Direct evidence for mRNA binding and post-transcriptional regulation by *Escherichia coli* aconitases

Yue Tang and John R. Guest

**Abstract**

*Escherichia coli* contains a stationary-phase aconitase (AcnA) that is induced by iron and oxidative stress, and a major but less stable aconitase (AcnB) synthesized during exponential growth. These enzymes were shown to resemble the bifunctional iron-regulatory proteins (IRP1)/cytoplasmic aconitases of vertebrates in having alternative mRNA-binding and catalytic activities. Affinity chromatography and gel retardation analysis showed that the AcnA and AcnB apo-proteins each interact with the 3'H\(_{\text{UTR}}\)s of *acnA* and *acnB* mRNA at physiologically significant protein concentrations. AcnA and AcnB synthesis was enhanced *in vitro* by the apo-aconitases and this enhancement was abolished by 3'H\(_{\text{UTR}}\) deletion from the DNA templates, presumably by loss of acn-mRNA stabilization by bound apo-aconitase. *In vivo* studies showed that although total aconitase activity is lowered during oxidative stress, synthesis of the AcnA and AcnB proteins and the stabilities of *acnA* and *acnB* mRNAs both increase, suggesting that inactive aconitase mediates a post-transcriptional positive autoregulatory switch. Evidence for an iron–sulphur-cluster-dependent switch was inferred from the more than threefold higher mRNA-binding affinities of the apo-aconitases relative to the holo-enzymes. Thus by modulating translation via site-specific interactions between apo-enzyme and relevant transcripts, the aconitases provide a new and rapidly reacting component of the bacterial oxidative stress response.

**Keywords:** conitase, mRNA stability, oxidative stress, post-transcriptional regulation, protein–mRNA interactions

---

**INTRODUCTION**

The aconitases convert citrate to isocitrate via cis-aconitate in the citric acid and glyoxylate cycles. They are monomeric enzymes containing single iron–sulphur clusters which are interconvertible between the catalytically active [4Fe–4S] forms and the inactive [3Fe–4S] and apo-enzyme forms (Beinert *et al.*, 1996). Crystallographic studies have shown that porcine heart mitochondrial aconitase contains four structural domains (Robbins & Stout, 1989). Domains 1, 2 and 3 are tightly packed around the iron–sulphur cluster, which is ligated to three cysteine residues in domain 3, and domain 4 is connected by an extended linker to domain 3 so as to form a deep active-site cleft between domain 4 and the rest of the protein. This is the structural archetype for all members of the aconitase protein family including the vertebrate bifunctional iron-regulatory proteins/cytoplasmic aconitases (IRP1/c-Acn) and the monofunctional iron-regulatory proteins (IRP2). The IRPs are cytoplasmic mRNA-binding proteins that co-ordinate the post-transcriptional synthesis of several important proteins concerned with iron metabolism in higher organisms (Hentze & Kuhn, 1996). They block the translation of some mRNAs, e.g. ferritin mRNA, by binding to iron-responsive elements (IREs) in the 5′ untranslated region (5′UTR) of the transcript; or alternatively they enhance the translation of other mRNAs, e.g. transferrin receptor mRNA, by increasing mRNA stability as a result of binding to IREs in the 3′ untranslated region of the transcript (Hentze & Kuhn, 1996; Hirling *et al*., 1994; Klausner & Rouault, 1996).

Abbreviations: IRE, iron-responsive element; IRP, iron-regulatory protein; MV, methyl viologen; NEM, N-ethylmaleimide; UTR, untranslated region.
there is a reciprocal iron-dependent relationship between mRNA-binding and aconitase activity such that the [4Fe–4S] protein is catalytically active whereas the apo-enzyme binds mRNA via the active-site cleft (Hentze & Kuhn, 1996). The RNA-binding activity of IRP1 is likewise increased in response to agents such as hydrogen peroxide and nitric oxide which induce oxidative stress and inactivate the enzyme (Pantopoulos & Hentze, 1995a, b).

In Escherichia coli, two aconitases (AcnA and AcnB) have been identified and the existence of a third aconitase (AcnC) has been inferred from the residual activity (<5% of wild-type) detected in an acnAB double mutant (Prodromou et al., 1992; Bradbury et al., 1996; Gruer et al., 1997b). Sequence comparisons show that AcnA is 53% identical to human IRP1 and 27–29% identical to mitochondrial aconitases, whereas AcnB is only 15–17% identical to AcnA, the IRPs and mitochondrial aconitases. The major IRE-binding site of IRP1 (DLVIDHSIQVD), identified by UV cross-linking (Basilion et al., 1994), is likewise conserved at nine of the eleven positions in AcnA but only at two positions in AcnB. It is also apparent that the domain organization of AcnB (4-1-2-3) is cyclically permuted relative to the arrangement (1-2-3-linker-4) found in other aconitases and the IRPs (Gruer et al., 1997a). Specific roles for AcnA and AcnB have been identified by physiological and enzymological studies with acn mutants (Gruer et al., 1997b), regulatory studies with acn–lacZ reporter fusions, and transcript analyses (Gruer & Guest, 1994; Cunningham et al., 1997). Thus, AcnB is the major aconitase synthesized during the exponential phase, whereas AcnA is a stationary-phase enzyme which is also specifically induced by iron and oxidative stress. The aerobic growth of an acnA mutant is unimpaired in glucose minimal medium, indicating that this lesion is fully complemented by acnb⁺, whereas the growth of an acnB mutant is severely impaired and only slightly improved by adding glutamate. Growth is virtually abolished in an acnAB double mutant, except in the presence of glutamate, where the spontaneous mutational inactivation of the citrate synthase gene (gltA) seems to eliminate the deleterious effects of citrate accumulation and leads to the proliferation of acnAB gltA triple mutants. The regulatory studies further showed that the acnA gene is expressed from independent σ28 and σ70 promoters, the latter being repressed by FNR and ArcA and activated (directly or indirectly) by CRP, Fur, iron, FruR and SoxRS, whereas the acnB gene is expressed from a single σ70 promoter that is repressed by ArcA, FruR and Fis, and activated by CRP.

