Role of the *Escherichia coli* FadR Regulator in Stasis Survival and Growth Phase-Dependent Expression of the *uspA*, *fad*, and *fab* Genes

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The increased expression of the uspA gene of $Escherichia\ coli$ is an essential part of the cell's response to growth arrest. We demonstrate that stationary-phase activation of the uspA promoter is in part dependent on growth phase-dependent inactivation or repression of the FadR regulator. Transcription of uspA is derepressed during exponential growth in fadR null mutants or by including the fatty acid oleate in the growth medium of FadR⁺ cells. The results of DNA footprinting analysis show that FadR binds downstream of the uspA promoter in the noncoding region. Thus, uspA is a member of the fadR regulon. All the fad-lacZ fusions examined (fadBA, fadL), and fadD) are increasingly expressed in stationary phase with kinetics similar to that of the increased expression of uspA. In contrast, β -galactosidase levels decrease during stationary phase in a fabA-lacZ lysogen, consistent with the role of FadR as an activator of fabA. The growth phase-dependent increased and decreased transcription of fad genes and fabA, respectively, is dependent on the status of the fadR gene. Cells carrying a mutation in the FadR gene (fadRS219N) that makes it nonderepressible exhibit a weak stationary-phase induction of uspA and fad genes. In addition, cells carrying fadRS219N survive long-term stasis poorly, indicating that FadR-dependent alterations in fatty acid metabolism are an integral and important part of the adaptation to stationary phase.

The universal stress protein, UspA, is a member of all starvation and stress stimulons so far studied in Escherichia coli except the cold shock response (18, 20). UspA is a cytoplasmic serine/threonine phosphoprotein that is phosphorylated during conditions that induce its synthesis (16). Mutant cells devoid of UspA are impaired in their ability to survive complete and prolonged growth inhibition caused by a variety of starvation and stress conditions, indicating that UspA has a general protective function related to the growth-arrested state (15, 20). In addition, mutations in uspA result in an abnormal excretion of acetate during growth of cells on glucose or gluconate, a feature leading to a diauxic growth pattern of uspA mutant cultures (19). Moreover, uspA mutations affect the timing of protein expression during prolonged periods of growth arrest, accelerating the changes in protein expression that normally occur (20). While the lack of UspA renders the cells less resistant to stasis, constitutively elevated levels of the protein significantly reduce the ability to recover from growth arrest (21).

The *E. coli fadR* gene product functions as a repressor of many unlinked genes and operons encoding proteins involved in long-chain fatty acid transport, activation, and β -oxidation. FadR binds specific operator sites upstream of the fatty acid degradative (*fad*) gene coding sequences to turn off transcription (5). However, FadR can also act as an activator of at least the *fabA* and *fabB* genes which are required for fatty acid biosynthesis (2, 4, 13). Derepression of *fad* genes occurs upon

growth of $E.\ coli$ in medium containing long-chain fatty acids (C_{14} to C_{18}). Such growth conditions result in a decrease in the expression of fatty acid biosynthetic (fab) genes (2). Long-chain acyl coenzyme A (acyl-CoA) thioesters, generated from exogenous fatty acids, are the effector molecules that regulate fatty acid metabolism at the level of FadR-dependent transcription by binding directly to FadR (24). The FadR-acyl-CoA thioester complex is unable to bind the operator sequence, resulting in transcription of the fad genes and the loss of activation of fabA transcription (5–7).

In this study, we demonstrate that stationary-phase induction of uspA is dependent on the status of the fadR gene and that the FadR regulon and uspA are induced concomitantly in stationary phase. Moreover, cells that carry a dominant non-derepressible fadR allele are impaired in their ability to survive long-term stasis. We discuss the possible physiological assignment of the regulon during growth arrest.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The $E.\ coli$ strains used in this work are listed in Table 1. Cultures were grown aerobically in liquid Luria-Bertani (LB) or M9 (26) medium in Erlenmeyer flasks in a rotary shaker. When required, the media were supplemented with oleate (0.5 mM), glucose (0.4%), glycerol (0.4%), acetate (0.4%), or L-serine (0.4%), and thiamine (10 mM). When appropriate, the media were supplemented with kanamycin (50 μ g/ml), carbenicillin (50 μ g/ml), tetracycline (20 μ g/ml), chloramphenicol (30 μ g/ml), and/or streptomycin (200 μ g/ml).

General methods. Plasmid DNA was purified by using Qiagen columns (Qiagen, Inc.) according to the protocol provided by the manufacturer. P1 transductions and plasmid transformations were performed as described previously by Miller (14) and Sambrook et al. (26). Culture samples were processed to produce extracts for resolution on two-dimensional polyacrylamide gels by the method of O'Farrell (22) with modifications (28).

