# Mapping of Two Loci Affecting the Regulation of Branched-Chain Amino Acid Transport in *Escherichia coli* K-12

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Two mutant loci resulting in derepression of, respectively, the L-leucinespecific transport system (lstR) and both the leucine-specific and the general branched-chain amino acid transport LIV-I systems (livR) were mapped by conjugation and transduction. Both livR and lstR were found to be closely linked to aroA at min 20 on the Escherichia coli genetic map. The merodiploid  $livR^+/$ livR displayed wild-type regulation of L-leucine transport, indicating that the livR product is a diffusible, negative controlling element for high-affinity leucine transport systems. Isogenic strains carrying lstR, livR, and wild-type transport alleles were compared for leucine uptake kinetic parameters and leucine-binding protein levels. The higher levels of leucine transport in the mutant strains under repressing conditions were generally due to increased high-affinity systems, which were accompanied by striking increases in the level of leucine-binding proteins.

The transport of L-leucine, L-isoleucine, and L-valine in Escherichia coli K-12 is complex, being mediated by at least three kinetically distinct systems (4, 18, 23). There are two highaffinity systems, the LIV-I and the leucine specific. The LIV-I system is characterized by a low  $K_m$  of transport for leucine, isoleucine, and valine and also shows affinity for L-threonine, Lalanine, L-serine, and L-homoserine (23). These high-affinity systems are repressible by leucine, apparently through leucyl-transfer ribonucleic acid (17). The leucine-specific system, which comprises about 20% of the high-affinity transport of leucine in wild-type E. coli K-12, can be identified by its resistance to competition by isoleucine. This system is also repressible by leucine and perhaps methionine (23). Both high-affinity systems are lost upon osmotic shock, coinciding with the loss of at least two binding proteins, one identified as the LIVbinding protein (16) and another the leucinespecific binding protein (L-binding protein) (3).

The low-affinity membrane-bound transport system for leucine has a 10-fold higher  $K_m$  for all three branched-chain amino acids, is resistant to osmotic shock, and is apparently nonrepressible (18, 23).

Previously this laboratory reported the isolation of two mutant strains of E. coli K-12 with derepressed levels of the high-affinity systems (18). Strain EO311 had derepressed levels of the LIV-I system and the leucine-specific system; strain EO318 had derepressed levels of the leucine-specific system. We have designated the mutant locus affecting the regulation of the LIV-I system in strain EO311 as livR and the mutant locus affecting the leucine-specific system in strain EO318 as lstR. In this paper we report the map positions of these two loci, the characteristics of strains diploid for the livR locus, and the transport and binding protein characteristics of isogenic strains carrying the wild-type, lstR, and livR alleles.

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### MATERIALS AND METHODS

**Bacterial strains.** The strains used were all derivatives of E. coli K-12 and are listed in Table 1. The leucine auxotrophs EO311 and EO318 were isolated as p-leucine utilizers as reported previously (18).

Materials. MOPS (morpholinopropane sulfonic acid), tricine [N-tris(hydroxymethyl)methyl glycine], and nutritional supplements were products of Sigma Corp. (St. Louis, Mo.). 5,5,5-Trifluoro-DL-leucine was obtained from Columbia Organic Chemicals Inc., Columbia, S.C., and DL-4-azaleucine was obtained from Nutritional Biochemicals Corp., Cleveland, Ohio.

Media. Minimal medium for growth of cultures used in transport assay was basal MOPS-salts medium (14) supplemented with 0.2% glucose and 50  $\mu$ g of required L-amino acids per ml except for L-leucine, which was used at 25  $\mu$ g/ml. Thymine and uracil were present at 50  $\mu$ g/ml, and vitamins were present at 1  $\mu$ g/ml. Luria broth without glucose (11)

