3-Sulphocatechol 2,3-dioxygenase and other dioxygenases (EC 1.13.11.2 and EC 1.14.12.--) in the degradative pathways of 2-aminobenzenesulphonic, benzenesulphonic and 4-toluenesulphonic acids in Alcaligenes sp. strain 0-1

Frank Junker, Thomas Leisinger and Alasdair M. Cook†

Author for correspondence: Alasdair M. Cook. Tel: +49 7531 88 4247. Fax: +49 7531 88 2966.

Alcaligenes sp. strain O-1 utilizes three sulphonated aromatic compounds as sole sources of carbon and energy for growth in minimal salts medium – benzenesulphonate (BS), 4-toluenesulphonate (TS) and 2-aminobenzenesulphonate (2AS). The degradative pathway(s) in 2AS-grown cells are initiated with membrane transport, NADH-dependent dioxygenation and meta ring cleavage. The specific activity of the NADH-dependent dioxygenation(s) varied with the growth phase and was maximal near the end of exponential growth for each growth substrate. Cells were harvested at this point from BS-, TS- and 2AS-salts medium. Cells grown with each sulphonated substrate could oxygenate all three compounds, but only 2AS-grown cells consumed 2 mol O₂ per mol 2AS or BS or TS. BS- and TS-grown cells consumed 2 mol O₂ per mol BS or TS but failed to oxygenate the product of oxygenation of 2AS, 3-sulphocatechol (3SC). These observations were repeated with cell extracts and we concluded that there were two sets of desulphonative pathways in the organism, one for 2AS and one for BS and TS. We confirmed this hypothesis by separating the degradative enzymes from 2AS-, BS- or TS-grown cells. A 2AS dioxygenase system and a 3SC-2,3-dioxygenase (3SC230) were detected in 2AS-grown cells only. In both BS- and TS-grown cells a dioxygenase system for BS and TS was observed as well as a principal catechol 2,3-dioxygenase (C230-III), neither of which was present in 2AS-grown cells. The 3SC230 was purified to near homogeneity, found to be monomeric (Mr, 42 000), and to catalyse 2,3-dioxygenation to a product that decayed spontaneously to sulphite and 2-hydroxymuconate. The 2AS dioxygenase system could cause not only deamination of 2AS but also desulphonation of BS and TS. The BS dioxygenase could desulphonate BS and apparently either desulphonate or deaminate 2AS. Strain 0-1 thus seems to contain two putative, independently regulated operons involving oxygenation and spontaneous desulphonation(s). One operon encodes at least the 2AS dioxygenase system and 3SC230 whereas the other encodes at least the BS/TS dioxygenase system and C230-III.

Keywords: Alcaligenes sp. strain O-1, desulphonation, multi-component dioxygenases, meta ring cleavage, convergent degradative pathways

INTRODUCTION

Four mechanisms of desulphonation of aromatic compounds have been elucidated, all of which involve oxygenation directly or indirectly (Cook & Leisinger, 1991; Junker et al., 1994). Two mechanisms involve...
Fig. 1. Initial steps in the degradative pathway(s) for BS, TS and 2AS in 2AS-grown cells of *Alcaligenes* sp. strain O-1. All the illustrated reactions have been observed (Thurnheer et al., 1990; Junker et al., 1994). No structure for the ring-cleavage product of 4-methylcatechol is shown, because we do not know whether 2,3- or 1,6-cleavage occurs. The presence of an arrow, e.g. 'transport', need not mean that a specific enzyme is present purely for that reaction.

