# Identification of livG, a Membrane-Associated Component of the Branched-Chain Amino Acid Transport in Escherichia coli

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Branched-chain amino acids are transported into Escherichia coli by two osmotic shock-sensitive systems (leucine-isoleucine-valine and leucine-specific transport systems). These high-affinity systems consist of separate periplasmic binding protein components and at least three common membrane-bound components. In this study, one of the membrane-bound components, livG, was identified. A toxic analog of leucine, azaleucine, was used to isolate a large number of azaleucine-resistant mutants which were defective in branched-chain amino acid transport. Genetic complementation studies established that two classes of transport mutants with similar phenotypes, livH and livG, were obtained which were defective in one of the membrane-associated transport components. Since the previously cloned plasmid, pOX1, genetically complemented both livH and livG mutants, we were able to verify the physical location of the livG gene on this plasmid. Recombinant plasmids which carried different portions of the pOX1 plasmid were constructed and subjected to complementation analysis. These results established that livG was located downstream from livH with about 1 kilobase of DNA in between. The expression of these plasmids was studied in minicells; these studies indicate that livG appears to be membrane bound and to have a molecular weight of 22,000. These results establish that livG is a membrane-associated component of the branched-chain amino acid transport system in E. coli.

Active transport in gram-negative bacteria is mediated mainly by two classes of transport systems: osmotic shockinsensitive systems and osmotic shock-sensitive systems (12, 20). Osmotic shock-insensitive systems appear to utilize only one membrane-bound component (12, 30). Osmotic shock-sensitive systems appear to be more complex and require several membrane components along with a periplasmic binding protein (1, 9, 12, 26). The periplasmic components are soluble proteins with binding activities for a specific substrate or set of substrates (12, 14, 21). It has been proposed that their major role in transport is to deliver the substrate to the membrane components by direct interaction of the binding protein substrate complex with at least one membrane protein (3, 26). This interaction may activate conformational changes of one or more of the membrane proteins, which results in the delivery of free substrate inside the cell (1, 3, 26). Most shock-sensitive transport systems have more than two membrane-associated components which are present in much smaller quantities than the binding proteins (2, 7, 9, 10, 12, 27, 28).

We have found that the transport of the branched-chain amino acids in *Escherichia coli* is carried out by two periplasmic binding protein-dependent, high-affinity transport systems designated the leucine-isoleucine-valine (LIV-I) and the leucine-specific (LS) transport systems (4, 13, 22, 24). In addition, a membrane-bound low-affinity system designated LIV-II is present (5). The structural genes for the LIV-binding protein (*livI*), the leucine-specific binding protein (*livI*), and one of the membrane-associated components (*livH*) were initially identified by using genetic approaches involving the mutator phage, Mu, to isolate transport mu-

tants (4). Mutants have also been identified for the LIV-II transport system (livP) (5). The genes for the shock-sensitive transport systems (LIV-I and LS) contained in a 13-kilobase (kb) EcoRI DNA fragment have been cloned into the pACYC184 plasmid vector, yielding the pOX1 plasmid (20). By using subcloning strategies combined with transposon insertion mutagenesis and DNA sequencing, the livJ, livK, and livH genes have been identified on the pOX1 plasmid (13, 21; R. Landick, Ph.D. thesis, The University of Michigan, Ann Arbor, Mich., 1983).

In this paper, we report the identification of an additional membrane-associated component, livG, which is required for both of the high-affinity periplasmic transport systems. For this study, we first isolated ethyl methanesulfonate (EMS)-induced mutations in various components of the high-affinity transport systems and then subjected them to genetic complementation studies. In a second approach, a series of derivative plasmids were constructed from pOX1, carrying various portions of the liv regulon which allowed us to both physically and functionally identify the livG gene. The livG gene product has been tentatively identified as a membrane-associated protein with an apparent molecular weight of 22,000, by using a minicell expression system.

## MATERIALS AND METHODS

Bacterial strains and phages. The bacterial strains used for these studies were all derivatives of E. coli K-12 and are listed in Table 1. Bacteriophage P1CMclr100 was used for transductions and was a gift of D. Friedman and L. Rosner.