Strain	Sex	Relevant genotype	Source
EO300	$\mathbf{F}^{-}$	Wild-type K-12	Rahmanian et al. (18)
EO311	$\mathbf{F}^{-}$	leu, trp, dlu, livR	Rahmanian et al. (18)
EO318	$\mathbf{F}^{-}$	leu, trp, dlu, lstR	Rahmanian et al. (18)
F15/EO311	F-prime	Chromosomal markers as EO311	Unselected sexductant of EO311 from F15/KL110, a K. B. Low strain
Hfr 6-1	Hfr	Point of origin in Low (10)	CGSC <sup>a</sup>
KL208	Hfr	Point of origin in Low (10)	CGSC
KL209	Hfr	Point of origin in Low (10)	CGSC
KL983	Hfr	Point of origin in Low (10)	CGSC
KL999	Hfr		CGSC
MA1079	Hfr	Point of origin in Low (10) thy <sup>+</sup> , recA; point of origin of KL16 (10)	W. Maas strain via K. B. Low
JC1552	F-	argG6, metB1, his-1, leu-6, trp- 31, mtl-2, xyl-7, malA1, gal-6, lacY1, or lacZ4, str-104, tonA2, tsx-1, λ <sup>R</sup> λ <sup>-</sup> , supE44	CGSC
AB1133	$\mathbf{F}^{-}$	ara-14, leu-6	CGSC
AB2829	$\mathbf{F}^+$	aroA354	CGSC
KL282	Hfr	serC13, serS16	CGSC
KL282-1	Hfr	serC13	EO300 transductant of KL282 to tem- perature resistance
KL188	$\mathbf{F}^{-}$	pyrD34	CĜSC
PLS-73		pyrD34	CGSC
AT3143	- F-	pdxC3	CGSC
AE9	F-	nalA, otherwise as JC1552	Spontaneous
AE18	Hfr	<i>leu-6, lstR;</i> point of origin of KL209 (10)	D-Leucine-utilizing transductant (from strain EO318) of an Hfr recombinant between KL209 and AB1133
AE20	$\mathbf{F}^{-}$	$met^+$ , otherwise as AE9	$met^+$ transductant of AE9, from EO300
AE30	F-	leu-6, lstR, trp, his, gal	<i>met</i> <sup>+</sup> recombinant of conjugation of AE18 and AE9
AE40	$\mathbf{F}^{-}$	<pre>metB<sup>+</sup>, thyA, araC, otherwise as in AE9</pre>	Sequential spontaneous mutants of AE9
AE41	F-	lstR, otherwise as in AE40	p-Leucine-utilizing transductant of AE40 from AE30
AE49	F-	aroA354, otherwise as in AE40	Penicillin-enriched aroA transductant of AE40 from AB2829
AE52	$\mathbf{F}^{-}$	livR, otherwise as AE40	aro <sup>+</sup> transductant of AE49 from EO311
KLF3/AE55	F-prime	F103 his+/his, livR	F103 from KLF3/JC1552; AE55 is an aro <sup>+</sup> transductant of AB2829 from EO311, made his by P2 eduction (8)
AE59	$\mathbf{F}^{-}$	<i>thyA</i> <sup>+</sup> , <i>recA</i> , otherwise as in AE49	$thy^+$ recombinant of MA1070 and AE49
AE62	F	$leu^+$ , otherwise as in AE40	<i>leu</i> <sup>+</sup> transductant of AE40 from EO300
AE63	F-	$leu^+$ , otherwise as in AE41	$leu^+$ transductant of AE41 from EO300
AE64	F-	aroA354, livR, thyA	gal <sup>+</sup> recombinant of conjugation be- tween KLF3/AE55 and AE49
AE65	F-	$recA$ , $thy^+$ , otherwise as in AE64	thy <sup>+</sup> recombinant of conjugation be- tween MA1079 and AE64
AE66	$\mathbf{F}^{-}$	<i>leu</i> <sup>+</sup> , otherwise as in AE64	leu <sup>+</sup> transductant of AE64 from EO300
AE68	F-	$leu^+$ , otherwise as in AE52	leu <sup>+</sup> transductant of AE52 from EQ300
AE80	$\mathbf{F}^{-}$	pdxC3	<i>aroA</i> <sup>+</sup> transductant of AB2829 from AT3143

TABLE 1. E. coli K-12 strains

" CGSC, E. coli Genetic Stock Center.

was used for routine growth of strains and for matings. Recombinant selective plates contained 1.5% agar, based on half-strength medium 56 (9), and were supplemented with nutrients at concentrations listed above for liquid media. Fluoroleucine was used at 0.5 to 5  $\mu$ g/ml, and azaleucine was used at 50 to 100  $\mu$ g/ml. Strains with the *aroA* mutation required L-phenylalanine, L-tyrosine, L-tryptophan, and the vitamins *p*-hydroxybenzoate, *p*-aminobenzoate, and 1,4-dihydroxybenzoate. Strains with *serC* 

mutations required both L-serine and pyridoxine; strains with pdxC required pyridoxine alone. Strains with pyrD required uracil.

Genetic crosses. Conjugations were carried out as recommended by Miller (12). Donor Hfr and recipient F<sup>-</sup> strains were used at  $3 \times 10^8$  cells/ml after two doublings in Luria broth and were mixed 10:1 (F<sup>-</sup>:Hfr) in a final volume of 4 ml in a 125-ml Erlenmeyer flask, usually unshaken. Matings in this study were not interrupted. Samples were drawn after 60 to 90 min of incubation at 37 C, diluted, and plated in 3 ml of soft agar (0.75%) on selective plates containing streptomycin (100  $\mu$ g/ml) as counterselective agent. F-prime transfers were done similarly. Transductions were carried out by using phage P1 CM clr100 (gift of D. Freidman from L. Rosner), a temperature-inducible phage (20) conferring chloramphenicol resistance upon lysogens. Lysate preparation through heat induction of lysogens was carried out as described (12). For transductions, 0.1 ml of a series of 10-fold dilutions of phage was incubated with 0.1 ml of recipient cells (washed and resuspended in 0.01 M MgSO<sub>4</sub>-CaCl<sub>2</sub>) at 37 C for 30 min. Controls of undiluted lysate and cells alone were treated similarly. Killing was reduced by adding 0.2 ml of 1 M sodium citrate (pH 7.0), and the contents of the tubes were plated out directly or in 3 ml of soft minimal agar on selective plates at 37 C. Colonies for analysis were chosen from the highest phage dilution that yielded a 20-fold or better excess of transductants over revertants. Unselected markers were scored by transferring recombinant colonies onto appropriate solid medium with a platinum wire (9) and replicating growing patches to selective plates. In some instances mentioned below, the variable quantity of inocula transferred by the velvet master introduced uncertainty in scoring; in this event, each patch was individually replicated by platinum wire.

