The second aconitase (AcnB) of Escherichia coli

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The second aconitase (AcnB) of Escherichia coli was partially purified from an acnA::kanR mutant lacking AcnA, and the corresponding polypeptide identified by activity staining and weak cross-reactivity with AcnA antiserum. The acnB gene was located at 2.85 min (131.6 kb) in a region of the chromosome previously assigned to two unidentified ORFs. Aconitase specific activities were amplified up to fivefold by infection with λacnB phages from the Kohara E. coli gene library, and up to 120-fold (50% of soluble protein) by inducing transformants containing a plasmid (pGS783) in which the acnB coding region is expressed from a regulated T7 promoter. The AcnB protein was purified to >98% homogeneity from a genetically enriched source (JRG3171) and shown to be a monomeric protein of M, 100000 (SDS-PAGE) and 105000 (gel filtration analysis) compared with M, 93500 predicted from the nucleotide sequence. The sequence identity between AcnA and AcnB is only 17% and the domain organization of AcnA and related proteins (1-2-3-linker-4) is rearranged in AcnB (4-1-2-3).

Keywords: aconitase, Escherichia coli, citric acid cycle, iron–sulphur proteins, protein domains

INTRODUCTION

Aconitase (EC 4.2.1.3) catalyses the reversible isomerization of citrate and isocitrate via cis-aconitate in the citric acid and glyoxylate cycles. Aconitases are monomeric enzymes having a single [4Fe-4S] centre that is essential, but has no redox function in catalysis. Recent studies have shown that Escherichia coli contains at least two aconitases, AcnA and AcnB (Gruer & Guest, 1994). The gene (acnA) encoding the first enzyme to be purified (AcnA, M, 97500) was located at 28 min (1350 kb) on the physical map of the E. coli chromosome, and AcnA specific activity could be amplified up to 200-fold in phagemid-containing strains.

The AcnA protein was purified to homogeneity from both wild-type and genetically amplified sources (Prodromou et al., 1991; Bennett et al., 1995). Studies with acnA–lacZ fusions have shown that expression of the acnA gene is subject to cAMP receptor protein (CRP)-mediated catabolite activation, ArcA- and FNR-mediated anaerobic repression, iron and Fur-dependent activation, superoxide and SoxRS-dependent activation and RpoS-mediated stationary-phase activation (Gruer & Guest, 1994; M. J. Gruer & J. R. Guest, unpublished observations). The primary structure of AcnA deduced from the nucleotide sequence of the acnA gene showed that the E. coli enzyme is structurally related to the porcine mitochondrial aconitase (Prodromou et al., 1992; Robbins & Stout, 1989). Some 19 of the 20 active-site residues, including three cysteine residues involved in ligand binding to the [4Fe-4S] centre, are conserved and the overall sequence identity is 29%. AcnA is also 20–21% identical to the bacterial and fungal isopropylmalate isomerases (IPMIs) which catalyse the reversible isomerization of α-isopropylmalate and β-isopropylmalate via dimethylcitraconate in the leucine biosynthetic pathway (Prodromou et al., 1992). The fungal IPMIs resemble AcnA and other aconitases in being monomeric and in containing four colinear structural domains with a long interdomain linker extending between domains 3 and 4. In contrast, bacterial IPMIs are heterodimers expressed from adjacent genes (leuCD) that encode products corresponding to the three N-terminal domains of AcnA and an unethered C-terminal domain, but no linker (Rosenthal & Calvo, 1990; Prodromou et al., 1992). The AcnA protein is even more closely related (53% identity) to the mammalian iron regulatory proteins (IRPs) or iron regulatory factors (IRFs). Depending on the availability of iron, these proteins exist either as apo-proteins or

Abbreviations: CRP, cAMP receptor protein; IPMI, isopropylmalate isomerase; IRP, iron-regulatory protein.
[Fe-4S]-containing holo-proteins, each form having a distinct and mutually exclusive function. During iron-depletion, IRP adopts the enzymically inactive apo-protein conformation, which regulates translation by binding to the iron-responsive elements (IREs) of relevant transcripts, whereas during iron-sufficiency IRP becomes the [Fe-4S]-containing cytoplasmic aconitase which no longer binds IRE. It is postulated that IRP may sense iron availability through the continuous assembly and disassembly of the [Fe-4S] cluster, and that assembly of the [Fe-4S] cluster induces a conformational change such that the IRE-binding site of IRP becomes inaccessible to RNA (Klausner & Rouault, 1993; Gray et al., 1993; Emery-Goodman et al., 1994; Hirling et al., 1994). Recently, a second [Fe-4S]-containing IRP (IRP₂) has been identified but in this case the holo-protein lacks aconitase activity (Henderson et al., 1993; Guo et al., 1994).