Oxygenases are enzymes which introduce one or two atoms of oxygen from molecular oxygen into a substrate which is usually organic. These enzymes are found in some ten subclasses defined by the Enzyme Commission (e.g. Webb, 1992). Of those subclasses, two types of dioxygenases are relevant to this paper, (i) the multi-component enzyme systems (EC 1.14.12.-) which typically initiate metabolic attack on inactive aromatic rings (Batie et al., 1992; Correll et al., 1992; Harayama et al., 1992; Mason & Cammack, 1992), and (ii) the somewhat simpler dioxygenases (EC 1.13.11.-) involved in ring-cleavage reactions (e.g. Lipscomb et al., 1992). The enzymes of EC 1.14.12.- can be further subdivided, but they all involve an electron transport chain of varying complexity which transfers electrons from NADH via a flavine and one or more [2Fe-2S] centres to the Rieske [2Fe-2S] centre on the oxygenase and then presumably to the mononuclear iron site where oxygen is activated prior to its attack on the named (non-reactive aromatic) substrate. By this means the aromatic substrate is activated and made available to less complex enzymes. The activation can lead to the spontaneous elimination of stable substituents (e.g. sulphono, halo, nitro, amino and ether functions) by generating unstable intermediates (a ring carbon carrying a hydroxyl group and a heteroatom) which lead to spontaneous loss of an anion (e.g. the good leaving groups sulphite, halide, nitrite, ammonia and phenolate/alkoxylate; Engesser et al., 1989; Junker et al., 1990).

Strain O-1 is known to utilize three sulphonated aromatic compounds as sole sources of carbon and energy for growth, orthanilate (2-aminobenzenesulphonate, 2AS), benzenesulphonate (BS) and 4-toluenesulphonate (TS) (Thurnheer et al., 1986). The degradative pathways for BS, TS and 2AS in cells grown in 2AS-salts medium have been elucidated and found to consist of transport, NADH-dependent dioxygenation and *meta* ring cleavage (Fig. 1) (Thurnheer et al., 1990; Junker et al., 1994; cf. Locher et al., 1993). Given the different mechanisms of desulphonation in Fig. 1, associated with either the NADH-dependent dioxygenase (BS and TS) or with the *meta* ring cleavage (2AS), it is unclear how many sets of enzymes are involved in the three pathways. One of those sets, however, that for the degradation of 2AS, is plasmid-encoded (Jahnke et al., 1990).
Some of these enzymes can eliminate several anions (cited in Junker et al., 1994), others are highly specific (e.g. Locher et al., 1991). The dioxygenases of EC 1.13.11.2 are also somewhat heterogeneous (Harayama et al., 1992; Happe et al., 1993) but they are "simple" in that only one type of subunit is involved. The activation of oxygen is at an iron centre, but no external reductant is needed. The reaction of catechol 2,3-dioxygenase opens the aromatic ring to yield a linear chain (e.g. Bayly & Barbour, 1984; Whittam et al., 1991; Junker et al., 1994).

We initially anticipated there to be several multi-component desulphonative dioxygenases in strain O-1, because of the different patterns of apparent specific activities against a range of substrates observed in cell-free extracts from cells grown with the different arylsulphonates (Thurnheer et al., 1986). The problems involved in quantifying the multi-component dioxygenases became apparent to us only later (Geary et al., 1990; Thurnheer et al., 1990; Locher et al., 1991; Bünz & Cook, 1993), when we realised that either constant protein concentrations had to be used in assays, or that additions of otherwise limiting components were required. This uncertainty in the measurement of activity leads to uncertainty in the attribution of activity to a putative enzyme. So we presumed that our initial data on rates in crude extracts (Thurnheer et al., 1986) were suspect, because they were obtained at various protein concentrations; correspondingly the hypothesis on the number of oxygenases was suspect. The realization that different desulphonative mechanisms (Thurnheer et al., 1990; Junker et al., 1994) are present (Fig. 1) brought us back to the hypothesis of multiple sets of enzymes.

We have now shown that Alcaligenes sp. strain O-1 has two sets of specifically induced, desulphonative pathways, and that the substrate ranges of the enzymes in these two pathways in part overlap.

METHODS

Materials. The preparation of 3SC (Junker et al., 1994), the sources of chemicals (Junker et al., 1994; Thurnheer et al., 1986, 1990), as well as the commercial columns for enzyme separation and the molecular mass markers (Locher et al., 1991) are described elsewhere.

Analytical methods. Spectrophotometric and optical density measurements (Thurnheer et al., 1986; one OD unit at 500 nm represented 160 mg protein 1⁻¹), reversed phase HPLC (Grosenbacher et al., 1986; Locher et al., 1989) and oxygen uptake measurements (Zamanian & Mason, 1987) were done with apparatus described previously. Protein from whole cells was measured in a Lowry-type assay (Kennedy & Fewson, 1968); protein in crude extracts and purifications was measured by the method of Bradford (1976). SDS-PAGE was done as previously described (Laemmli, 1970; Locher et al., 1991). The N-terminal amino acid sequence of a blotted protein was determined by automated Edman degradation (Locher et al., 1991).

