Enzymic Basis For Leakiness of Auxotrophs for Phenylalanine in *Pseudomonas aeruginosa*

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The dual enzymic routes for phenylalanine biosynthesis that exist in *Pseudomonas aeruginosa* complicate the isolation of phenylalanine auxotrophs. Mutants blocked in each of the various phenylalanine-pathway steps are essential for full appreciation of the physiological nature and gene–enzyme relationships of this biochemical system. A leaky phenylalanine-requiring mutant of *P. aeruginosa* (PAT1051) was found to lack the bifunctional P-protein (chorismate mutase–prephenate dehydratase), but retained the monofunctional isozyme species of chorismate mutase (chorismate mutase-F) as well as cyclohexadienyl dehydratase (components of the arogenate ‘overflow’ route to phenylalanine). This is the first mutant of *P. aeruginosa* shown to be deficient in any enzyme specific for phenylalanine synthesis. It is concluded that although the arogenate pathway has the demonstrated potential to overproduce phenylalanine, the substrate levels normally available to the arogenate pathway in the wild-type are inadequate to satisfy the full metabolic demand for phenylalanine.

**INTRODUCTION**

In *Pseudomonas aeruginosa* two enzymic routes exist for the biosynthesis of phenylalanine (Patel et al., 1977). This biochemical duality (depicted in Fig. 1), widely distributed in Gram-negative bacteria (Jensen, 1985), complicates the selection by conventional techniques of phenylalanine auxotrophs since the blocking of one enzymic route to phenylalanine is masked by the presence of the alternative route. Despite this biochemical basis for ‘reluctant auxotrophy’ (Patel et al., 1978), nutritional mutants requiring phenylalanine to sustain wild-type growth rates have been reported. Whitaker et al. (1982b) found that several leaky auxotrophs having a requirement for phenylalanine were deficient in one of five aminotransferases capable of functioning in phenylalanine and tyrosine synthesis. Waltho (1972) described six transduction groups of phenylalanine-responding mutants. Mutants belonging to five of these groups possessed chorismate mutase and prephenate dehydratase activities resembling those seen in the wild-type; they were thought to possess partial enzyme blocks in the common aromatic pathway, although specific enzyme deficiencies were not demonstrated. This assumption is indeed consistent with our finding that in *P. aeruginosa*, phenylalanine is the first of the aromatic amino acids to become rate-limiting to growth when flux through the common aromatic pathway is decreased. Thus, when moderately inhibitory concentrations of glyphosate, a specific inhibitor of 5-enolpyruvylshikimate-3-phosphate synthase (Steinrücken & Amrhein, 1980), are administered to cultures of *P. aeruginosa*, phenylalanine alone restores the wild-type growth rate (A. Berry, unpublished data).

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Abbreviation: DAHP, 3-deoxy-δ-arabino-heptulosonate 7-phosphate.

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The *P. aeruginosa* transduction group PHE IV of Waltho (1972) was reported to consist of a mutant type lacking chorismate mutase. In view of the complexity of phenylalanine biosynthesis in *P. aeruginosa*, not known in 1972, a single mutation abolishing chorismate mutase is not likely. Since two chorismate mutase isozymes exist in *P. aeruginosa*, it seemed more probable that the quantitatively major isozyme, chorismate mutase-P (see Fig. 1), had been eliminated by mutation. This possibility seemed particularly likely since chorismate mutase-P channels chorismate toward phenylalanine (Calhoun et al., 1973a). In this report we show that a representative mutant of the PHE IV transduction group is not only deficient in chorismate mutase-P, but also lacks the second P-protein activity, prephenate dehydratase. This is the first mutant of *P. aeruginosa* known to be defective in any enzyme specific for phenylalanine synthesis, and the prospects for exploiting this mutant to probe the physiological nature of phenylalanine biosynthesis in *P. aeruginosa* are discussed.

**METHODS**

*Strains and culture conditions.* *P. aeruginosa* strains PAT2 (wild-type) and PAT1051 [a member of the PHE IV transduction group of phenylalanine-responding mutants (Waltho, 1972)] were obtained from B. W. Holloway (Department of Genetics, Monash University, Clayton, Victoria 3168, Australia). Cultures were grown at 37 °C with shaking at 300 r.p.m. in the minimal-salts medium described by Fischer et al. (1986). Glucose (0.5%, w/v) was used as the carbon source, and the medium was additionally supplemented with 50 μg L-phenylalanine ml⁻¹ wherever indicated.