In *E. coli*, a second aconitase (AcnB) was detected as the residual activity of a mutant (JRG2789) in which the chromosomal acnA gene had been disrupted by inserting a *kan<sup>R</sup>* cassette (Gruer & Guest, 1994). This enzyme resembled other iron–sulphur containing aconitases in being unstable and reactivatable by ferrous ions under reducing conditions. The present paper describes the partial purification of AcnB from the acnA::*kan<sup>R</sup>* mutant, the identification of the *acnB* gene, and the purification of AcnB from amplified sources to near homogeneity. AcnB is related to AcnA (17% sequence identity) and shows an interesting domain rearrangement (4-linker-1-2-3) relative to AcnA and other aconitases (1-2-3-linker-4).

**METHODS**

**Bacterial strains, plasmids, phagemids and bacteriophages.** The routine transformation host for all phagemids was *E. coli* DH5α (Hanahan, 1983); derivatives of *E. coli* DH1 (Sambrook et al., 1989) containing two plasmids (pGS799 and pGS801, see below) were used in specific growth tests, DH1(pGS801) being designated JRG3171; JRG3099 is a transformant of *E. coli* BL21/DE3 containing the expression plasmid, pGS783 (see below). The *acnA::*kan<sup>R</sup>* mutant (JRG2789) is a derivative of *E. coli* W3110 in which a 1.04 kb segment of the chromosomal *acnA* coding region is replaced by a 1.3 kb *kan<sup>R</sup>* cassette (Gruer & Guest 1994). Phages J4D12, J4E11 and J4C11 (112–114 in the mini-set of Kohara et al., 1987) were used in infection experiments and to generate the pUC18-derivatives pGS799 and pGS801 (Fig. 1). The *acnB* expression plasmid (pGS783) was constructed by PCR-amplification of the *acnB* coding region from JRG2789 chromosomal DNA with the simultaneous conversion of the GTG start codon to ATG and the incorporation of flanking *BamHI-NdeI* and *HindIII-PstI* sites, with the aid of two primers, S328, GCTG20261, and S329, TTGGATCCATAACGCGCTCAGTTCTTCACTCATTACAAGCGCGGGCAGCAG (numbered according to Fujita et al., 1994, with superscripted mismatches). The PCR reactions contained 200 ng DNA, 100 pmol of each primer and 2 U pfu DNA polymerase. The major amplified product (2.9 kb) was purified and cloned as a *BamHI-PstI* fragment into pUC119. Transformants were screened by sequencing single-stranded phagemid DNA from the universal priming site using the dideoxy chain-termination method. The 2.9 kb *acnB* coding region was then subcloned downstream of the IPTG-inducible T7 promoter in pET21a (Novagen), as an *NdeI-HindIII* fragment from a representative phagemid (pGS782), to generate the expression plasmid pGS783.

**Microbiological methods.** The routine complex medium was single-strength L broth (unless stated otherwise) and a rich maltate medium was used for specific purposes (Prodromou et al., 1991). A citrate-free minimal medium (Cole & Guest, 1980) was used with glucose (11 mM) as carbon source and supplements of thiamine hydrochloride (5 μg ml<sup>–1</sup>) and iron citrate (16 μM) or citric acid (0.2%, w/v). Cultures were grown aerobically at 37 °C, with ampicillin (100 or 200 μg ml<sup>–1</sup>) or kanamycin (20 μg ml<sup>–1</sup>) when necessary. For enzyme assays with small-scale cultures (1.5 ml), the bacteria were resuspended in 600 μl Tris/citrate buffer (20 mM, pH 8.0) at 0 °C and disrupted by two ultrasonic treatments (10 s at 10 MHz separated by a 30 s interval) in two 300 μl aliquots, followed by the removal of cell debris by centrifuging (10000 g for 5 min). To stabilize aconitase activity, dithiothreitol (5 mM), (NH₄)₂Fe(SO₄)₂ (1 mM) and Tris/HCl (50 mM, pH 8.0) were added to one aliquot prior to disruption and immediate assay, the other aliquot being used for protein estimation.

**Analysis of acnB expression in *λ*-infected bacteria and phagemid transformants.** Aerobic cultures of *E. coli* JRG2789 were grown to OD<sub>600</sub> 0.4 in 20 ml L broth containing 0.2% maltose and 10 mM MgSO<sub>4</sub>, and phages (m.o.i. of 3) adsorbed for 15 min at 37 °C without shaking. Shaking was then restarted (time zero) and samples (1.5 ml) were pelleted (15000 g for 2 min) and stored at −20 °C prior to analysis. Transformants of *E. coli* DH1 and BL21/DE3 were grown with shaking at 37 °C in 50 ml L broth containing ampicillin (200 μg ml<sup>–1</sup>) and sampled as described above. Expression of *acnB* from pGS783 was induced with IPTG (40 μg ml<sup>–1</sup>) 2 h before the cultures were harvested.

**Aconitase assay and reactivation.** Aconitase was assayed spectrophotometrically at 20 °C by following the formation of cis-aconitate from isocitrate at 240 nm, using an absorption coefficient of 3.6 mM<sup>–1</sup> cm<sup>–1</sup> (Kennedy et al., 1983). One unit (U) corresponds to 1 μmol cis-aconitate formed min<sup>–1</sup>. Protein concentrations were determined by the Bio-Rad micro-assay procedure. Where stated, aconitase was reactivated by incubating samples with dithiothreitol (5 mM), (NH₄)₂Fe(SO₄)₂ (1 mM) and Tris/HCl, (50 mM, pH 8.0) for 20 min at 20 °C. Staining for aconitase activity in non-denaturing polyacrylamide gels was performed by coupling the conversion of cis-aconitate to isocitrate to a colorimetric test for isocitrate dehydrogenase (Prodromou et al., 1991).