The organism, its growth and the preparation of cell extracts. Alcaligenes sp. strain O-1 (DSM 6325; Thurnheer et al., 1986; Jahnke et al., 1990) was maintained, grown and harvested under optimal conditions for the multi-component dioxygenase, as described elsewhere (Junker et al., 1994) or as indicated in the Results section. Cell pellets were suspended in extraction buffer (50 mM Tris/HCl, pH 7.5, containing 20%, v/v glycerol), brought to 1 mM PMSF, disrupted in a French pressure cell and centrifuged to remove particulate material (Junker et al., 1994). The supernatant fluid, called crude extract, was used for some experiments. Otherwise, crude extract treated with streptomycin sulphate to remove nucleic acids (Junker et al., 1994) was used for enzyme purifications.

Enzyme assays. 2-Aminobenzenesulphonate 2,3-dioxygenase system (2ASDOS), benzensulphonate 1,2-dioxygenase system (BSDOS) or toluene-4-sulphonate 3,4-dioxygenase system (TSDOS) in crude extract were routinely assayed at 30 °C as the rate of 2AS-dependent (or BS-dependent or TS-dependent) oxygen uptake in 0.25 ml reaction mixtures. These contained about 12 μmol Tris/HCl, pH 7.5, 0.5 mg protein, 150 nmol NADH and 750 nmol O₂ (or BS or TS) with which the reaction was being examined, 20 nmol portions of sulphonate were used. When components of these multi-component enzyme systems were being located in separated protein fractions, 80 μl portions of different fractions were used, irrespective of the protein concentration. The location of the components could be deduced from their colour (cf. Mason & Cammack, 1992), the reductase being yellow and the oxygenase red. The reaction of the reductase was confirmed by its reaction with dichlorophenol indophenol and with cytochrome c (cf. Locher et al., 1991).

3-Sulphocatechol 2,3-dioxygenase (3SC230) and catechol 2,3-dioxygenase(s) (C230) in crude extract were routinely assayed photometrically (374 nm) at room temperature as formation of 2-hydroxymuconate semialdehyde from catechol (cf. Bird & Cain, 1974). Reaction mixtures (0.5 ml) contained 23 μmol Tris/HCl, pH 7.5, about 50 μg protein, 10 nmol FeSO₄, and 5 μmol catechol with which the reaction was started. When fractions from protein separations were examined, no attempt was made to set the protein concentration in the assay. These assays could also be quantified as substrate-dependent oxygen uptake rates with (substituted) catechols. When stoichiometry was being examined, 20 nmol portions of catechols (catechol, 4-methylecatechol or 3SC) were used in determinations of oxygen uptake.

Representative experiments involving oxygen uptake were tested for substrate disappearance and product formation. The reaction mixture was brought to pH 2 with HCl to precipitate protein, which was removed by centrifugation. The supernatant fluid was examined by HPLC.

Separation and purification of enzymes. The enzymes studied were all sensitive to oxygen, so all solutions were sparged with oxygen-free nitrogen or helium prior to use. Where appropriate, solutions were protected by a blanket of nitrogen. Preliminary experiments were done with streptomycin-treated extracts of 2AS-grown cells. We were unable to use a precipitation with ammonium sulphate to treat the extract, because there was extreme loss of activity (≥ 60%) which was not regained on reconstituting the fractions. Further, this extract was a stable solution of proteins which was unaltered visually by freezing and thawing, but during (or immediately after) anion-exchange chromatography many proteins precipitated. We thus chose a robust and relatively inexpensive DEAE-Sepharose column for the first separative step.

Step 1. The DEAE column (15 × 2.6 cm) at 4 °C was used in an
identical manner with extracts obtained from 2AS-, BS- or TS-grown cells. The column was equilibrated with 50 mM Tris/HCl buffer, pH 7.5, containing 0·1 mM DTT (buffer A) at 2·5 ml min⁻¹ and extract (400 mg protein) was applied to the column; 8 ml samples were collected. Buffer A was pumped for about 30 min when a linear gradient to 60% buffer B (1 M Tris/HCl, pH 7·5 containing 0·1 mM DTT) over 3 h was started.