*Preparation and fractionation of crude extracts.* Crude, desalted extracts were prepared as described by Byng et al. (1980) by sonication at 4 °C of cells harvested from 1 litre cultures. The extract buffer used was 50 mM-EPES/KOH, pH 7.9 (at 25 °C). Fractionation of crude extracts was done using 1.5 × 20 cm columns of DEAE-cellulose (Whatman DE52) equilibrated with the buffer used for extract preparation (see above). Samples of crude extracts (72.9 mg protein and 84.8 mg protein for PAT2 and PAT1051, respectively) were applied to separate columns, which were then washed with 100 ml of equilibration buffer to remove unbound proteins. The bound proteins were then eluted with linear gradients of KCl (0–0.5 M), prepared in 300 ml of equilibration buffer.

*Enzyme assays.* Chorismate mutase and prephenate dehydratase were assayed as described by Patel et al. (1977), while arogenate dehydratase was assayed using the HPLC method of Zamir et al. (1985). All reactions were done at 37 °C. Reaction mixture components are specified in the appropriate Figure and Table legends. Protein concentrations were estimated using the method of Bradford (1976).
**Materials.** L-Arogenate was prepared according to Zamir et al. (1980, 1983). Barium prephenate was isolated from the culture supernatant of a tyrosine auxotroph of *Salmonella typhimurium* (Dayan & Sprinson, 1970) and was converted to the potassium salt with 50 mM-potassium phosphate buffer (pH 7.0) before use. Chorismate was isolated from the medium of the triple auxotroph *Klebsiella pneumoniae* 62-1, and was purified as the free acid (Gibson, 1970). All other chemicals and Sephadex G-25 were obtained from Sigma. Protein assay reagent was obtained from Bio-Rad.

**RESULTS AND DISCUSSION**

*Comparison of enzyme properties of PAT1051 with the wild-type*

The wild-type parent strain, PAT2, is identical to *P. aeruginosa* PA01 with respect to the enzymic arrangement and allosteric regulation of the aromatic amino acid pathway (A. Berry, unpublished observations). Thus, the published reports on aromatic-pathway enzymology in *P. aeruginosa* PA01 (Calhoun et al., 1973a, b; Fiske et al., 1983; Jensen et al., 1973; Patel et al., 1977, 1978; Stenmark-Cox & Jensen, 1975; Whitaker et al., 1982a, b) are also typical of the data obtained for *P. aeruginosa* strain PAT2.

PAT1051 possessed distinctly low activities for both chorismate mutase and prephenate dehydratase, compared to the wild-type parent PAT2 (Table 1). Since the bifunctional P-protein accounts for the major portion of chorismate mutase activity as well as for the majority of prephenate dehydratase in crude extracts, the data of Table 1 suggested a deficiency of the bifunctional P-protein (chorismate mutase-prephenate dehydratase; see Fig. 1). The residual chorismate mutase activity present in crude extracts of PAT1051 was that expected for the chorismate mutase-F isozyme, while the residual prephenate dehydratase activity could be due to cyclohexadienyl dehydratase. Cyclohexadienyl dehydratase, originally denoted arogenate dehydratase (Patel et al., 1977), recognizes both prephenate and L-arogenate as substrate (Jensen & Fischer, 1986; Fischer & Jensen, 1986). Partially purified cyclohexadienyl dehydratase exhibits a ratio of about five in comparison of activity with prephenate or L-arogenate as substrate under standard assay conditions, consistent with the ratio seen for these activities in mutant PAT1051.

The prephenate dehydratase component of the P-protein from wild-type PAT2 is inhibited more than 90% under standard assay conditions at phenylalanine concentrations exceeding 6.04 mM (Fig. 2, inset). Fig. 2 gives the inhibition data obtained with a crude extract of PAT2, showing inhibition to level off at 55–60%. This result is consistent with expectations for a crude extract containing a mixture of feedback-resistant enzyme molecules (cyclohexadienyl dehydratase) and feedback-sensitive enzyme molecules (prephenate dehydratase). If the

