# Regulation of the Salmonella typhimurium aroF gene in Escherichia colit

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The Salmonella typhimurium aroF gene, encoding the tyrosine-sensitive 3-deoxy-D-arabino-heptulosonate 7-phosphate (DAHP) synthase, was localized to a chromosomal PstI fragment by Southern blotting with an Escherichia coli aroF probe. This fragment was cloned by screening a plasmid library for complementation of an E. coli aroF mutant. The nucleotide sequence of S. typhimurium aroF was determined and compared with its  $E.$  coli homolog. The nucleotide sequences are  $85.1\%$  identical, and the corresponding amino acid sequences are 96.1% identical. The E. coli genes encoding the three DAHP synthase isoenzymes are evolutionarily more distant from one another than are the homologous aroF genes of E. coli and S. typhimurium. The S. typhimurium aroF regulatory region contains three imperfect palindromes, two upstream of the promoter and one overlapping the promoter, that are nearly identical to operators aroFol, aroFo2, and TyrR box 1 of E. coli. The aroFol and aroFo2 sequences of the two organisms are each separated by three turns of the DNA helix with no sequence similarity. The 5' ends of the *aroF* transcripts for both organisms contain untranslated regions with potential stem-loop structures. Translational fusions of the aroF regulatory regions to lacZ were constructed and then introduced in single copy into the  $E$ . coli chromosome.  $\beta$ -Galactosidase assays for tyrR-mediated regulation of  $arcF$ -lacZ expression revealed that the  $E$ . coli TyrR repressor apparently recognizes the operators of both organisms with about equal efficiency.

In bacteria and plants, aromatic amino acid biosynthesis proceeds by the common aromatic or shikimate pathway, which delivers chorismate to the terminal pathways to generate phenylalanine, tyrosine, and tryptophan (18, 32). In Escherichia coli, carbon flow through the shikimate pathway is controlled by modulation of the first enzyme, the 3 deoxy-D-arabino-heptulosonate 7-phosphate (DAHP) synthase (EC 4.1.2.15) (31). This enzyme catalyzes the condensation of phosphoenolpyruvate and erythrose 4-phosphate to DAHP (41).

In E. coli and Salmonella typhimurium there are three DAHP synthase isoenzymes that can be distinguished by their regulatory properties. The tyrosine-, phenylalanine-, and tryptophan-sensitive isoenzymes are encoded by the unlinked genes aroF, aroG, and aroH, respectively (11, 32). The nucleotide sequences of the E. coli aroF, aroG, and aroH genes have been determined (9, 33, 38). Expression of aroF and  $aroG$  is repressed by the tyrR gene product, the Tyr repressor, complexed to tyrosine or phenylalanine, respectively. Regulatory mutants with lesions linked to aroF were isolated (7, 12). Nucleotide sequence analysis localized the lesions to three operator boxes, designated aroFol and  $arofo2$  (12) and TyrR box 1 (7). These boxes are 20base-pair (bp) imperfect palindromes; aroFol and aroFo2 are of similar sequences, located upstream of the aroF promoter, and are separated by three turns of the DNA helix (12). The TyrR box <sup>1</sup> overlaps the promoter and shows some sequence similarity with  $arofol$  and  $arofol$  (10). The regulatory region of aroG features only one operator box, which is similar to aroFol and aroFo2 (12).

In wild-type  $E.$  coli grown in minimal media, the phenylalanine-sensitive DAHP synthase contributes 80% and the tyrosine-sensitive DAHP synthase contributes 20% of the

total enzyme activity  $(42)$ . However, the *aroF* promoter scores much higher than the *aroG* promoter on a scale of relative promoter strength (12, 15), a fact that may only be reflected in tyrR strains. Thus, repressor binding to the  $arcG$ operator may be less efficient than to the aroF operator. In wild-type S. typhimurium grown in minimal media, the phenylalanine-sensitive DAHP synthase may also be the major isoenzyme (20), although the tyrosine-sensitive isoenzyme has been reported to predominate under these conditions (22). A comparison of the nucleotide sequences for the aroF genes of  $E$ . coli and  $S$ . typhimurium was expected to offer some insight into an explanation for the potentially different expression levels. This report describes the cloning and characterization of aroF from S. typhimurium, the construction of E. coli and S. typhimurium aroF-lacZ translational fusions, and studies on the regulation of S. typhimurium aroF in E. coli.