It is not known whether any bacterial aconitases have regulatory functions like the IRPs. Aconitase is needed in Bacillus subtilis for sporulation but its mode of action is unclear (Dingman et al., 1987), and in Xanthomonas campestris a gene (rpfA) which regulates pathogenicity factor production has recently been shown to encode an aconitase that may serve as an iron-responsive transcription regulator (Wilson et al., 1998). The present work shows that the apo-forms of AcnA and AcnB of E. coli each bind to the 3’UTRs of acnA mRNA and acnB mRNA and enhance the synthesis of both proteins (in vitro), presumably by increasing mRNA stability. A direct correlation between enzyme inactivation, increased Acn protein synthesis and acn mRNA stability, observed with oxidatively stressed cultures, further indicates that mRNA binding by bacterial aconitases represents a novel component of the oxidative stress response. Brief accounts of these findings were communicated previously at a Gordon Research Conference (Microbial Response to Stress, 1998) and the World Congress on Iron Metabolism (BioIron’99 Abs 5).

**METHODS**

**Bacterial strains, plasmids and basic methods.** The strains of E. coli and plasmids used as enzyme sources, genes and RNA, are listed in Table 1. An acnA expression plasmid, pGS1203, analogous to pGS783, the pET21a-derived acnB expression plasmid (Bradbury et al., 1996), was constructed by first replacing the 1.6 kb KpnI–SalI fragment of pGS447 by a 369 bp PCR-amplified fragment containing the 5’ end of the acnA coding region flanked by KpnI and NdeI sites at one end and the acnA SalI site at the other end, to generate pGS1202. The sequence of the cloned region was confirmed and the entire 3.2 kb NdeI–HindIII fragment was then transferred to pET21a to generate pGS1203. Cultures were grown aerobically in 2 × YT medium at 37 °C with ampicillin (100 µg ml⁻¹, when required) for plasmid isolation (Promega) and aconitase production. Protein was assayed by the Bio-Rad method with bovine serum albumin (BSA) as standard: non-denaturing PAGE and SDS-PAGE were performed according to Sambrook et al. (1989). DNA was prepared and manipulated by standard procedures (Sambrook et al., 1989). Total bacterial RNA was extracted from early-stationary-phase cultures (unless otherwise specified) by the hot acid phenol method (Aiba et al., 1981). The RNA (100 µg) was resuspended in 50 µl buffer containing 20 mM Tris–HCl (pH 8.0), 10 mM MgCl₂, 1 mM DTT and 10% (v/v) glycerol before adding 20 U RNAse-free DNase I and incubating at 37 °C for 30 min to remove traces of DNA. The RNA was then phenol-extracted and precipitated with ethanol. Western blotting was according to Gruer et al. (1997b) using enzyme-specific polyclonal antisera. Samples containing 15 µg protein were fractionated by SDS-PAGE (10% gel) and electroblotted to Hybond-C (Amersham) for staining and subsequent quantitative densitometry using a Bio-Profil (Vilber Lourmat) with 1D and 2D analytical software.

**Overproduction, purification and assay of aconitases.** The enzymes were purified from shaken cultures (2 × 500 ml) of JRG4004 (acnA) and JRG3069 (acnB) grown to OD₅₆₀ 0.5 at 37 °C before induction with IPTG (1 mM) for 4 h at 25 °C (AcnA) or 2 h at 37 °C (AcnB). The bacteria were resuspended in 20 ml Tris–HCl buffer (40 mM, pH 8.0) for preparing ultrasonic extracts. Ammonium ferrous sulphate (0.25 mM, final concn) and DTT (2.5 mM, final concn) were added to the cell-free extracts to reactivate the aconitases. Extract containing 150 mg protein in 20 ml was loaded on a DEAE Sepharose column (16 × 35 mm), washed with 5 vols Tris–HCl (40 mM, pH 8.0), and eluted with a linear gradient (300 ml, 0–0.2 M sodium citrate in the same buffer) using an FPLC system (Pharmacia). Coloured fractions containing AcnA or AcnB were pooled and ammonium sulphate (1.7 M, final
concn) was added before applying to an Ether 650S column (16 × 40 mm). The column was washed with 5 vols 40 mM Tris/HCl (pH 8.0) containing 1.7 M ammonium sulphate and the Acn protein was eluted with a linear gradient (200 mM, 1-70 M ammonium sulphate in the same buffer). Typically about 30 mg Acn protein (>95% pure) was recovered.