**Purification of AcnB from an unamplified source, JRG2789.** AcnB was purified from JRG2789 (acnA::*kan<sup>R</sup>*) grown with vigorous aeration at 37 °C for 7 h in five 2 l flasks, each containing 400 ml double-strength L broth inoculated to give a starting OD<sub>600</sub> 0.1. The bacteria were harvested (4000 g for 30 min) and resuspended (0.5 g wet wt ml<sup>–1</sup>) in MES/Mg buffer (10 mM MES/KOH, pH 6.0, with 5 mM MgCl<sub>2</sub>) before disruption by three passes through a French pressure cell (15000 p.s.i., 103.5 kPa). Extracts were clarified by centrifuging (25000 g for 30 min and 18000 g for 2 h). All steps were performed at 4 °C. The cell extracts were fractionated with (NH₄)₂SO₄ at 45–65% saturation, the pellets (15000 g for 30 min) being solubilized in 10 ml Tris/citrate buffer (20 mM Tris/citrate, pH 8.0) containing 0.25 M (NH₄)₂SO₄. This was loaded on a Phenyl Sepharose CL-4B column (5 × 10 cm) equilibrated with Tris/citrate buffer containing 0.25 M (NH₄)₂SO₄, washed with 200 ml of the same buffer, and eluted with a 600 ml linear gradient of 0.25–0.00 M (NH₄)₂SO₄ in
Tris/citrate buffer followed by buffer alone, at a flow rate of 5 ml min⁻¹. Fractions (15 ml) containing aconitase activity eluted 440 ml after applying the gradient. These fractions were pooled and loaded on a Procion Green HE4BD dye affinity column (2 x 30 cm) equilibrated with MES/Mg buffer, washed with 50 ml MES/Mg buffer, and eluted with an 800 ml linear gradient of 0–100 mM citrate in MES/Mg buffer at a flow rate of 2 ml min⁻¹. Aconitase activity emerged at a citrate concentration of 35 mM.

**Purification of AcnB from a genetically amplified source, JRG3171.** Inocula developed from a freshly prepared pGS801 transformant of DH1 (JRG3171) were used to grow double-strength L broth cultures (5 x 400 ml) with iron citrate (50 µM) and ampicillin (200 µg ml⁻¹) for 15 h from a starting OD₆₅₀ 0.05, prior to harvesting and extraction, as above. The pellets from (NH₄)₂SO₄ fractionation (45–65 % saturation) were combined and manipulated by standard procedures (Sambrook et al., 1989). The mixed oligonucleotides used in attempts to PCR-amplify and clone acnB sequences were either blunt-ended using DNA polymerase (Klenow fragment) or digested with Sau3A for cloning into HincII- or BamHI-digested pUC118, respectively. Artificially created sites are shown in brackets.

**Western blotting.** Samples fractionated by SDS-PAGE were transferred to nitro-cellulose using a Bio-Rad Transblot Electrophoretic Transfer Cell and the blots were immuno-stained with biotinylated anti-AcnA serum as the primary antiserum, biotinylated anti-galE serum, and peroxidase complex, according to the method of Prodromou et al. (1991). Biotinylated size markers were created sites are shown in brackets.

**DNA manipulation and related methods.** DNA was prepared and manipulated by standard procedures (Sambrook et al., 1989). The mixed oligonucleotides used in attempts to PCR-amplify and clone acnB sequences were: S325, ATG(TC)TIGA(GA)-GA(GA)TA(TC)(AC)G(GTGC)AA(AG); S326, PSFGG(GA)-TT(AGTC)(GC)(TA)(AGTC)G(TA)(GA)TT(AGTC)G(TA)(AGCA)(AGTC)(GC)(AT)⁵⁰⁵⁵; and S327, ¹⁷⁴⁶ CC(AGTC)-GG(AGTC)(AT)(CG)(ATGC)G(GTGC)(TA)(CT)G(TG)(ATGC)CA(TC)CA(AG)(AG)²⁵⁸¹ (coordinates refer to the ainA gene; Prodromou et al., 1992). Amplified products were either blunt-ended using DNA polymerase (Klenow fragment) or digested with Saa3A for cloning into HindIII- or BamHI-digested pUC118, respectively. DNA-directed tran-
scription–translation was performed with an *E. coli* S30 extract (Promega) and circular dsDNA. Reaction mixtures contained (in 50 µl): DNA, 1·5 µg; [35S]-L-methionine, 15 µCi (555 kBq, 12 pmol); amino acid mixture minus methionine, 0·1 mM; and S30 extract plus premix, according to manufacturer’s instructions. Radiolabelled polypeptides were fractionated by SDS-PAGE and detected autoradiographically.