# MATERIALS AND METHODS

Bacteria, plasmids, and bacteriophages. Bacteria, plasmids, and bacteriophages used in this study are listed in Table 1. E. coli GKM41 (aroF hsdR4) was derived from JM105 (47) in two steps by using phage P1 grown on NK5161  $(tyrA::Tn10)$  (23) and YS482 (aroF), respectively. Strain YS482, isolated by R. L. Somerville, is a one-step UVinduced mutant of wild-type strain W1485. Strain YS482 lacks tyrosine-sensitive DAHP synthase. Strain GKM41 grows on minimal salts medium, but not on such medium supplemented with phenylalanine and tryptophan. Strain SP564, a CSH63 derivative obtained by transduction with phage P1 grown on PLK 1336, has a  $Tn10$  insertion near tyrR but is Tet<sup>r</sup> and TyrR<sup>+</sup>. Strain SP564-1, an SP564 derivative from which the TnJO has been excised by the method of Bochner et al.  $(4)$ , is Tet<sup>s</sup> and TyrR<sup>-</sup>. Bacteria were grown in either Luria broth,  $2 \times$  YT (29), or minimal salts medium supplemented with vitamin  $B_1$  and biotin (45). Cells were plated on nutrient agar or M9 agar supplemented with

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Designation	Relevant genotype and genes	Source or reference	
S. typhimurium			
LT <sub>2</sub>	Wild type	R. L. Somerville	
E. coli			
<b>YS482</b>	aroF	R. L. Somerville	
<b>JM101</b>		28	
<b>JM105</b>	hsdR4	47	
D <sub>1210</sub>	lacI <sup>q</sup>	24	
<b>NK5161</b>	tryrA::Tn10	N. Kleckner via R. L. Somerville	
<b>GKM41</b>	$arcF$ hsdR4	This study	
<b>MC4100</b>	$\Delta(\text{argF-lac})U169$	40	
CSH <sub>63</sub>	$\Delta (lac-pro)$	29	
<b>PLK1336</b>	zci-223:: $\text{Tr}10$ near min 28	P. L. Kuempel via R. L. Somerville	
SP <sub>564</sub>	$\Delta (lac-pro)$ zci-233::Tn10	R. L. Somerville	
SP564-1	$\Delta (lac$ -pro) tyrR	R. L. Somerville	
<b>Plasmids</b>			
pKGW	E. coli endonuclease	24	
pMLB1034	Promoterless lacZ	40	
DCG201	E. coli aroF	12	
pGM59	S. typhimurium aroF	This study	
<b>Bacteriophages</b>			
M13mp18, M13mp19	lacα	28	
M13GME	E. coli aroF regulatory region	This study	
M13GMS	S. typhimurium aroF regulatory region	This study	
<b>ARZ5</b>	Promoterless 'lacZ,' bla	R. Zagursky via H. Zalkin	
λGME4	E. coli aroF-lacZ fusion	This study	
λGMS4	S. typhimurium aroF-lacZ fusion	This study	

TABLE 1. Bacterial strains, plasmids, and phages used in this study

vitamin  $B_1$  and biotin (29) and, where indicated, with 20  $\mu$ g of amino acids per ml, 50  $\mu$ g of ampicillin per ml, 20  $\mu$ g of kanamycin per ml, and  $15 \mu g$  of tetracycline per ml. M13 phages were plated in Luria broth top agar containing 40  $\mu$ g of 5-bromo-4-chloro-3-indolyl-p-D-galactopyranoside (X-Gal) per ml and 28  $\mu$ g of isopropyl- $\beta$ -D-thiogalactopyranoside (IPTG) per ml. A phages were plated in tryptone top agar on minimal plates supplemented with 40  $\mu$ g of X-Gal per ml.

Chemicals. Phosphoenolpyruvate (5) and erythrose 4 phosphate (39) were synthesized and assayed as described previously. Kanamycin, chloramphenicol, tetracycline, tyrosine, phenylalanine, and tryptophan were from Sigma Chemical Co.; nutrient agar was from Difco Laboratories; agarose and IPTG were from Bethesda Research Laboratories, Inc.; X-Gal, dideoxynucleotides, and 7-deaza-dGTP were from Boehringer Mannheim Biochemicals;  $[\alpha^{-32}P]$ dCTP was from ICN Radiochemicals or Amersham Corp.; low-melting-point agarose was from FMC Corp., Marine Colloids Div.; redistilled phenol and 0.2-mm wedge-shaped sequencing gel spacers were from International Biochemical, Inc.; deoxynucleotides were from Pharmacia, Inc.; acrylamide, hydroxylapatite-HTP, and protein molecular weight markers were from Bio-Rad Laboratories; and formamide and propanediol were from Eastman Kodak Co. Oligonucleotides were synthesized by the Purdue Laboratory for Macromolecular Structure.