The apo-aconitases were obtained by incubating holo-enzyme solutions with a 50-fold molar excess of EDTA (pH 8.0) and a 20-fold molar excess of potassium ferricyanide for 6 min at 4 °C according to Kennedy & Beinert (1988) and the protein was recovered using a small Ether 650S column, as above. Aconitase activity was measured according to Kennedy et al. (1983): one unit of activity represents 1 µmol cis-aconitate formed min⁻¹. The native (as isolated) aconitases retained 6–20% of the specific activity and approximately 70% of the iron of the fully reconstituted enzymes whereas the apo-proteins were completely lacking in iron and totally inactive.

**Affinity chromatography and RT-PCR procedures.** Apo-AcnA and apo-AcnB (500 µg protein) were coupled to 100 µl of activated CH Sepharose 4B (Pharmacia) by incubating for 16 h at 4 °C in 0.1 M NaHCO₃ buffer pH 8.0 containing 0.5 M NaCl (final vol., 500 µl). The affinity matrices were washed exhaustively and blocked with 0.1 M Tris/HCl (pH 7.5), whereafter 20 µl samples were used to prepare affinity columns. Columns were equilibrated with 20 µg total MC4100 RNA in 200 µl RNA-binding buffer (10 mM HEPEs pH 7.5, 3 mM MgCl₂, 0.1% Triton X-100, 400 mM KCl and 5% glycerol), as well as 1% β-mercaptoethanol, 200 µg BSA, 150 U RNasin (Promega) and 20 µl tRNA, for 30 min at 20 °C, and then washed with 5 ml of the RNA-binding buffer to remove unbound RNA. Bound RNA was eluted with 50 µl 0.3 M sodium acetate buffer (pH 4.5) and precipitated with ethanol.

Reverse transcription was performed with all of the affinity-enriched RNA sample or 2 µg of total RNA in 2 µl, mixed with 1 µ reverse primer (40 µM), 1 µ reverse transcriptase buffer (10 ×), and 2 µl dNTPs (5 mM each), and heated to 70 °C for 10 min before cooling slowly to room temperature. Thereafter, 0.5 µl sodium pyrophosphate (80 mM), 0.5 µl actinomycin D (500 µg ml⁻¹), 1 µl placental ribonuclease inhibitor (Promega), 1 µl AMV reverse transcriptase (25 U; NBL) and H₂O to 10 µl were added and the mixtures were incubated at 37 °C for 90 min before the product DNA was precipitated by adding 1 µl sodium acetate (3 M; pH 4.5) and 50 µl ethanol. Specific segments of cDNA (~300–600 bp copied from relevant coding regions) were then amplified by PCR with Tag DNA polymerase (Promega) over 25 cycles using appropriate forward primers and the same reverse primers, and the products were analysed by agarose gel electrophoresis (1.5 %) using a 0.25–100 kb GeneRuler (MBI Fermentas) for calibration.

RNA Sepharose affinity columns were prepared by cross-linking in vitro transcription products of the A5 and B5 3'UTR fragments, or 0.5 µg tRNA, to 100 µl activated CH Sepharose as described above. The columns were equilibrated with Acn protein (50 µg in 200 µl RNA-binding buffer) for 15 min at 20 °C before washing (2 × 2 ml RNA-binding buffer), eluting with successive 2 × 300 µl aliquots of the same buffer containing increasing concentrations of NaCl (0.1–3 M), and analysing the fractions by SDS-PAGE.

**Gel retardation analysis.** RNA probes were first denatured by heating at 70 °C for 5 min and cooled rapidly. RNA-binding reactions were performed in 1 µl 10 × RNA-binding buffer (100 mM HEPEs pH 7.5, 30 mM MgCl₂, 1% Triton X-100, 400 mM KCl and 50%, glycerol), 1 µl β-mercaptoethanol, 200 µg BSA, 150 U RNasin (Promega) and 20 µl tRNA, for 30 min at 20 °C, and then washed with 5 ml of the RNA-binding buffer to remove unbound RNA. Bound RNA was eluted with 50 µl 0.3 M sodium acetate buffer (pH 4.5) and precipitated with ethanol.

Reverse transcription was performed with all of the affinity-enriched RNA sample or 2 µg of total RNA in 2 µl, mixed with 1 µ reverse primer (40 µM), 1 µ reverse transcriptase buffer (10 ×), and 2 µl dNTPs (5 mM each), and heated to 70 °C for 10 min before cooling slowly to room temperature. Thereafter, 0.5 µl sodium pyrophosphate (80 mM), 0.5 µl actinomycin D (500 µg ml⁻¹), 1 µl placental ribonuclease inhibitor (Promega), 1 µl AMV reverse transcriptase (25 U; NBL) and H₂O to 10 µl were added and the mixtures were incubated at 37 °C for 90 min before the product DNA was precipitated by adding 1 µl sodium acetate (3 M; pH 4.5) and 50 µl ethanol. Specific segments of cDNA (~300–600 bp copied from relevant coding regions) were then amplified by PCR with Tag DNA polymerase (Promega) over 25 cycles using appropriate forward primers and the same reverse primers, and the products were analysed by agarose gel electrophoresis (1.5 %) using a 0.25–100 kb GeneRuler (MBI Fermentas) for calibration.