Construction of λ-uspA-lacZ lysogens and uspA::lacZ-Km^r insertional mutants. A uspA::lacZ-Km^r fusion was constructed by inserting the SaII fragment of plasmid pKOK5 (10) containing lacZ-Km^r into the SaII site of uspA carried on plasmid pTN6093 (19) (Fig. 1A). A fragment containing the uspA::lacZ-Km^r fusion was integrated into the chromosome of the E. coli recD K4633 by linear

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TABLE 1. Bacterial strains and plasmids

Strain or	Relevant genotype or	Source or	
plasmid	characteristic	reference	
E. coli strains			
MC4100	F^- araD139 Δ (argF-lac)U169	M. Giskov	
11101100	rpsL 150 relA1 flbB5301	W. Gibkov	
	deoC1 ptsF25 rbsR		
TN4100	MC4100 uspA::lacZ-Km ^r	This work	
TN4111	TN4100 Δcya-851	This work	
TN4112	TN4100 Δcrp-96 zhd-732::Tn10	This work	
TN4131	TN4100 fadR::Tn10	This work	
TN4132	TN4131/pCD126	This work	
TN4133	TN4100/pRW22	This work	
TN4134	TN4100/pACYC177	This work	
TN4135	TN4100 ΔarcA::Tn10	This work	
TN4136	TN4100 Δhns-206::bla	This work This work	
TN4137	TN4100 dam-13::Tn9		
TN4138 TN4139	TN4100 fis-767 TN4100 soxS::Tn10	This work This work	
TN4140	TN4100 5025.: THT0 TN4100 fur::Tn10	This work This work	
TN4140	TN4131/F' proAB lacIq	This work	
211111	$\Delta(lacZ)$ M15	Tino worn	
TN4142	TN4141/pTN223	This work	
AF634	MC4100 $\lambda\Phi(uspA-lacZ)$	This work	
AF637	AF634 himA::kan	This work	
AF638	AF634 relA251 spoT207	This work	
AF639	AF634 relA251	This work	
AF640	AF634 fadR::Tn10	This work	
AF641	AF634 uspA::kan	This work	
W3110	In(rrnD-rrnE)1	F. C. Neidhardt	
ZK126	W3110 ∆ <i>lac</i>	R. Kolter	
TN31	ZK126 uspA::lacZ-Km ^r	This work	
LS1345	MC4100 $\lambda\Phi(fadB-lacZ)$	DiRusso et al. (6)	
LS1346	LS1345 fadR::Tn10	DiRusso et al. (6)	
LS1349 LS1350	MC4100 $\lambda\Phi(fadL\text{-}lacZ)$ LS1349 $fadR$::Tn10	DiRusso et al. (6) DiRusso et al. (6)	
PN428	C600 $\lambda\Phi(fadD-lacZ)$	Black et al. (3)	
PN424	PN428 fadR::Tn10	Black et al. (3)	
LS1347	MC4100 $\lambda\Phi(fabA-lacZ)$	DiRusso et al. (6)	
LS1348	LS1347 fadR::Tn10	DiRusso et al. (6)	
RW11	fadR::Tn10	C. C. DiRusso	
K4633	recD	D. Friedman	
K5302	himA::kan	D. Friedman	
RH76	MC4100 Δ <i>cya-851</i>	R. Hengge-Aromis	
BD4100	MC4100 Δ <i>crp96 zhd-732</i> ::Tn <i>10</i>	R. Hengge-Aromis	
MG1655	Wild type	Laboratory stock	
LN1655	MG1655 Δfis-767	L. Nilsson	
GM2929	dam-13::Tn9	M. G. Marinus	
TN3151	W3110 uspA::kan	T. Nyström	
PD32	MC4100 Δhns-206::bla	V. deLorenzo	
CF1948	W3110 ΔrelA251 ΔspoT207	M. Cashel	
QC1554 QC2085	soxS::Tn10 ΔarcA::Tn10	D. Touati D. Touati	
QC2003	∆u, c/1 1 111 0	D. Iouan	
Plasmids			
pTN6091	5-kb PstI fragment harboring	T. Nyström	
r	uspA	· y · · · · · · · · ·	
pTN6093	2.3-kb <i>Kpn</i> I- <i>Pst</i> I fragment	T. Nyström	
•	harboring uspA	•	
pTN6095	XhoI-PstI fragment harboring	T. Nyström	
	uspA		
pKOK5	lacZ-Km ^r operon fusion vector	Kokotek and Lotz (10)	
pTL61T	lacZ operon fusion vector	Linn and St. Pierre (11)	
pCD126	Wild-type fadR on pACYC177	C. C. DiRusso	
pRW22	fadRS219N on pACYC177	C. C. DiRusso	
pTN223	P_{tac} -uspA operon fusion	Nyström and Neidhardt	
n A E602	1.2 kb Cacli Dati from the	(21)	
pAF602	1.2-kb SacII-PstI fragment har- boring usn4	A. Farewell	
	boring uspA		

transformation and subsequently moved by P1 transduction into strain MC4100 (Fig. 1A). The lack of UspA synthesis in these strains was confirmed by two-dimensional gel electrophoretic analysis.