Enzymes. Restriction enzymes and T4 DNA ligase were from Bethesda Research Laboratories or New England BioLabs, Inc.; restriction enzyme and ligase buffers and exonuclease III were from Bethesda Research Laboratories; Klenow fragment of DNA polymerase <sup>I</sup> was from Boehringer Mannheim Biochemicals or New England BioLabs; Klenow sequencing kit, T4 polynucleotide kinase, and mung bean nuclease were from New England BioLabs; Sequenase (T7 DNA polymerase) and Sequenase sequencing kit were from U.S. Biochemicals Corp. DAHP synthase was assayed as described previously (37).

DNA manipulations. Chromosomal DNA was isolated by the method of Saito and Miura (34); agarose gel electrophoresis, ligations, plasmid and phage miniscreens, electroelutions, and transformations were as described previously (1).

DNA probes were labeled by nick translation with  $\int \alpha^{-3}$ dCTP. Genome blotting to dried agarose gels (44) was performed under optimal hybridization conditions (2, 27). S. typhimurium DNA, digested with restriction endonuclease PstI and size fractionated, was ligated into pKGW (24). The resulting recombinant plasmids were grown in E. coli D1210 on M9 medium containing kanamycin and IPTG. Amplification and preparation by the method of Birnboim (3) yielded the recombinant plasmid pGM59.

Site-directed mutagenesis with an Amersham kit was performed by the method of Sayers et al. (36). Mutants were screened by SmaI digestion, and the mutations were confirmed by nucleotide sequence analysis.

Nucleotide sequence analysis. For nucleotide sequence analysis by the dideoxy method (35), pGM59 fragments were cloned into phages M13mpl8 and M13mpl9. One subclone was generated by ligating the 900-bp BssHII fragment previously filled in with DNA polymerase <sup>I</sup> (Klenow fragment) (1) into the SmaI site of M13mpl8. The resulting clone had the same orientation as lacZ. This recombinant was used for exonuclease III digestion (16). Both replicative-form and single-stranded phages were prepared, and the extent of the deletions was determined by agarose gel electrophoresis and DNA sequencing. Two plasmids with deletions of <sup>150</sup> and 350 bp were used to complete the nucleotide sequence analysis of the BssHII fragment.

 $arcF$ -lacZ fusions. The  $arcF$  regulatory regions of E. coli and S. typhimurium were subcloned into pMLB1034. The resulting recombinant plasmids were used to transform E.



FIG. 1. Restriction endonuclease cleavage maps for the E. coli and S. typhimurium aro F genes and cloning scheme for S. typhimurium aroF.

 $\text{coll}$  MC4100, yielding Lac<sup>+</sup> derivatives that served as hosts to prepare lysates of  $\lambda$ RZ5 (14) by the method of Silhavy et al. (40). Recombinant phages carrying functional  $lacZ$  genes were identified as blue plaques on X-Gal-agar. Six lysogens from each  $\lambda$ RZ5 derivative carrying aroF-lacZ fusions were purified and shown to be single lysogens  $(40)$ . The  $\beta$ galactosidase activities in extracts of the lysogens were measured and are given in Miller units (29).

Purification of S. typhimurium tyrosine-sensitive DAHP synthase. E. coli GKM41 carrying plasmid pGM59 was grown to late log phase in minimal salts medium containing ampicillin. The cells were harvested by centrifugation and resuspended in <sup>50</sup> mM potassium phosphate (pH 6.5) containing 2% propanediol and <sup>2</sup> mM phosphoenolpyruvate (buffer A). The cells were disrupted in an Aminco French pressure cell at  $20,000$  lb/in<sup>2</sup>; cell debris was removed by centrifugation for 60 min at 25,000  $\times$  g. The supernatant was treated with 0.1 volume of 2% protamine sulfate, and the precipitate was removed by centrifugation for 75 min at 30,000  $\times$  g. The supernatant was applied to a BioGel hydroxylapatite-HTP column equilibrated with buffer A. Protein was eluted from the column with a 500-ml linear gradient of 0.05 to 0.5 M potassium phosphate containing 2% propanediol and <sup>2</sup> mM phosphoenolpyruvate.

Protein was quantitated by the procedure of Lowry et al.

(26); gel electrophoresis was performed on a sodium dodecyl sulfate-10% polyacrylamide gel by the method of Laemmli (25); the proteins in the gel were detected by silver staining (46). Amino acid sequencing was carried out on an Applied Biosystems model 470A gas-phase sequenator with a model 120A analyzer (17).

### **RESULTS**

Genome blotting. S. typhimurium LT2 chromosomal DNA was digested separately with five restriction endonucleases. The digestion products were subjected to agarose gel electrophoresis. The dried gel was probed with E. coli DNA encoding tyrosine-sensitive DAHP synthase. Plasmid  $pCG201$  (38) carries the E. coli aroF gene (Fig. 1) totally contained within two DdeI fragments of 714 and 796 bp. The autoradiogram of a gel probed with the 796-bp DdeI fragment is shown in Fig. 2A. The 714-bp DdeI probe gave an identical pattern. PstI digestion yielded a 5.5-kilobase (kb) fragment that hybridized with both the 796- and 714-bp DdeI probes and was predicted to contain the entire aroF gene of S. typhimurium.