Materials. Isocitrlic acid, tetrathionate tetratolium, phenazine methosulphate, isocitrate dehydrogenase, biotinylated molecular mass markers and molecular mass markers for gel filtration were obtained from Sigma. Alkaline phosphatase, NADP and (Promega) and circular dsDNA. Reaction mixtures contained transcription–translation was performed with an *E. coli* S30 extract (Promega) and circular dsDNA. Reaction mixtures contained (in 50 µl): DNA, 1·5 µg; [35S]-L-methionine, 15 µCi (555 kBq, 12 pmol); amino acid mixture minus methionine, 0·1 mM; and S30 extract plus premix, according to manufacturer’s instructions. Radiolabelled polypeptides were fractionated by SDS-PAGE and detected autoradiographically.

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RESULTS

Partial purification and characterization of AcnB from *E. coli* JRG2789 (acnA::kanR)

The second aconitase (AcnB) of *E. coli* was purified from a genetically engineered *acnA::kanR* mutant (JRG2789) which lacks AcnA (Gruer & Guest, 1994). Previous studies had indicated that AcnB is not an abundant enzyme, so different growth media were tested in an attempt to find an enriched source for use as starting material. The maximum specific activities observed throughout late-exponential and early-stationary phases were [U (mg protein)⁻¹ in ultrasonic extracts]: L broth (0·39); double-strength L broth (0·32); rich malate (0·31); glucose minimal (0·10); glucose minimal plus iron (0·18); and glucose minimal plus citrate (0·14). Early-stationary-phase double-strength L broth cultures were chosen to take advantage of the higher growth yield and the relatively high specific activity.

A typical AcnB purification is summarized in Table 1 and an SDS-PAGE fractionation of samples taken at each stage is shown in Fig. 2(a). Major activity losses occurred during the chromatographic steps, the overall yield (0·3%) and purification factor (1·6-fold) being particularly low. However, it is likely that the low specific activities are partly due to irreversible inactivation rather than loss of AcnB protein. Oxidative degradation of the iron–sulphur cluster during purification probably generates an inactive but co-purifying apo-protein which lowers the specific activity of the final product.

Although partially purified AcnB contained several polypeptide components, only one (*M*₉ 100000) was continuously enriched and had a fractionation pattern that correlated with the activity profile in the final stages of purification (Fig. 2a). The same polypeptide was one of three that cross-reacted, albeit weakly, with anti-AcnA serum in Western blotting. Activity staining of partially purified samples of the enzyme after fractionation in non-denaturing polyacrylamide gel (Fig. 2b, lane 5) further showed that the active band contains a polypeptide of *M*₉ 97 500. In direct comparisons, the mobility of the putative AcnB polypeptide was slightly less than that of the purified AcnA protein (*M*₉ 97 5000). A native *M*₉ of 105 000 (+ 8000) was determined for the AcnB protein by monitoring the

Table 1. Purification of aconitase B from *E. coli*

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Total activity units (reactivation factor)</th>
<th>Total protein (mg)</th>
<th>Specific activity [U (mg protein)⁻¹]</th>
<th>Yield (%)</th>
<th>Purification factor</th>
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<tbody>
<tr>
<td>JRG2789</td>
<td>240 (1·0)</td>
<td>900</td>
<td>0·26</td>
<td>100</td>
<td>1·0</td>
</tr>
<tr>
<td>(NH₄)₂SO₄ Ppn</td>
<td>160 (1·6)</td>
<td>325</td>
<td>0·49</td>
<td>67</td>
<td>1·8</td>
</tr>
<tr>
<td>Phenyl Sepharose CL-4B</td>
<td>4·6 (1·0)</td>
<td>19</td>
<td>0·24</td>
<td>1·9</td>
<td>0·9</td>
</tr>
<tr>
<td>Procion Green HE4BD</td>
<td>0·64 (1·0)</td>
<td>1·5</td>
<td>0·43</td>
<td>0·3</td>
<td>1·6</td>
</tr>
<tr>
<td>JRG3171/DH1(pGS801)</td>
<td>4300 (1·0)</td>
<td>700</td>
<td>6·1</td>
<td>100</td>
<td>1·0</td>
</tr>
<tr>
<td>(NH₄)₂SO₄ Ppn</td>
<td>2400 (1·0)</td>
<td>490</td>
<td>4·9</td>
<td>56</td>
<td>0·8</td>
</tr>
<tr>
<td>Procion Green HE4BD</td>
<td>1330 (83)</td>
<td>60</td>
<td>22·2</td>
<td>30</td>
<td>3·6</td>
</tr>
<tr>
<td>Procion Blue HERD</td>
<td>350 (21)</td>
<td>9·2</td>
<td>38·0</td>
<td>8</td>
<td>6·2</td>
</tr>
</tbody>
</table>
AcnB, the second aconitase of *E. coli*