A plasmid-borne P_{uspA} -lacZ fusion was constructed by inserting the Sma1-Sal1 fragment of pTN6095 (Xho1-uspA-Pst1 [18]), containing the uspA promoter, into plasmid pTL61T (11) (Fig. 1B) digested with the same enzymes. The uspA-lacZ fusion was recombined into the chromosome by the method of Simons et al. (27) using λ phage RS45 (Fig. 1B). One strain, AF634, monolysogenic for λ phage carrying P_{uspA} -lacZ was isolated for further studies.

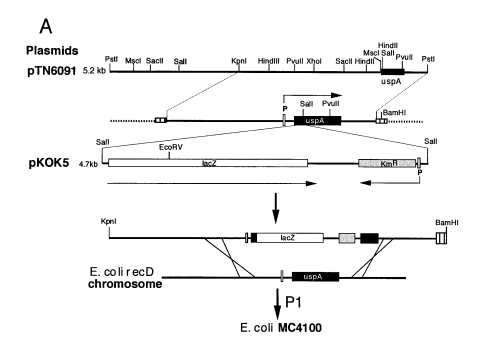
Measurements of β-galactosidase activity. The β-galactosidase levels were measured as previously described by Miller (14) with modifications (1). Samples were measured spectrophotometrically at 420 nm (β-galactosidase) and 550 nm to correct for remaining cell debris. The β-galactosidase activity is expressed as follows: $[OD_{420} - (1.75 \times OD_{550})]/(OD_{600}$ culture × reaction time × volume), where OD_{420} is the optical density at 420 nm.

FadR-uspA footprinting with DNase I. A 377-bp fragment including positions –214 to +163 of uspA was amplified from plasmid pAF602 using the PCR and oligonucleotides 5' GTCAGAATTCTCCCGATACGCTGCCAATCAGTT and 5' TCGAGGATCCAGGTCGACCGCGATGAGAATGTGT as primers. The underlined nucleotides include those added to generate EcoRI and BamHI restriction sites, respectively. The amplified fragment was cloned into pUC19 to generate pMD107-19. The insert was verified by DNA sequencing. For DNase I footprinting of the template strand, the reverse primer was phosphorylated with [y-³²P]ATP (7,000 Ci/mmol) and polynucleotide kinase. The PCR primers used for the coding strand were 5' TAAGCAAGGCGGATTGA and 5' CCGCGAT GAGAATGTGTT. Thirty-five cycles of PCR were performed; 1 PCR cycle consists of 1 min at 94°C, 1 min at 36°C, and 1 min at 72°C. The fragment of interest was gel purified. DNA was treated with FadR and DNase I as described previously by DiRusso et al. (5).

RESULTS

The uspA gene is transcriptionally activated during entry of cells into stationary phase. β-Galactosidase activities were determined for E. coli strains lysogenic for λ phage carrying P_{uspA} -lacZ fusions (strain AF634) and a strain (TN4100) in which uspA::lacZ was integrated by double homologous recombination into the normal uspA location on the chromosome. The uspA promoter was induced in early stationary phase, regardless of whether the fusion was located at the λatt site or the normal *uspA* site on the chromosome (Fig. 2A). The same results were obtained when the uspA::lacZ fusion was inserted into the chromosome of *E. coli* W3110∆*lac* instead of MC4100 (not shown). The induction was independent of functional UspA, since strain AF634, made uspA::kan (AF641), exhibited β-galactosidase levels during exponential growth and in stationary phase identical to those of the parent (Fig. 2A). The induction upon entry into stationary phase of the uspA-lac fusions was approximately fourfold. This is lower than the induction ratio previously reported when the UspA protein was examined (18, 20), indicating that perhaps there is posttranscriptional as well as transcriptional regulation of uspA. The transcriptional levels of uspA-lacZ were independent of growth rate (Fig. 2B), as are the levels of UspA synthesis (18).