PstI-digested S. typhimurium LT2 DNA was subjected to gel electrophoresis with low-melting-point agarose. Fractions containing DNA fragments of different sizes were isolated by phenol extraction of gel slices melted at 65°C. A sample of each fraction was subjected to agarose gel electrophoresis and probed as before with the 796-bp DdeI fragment (Fig. 2B). DNA of fraction <sup>3</sup> was used in the cloning experiments.

Cloning of S. typhimurium aroF. The positive selection vector pKGW (24) was chosen to clone S. typhimurium aroF. This plasmid carries the gene encoding the restriction endonuclease EcoRI under the control of the lacUV5 promoter (Fig. 1). This high-copy-number plasmid is lethal to wild-type cells, but can be propagated in  $lacI<sup>q</sup>$  cells grown in the absence of inducer. Upon induction of the lacUV5 promoter with IPTG, overproduction of EcoRI becomes lethal to the host cells, unless the coding sequence of  $EcoRI$ is interrupted. The PstI or  $BgIII$  sites within the  $EcoRI$ coding sequence are suitable cloning sites.

Size-fractionated PstI-digested S. typhimurium DNA was cloned in plasmid pKGW. Transformants of E. coli D1210  $(lacI<sup>q</sup>)$  were selected in the presence of IPTG and kanamycin. A library of recombinants was generated, amplified, and screened by transformation of the aroF strain GKM41.



FIG. 2. Genome blotting of S. typhimurium DNA. (A) S. typhimurium DNA was digested with the indicated restriction endonucleases, and the digests were subjected to agarose gel electrophoresis. The dried gel was hybridized with the 796-bp DdeI probe from E. coli (Fig. 1). (B) PstI-digested S. typhimurium DNA was size fractionated by agarose gel electrophoresis; samples of fractionated DNA fragments were again subjected to agarose gel electrophoresis and treated as in panel A.



FIG. 3. Sequencing strategy for S. typhimurium aroF. The box indicates the coding region of aroF. Arrows mark the extent and direction of nucleotide sequence runs. The two clones of the exonuclease III-digested BssHII fragments are marked with asterisks.

Selection was made for growth on minimal salts medium supplemented with phenylalanine and tryptophan. An aroFcontaining plasmid designated pGM59 (Fig. 1) was isolated and shown to contain aroF by hybridization and DAHP synthase assay of cell extracts from plasmid-bearing strains.

Nucleotide sequence analysis of S. typhimurium aroF. The 5.5-kb PstI insert of plasmid pGM59 was subjected to restriction analysis (Fig. 1). By hybridization analysis, aroF was localized within the 5.5-kb PstI fragment to a 2-kb EcoRV-HincII fragment. Figure 3 shows the strategy that