Assay of transport and binding activities. Cultures of cells for transport assays were harvested in exponential phase by centrifugation and washed at 4 C three times with unsupplemented MOPS-salts medium. The cells were resuspended in this solution and incubated for 5 min at 37 C. The uptake reaction was started by adding the bacterial suspension to a solution of the radioactive amino acids. Uptake was measured by removing 0.5-ml samples at appropriate time intervals, filtering these through a 24-mm nitrocellulose filter (Millipore HA, Millipore Corp., Bedford, Mass.), and washing immediately with 5 ml of 37 C 10 mM potassium phosphate (pH 7.2) containing 0.1 mM MgSO4. Radioactivity was determined in a Packard liquid scintillation spectrometer with a standard scintillator solution. The rate of uptake was linearly proportional to the dry weight of cells in the range used and remained constant during the time each experiment required. The quantity of cells used was estimated from the absorbance at 420 nm, using a Zeiss PMQ II spectrophotometer. An absorbance of 1.0 equaled 0.16 mg/ ml. drv weight.

Experiments to determine the kinetics of transport were made by measuring the transport velocity of leucine at 13 substrate concentrations from  $8.5 \times$ 

 $10^{-8}$  to  $5 \times 10^{-6}$  M. These data were plotted by the Lineweaver-Burk method, and the slope and intercept of the linear portions were determined by least-squares analyses. When biphasic kinetics were observed, the slope and intercept from the two linear portions of the line were used to calculate two  $K_m$  and two  $V_{max}$  values, using the method of Neal (13). The four kinetic values obtained were checked by recalculating the expected velocity by the following equation:

$$v_{obs} = \frac{V_1(S)}{K_1 + (S)} + \frac{V_2(S)}{K_2 + (S)}$$

Further refinement of the kinetic parameters was performed by iterative procedures until the calculated values closely approximated the observed.

The isolation and quantitation of protein with leucine-binding activity from osmotically shocked cells was performed as described previously (15).

Rapid assay for transport. For the large-scale assay of L-leucine transport phenotypes among progeny of genetic manipulations, a rapid variant of the standard assay above was devised. MOPS minimal medium was prepared with supplements as required, except that 0.04% glucose was the carbon source and L-leucyl-L-alanine (15  $\mu$ g/ml) was the source of leucine for both Leu<sup>+</sup> and Leu<sup>-</sup> strains. Two-milliliter volumes of this medium were dispensed into test tubes (17 by 125 mm), inoculated with small portions of recombinant colonies, and incubated overnight at 37 C with shaking. Regardless of inoculum size or growth rate, all cultures ceased growth at the same density when glucose was exhausted. Glucose was then added back by a calibrated dropping bottle (Microdels, Microbiological Associates, Rockville, Md.) to 0.2%, and the tubes were incubated for 20 min to allow partial recovery. Chloramphenicol was added similarly (150  $\mu$ g/ml), and the tubes were placed at 37 C for assay. Standard strains were always included as internal controls. Cell samples (0.5 ml) were then added directly to 0.5 ml of unsupplemented MOPS salts containing L-[3H]leucine at a final concentration of 0.75  $\mu$ M. After 20 s of incubation, 0.5 ml of the mixture was rapidly filtered, washed, and counted as above. Reproducibility of this test was assayed by using 10 independent tubes for each of the standard wild-type and transport mutants examined in this report. The values obtained for the wild type (strain AE40) were a mean of 204 counts/min, standard error = 4%; for strain AE52 carrying the derepressed LIV-I transport allele livR, a mean of 2,668, standard error = 17%; for strain AE41 carrying the derepressed leucine-specific transport allele lstR, a mean of 1,821, standard error = 6%. The range of the different transport phenotypes did not overlap under these conditions.

Plate phenotypes of transport mutants. The original strains EO311 and EO318 were found to be multiple mutants (see Results). When strains carrying only the transport mutations were constructed by transduction, the phenotypes changed from those previously reported (18). Strains carrying *livR* were scored in a *leu* strain background by their ability to utilize 200 but not 100  $\mu$ g of *D*-leucine per ml as a source of L-leucine. The wild type cannot use D- leucine at 200  $\mu$ g/ml. In contrast, strains carrying *lstR* are able to utilize 100 or 200  $\mu$ g of p-leucine per ml and are easily distinguished on the basis of this property. In leu<sup>+</sup> strain backgrounds where D-leucine utilization cannot be assayed, strains with either lstR and livR mutations can be recognized by their sensitivity to 100  $\mu g$  of DL-azaleucine per ml (the wild-type strain is resistant to this level). Replica platings were used to distinguish wild-type from either lstR- or livR-carrying strains, although the latter two lesions cannot be distinguished from each other by this technique. They are best distinguished by direct transport assay for leucine uptake (see Rapid uptake assay, above). Strains requiring supplementation with the aromatic amino acids L-tyrosine, L-phenylalanine, and L-tryptophan posed special problems, since these three amino acids together at 50  $\mu$ g/ml completely prevented p-leucine utilization by otherwise competent strains; reciprocally, the presence of 200  $\mu g$  of p-leucine per ml prevented use of exogenous aromatic amino acids in aro strains. In leu<sup>+</sup> backgrounds the aromatic amino acids protected sensitive cells from azaleucine, but not trifluoroleucine, inhibition. Accordingly, when aro strains were used or screened for transport mutations, trifluoroleucine sensitivity was used to detect the *livR* or *lstR* alleles. Replica plating from velvet masters could not be reliably used, however, since variable inocula obscure sensitivity determinations due to an adaptation of cells to the analogue (19). Instead, each colony was individually streaked on trifluoroleucine-supplemented medium with a platinum wire and scored after overnight incubation. Useful trifluoroleucine concentrations varied from 0.4 to 5  $\mu$ g/ml, reflecting inherent strain background differences in sensitivity to this analogue.