**Table 1. Altered enzyme activities in strain PAT1051**

Crude extracts were prepared in 50 mM-EPPS buffer (pH 7.9 at 25 °C) as described in the text. All reaction mixtures contained 50 mM-EPPS buffer (pH 7.9 at 25 °C), substrate (at the concentration specified below), and crude-extract protein in a final volume of 200 μl. Chorismate mutase, prephenate dehydratase and cyclohexadienyl dehydratase were assayed using final substrate concentrations of 1.0 mM-chorismate, 1.0 mM-prephenate, and 1.1 mM-L-arogenate, respectively.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Arogenate dehydratase</th>
<th>Chorismate mutase</th>
<th>Prephenate dehydratase*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- Phe</td>
<td>+ Phe</td>
<td>- Phe</td>
</tr>
<tr>
<td>PAT2</td>
<td>3.6</td>
<td>31.0</td>
<td>66.0</td>
</tr>
<tr>
<td>PAT1051</td>
<td>2.8</td>
<td>4.2</td>
<td>15.3</td>
</tr>
</tbody>
</table>

* Determined in the absence (− Phe) or presence (+ Phe) of 100 μM-L-phenylalanine.
prephenate dehydratase activity present in PAT1051 is exclusively attributed to cyclohexadienyl dehydratase, then one would expect insensitivity to inhibition by phenylalanine because only the P-protein dehydratase of P. aeruginosa is sensitive to inhibition by phenylalanine (Calhoun et al., 1973a; Patel et al., 1977; Zamir et al., 1985). The data given in Table 1 show that the latter expectations were borne out, although prephenate dehydratase activity in PAT1051 was inhibited about 11% by phenylalanine. The error in the assay is no greater than 5%, and three different experiments gave a mean value of about 15% for inhibition of total prephenate dehydratase activity in PAT1051. Inhibition by antibody raised against purified P-protein from Acinetobacter calcoaceticus (courtesy of S. Ahmad) gave results consistent with as much as 14% of the prephenate dehydratase activity of PAT1051 being due to P-protein dehydratase. If so, PAT1051 may express as much as 5% of the normal P-protein dehydratase. Mutant PAT1051 is very stable and has never been observed to revert, even though selection for revertants during slow growth on minimal medium would be very strong. Perhaps a multi-site deletion has abolished the promoter and/or the promoter-proximal chorismate mutase portion of the P-protein gene, leaving a weak internal promoter that can account for minor expression of a feedback-sensitive prephenate dehydratase. Thus far, all attempts to find any dehydratase activity separable from cyclohexadienyl dehydratase in PAT1051 have yielded negative results (see next section).

The specific activity of cyclohexadienyl dehydratase was consistently lower in PAT1051 than in PAT2. In three different experiments PAT1051 possessed about 70% of the activity found in PAT2. (The error in the assay of cyclohexadienyl dehydratase with L-arogenate was no greater than 7%.) This phenomenon was not related to faster growth of PAT2 than of PAT1051 in minimal medium since phenylalanine-grown cultures gave similar results. The data also indicated that phenylalanine excess does not repress and phenylalanine limitation (PAT1051 grown without phenylalanine) does not derepress cyclohexadienyl dehydratase.
Fig. 3. Elution profiles of chorismate mutase isoenzymes, prephenate dehydratase and cyclohexadienyl dehydratase (CDT) from *P. aeruginosa* PAT1051 (a) and PAT2 (b) following fractionation of crude extracts by DEAE-cellulose chromatography as described in the text. Prephenate dehydratase activity (●) and chorismate mutase activity (○) are expressed as phenylpyruvate absorbance at 320 nm. For convenience, CDT activity peaks were initially located using prephenate as substrate. The identities of these peaks were then confirmed by assaying peak fractions using L-arogenate as substrate. The specific activities for the CDT peak fractions (using L-arogenate as substrate) were 44 and 25 nmol product formed min⁻¹ (mg protein)⁻¹ for PAT2 and PAT1051, respectively. The prephenate dehydratase (P-protein) from PAT2 was totally unreactive with L-arogenate as substrate. All reaction mixtures contained 50 mM-EPPS buffer (pH 7.9 at 25 °C), a 1.0 mM final concentration of the assay substrate and eluate protein in a final volume of 200 μl. . ., Distribution of protein measured as absorbance at 280 nm; ---, KCl gradient.

Chromatographic resolution of enzymes of phenylalanine biosynthesis from PAT2 and PAT1051

The lack of the P-protein in PAT1051 was confirmed by fractionating crude extracts by DEAE-cellulose chromatography. Fig. 3 shows a comparison of the elution profiles for chorismate mutase, prephenate dehydratase and cyclohexadienyl dehydratase from PAT2 (b) and PAT1051 (a). Total chorismate mutase activity from PAT2 eluted as two distinct peaks, a minor peak (isozyme chorismate mutase-F) and a major peak (isozyme chorismate mutase-P). Chorismate mutase-P co-eluted exactly with prephenate dehydratase, consistent with their joint physical association as a bifunctional P-protein (Patel *et al.*, 1977). Cyclohexadienyl dehydratase from PAT2 eluted earlier than the P-protein in the KCl gradient.