A T C<br>ATGCAAAAAGACGCGCTGAATAACGTACOTATLACCGATGAAC TGCAAAAAGACGCGCTGAATAACGTACGTATCACCGATGA T T A A UAGGTATTAATGACGCOGGAGCAG Q V L M T P E Q 22 EU<br>GAAA CAACAA T TG C G TAA GAARAA T TG C G TAA GAARAA T TG C G TAA GAARAA TAACAA TAACAA TAACAA TAACAA TAACAA TAA CTTAAAGCGGCCTTTCCCTTGAGCCTGGCGCAGGAAGCCCAGATAGCGCAGTCCCGGG<u>GAAT</u>CATT L K A A F P L 8 L Q Q B A Q L A Q E A G R G L L 44  $A \cdot T$  C  $C \cdot T$  GC  $A \cdot T$  C  $T$ TCTGACATTATTGCCGGGCGCGATCCG1GTCCGTGT TCTTTCCACGAT <sup>S</sup> <sup>D</sup> <sup>I</sup> <sup>I</sup> A G <sup>R</sup> <sup>D</sup> <sup>P</sup> <sup>R</sup> <sup>L</sup> <sup>L</sup> <sup>V</sup> <sup>V</sup> <sup>C</sup> G <sup>P</sup> <sup>C</sup> <sup>S</sup>I <sup>H</sup> <sup>D</sup> <sup>66</sup>  $\mathbf{G}$  ,  $\mathbf{T}$  , and the set of  $\mathbf{T}$ CCTGAAACCGCTCTGGAATATGCCCGTCGATTTAAAGCCCTTGCCGCAGAGGTCAGCGATAGCCTC <sup>P</sup> I <sup>T</sup> <sup>A</sup> <sup>L</sup> <sup>E</sup> <sup>Y</sup> <sup>A</sup> <sup>R</sup> <sup>R</sup> FK <sup>A</sup> <sup>L</sup> <sup>A</sup> <sup>A</sup> <sup>Z</sup> <sup>V</sup> S <sup>D</sup> <sup>S</sup> <sup>L</sup> <sup>88</sup>  $A$  C T C T  $\ldots$  T  $A$ TATCTGGTAATGCGCGTCTATTTTGAAAAGCCGCGAACTACCGTCGGCTGGAAAGGGCTGATTAAC Y L V N R V Y F Z K P R T T V G W K G L <sup>I</sup> N 110  $C$  T  $T$   $A$   $C$  C G  $C$   $A$   $A$  G  $C$  T GATCCTCACATGGATGGCT CA TAGAT CTG <sup>D</sup> <sup>P</sup> ND <sup>G</sup> SF DVF AG LQI AGR <sup>L</sup> <sup>L</sup> <sup>t</sup> <sup>132</sup> G AC C G A T GAACTGGTGAATATGGGGTTGCCATTGGCGACCGAATCCTTGGATCCGAACAGCCCGCAATACCTG AACTGGTGAATATGGGGTTGCATTGGGAACGGAACGGTTGGATCGGAACAGCCGCAATACCTG<br>E L V N M G L P L A T E A L D P N S P Q Y L 154 A A T T T T T G T T T T GGCGATCTGTTTAGCTGGTCGGCGATAGGGGCGCGCACAACCGAAACCCACCGCGAAATG G D L F S W 176 A I G A R T T E S Q T H R E M 176 C C G <sup>C</sup> <sup>T</sup> GT C C <sup>T</sup> <sup>A</sup> <sup>A</sup> COTCOOL CONTROLS COOLSTER AND CORPORATE CONSIDERATION OF SALT A TAST 198  $T$  $\frac{1}{2}$   $\frac{1}{2}$  TATTGCAAACCCAGGGAAATCCGCAGGGAAGTATCTCGGGGGGAAAGCGCGAACTAT<br>TATTGCAAACCCAGGAAATCGGGAAGGCCATGAATCTCGGGGGGGAAAAGCGCGAACTAT <sup>T</sup> <sup>O</sup> <sup>T</sup> <sup>G</sup> <sup>A</sup> <sup>G</sup> <sup>C</sup> <sup>G</sup> <sup>T</sup> AGCCCGGCAGATOTCGCTCAGTGTGAAAAAGAGATGGACAGGCGGOACTAC1TCCTTCGCTGATG <sup>S</sup> <sup>P</sup> <sup>A</sup> <sup>D</sup> <sup>V</sup> <sup>A</sup> <sup>Q</sup> <sup>C</sup> <sup>I</sup> <sup>K</sup> <sup>Z</sup> <sup>N</sup> <sup>Z</sup> <sup>Q</sup> <sup>A</sup> <sup>0</sup> <sup>L</sup> <sup>R</sup> <sup>P</sup> <sup>8</sup> <sup>L</sup> <sup>N</sup> <sup>264</sup> <sup>C</sup> <sup>C</sup> <sup>T</sup> <sup>T</sup> <sup>T</sup> <sup>O</sup><sup>C</sup> <sup>A</sup> <sup>C</sup> GTAGATGCAGTCATOGTAACTCCAATAAAGATTATGCCCGCCAGCCAGCCOTTGCcGAATCTGTG <sup>V</sup> <sup>D</sup> <sup>C</sup> <sup>S</sup> <sup>H</sup> <sup>0</sup> <sup>N</sup> <sup>S</sup> <sup>N</sup> <sup>K</sup> <sup>D</sup> <sup>Y</sup> <sup>R</sup> <sup>R</sup> <sup>Q</sup> <sup>P</sup> <sup>A</sup> <sup>V</sup> <sup>A</sup> <sup>Z</sup> <sup>S</sup> <sup>V</sup> <sup>286</sup>  $T$  A C  $C$   $T$   $T$   $C$  G  $C$   $C$   $C$ GTTGCGCAGATTAAAGATGGCAATCGTTCAATCATTIGCTTAATGATTGAAAGTAATATTCATGAG <sup>V</sup> <sup>A</sup> <sup>Q</sup> <sup>I</sup> <sup>K</sup> <sup>D</sup> <sup>G</sup> <sup>N</sup> <sup>R</sup> <sup>S</sup> <sup>I</sup> <sup>I</sup> <sup>0</sup> <sup>L</sup> <sup>N</sup> <sup>I</sup> <sup>1</sup> <sup>8</sup> <sup>N</sup> <sup>I</sup> <sup>H</sup> <sup>1</sup> <sup>308</sup>  $C$ ,  $T$ ,  $T$ ,  $C$ ,  $T$ ,  $A$ ,  $C$ ,  $T$ ,  $A$ ,  $C$ ,  $C$ ,  $C$ GGTAACCAGTCTTCCGAACAGCCGCGCAGCGAAATGAAGTATGGCOTTCCGTCACCGATGCTTCT <sup>G</sup> <sup>N</sup> <sup>Q</sup> <sup>S</sup> <sup>S</sup> <sup>E</sup> <sup>Q</sup> <sup>P</sup> <sup>R</sup> <sup>S</sup> <sup>Z</sup> <sup>M</sup> <sup>K</sup> <sup>Y</sup> <sup>0</sup> <sup>V</sup> <sup>S</sup> <sup>V</sup> <sup>T</sup> <sup>D</sup> <sup>A</sup> <sup>C</sup> <sup>330</sup> A T C G C C C A G A ATTAGCTGGGAGATGACGATGCCCTGTTACGTGAAATTCATAAGATTTGAGGGGAGCTGGC EXPRESS DESCRIPTION OF THE RESERVE OF THE RESERVE<br>I SO THE RESERVE OF THE RESERVE OF<br>TH CT<br>GTGCGCGTCGCATAA<br>|V R V A 356 A