## RESULTS

Direct selection of transport phenotypes. The mutant strains EO311 and EO318 were originally isolated as p-leucine utilizers (Dlu<sup>+</sup>) (18). The threshold concentrations of *D*-leucine for growth of the mutant strains on solid media were determined to be 60 and 40  $\mu$ g/ml, respectively. In contrast, the parental Dlu<sup>-</sup> leucine auxotroph cannot utilize 200  $\mu$ g of p-leucine per ml. Attempts to directly transduce the Dlu<sup>+</sup> phenotype associated with increased L-leucine transport using P1 phage grown on the two mutants met with limited success. When the donor was strain EO311, all Dlu+ transductants had wild-type transport activity for L-leucine, which was normally repressible, indicating that a nontransport mutation, dlu, existed in strain EO311, which permitted utilization of Dleucine in L-leucine auxotrophs. When the donor was strain EO318, greater than 90% of all directly selected Dlu<sup>+</sup> transductants also had normal L-leucine transport regulation and were presumably dlu mutants; however, several Dlu<sup>+</sup> transductants did show the derepressed transport phenotype. These transductants had a threshold for p-leucine utilization of 80  $\mu$ g/ml compared with nontransport *dlu* transductants, which had a threshold of 40  $\mu$ g/ml. Intervening growth of transductants from either donor strain before plating (to overcome phenotypic lag) had no effect on the low efficiency of recovery of p-leucine utilizers with derepressed transport; accordingly, we do not generally use p-leucine utilization to select for derepressed transport alleles.

The leucine prototrophic derivatives of the Dlu<sup>+</sup> strains EO311 and EO318 are sensitive to either 100  $\mu$ g of azaleucine per ml or 1  $\mu$ g of trifluoroleucine per ml; the Leu<sup>+</sup> Dlu<sup>-</sup> parental strain is resistant. Attempts to transduce out the transport mutations with phage grown on the transport wild type by selection for analogue resistance were unsuccessful due to high background growth on plates; these findings have forced us to treat the transport phenotypes as unselected characters in the crosses reported below. We do use the phenotypes of D-leucine utilization or analogue sensitivity as screening agents, depending on the genetic background of recipient strains.

Segregation of *dlu* and *livR* from strain EO311. The chromosome of strain EO311 (dlu livR) was mobilized by sexduction with the Fprime F15 (Fig. 1) and used as a donor in the cross described in Table 2. Two classes of Dleucine utilizers were recovered from selected  $his^+$  recombinants: those with a high efficiency of utilization of p-leucine but normally regulated L-leucine transport (dlu mutants) and those with low efficiency of utilization but carrying the derepressed leucine transport marker (livR). The relative frequency of the unselected livR, dlu, and gal (3/200) alleles among the  $his^+$  recombinants indicated that livR and dlu were distal to trp but proximal to gal with respect to the point of origin of F15 (Table 2). The dlu allele evidently maps closer to his than does livR.

**Mapping of** *lstR* **by conjugation.** The *leu* Hfr AE18 (Table 1) was constructed by direct transduction to D-leucine utilization with strain EO318 as donor. The ability to use 80  $\mu$ g of Dleucine per ml as a source of L-leucine was transduced together with derepressed leucinespecific transport activity (*lstR*). A preliminary uninterrupted mating with a suitably marked *leu* recipient revealed a pattern of distribution of *lstR* (identified both by D-leucine utilization and derepressed leucine transport) among other unselected markers that placed the locus between min 18 and 25 on the genetic map (data not shown). A multiply marked *lstR* F<sup>-</sup> recom-



FIG. 1. Simplified map of the E. coli chromosome showing the positions of relevant genes and points of origin and direction of transfer of Hfr and F-prime strains used in this study (10). Genetic symbols are those of Taylor and Trotter (21), except for lstR and livR in the expanded aroA region. The relative positions of livR, lstR, serC, pdxC, and aroA were determined from this study.

 

 TABLE 2. Segregation of two classes of D-leucineutilizing recombinants from the cross F15/EO311

 His<sup>+</sup> Gal<sup>+</sup> Dlu<sup>+</sup> Str<sup>s</sup> × AE20 F<sup>-</sup> His<sup>-</sup> Gal<sup>-</sup> Dlu<sup>-</sup> Str<sup>R</sup>

Selected	Dlu <sup>+</sup> recombinants <sup>a</sup>			<b>D-Leucine</b>	
marker (no. scored)	Туре	No. scored	L-Leucine transport <sup>o</sup>	threshold <sup>r</sup> (µg/ml)	
His <sup>+</sup> (200)	dlu	15	$0.05 \pm 0.02$	40	
	livR	9	$0.5 \pm 0.2$	150	

" Screened for ability to utilize 200  $\mu g$  of D-leucine per ml.