Induction of uspA is dependent on the status of the fadR **gene.** To assess whether known regulatory genes control P_{uspA} activity, mutations in such genes were transduced to strains AF634 and TN4100 and their effects on β-galactosidase activity were determined. Regulators tested included cya, crp, soxS, rpoH, arcA, fur, relA, fis, dam, himA, oxyR, rpoS, and hns (Fig. 3). No significant effects on growth phase-dependent uspAlacZ expression were found except for the fadR mutation. The crp and the cya mutations significantly increased expression from P_{uspA} both during exponential growth and in stationary phase (Fig. 3). The activity of the *uspA* promoter was reduced in cells carrying a himA insertion mutation, but like mutations in crp and cya, the effects were seen in both logarithmic growth and stationary phase, and the ratio of β-galactosidase levels in stationary phase to that in exponential growth was very similar to this ratio in the wild-type strain. In other words, wild-type as well as cya, crp, and himA strains had a four- to fivefold induction of uspA-lac in stationary phase over that of log phase.



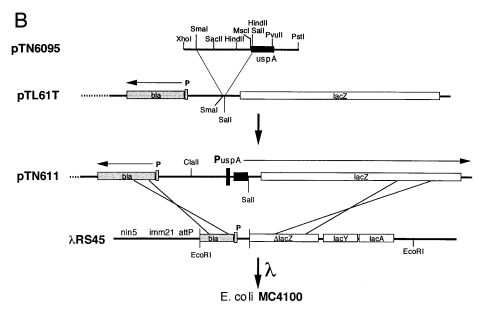


FIG. 1. Schematic drawing of the construction of P_{uspA} -lacZ fusions integrated at the normal uspA location on the chromosome (A) or at the λatt site (B). The figure is not drawn to scale. (A) The uspA:lacZ-Km^r into one was constructed by inserting the Sall fragment of plasmid pKOK5 (10) containing lacZ-Km^r into the Sall site of uspA carried on plasmid pTN6093 (19). The correct orientation of lacZ with respect to the uspA promoter was confirmed by digesting the resulting plasmid (pTN6099) by BamHI and EcoRV. This plasmid was linearized by Kpnl-BamHI digestion, and the uspA-lacZ-Km^r fusion was integrated into the chromosome of the E. coli recD K4633 by linear transformation and subsequently moved by P1 transduction into strain MC4100. (B) A plasmid-borne P_{uspA} -lacZ fusion was constructed by inserting the Smal-Sall fragment of pTN6095, containing the uspA promoter, into plasmid pTL61T (11) digested with the same enzymes. In the resulting plasmid (pAF632), lacZ is transcribed from the uspA promoter. λ phage RS45 (27) contains the λ end of the λ end of the λ end of λ end of λ in general plasmid, and recombination between the plasmid and phage. λ RS45 were identified by plating the lysate on MC4100 in medium containing X-Gal (5-bromo-4-chloro-3-indolyl- β -p-glucuronic acid).

In contrast, mutations in fadR significantly affected growth phase-dependent expression from the uspA promoter. The fadR null mutant exhibited significantly higher levels of β -galactosidase in the uspA-lacZ fusion during exponential growth over that of the isogenic parent, and there was only a small degree of induction upon entry into stationary phase (1.6-fold)

(Fig. 4A). This phenotype was complemented with plasmid pCD126 carrying the wild-type *fadR* allele (Fig. 4A). Levels of β-galactosidase were also significantly higher during exponential growth in the FadR⁺ strain when the long-chain fatty acid oleate was added to the medium (Fig. 4B). No effect was observed with the addition of glucose or glycerol (Fig. 4B). In

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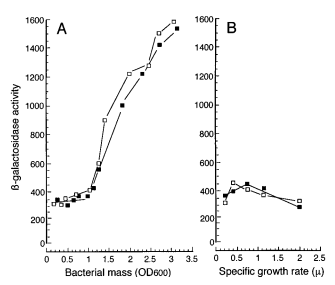


FIG. 2. (A) Expression of the *uspA* promoter in *λuspA-lacZ* lysogens AF634 (wild type) and AF641 (*uspA::kan*) during exponential growth and in stationary phase. β-Galactosidase activity is plotted as a function of cell mass (optical density at 600 nm [OD₆₀₀]). Cell growth typically started to cease at an OD₆₀₀ of 1.0 in the growth conditions examined (LB medium with glucose, 37°C, aerobic conditions). (B) Expression of the *uspA* promoter during exponential growth of strains AF634 and AF641 in minimal medium supplemented with glucose plus amino acids, glucose, glycerol, succinate, or L-serine. β-Galactosidase levels are plotted as a function of growth rate which is expressed as μ , the first-order growth rate constant. Symbols: \square , AF634; \blacksquare , AF641.

addition, cells carrying a transdominant nonderepressible mutation in fadR (change of Ser-219 to Asn [fadRS219N] [24]) on plasmid pACYC177 exhibited weaker induction of uspA than did cells carrying the wild-type allele or vector alone (Fig. 4C). Taken together, these observations suggest that the increased expression of uspA is in part an effect of FadR-dependent derepression during entry of cells into stationary phase. However, regulatory factors other than FadR are likely to be involved in the induction of uspA, since a fadR null mutant still exhibits some induction of uspA-lacZ upon entry into stationary phase.