116 - 77 - 66 - 43 - 30 - 17

**Fig. 2.** Purification and activity staining of AcnB. (a) Purification of AcnB from an unenriched source, JRG2789. Coomassie-Blue-stained SDS-PAGE (15% gel) showing samples of active fractions at different stages. Lanes: 1, cell-free extract; 2, ammonium sulphate precipitate; 3, Phenyl Sepharose; 4, Procion Green HE48D. (b) Activity staining of AcnB. Lanes: 5, activity-stained gel containing the lane 4 sample of partially purified AcnB fractionated in non-denaturing PAGE (10% gel); lane 6, SDS-PAGE (15% gel)-fractionated and Coomassie-Blue-stained gel containing the enzymically active band (arrowed) excised from lane 5. (c) Purification of AcnB from a genetically enriched source JRG3171, DH1(pGS801). Coomassie-Blue-stained SDS-PAGE (15% gel) with active fractions from cell-free extract (lane 7), ammonium sulphate precipitate (lane 8), Procion Green HE48D (lane 9), Procion Blue HERD (lane 10).

elution of enzyme activity in quantitative gel filtration studies.

Several preparations of AcnB purified by Procion Green chromatography (Table 1; Fig. 2a) were fractionated by SDS-PAGE to determine the N-terminal amino acid sequence of the polypeptide tentatively identified as AcnB by activity staining and by Western Blotting. In all cases the *M*<sub>r</sub> 100000 polypeptide had the following N-terminal sequence: MLEEYRKxAE, where *x* denotes an unidentified residue. Two other immunopositive polypeptides of *M*<sub>r</sub> 46000 and 35000 in partially purified AcnB had N-terminal sequences that were identical to isocitrate dehydrogenase over 24 residues, indicating that they correspond to isocitrate dehydrogenase and a C-terminally degraded form, respectively. Their appearance suggests that the protein originally used to raise the AcnA antiserum was contaminated with isocitrate dehydrogenase. Indeed it would seem that isocitrate dehydrogenase co-purifies to some extent with both AcnA and AcnB.

**Identification of the acnB gene**

Preliminary attempts to clone the *acnB* gene by a PCR amplification strategy using DNA from the *acnA::kan<sup>R</sup>* mutant (JRG2789) and mixed oligonucleotide primers encoding the N-terminus of AcnB (S325) and two highly conserved internal regions of the aconitase-IPMI family that are deleted in JRG2789 (S326 and S327), failed. None of the amplified products resembled *acnA* sequences. With hindsight it is apparent that there is insufficient similarity between AcnA and AcnB (even in the most conserved regions) for this approach to have been successful (e.g. there are identical residues at only three out of eight and five out of nine positions in the chosen internal regions).

Subsequent database searches revealed that the N-terminal sequence of the putative AcnB protein is encoded by an internal segment of a 408 bp unidentified ORF (*yacl* in the Swiss-Prot database; 38 in Fujita *et al.*, 1994, GenBank D26562). This short segment of DNA was located immediately upstream of a 2319 bp ORF (*yacJ* in the Swiss-Prot database; 40 in Fujita *et al.*, 1994) part of which had been recognized as encoding a product that is 23% identical to a fungal isopropylmalate dehydratase (Fujita *et al.*, 1994). It was realized that if the two ORFs (*yacl* and *yacJ*) were fused by a frame-shifting single base-pair insertion in the upstream sequence, the resultant reading frame would encode a polypeptide having the same N-terminal sequence and approximately the same *M*<sub>r</sub> (93500) as AcnB (Berlyn *et al.*, 1996). The putative *yacl* gene, hereinafter designated *acnB*, is located at 2-85 min (131·6 kb) in the physical map of the *E. coli* chromosome (Fig. 1) and it is flanked by two further ORFs (*yacH* and *yacL*), which are in turn situated between the *lpdA* and *speD* genes.
Studies with putative \( \lambda \text{acnB} \) phages and characterization of phagemids containing the \( \text{acnB} \) gene

The first direct evidence that the \( \text{acnB} \) gene is located in the \( \text{yacJ} \) region was obtained by infecting JRG2789 with selected phages from the \( \lambda \)-\( \text{E. coli} \) W3110 gene library (Kohara et al., 1987). Aconitase activity was significantly and reproducibly increased about four- to sixfold after infection with \( \lambda \text{E}11 \) and \( \lambda \text{C}11 \) which contain the \( \text{yacJ} \) region (Fig. 1), but not with a phage \( \lambda \text{D}12 \) that lacks this region (Fig. 3a). There was also a reproducible correlation between the orientation of \( \text{acnB} \) \((\text{yacJ})\) in the phage genome and the early or late increase in aconitase activity (Fig. 3a). Thus, activity increased 20 min earlier with \( \lambda \text{C}11 \), where \( \text{acnB} \) should be transcribed from the early phage promoter \( (\text{AP}_1) \), than with \( \lambda \text{E}11 \), where \( \text{acnB} \) has the same polarity as the late promoter \( (\text{AP}_2) \).