RNA Sepharose affinity columns were prepared by cross-linking in vitro transcription products of the A5 and B5 3'UTR fragments, or 0.5 µg tRNA, to 100 µl activated CH Sepharose as described above. The columns were equilibrated with Acn protein (50 µg in 200 µl RNA-binding buffer) for 15 min at 20 °C before washing (2 × 2 ml RNA-binding buffer), eluting with successive 2 × 300 µl aliquots of the same buffer containing increasing concentrations of NaCl (0.1–3 M), and analysing the fractions by SDS-PAGE.

**Gel retardation analysis.** RNA probes were first denatured by heating at 70 °C for 5 min and cooled rapidly. RNA-binding reactions were performed in 1 µl 10 × RNA-binding buffer (100 mM HEPEs pH 7.5, 30 mM MgCl₂, 1% Triton X-100, 400 mM KCl and 50%, glycerol), 1 µl β-mercaptoethanol, 200 µg BSA, 150 U RNasin (Promega) and 20 µl tRNA, for 30 min at 20 °C, and then washed with 5 ml of the RNA-binding buffer to remove unbound RNA. Bound RNA was eluted with 50 µl 0.3 M sodium acetate buffer (pH 4.5) and precipitated with ethanol.

Reverse transcription was performed with all of the affinity-enriched RNA sample or 2 µg of total RNA in 2 µl, mixed with 1 µ reverse primer (40 µM), 1 µ reverse transcriptase buffer (10 ×), and 2 µl dNTPs (5 mM each), and heated to 70 °C for 10 min before cooling slowly to room temperature. Thereafter, 0.5 µl sodium pyrophosphate (80 mM), 0.5 µl actinomycin D (500 µg ml⁻¹), 1 µl placental ribonuclease inhibitor (Promega), 1 µl AMV reverse transcriptase (25 U; NBL) and H₂O to 10 µl were added and the mixtures were incubated at

<table>
<thead>
<tr>
<th>Table 1. Strains of E. coli and plasmids</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strain or plasmid</strong></td>
</tr>
<tr>
<td>------------------------</td>
</tr>
<tr>
<td>W3110</td>
</tr>
<tr>
<td>DH1</td>
</tr>
<tr>
<td>DH5x</td>
</tr>
<tr>
<td>MC4100</td>
</tr>
<tr>
<td>DJ901</td>
</tr>
<tr>
<td>BL21/D3</td>
</tr>
<tr>
<td>JRG2387</td>
</tr>
<tr>
<td>JRG3099</td>
</tr>
<tr>
<td>JRG317</td>
</tr>
<tr>
<td>JRG4004</td>
</tr>
<tr>
<td>pGS1203</td>
</tr>
</tbody>
</table>
2.5 mM), 1 µl BSA (20 mg ml⁻¹), 2 µl [α-32P]UTP (3000 Ci mmol⁻¹; ICN), 2 µl T3 RNA polymerase (Promega; 40 U) and water to 20 µl. After incubation at 37 °C for 90 min, 10 U RNase-free DNase I was added to remove the DNA templates, and labelled RNA was purified by phenol and ethanol precipitation.

**In vitro transcription-translation.** DNA (1 µg), either uncut plasmid DNA or gel-purified PCR-amplified DNA (Expand High Fidelity PCR System, Boehringer), was mixed with 3.75 µl S30 extract (linear), 0.25 µl [35S]methionine [1000 Ci mmol⁻¹; ICN], 1.25 µl amino acid mixture minus methionine, 5 µl S30 premix, aconitase (where stated) and H₂O to 12.5 µl, before incubating at 37 °C for 90 min (Promega, *E. coli* S30 Extract System for Linear Templates). When used, NEM was incubated with the aconitase (30 min at 4 °C in 2.5 µl) before completing the reaction mixtures. Translation products were precipitated with 50 µl acetone and resuspended in loading buffer for SDS-PAGE fractionation, autoradiography and quantitative analysis of 35S-labelled components (c.p.m.), which were excised from the gels and counted in vials containing 3 ml Safefluor S (Lumac-LSC) using a Beckman LS 1801 Liquid Scintillation System.

**Methyl-violelogen-induced oxidative stress.** The effects of methyl viologen (MV) on Acn activity and Acn protein synthesis were studied with cultures of *E. coli* DJ901 (∆sosX) grown for 2 h to exponential phase. Rifampicin (200 µg ml⁻¹) was added immediately by MV and incubation for a further 1 h before measuring total aconitase activity (without reactivation) and immunoblotting with specific anti-Acn sera. The effects of MV on *acn* mRNA content and stability were studied by adding MV (0.3 mM) 10 min before rifampicin (200 µg ml⁻¹), to cultures of W3110. Samples taken at 0, 2, 5, 10 and 20 min were chilled immediately to 4 °C in liquid N₂ for RNA extraction and quantitative Northern blotting analysis. The RNA samples (25 µg) were fractionated in 1% agarose formaldehyde gel, transferred to Hybond-N (Amersham), and hybridized with [α-32P]dCTP-labelled probes according to the ‘Ready to Go’ labelling-kit instructions (Pharmacia). The *acnA* and *acnB* probes containing the intact 2.5 kb (*acnA*) and 2.8 kb (*acnB*) coding regions, were generated by PCR and the relative densities of hybridized bands were measured with an Instant Imager (Electronic Autoradiography; Packard).