Media and chemicals. Cells for transport assays and osmotic shock treatment were grown in MOPS (morpholino-propanesulfonic acid) minimal medium (17) or Vogel-Bonner medium (29) supplemented with 0.2% glucose and 50  $\mu$ g of each of the required amino acids per ml except for leucine, which was present at 25  $\mu$ g/ml. Thymine was present at 50  $\mu$ g/ml, and pyridoxine-hydrochloride was present at 1  $\mu$ g/ml when required. Luria broth without glucose (18) was supplemented with 50  $\mu$ g of thymine LBT per ml and used for

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TABLE 1. Strains used in this study

Strain	Relevant genotype	Source
AE84	argG6 hisG1 trp-31 thyA746	Anderson <sup>a</sup>
	malA1 rpsL104 mtl-2 araC601	
	tonA2 lacY1 supE44 gal-6	
A E940201	gyrA260 xyl-7 pdxC3 livR	This sands
AE840201	livG; otherwise as AE84	This study
AE840203	livK; otherwise as AE84	This study
AE840212	livH; otherwise as AE84	This study
AE114	recA lstR livH::Mu thyA <sup>+</sup> ; otherwise as AE84	Anderson
AE126	mal <sup>+</sup> ; otherwise as AE84	By transduction
AE179	recA Tet <sup>r</sup> ; otherwise as AE84	By transduction
AE300	mal <sup>+</sup> glpD; otherwise as AE840203	By transduction
AE301	mal <sup>+</sup> glpD; otherwise as	Transductant from
	AE840212	line 8 glpD malT <sup>+</sup>
AE302	$mal^+$ $glpD$ ; otherwise as	Transductant from
	AE840201	line 8 <i>glpD</i> malT <sup>+</sup>
AE305	snl::Tn10; otherwise as AE300	From K230 by transduction
AE306	snl::Tn10; otherwise as AE301	From K230 by transduction
AE307	snl::Tn10; otherwise as AE302	From K230 by transduction
AE404	recA thy <sup>+</sup> ; otherwise as AE301	By mating
AE405	recA thy <sup>+</sup> ; otherwise as AE302	By mating
KL16-99	recAl relAl thi-l deoBl 3 HFr	From CGSC <sup>b</sup>
JM101	$lacI^{q}$ , ZM15 $traDI$ , F' $\Delta(lac-pro)$	From Bethesda
	,	Research
		Laboratories
F'104-5/86	F' mal <sup>+</sup> lin <sup>+</sup>	From F'140

<sup>&</sup>lt;sup>a</sup> Reference 4.

routine growth of strains. Plates for selecting mutants, transductants, and sexductants contained 1.5% agar and were based on Vogel-Bonner medium supplemented with the same concentrations of nutrients as described above. Ampicillin was used at 25 μg/ml. EMS was purchased from Eastman Organic Chemicals, Rochester, N.Y. <sup>14</sup>C- and <sup>3</sup>H-amino acids were from New England Nuclear Corp., Boston, Mass. L-[<sup>35</sup>S]methionine was from Amersham Corp., Arlington Heights, Ill. 5-Bromo-4-chloro-3-indolyl-β-D-galactoside and isopropyl-β-D-pyranothiogalactoside were from Bethesda Research Laboratories, Inc., Gaithersburg, Md.

Genetic techniques. All phage transductions were carried out with lysates of phage P1CMclr 100, as described previously (6). Matings were performed by the replica mating technique by the procedure of Miller (16). Homogenotes of several of the mutants were made by the following procedure. Mutant malT strains were mated with F' 140-5/86, containing the malT locus which is closely linked to the liv mutations (4). Sexductants were grown overnight in maltose minimal medium to ensure retention of the F'. Strains carrying the liv mutation on both the F' and the chromosome were selected by plating on minimal plates containing either maltose or glycerol and 0.1 µg of L-valine per ml. Valine-resistant colonies were plate mated with a recA F<sup>-</sup> strain to recover the recombinant F'. Resultant sexductants were purified and used for complementation studies.

