The Essential Role of Diaminopimelate Dehydrogenase in the Biosynthesis of Lysine by Bacillus sphaericus

By P. J. White

Department of Microbiology, The University, Western Bank, Sheffield S10 2TN, U.K.

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Extracts of Bacillus sphaericus NCTC 9602 catalysed the formation of meso-diaminopimelate from aspartic β-semialdehyde plus pyruvate, or from dihydrodipicolinate, even though no activities of tetrahydrodipicolinate acetylase (or succinylase) nor N-acetyl-(or N-succinyl-)L-LL-diaminopimelate deacylase nor diaminopimelate epimerase were found. However, meso-diaminopimelate D-dehydrogenase was present, and had very high activity at pH 7.5 in the direction of synthesis of meso-diaminopimelate from tetrahydrodipicolinate. A lysine-requiring mutant of B. sphaericus lacked diaminopimelate dehydrogenase, and this enzyme reappeared in a revertant that grew without lysine. Other lysine-requiring auxotrophs were defective in dihydrodipicolinate synthase or dihydrodipicolinate reductase or diaminopimelate decarboxylase, but had diaminopimelate dehydrogenase. Diaminopimelate dehydrogenase is not important in the assimilation of ammonia. Mutants that lack this enzyme or else cannot make one of its substrates (tetrahydrodipicolinate) still grow rapidly in minimal medium (plus 0.7 mM-L-lysine) containing ammonium chloride (36 mM) as the only major source of nitrogen. The wild-type grew with L-glutamine, but not with glutamate or lysine as sole source of nitrogen.

INTRODUCTION

Bacteria that can make lysine do so by the ‘diaminopimelate pathway’, which is named after a characteristic intermediate, 2,6-diaminopimelic acid. This metabolic route was first elucidated in Escherichia coli (reviewed by Meister, 1965) in which succinylated intermediates are used at certain steps (Fig. 1). Later, Sundharadas & Gilvarg (1967) and Weinberger & Gilvarg (1970) showed that acetylated intermediates were employed in all the species (six) of the genus Bacillus that they examined; succinylated compounds were not metabolized.

In this paper another variation of the diaminopimelate pathway is reported. In Bacillus sphaericus, Δ1-tetrahydrodipicolinate is converted by a single enzymic step to meso-diaminopimelate (Fig. 1) while acylated intermediates and L-diaminopimelate are not used at all. The enzyme which catalyses this step (meso-diaminopimelate D-dehydrogenase) has been extensively studied (Misono & Soda, 1980). Misono et al. (1979) proposed the new pathway and suggested that it might play a secondary role in lysine formation by B. sphaericus, but these possibilities were not tested experimentally.

Bacillus sphaericus was one of the first bacteria to be assayed for diaminopimelate decarboxylase and diaminopimelate epimerase (Powell & Strange, 1957). The latter enzyme was not detected, though its absence did not seem remarkable because the function of the enzyme in lysine synthesis by other bacteria (Fig. 1) was not clear at the time, and the ability of B. sphaericus to make lysine had not been shown.

The activity of diaminopimelate epimerase in B. sphaericus was re-examined when this organism was found to grow with acetate as sole source of carbon (White & Lotay, 1980). The enzyme was absent, even though diaminopimelate decarboxylase was present with relatively high activity. The lack of the epimerase means that neither of the established variants of the diaminopimelate pathway can operate in B. sphaericus, because diaminopimelate decarboxylase...
Fig. 1. The pathway of lysine biosynthesis in *B. sphaericus*, compared with routes in other bacteria. The bold arrows (reactions 1 to 6) indicate the pathway in *B. sphaericus*. Acetylated compounds are used by *B. megaterium*, and *E. coli* uses succinylated intermediates (shown in brackets) in reactions 7 to 9. The enzymes are: 1, aspartokinase; 2, aspartic β-semialdehyde dehydrogenase; 3, dihydrodipicolinate synthase; 4, dihydrodipicolinate reductase; 5, *meso*-diaminopimelate N-dehydrogenase; 6, diaminopimelate decarboxylase; 7, tetrahydrodipicolinate acetylase (or succinylase); 8, *N*-acetyl-(or *N*-succinyl)-aminoketopimelate:glutamate aminotransferase; 9, *N*-acetyl-(or *N*-succinyl)-diaminopimelate deacylase; 10, diaminopimelate epimerase; 11, homoserine dehydrogenase; 12, dihydrodipicolinate dehydrogenase.
is entirely specific for *meso*-diaminopimelate. It therefore seemed possible that the proposed secondary route of lysine synthesis (Misono *et al*., 1979) was in fact the principal route in *B. sphaericus*. Further evidence to support this view is given below.

