Xsc_{sg1}-f</td>
<td>GCGCCGGGATACGACATCGGG</td>
<td>726 with Xsc_{sg1}-r</td>
<td>This paper</td>
</tr>
<tr>
<td>Xsc_{sg1}-r</td>
<td>GGGGTTGTAGAAGTCCACCTG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Xsc_{sg3}-f</td>
<td>TGGGCTACACCGGGTGCGAA</td>
<td>860 with Xsc_{sg3}-r</td>
<td>RHA1 genome sequence</td>
</tr>
<tr>
<td>Xsc_{sg3}-r</td>
<td>GCCSGGACGTGGATATTTGG</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Xsc_{RT-f}</td>
<td>GATCATCAGRTGCTGACGCA</td>
<td>420 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
</tr>
<tr>
<td>Xsc_{RT-r}</td>
<td>YIGGACATCGRTGCTGACGCA</td>
<td>118 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
</tr>
<tr>
<td>TauD_{uni}-f</td>
<td>CAAGGTCTGKTYYTSCG</td>
<td>331 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
</tr>
<tr>
<td>TauD_{uni}-r</td>
<td>YIGGACATCGRTGCTGACGCA</td>
<td>262 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
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<tr>
<td>TauD_{uni}-f</td>
<td>CATCCGGGCTCGTGACGAT</td>
<td>155 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
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<tr>
<td>TauD_{uni}-r</td>
<td>CATCCGGGCTCGTGACGAT</td>
<td>155 with TauD_{uni}-r</td>
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<td>TauD_{uni}-f</td>
<td>TGCCAACCTGGCGGATCGTG</td>
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<tr>
<td>TauD_{uni}-r</td>
<td>TGCCAACCTGGCGGATCGTG</td>
<td>155 with TauD_{uni}-r</td>
<td>RHA1 genome sequence</td>
</tr>
</tbody>
</table>

### Table 2. Growth of *R. opacus* ISO-5 in different media, with data on the presence of Tpa, Ald and Xsc detected as enzyme activity and by SDS-PAGE, and data on the transcript of the xsc gene

The acetate-salts medium (control experiment) contained ammonium ion as nitrogen source and sulfate ion as sulfur source.

<table>
<thead>
<tr>
<th>Growth medium containing taurine as sole source of:</th>
<th>Carbon</th>
<th>Nitrogen</th>
<th>Sulfur</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molar growth yield*</td>
<td>6·0</td>
<td>5·4</td>
<td>56·3</td>
</tr>
<tr>
<td>Activity of Tpa [mkat (kg prot.)⁻¹]</td>
<td>ND</td>
<td>2·4</td>
<td>0·7</td>
</tr>
<tr>
<td>Induced 50 kDa protein (Tpa)†</td>
<td>ND</td>
<td>+ +</td>
<td>+</td>
</tr>
<tr>
<td>Activity of Ald [mkat (kg prot.)⁻¹]</td>
<td>ND</td>
<td>10·2</td>
<td>2·3</td>
</tr>
<tr>
<td>Induced 41 kDa protein (Ald)†</td>
<td>ND</td>
<td>+ +</td>
<td>+</td>
</tr>
<tr>
<td>Activity of Xsc [mkat (kg prot.)⁻¹]</td>
<td>ND</td>
<td>3·0</td>
<td>1·5</td>
</tr>
<tr>
<td>Induced 64 kDa protein (Xsc)†</td>
<td>ND</td>
<td>+ +</td>
<td>+</td>
</tr>
<tr>
<td>Detected xsc mRNA†</td>
<td>ND</td>
<td>+ +</td>
<td>+</td>
</tr>
</tbody>
</table>

ND, Not detected.

*Mean molar growth yields are about 6 g protein (mol C)⁻¹, 50 g protein (mol N)⁻¹ and 3000 g protein (mol S)⁻¹ (Cook, 1987).

†Intensity of bands observed by SDS-PAGE or RT-PCR: + +, strong signals; +, weak signals.
recovered as sulfate, which was released concomitantly with growth, and ammonium ion was recovered in the growth medium (not shown). Traces of transient sulfite were also detected in the medium (not shown). *Rhodococcus* sp. strain RHA1 behaved similarly (not shown), which shows that the growth we observed previously (Cook & Denger, 2002; Ruff *et al.*, 2003) reflected overall mass balances.