FIG. 4. Nucleotide sequence of the S. typhimurium aroF (Gen-Bank no. M31302) and amino acid sequence of the encoded tyrosinesensitive DAHP synthase (middle two lines). The sequence variations to the  $E.$  coli aro $F$  coding region (upper line) and the  $E.$  coli DAHP synthase (lower line) are also shown.

was used to obtain the nucleotide sequence of the 2-kb EcoRV-HincII fragment. The nucleotide sequence reveals a 1,068-bp open reading frame that is similar in sequence to  $arcF$  of  $E.$  coli (Fig. 4).

Purification of S. typhimurium tyrosine-sensitive DAHP synthase and amino acid sequence analysis. The tyrosinesensitive DAHP synthase was purified from extracts of E. coli GKM41 (aroF) carrying the S. typhimurium aroF plasmid pGM59. A two-step procedure yielded enzyme of near electrophoretic homogeneity (Fig. 5). The protein was subjected to N-terminal amino acid sequence analysis. The N-terminal 14 amino acid residues are identical to those predicted from the nucleotide sequence of the 1,068-bp open reading frame. Thus, we have cloned and sequenced the coding region of the  $S$ . typhimurium aro $F$  gene.

Regulation of S. typhimurium aroF in E. coli. To initiate a functional analysis of the S. typhimurium aroF regulatory region, we constructed two very similar lacZ fusions to the homologous regions of the E. coli and the S. typhimurium aroF genes; the fusions contain identical coding regions. Site-directed mutagenesis was used to generate two SmaI sites covering codons 4 and <sup>5</sup> of the two genes (Fig. 6). Two M13 constructs served as templates for the mutagenesis: M13GME contains the 674-bp *EcoRV-NdeI E. coli* fragment (Fig. 1) that encompasses 334 bp upstream of the E. coli aroF transcription start. M13GMS contains the 434-bp EcoRV-SmaI S. typhimurium fragment (Fig. 1) that covers 313 bp upstream of the predicted S. typhimurium aroF transcription start. Both fragments were inserted into the SmaI site of M13mpl8. After mutagenesis, new SmaI sites were detected by restriction analysis and confirmed by nucleotide sequence analysis. The EcoRI-SmaI fragment that contains 18 bp of the M13 polylinker <sup>5</sup>' to the new E. coli EcoRV-SmaI fragment of 397 bp and the EcoRI-SmaI fragment that contains the same polylinker fragment <sup>5</sup>' to the new S. typhimurium EcoRV-SmaI fragment of 376 bp were subcloned into the EcoRI and SmaI sites of the polylinker of plasmid pMLB1034, which carries a promoterless lacZ gene, yielding aroF-lacZ translational fusions. In the two recombinant pMLB1034 derivatives, codons 4 of the E. coli and the S. typhimurium aroF were fused to codon 6 of lacZ. The inserts of the recombinant plasmids were confirmed by restriction and nucleotide sequence analyses.



FIG. 5. Purification of S. typhimurium tyrosine-sensitive DAHP synthase. Protein samples were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis, and proteins were detected by silver staining. Lanes: 1, molecular weight marker; 2, 1.4  $\mu$ g of protein after treatment of the extract with protamine sulfate; 3, 2.8  $\mu$ g of protein after ammonium sulfate precipitation; 4, 0.6  $\mu$ g of protein from the hydroxylapatite pool.