Transport and binding assays. Routine transport assays of the indicated amino acids were carried out as described previously (4, 6). We also applied the following version of the rapid transport assay. Cells were grown overnight in 0.04% glucose–MOPS minimal medium to arrest growth at a constant, low cell density. The cells were harvested by centrifugation, washed four times with MOPS minimal medium, and suspended in MOPS with 0.2% glucose. The transport activity of the cells was determined by measuring the uptake of 0.5  $\mu$ M L-[ $^3$ H]valine or 0.1  $\mu$ M L-[ $^3$ H]leucine. Amino acid-binding activities of osmotic shock fluids were determined by equilibrium dialysis with 0.1  $\mu$ M L-[ $^3$ H]leucine or L-[ $^3$ H]isoleucine as previously described (23). Preparation of osmotic shock fluids was performed by the procedure of Neu and Heppel (19).

Isolation of EMS-induced mutants. The liv transport mutants were isolated from strain AE84 by mutagenesis with EMS and selection on plates for growth in the presence of the toxic analog azaleucine. EMS mutagenesis was carried out by the procedure of Miller (16). Azaleucine-resistant mutants were selected by plating dilutions of mutagenized cells on plates containing a concentration gradient of azaleucine. Mutants were also selected by plating untreated cells on minimal plates containing 100 µg of D,L-azaleucine per ml with a disk containing EMS in the center of the plate. In all cases, small colonies barely discernible above the background growth were picked, purified, and assayed for [3H]leucine and [14C]proline transport by the rapid transport assay. Decreases in [3H]leucine uptake relative to [14C]proline uptake indicate specific transport mutants for the LIV-I system.

Fifty-four azaleucine-resistant mutants were screened by the rapid transport assay. Roughly half (40%) had less than 50% of parental leucine transport while retaining normal proline transport. These mutants were subjected to osmotic shock treatment, and the periplasmic proteins were analyzed by sodium dodecyl sulfate (SDS)-polyacrylamide slab gel electrophoresis. Two classes of mutants were obtained, based upon the presence or absence of the binding proteins. One class had both LIV- and LS-binding proteins present (class I). The other class of mutants was missing the LS binding protein (class II).

DNA manipulations. Restriction endonucleases, T4 ligase, and DNA polymerase Klenow fragment were obtained from Bethesda Research Laboratories and from New England Nuclear Corp. Restriction endonuclease digestions were performed as described in the instructions of the suppliers. DNA ligation, transformation, and filling-in reactions were performed as described by Maniatis et al. (15). Procedures for plasmid DNA isolation have been described previously (8). Plasmid DNA fragments were analyzed by electrophoresis on horizontal 0.8% agarose gels or on 5% polyacrylamide gels as described previously (25).

Isolation of minicells. Minicells were purified from transformed *E. coli* minicell-producing strain X1411 grown in MOPS-rich medium as described previously (20). Purified minicells (200 to 400 μl; optical density at 420 nm, 1.0) were centrifuged, suspended in 100 μl of MOPS complete medium without L-methionine, incubated for 20 min to decrease background mRNA levels, and then labeled with L-[35S]methionine for 45 min. Labeled minicells were washed, suspended in 50 μl of sample buffer (11), and subjected to SDS-polyacrylamide gel electrophoresis by the Laemmli procedure (11). Fluorography enhancement reagents were obtained from New England Nuclear Corp.

For the minicell fractionation, labeled minicells were treated with 50 mM Tris (pH 8) containing 25 mM EDTA and 500 µg of lysozyme per ml for 15 min on ice and disrupted by

<sup>&</sup>lt;sup>b</sup> CGSC, E. coli Genetic Stock Center, Yale University, New Haven, Conn.

1198 NAZOS ET AL. J. BACTERIOL.

sonication. Minicell membrane preparations were separated from the cytoplasmic-periplasmic fraction by centrifugation at  $100,000 \times g$  for 16 h.