The BLAST algorithm was used to make comparisons of genes encoding components of pathways of taurine metabolism (see Introduction and Fig. 1) with preliminary sequence data from the genome of strain RHA1. A single putative *xsc* gene was found. It was not contiguous (about 20 kbp distant from the *tauR-tpa-ald* quasi-cluster; see below) with genes known or presumed to be relevant to the regulation, transport or metabolism of taurine. PCR primers derived from the sequence of the transport or metabolism of taurine. PCR primers derived from the sequence of the *xsc* gene in strain RHA1 (*xsc*RHA1) allowed a fragment of DNA from strain ISO-5 to be amplified and sequenced. The sequence of the *xsc*ISO-5 gene shared 97 % identical positions with *xsc*RHA1. This *xsc* gene allowed amplification of the anticipated product from the consensus PCR primers for *xsc*-subgroup 1 (Table 1) but not from primers for subgroup 3 (Table 1). There is negligible identity between the sequences of subgroup 1 and the few known members of subgroup 2, which is found in low G+C content Gram-positive bacteria, so the latter subgroup is also not represented. Further analyses of the genome in strain RHA1 indicated the presence of homologues of (i) a gene (*tauR*) postulated to encode regulation of taurine metabolism, (ii) three possible genes (*tpa*) encoding taurine–pyruvate amino-transferase, (iii) two possible genes (*ald*) encoding alanine dehydrogenases, (iv) two possible genes (*pta*) encoding phosphate acetyltransferases, and (v) many possible ATP-binding-cassette transporters (*tauBC*); the latter were not further examined. There was one apparent cluster, potentially *tauR-ald-tpa*, which was sequenced in strain ISO-5 as indicated above for the *xsc* gene. Homologues of the genes (*tauXY*) presumed to encode taurine dehydrogenase (Brüggemann *et al.*, 2004) were either absent (*tauX*), or shared a low level of identity (*tauY*; 28 % identical positions).

Strain ISO-5 was grown with acetate or taurine as sole source of carbon, and cell extracts were prepared. Inducible formation of *Tpa* [to 2·4 mkat (kg protein)]−1, *Ald* [to 10·2 mkat (kg protein)]−1 and *Xsc* [to 3·0 mkat (kg protein)]−1 (Table 2) was detected in taurine-grown cells; *Pta* was apparently constitutive [0·1 mkat (kg protein)]−1 and was not examined further. No taurine dehydrogenase was detected, as predicted from the absence of *tauXY* (see above). Proteins in these extracts were separated by SDS-PAGE, which showed the strong induction of three proteins (molecular masses of 50, 41 and 64 kDa) in the taurine-grown cells (Fig. 2); similar data were obtained with strain RHA1 (not shown). Enzyme purification from strain ISO-5 (see below) showed these proteins to be *Tpa* (50 kDa), *Ald* (41 kDa) and *Xsc* (64 kDa) (Fig. 2); the molecular masses corresponded to those derived from the gene sequences (see below).

A single *Tpa* eluted (at about 140 mM Na2SO4 in the gradient) at >95 % purity from the anion-exchange column (Fig. 2); the specific activity of this material [15·0 mkat (kg protein)]−1 represented about a sixfold purification. The identity of the enzyme, suggested by the routine assay by HPLC in which taurine disappeared and alanine was formed, was confirmed by following the reverse reaction in which sulfoacetaldehyde disappeared. The apparent *Km* value for taurine was calculated to be 42 ± 2 mM. The N-terminal heptapeptide was determined to be MDVTELR. This was identical with the sequence deduced from *tpa*ISO-5, assuming that the GUG codon (nominally encoding valine) encodes fMet (Kozak, 1999) under these conditions. This then allowed *tpa*RHA1 to be identified and distinguished from the other candidate genes to encode *Tpa* proteins (N-terminal sequences MTTVTNS or MSMASAL) of strain RHA1. Gene *tpa*ISO-5 was 1371 bp long.

A single *Ald* eluted (at about 130 mM Na2SO4 in the gradient) at >99 % purity from the anion-exchange.
The specific activity of this material (47.8 mkat (kg protein)^{-1}) represented about a fivefold purification. The identity of the enzyme, indicated photometrically by the alanine-dependent reduction of NAD^+ to NADH, was confirmed by the reverse reaction in which ammonium ion, pyruvate and NADH were essential to form alanine, which was quantified, after derivatization, by HPLC. The apparent \( K_m \) value for alanine was determined to be 3.9 ± 0.5 mM. The N-terminal amino acid sequence was determined to be MRIGIPK, which was identical to that derived from the nucleotide sequence of \( ald_{SO-5} \) and to the sequence data of one RHA1 gene, which we annotated \( ald \). The other candidate gene in strain RHA1 was derived to contain a different N-terminus (MKIGIPR); this allele presumably encodes an Ald for a different pathway.