The effect of *fadR* mutations on *uspA* expression suggests that UspA may have a role in fatty acid metabolism. A *uspA* null mutant grows on long-chain fatty acids, such as oleate, as the sole source of carbon, albeit with a somewhat slower growth rate than the isogenic wild-type MC4100 (Fig. 5). Therefore, *uspA* does not appear essential for fatty acid degradation, but it may be involved in fatty acid biosynthesis and/or membrane composition and integrity as discussed below.

FadR protein binds upstream of the uspA coding region. To test if the effect of FadR on uspA expression was direct, we analyzed the binding of FadR to uspA by DNase I footprinting. As shown in Fig. 6, FadR clearly binds upstream of the uspA coding region at a position +88 to +104 relative to the start of transcription. Examination of the sequence shows that the region where FadR binds has sequence similarity to other FadR binding sites (Table 2). Thus, uspA should be considered part of the fadR regulon. A second site (positions -4 to +13) also shows some weak FadR binding (Fig. 6), but this site exhibits little homology to the other FadR binding sites.

The fadR regulon is derepressed in stationary phase. The results in the preceding sections indicate that the FadR repressor is inactivated or is itself repressed during entry of cells into stationary phase. If so, the fad genes and operons, derepressed

by fatty acids and repressed by FadR, should also be increasingly expressed in stationary phase while the expression of the fatty acid biosynthetic genes activated by FadR should be reduced. We examined β -galactosidase levels in three strains lysogenic for λ phage carrying lacZ fusions to the fadBA, fadD, and fadL promoters. The fadBA operon encodes two proteins of the β -oxidation multienzyme complex; fadD encodes acyl-CoA synthetase, and fadL encodes the outer membrane fatty acid transport protein (2). We found that all fad fusions examined were increasingly expressed in stationary phase with kinetics similar to that of expression of uspA (Fig. 7). Comparison of growth phase-dependent expression in the wild-type and fadR mutant demonstrated that the increased expression was at least partly dependent on the fadR gene (Table 3). In the medium used, fadBA and fadD expression was also dependent on cya and crp (not shown), while no effects of mutations in these genes could be observed in the fadL-lacZ fusion. These results are consistent with previous work showing that fadB and fadD are regulated by cyclic AMP receptor protein while fadL is not (3, 6).

We also examined β -galactosidase levels in a strain lysogenic for λ phage carrying a lacZ fusion to the fabA gene encoding β -hydroxydecanoyl-acyl carrier protein dehydratase which is required for unsaturated fatty acid biosynthesis (2). β -Galactosidase levels decreased in the fabA-lacZ lysogen in stationary phase (Fig. 7), consistent with inactivation of FadR. fabA expression was much lower during logarithmic growth in the fadR background and was not further regulated by growth phase transition (Table 3). As expected, no effects on fabA expression were observed by mutations in either cya or crp (not shown) (6).

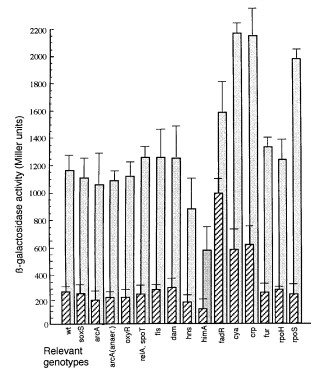


FIG. 3. Effects of a number of regulatory mutations on the expression of the uspA promoter. β -Galactosidase levels were determined during logrithmic growth (hatched bars) and approximately 1 h after growth ceased (gray bars). Cells were grown in LB medium with glucose. The strains used are described in Table 1. wt, wild type (MC4100); anaer., anaerobic.