#### RESULTS.

Phenotypes of EMS-induced liv transport mutants. In a wild-type E. coli K-12 strain with normal regulation of leucine transport, the kinetics of leucine transport yield biphasic reciprocal plots due to the presence of low-affinity (LIV-II) and high-affinity (LS and LIV-I) transport systems (24). In livR<sup>-</sup> strains, however, derepression of the LIV-I and LS systems increases high-affinity transport and largely masks the biphasic nature of the kinetic plots, so that the LIV-II contribution to the total leucine transport is essentially negligible at low leucine concentrations (6). LIV-I transport mutants show kinetics characteristic of having only the low-affinity LIV-II transport present. Two classes of LIV-I transport mutants resistant to azaleucine were selected as described in the Materials and Methods section. When the kinetics of uptake in the azaleucine-resistant mutant strains AE840201 and AE840212 of class I and AE840203 of class II were examined, we found only the LIV-II, or low-affinity, system present. Values for the kinetic parameters,  $K_m$  and  $V_{\text{max}}$ , derived from Lineweaver-Burke plots of leucine transport of the various mutant classes are summarized in Table 2. The lower  $K_m$  value for strain liv302 may result from a partially defective membrane component. Also shown in Table 2 is the measurement of the binding activity in osmotic shock fluids from the mal<sup>+</sup> derivatives of azaleucine-resistant mutant strains AE840201, AE840203, and AE840212 and the parent strain AE84. The results show that the shock fluid from the mutant strain AE300 (AE840203 mal<sup>+</sup>) exhibits leucine-binding activity which can be completely inhibited by isoleucine, indicating a lack of functional LS binding protein. In addition, the shock fluids from mutant strains AE301 and AE302 of class I appear to have normal leucine-binding activity, indicating lesions in the nonbinding protein components of the transport system. These conclusions were further supported by examining the presence of the binding proteins from the shock fluids of the above strains by polyacrylamide gel electrophoresis (data not shown).

It appears that these EMS-induced azaleucine-resistant mutants show phenotypes similar to some of the Mu phage induced mutations in LIV-I isolated previously in this laboratory (4). Azaleucine-resistant mutant strain AE300 gives a

TABLE 2. Summary of kinetic parameters in LIV mutants

Trans- port genotype	$V_{ m max}$ (nmol/min per mg) $^a$	$K_m(\mu M)^a$	L-leucine binding activity <sup>b</sup> (nmol/mg of protein)		
			Without isoleucine	Plus isoleucine	
liv <sup>+</sup>	8.80	0.5	0.91	0.18	
livK300	2.66	8.4	0.71	_c	
livH301	2.19	6.8	0.60	0.06	
livG302	1.14	2.5	0.70	0.20	

 $<sup>^</sup>a$  Uptake was measured with 0.1  $\mu M$  L-[³H]leucine in strains AE84, AE840203, AE840212, and AE84201, respectively.

TABLE 3. Complementation analysis of liv mutations<sup>a</sup>

Recipient strains	Mutations on F140-5/86 (nmol/min per mg)				Uptake of haploids
Recipient strains	livK300	livH301	livH302	liv <sup>+</sup>	(nmol/min per mg)
AE306 livH301	0.03	0.05	1.03	0.52	0.08
AE307 livH302	0.07	1.00	0.02	0.44	0.05
AE305 livK300	0.04	0.05	0.05	1.14	0.08
AE114 livH::Mu	0.07	0.10	0.26	0.76	0.16
AE84 liv <sup>+</sup>	<u>_</u> b				1.1

<sup>&</sup>lt;sup>a</sup> Assayed by uptake of 0.5 μM L-[<sup>3</sup>H]valine.

LivK phenotype, while strains AE301 and AE302 have phenotypes similar to that reported for *livH* (4). The mutations in these strains were referred to initially as *liv301* and *liv302*.

The EMS-induced mutants were located by F' mapping by using F' 140-5/86, which covers the *liv* region near *malT* at min 74 on the *E. coli* chromosome, a region in which the previous Mu-induced leucine transport mutants were found to map (4). Complementation data (Table 3) indicate that all of the mutations are complemented by F' 140-5/86.