Separated fractions from BS-grown cells were tested for BSDOS and C23O, fractions from TS-grown cells for TSDOS and C23O, and fractions from 2AS-grown cells for 2ASDOS, C23O and 3SC23O. Attempts to concentrate proteins by membrane filtration led to extensive loss of activity, so active fractions were routinely desalted with a Sephadex G-25 column equilibrated with buffer C (50 mM Tris/H₂SO₄, pH 7.5 containing 0·1 mM DTT) and used immediately. Occasionally, pooled fractions maintained on ice were brought to 80% saturation with (NH₄)₂SO₄ to precipitate proteins which were collected by centrifugation, taken up in a small volume of buffer C and desalted.

Step 2. A pre-packed anion exchanger column (Mono Q; 10 x 100 mm) was routinely equilibrated at room temperature with buffer C at 0·7 ml min⁻¹ and 1 ml samples were collected. A sample of 2 to 5 ml of (3S)C23O from step 1 was loaded on to the column and a first linear gradient to 10% buffer D (buffer C containing 1 M Na₂SO₄) was applied over 10 min. This was followed by a second linear gradient to 20% buffer D over 30 min, and a step gradient to 100% buffer D for 10 min. A step gradient to 100% buffer C for 15 min regenerated the initial conditions. Fractions were tested for C23O activity. Active fractions were pooled and used immediately and without further treatment in step (3).

C23O-I was examined in a modified step (2) in which all 50 mM buffers were replaced by 20 mM buffers. BS(TS)DOS from step (1) was separated under routine conditions, where no single fraction catalysed BS (TS)-dependent oxygen uptake, but combinations of fractions were active.

Step 3. A gel-filtration column (Superose-6; 10 x 300 mm) was equilibrated at room temperature with buffer C containing 150 mM Na₂SO₄ at 0·6 ml min⁻¹, and 1 ml fractions were collected. Preparative separations were done with 2 ml portions applied to the column. Values for native M, were obtained by chromatographing 50 µl portions and interpolating data into standard curves generated by chromatographing 50 µl of standard proteins.

RESULTS

Growth of Alcaligenes sp. strain O-1 in BS- and TS-salts medium

Strain O-1 grew in BS-salts medium (μ = 0·16 h⁻¹; Fig. 2) and growth was concomitant with substrate utilization (not shown). The BS-dependent uptake of oxygen by washed cells from the culture, an assay for the putative BSDOS, was initially undetectable, increased to a maxi-

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**Table 1.** Comparison of oxygen uptake rates with various sulphonate substrates in differently grown cells of Alcaligenes sp. strain O-1 and in extracts from these cells

<table>
<thead>
<tr>
<th>Growth substrate</th>
<th>Ratios of oxygen uptake rates with individual substrates under different conditions</th>
<th>Oxygen uptake rate with BS [mkat (kg protein)⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BS:2AS</td>
<td>BS:TS</td>
</tr>
<tr>
<td>2AS</td>
<td>0·4</td>
<td>0·9</td>
</tr>
<tr>
<td>BS</td>
<td>2·6</td>
<td>2·0</td>
</tr>
<tr>
<td>TS</td>
<td>2·9</td>
<td>1·8</td>
</tr>
</tbody>
</table>
Table 2. Stoichiometry of oxygen uptake with different substrates by whole cells of *Alcaligenes* sp. strain 0-1 grown with one of three sulphonated substrates.

The values represent the mean of three determinations. No assay differed by more than 10% from the data shown.

<table>
<thead>
<tr>
<th>Growth substrate</th>
<th>Stoichiometry of oxygen consumption with different substrates (mol O₂ (mol substrate)⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2AS</td>
</tr>
<tr>
<td>2AS</td>
<td>2-0</td>
</tr>
<tr>
<td>BS</td>
<td>1-0</td>
</tr>
<tr>
<td>TS</td>
<td>NA*</td>
</tr>
</tbody>
</table>

*NA* No assay; the TS-grown cells had a high rate of endogenous respiration and no accurate data could be obtained for the poor substrate 2AS.