A single Xsc eluted (at about 160 mM Na_2SO_4 in the gradient) at about 90% purity from the anion-exchange column; the specific activity of this material (30 mkat (kg protein)^{-1}) represented about a tenfold purification. The purification was improved (Fig. 2, lane 5) by subsequent anion-exchange chromatography under slightly different conditions (not shown). The enzyme was identified by observing that sulfaoxaldehyde, phosphate and ThDP were essential to generate sulfite and acetyl phosphate. The apparent \( K_m \) value for sulfaoxaldehyde was calculated to be 2.6 ± 0.6 mM, typical for the subgroup 1 enzymes. The N-terminal amino acid sequence was determined to be XXXXNGADR. Allowing for cleavage of the initiatory methionine, an identical pentapeptide at the correct position was derived from the nucleotide sequence of the putative \( xsc_{SO-5} \) and from the sequence data of strain RHA1. The complete sequence of \( xsc \) of strain ISO-5 (1809 bp) is amongst the longest of the 21 known \( xsc \) sequences; only the gene from *Achromobacter xylosoxidans* NCIMB 10751 is longer (1812 bp).

RT-PCR analysis showed that mRNA was transcribed from the \( xsc \) gene during growth with taurine as a carbon source (Fig. 3). The gene was not transcribed when acetate was the carbon source (Table 2). It was also not transcribed when either sulfate or taurine was the sole source of sulfur (Table 2).

Several separate loci are required for taurine dissimilation. The \( tauR-ald-tpa \) cluster involves divergent transcription of the \( ald \) and \( tpa \) genes, which are distant from the unknown genes encoding transport, from \( xsc \) and from \( pta \). This situation differs from that in several \( z \)-Proteobacteria, where the genes are clustered (Brüggemann et al., 2004).

**Nitrogen-limited growth with taurine also initiated by transamination and desulphonation**

*R. opacus* ISO-5 utilized taurine as sole source of nitrogen for aerobic growth with a molar growth yield that showed mass balance for nitrogen (Table 2) and a specific growth rate (\( \mu = 0.1 \) h^{-1}) which allow the specific utilization rate for taurine to be calculated as 0.5 mkat (kg protein)^{-1}. The

### Sulfur-limited growth with taurine independent of Xsc

Strain ISO-5 utilized taurine as the sole source of sulfur for aerobic growth (\( \mu = 0.11 \) h^{-1}) with the standard molar growth yield (Table 2); this represents a specific degradation rate of 10 \( \mu \)kat (kg protein)^{-1}. Similar data were obtained for *Rhodococcus* sp. strain RHA1 (not shown). No Xsc was detected in these cells, no induction of Xsc was detected by SDS-PAGE (Fig. 2) and no mRNA from the \( xsc \) gene was detected (Table 2). So the utilization of taurine was independent of Xsc, as predicted in the Introduction.

The predicted desulphonation reaction under these metabolic conditions is catalysed by taurine dioxygenase, TauD (see Introduction). Cells of strain ISO-5 grown with limiting sulfate as the sulfur source did not catalyse disappearance of taurine (Fig. 4). In contrast, cells grown with taurine as the sole sulfur source displayed oxygen-dependent taurine disappearance (Fig. 4) with a specific activity of 9.3 \( \mu \)kat (kg protein)^{-1}. This was interpreted as preliminary evidence for the presence of taurine dioxygenase, TauD. The genome of strain RHA1 was found to contain a major candidate for a \( tauD \) gene (no gaps) with three minor candidates (with gaps). In addition, a putative \( atsK \) gene, also encoding a
were set up with primer tauDuniv-r for reverse transcription, during growth of strain ISO-5 with taurine as sulfur source. The gene shared two common segments of sequence that allowed a PCR primer pair (tauD univ-f and tauD univ-r) (Table 1) to be generated, which should amplify a fragment from each of these four genes in strain RHA1, and from the tauD gene from Pseudomonas putida, Pseudomonas aeruginosa or E. coli. The amplification yielded two bands, one with the expected size range (420 nt) and one (530 nt) which was generated with one primer alone (tauD univ-r). We developed specific forward primers for each candidate gene (tauD cand1-f to tauD cand4-f) (Table 1), but we had to use tauD univ-r as the reverse primer in each case. The 420 nt band was used as a template, and the amplification products we obtained (118, 155, 262 and 331 nt; not shown) showed that all four major and minor candidates for the tauD gene were present in strain ISO-5 (see Table 1). RT-PCR experiments to detect mRNA transcribed from a tauD-candidate during growth of strain ISO-5 with taurine as sulfur source were set up with primer tauD univ-r for reverse transcription, and with the primer pair tauD univ-f plus tauD univ-r for subsequent PCR. This yielded neither a specific 420 nt fragment nor a non-specific 530 nt fragment (not shown). We suspected that this particular mRNA is transcribed only in low levels (corresponding to the low specific activity of TauD, see above), and we desisted from further RT-PCR experiments.