Two derivatives of pUC118 containing the \( \text{acnB} \) \((\text{yacJ})\) gene of \( \lambda \text{E}11 \) were constructed by subcloning the 7.4 kb \( \text{PsI} \) fragment containing the \( \text{lpdA} \text{--yacH--acnB} \) region to generate pGS799, and the 6.5 kb \( \text{PvuII} \) fragment containing the \( \text{yacH} \text{--acnB} \text{--yacL} \) region to generate pGS801 (Fig. 1). In each case, the aconitase specific activities of the corresponding DH1 transformants were up to 10-fold higher than those of the vector control during early-stationary phase (Fig. 3b). SDS-PAGE analysis clearly showed that a polypeptide of the size expected for AcnB \((M_\text{r} 100000)\) is enriched in cell-free extracts of DH1(pGS799) and DH1(pGS801) prepared from late-exponential-phase cultures (Fig. 4a). Another polypeptide corresponding to the \( \text{lpdA} \) gene product, lipoamide dehydrogenase \((M_\text{r} 55000)\), was enriched in DH1(pGS799) but not in DH1(pGS801) which lacks the complete \( \text{lpdA} \) gene (Fig. 4a). No enriched products corresponding to the \( \text{yacH} \) or \( \text{yacL} \) genes were detected. Densitometric analyses of Coomassie-Blue-stained gels indicated that the putative AcnB protein amounts to 30% and 50% of soluble cell protein in the pGS799 and pGS801 transformants compared with <1% in the untransformed host, DH1. The degree of amplification is greater than might be expected from the increase in AcnB activity, suggesting that a large proportion of the enzyme accumulates in an inactive state. Attempts to reactivate this protein were unsuccessful.

In parallel \textit{in vitro} transcription-translation experiments, pGS799 and pGS801 each directed the synthesis of three major labelled polypeptides in addition to those produced in control reactions (Fig. 4b). The \( \text{acnB} \) product \((M_\text{r} 100000)\) and putative \( \text{yacH} \) product \((M_\text{r} 41000)\) were common to both, whereas products corresponding to the \( \text{lpdA} \) \((M_\text{r} 55000)\) and \( \text{yacL} \) \((M_\text{r} 11000)\) genes were associated only with pGS799 and pGS801, respectively.
The minor labelled products probably represent in vitro artefacts or degraded forms of the major polypeptides.

**Aconitase amplification from the acnB coding region**

The above observations are consistent with the proposal that AcnB is encoded by the combined yacI reading frame. To confirm this possibility, single-stranded pGS782 phagemid DNA was used to sequence a 393 bp DNA segment extending from the proposed translational start of acnB in yacI (coordinate 20231; Fujita et al., 1994, GenBank D26562) up to a point 117 bp within the downstream yacI reading frame (coordinate 20620). A single C insertion was detected immediately after coordinate 20455, indicating that the yacI reading frame is indeed contiguous with yacI. Relative to the previous sequence, an extra alanine codon (CGC, coordinates 20344–20346) and a silent T for C replacement (coordinate 20383), were also detected within the sequenced region (GenBank accession no. U41560, for revised sequence). The proposed acnB coding region starts with a GTG (formyl-methionine) codon preceded by a well-placed RBS (GAGCGAGGAGAACCGTCGTG). It extends for a total of 865 amino acid codons until terminated by a UAA stop codon (coordinate 22826–22828). The codon usage is typical of an *E. coli* coding region. Furthermore, the low proportion of modulatory codons (0·58%) combined with a strong preference for optimum energy codons (60·6%) suggests that acnB is strongly expressed compared with acnA, where the corresponding values (3·8 and 44·6%) are indicative of weak to moderate expression (Grosjean & Fiers, 1982; Prodromou et al., 1992). The size of the predicted gene product (M, 93500) is slightly less than that estimated from the electrophoretic mobility of AcnB. The coding region is flanked by a 593 bp upstream non-coding region containing several possible acnB and yacH promoters (Fig. 1) and by a 221 bp downstream region encoding a potential Rho-independent terminator, i.e. a GC-rich region of hyphenated dyad symmetry followed by a T(U)-rich sequence, ATGC-CCGGAGGATGATTCGCAGCGCATTTTTTTATT (coordinates 22861–22898; the non-complementary base in the GC-rich stem 1, the loop (9AG), and all of the bases in the T-rich sequence, are superscripted).

The gene was conclusively identified as the acnB gene by subcloning the PCR-amplified coding region downstream of the IPTG-regulated T7 promoter of pET21a, and showing that relative to vector-containing controls, aconitase activity could be raised by factors of 5- to 27-fold in JRG3099, a transformant of BL21/ADE3 containing the corresponding expression plasmid (pGS783, see Methods). The wide range of specific activities [2–11 U (mg protein)−1] observed after inducing cultures in mid-exponential phase (OD650 0·7), sharply contrasted with the almost uniform amplification of AcnB protein to 48% of soluble protein in Coomassie-Blue-stained SDS-PAGE fractionated samples (not shown). As with DH1(pGS799) and DH1(pGS801), it would appear that much of the amplified AcnB protein accumulates in an...
inactive form. The proportion of inactive AcnB protein was extremely variable and attempts to reactivate samples to a uniformly high activity failed.