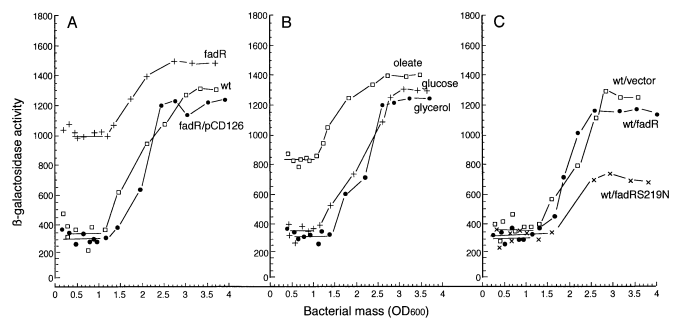


FIG. 4. (A) Effects of a *fadR* mutation on the levels of expression of the *uspA* promoter. β-Galactosidase levels were determined for strain TN4100 (wild type [wt]), TN4131 (*fadR*), and TN4131 carrying the wild-type *fadR* allele on plasmid pCD126 (fadR/pCD126). Cells were grown in LB medium with glucose. (B) Effect of including the fatty acid oleate in the growth medium on the levels of expression of the *uspA* promoter. The wild-type strain TN4100 was grown in LB medium supplemented with either oleate, glucose, or glycerol. (C) Effects of the transdominant nonderepressible mutation *fadR*S219N on the levels of expression of the *uspA* promoter. β-Galactosidase levels were determined for strain TN4100 carrying the nonderepressible *fadR*S219N allele (wt/fadRS219N), the wild-type allele (wt/fadR) or vector alone (wt/vector). Cells were grown in LB medium with glucose. The same results were obtained in a *fadR* null mutant carrying the *fadR*S219N plasmid as were obtained with a *fadR*⁺ cell (data not shown).

Derepression of the *fadR* **regulon is important for stasis survival.** To examine whether a failure to derepress the *fadR* regulon affects the cell's ability to survive stasis, we compared stationary-phase survival of cells carrying either the nonderepressible *fadR*S219N allele, the wild-type allele, or a vector plasmid. As depicted in Fig. 8, cells carrying *fadR*S219N were more sensitive to long-term stasis than cells carrying vector or the wild-type *fadR* allele. The same results were obtained whether the strains carrying the plasmids were *fadR*⁺ or *fadR* (data not shown). Because it is known that *uspA* mutants survive poorly in stationary phase (20), we examined whether the

poor ability of *fadR*S219N cells to survive was the result of their reduced ability to increase *uspA* expression during entry into stationary phase (Fig. 4C). We approached this question by transforming the cells carrying the *fadR*S219N plasmid with the compatible plasmid pTN223, which is pBR322 carrying an operon fusion between the *tac* promoter and *uspA* (21). Cells were starved in the presence of 50 μM IPTG (isopropyl-β-D-thiogalactopyranoside) which induces UspA synthesis eightfold (21), an induction level comparable to that reached in wild-type cells subjected to different starvation and stress conditions. However, *uspA* induction could not alleviate the harmful effect of the *fadR*S219N mutation (Fig. 8).

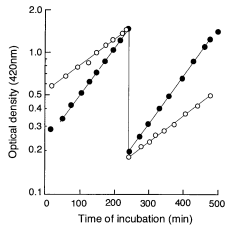


FIG. 5. Effect of uspA mutation on the growth of $E.\ coli$ on the fatty acid oleate. The wild-type strain MC4100 (\bullet) and its isogenic uspA::kan derivative (\bigcirc) were grown aerobically in oleate (0.5 mM) minimal M9 medium at 37°C. The optical density at 420 nm is plotted on a logarithmic scale as a function of time. The vertical line indicates dilution of the culture.

DISCUSSION

We report here that the expression of *uspA* is dependent on the status of the fadR gene. A fadR null mutant exhibits derepressed expression levels of uspA during exponential growth, while cells carrying a transdominant nonderepressible fadR allele increase uspA expression poorly during entry into stationary phase. The effects of fadR mutations on uspA expression suggest that UspA may have a role in fatty acid metabolism; all genes demonstrated to be regulated by FadR so far have assignments in fatty acid uptake, activation, or metabolism. However, a uspA null mutant grows on long-chain fatty acids, such as oleate, as the sole source of carbon, albeit with a somewhat slower growth rate than that of the isogenic wild type. Therefore, uspA does not appear essential for fatty acid degradation. Alternatively, UspA may be involved in fatty acid or membrane lipid biosynthesis. If so, the fact that FadR appears to repress uspA expression suggests that UspA may function as a governor that either reduces the pace of biosynthesis or affects the overall composition of membrane fatty acids. Several observations indicate that UspA may be involved in 6448 FAREWELL ET AL. J. BACTERIOL.