<sup>b</sup> Uptake of 0.75  $\mu$ M L-leucine, in nanomoles per minute per milligram of cells. The cells were grown in medium containing 25  $\mu$ g of L-leucine per ml. Error measurement is standard deviation.

<sup>c</sup> Plate concentration of p-leucine permitting growth of *leu* recombinants.

binant from this cross was used as a recipient in matings with Hfr's that transferred this region early. The gradient of transmission of  $lstR^+$ (scored by loss of D-leucine utilization and derepressed leucine transport) with respect to other unselected markers is shown in Fig. 2. No  $lstR^+$ recombinants were detected among 400  $trp^+$  selected recombinants when the donor was the Hfr KL99; this indicated that lstR was located distal to the point of origin of KL99, between min 20 and 22 on the genetic map (21), in the same general area as livR.

Cotransduction of *lstR* and *livR* with loci in the aroA region. Two-factor crosses were carried out by using donors grown on the transport mutants (Table 3). Reciprocal crosses were not possible since the transport phenotypes could not be reliably selected. Plate transport phenotypes of transductants were verified by L-leucine transport assay on at least 10 members of each phenotypic class. Crosses 1 to 4 show that livR is weakly linked to pyrD and closely linked to aroA and serC. One of the  $aro^+$ , livR transductants of cross 2 was further transduced to leucine auxotrophy by making use of the linkage of leu with araC, which is a selectable marker (2). The resulting leu, livR transductant showed marginal utilization of p-leucine (200  $\mu$ g/ml), confirming the identity of the latter character, azaleucine sensitivity, and derepressed L-leucine transport with the livR locus.

Crosses 5 and 10 show that lstR is also weakly linked to pyrD; however, anomalous results were obtained when linkage to *aroA* or *serC* was scored. In crosses 6, 7, and 8, no linkage with *aroA* or *serC* was detected when strain EO318 or AE30 was the donor. Twentyfive *aroA*<sup>+</sup> or *serC*<sup>+</sup> azaleucine-resistant transductants from these crosses were tested for derepressed leucine transport, on the possibility that azaleucine sensitivity and the derepressed transport phenotype might have segregated.



FIG. 2. Gradient of transmission of unselected markers in crosses of the Hfr strains KL983, B7, Hfr6, and KL99 (all wild type, Str<sup>s</sup>, and D-leucine nonutilizers [Dlu<sup>-</sup>]) with the recipient  $F^-$  strain AE30 (leu, trp, gal, his, lstR, [Dlu<sup>+</sup>], Str<sup>R</sup>). In each case an early transferred marker was selected in a 120-min uninterrupted mating: his<sup>+</sup> (KL983), trp<sup>+</sup> (B7 and KL99), gal<sup>+</sup> (Hfr6). Three hundred recombinants (open symbols) from each cross were scored for distal donor genes (open symbols) and loss of Dleucine utilization (closed symbols). The arrowhead indicates the point of origin of strain KL99; no Dlu<sup>-</sup> recombinants were scored from this cross.

However, no lstR transductants were detected. Cross 9, however, shows that when strain AE41 was the donor, 50% cotransduction of lstR and aroA was obtained. Since the biochemical properties (derepressed leucine-specific transport, elevated leucine-specific binding protein) of all three donor strains were similar, we have ruled out the possibility that the lstR mutation in strain AE41 arose de novo during direct transduction of this strain to *D*-leucine utilization. Rather, since strain EO318 was a spontaneous mutant (18), we suspect that the mutational event may have introduced local interference for crossover between closely linked markers in the aroA region and that the successive direct transductions that led to the construction of strain AE41 (Table 1) removed this interference. At present we are accepting the linkage relationships of strain AE41 as indicative of the map position of lstR. Accordingly, since both livR and lstR have similar cotransduction frequencies with nearby markers, they must themselves be closely linked.

Three-point crosses. To determine the order of *livR* with respect to the closely linked markers *aroA*, *serC*, *pdxC*, and *pyrD*, threepoint crosses were carried out by using the double mutant AE64 (*aroA livR*), constructed by screening *gal*<sup>+</sup> recombinants from a conjugation of strains KLF3/AE55 and AE49. This strain was transduced to leucine prototrophy since *D*-leucine and the aromatic amino acids reciprocally interfere with each other's utilization (see Materials and Methods). This strain was used as both donor and recipient for the crosses listed in Table 4. Cross 1 establishes that the order *livR aroA pyrD* is consistent with the transduction data (Fig. 3). Crosses

Cross —	Relevant genotype			Percentage with unselected	
	Donor	Recipient	<ul> <li>Selected marker</li> </ul>	donor marker	
1	EO311 livR	PLS-73	pyrD+	3 (12/400) <sup><i>a</i></sup>	
2	EO311 livR	AB2829	aroA+	73.5 (294/400) <sup>a</sup>	
3	EO311 livR	KL282-1	$serC^+$	49 (196/400) <sup>a</sup>	
4	EO311 livR	KL282-1	serC+	52 (52/100) <sup>a</sup>	
5	EO318 lstR	PLS-73	$pyrD^+$	$2 (8/400)^a$	
6	EO318 lstR	KL282-1	serC+	$<1 (0/400)^a$	
7	EO318 lstR	AE49	aroA+	<1 (0/100) <sup>b</sup>	
8	AE30 lstR	AB2829	aroA +	$<1 (0/400)^{a}$	
9	AE41 lstR	AE49	aroA+	50 (37/74) <sup>b</sup>	
10	AE41 lstR	KL188	$pyrD^+$	$3.8 (15/400)^a$	
11	AT3143 $pdxC$	AB2829	aroA+	>95 (196/197)	
12	KL282-1 serC	AB2829	aroA+	73.5 (146/199)	