**RESULTS**

**Specific RNA binding by AcnA and AcnB**

In view of the close structural relationship between AcnA and IRP1, the possibility that one or both of the *E. coli* aconitases might bind to specific mRNAs and modulate protein synthesis at the post-transcriptional level was investigated. Direct evidence for mRNA binding by AcnA and AcnB was accordingly sought using a combination of affinity chromatography and RT-PCR. Affinity columns prepared from CH Sepharose coupled to apo-AcnA and apo-AcnB were equilibrated with total bacterial RNA purified from MC4100, and washed free of unbound RNA before eluting and precipitating bound RNA. The presence of specific mRNAs in the affinity-purified material was then investigated by RT-PCR using pairs of primers designed to amplify specific segments of the coding regions of selected genes. The efficacy of the primers was established by showing that products of the predicted size were amplified when total RNA was used (Fig. 1). The experiments showed that *acnA* mRNA and *acnB* mRNA bind to both types of column but not to the unliganded matrix (Fig. 1a, b). No *acn* mRNA was detected when reverse transcriptase was omitted from the amplification reactions or when no RNA was added to the affinity matrices at the first stage (not shown). The specificity of the interaction with the *acnA* and *acnB* transcripts was inferred from the failure to detect the highly abundant transcript of a related citric acid cycle gene (*sucD*) or transcripts of the iron-regulated bacterioferritin gene (*bfr*) in the bound RNA fraction, even though these transcripts were present in the initial samples of total bacterial RNA (Fig. 1c, d). It was therefore concluded that apo-AcnA and apo-AcnB each interact with some degree of specificity to *acnA* and *acnB* mRNA. This could be mediated by one or more sequence motifs that are shared by both of the *acn* transcripts and presumably by other untested transcripts, but not by the *sucD* or *bfr* transcripts. Attempts were made to identify other bound transcripts by making cDNA libraries of the affinity-purified RNA in pUC18. However, this was frustrated by the high proportion of clones (all of 20 tested) that contained 16S rRNA or 23S rRNA sequences, which are presumably bound non-specifically. No attempts were made to compare the RNA-binding activities of affinity columns made with apo-protein versus native protein because the latter invariably contain some apo-protein which would react positively.

**Location of Acn-binding sites in acn transcripts**

Local homology searches in the regions flanking and overlapping the *acnA* and *acnB* genes failed to reveal conserved sequences or secondary structural features like IREs that might serve as Acn-binding sites. The presence and location of such sites was investigated by gel retardation analysis using both aconitases and overlapping segments of the 5’ and 3’UTRs of both *acn* transcripts derived by transcription in vitro from two plasmids, pGS447 (*acnA*) and pGS801 (*acnB*), as shown in Fig. 2(a). Specific retardation by apo-AcnA and apo-AcnB was only observed with two RNA fragments, A5 and B5, representing the 3’UTRs of the *acnA* and *acnB* transcripts (Fig. 2b). No specific retardation was observed with any of the 5’UTR fragments shown in Fig. 2(a). Likewise, subsegments of the 3’UTR fragments (A4, A6, B3 and B4) were not retarded, suggesting that more than one binding site is required or that long molecules are essential for AcnA and AcnB binding. The retardation patterns also indicated that more than one protein molecule can be bound (Fig. 2b).

Apo-AcnA had a lower affinity (Kₐ approximately 8 µM) for both A5 and B3 compared to A5-B3 A5-B3 (Kₐ approximately 1.3 µM), where Kₐ denotes the protein concentration giving half-maximal initial retardation. Although relatively low, the binding affinities are physiologically significant when related to the intracellular concentrations of aconitase deduced from specific activity measurements and Western blotting analysis: 1–6 µM for AcnA and 2–12 µM for AcnB, at different stages of the growth cycle (Gruer et
Fig. 1. RNA binding to immobilized AcnA and AcnB apo-proteins. The presence of specific transcripts in affinity-purified RNA was detected after RT-PCR amplification with gene-specific primers by agarose gel electrophoresis. The samples of bound RNA were eluted from unliganded Sepharose (s), AcnA-Sepharose (As) and AcnB-Sepharose (Bs). Total unfractionated RNA (T) was used as a control. The predicted sizes of the products (shown below) were confirmed by comparison with standard markers (M). The primers were designed to amplify specific segments of the coding regions of the following genes: (a) acnA (354 bp); (b) acnB (370 bp); (c) sucD (518 bp); (d) bfr (431 bp).