The stoichiometry of oxygen uptake for different substrates also showed differences between 2AS-grown cells, on the one hand, and BS- and TS-grown cells, on the other (Table 2). Whereas BS or TS caused the utilization of 2 mol O₂ (mol sulphonate)⁻¹, and 1 mol O₂ (mol catechol or 4-methylcatechol)⁻¹ under all conditions, 2AS caused the uptake of 2 mol O₂ (mol 2AS)⁻¹ solely in 2AS-grown cells. BS- or TS-grown cells catalysed the consumption of only 1 mol O₂ (mol 2AS)⁻¹ (where measurable) and the corresponding catechol, 3SC, was not oxygenated. Similar data were obtained with crude extracts (not shown).

We thus hypothesized that there are two NADH-dependent dioxygenases, 2ASDOS and BSDOS, and that the latter is identical with TSDOS. It was unclear whether both may be present under some conditions. There must also be two meta cleavage enzymes, one of which does not desulphonate 3SC.

**Oxygen uptake by whole cells and cell extracts**

Oxygen uptake by optimally grown whole cells from 2AS-, BS-, TS- or succinate-salts medium was examined with the substrates 2AS, BS, TS and succinate. Succinate-grown cells showed significant oxygen uptake with succinate, but they showed no oxygen uptake with 2AS, BS or TS. The enzymes required for degradation of the arylsulphonates are thus inducible, in agreement with earlier data (Thurnheer et al., 1986).

2AS-, BS- and TS-grown cells showed two patterns of oxygen uptake (Table 1). 2AS-grown cells had a high uptake rate with 2AS compared with BS and TS (ratio BS:2AS of 0-4). In both BS- and TS-grown cells, BS led to the highest uptake rate and 2AS the lowest rate of uptake (ratio BS:2AS of 2:6 to 2:9). The ratios of different activities (Table 1) indicate that BS- and TS-cells could have one enzyme complement, probably expressed at different levels in BS- and TS-cells (Table 1, right-hand column), whereas 2AS-cells have a different set of enzymes. The oxygen uptake rates by cell extracts displayed analogous behaviour to whole cells, which supports the conclusion from experiments with whole cells (Table 1).

The stoichiometry of oxygen uptake for different substrates also showed differences between 2AS-grown cells, on the one hand, and BS- and TS-grown cells, on the other (Table 2). Whereas BS or TS caused the utilization of 2 mol O₂ (mol sulphonate)⁻¹, and 1 mol O₂ (mol catechol or 4-methylcatechol)⁻¹ under all conditions, 2AS caused the uptake of 2 mol O₂ (mol 2AS)⁻¹ solely in 2AS-grown cells. BS- or TS-grown cells catalysed the consumption of only 1 mol O₂ (mol 2AS)⁻¹ (where measurable) and the corresponding catechol, 3SC, was not oxygenated. Similar data were obtained with crude extracts (not shown).

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**Dioxygenases in strain O-1 grown in 2AS-, BS- or TS-salts medium**

Extract from cells grown optimally in 2AS-, BS- or TS-salts medium was applied to the DEAE anion-exchange column and the proteins were separated under identical conditions. Two patterns were observed. In extracts of 2AS-grown cells, the 2ASDOS was separated into two fractions, iv (red in colour) and v (yellow in colour) in Fig. 3(c), and one catechol 2,3-dioxygenase, 3-sulphocatechol-2,3-dioxygenase (3SC230, vi in Fig. 3(c)) was detected. No other oxygenase was observed. The fractions of 2ASDOS were attributed to a reductase (fraction v), which reduced cytochrome c and dichlorophenolindophenol, and to an oxygenase (fraction iv) which was active only in the presence of the reductase. The separations of proteins from BS- and TS-grown cells gave chromatograms which were apparently identical with one another (Fig. 3a, b). The putative BSDOS (TSDOS) eluted in one peak (iii in Fig. 3a, b) with different
The multi-component dioxygenases

The separated 2ASDOS oxygenated the three tested substrates, 2AS, BS and TS, and one product was observed in each case (3SC, catechol and 4-methylcatechol, respectively). These products were identified by co-chromatography (HPLC) with the appropriate authentic material and by identity of the appropriate UV spectra with the spectrum of the standard (Junker et al., 1994; see Thurnheer et al., 1990 for representative spectral data). The enzyme system thus has a wide substrate range and is apparently capable of causing the removal of either the amino substituent or a sole sulphono substituent on the ring.