**METHODS**

*Organisms*. *Bacillus sphaericus* NCTC 9602 was maintained on slopes of nutrient agar, which were incubated at 30 °C and kept at 2 °C. *Bacillus megaterium* NCIB 7581 and *E. coli* W (ATCC 9637) were maintained similarly, but the slopes were incubated at 37 °C.

*Media*. The defined medium A3 was used for most experiments with *B. sphaericus*. This medium contains sodium acetate. 3H2O (5 g l⁻¹) and trisodium citrate. 2H2O (20 mg l⁻¹) as the only sources of carbon, and it was prepared as described by White & Lotay (1980). In a few experiments with *B. sphaericus* nutrient broth was used. *Bacillus megaterium* and *E. coli* were grown in medium A1 (White, 1972) with biotin and trisodium citrate added for *B. megaterium* (White, 1979).

*Growth of cultures*. Batch cultures of *B. sphaericus* were grown at 30 °C in liquid medium, and growth was assessed turbidimetrically as described by White & Lotay (1980). Cultures of *E. coli* and *B. megaterium* were grown similarly, but at 37 °C.

*Enzyme assays*. Bacterial extracts were made by passage of suspensions through a pressure cell (Milner *et al*., 1950) at 140 MPa and assayed for protein (by the Lowry method) as described by Chatterjee & White (1982). Procedures for assay of individual enzymes were as listed by Chatterjee & White (1982), with the following additional methods: dihydrodipicolinate synthase (EC 4.2.1.52) was sometimes assayed by the method of Yamakura *et al.* (1974); dihydrodipicolinate reductase (EC 1.3.1.26) was measured as described by Kimura (1975), with dihydrodipicolinate (about 2 μmol L-isomer per assay mixture) that had been made by non-enzymic reaction between pyruvate and aspartic β-semialdehyde in 1 M-KOH (see below); *meso*-diaminopimelate d-dehydrogenase was assayed in the direction of synthesis of diaminopimelate, with L-2-amino-6-ketopimelate, NH₄⁺ and NADPH as substrates, at pH 7.5 (Misono & Soda, 1980); N-succinyl-L-L-diaminopimelate desuccinylase (EC 3.5.1.18) was assayed as described by Kindler (1962), but on half-scale. Some variations of these general procedures for preparation of extracts and for enzymic assays are given in Results.

*Synthesis of diaminopimelate from aspartic β-semialdehyde plus pyruvate*. Formation of diaminopimelate by extracts of *B. sphaericus* took place in a system derived from that used to measure its synthesis with *E. coli* (Gilvarg, 1962). The following were mixed in a small test tube (100 × 10 mm) to a total volume of 0.9 ml (before adding extract): Tris/HCl buffer pH 7.5 (200 pmol); NADPH (0.1 pmol); glucose-6-phosphate dehydrogenase (10 μl); NH₄Cl (200 μmol); L-aspartic β-semialdehyde (2 μmol L-isomer); sodium pyruvate (10 μmol). The mixture was equilibrated at 30 °C for 5 min while N₂ was passed above the liquid surface (to minimize enzymic oxidation of NADPH with dissolved O₂). Extract (0.1 ml; about 1 to 2 mg protein) was added, and a sample (0.2 ml) was withdrawn at once from the mixture and put into 0.6 ml ethanol. Again N₂ was passed over the enzymic reaction mixture at 30 °C for 2 min and the tube was then capped and incubated for 1 h (from the time of adding the extract). A further sample of the mixture (0.2 ml) was finally added to ethanol (0.6 ml). Material precipitated by the ethanol was removed by centrifugation. Part of the supernatant liquid (0.4 ml) was evaporated to dryness at 40 °C under diminished pressure, then taken up in water (0.1 ml). Acetic acid (0.8 ml) and ninhydrin reagent (0.1 ml) were added, and this mixture was incubated at 37 °C for 90 min before reading the absorbance at 440 nm as a measure of diaminopimelate (Work, 1957); the ninhydrin reagent was 50 mg ninhydrin in 0.8 ml 0.6 M orthophosphoric acid plus 1.2 ml glacial acetic acid. Another portion of the supernatant liquid (0.1 ml) from the ethanol precipitation was spotted on to paper to test chromatographically for the presence of diaminopimelate (see below). Control mixtures were incubated without extract, and without individual substrates (see Results).