Fig. 2. Location of the Acn-binding regions of acnA mRNA and acnB mRNA by gel-retardation analysis. (a) Diagram showing segments of the 5′ and 3′UTRs of the acnA and acnB transcripts (A1–A6 and B1–B5) synthesized by transcription (in vitro) with T3 RNA polymerase using templates that were PCR-amplified from acnA and acnB plasmids using different pairs of oligonucleotide primers (converging arrows). The angled 5′ extremities denote the T3 promoter sequences incorporated by the forward primers, and the total length of each amplified molecule (bp) is indicated. (b) Gel retardation by apo-Acn proteins of radiolabelled RNA fragments (i) A5 and (ii) B5: lanes 1 and 4, no protein; lane 2, 12 µg apo-AcnA; lane 3, 24 µg apo-AcnA; lane 5, 3 µg apo-AcnB; lane 6, 5 µg apo-AcnB. No other fragments or subfragments were specifically retarded by AcnA or AcnB.

al., 1997b). The shifts were correspondingly smaller for AcnA than those induced by AcnB (Fig. 2b). This could reflect inherently different Acn-binding affinities based on binding-site recognition. Alternatively, the presence of 124 additional amino acid residues in domain 4 of AcnB and correspondingly fewer in domains 1–3 compared to AcnA (Gruer et al., 1997a) might have a greater effect on the mobility of the apo-AcnB–RNA complex, especially if RNA is bound in the active-site cleft.

In order to ensure that the natural acn transcripts extend across the regions of Acn binding, RT-PCR was used with total E. coli RNA to provide evidence for transcription of the corresponding 3′UTRs. Two series of reverse primers designed to generate and amplify cDNAs from 169, 269 and 336 bp downstream of the acnA stop codon, and 81, 199, 308, 482 and 528 bp downstream of the acnB stop codon, were paired with forward primers hybridizing within the respective coding regions. Amplified products were obtained in every case except in the controls lacking reverse transcriptase. Thus it was concluded that even though the 3′ transcription termini were not defined, they are located downstream of the longest 3′UTR segments (A5 and B5) studied here.

Effects of the AcnA and AcnB proteins on acn gene expression in vitro

The regulatory effects of IRPs have been observed in transcription–translation systems in vitro (Kim et al., 1996; Gray et al., 1993). Analogous studies with an E.
apo-Acn concentrations as low as 0 µM; 240 µg protein ml⁻¹) except in one case where higher concentrations (≥24 µM; 240 µg protein ml⁻¹) of apo-AcnB were needed to stimulate acnB expression. In the same experiments, expression of the plasmid-encoded bla and yacH genes was unaffected by the aconitases (Fig. 3). No incorporation in the Acn region was observed with vector controls, nor was any enhancement observed with comparable concentrations of bovine serum albumin. It was therefore concluded that the Acn apo-proteins specifically enhance expression of the acn genes, possibly at the translational level by acn transcript stabilization (via 3'UTR binding), as proposed for IRP1 and transferrin receptor mRNA.

Role of the 3' UTR in translational enhancement and mRNA binding

In order to confirm that the 3'UTRs of the acn mRNAs are needed for Acn-mediated translational enhancement and to locate potential Acn-binding sites, two series of PCR-amplified fragments containing acn genes that terminate at different 3' positions were used as linear templates in in vitro transcription–translation reactions (Fig. 4). With acnA a 3'UTR of between 71 and 169 bp was needed for translational enhancement by the apo-Acn proteins; with acnB an 80 bp 3'UTR was sufficient to detect enhancement at high protein concentrations but better enhancement was observed with a 297 bp 3'UTR. Further support for the existence of specific interactions between the Acn apo-proteins and acn 3'UTRs was obtained using affinity columns prepared by cross-linking the 3'UTRs (A5 and B5 in Fig. 2) and tRNA to CH Sepharose (see Methods). Apo-AcnA and apo-AcnB were bound to the A5 and B5 Sepharose matrices in RNA-binding buffer and released by adding salt to the eluting buffer but neither protein was bound to tRNA Sepharose (data not shown).

Effects of treating the Acn proteins with NEM

The RNA-binding activity of IRP1 is extremely sensitive to alkylation of the highly conserved Cys-357 residue with NEM (Hirling et al., 1994). In contrast, the RNA-binding activities of both Acn apo-proteins were impaired only by treatment with NEM at low concentrations (0-01-0-06 mM) and were unaffected or even augmented with higher NEM concentrations (0-6-10 mM) that would completely abolish IRP1 activity. Nevertheless, these observations served to demonstrate that there is a direct correlation between the mRNA-binding and in vitro translation-enhancing activities of the Acn proteins because acnA and acnB expression was impaired with apo-proteins treated with 0-01 and 0-1 mM NEM but augmented by treatment with 1-0 mM NEM (data not shown).

Effects of stress factors on Acn enzyme activity, Acn synthesis and acn mRNA stability, in vivo

Oxidative stress factors abolish the catalytic activity with the concomitant acquisition of the mRNA-binding and stabilizing activities of IRP1 (Pantopoulos &
Fig. 4. Stimulation of in vitro transcription–translation by Acn apo-proteins using acn genes with nested 3′UTR deletions as templates. PCR amplification was used to generate acn genes that extend to different positions beyond the stop codons (distance from stop codon indicated in bp; total length of template ∼3–5 kb). These served as linear acnA (a) or acnB (b) templates in in vitro transcription–translation reactions with different Acn proteins (µg protein ml⁻¹): no Acn (lane 1); apo-AcnA, 120 (lane 2) and 1200 (lane 3); apo-AcnB, 120 (lane 4) and 1200 (lane 5). The relative extents of Acn labelling are shown below each autoradiograph, where 10 represents 70–150 c.p.m. per band above background (25 c.p.m.). The converging arrows (A1–3 and B1–3) denote potential stem–loops in the transcripts.

