Organization of Interrelated Aromatic Metabolic Pathway Enzymes in 
Acinetobacter calco-aceticus

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INTRODUCTION

In Neurospora crassa and several other fungi, five sequential steps in the common aromatic biosynthetic pathway are catalysed by a multi-enzyme complex, encoded in a gene cluster (Giles, Case, Partridge & Ahmed, 1967a; Ahmed & Giles, 1969). In contrast, in bacteria the corresponding enzymes are for the most part separable, of low molecular weight and not encoded in a gene cluster (Berlyn & Giles, 1969). It has been suggested that the occurrence of the complex is related to the existence of a hydroaromatic degradative pathway which shares three metabolites and one enzymatic reaction with the aromatic biosynthetic pathway; competition for substrates would be alleviated by channelling the biosynthetic intermediates within the multi-enzyme complex (Giles, Partridge, Ahmed & Case, 1967b). However, the multi-enzyme complex and the degradative pathway do not always occur together (Ahmed & Giles, 1969; Berlyn & Giles, 1969). The reports by Canovas and co-workers (e.g. Tresguerres, deTorrontegui & Canovas, 1970a; Ingledew, Tresguerres & Canovas, 1971) concerning the occurrence and regulation of the hydroaromatic pathway in the moraxella species, Acinetobacter calco-aceticus, led us to examine the physical state of the biosynthetic enzymes in this organism. The pathways (cf. Giles et al. 1967b) are shown overleaf: the upper reaction sequence (a) leads to the eventual formation of acetyl CoA and succinate and the lower sequence (b) to the aromatic amino acids and vitamins.

METHODS

Acinetobacter calco-aceticus strain 73 originated from the Collection of the Department of Bacteriology and Immunology, University of California, Berkeley, U.S.A., and was obtained from Dr José Canovas, Instituto de Biología Celular, C.S.I.C. Madrid, Spain. Bacteria were grown in 500 ml of liquid minimal medium as used for Pseudomonas (Ornston & Stanier, 1966) containing either 20 mM-succinate, 10 mM-quininate or a combination of 20 mM-succinate and 2 mM-quininate in 2800 ml Fernbach flasks on a rotary shaker at 30 °C.

The extraction buffer was 0·1 M-potassium phosphate, pH 7·4, containing 2 × 10⁻³ M-dithiothreitol and 10⁻³ M-ethylenediaminetetraacetic acid. In some cases 0·1 M-KCl was included in both the buffer and the sucrose gradients. Several extraction procedures were used: bacteria suspended in buffer were subjected to sonic disintegration; frozen pellets were broken in a Hughes pressure cell at 98 MN/m² (17000 lb/in²) and suspended in buffer; bacterial suspensions were treated with lysozyme and subjected to osmotic shock as described by Tresguerres et al. (1970a). After disruption, suspensions were centrifuged at 27000g for 30 min and the supernatant was either dialysed and used as a crude enzyme preparation or treated with 1/10 volume of 1·4% (w/v) protamine sulphate, fractionated

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Quinic acid (QA) \[\text{HO} \text{COOH}\]
Shikimic acid (SA) \[\text{HO} \text{COOH}\]
Protocatechuic acid (PCA) \[\text{HO} \text{COOH}\]

**RESULTS**

The sedimentation patterns for the enzymes are shown in Fig. 1. The various methods of extraction and the use of crude or fractionated preparations did not alter the observed patterns. These enzymes occurred *in vitro* as distinct, rather low molecular weight proteins rather than as a multi-enzyme complex.

Two dehydroquinase isozymes were observed: a light, heat-labile enzyme found in quinate-grown bacteria (Fig. 1a) and a heavy, heat-stable enzyme found in both quinate- and succinate-grown bacteria (Fig. 1b). The average molecular weight values based on calculations from several sucrose gradients were: DHQ synthetase, 60000; biosynthetic dehydroquinase, 125000; inducible, degradative dehydroquinase, 41000; DHS reductase, 26000; shikimate kinase, 13000; EPSP synthetase, 44000.
DISCUSSION

It has been suggested (Tresguerres, Ingledew & Cánovas, 1972) that in *Acinetobacter calco-aceiticus* the regulatory mechanism necessary for protecting aromatic biosynthetic intermediates in the presence of inducible enzymes catalysing their degradation may involve a multi-enzyme complex of the biosynthetic enzymes, similar to those found in fungi by Giles and co-workers. Our data do not support this suggestion. Under conditions in which the polyaromatic biosynthetic enzyme complex is readily demonstrable in fungi (Giles et al. 1967a; Ahmed & Giles, 1969), the *A. calco-aceiticus* enzymes, like those of other bacteria, including species with and without the degradative pathway (Berlyn & Giles, 1969), were separable. The molecular weight values were similar to those for other bacteria included in that study, resembling most closely those for *Pseudomonas aeruginosa* and *Streptomycetes coelicolor*. Although Tresguerres et al. (1972) observed a changed pattern of differential utilization of exogenous and DHQ-derived DHS in extracts prepared by sonic or lysozyme treatment, we observed similar sedimentation patterns of the biosynthetic enzymes extracted...
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by the two procedures. Under a variety of extraction conditions the enzymes were not aggregated in vitro.