Complementation studies of the liv transport mutants. To determine the number of the genes that can be assigned to branched-chain amino acid transport from the EMS-induced mutations, genetic complementation studies were performed. Homogenotes from the mutant strains AE300 (livK300), AE301 (livH301), and AE302 (livH302) were isolated as described in Materials and Methods and used for mating with recA derivatives of each of the mutant strains. The complementation properties of the resultant sexductants were examined by measuring L-valine transport activity, and the results are presented in Table 3. Mutations livH301 and livH302, both from class I, were found to complement one another. Although not conclusive, these results suggest that mutants originally designated as livH301 and livH302 represent two distinct genes, which were tentatively named livH and livG, respectively. There was the possibility, however, that intracistronic complementation occurred within the livH gene. To confirm the existence of the livG gene, we took advantage of the availability of the pOX1 plasmid which contained all the genes for LIV-I and LS transport systems (20). We constructed a number of recombinant plasmids containing a subset of pOX1 DNA fragments representing different sequences downstream from the livK gene and attempted to map functionally and physically the livG gene as described below.

We also found that the livK300 mutation failed to complement either the livH301 or the livG302 mutation and that the livH::Mu mutation was not complemented by the livH301 mutation and only weakly complemented by the livG302 mutation. These complementation patterns can be attributed to polar effects due to a point mutation in livK or to the insertion of the Mu element in livH, respectively, leading to the conclusion that the livK, livH, and livG genes are part of the same transcriptional unit (see Discussion).

Identification and cloning of livG. Previous work reported from this laboratory has shown that the 13-kilobase (kb) EcoRI DNA fragment contained in the pOX1 plasmid carries the genetic region for the branched-chain amino acid transport (20). Moreover, the exact locations of the livJ, livK, and livH genes (Fig. 1) have been defined by using subcloning strategies combined with DNA sequence analysis (13, 21;

 $<sup>^</sup>b$  LS binding protein activity was determined by equilibrium dialysis of crude osmotic shock fluids in 2.5  $\mu M$  L-{ $^3H$ ]leucine plus 200  $\mu M$  L-isoleucine. LIV-binding activity was calculated as the activity in 2.5  $\mu M$  L-( $^3H$ ]leucine minus LS-binding protein activity. These results were measured with strains AE126, AE300, AE301, and AE302.

<sup>&</sup>lt;sup>c</sup> —, Not determined.

b -, Not determined.

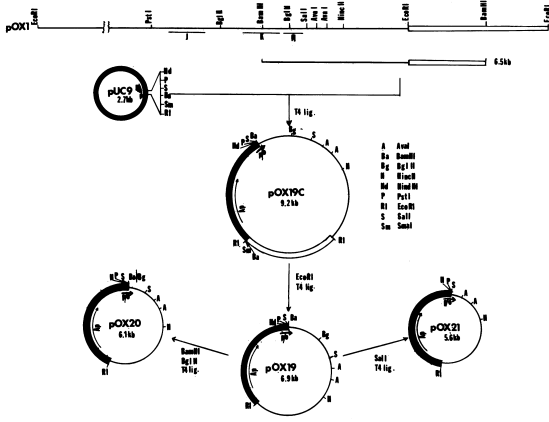


FIG. 1. Construction scheme for plasmids pOX19, pOX20, and pOX21. Plasmid pOX19C was derived by inserting the 6.5-kb BamHI DNA fragment from pOX1 into the single BamHI site of the pUC9 plasmid vector. Plasmid pOX19C was digested with EcoRI and religated, which deleted the smaller EcoRI fragment and gave plasmid pOX19. Digestion of pOX19 with Sall and subsequent religation produced pOX21. In addition, pOX19 was digested with BamHI and BglII and religated to produce plasmid pOX20. The double open line represents the portion of the pACYC184 vector of plasmid pOX1, the double solid line represents the plasmid vector pUC9, and the single line indicates E. coli chromosomal DNA.

Landick, Ph.D. thesis). Because of the physical map positions of livK and livH and the potential polar effects of livK300, livH301, and livH114::Mu mutations, we assumed that the livG gene should map downstream from the livH gene. We could eliminate the region on pOX1 upstream from the livJ gene as a potential location of the livG gene since a new operon, htpR, has been recently shown to map immediately upstream from the livJ gene (18).