To measure synthesis of diaminopimelate from dihydrodipicolinate the procedure was similar, except that aspartic β-semialdehyde and pyruvate were omitted from the enzymic reaction mixture, and instead dihydrodipicolinate (prepared non-enzymically; about 2 μmol L-isomer) was added.

*Chromatography*. Descending chromatograms on Whatman no. 1 paper were developed with the solvent methanol/pyridine/11.6 M-HCl/water (80:10:2.5:17.5; by vol); Rhuland *et al.*., 1955). After drying the paper, spots were revealed with ninhydrin. Use of the above solvent, followed by ninhydrin, leads to a diaminopimelate spot with a very characteristic dull green colour, which changes in about 1 h to bright yellow. The solvent also separates the LL-isomer of diaminopimelate from the meso-isomer and DD-isomer; lysine migrates further than any isomer of diaminopimelate and gives a brown spot.

*Isolation of lysine-requiring mutants of B. sphaericus*. A series of 250 ml conical flasks containing medium A3 (25 ml per flask) was inoculated with *B. sphaericus* (wild-type). The size of the inoculum ranged from 2 ml to 0.1 ml of an aqueous suspension of organisms (10⁶ ml⁻¹) from a nutrient agar slope. The flasks were incubated overnight at 30 °C on a mechanical shaker. The following day a flask was chosen in which the bacteria were in the early exponential phase of growth (EEL colorimeter reading about 0.5; roughly 5 × 10⁶ organisms ml⁻¹), and 5 ml of the
culture was put into a test tube. To this was added $N$-methyl-$N'$-nitro-$N$-nitrosoguanidine (0.5 mg, as a freshly prepared aqueous solution) and the suspension was rocked at 30 °C for 10 min. A portion of this (0.2 ml) was then added to 50 ml nutrient broth which was incubated overnight with shaking, to allow phenotypic expression. The following day the organisms were starved by adding 0.05 ml of the broth culture to 5 ml medium A3, and rocked at 30 °C for 4 h. Benzyl penicillin (5 μg, as an aqueous solution) was added, and the incubation was continued for 1 h. The bacteria were centrifuged and resuspended in 50 ml medium A3 plus L-lysine (100 μg ml$^{-1}$) which was incubated overnight with shaking to allow outgrowth of lysine-requiring mutants. The next day the organisms from this culture were starved in medium A3 and treated with benzyl penicillin as before. Instead of centrifuging, the culture was diluted, plated on to nutrient agar and incubated overnight. Generally, 0.1 ml of a 100-fold dilution gave about 100 colonies. Prospective lysine-requiring auxotrophs were recognized by replica-plating on to medium A3 and A3 plus lysine. The mutants were further screened by tests on solid and liquid media. Some variations of the selection procedure are described in Results.

Chemicals. DL-Aspartic β-semialdehyde was made by ozonolysis of DL-allyl glycine (Black & Wright, 1955a). After ion-exchange chromatography of the reaction mixture, the fractions which contained aspartic β-semialdehyde were identified by the formation of a purple adduct with 2-aminobenzaldehyde when the fractions were incubated with pyruvate and a crude extract of E. coli. The following were mixed in a small test tube: sodium phosphate buffer pH 7.4 (100 μmol); sodium pyruvate (10 μmol); 2-aminobenzaldehyde (0.5 mg); column fraction (0.1 ml, neutralized with NaHCO$_3$); extract (about 1 mg protein); water to 1 ml. The mixture was incubated at 30 °C for 30 min before the absorbance at 540 nm was measured. Fractions that contained aspartic β-semialdehyde were pooled and the concentration of this compound (β-isomer) was found by the aspartic β-semialdehyde-dependent oxidation of NADH which is catalysed by homoserine dehydrogenase (Black & Wright, 1955b). A crude extract of E. coli was satisfactory as a source of this enzyme, provided that the reaction was done in an anaerobic cuvette (N$_2$ atmosphere), so that there was no oxidation of NADH by the extract until the substrate was added. Aspartic β-semialdehyde was kept at −60 °C as a solution in 4 M-HCl.