The multi-component oxygenase from BS- or TS-grown cells could also oxygenate the three substrates tested. The product from BS (TS) was catechol (4-methylcatechol), which was identified by co-chromatography and UV spectra (see previous paragraph). In the case of 2AS, two products were detected in low yield. One product was 3SC, identified by co-chromatography and by UV spectra. The other product was presumably the unstable 3-aminocatechol, which we deduced from its chromatographic behaviour (retention time about 4.8 min) and UV spectrum (λmax 243 nm; λmax 272 nm) because these compare with data we generated earlier (4.6 min: 243 nm; 272 nm: Thurnheer et al., 1990). We have thus the same enzyme system (BSDOS) in both BS- and TS-grown cells. BSDOS also appears to have a wide substrate range and to catalyse both desulphonation and deamination.

The purification of 3SC23O and the separation of other meta cleavage enzymes

3SC23O could be purified to about 90% purity in a three-step procedure (Fig. 4, Table 4). The enzyme would appear to be a monomer (Table 3) with the following N-terminal amino acid sequence: Met-Gln-Val-Arg-Leu-Ile-Leu-X-Ser-His-Ser-Pro-Leu-Met-Leu-Lys-Glu-Met-Pro-Thr-Leu-Pro-. This sequence was not detected in databases (GenEMBL and SwissPro). The enzyme is very labile and shortly after the purification the activity is permanently lost.
Table 3. Properties of the four catechol meta cleavage enzymes observed under the experimental conditions used

<table>
<thead>
<tr>
<th>Catechol 2,3-dioxygenase*</th>
<th>Elution from anion exchanger†</th>
<th>Mₐ values‡</th>
<th>Substrate spectrum§</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DEAE (Tris/HCl)</td>
<td>Mono Q (Tris/H₂SO₄)</td>
<td>Native</td>
</tr>
<tr>
<td>From 2AS-grown cells</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3SC230 (100%)</td>
<td>320 mM</td>
<td>130 mM</td>
<td>46000</td>
</tr>
<tr>
<td>From BS-grown cells</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C230-I (1 %)</td>
<td>None</td>
<td>20 mM</td>
<td>ND</td>
</tr>
<tr>
<td>C230-II (&lt; 1 %)</td>
<td>None</td>
<td>100 mM</td>
<td>ND</td>
</tr>
<tr>
<td>C230-III (98 %)</td>
<td>140 mM</td>
<td>110 mM</td>
<td>79000</td>
</tr>
<tr>
<td>From TS-grown cells</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C230-II (7 %)</td>
<td>None</td>
<td>100 mM</td>
<td>87000</td>
</tr>
<tr>
<td>C230-III (93 %)</td>
<td>140 mM</td>
<td>110 mM</td>
<td>79000</td>
</tr>
</tbody>
</table>

ND, Not determined.
* Numbers in parentheses are percentages of the total activity of meta-cleavage enzyme present, measured with catechol as substrate.
† The values refer to the concentration of the gradient at which the peak-maximum eluted.
‡ The values were obtained by gel-filtration chromatography (native) and SDS-PAGE (denatured).
§ Abbreviations: CAT, catechol; 3SC, 3-sulphocatechol; 3MC, 3-methylcatechol; 4MC, 4-methylcatechol; 23B, 2,3-dihydroxybenzoate. The numbers refer to the relative specific activities of the enzyme preparations with 100 representing the specific activity with catechol as a substrate.

Table 4. Purification of 3-sulphocatechol 2,3-dioxygenase

The table is representative of six purifications.

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Vol. (ml)</th>
<th>Total activity (µkat)</th>
<th>Protein (mg ml⁻¹)</th>
<th>Specific activity [µkat (kg protein)⁻¹]</th>
<th>Yield (%)</th>
<th>Purification (-fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude extract</td>
<td>25</td>
<td>2.3</td>
<td>17</td>
<td>5.5</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>DEAE</td>
<td>47</td>
<td>1.9</td>
<td>1.8</td>
<td>22</td>
<td>83</td>
<td>4:1</td>
</tr>
<tr>
<td>Mono Q</td>
<td>20*</td>
<td>1.7</td>
<td>0.94</td>
<td>92</td>
<td>75</td>
<td>17</td>
</tr>
<tr>
<td>Superose-6</td>
<td>20*</td>
<td>1.0</td>
<td>0.60</td>
<td>87</td>
<td>45</td>
<td>16</td>
</tr>
</tbody>
</table>

* Multiple portions of the preceding pool were chromatographed.