 TABLE 3. Cotransduction frequencies of markers in the aroA region

<sup>a</sup> Derepressed transport phenotype scored by sensitivity to 100  $\mu$ g of pL-azaleucine per ml.

<sup>b</sup> Derepressed transport phenotype scored as ability to utilize p-leucine at 200  $\mu$ g/ml.

<b>TABLE 4.</b> Distribution of progeny of three-factor crosses by tra
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Cross	Strains (relevant genotype) <sup>o</sup>	Selected marker (no. scored)	Recombinant class	Percent total trans- ductants
1	AE64 (aroA livR) D	<i>pyrD</i> <sup>+</sup> (388)	aroA+ livR+	90.7
	×		aroA $livR^+$	3.4
	KL188 $(pyrD)$ R		$aroA^+ livR$	<1 (0/388)
			aroA livR	5.9
2(A)	AE64 (aroA $livR$ ) D	$pdxC^{+}$ (393)	aroA+ livR+	3.8
	×	•	$aroA^+ livR$	1.8
	AE80 $(pd \ x \ C)$ R		aroA $livR^+$	30.7
	•		aroA livR	63.6
<b>(B)</b>	AT3143 $(pd \ x \ C)$ D	aroA + (392)	$pdxC^+ livR^+$	0.5
	×		$pdxC^+ livR$	3.8
	AE66 $(aroA \ livR)R$		$pdxC livR^+$	71.9
			pdxC livR	23.7
3(A)	AE64 (aroA livR) D	serC <sup>+</sup> (397)	aroA+ livR+	11.6
	×		$aroA^+ livR$	5.3
	KL282-1 (serC) R		aroA $livR^+$	37.3
			aroA livR	45.8
<b>(B)</b>	KL282-1 (serC) D	aroA + (296)	ser $C^+$ liv $R^+$	2.4
	×		$serC^+$ livR	22.3
	AE66 (aroA $livR$ )R		serC $livR^+$	51.0
			serC livR	24.3

<sup>a</sup> Possible orders of the appropriate loci are given in Fig. 3.

<sup>b</sup> Symbols: D, donor; R, recipient.

# ORDER 1





FIG. 3. Possible configurations of loci involved in the transductions of Table 4. Horizontal lines indicate donor or recipient chromosomes, depending on the direction of the cross. Lower-case letters and dashed lines designate crossover intervals between genes.

2(A) and 2(B) were reciprocal, treating *livR* as unselected donor marker or unselected recipient marker. The least frequent class of recombinants will be that requiring four crossover events (Fig. 3). The order indicated by the data is  $livR \ pdxC \ aroA$ . Similarly, crosses 3(A) and 3(B) showed a four-crossover class consistent only with the order livR serC aroA (Fig. 3). Since both pdxC and serC were middle markers with respect to aroA and livR, and pdxC was more closely linked to aroA than is serC (Table 3, crosses 11 and 12), the probable order of markers in this region is livR serC pdxC aroA pyrD (Fig. 4). This order does, however, predict a greater frequency of contransduction of serCwith livR than aroA with livR; the opposite is found (Table 3, crosses 2 to 4), indicating that livR might be somewhat deleterious in certain strain backgrounds and be underrepresented in progeny of some crosses.

Linkage of lstR and livR. Cotransduction data from Table 3 suggested that lstR and livRwere closely linked. A three-point cross was done, using the *aroA* livR mutant as recipient and lstR as donor;  $aroA^+$  transductants were selected and screened for lstR and livR by their differing capability to utilize D-leucine (Table 5). The data show that both mutant transport phenotypes were recovered, but no wild-type recombinants (D-leucine nonutilizers) were detected; this implies a very close linkage of the two mutant loci or an order (*livR lstR aroA*) that would require four crossovers to generate a wild-type recombinant. Reciprocal crosses are necessary to clarify this point. At present, the position of *lstR* is not uniquely defined (Fig. 4).

**Diploid analysis.** The F-prime F106, with point of origin between pyrD and pyrC (9), was found to cover the *aroA* region (Fig. 1). The *recA* derivatives of strains AE49 (*aroA*) and AE64 (*aroA livR*) were used as recipients for KLF6/KL181 in liquid matings, and the resulting *aroA*<sup>+</sup> merodiploids were tested for transport phenotype (Table 6). The heterozygous merodiploid *livR*<sup>+</sup>/*livR* showed wild-type repressed leucine transport; when the episome was lost, the strain regained both derepressed leucine transport and the *aroA* character, thereby demonstrating that the F-prime carried a dominant wild-type *livR*<sup>+</sup> allele. Accord-



**FIG. 4.** Order of loci in the aroA region. Numbers are cotransduction frequencies with the point of each arrow designating the selected marker. Numbers in parentheses are averages of reciprocal crosses reported by Cronan et al. (1). The precise position of lstR with respect to livR and other markers is not established. Not drawn to scale.