2-Aminobenzoic acid was obtained from Fluka. Some batches of this material were polymerized; only material that will dissolve in 20% (v/v) aqueous ethanol to give a solution containing 5 mg ml$^{-1}$ is suitable for the assays described here. To prepare dihydrodipicolinate, a solution (2 ml) of DL-aspartic β-semialdehyde in 4 M-HCl (containing 64 μmol L-aspartic β-semialdehyde) at 20 °C was added slowly, with mixing, to a solution (1.1 ml) of sodium pyruvate (135 μmol) in 11 M-KOH (Kimura, 1974). The mixture, which became yellow after about an hour, was left to stand overnight. The pH was lowered by cautious addition of 6 M-HCl, and excess acid removed by adding solid NaHCO$_3$ until effervescence stopped. This solution was used at once in assays for dihydrodipicolinate reductase and in measurements of synthesis of diaminopimelate by extracts; its concentration was assumed to be about 10 μmol l$^{-1}$ dihydrodipicolinate ml$^{-1}$. A new preparation was made freshly each day it was needed, usually on one-quarter of the scale described above.

L-Δ$^1$-Tetrahydropicolinate (L-2-amino-6-ketopimelate) was made enzymically from meso-diaminopimelate by the action of diaminopimelate dehydrogenase, in a modification of the procedure of Misono et al. (1979) which was adopted to economize on use of NADP. The following were mixed in a 500 ml conical flask: NaHCO$_3$/Na$_2$CO$_3$ buffer, pH 10.5 (8 mmol); NADP (16 mg); phenazine ethosulphate (8 mg); dichlorophenol-indophenol (oxidized; 50 mg); diaminopimelic acid [4 mmol; synthetic; 50% (w/w) meso-isomer]; water to 200 ml (final volume after adding enzyme). The solution was stirred at 37 °C and diaminopimelate dehydrogenase (10 units; 10 mg protein) was added. When the blue colour of the reaction mixture disappeared, more dichlorophenolindophenol (50 mg) was added as an aqueous solution (10 mg ml$^{-1}$). Additions were made at intervals over about 2 h until the reaction mixture was permanently blue and smelled of ammonia. The pH was brought to 1 by adding 6 M-HCl, and the precipitated material was removed by centrifuging. The supernatant liquid was applied to a column of Dowex 50, which was eluted as described by Misono et al. (1979). Fractions (10 ml) were tested (0.13 ml per assay) for ninhydrin-positive material by the method of Gilvarg (1958) with a heating time of 20 min. Fractions which were positive were then tested with 2-aminobenzaldehyde: each fraction (0.4 ml, neutralized with NaHCO$_3$) was mixed with 1 M-imidazole buffer, pH 7.4 (0.1 ml) and 0.5 ml 2-aminobenzaldehyde [1 mg ml$^{-1}$ in 20% (v/v) aqueous ethanol] then incubated at 30 °C for 1 h before measuring the absorbance at 440 nm. Fractions 10 to 20 (eluted by 1 M-HCl) were positive with 2-aminobenzaldehyde and the absorption spectrum (in the visible range) of the adduct was almost identical to that of the adduct formed by authentic Δ$^1$-tetrahydropicolinate (Farkas & Gilvarg, 1965). No other fractions gave a visible colour with 2-aminobenzaldehyde. Fractions 10 to 20 were also the only fractions which gave a colour with 2,4-dinitrophenylhydrazine: each fraction (0.5 ml) was mixed at 37 °C with 0.1 ml of a solution of 2,4-dinitrophenylhydrazine (1 mg ml$^{-1}$) in 2 M-HCl, and 0.4 ml water. The absorbance at 415 nm was measured after 10 min. The pooled fractions 10 to 20 were dried by evaporation under diminished pressure. The yield of crude 2-amino-6-ketopimelate was only 20 mg. The material was taken up in water (to give a concentration of 10 μmol ml$^{-1}$), and kept at −15 °C.
Diaminopimelate dehydrogenase was partly purified from *B. sphaericus* (grown in nutrient broth) as described by Chatterjee & White (1982). Glucose-6-phosphate dehydrogenase (from yeast) was from Boehringer.

Isomers of diaminopimelate were prepared as described by Saleh & White (1976). N-Succinyl-LL-diaminopimelate acid was prepared from LL-diaminopimelate (200 mg) as described by Gilvarg (1959).

**RESULTS**

**Assays for diaminopimelate epimerase and decarboxylase**

*Bacillus sphaericus* NCTC 9602 was grown in medium A and the washed bacteria were broken (see Methods). The supernatant liquid (extract) obtained after centrifugation of the debris at 20000 g for 20 min contained diaminopimelate decarboxylase [50 nmol CO₂ min⁻¹ (mg protein)⁻¹] which was measured manometrically (White, 1971). In the same extract no diaminopimelate epimerase could be detected manometrically (White et al., 1969). There was still no activity when broken organisms were assayed without preliminary centrifugation.