To locate the livG gene physically in the liv regulon, we constructed recombinant plasmids pOX19, pOX20, and pOX21 from pOX1, with variable start points within or downstream from the livH gene, by applying the strategy shown in Fig. 1. DNA samples were isolated from plasmids pOX19C and pOX19C2, which carried the 6.55-kb BamHI DNA fragment cloned into pUC9 vector in both orientations, with respect to the lac promoter. Plasmid pOX19C, which was shown by restriction enzyme analysis (data not shown) to carry the livH and livG genes in the correct orientation, was used for further constructions. We were able to eliminate the pACYC184 DNA sequences (vector of pOX1) in plasmid pOX19C by digesting it with EcoRI and religating the mixture at a low DNA concentration to produce the pOX19 plasmid (Fig. 1). Plasmid pOX19 carries the 4.2-kb BamHI-EcoRI DNA fragment from pOX1 cloned into the pUC9 vector. Moreover, both the BamHI and SmaI sites of the pUC9 plasmid were deleted when the 2.3-kb EcoRI fragment was removed. The presence of a single BamHI site,

a Bg/II site (in the liv locus), and two Sa/I sites (one in the vector linker and one in the liv locus) allowed us to perform the following constructions. The pOX19 plasmid DNA was further digested with Sa/II and religated to eliminate the liv H gene contained within the 1.3-kb Sa/II fragment, yielding the pOX21 plasmid. In an alternative approach to the preparation of plasmid pOX20, the pOX19 plasmid was cut with BamHI and Bg/II and then religated to destroy the 5' end of the liv H gene by eliminating a 0.85-kb BamHI-Bg/II fragment, yielding the pOX20 plasmid. The 1.3-kb BamHI-Sa/II DNA fragment from pOX1, which carries the liv H gene, was cloned into pBR322 cut with BamHI and Sa/II and gave the pOX14 plasmid (manuscript in preparation).

The transport gene components contained in the pOX14, pOX19, pOX20, and pOX21 plasmids were determined by transforming the AE404 livH301 recA and AE405 livG302 recA transport-defective strains with these plasmids and by testing their ability to restore L-leucine transport activity. The results obtained from these complementation studies, along with a restriction map of the BamHI-EcoRI restriction fragment from pOX1, are summarized in Fig. 2. As shown in the figure, plasmid pOX14 was able to complement the livH301 mutation (strain AE404) but failed to complement the livG302 mutation (strain AE405). Alternatively, plasmids pOX20 and pOX21, which do not contain the livH gene, were able to complement the livG302 mutation but failed to complement the livH301 mutation. These results, combined

1200 NAZOS ET AL. J. BACTERIOL.

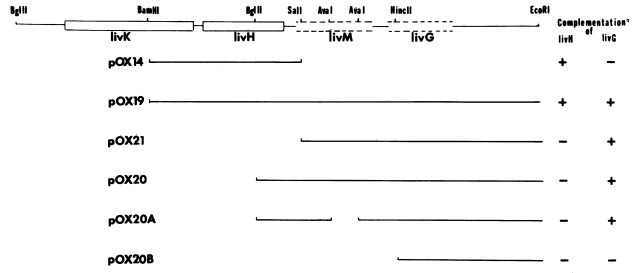


FIG. 2. Complementation analysis of livH and livG genes by various pOX plasmids. The solid lines indicate the portion of the Bg/II-EcoRI fragment shown at the top that is contained in the various plasmids listed on the left. On the right, the positive or negative results of the complementation analysis with these plasmids and the livH and livG strains are shown. Complementation was determined by measuring transport of 0.1 µM L-[3H]leucine in plasmid-transformed livH mutant strain AE404 and livG mutant strain AE405.

with the known physical location of the livH gene on pOX1 derived from DNA sequence information, strongly suggest that the livG302 mutation represents a new gene which we have named livG and which is located downstream from the previously identified livH gene.