It thus seems likely that TauD is involved in the desulfonation of taurine under sulfur-limiting conditions, but it is currently impracticable to identify which of the candidate genes encodes this TauD.

Differences between R. opacus ISO-5 and Rhodococcus sp. strain RHA1

We have sequenced five genes in strain ISO-5 (tauR, ald, tpa, xsc and putative atsK) and we observed about 95% identical positions in the corresponding genes in strains ISO-5 and RHA1. In addition, the four candidate tauD genes were detected in strain ISO-5 using the primers developed for strain RHA1, so high levels of identity are obviously present here, too. We then wondered whether we could detect any significant differences between the organisms. One difference was that PCR primers from the 3′ end of the tauR RHA1 gene or further downstream gave no amplification product with any potential reverse primer from the 5′ end of the gene. Analyses with the BLAST algorithm indicated potential ssuD, asfAB, asaA and atsK genes on the RHA1 genome, and the corresponding sulfur substrates (see Methods), arylsulfonates (ssu genes), arylsulfonates (asaA genes), alkylsulfonates (asaA and atsK genes), and putative (phenyl)propionate and 4-(phenyl)butyrate, but they differed in that only strain RHA1 could utilize phenylacetate.

DISCUSSION

The dissimilation of taurine via Tpa is widespread in Gram-positive and Gram-negative bacteria, and the pathway in aerobes is now known to end at acetyl-CoA, where it enters the Krebs and glyoxylate cycles (Fig. 1) (Cook & Denger, 2002). Activity of each of the four metabolic enzymes involved, Tpa, Ald, Xsc and Pta, can be assayed in R. opacus ISO-5 and Rhodococcus sp. strain RHA1. Whereas Tpa, Ald and Xsc are obviously induced to specific activities [2–10 mkat (kg protein)]−1 which can essentially explain the growth rate [3·1 mkat (kg protein)]−1, we wonder whether we may have suboptimal conditions for the assay of the apparently constitutive Pta.

We have confirmed directly that taurine nitrogen is a sole source of nitrogen for growth of two Rhodococcus spp. (compare Chien et al., 1999) in the presence of excess carbon source. The organisms use the same degradative pathway [Tpa, Ald, Xsc (Table 2), and presumably Pta], because sulfate is released, and involve the only xsc gene (Fig. 3), so presumably the same regulator (putative TauR) is involved. TauR obviously controls the level of expression economically, because different levels of expression are found appropriate to the different fluxes that are required [3·0 and 1·5 mkat (kg protein)]−1].

Another aspect of the dissimilation of taurine that requires regulation is the excretion of sulfate and sulfate on the one hand, and ammonium ion on the other (Fig. 1). This control will require modulation when the cells utilize taurine as a sole source of nitrogen, where the same regulator, TauR, presumably in cooperation with other factors, will presumably prevent the export of ammonium
ion, while still excreting sulfate. Aspects of nitrogen control are discussed elsewhere (e.g. Magasanik, 1996).

Tpa [EC 2.6.1.77] has been purified only once before (Laue & Cook, 2000a), but a gene whose product encodes the Tpa involved in the anaerobic assimilation of taurine-sulfur in *Rhodobacter capsulatus* has been identified (Masepohl et al., 2001). These three gene products, Tpa\(_baw\), Tpa\(_bc\) and Tpa\(_is\) (with the almost identical Tpa\(_RHA1\)), belong to COG1061, according to the Domain Search Tool of the NCBI BLAST server, but the levels of identity amongst them is low: Tpa\(_RHA1\), Tpa\(_bw\) and Tpa\(_bc\) share 99%, 32% and 30% identical positions, respectively, with Tpa\(_is\), while Tpa\(_bw\) and Tpa\(_bc\) share 57% identical positions. This poor correlation of sequence with function suggests that, of the three *tpa*-like genes in strain RHA1, only that annotated *tpa* actually encodes a Tpa. Heterogeneity in aminotransferases is, apparently, not unusual (Mehta & Christen, 2000). The initial codon of the *tpa* gene, GUG, obviously encodes Met. This codon is frequently used in strain RHA1 (W. W. Mohn, personal communication) so *Rhodococcus* sp. strain RHA1 presumably uses this initial codon much more commonly than the 10% reported for prokaryotes in a review of initiation of translation (Kozak, 1999).