Neither broken organisms nor an extract could form meso-diaminopimelate (or lysine) from LL-diaminopimelate (shown by chromatography) in the system of Antia et al. (1957), even after incubation for 18 h at 30 °C. These experiments were done at pH 5 (sodium acetate buffer), pH 6, pH 7 (phosphate buffer), pH 8 (phosphate buffer or Tris/HCl buffer) and pH 9 (borate buffer), with and without an activator (1 nm-2,3-dimercaptopropan-1-ol). Bacteria dried with acetone (Antia et al., 1957) were also inactive in these tests. Finally, a lysozyme digest (of organisms harvested in the early part of the exponential phase of growth) was tested by manometric and chromatographic methods, but no diaminopimelate epimerase was found. In all these experiments, an extract of *B. megaterium* served as a control to show that diaminopimelate epimerase could be detected by the assay systems.

Extracts were tested in the spectrophotometric assay system for diaminopimelate dehydrogenase, with LL-diaminopimelate as substrate, at various pH values between 6.5 to 10.5, at all of which diaminopimelate dehydrogenase showed easily measurable activity with meso-diaminopimelate. In no case was an increase of absorbance seen at 340 nm, which implies that the LL-isomer could not be epimerized. Addition of meso-diaminopimelate or an extract of *B. megaterium* to the test systems led to a rapid rise of absorbance at 340 nm.

**Other enzymes of the pathway to diaminopimelate**

All the enzymes needed to form tetrahydrodipicolinate from aspartate plus pyruvate were present in an extract of *B. sphaericus* (Table 1). There was, however, no reaction between 2-amino-6-ketopimelate (tetrahydrodipicolinate) and acetyl-CoA or succinyl-CoA, and no deacylation of N-acetyl- or N-succinyl-LL-diaminopimelate in the presence of the extract. On the other hand, diaminopimelate dehydrogenase had very high activity in the extract, whether measured in the direction of synthesis of meso-diaminopimelate (at pH 7.5) or in the direction of oxidation of this compound (at pH 10.5).

**Synthesis of diaminopimelate from aspartic β-semialdehyde**

Incubation of an extract of *B. sphaericus* 9602 with aspartic β-semialdehyde plus pyruvate (under conditions described in Methods) led to the formation of diaminopimelate, which was assayed colorimetrically, and identified as the meso-isomer by chromatography. After 1 h at 30 °C about 0.2 µmol had been formed by 1 mg protein. No synthesis occurred when the above two substrates or NADPH were omitted. Formation of diaminopimelate from dihydrodipicolinate (in place of aspartic β-semialdehyde plus pyruvate) was shown by chromatography; omission of NH₄Cl diminished (by about half) the size and intensity of the spot on the final chromatogram due to meso-diaminopimelate. In no experiment was any synthesis of lysine detected by chromatography. This could have resulted from the absence of pyridoxal phosphate and the slightly alkaline incubation conditions, both of which would have limited the activity of diaminopimelate decarboxylase.
Table 1. Activities of enzymes of lysine biosynthesis in *B. sphaericus* NCTC 9602

Organisms were grown in medium A3 in the absence or presence of L-lysine (100 mg l\(^{-1}\)) and extracts were made and assayed for enzymes as described in Methods. All the assays were done at 30 °C and no attempt was made to alter the published procedures (in order to record maximal activities for *B. sphaericus*). Activities are in nmol min\(^{-1}\) (mg protein\(^{-1}\)) except for dihydrodipicolinate synthase, which was assayed by the method of Shedlarsky (1971) and the activity is given as units of enzyme (mg protein\(^{-1}\)); the unit represents about 0.5 nmol min\(^{-1}\).

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<td>25</td>
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<tr>
<td>(Homoserine dehydrogenase)</td>
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<td>Diaminopimelate dehydrogenase pH 10.5(\dagger)</td>
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<td>711</td>
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<tr>
<td>Diaminopimelate decarboxylase</td>
<td>77</td>
<td>57</td>
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</table>

ND, Not determined.

\(\ast\) 2-Amino-6-ketopimelate plus NH\(_4\)Cl as substrates.

\(\dagger\) meso-Diaminopimelate as substrate.

**Lysine-requiring mutants of *B. sphaericus***

Because diaminopimelate itself is not needed as a structural component of vegetative peptidoglycan in *B. sphaericus*, auxotrophs can be isolated that are blocked at various steps in the synthesis of diaminopimelate yet which grow in minimal medium supplemented only with lysine. The selection of a mutant that requires lysine and still has all the enzymes of the pathway, except for diaminopimelate dehydrogenase, would indicate that this latter enzyme does have an essential role in the biosynthesis of lysine.