3SC230 was the only catechol dioxygenase detected in 2AS-grown cells. It was not detectable in extracts of BS- or TS-grown cells (Fig. 3). Extracts of BS- or TS-grown cells, in contrast, contained multiple C230s, none of which oxygenated 3SC (Fig. 3, Table 3). The major portion of the activity in each case was termed C230-III, which we presume to be identical in BS- and TS-cells, because the chromatographic behaviour on two columns, the Mₐ-values for native and denatured protein, and the substrate ranges were found to be identical.

The minor portions of C230 were termed C230-I and C230-II. None of the four catechol dioxygenases has the same pattern of substrate turnover (Table 3). C230-I was detected in extracts of BS-grown cells only. C230-II probably occurs in both BS- and TS-grown cells, but there was insufficient activity in extracts of BS-grown cells to confirm this. The level of activity of C230-II in TS-grown cells leads us to believe that the enzyme is involved in the degradation of substituted catechols in another pathway and that it has been induced by traces of, e.g. methylcatechol or even catechol.

DISCUSSION

2ASDOS and 3SC230 were not synthesized by Alcaligenes sp. strain O-1 in the absence of 2AS; the enzymes were induced specifically in 2AS-salts medium. Similarly,
BSDOS and C23O-III were induced specifically in BS- (or TS-) salts medium. The regulation of enzyme synthesis is thus highly specific. Given that other multi-component oxygenases and the corresponding ring cleavage enzymes are often encoded in operons (Harayama et al., 1992), we hypothesize that two independent operons encoding desulphonative pathways may be present in Alcaligenes sp. strain O-1.

In contrast to the high specificity of induction of these two putative operons, the enzymes they encode displayed a lower specificity. It was this phenomenon, indeed, which led us to examine the enzymes (see Introduction). It is now clear that 2ASDOS is a completely separate entity from BSDOS. The oxygenase components of the two systems elute from the DEAE column at 165 mM Tris and 200 mM Tris, respectively (Fig. 3), and it is the oxygenase component of the enzyme system that interacts with the substrate which is oxygenated. We thus see from Table 1 that BSDOS in crude extract of BS-grown cells oxygenates BS twice as fast as 2AS. 2ASDOS in crude extract oxygenates BS and AS at similar rates. The apparent specific activity of 2ASDOS is apparently too low (Table 1) to account for the oxygen uptake rate by whole cells, but the non-linear response of this multi-component system to protein concentration masks a higher real specific activity (Thurnheer et al., 1990; cf. Locher et al., 1991). Just as 2ASDOS oxygenates BS (or TS) to catechol (or 4-methylcatechol) with unit stoichiometry (Thurnheer et al., 1990), BSDOS oxygenates BS (or TS) to catechol (or 4-methylcatechol) with unit stoichiometry (this paper and F. Junker, unpublished data). The reductases from 2ASDOS and BSDOS also elute reproducibly at different positions on the gradient (250 and 200 mM, respectively). The main meta cleavage enzymes (3SC23O and C23O-III; Fig. 3, Table 3) in the two putative operons are also obviously different.

Whereas 2ASDOS or BSDOS catalyses the first metabolic reaction in the degradation of arylsulphonates by strain O-1, the first specific interaction of, e.g. 2AS with the cell is transport (Thurnheer et al., 1990; cf. Locher et al., 1993). We have no indication of how many transport systems are present or how these systems are regulated.