Hentze, 1995a, b; Constable et al., 1992). The conversion of active bacterial aconitases to the inactive [3Fe–4S] or apo-enzyme forms could likewise provide a comparable stress-responsive switch for stabilizing acn and other relevant transcripts in vivo, especially as the aconitase activity of E. coli is known to be sensitive to oxidative stress (Gardner & Fridovich, 1992; Gardner et al., 1997). Accordingly, the effect of MV-induced oxidative stress on aconitase activity and Acn protein synthesis was investigated in growing cultures of E. coli. Strain DJ901 (∆soxR) was chosen because it allowed translation to be studied during oxidative stress without interference from SoxRS-dependent activation of acnA transcription (Grüer & Guest, 1994; Cunningham et al., 1997). In a typical experiment the total aconitase activity decreased by 60% whilst the AcnA and AcnB proteins increased three- to fourfold in response to added MV (Fig. 5). Similar reciprocal effects were observed with H₂O₂ (0.5 mM) and manganous ions (50 µM). These findings are therefore consistent with the view that inactivated aconitases enhance the synthesis of Acn proteins in vivo, presumably by stabilizing the corresponding mRNAs.

Direct evidence that oxidative stress stabilizes the acnA and acnB transcripts was obtained by measuring their half-lives in cultures of W3110 that had been treated with and without MV for 10 min before adding rifampicin to block transcription initiation (see Methods). After the anticipated MV-induced increase in acnA transcription (Cunningham et al., 1997), the half-life of acnA mRNA increased from 4 min to 17 min and that of acnB mRNA increased from 7 min to 15 min, relative to
untreated controls (Fig. 6). The results clearly show that oxidative stress increases acn transcript stability and hence transcript concentration in the absence of de novo transcription. They also confirm that oxidative-stress-mediated translational enhancement, presumably by apo-Acns, occurs in vivo as well as in vitro.

**DISCUSSION**

The close structural relationship between *E. coli* AcnA and the bifunctional IRP1s of higher organisms prompted a series of *in vitro* and *in vivo* experiments which revealed the existence of a post-transcriptional mechanism for positive autoregulation of AcnA and AcnB synthesis by their apo-proteins. It was further inferred from the effects of MV that conversion of the active holo-enzyme into a site-specific RNA-binding apo-protein by oxidative [4Fe–4S]-cluster degradation provides a stress-responsive regulatory switch, as in the case of IRP1. In bacteria, where stress responses are primarily controlled at the transcriptional level by specific regulators like Fur, SoxRS, OxyR and FNR, the Acn-mediated system offers a rapidly responding mechanism for modulating translation that may be of much greater general significance than simply controlling Acn synthesis. However, there are significant differences between the IRP1 and bacterial Acn systems, particularly with respect to mRNA-binding affinity, binding-site specificity, and potential physiological functions.

Early attempts to detect AcnA binding to mammalian IREs by gel retardation were unsuccessful (R. D. Klausner & J. R. Guest, unpublished) and the same was found with MICP, the AcnA of *Legionella pneumophila* (Mengaud & Horwitz, 1993). Three IRE-like sequences of unknown significance have been found in *E. coli*.
Fig. 8. Potential secondary structures of the acnA and acnB 3'UTRs. Putative stem–loops, denoted A1–3 and B1–3 as in Fig. 5, are shown with the corresponding free energies. The sequences are numbered from the bases that immediately follow the stop codons; identical or related motifs shared by both sequences are shown in bold-face, boxed or highlighted.

The binding affinities of apo-AcnA and apo-AcnB for the acn 3'UTRs ($K_D \approx 8.0$ and $1.3 \mu M$, respectively) are much lower than have been reported for IRP1 and IRE ($K_D = 5–50 \mu M$; Butt et al., 1996). Nevertheless, relative to the intracellular concentrations of AcnA and AcnB (6 and 12 $\mu M$ maximum, respectively), the apo-protein binding affinities are entirely compatible with the operation of two sensitive physiologically poised switches. Higher affinities might indeed be deleterious to the growth of *E. coli*. In view of its great instability, AcnB could provide a particularly sensitive mechanism for stress-induced apo-enzyme formation and mRNA binding/stabilization. It is possible that the observed mRNA-binding affinities are underestimated because the formation of stem–loops that might be needed for optimal apo-Acn binding could have been limited by the use of denatured and rapidly cooled RNA in the gel retardation experiments. It is also relevant that longer 3'UTRs are needed for retardation than for detecting translational enhancement in *vitro*. Thus, the shorter subfragments, A4 and B3, were not retarded (Fig. 2) even though they contain sufficient of the 3'UTR to enhance AcnA and AcnB synthesis from DNA-derived transcripts in transcription–translation tests (Fig. 4). This could mean that the 3'UTRs contain several apo-Acn-binding sites, not all of which are located in the minimal fragments, or that secondary structural elements form more readily in longer RNA molecules.