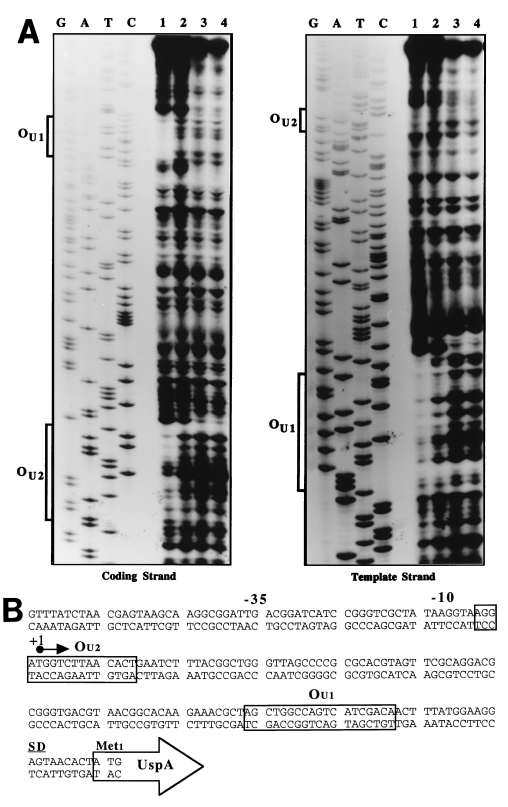


FIG. 6. (A) Identification of the FadR binding site within uspA and the sequence of the uspA promoter region. The products of sequencing reaction using the universal M13 reverse primer and pMC107-19 as a template are identified as G, A, T, and C. Lanes 1, 2, and 3 contain 4 mM, 400 nM, and 40 nM FadR, respectively. Lanes 4 have no added FadR. (B) Sequence of the uspA promoter region. The -35 and -10 regions of the uspA promoter, the transcription start site (small arrow), the FadR binding sites (O_{U1} and O_{U2}), the Shine-Dalgarno sequence (SD), and the first codon of uspA (large arrow) are indicated.

TABLE 2. Identified and predicted FadR binding sites

Gene	Sequence ^a		No. of bases in common with ^b :	
	•	O_{B}	O_A	
fadB	ATCTGGTACGACCAGAT	+17	(17)	10
fadL1	AGCTGGTCCGACCTATA	-9	11	11
fadL2	CACTGGTCTGATTTCTA	+16	7	11
fadD1	AGCTGGTATGATGAGTT	-29	12	9
fadD2	GGCTGGTCCGCTGTTTC	-115	7	8
fadE	AAGTGGTCAGACCTCCT		10	11
fabA1	AACTGATCGGACTTGTT	-31	10	(17)
fabB	GGCTGATCGGACTTGTT	-31	9	15
uspA	AGCTGGCCAGTCATCGA	+104	8	9
Consensus sequence	$\mathbf{ANCTGGTCNGANC}_{\mathbf{A}}^{\mathtt{T}}\mathbf{GTN}$			

^a Sequence data were taken from Black and DiRusso (2) or unpublished results (4). The position relative to the start of transcription as determined by primer extension analysis is shown. The *fadE* transcriptional start site has not been mapped.

membrane integrity. We have noted that uspA mutants are extremely sensitive to weak acids and exhibit an altered pH range for growth. Also, uspA mutants exhibit a diauxic type of growth when grown on glucose or gluconate. This phenotype was found to be the result of significant amounts of acetate being excreted into the medium of the mutant strain which, after an extended lag, was able to grow on this acetate (19). The excretion of acetate may be the result of an increased leakiness of the membrane of the mutant. In addition, it has been observed that UspA copurifies with the β -ketoacyl-acyl carrier protein synthase I (product of fabB) (23). These observations have provided the impetus to further explore possible alterations in membrane lipid composition due to the lack or overproduction of UspA.

The degradation of membrane constituents is an integral part of the dwarfing process of nondifferentiating bacteria subjected to starvation for exogenous carbon and energy. This

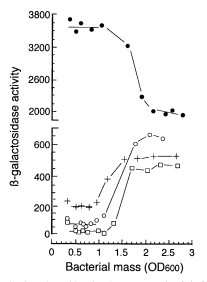


FIG. 7. Levels of β -galactosidase in λ lysogens carrying fadB-lacZ (\Box) , fadD-lacZ (\bigcirc) , fadL-lacZ (+), and fabA-lacZ (\bullet) during growth and in stationary phase. Consult the legend to Fig. 2 for details.

TABLE 3. Transcription levels of FadR-dependent genes during exponential growth and in stationary phase

	exponential growth and in stationary phase							
Strain	Relevant genotype	Growth phase	β-Galacto- sidase activity ^a	Induction ratio ^b				
LS1345	$\lambda\Phi(fadB ext{-}lacZ)$	Log Stationary	35 (11) 455 (21)	13.0				
LS1346	$fadR \ \lambda \Phi(fadB-lacZ)$	Log Stationary	467 (56) 1,077 (81)	2.3				
LS1349	$\lambda\Phi(fadL ext{-}lacZ)$	Log Stationary	206 (32) 505 (23)	2.5				
LS1350	$fadR \ \lambda \Phi(fadL\text{-}lacZ)$	Log Stationary	493 (21) 827 (73)	1.7				
PN428	$\lambda\Phi(fadD ext{-}lacZ)$	Log Stationary	61 (8) 651 (32)	10.7				
PN424	$fadR \ \lambda \Phi(fadD\text{-}lacZ)$	Log Stationary	341 (115) 1,007 (56)	3.0				
LS1347	$\lambda\Phi(fabA-lacZ)$	Log Stationary	3,680 (131) 1,140 (103)	0.31				
LS1348	$fadR \ \lambda \Phi(fabA-lacZ)$	Log Stationary	580 (40) 500 (35)	0.86				