To localize the livG gene more accurately within the cloned 2.9-kb SalI-EcoRI DNA fragment, we constructed a number of internal deletions in the pOX20 plasmid and examined the ability of these deletion plasmids to complement the AE405 livG mutant strain. To make these plasmids, the pOX20 plasmid was digested with AvaI restriction endonuclease, treated with the Klenow enzyme to create blunt ends, and religated to produce the pOX20A plasmid which carries a 300-bp AvaI deletion approximately 300 bp downstream from the livH gene (Fig. 2). In addition, we took advantage of the presence of two HincII restriction sites in the pOX20 plasmid (Fig. 1), one in the pUC9 polylinker region and a second one about 1.1 kb downstream from the livH gene, to produce another deletion plasmid. The pOX20 plasmid was cut with HincII restriction endonuclease and religated to give the pOX20B plasmid which carries a 1,000base-pair (bp) deletion ending approximately 1 kb downstream from the livH gene. These plasmids were used to transform the AE405 (livG) mutant strain and were tested for their ability to restore L-leucine high-affinity transport activity. Plasmid pOX20A complemented the livG302 mutation whereas plasmid pOX20B failed to complement it (Fig. 2). From these results, we concluded that the livG gene begins 600 to 1,000 bp downstream from the livH gene, within the Aval-HincII DNA fragment. We currently are determining the DNA sequence of this region. We also attempted to identify the livG gene product by examining the pOX19-, pOX20-, and pOX21-encoded proteins in minicells harboring these plasmids. The polypeptides were labeled with L-[35S]methionine, separated by SDS-polyacrylamide gel electrophoresis and visualized by autoradiography. Minicells containing the plasmid pOX19 produced two proteins with molecular weights of approximately 27,000 and 22,000 which were absent from the pUC9-containing minicell background. The 27,000-molecular-weight protein (27K protein) was also synthesized by the pOX20 plasmid containing minicells (data

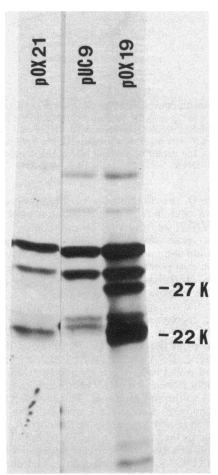


FIG. 3. Fluorogram of 12.5% SDS-polyacrylamide gel electrophoresis, illustrating L-[ $^{35}$ S]methionine-labeled proteins synthesized in minicells carrying the indicated plasmids. The unlabeled molecular weight markers (not shown) included bovine serum albumin ( $M_r$ , 67,000), ovalbumin (45,000), pepsin (34,700), trypsinogen (24,000), and lysozyme (14,300).

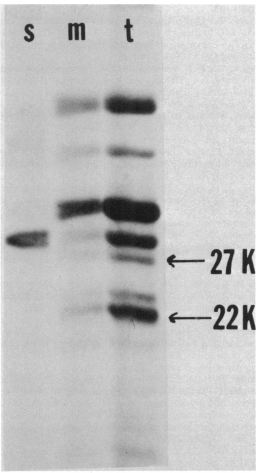


FIG. 4. Fluorogram of 12.5% SDS-polyacrylamide gel electrophoresis, illustrating the cellular fractionation of L-[35S]methionine-labeled proteins synthesized by plasmid pOX19 in minicells. Lane s contains the soluble fraction (cytosol plus periplasm); lane m contains the membrane fraction, approximately 50% of the total sample; lane t contains unfractionated minicells.

not shown) but was not made in detectable amounts by pOX21 plasmid containing minicells (Fig. 3). The small (22K) polypeptide, however, was present in pOX21containing minicells. It appears that neither the 27K protein nor the 22K protein is synthesized by pOX1-containing minicells in detectable levels (20). We assumed that the reason that both proteins are very poorly expressed by pOX1-containing minicells is that they are under the normal chromosomal promoter control. The fact that the 27K protein was not encoded by the pOX21 plasmid, even though this plasmid has livG complementation activity, makes this polypeptide an unlikely candidate for the livG product, leaving the 22K protein the most likely candidate. We tentatively conclude that the 27K protein may be the product of an additional gene (livM) lying between the livH and livG genes (see Discussion).