It can be calculated that an iron concentration of 8.5 μM would be needed to provide BL21/ADE3(pGS783) with sufficient iron for the synthesis of active AcnB at 48% of soluble protein. A deficiency in iron might thus limit the synthesis of active AcnB because L broth contains only 17 μM iron (S. C. Andrews & A. J. Hudson, personal communication). After growth and induction in L broth supplemented with 50 μM iron citrate, the range of AcnB specific activities increased about two- to fourfold [to 3.5–41 U (mg protein)^{-1}, which is 9–120-fold higher than the vector-containing control]. This suggests that for equivalent amounts of protein, a greater proportion of AcnB produced in BL21/ADE3(pGS783) is in a reactivatable form when the iron concentration in the medium is increased. The specific activities were still very variable and again could not be reactivated to a uniform level. Nevertheless, iron citrate (50 μM) was added thereafter to the medium used for the amplification of AcnB. Raising the iron concentration to 100 μM produced no further increase in AcnB content or specific activity. Likewise, no increase in AcnB activity was observed when extra iron was added to the growth medium of DH1(pGS799) or DH1(pGS801) cultures, further indicating that most of the inactive protein is not in the reactivatable [3Fe-4S] form. The reason why DH1(pGS799) and DH1(pGS801) produced greater proportions of unreactivated AcnB than BL21/ADE3(pGS783) is not known, but it may reflect differences in iron–sulphur centre assembly or stability under different culture conditions.

### Purification of AcnB from *E. coli* JRG3171, DH1(pGS801)

*E. coli* JRG3171, a pGS801 transformant of DH1 containing multiple copies of the *acnB* gene, was chosen as the source for purifying AcnB because it gave consistently high specific activities and because AcnB was the only protein that was significantly overproduced in this strain. A typical purification is shown in Table 1 and an SDS-PAGE fractionation of samples taken at each step is shown in Fig. 2(c). The AcnB protein was purified to ≥ 98% homogeneity with an overall yield of 8% and a purification factor of 62-fold. Unlike the material from the unenriched source, the enzyme eluted at the final stage could be reactivated some 21-fold to give a final specific activity of 38 U (mg protein)^{-1}. The discrepancy apparent from the need for a sixfold enrichment to purify a protein that is already 50% pure in the starting material again suggests that there are different relative proportions of active, reversibly inactive and irreversibly inactive enzyme in different samples.

### DISCUSSION

Despite its inherent instability, the second *E. coli* aconitase (AcnB) was partially purified from an *acnA* mutant in extremely low but sufficient yield to permit identification and N-terminal sequence analysis of the polypeptide and subsequent isolation of the *acnB* gene. During purification no evidence was obtained for the presence of further aconitate activities in *E. coli*, but their existence cannot be excluded, especially as this organism is known to express three differentially regulated fumarases (Guest, 1992). Whether there are more than two aconitases is now being addressed by constructing an *acnA* *acnB* double mutant. Like other aconitases, AcnB is monomeric. Its M<sub>r</sub> (93500–105000 by different methods) most closely resembles those of AcnA (M<sub>r</sub> 97500) and the IRP (M<sub>r</sub> 98000), which are significantly larger than the mammalian and yeast mitochondrial enzymes (M<sub>r</sub> 79000–83000), but smaller than the *Bacillus subtilis* and *Salmonella typhimurium* enzymes (M<sub>r</sub> 120000 approx.).

Overproduction to 50% of soluble cell protein allowed AcnB to be purified essentially to homogeneity. The specific activity of pure reactivated AcnB [38 U (mg protein)^{-1}] is somewhat lower than that of reactivated AcnA [59 U (mg protein)^{-1}] purified from a comparably amplified source under similar conditions (Bennett *et al.*., 1995). However, such comparisons must be interpreted with care because a significant proportion of both the amplified-crude and purified AcnA and AcnB proteins could not be reactivated by incubating with ferrous ions under reducing conditions (Bennett *et al.*, 1995; M. J. Gruer & J. R. Guest, unpublished observations). Much of the AcnA protein would thus seem to be produced in an irreversibly inactivated apo-form, in addition to being irreversibly degraded during purification. It has been proposed that the iron released in the conversion of [4Fe-4S] to [3Fe-4S] clusters under aerobic conditions may generate hydroxyl radicals that inflict irreversible enzyme damage, because a smaller proportion of AcnA is irreversibly degraded after anaerobic purification (Bennett *et al.*, 1995). It will be interesting to determine whether the NifS protein, which catalyses the formation of iron–sulphur clusters using cysteine-derived enzyme-bound persulphide as the sulphide donor, will reactivate AcnB, as has been observed for apo-nitrogenase and FNR (Zheng & Dean, 1994; Khoroshilova *et al.*, 1995).

The *acnB* gene is flanked by two unidentified ORFs (*yaeH* and *yaeL*) which in turn are located between the pyruvate dehydrogenase operon (*pdfB–aceEF–lpdA*) and the S-adenosylmethionine decarboxylase gene, *speD* (Fig. 1). No clues about the functions of *yaeH* and *yaeL* could be gleaned from database searches for potential homologues. However, their locations suggest that they may have functions connected with the CAC and this possibility is being investigated.