TABLE 5. Linkage relationships among the markers aroA, lstR, and livR

Cross (relevant genotype)	Selected marker (no. scored)	Recombinant class <sup>a</sup>	Percentage of total
Donor: AE41 (lstR)	aroA + (391)	lstR livR+	72
		$lstR^+ livR$	28
Recipient: AE64 (aroA livR)		lstR livR	b
		$lstR^+$ $livR^+$	<1 (0/391)

" lstR scored as ability to utilize 100  $\mu$ g of D-leucine per ml; livR scored as ability to use 200  $\mu$ g of D-leucine per ml by not 100  $\mu$ g/ml. The wild type is unable to use D-leucine at 200  $\mu$ g/ml.

<sup>b</sup> Unknown phenotype.

Strain	Relevant genotype	L-Leucine uptake <sup>a</sup>	Percentage of haploid activity	
AE59	aroA livR <sup>+</sup> recA	$0.068 \pm 0.002$		
KLF6/AE59	aroA+ livR+/aroA livR+ recA	$0.056 \pm 0.001$	82	
AE65	aroA livR recA	$0.663 \pm 0.064$		
KLF6/AE65	aroA+ livR+/aroA livR recA	$0.090 \pm 0.006$	6	
AE59-1 <sup>b</sup>	aroA $livR^+$ recA	$0.074 \pm 0.002$		
AE65-1 <sup>b</sup>	aorA livR recA	$0.460 \pm 0.034$		

TABLE 6. L-Leucine transport activity in haploid and merodiploid strains

<sup>a</sup> L-Leucine uptake measured at 0.75  $\mu$ M L-leucine, given as nanomoles per minute per milligram (dry weight) of cells. The cells were grown in minimal medium plus repressing (25  $\mu$ g/ml) amounts of L-leucine. Error measurement is standard deviation of triplicate samples.

<sup>b</sup> Merodiploids cured of the episome by acridine orange treatment (7).

ingly, livR corresponded to a negative controlling element responsible for a diffusible substance regulating expression of the LIV-I system. No gene dose effect was evident in the wild-type merodiploid  $livR^+/livR^+$ ; this does not rule out the possibility that structural genes may be present on the episome, since the strain was grown under repressing conditions where the wild type has almost undetectable levels of the LIV-I system.

Characterization of isogenic transport mutants strains. The isogenic strains AE62, 63, and 68 were constructed (Table 1) in order to provide well-marked starting strains for further acquisition of mutations in the branchedchain amino acid transport system and to determine the physiological effects of derepressed transport. The kinetic properties of the  $leu^+$ derivatives of these strains are shown in Table 7. The wild-type strain AE62 demonstrated the repression of the low- $K_m$  LIV-I system when the cells were grown with excess branched-chain amino acids; the level of the high- $K_m$  system (LIV-II) was not significantly affected. In comparison with the wild type, both the livR and lstR strains showed higher  $V_{max}$  values for leucine transport and higher binding protein levels regardless of growth conditions. Under repressing conditions, the lstR strain AE63 was derepressed in the leucine-specific binding protein and the livR strain was derepressed in the LIV-binding protein. When either of these strains was grown under nonrepressing conditions (Table 7), additional increases in the overall  $V_{max}$   $(V_1 + V_2)$  of leucine transport and binding protein levels occurred. The relative contributions of LIV-I and LIV-II in the case of AE63 (lstR) under repressing conditions could not be determined. A single  $K_m$  value was derived from a least-squares analysis of the kinetic data (Table 7). Under nonrepressing conditions, however, strain AE63 (lstR) showed biphasic kinetics with an apparent threefold increase in the LIV-II system over the wild-type or strain AE68 (livR) level.

TABLE 7. L-Leucine transport characteristics of isogenic strains with mutant and wild-type transport loci, grown in minimal MOPS medium in the presence and absence of the branched-chain amino acids

Growth condi- tions	Kinetic parame- ters <sup>a</sup>	Strain AE62 (wild type)	Strain AE63 (lstR)	Strain AE68 (livR)
L-Leucine, L- isoleucine,	K <sub>1</sub>	0.13	0.83	0.2
and L-va-	<i>K</i> <sub>2</sub>	2.60		2.5
line (each	V,	0.13		0.7
at 2.5 μg/	•		2.32	
ml)	$V_2$	0.80		0.9
	Leucine-binding	< 0.02	0.72 <sup>r</sup>	1.14
	protein			
No addition	K,	0.17	0.2	0.4
	$K_2$	1.7	3.7	5.0
		0.92	1.5	2.4
	$V_2$	0.62	1.9	0.7
	Leucine-binding protein	1.11	1.62	1.61

" Parameters resolved from biphasic Lineweaver-Burk plots as described in the text.  $K_1$  and  $K_2$  refer, respectively, to the apparent  $K_m$  values of the LIV-I and LIV-II leucine uptake systems (micromolar).  $V_1$ and  $V_2$  are the respective  $V_{max}$  values in nanomoles of L-leucine per minute per milligram (dry weight) of cells. Leucine-binding protein (nanomoles of L-leucine bound per milligram of crude shock fluid protein) was measured after osmotic shock treatment of logarithmically growing cells as described previously (15).

