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# Genetic Analysis of the Glutamate Permease in Escherichia coli K-12<sup>1</sup>

MENASHE MARCUS AND YEHESKEL S. HALPERN

Department of Bacteriology, The Hebrew University-Hadassah Medical School, Jerusalem, Israel

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The glutamate permeation system in Escherichia coli K-12 consists of three genes: gltC, gltS, and gltR. The genes gltC and gltS are very closely linked, and are located between the pyrE and tna loci, in the following order: tna, gltC, gltS, pyrE; gltR is located near the metA gene. The three glt genes constitute a regulatory system in which gltR is the regulator gene responsible for the formation of a repressor, gltS is the structural gene of the glutamate permease, and gltC is most probably the operator locus. The synthesis of glutamate permease is partially repressed in wild-type K-12 strains, resulting in the inability of these strains to utilize glutamate as the sole source of carbon. Derepression due to mutation at the gltC locus enables growth on glutamate as a carbon source both at 30 C and at 42 C. Temperature-sensitive gltR mutants capable of utilizing glutamate for growth at 42 C but not at 30 C were found to be derepressed for glutamate permease when grown at 42 C and partially repressed (wild-type phenotype) upon growth at 30 C. These mutants produce an altered thermolabile repressor which can be inactivated by mild heat treatment (10 min at 44 C) in the absence of growth.

In previous papers from this laboratory (9, 10, 17), evidence was presented showing that a permeability barrier was responsible for the inability of wild-type strains of Escherichia coli K-12 to grow on glutamate as the sole source of carbon and energy. The location of a gene (gltC) determining or regulating glutamate permeation was established by conjugation (17). The mutated allele of this gene,  $gltC^{\circ}$  (previously designated  $gltC^+$ ), enables the mutants to grow on glutamate, thanks to an increased rate of glutamate uptake and a higher capacity for glutamate accumulation, as compared with the wild-type strains carrying the gltC+ allele (previously named  $gltC^{-}$ ). On the basis of these data, gltC could be the structural gene of the glutamate permease or a gene which regulates the rate of synthesis of the glutamate permease.

The purpose of this report is to show that the glutamate permease system consists of at least three components: the structural gene, gltS, determining the synthesis of glutamate permease, and two regulatory genes, gltR and gltC. The gltR gene specifies the formation of a repressor which represses the synthesis of glutamate permease in wild-type strains; the gltR locus is not linked to the structural gene. The gltC

<sup>1</sup> This work was part of a Ph.D. thesis submitted by the senior author to the Senate of the Hebrew University.

locus determines the sensitivity of glutamate permease synthesis to the product of the gltR gene; gltC is very closely linked to the structural gene of glutamate permease.

## MATERIALS AND METHODS

Microorganisms. The strains used throughout this work were all derivatives of  $E.\ coli$  K-12. A list of the strains used is given in Table 1.

Growth media. The liquid minimal medium in which the bacteria were grown and the solid minimal medium used for the selection of recombinants have already been described (17). These media were supplemented with a carbon source as required (glucose, 0.2%; glutamate, 0.5%; lactose, 0.5%; mannitol, 0.5%; tryptophan, 0.3%; succinate, 0.5%; and xylose, 0.5%), amino acids (arginine, histidine, isoleucine, leucine, methionine, threonine, and valine, where required, 25  $\mu$ g of the L isomer per ml, except where otherwise stated), thiamine hydrochloride (where required, 1  $\mu$ g/ml), and thymine and uracil (where required, 20 µg/ml). Mating broth containing Casamino Acids and yeast extract was that described by Hayes (12). Transduction broth (T broth) contained, at pH 7.0: 1% tryptone (Difco), 0.5% yeast extract, 0.5% NaCl, and 5 × 10<sup>-3</sup> M CaCl<sub>2</sub> (the CaCl<sub>2</sub> was autoclaved separately). Solid medium for phage P1kc titration contained transduction broth supplemented with 0.1% glucose and was solidified with 1% agar for the bottom layer and with 0.6% agar for the top layer. Stocks were maintained as described earlier (17).

Growth experiments. Young bacterial cultures from

TABLE 1. Strains used

Designation	Mat-	Grow Gluta						Rel	evan	t gen	otype	a							
	type	30 C	42 C	gltR	gltC	gltS	arg	his	ilv	leu	met	thr	thi	pyr E	lac	mtl	tna	xyl	st
CS101 CS7 CS8 CS2 <sup>TC</sup> CS3 <sup>TC</sup> CS4 <sup>TC</sup> CS7/50 P678	Hfr Hfr Hfr Hfr Hfr Hfr F	- + + - - -	- + + + +	+ + + TL TL TL +	+ c c + + + c -	+++++	+		+	++++		++++	++++++	+	++++++	+	++ +	+	SSSSS
C600 GR103 DF1933 AB1450 KL141	F- F- F- F- F-	_ _ _ _	_ _ _ _	+++++	+ + + + + c	++++-	ЕСВН-	_		_	+ + <b>A</b> -	_		+		_	+	_	F F F