There was only one candidate *xsc* gene in strain RHA1, though candidate genes for other ThDP-coupled enzymes, e.g. *ilvB*, were detected (see Ruff et al., 2003). As with Tpa and Ald, the N-terminal amino acid sequence of Xsc obtained with microsequencing confirmed the tentative identification of the corresponding gene. The *xsc\(_RHA1\)* gene (and *xsc\(_ISO-5\)*) shared 78–87% identical positions with the other nine genes in subgroup 1 of *xsc*, and the appropriate portion of the gene was amplified with the consensus primers for subgroup 1 (Table 1): these data confirm and expand the unpublished data in our earlier attribution (Ruff et al., 2003). Up till now, we had the impression that subgroup 1 of *xsc* was restricted to *β*-Proteobacteria. Obviously the subgroup is also found in Gram-positive organisms, which have a similar G+C content to that in the *β*-Proteobacteria.

Just as there was only one candidate for the *xsc* gene in strain RHA1, there was just one candidate for the putative regulator gene, *tauR* (dendrogram in Brüggemann et al., 2004) and the level of identical positions in the corresponding section of incomplete *taurISO-5* was 95%. If *TauRISO-5* is closely related to homologues, the pattern of regulation of genes is somewhat different. In the *α*-Proteobacteria, the regulated genes are located in a cluster that could be an operon (in *Sinorhizobium meliloti*), whereas the cluster in the *β*-Proteobacteria (e.g. *Burkholderia fungorum* LB400) is a less structured entity (see Brüggemann et al., 2004). In strains RHA1 and ISO-5 there is a different start of transcription for *tpa, ald*, *xsc* and transport, while *pta* may be differently regulated. As this is the first transamination pathway for taurine dissimilation to be explored on a sequenced genome, it is possible that this non-contiguous pattern is normal. However, preliminary annotation of the genome of *Rhodococcus* sp. strain RHA1 indicates that most pathways are encoded on clustered genes (W. W. Mohn, unpublished data).

*R. opacus* ISO-5 was isolated for its ability to desulfonate quantitatively a multitude of complex sulfonated aromatic compounds (Schleheck et al., 2003) as sulfur sources for growth, and *Rhodococcus* sp. strain RHA1 can also desulfonate many aromatic sulfonates (listed in Methods). Genes corresponding to the known desulfonation complex of desulfonation enzymes (Ssu and Ats) are present in strain RHA1. Several other desulfonation or sulfoesterase enzymes must be present in both organisms, and the corresponding genes can be detected in the RHA1 genome. These enzymes were initially found in Gram-negative bacteria, each encoded in a cluster which also encodes an ABC transport system (Kertesz, 2000, 2001; Kertesz & Kahnert, 2001). Many ABC transporter systems can be detected in the RHA1 genome, sometimes with a potential A gene (periplasmic binding protein), and often without. In no case is a potential ABC transporter encoded adjacent to genes encoding enzymes to release sulfite of sulfate from a source of sulfur. This situation resembles that seen above for the dissimilation of taurine.

There is mass balance for the assimilation of taurine-sulfur (Table 2) and the specific degradation rate of taurine calculated under these conditions [10 μkat (kg protein\(^{-1}\)] can be observed in whole cells (Fig. 4). Xsc is not involved in this assimilation (see Table 2), and we interpret the enzymic and PCR data to indicate the presence of TauD. Independent sets of metabolism for quantitative dissimilation of taurine-carbon or the quantitative assimilation of taurine-sulfur are thus confirmed, which is scarcely surprising given the need for a high turnover of taurine at high concentrations [0.5–5 μkat (kg protein\(^{-1}\) at 2–10 mM taurine] or a low turnover at low concentrations [10 μkat (kg protein\(^{-1}\) at 30 μM]. When taurine is serving as a source of sulfur under anoxic conditions, however, TauD (taurine dioxygenase) cannot be functional, and a *tpa* gene with an *xsc* gene have been shown in genetic experiments to be involved in desulfonation in *Rhodobacter capsulatus* (Masepohl et al., 2001), while another group has found a *Tpa* under corresponding conditions in *Clostridium pasteurianum* (Chien et al., 1997), so presumably an Xsc is involved in the desulfonation. It remains to be seen whether the kinetic properties of these desulfonative enzymes involve a higher affinity for the sulfonate than that (about 5 mM) found in the subgroup 1 enzymes examined to date (Ruff et al., 2003).

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