Our observation of two distinct peaks on sucrose gradients of quinate-grown bacteria, together with the inducibility and heat-lability of the lighter peak, are consistent with the previous observations by Ingledew et al. (1971) of a quinate-induced isozyme distinguishable from synthetic dehydroquinase by thermal inactivation kinetics. Two dehydroquinase isozymes were described first in *Neurospora crassa* (Giles et al. 1967b) and subsequently in several fungi (Ahmed & Giles, 1969). The inducible dehydroquinase of *Acinetobacter calco-aceticus* sedimented at a position corresponding to a molecular weight of approximately 40000. The enzyme which catalyses the preceding step in the degradative pathway, quinate (shikimate) dehydrogenase was found by Tresguerres, de Torrontegui, Ingledew & Cánovas (1970b) to be associated with a particulate fraction. Although we did not study this enzyme, in a single assay we found the enzyme in the crude extract (of quinate-grown bacteria), but not in gradients after high-speed centrifugation. Thus the dehydroquinase and dehydrogenase of the degradative pathway were also separable in vitro. However, the association of dehydrogenase with a particulate fraction suggests the possibility of a specialized localization of this reaction sequence within the bacterium.

Several lines of evidence have confirmed the existence of mechanisms regulating the flow of intermediates through the two pathways in *Acinetobacter calco-aceticus*: biosynthesis of aromatic amino acids continues in the presence of high levels of the hydroaromatic degradative enzymes, which can be induced gratuitously by *p*-hydroxybenzoic acid or PCA or maintained constitutively in PCA-accumulating mutants (Cánovas, Wheelis & Stanier, 1968); the biosynthetic dehydroquinase does not catalyse the degradation of exogenously supplied quinate in mutants lacking inducible dehydroquinase (Ingledew et al. 1971); the rate of formation of PCA from exogenously supplied DHS is greater than the rate of the DHS reductase reaction, whereas with DHS derived from conversion of exogenous DHQ by the biosynthetic dehydroquinase, the DHS reductase rate exceeds the DHS dehydrase rate at low substrate concentrations, with bacteria disrupted by lysozyme-osmotic shock treatment (Tresguerres et al. 1972). The induction of PCA-dependent enzymes in PCA oxygenase mutants grown on succinate at 30 °C suggests some leakage from the biosynthetic pathway, leading to PCA accumulation at this temperature; at 20 °C this does not occur (Cánovas et al. 1968).

In the absence of stable multi-enzyme complexes as found in fungi, alternate means of regulation in bacteria must be considered. In *Acinetobacter calco-aceticus* end-product induction by PCA of the hydroaromatic enzymes, as found by Cánovas et al. (1968) and Ingledew et al. (1971), ensures that the biosynthetic intermediates do not induce enzymes which would catalyse their degradation, since only at very high levels of (exogenously supplied) quinate or shikimate in an uninduced wild-type bacterium will enough PCA be formed to induce the early degradative pathway enzymes. This mechanism and its occurrence in other pathways degrading important biosynthetic compounds is discussed by Cánovas et al. (1968). However, this method for restricting induction of the hydroaromatic enzymes is not universal among bacteria. The hydroaromatic pathway in *Pseudomonas putida* appears to be induced by DHS or its immediate precursors, rather than by PCA (Wheelis & Stanier, 1970). This is similar to the substrate-induction implicated by Rines (1969) for *Neurospora crassa*. In *Aerobacter aerogenes*, the mode of induction of this pathway has not been elucidated. The continued synthesis of aromatic amino acids in bacteria with fully induced degradative enzymes may rely on favourable differences in reaction velocities and binding affinities of the enzymes for the common intermediates – although under many conditions
DHS utilization appears to be an exception in this regard, and feedback inhibition of DHS dehydrase by PCA may be important (Tresguerres et al. 1972)—or it may involve a channeling mechanism not characterized by aggregation in vitro. It is possible that physical compartmentalization of the components of the two pathways may predominate, but be destroyed after disruption; indeed, stability of the known aromatic biosynthetic multi-enzyme complexes and activities of their individual components have been found to vary in different organisms (S. I. Ahmed and N. H. Giles, unpublished; Berlyn et al. 1970).

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REFERENCES