3SC23O catalyses the standard C23O reaction (Table 3; Junker et al., 1994) and is inhibited by 3-chlorocatechol (Junker et al., 1994), and would therefore appear to be a normal meta cleavage enzyme (EC 1.13.11.2). Most C23Os, however, do not seem to oxygenate 3SC (Table 3). If 3-chlorocatechol inhibits a C23O by oxygenation to the acyl chloride which derivatizes and permanently inhibits the enzyme (Bartels et al., 1984), why does the highly unstable acyl sulphonate (sulphonoaldehyde) not display the same inhibitory effect? Junker et al. (1994) produced mg-amounts of product from 3SC with 3SC23O, whereas µg-amounts of 3-chlorocatechol are inhibitory. Indeed, 3SC23O in crude extract, though temporarily inhibited by 3-fluorocatechol, recovers from the inhibition (A. M. Cook, unpublished). This information raises the question of the extent of inhibition of a C23O by derivatization (Bartels et al., 1984) and by chelation of the iron cofactor (Klecka & Gibson, 1981). We wonder whether the acyl sulphonate is stable in the active site, and is only unstable in solution. 3SC23O is also unusual amongst C23Os in being monomeric (Table 3). We know of only one other monomeric C23O, that in Spingomonas sp. strain RW1 (Happe et al., 1993), others being homomultimeric (Harayama et al., 1992; Kataeva & Golovleva, 1990). We plan to clone the gene encoding this seemingly unusual enzyme and, if appropriate from the sequence, attribute the enzyme to one of the known families of meta cleavage enzymes (cf. Harayama et al., 1992).

The instability of 3SC23O prevented extensive studies of its properties. Lipscomb & Orville (1992) have several suggestions for stabilizing these enzymes, and Happe et al. (1993) achieved satisfactory stability of a C23O by purifying under anaerobic conditions. Despite the poor stability of 3SC23O, the specific activity observed in crude extracts with catechol as substrate [5.5 mkat (kg protein)⁻¹; Table 4], which represents about 3-9 mkat (kg protein)⁻¹ with 3SC (Table 3), is sufficient to explain the flux through growing cells, which we calculate to be 0.8 mkat (kg protein)⁻¹ from the growth rate and the growth yield (cf. Junker et al., 1994). In agreement with this calculation, whole cells in suspension display oxygen uptake rates of this order of magnitude (Table 1).

A common factor amongst 3SC23O (EC 1.13.11.1–), 2ASDOS and BSDOS (both presumably EC 1.14.12.1–) is the apparently spontaneous elimination reaction (loss of the anion sulphite, ammonia and sulphite, respectively) induced by oxygenation in the named reaction (cf. Junker et al., 1994). The spontaneous reaction arises from the unstable formal intermediate generated by oxygenation. This intermediate is a cis-diol, one of whose carbon atoms is also bound to a heteroatom which is simultaneously a good leaving group. Spontaneous loss of the good leaving group is simultaneously a rearomatization to the catechol (Locher et al., 1991; Junker et al., 1994). We have previously discussed the possibility that the spontaneity may be only apparent, because we do not know to what extent specific catalysis or spontaneous decay cause loss of anion (Bünz & Cook, 1993; Junker et al., 1994). The data presented here argue more for spontaneity. The 2ASDOS fractions can cause not only deamination but also desulphonation, and the BSDOS fractions can cause not only desulphonation but also possibly deamination. The question can only be better resolved when pure enzymes become available.

Strain O-1 is not the only organism to catalyse two desulphonations. Wittich et al. (1988) describe a Moraxella sp. which obviously contains both a dioxygenase and a putative monooxygenase involved in desulphonation, though these activities do not seem to be stable in cell-free extracts. Strain O-1 thus remains the only organism where several desulphonations can be examined at the enzymic level.
With some notable exceptions, multiple-component (di-) oxygenases have seldom been examined at the biochemical level (Mason & Cammack, 1992). One reason for this is probably the oxygen-sensitive nature of the enzymic components. Another is the sometimes low yield of enzyme on disruption of cells (Table 1). Yet another is probably the large amount of enzyme required in assays (2 mg ml^{-1} in this work; cf. Thurnheer et al., 1990; Geary et al., 1990). We presume that another difficulty is harvesting the cells at the correct time. Fig. 2 shows the relatively sharp spike in specific activity of BSDOS (whether from BS- or TS-grown cells). The same behaviour is seen with 2ASDOS (Junker et al., 1994; cf. Jahnske et al., 1993) where unpublished data show the optimal timing to be in a 2 h window just before the end of growth, similar to data from Bünz & Cook (1993). Moodie et al. (1990) have localized the effect in benzoate dioxygenase to loss of the reductase.

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