Computer-based sequence comparisons using DIAGON (Staden, 1982) failed to reveal any sustained similarities between AcnB and AcnA or other members of the Acn/IRP/IPMI family. However, several conserved active-site residues and motifs were detected in AcnB by visual inspection, but these could only be aligned with their counterparts in AcnA, by cyclically permuting the domain organization from 1–2–3–4 in AcnA to 4–1–2–3 in AcnB. This rearrangement is illustrated in Fig. 5 where the domain corresponding to the C-terminal domain in
Aconitase (AcnA) and AcnB, the second aconitase of E. coli

AcnA (domain 4) becomes the N-terminal domain in AcnB. The long linker connecting domains 3 and 4 in AcnA is absent in the arrangement proposed for AcnB, where domain 4 is joined directly to the rest of the molecule via domain 1 (Fig. 5). Members of the Acn/IRP/IPMI family thus exhibit three distinct ways of achieving similar overall structural conformations, domain 4 being situated either at the N- or C-termini of single polypeptides, or provided by a completely independent polypeptide (Fig. 5). The present arrangements in AcnA and AcnB may have arisen by the fusion of two ancestral genes which had previously come together in different relative orders, one of which corresponds to the current bacterial IPMI (leuCD) situation. An alternative evolutionary route for generating a cyclically permuted coding region might have involved tandem duplication of the ancestral gene(s) followed by the deletion of flanking DNA. The relationship between either AcnA or AcnB and the two-subunit bacterial IPMI resembles the relationship between the dimeric iron–sulphur containing fumarases (FumA or FumB) and the tetrameric L-tartrate dehydratase (Ttd), where the amino acid sequence of each fumarase corresponds to the combined TtdA and TtdB sequence, the overall sequence identity being 23–25% (Reaney et al., 1993).

A multiple amino acid sequence alignment was made by repositioning different C-terminal segments of the AcnB sequence at the N-terminal end and making computer-based comparisons with selected Acn, IRP and IPMI sequences (Fig. 6). A computerized version of a previous multiple alignment (Prodromou et al., 1992) was first constructed using FILEUP (Higgins & Sharp, 1989) and LINEUP (Gribskov et al., 1990), and was then optimally aligned with AcnB using PROFILEMAKE and PROFILEGAP (Schwartz & Dayhoff, 1979). The alignment was ultimately refined by inspection. Some 17 of the 20 active-site residues assigned in porcine mitochondrial aconitase are conserved in AcnB, including the three cysteine residues involved in ligand binding to the iron–sulphur centre (Fig. 6). Most of the secondary structural elements in porcine aconitase appear to be conserved in AcnB and the insertion-deletions are located in superficial loops and turns. The main differences concern the absence of a sequence corresponding to the long interdomain linker in AcnB, the absence of three helical segments from the N-terminal region of domain 1, and the presence of 120 unaligned N-terminal residues connected to domain 4 in AcnB (Fig. 6). The overall amino acid sequence identities between AcnB and porcine aconitase (15%), AcnA (17%), IRP (17%), and the IPMI of Mucor circinelloides (18%) and S. typhimurium (15%), were deduced using DISTANCES (University of Wisconsin Genetics Group). The corresponding relationships with AcnA are: porcine aconitase (29%); AcnB (17%); IRP (53%); and the IPMI of M. circinelloides (21%) and S. typhimurium (21%), indicating that AcnB is only distantly related to other members of the family.

Possible reasons for the duplication of metabolic genes and enzymes including citrate synthase, succinate dehydrogenase and fumarase, have been discussed previously (Gruer & Guest, 1994; Guest, 1995). AcnA and AcnB appear not to have mutually exclusive functions because AcnB successfully complements the AcnA deficient in the acnA::kanR mutant. It will be interesting to ascertain whether the converse is true in an acnB mutant. The codon preferences suggest that the acnB gene is more strongly expressed than acnA. It seems likely that acnB will exhibit a different pattern of regulatory responses to acnA, which is subject to ArcA- and FNR-mediated repression and CRP-, SoxRS-, RpoS-, and Fur-mediated activation, and it has been suggested that AcnB might be preferentially used in the glyoxylate cycle (Gruer & Guest, 1994; M. J. Gruer & J. R. Guest, unpublished...
observations). It may be significant that the genes encoding the aerobic fumarase (FumA) and aconitase (AcnA) are activated by iron and Fur, because this could provide a mechanism for controlling citric acid cycle activity in response to iron availability. It is also possible that both of the aconitases might function as an iron-responsive translational regulator analogous to mammalian IRP. Current work is aimed at studying the regulation of acnB gene expression using acnB-lacZ fusions, constructing and defining the phenotype of an acnB mutant, and investigating the properties of an acnA acnB double mutant. This approach might identify specific metabolic roles for AcnA and AcnB, and explain why E. coli contains two aconitase isoenzymes.

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