Lysine-requiring auxotrophs of *B. sphaericus* were isolated (see Methods) and extracts were assayed for various enzymes (Table 2). Since none of the mutants required homoserine (or methionine plus threonine), it was unlikely that aspartokinase or aspartic β-semialdehyde dehydrogenase would be absent, and indeed these enzymes were present in every mutant that was tested.

One mutant (M1) had no diaminopimelate dehydrogenase, whether assayed in the direction of reduction of NADP\(^{+}\) (at pH 10.5) with meso-diaminopimelate as substrate, or in the opposite direction (at pH 7.5) with 2-amino-6-ketopimelate as substrate. All of the other enzymes needed to make tetrahydrodipicolinate were present, as was diaminopimelate decarboxylase. An extract of mutant M1 was unable to form diaminopimelate from dihydrodipicolinic acid. When liquid minimal medium A3 was inoculated with mutant M1, turbidity appeared after 3 d at 30 °C. On transfer to fresh minimal medium these revertants grew as quickly as the wild-type, and an extract of revertant organisms had considerable activity [125 nmol NADPH formed min\(^{-1}\) (mg protein\(^{-1}\))] of diaminopimelate dehydrogenase.

All of the other mutants were deficient at one of the other enzymic steps between aspartic β-semialdehyde and lysine (Table 2). Using substrates that had been made chemically proved very advantageous in the assay for dihydrodipicolinate reductase. After the incubation of aspartic β-semialdehyde overnight in 1 M-KOH (with pyruvate) very little of this compound remains (Kimura, 1975). In contrast, when dihydrodipicolinate is generated enzymically, the residual aspartic β-semialdehyde may be a substrate for homoserine dehydrogenase, so that assays for dihydrodipicolinate reductase give erroneously high results with a crude extract. Inactivation of homoserine dehydrogenase by heating briefly to 70 °C (Tamir, 1971) was unsatisfactory with extracts of *B. sphaericus*, almost all the protein was coagulated, and no activity of
Table 2. Activities of certain enzymes of lysine biosynthesis in lysine-requiring mutants of B. sphaericus

Mutants were successfully isolated in eight separate experiments; five other experiments gave no lysine-requiring mutants. The various mutants were grown in medium A3 plus lysine, and extracts were made and assayed for enzymes as described in Methods. The procedure of Shedlarsky (1971) was used for dihydrodipicolinate synthase; diaminopimelate decarboxylase was measured manometrically with pyridoxal phosphate added. Diaminopimelate dehydrogenase was assayed at pH 10.5, with meso-diaminopimelate as substrate.

<table>
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<tr>
<th>Experiment no.</th>
<th>Mutant(s) isolated in experiment</th>
<th>Aspartokinase</th>
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<th>Dihydrodipicolinate synthase</th>
<th>Dihydrodipicolinate reductase</th>
<th>Diaminopimelate dehydrogenase</th>
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+, Activity similar to that of wild-type (grown with lysine); (+), activity less than half that of wild-type; -, no activity found; ND, not determined.

* Outgrowth of mutants was from nutrient broth (rather than medium A3 plus lysine). Recognition of potential mutants was done on a mixture of L-amino acids, with and without lysine. No mutants were obtained which required lysine plus a second amino acid.

† Outgrowth and recognition of mutants was on medium A3 plus histidine, glutamate and lysine (see Results). No mutants were obtained which required glutamate and/or histidine in addition to lysine.

‡ Four other isolates with this phenotype from experiment 6.
§ Ten other isolates with this phenotype from experiment 7.
dihydrodipicolinate reductase was left in the supernatant liquid. Two mutants, MW2 and B2, seemed at first to lack none of the enzymes of the lysine pathway. In the assay of Shedlarsky (1971) mutant MW2 had slight activity of dihydrodipicolinate synthase, but the extract was inactive in the more precise assay system of Yakamura et al. (1974). The extract made diaminopimelate from dihydrodipicolinate, but there was no synthesis from aspartic β-semialdehyde (plus pyruvate). Mutant B2 could form diaminopimelate from aspartic β-semialdehyde just as well as the wild-type, and seemed to have all the enzymes necessary to make lysine. However, in this mutant, diaminopimelate decarboxylase proved to be abnormal. When measured with pyridoxal phosphate in the assay system the activity was 27 nmol min⁻¹ (mg protein)⁻¹; with pyridoxal or with no cofactor the activity was zero. After growth in medium A3 plus lysine the wild-type activities were [nmol min⁻¹ (mg protein)⁻¹]: 57 with pyridoxal phosphate; 44 with pyridoxal; and 31 with no cofactor (cf. Meadow & Work, 1958a).