GCCATC ATG CAA AAA CCC GGG GAT CCC GTC codon <sup>#</sup>9 of *lacZ* MET GLN LYS PRO|ALA ASP PRO VAL



FIG. 6. Construction of aroF-lacZ fusions. The sequences around the site of mutagenesis are identical for E. coli and S. typhimurium. Through three single-base-pair changes a new SmaI site is obtained overlapping  $aroF$  codons 4 and 5.

Phage  $\lambda$ RZ5 was used to place the *aroF-lacZ* fusions in single copy at the  $\lambda$  *att* site on the *E*. *coli* chromosome (14). 3-Galactosidase activities in the resulting lysogens grown on media with and without tyrosine were determined (Table 2). Cells of  $tyrR<sup>+</sup>$  strains grown in minimal-salts tyrosine medium show very similar, low enzyme values, indicating that the E. coli Tyr repressor can regulate the expression of both the E. coli and the S. typhimurium aro F gene. Since the aro  $F$ genes differ markedly for the nucleotide sequence in the three helix turns that separate  $arcFol$  and  $arcFol$  (12), our results also suggest that the sequence in this region is not

TABLE 2. Regulation of S. typhimurium aroF in E. coli

		<b>B-Galactosidase activity</b>	
Lysogen	200 μM tyrosine supplement	Miller units	Relative activity
SP564(λGME4)	+	$0.32 \pm 0.02$	
SP564(λGME4)		$2.3 \pm 0.1$	
SP564-1(λGME4)		$25.3 \pm 1.3$	80
SP564-1(λGME4)		$26.0 \pm 2.1$	81
SP564(λGMS4)		$0.3 \pm 0.09$	
SP564(λGMS4)		$4.1 \pm 0.1$	13
SP564-1(λGMS4)		$49.4 \pm 3.0$	155
SP564-1(λGMS4)		$46.2 \pm 4.4$	145

critical for repressor function. Under conditions of derepression,  $\beta$ -galactosidase activity from the S. typhimurium aro  $F$ lacZ fusion is about twice as high as the activity in the corresponding strain carrying the E. coli fusion (Table 2).

#### DISCUSSION

S. typhimurium chromosomal DNA contains <sup>a</sup> 5.5-kb PstI fragment that strongly hybridizes with E. coli aroF probes. The 5.5-kb PstI fragment was cloned into the positive selection vector pKGW. The recombinant plasmid pGM59 complements the E. coli aroF mutant strain GKM41. A restriction map of the 5.5-kb insert of pGM59 was obtained, and the position of aroF on a 2-kb fragment within the insert was determined by hybridization analysis.

The nucleotide sequence of this 2-kb fragment contains a 1,068-bp open reading frame with extensive sequence similarity to the E. coli aroF gene (38). The two DNA sequences are 85.1% identical. Several other genes have been sequenced from both  $S$ . typhimurium and  $E$ . coli, most notably the genes of the trp operon (8, 19, 30, 48). The trp operon genes showed between 75 and 84% sequence identity. The degree of relatedness of the two *aroF* genes is within the range at the upper end of the identity scale, perhaps reflecting the highly evolved role of DAHP synthase as <sup>a</sup> control enzyme for the common aromatic amino acid biosynthetic pathway (31).

The 85.1% identity between the two  $arcF$  genes contrasts with about 50% identity between E. coli aroF and the other two E. coli genes encoding DAHP synthases (9, 33, 38). The three E. coli genes  $arcF$ ,  $arcG$ , and  $arcH$  are evolutionarily more distant than the homologous S. typhimurium and E. coli aroF genes, suggesting that the three DAHP synthase isoenzymes existed before the genera Escherichia and Salmonella diverged.

The S. typhimurium aro F gene encodes a protein of 356 amino acid residues. S. typhimurium tyrosine-sensitive DAHP synthase was purified, and the amino acid sequence predicted by the nucleotide sequence was confirmed through N-terminal sequence analysis. The amino acid sequences of the tyrosine-sensitive DAHP synthase from S. typhimurium and E. coli are 96.1% identical (Fig. 4). There are only 14 amino acid residue differences between the two proteins. Eleven of these differences are localized to three small regions of the protein, residues 32 to 43, 125 to 132, and 345 to 353. Within these regions the sequence similarity is only 55.6, 66.3, and 66.7%, respectively. Apparently, none of these differences have a marked influence on the kinetic or stability properties of the enzyme. The apparent  $V_{\text{max}}$  values for both enzymes are very similar, and both enzymes are subject to rapid inactivation when cells approach the stationary phase of growth (G. K. Muday and K. M. Herrmann, unpublished data), a phenomenon previously described for the E. coli enzyme (43).