 $^{^{\}prime\prime}$ Values are expressed as described in Materials and Methods under "Measurements of $\beta\text{-galactosidase}$ activity". Numbers in parentheses are standard deviations.

process generates small, coccoid cells (e.g., see references 17 and 25), and it includes extensive degradation of endogenous membrane phospholipids (9, 12), an activity proposed to provide the cell with carbon and energy for maintenance requirements. The derepression of the fadR regulon (either by inactivation or repression of FadR) during entry of cells into stationary phase suggests that this regulon, apart from being required for growth on exogenous long-chain fatty acids, may be involved in providing the growth-arrested cell with endogenous carbon and energy during dwarfing. The derepression of the regulon would make physiological sense in that fatty acids generated from degradation of membrane lipids during dwarfing could be scavenged by the activity of the acyl-CoA synthetase (product of the fadD gene) to generate acyl-CoA. Acyl-CoA is further catabolized by the β-oxidation enzymes (encoded by fadBA, fadE, fadFG, and fadH) to generate acetyl-CoA, a source of carbon and energy. Indeed, cells carrying the nonderepressible fadRS219N allele were more sensitive to long-term stasis than the isogenic parent (Fig. 8A). This mutated form of FadR has lost its ability to bind the effector molecule acyl-CoA but remains capable of binding DNA, repressing fad genes, and activating fab genes (24). Starvation also caused significant lysis of fadRS219N cells (Fig. 8B) which may be due in part to an altered membrane lipid composition.

The fadR regulon may be involved in the alterations in membrane phospholipid composition that take place during growth arrest. Specifically, fadR null mutants are known to synthesize more cardiolipin (diphosphatidylglycerol) and less phosphatidylglycerol (29). Thus, derepression of the fadR regulon during stationary phase would presumably result in increased levels of membrane cardiolipin; a feature typical of stationary-phase cells of E. coli (8). It may be argued that the

^b Number of bases in common with the binding sites of fadB (O_B) or fabA1 (O_A).

^b Induction ratios are expressed as the β-galactosidase activity levels obtained in stationary phase divided by β-galactosidase activity levels for cells in the exponential phase of growth.

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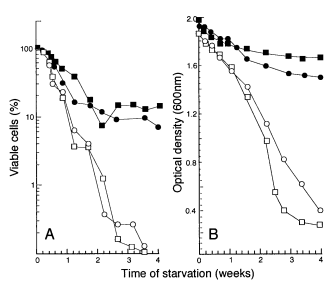


FIG. 8. Survival of cells carrying the vector (\bullet), the transdominant nonderepressible mutation fadRS219N (\Box), the wild-type fadR allele (\blacksquare), or fadRS219N together with pTN223 (\bigcirc). Strains were grown aerobically in LB medium at 37°C. After growth ceased, incubation was monitored for 4 weeks under the same conditions. (A) Viable cells. Viable cells were counted as colonies plated on LB medium after the appropriate dilution. (B) Optical density at 600 nm was read. One hundred percent viability corresponds to the number of viable cells counted 5 h after growth (measured by optical density) was arrested. At this time there was no further increase in cell number due to reductive division.

fadRS219N mutant would contain less cardiolipin in stationary phase because of its inability to derepress the fadR regulon. Cardiolipin is one of the major phospholipids in E. coli, but its essentiality and specific functions are unclear. It is mainly synthesized by cardiolipin synthase, encoded by cls, whose activity increases about 10-fold in stationary phase, while the activities and levels of other enzymes of phospholipid biosynthesis decrease (8). A null cls mutant is able to grow but survives stationary phase poorly (8). Thus, it is possible that the sensitivity of fadRS219N mutants to stasis is related to cardiolipin metabolism rather than a reduced ability to utilize endogenous fatty acids for maintenance requirements. However, the role of FadR in cls transcription or cardiolipin synthase activity remains to be elucidated. Future work will also include examination of the importance of alterations in membrane composition under the diverse stresses which are known to induce UspA and whether FadR regulation of UspA is sufficient to induce the protein in all these conditions. The mechanism by which FadR-dependent regulation in stationary phase occurs will also be a subject of future study; preliminary work indicates that FadR may not be responding to acyl-CoA thioesters synthesized by fadD, thus either FadR is itself repressed or is responding to a signal molecule synthesized by a novel fadDindependent mechanism in stationary phase. The role of UspA in growth phase alterations of membrane composition also remains to be elucidated.

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