Several of the mutants that were blocked before diaminopimelate (CW4, CW5, CW6, MW1, MW2, MW3 and MW4) were tested for growth in liquid medium A3 plus diaminopimelate (meso-isomer, or LL-isomer or synthetic) at various concentrations up to 1 mg ml⁻¹. None showed appreciable growth within 4 d. 2-Amino-6-ketopimelate (1 mM) failed to support growth of mutants M1, P4, K2, A1, B2, B3, B4 on solid medium A3.

Nitrogen requirements of B. sphaericus

When each lysine-requiring mutant was obtained it was first screened for activity of diaminopimelate dehydrogenase at pH 10-5 with meso-diaminopimelate as substrate. Some 20 auxotrophs were isolated before one was found which lacked this enzyme. It seemed possible that the enzyme might be needed for assimilation of ammonia (see later) so that a mutant lacking the dehydrogenase might require some organic source of nitrogen, and might fail to grow in medium A3 plus lysine.

Neither L-lysine nor L-glutamate allowed growth of the wild-type in medium A3 from which ammonium ions were omitted (White & Lotay, 1980). Nitrate and urea were also ineffective, but a mixture of six L-amino acids (aspartate, glutamate, proline, phenylalanine, arginine and histidine; each at 200 mg l⁻¹) did support growth. Progressive omissions showed that L-glutamate plus L-histidine (400 and 200 mg l⁻¹, respectively) could replace ammonium ions, and some mutants were then isolated from medium A3 supplemented with these two amino acids and lysine (Table 2).

Subsequently, L-glutamine (or L-asparagine) proved to be adequate as a single source of nitrogen for B. sphaericus 9602. The lowest concentration of L-glutamine that did not restrict growth was 1 g l⁻¹, which represents 190 mg nitrogen l⁻¹. At least 11 mM-NH₄Cl (154 mg nitrogen l⁻¹) was needed to support maximal growth when this salt was sole source of nitrogen. If nitrogen makes up 15% of the bacterial dry weight, then growth to the observed maximal density (1 g dry wt l⁻¹) should require a supply of 150 mg nitrogen l⁻¹.

In contrast, the lowest concentration of L-lysine that supported full growth of all the mutants tested was 50 mg l⁻¹, which represents only 10 mg nitrogen l⁻¹. If protein makes up 50% of the bacterial dry weight and peptidoglycan represents a further 10%, then 1 g dry wt of organisms will contain about 50 mg of lysine residues. It seems very probable, therefore, that lysine in the medium is a source of this amino acid for biosynthesis, but that lysine is not a general source of nitrogen.

Non-involvement of diaminopimelate dehydrogenase in the assimilation of ammonia by B. sphaericus

White & Lotay (1980) found that glutamate dehydrogenase was present in B. sphaericus 9602, while glutamate synthase was absent. Diaminopimelate dehydrogenase catalysed the reductive amination of a carbon skeleton (2-amino-6-ketopimelate) in forming diaminopimelate, and transamination between diaminopimelate and some α-keto acids has been claimed to occur in B. sphaericus (Meadow & Work, 1958b). Hence, diaminopimelate dehydrogenase might play a part in the assimilation of ammonia by these bacteria.

The properties of the lysine-requiring mutants make this hypothesis unlikely. Mutant M1
Lysine biosynthesis in Bacillus sphaericus

Lacks diaminopimelate dehydrogenase but it grows in medium A3 supplemented only with lysine. All of the mutants blocked at dihydrodipicolinate synthase or dihydrodipicolinate reductase must be unable to form 2-amino-6-ketopimelate, yet all grow in medium A3 plus lysine, in some cases as quickly as the wild-type.

Attempts to detect transamination between synthetic diaminopimelic acid and 2-oxoglutarate, catalysed by an extract of B. sphaericus 9602 in the system of Meadow & Work (1958b), have been completely unsuccessful.

After dialysis, partially purified diaminopimelate dehydrogenase catalysed an anaerobic oxidation of NADPH at pH 7-5 that was dependent on the presence of both 2-amino-6-ketopimelate and NH₄Cl. There was no reaction when L-glutamine (up to 5 mM) was used in place of NH₄Cl. The enzyme, therefore, does not seem able to function, at low concentrations of glutamine, as a counterpart of glutamate synthase (Meers et al., 1970).