The codon usage of the two  $arof$  genes is relatively close to the overall codon usage of E. coli, as well as to each other. It is, however, quite interesting that in both *aroF* genes rare codons are used for the aromatic amino acids tyrosine and phenylalanine. In E. coli, 40% of all tyrosine codons are TAT; for the  $aroF$  genes, five of seven  $E$ .  $coli$  and six of seven S. typhimurium codons are TAT. Likewise in E. coli, 37% of all phenylalanine codons are TTT; for the aroF genes of both organisms, seven of seven codons are TTT. Also, several other rare codons are used slightly more frequently in the S. typhimurium gene, for example, TTA and TTG for



FIG. 7. Comparison of aroF regulatory regions of E. coli and S. typhimurium. The complete E. coli sequence and the S. typhimurium variations are given. Brackets designate operator boxes aroFol, aroFo2, TyrR box 1, and a palindromic sequence of unknown significance. Identified operator mutations (7, 12) are indicated by the arrows above the sequences.

leucine, ATA for isoleucine, and CGA and CGG for arginine.

A more quantitative analysis showed <sup>a</sup> correlation of optimal codon usage to cellular levels of the protein products (21). When this analysis is applied to the two *aroF* genes, the frequencies are 0.61 and 0.64 for S. typhimurium and E. coli, respectively. The E. coli aroF value is close to 0.63, 0.65, and 0.66 for the E. coli trpA, trpD, and trpE genes, respectively. The frequencies of optimal codon usage for S. typhimurium trpA, trpB, and trpD are all  $0.63$ , slightly lower than the  $E$ . coli trp gene values, just as the  $S$ . typhimurium aro $F$ gene has a lower value than the E. coli aroF gene. This result is in good agreement with a few other examples (13) that show higher frequencies of optimal codon usage in E. coli, suggestive of a slightly tighter codon preference.

The regulatory regions of the two  $arcF$  genes were also compared. For the E. coli gene, the start of transcription has been determined (12), a strong promoter has been identified (12), and three operator boxes have been defined by mutational analysis  $(7, 12)$ . The aroF regulatory region of S. typhimurium shows a high degree of sequence identity to the corresponding E. coli sequences of defined regulatory function, but very little similarity in the sequences outside the defined regions (Fig. 7). In all  $E$ . coli aro $F$  operator mutations identified thus far, the changes are in nucleotides that are identical for both wild-type organisms (Fig. 7). The S. typhimurium promoter was assigned on the basis of its similarity in sequence and location to the E. coli promoter. The promoters are very similar and very strong, with Mc-Clure algorithm scores of 68.6 and 69.2 for the S. typhimu*rium* and the  $E$ . *coli* promoter, respectively  $(15)$ .

Three operator boxes were identified within the S. typhimurium regulatory region, based on their similarity to the E. coli boxes. The sequence and location of the boxes and the distance between them are very similar for both genes (Fig. 7). The distance between aroFol and aroFo2 corresponds to three turns of the helix. Although the distance between these operator boxes is preserved, the nucleotide sequence in the region between the boxes shows much less similarity, indicating that the distance, if not the sequences, between the boxes may be important for proper operator recognition by the tyrR product.

The high degree of similarity between the regulatory regions of the two  $aroF$  genes suggests that the levels of transcription and translation of these two genes should be very similar. To compare the control features for both aroF genes, we fused their regulatory regions to lacZ and measured the B-galactosidase levels in strains carrying the translational fusions in single copy. Under derepression conditions, expression of the S. typhimurium promoter is about twice that of the E. coli promoter. This difference could account in part for the higher levels of the tyrosine-sensitive DAHP synthase reported for wild-type S. typhimurium (22). Under repression conditions, the enzyme values were about the same. These results indicate that the E. coli Tyr repressor recognizes both E. coli and S. typhimurium aroF operators with about equal efficiency. Since the sequences that separate *aroFol* and *aroFo2* in the two organisms show no similarity, one may assume that the sequences within these three turns of the DNA helix cannot be part of the recognition signal for repressor binding. A preliminary study with <sup>a</sup> multicopy plasmid system suggested that the distance between E. coli aroFol and aroFo2 is also not critical for repressor recognition (6). However, the separation of the two operator boxes by three turns of a helix for both organisms suggests an important role for this portion of the regulatory region. The regulatory region of the E. coli aroL gene, which is also controlled by the tyrosine repressor, also has three operator boxes, arranged in the same way as the aroF boxes (10). The distance between the two upstream aroL boxes is again three turns of the helix, yet the nucleotide sequence of the spacer shows no similarity to the corresponding aroF E. coli or S. typhimurium sequences.

A new potential regulatory region is also seen when the  $E$ . coli and S. typhimurium aroF gene sequences are compared. In the untranslated portion of the transcript for both regulatory regions, a palindromic sequence is found with a potential for stem-loop formation. The regulatory significance of these palindromes will be addressed through site-directed mutagenesis.

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