To determine the cellular location of the *livG* protein, pOX19 plasmid-containing minicells were labeled and fractionated into periplasmic, cytoplasmic, and membrane fractions as described in Materials and Methods. The labeled polypeptides of each fraction were analyzed on a 12.5% SDS-polyacrylamide gel. We found that under the conditions used for fractionation, both the 22K (*livG*) protein and

the 27K (*livM*) protein were associated predominantly with the membrane fraction (Fig. 4).

## **DISCUSSION**

We have previously shown that high-affinity branchedchain amino acid transport in E. coli requires two periplasmic binding proteins and at least one membrane-associated component (4). In those studies, the mutator phage Mu was used to induce mutants in the high-affinity branched-chain amino acid transport in E. coli, but because of the nature of the mutations, we were not able to perform complementation studies. In the present study, we have used additional genetic approaches combined with recombinant DNA techniques, to identify one of the membrane-associated components of this system, livG. A combination of less stringent selective conditions and the use of derepressed starting strains made it possible to isolate EMS-induced mutants in the three previously identified genes, livH, livJ, and livK, and in a new gene, livG. The tentative identification of the livG mutation was shown by genetic complementation studies. Since livH and livG mutations have similar phenotypes, they are difficult to distinguish. Similar results have been reported for the histidine transport system in Salmonella typhimurium (2, 10). We were able to use the cloned liv regulon to confirm the results of the genetic studies. The livG gene was mapped both functionally and physically by using a number of recombinant plasmids carrying different portions from the pOX1 plasmid. We found that plasmid pOX20, which carries a defective livH gene; plasmid pOX21, which lacks the livH gene; and plasmid pOX20A, which carries a defective livH gene and a 300-bp AvaI deletion approximately 300 bp downstream from the livH gene, all contain a functional livG component. These results suggested that livG defines a separate component for the highaffinity branched-chain amino acid transport systems, a component which maps downstream from the livH gene. The fact that plasmid pOX20A carried livG complementing activity and plasmid pOX20B (which lacks a 1.1-kb HincII fragment) did not suggests that the livG gene begins approximately 600 to 1,000 bp downstream from the livH gene, within the AvaI-HincII fragment.

A DNA sequence with an open reading frame lies between the livH and livG gene with a coding capacity for a protein with a molecular weight of approximately 30,000. This observation, taken together with the results from the minicell expression experiments, suggests that an additional gene may be located between the livH and livG and that the 27K protein present in the pOX19- and pOX20-containing minicells may be the product of this gene. The additional putative gene was named livM and is currently under further investigation. The 22K protein produced by the pOX19-, pOX20-, and pOX21-containing minicells is tentatively identified as the livG gene product since its presence corresponds to the livG-complementing activity of these plasmids. Both livG and the putative livM gene products appear to be membrane associated. The existence of multiple components for the shock-sensitive, high-affinity, branched-chain amino acid transport systems appears to be a common feature for other binding protein-dependent transport systems (1, 13, 30).

A common property of the membrane components of shock-sensitive transport systems is that they are often expressed at very low levels (2, 9, 27, 28). We attempted to amplify the expression of the livG gene by cloning the gene after a strong, controllable promoter, such as the lac promoter. We found that the 22K and 27K proteins are both

1202 NAZOS ET AL. J. BACTERIOL.

expressed at higher levels from the pOX19 plasmid, which is under *lac* promoter control, than from the pOX1 plasmid, which is under chromosomal control (21).

As shown by the complementation studies, the livK mutation that we have examined in strain AE300 failed to complement either the livH or the livG mutations. Although the nature of the mutation in the livK gene is not known, it appears to be a polar mutation since the selection scheme we used demanded that it also be defective in the LIV-I transport system. These and other results suggest that livK, livH, and livG genes are located on the same transcriptional unit (unpublished data).

We are currently studying the nature of the membrane-associated transport components by taking advantage of the amplified expression of the cloned genes. For these studies, we have constructed a number of  $\beta$ -galactosidase gene fusions to the livH, livM, and livG genes.

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