Attempts to identify potential binding sites for the Acn apo-protein by comparing the sequences and secondary structural features of the two acn 3'UTRs revealed two paired motifs, AA–ACACAUUGC and CAUUUU, that flank putative stem–loops A2 and B2 in each 3'UTR (Fig. 8). Their locations correlate precisely with the minimal requirements for detecting translational enhancement, 169 > A > 71 and 80 > B > 0 bp, for the respective acnA and acnB 3'UTRs (Fig. 4). There is also a sequence related to the first of these motifs downstream of B3 (Fig. 8), which might contribute to a second binding site and simultaneously explain why a longer 3'UTR (279 bp) improves translational enhancement with acnB. If significant, these motifs should be regarded as common features of potentially different AcnA- and AcnB-binding sites because there is no evidence that the two apo-proteins bind to identical sites even though they exhibit overlapping regulatory activities. These sites are clearly different from IREs, where sequence conservation is largely confined to the loops of the stem–loops recognized by IRPs. Searching the *E. coli* genome for pairs of motifs analogous to those flanking the A2 and B2 stem–loops, but with variable hyphenation, revealed no identities. Sequences differing at only one position
exist but there were no preferred locations or functional relationships among the corresponding genes.

The view that the regulatory switch is mediated by the conversion of holo-enzyme to apo-enzyme is supported by the more than threefold and threefold higher affinities of the apo-proteins relative to the native proteins for both acn 3′UTRs (Fig. 7). The real activation thresholds are likely to be considerably higher than threefold because the native (as isolated) enzymes invariably contain some apo-protein and both enzymes are very susceptible to aerobic inactivation and iron–sulphur cluster degradation. This is particularly so for AcnB, which is inactivated (and reactivated) seven (and ten) times faster than AcnA (A. Bradbury & J. R. Guest, unpublished observations). It is known that transcript processing and degradation plays an important part in regulating gene expression in E. coli and that mutations affecting various ribonucleases lower the rate of mRNA decay (Arraiano et al., 1997; Kushner, 1996). The apo-aconitases could thus regulate Acn synthesis by binding to the 3′UTRs of the acn mRNAs to block transcript processing and 3′exoribonuclease progression. Indeed, the half-lives of both acn mRNAs were here shown to increase two- to threefold during MV-induced oxidative stress, which is known to convert aconitase to its apo-form. It is also conceivable that, like IRP, apo-Acn could block or impair translation by binding to the 5′UTRs of specific mRNAs. In this context it interesting to note that oxidative stress lowers the half-life of soxS mRNA from 3.2 to 1.2 min (Ding & Demple, 1997) because this could be mediated by apo-Acn formation.

The overlapping (cross-reacting) feature of the proposed AcnA/B autoregulatory system probably ensures that despite variations in relative amounts of the two acn transcripts (and their products) in different growth phases, Acn synthesis will increase in response to oxidative stress. However, in view of the sixfold higher affinity of apo-AcnB for acn 3′UTR fragments relative to apo-AcnA, it is envisaged that AcnB will play the major role in mounting a particularly sensitive response to oxidative stress in growing cultures by enhancing the synthesis of the stabler AcnA (as well as AcnB) for survival and recovery purposes. The proposed autoregulatory mechanisms also provide a very plausible explanation for previous observations that AcnA and AcnB amplification from pUC-derived plasmids greatly exceeds the amounts expected from copy number alone (i.e. because accumulating apo-Acn stabilizes the acn transcripts).

The present work has revealed a hitherto undetected mechanism for modulating bacterial gene expression at the translational level based on the redox-sensitivity of the aconitases. It offers a rapid initial response to oxidative stress that is activated before the longer-term SoxRS and OxyR transcriptional regulatory mechanisms take effect (Hidalgo & Demple, 1996). It is switched by the interconversion of catalytic and RNA-binding forms of aconitase just like IRP1-mediated post-transcriptional regulation. Indeed, it could be speculated that RNA binding is an inherent property of the aconitases leading to the evolution of the high-affinity IRP1 system of higher organisms, where regulating gene expression from specific relatively stable transcripts is more important than in bacteria. It is also interesting that an iron–sulphur cluster serves as the sensory cofactor as in the SoxR- and FNR-mediated switches. The Acn system has so far been shown to autoregulate aconitase synthesis but it could perform a more general role in mounting the oxidative stress response, and possibly in regulating host–pathogen interactions and bacterial differentiation, where aconitases have been implicated. Future work will be aimed at characterizing the Acn regulatory system and the effects of stress factors (using a semi-synthetic reporter gene), defining the molecular basis of the Acn–mRNA interactions, and identifying other Acn-regulated genes.

ACKNOWLEDGEMENTS

We thank Simon Andrews and Louise Cunningham for supplying bfr- and sucD-specific primers, and Jeff Green, Colin Scott and Guanghui Wu for helpful advice and discussions. The work was funded by a project grant from the Wellcome Trust (J.R.G.).

REFERENCES


Expression in *Escherichia coli* soxRS

Acad Sci USA 93 operated by iron, nitric oxide, and oxidative stress.

translational repressor and an aconitase: a functional assay for *Escherichia coli* activated by superoxide generating agents in


Received 11 June 1999; revised 27 July 1999; accepted 2 August 1999.