DISCUSSION

Bacillus sphaericus NCTC 9602 grows with acetate as sole source of carbon (White & Lotay, 1980), and so must make the lysine which occurs in its vegetative peptidoglycan (Hungerer & Tipper, 1969) and in proteins. Aspartokinase, aspartic β-semialdehyde dehydrogenase, dihydrodipicolinate synthase, dihydrodipicolinate reductase and diaminopimelate decarboxylase are all readily detected in an extract of B. sphaericus. Furthermore, the extract can make meso-diaminopimelate from its precursors aspartic β-semialdehyde plus pyruvate, or from dihydrodipicolinate. However, the absence of diaminopimelate epimerase from B. sphaericus implies that neither of the established routes from tetrahydrodipicolinate to meso-diaminopimelate can operate in this organism. No enzyme was found in B. sphaericus that could catalyse a reaction between acetyl- or succinyl-CoA and tetrahydrodipicolinate, and crude extracts of these bacteria did not deacetylate N-acetyl- or N-succinyl-LL-diaminopimelate. This latter observation makes it unlikely that epimerization of an acylated LL-diaminopimelate occurs (to form a derivative of meso-diaminopimelate), followed by the action of a novel deacylase, to yield meso-diaminopimelate.

The enzyme meso-diaminopimelate D-dehydrogenase reductively aminates tetrahydrodipicolinate at pH 7-5 to produce meso-diaminopimelate in a single step, and the activity of this enzyme in B. sphaericus is much higher than any of the other enzymic activities in the lysine pathway. Bacillus sphaericus cannot readily take up diaminopimelate (mutants blocked in the synthesis of lysine before diaminopimelate are unable to grow with this amino acid in minimal medium) and so it is unlikely that diaminopimelate dehydrogenase is formed for the purpose of oxidizing diaminopimelate, even though such a reaction is readily catalysed by extracts (Antia & Work, 1961). A function for this enzyme in the assimilation of ammonia is also doubtful (see Results). A role in the synthesis of meso-diaminopimelate appears much more probable.

Loss of diaminopimelate dehydrogenase by mutation caused a requirement for lysine, and the enzyme reappeared in a revertant that grew in minimal medium. All other lysine-requiring mutants (that have diaminopimelate dehydrogenase) are defective in one of the three enzymic steps in the suggested pathway (Fig. 1) between aspartic β-semialdehyde plus pyruvate, and lysine. A mutant lacking aspartokinase or aspartic semialdehyde dehydrogenase would fail to make homoserine, and so could not be selected by screening for the single requirement for lysine.

Lysine-requiring mutants of B. sphaericus have already been obtained by other workers (Imae & Strominger, 1976; Linnett & Tipper, 1976). These mutants were divided into three groups, those blocked at or before dihydrodipicolinate synthase, those blocked between dihydrodipicolinate and diaminopimelate, and those lacking diaminopimelate decarboxylase. This division was based on the properties of the spores formed by the mutants, and the individual enzymes were not examined.

Synthesis of meso-diaminopimelate by a single enzymic step from tetrahydrodipicolinate seems to offer advantages over the established pathways. No acetyl- or succinyl-CoA is needed, and four enzymes (Fig. 1, numbers 7 to 10) are not required. Perhaps the operation of four enzymes allows a more refined control of metabolic flow, but very little has been reported on the regulation of this part of the diaminopimelate pathway in any bacteria.
In spite of its apparent advantages, the synthesis of meso-diaminopimelate via diaminopimelate dehydrogenase does not seem, on the basis of rather limited surveys, to be of wide occurrence. Misono et al. (1979) assayed 30 species of bacteria and found only four with appreciable activity of diaminopimelate dehydrogenase, while all the species examined (12) by Weinberger & Gilvarg (1970) deacylated either N-succinyl- or N-acetyl-LL-diaminopimelate.

In certain eukaryotic organisms (euglenoids, higher fungi and some lower fungi) lysine is made by an entirely different route, in which 2-aminoadipic acid is a characteristic intermediate (see review by Meister, 1965). Vogel et al. (1970) have considered how this pathway and the diaminopimelate pathway might have evolved separately. If, as Vogel et al. (1970) proposed, the diaminopimelate pathway was evolved in ancient times by ancestral forms of modern bacteria and plants, then natural selection might have been expected to favour the most efficient of the three variants of this pathway, and to have eliminated the other two.

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