<sup>b</sup> Only one system detected.

<sup>c</sup> Determined to be 80% leucine-specific binding protein by resistance of L-leucine binding to L-isoleucine inhibition.

 $^d$  Determined to be >90% LIV-binding protein by sensitivity to L-isoleucine inhibition of L-leucine binding.

### DISCUSSION

The mutant loci lstR and livR apparently govern two different patterns of regulation of branched-chain amino acid transport. Although both show derepressed L-leucine transport as reported previously (18), the pattern of derepression of the livR mutant permits growth only on high levels (200  $\mu$ g/ml) of p-leucine, whereas that of lstR permits growth on at least a twofold lower concentration of the p-amino acid. Since the latter lesion leads to a striking increase of the leucine-specific binding protein whereas livR derepresses the LIV-binding protein, we tentatively conclude that p-leucine can enter the cell through the leucine-specific system. Although binding studies have not been reported for *D*-leucine with the leucine-specific binding protein, it is noteworthy that the best inhibitor of the leucine-specific protein is racemic trifluoroleucine (3); it is possible that a considerable portion of the inhibition of L-leucine binding is due to the p-isomer. It is also clear that mutant loci other than those affecting L-leucine transport lead to the Dlu<sup>+</sup> phenotype, since the original mutant strains EO311 and EO318 have been shown to segregate Dlu+ strains with wild-type transport properties. Our finding that the aromatic amino acids antagonize **D**-leucine utilization and vice versa suggests an affinity of D-leucine for an aromatic transport system. Harrison et al. (6) have shown that L-azaleucine enters by both the aromatic permease (aroP) and by the leucine uptake systems.

Both lstR and livR map at min 20 on the E. coli genetic map (21), a site different from the previously reported mutant locus affecting branched-chain amino acid transport or biosynthesis. Preliminary mapping studies involving crosses between the strains carrying the two mutations indicate that the loci are too closely linked to readily detect crossover between them; this result is complicated by the difference in cotransduction frequencies of lstR with aroA and serC, depending on the source of donor phage carrying lstR. The simplest explanation for this phenomenon is that the original mutant EO318 was generated by an event that caused interference with crossover near aroA, such as an inversion or insertion. During subsequent sequential transductions of lstR by direct selection for the marker this interference was lost, resulting in linkage patterns that do not differ substantially from those of livR.

The recent report of Wood (23) on the kinetic parameters of L-leucine transport in a different K-12 strain (strain 7) provides a source of comparison for our own results on mutant and wildtype isogenic strains. We have confined our kinetics to L-leucine concentrations where nonspecific leucine transport would not be seen (less than 5  $\mu$ M). Since we have not separated leucine-specific from LIV-I transport, (Table 7), we have combined Wood's separate values for the two systems reported in Table 2 of reference 23 into one high-affinity system for purposes of comparison. Our values for the wild-type strain yield 14% high-affinity activity (0.13/0.13 +0.80; Table 7) and 86% low-affinity (LIV-II) activity in cells under repressing conditions compared with 40 and 60% in Wood (Table 2 in reference 23). For derepressed cells, our values are 60% high-affinity transport (0.92/0.92 +0.62) and 40% LIV-II versus 80 and 20%, respectively, in Wood (23). Strain differences could easily account for these discrepancies, since we have found that different K-12 strains have varying basal levels of L-leucine transport (S. C. Quay and J. J. Anderson, unpublished data).

The data of Table 7 confirm that as a consequence of livR mutation the levels of both LIVbinding protein and the LIV-I transport system are elevated over the wild-type levels during branched-chain amino acid repression, as reported previously for the original livR mutant, strain EO312 (18). However, unlike the latter strain, the elevated levels in strain AE68 (livR) are derepressed further when the repressing amino acids are withdrawn, suggesting that in this new strain background partial repression can still be observed.

The level of the LIV-II system did not vary appreciably in either wild-type or livR strains as a result of branched-chain amino acid supplementation (Table 7). However, in strain AE63 (lstR) under nonrepressing conditions, approximately 60% of the increase in leucine transport over wild type was apparently due to an elevated LIV-II system. Whether or not this is also true of strain AE63 (lstR) under repressing conditions could not be determined, since the single  $K_m$  value obtained for leucine transport is intermediate between the LIV-I and LIV-II systems. Such an intermediate value may represent a mutationally generated new system or simply a fortuitous combination of two or more separate systems whose additive Lineweaver-Burk kinetics are linear within experimental error. The close linkage of lstR and livR, each governing different patterns of expression of branched-chain amino acid transport, suggests that the associated region is genetically complex.

The finding that livR behaves in merodiploids as a recessive character suggests that the wild-type allele  $livR^+$  is responsible for the production of a diffusible, negative control element that effects repression of the LIV-I system. Presumably this element is associated with the repressing signal generated by interaction of L- leucine with the leucyl-transfer ribonucleic acid synthetase (17).

Present work is directed to the acquisition of appropriate mutants to determine the location of structural genes for the binding proteins. Positions of loci determining other elements in branched-chain amino acid transport have been reported (5), but none of these map near the regulatory loci reported here.

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