Only one peak of chorismate mutase activity was recovered from PAT1051 (Fig. 3a), which corresponded to the chorismate mutase-F isozyme. Cyclohexadienyl dehydratase activity was detected in the gradient fractions, as was expected from the presence of this activity in crude extract (see Table 1). Both activities of the P-protein were absent in PAT1051. Additional enzymic deficiencies were not detected in PAT1051 (data not shown). Thus, this strain resembled the wild-type with respect to the presence of two regulatory isozymes of 3-deoxy-D-arabino-heptulosonate-7-phosphate (DAHP) synthase, dehydroshikimate reductase, 5-enolpyruvylshikimate-3-phosphate synthase, prephenate dehydrogenase, arogenate dehydrogenase
and five separate aromatic aminotransferase activities (Calhoun et al., 1973a, b; Fischer et al., 1986; Fiske et al., 1983; Jensen et al., 1973; Patel et al., 1977, 1978; Stenmark-Cox & Jensen, 1975; Whitaker et al., 1982a, b).

The prephenate dehydratase component of the *P. aeruginosa* P-protein is activated by L-tyrosine (Calhoun et al., 1973a). In *Xanthomonas campestris* L-tyrosine not only activates the prephenate dehydratase of the P-protein, but is also required for stability (Whitaker et al., 1985). This and the observation that L-tyrosine stimulates the growth rate of *P. aeruginosa* PAT1051, suggested that PAT1051 might possess a labile P-protein that could be protected by L-tyrosine. However, an elution profile identical to that shown in Fig. 3(a) was obtained when a crude extract of PAT1051 was prepared and fractionated in the presence of 1.0 mM-L-tyrosine (data not shown).

**Biochemical and physiological implications**

We have shown that *P. aeruginosa* PAT1051 is deficient in the bifunctional P-protein (chorismate mutase–prephenate dehydratase), but retains both the minor isozyme of chorismate mutase (chorismate mutase-F) and cyclohexadienyl dehydratase. The joint elimination of both chorismate mutase-P and prephenate dehydratase as the consequence of a single mutation supports the contention that both catalytic activities are the property of a single protein. This bifunctional protein appears to be distributed throughout two of three major superfamilies of the Gram-negative bacteria (Ahmad & Jensen, 1986).

PAT1051 marks a starting point for strategies of selection for other mutant deficiencies in the post-prephenate portion of the aromatic pathway (see Fig. 1). For example, elimination of the P-protein in PAT1051 leaves only one route to phenylalanine intact (via cyclohexadienyl dehydratase). It should now be possible to isolate tight phenylalanine auxotrophs as single-step mutants by eliminating cyclohexadienyl dehydratase. It may then be possible to study mutants that are deficient only in cyclohexadienyl dehydratase following restoration of the P-protein gene. Secondly, the absence of chorismate mutase-P in PAT1051 leaves only chorismate mutase-F to provide all of the prephenate molecules needed for phenylalanine and tyrosine synthesis. Hence, a mutation that eliminates chorismate mutase-F in the genetic background of PAT1051 should yield a phenotype of auxotrophy for both phenylalanine and tyrosine. Nutritional and physiological studies of mutants lacking each of the pathway enzymes will provide insight into the significance of the dual-pathway arrangement which has proven to be widely distributed among Gram-negative bacteria (Jensen, 1985).

It has been shown that when DAHP synthase-tyr, the tyrosine-sensitive major isozyme of DAHP synthase in *P. aeruginosa* (Whitaker et al., 1982a), loses sensitivity to allosteric control by mutation, phenylalanine is overproduced owing to the absence of end-product control for the ‘overflow’ pathway (consisting of chorismate mutase-F and cyclohexadienyl dehydratase) (Fiske et al., 1983). Presumably, wild-type cells do not excrete phenylalanine because the overflow pathway requires elevated levels of phenylalanine-pathway precursors, a circumstance prevented by the controlling influence of tyrosine upon DAHP synthase-tyr. Our present results show that in the absence of the P-protein route through phenylpyruvate, the overflow pathway through L-arogenate and/or phenylpyruvate is not even adequate to provide sufficient phenylalanine to sustain the normal wild-type rate of growth.

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**REFERENCES**