<sup>a</sup> The symbols used are based on the nomenclature of Demerec et al. (5). The symbols arg, his, ilv, leu, met, thr, thi, and pyrE designate genes which determine and regulate the synthesis of arginine, histidine, isoleucine-valine, leucine, methionine, threonine, thiamine, and uracil, respectively. Plus and minus signs indicate the ability or inability, respectively, to synthesize the amino acid, vitamin, or pyrimidine. A capital letter following the three-letter abbreviation indicates the specific gene or genes mutated among those sharing the same symbol. The genes lac, mtl, tna, and xyl determine and regulate the utilization of lactose, mannitol, tryptophan, and xylose, respectively. Plus and minus signs indicate, respectively, the ability and inability to utilize the given substrate. The genes gltC gltR, and gltS determine and regulate the synthesis of the glutamate permease. The gene gltC is most probably the operator gene in the glutamate permease system. Its gltC+ allele is present in the wild-type strains which do not grow on glutamate (Glut- phenotype). The gltC7 and gltC8 alleles are present in mutants which grow on glutamate (Glut<sup>+</sup> phenotype). The gene gltR is the regulator gene determining the synthesis of repressor, and gltR+ is the wild-type allele which determines the synthesis of an active repressor resulting in a Glut-phenotype in wild-type strains. The symbol gltRTL represents the mutated allele. It determines the synthesis of a thermolabile repressor and thus gives a Glut+ phenotype at high growth temperature (42 C) and a Glut phenotype at low growth temperature (30 C). The gene gltS is the structural gene of glutamate permease; gltS+ indicates the allele in wild-type strains which determines the synthesis of a permease with a high affinity for glutamate ( $K_{\rm m}=5$  to  $10\times10^{-6}$  M), and  $gltS^-$  indicates the mutated allele in mutant strains which do not grow on glutamate. This allele determines the synthesis of a permease with a low affinity for glutamate  $(K_m = 10^{-4} \text{ M})$ . The str genes determine the response to streptomycin (S and R indicate susceptibility and resistance, respectively).

fresh agar slants were inoculated into media containing 0.5% glutamate as the sole source of carbon (enriched according to the requirements of the strain used) in conical flasks provided with side arms to allow direct turbidity measurements. The bacteria were incubated in a shaking water bath at the desired temperature. Growth was followed by measuring the changes in optical density in a Klett-Summerson photoelectric colorimeter, with the use of filter no. 42. Growth rates were determined graphically from semilogarithmic plots of optical density versus time of incubation.

Mutagenesis and selection of mutants. Mutations were induced by treatment with ultraviolet light (UV) or with N-methyl-N'-nitro-N-nitrosoguanidine (NG; 1). The treated cultures were incubated overnight in test tubes containing nutrient broth in a shaking water bath at 37 C, to allow phenotypic expression. Glut<sup>TC</sup> mutants (capable of growing on glutamate at 42 C but not at 30 C) were selected from strain CS101

treated with UV as follows. A 5-ml amount of the bacterial culture was centrifuged and washed with saline. The cells were suspended in 100-ml conical flasks to a density of  $2\times 10^7$  per ml in 40 ml of minimal medium which contained glutamate as the sole source of carbon and energy and was enriched with methionine. The flasks were incubated in a shaking water bath at 42 C until the bacterial density increased to  $2\times 10^9$  cells per ml. Only one mutant was selected from each culture. Forty independently isolated mutants were purified twice on plates of glutamate-minimal medium, and their ability to grow on glutamate at 42 C and at 30 C was tested by the replica method. Three mutants grew on glutamate at 42 C but not at 30 C.

Glut<sup>-</sup> mutants (unable to grow on glutamate at any temperature) were selected from strains CS7 and CS8 treated with UV or NG as follows. Samples (5 ml) from each of the two cultures were centrifuged, washed with saline, and resuspended in succinate-

minimal medium, enriched with methionine, to a density of 2 × 10<sup>7</sup> cells per ml. The cultures were grown in this medium to a density of  $2 \times 10^9$  cells per ml to reduce the frequency of auxotrophic mutants. The bacteria were centrifuged, washed, and resuspended in glutamate-minimal medium enriched with methionine. The cultures were treated with penicillin according to the methods of Gorini and Kaufman (7) and Lubin (16), as described by Dar-Guttman et al. (3), to enrich for Glut mutants. After this enrichment, the bacteria were grown in a succinate-minimal medium, enriched with methicnine, to increase the total number of mutants, and another penicillin cycle was performed. Appropriate dilutions were spread on petri dishes with Tryptose Blood Agar Base. These plates served as master plates for replication onto minimal glucose, succinate, and glutamate media, enriched with methionine. Only colonies which grew on glucose and succinate media but did not grow on the glutamate medium were isolated and further studied.

Mating experiments. Matings were performed as described by us earlier (17).

Transduction experiments. Phage P1kc was used. Lysates were prepared as follows. Logarithmic cultures of the donor strains, at a density of 108 cells per ml in T broth containing 5  $\times$  10<sup>-3</sup> M CaCl<sub>2</sub> and 0.1% glucose, were infected with the phage at a multiplicity of 0.1 and incubated with aeration for 3 hr at 37 C. At that time, considerable lysis could be observed. A few drops of chloroform were added to the culture, and incubation was continued for 10 min more. The bacterial debris was removed by centrifugation at low speed  $(5,000 \times g)$  for 10 min. The supernatant liquid was decanted and stored at 4 C in sterile test tubes to which one drop of chloroform was added. By this procedure, lysates with a titer of  $5 \times 10^{10}$  infective particles per ml were easily obtained. E. coli C600 served as the indicator strain. Transduction was performed as follows. The recipient strains were grown in T broth with aeration at 37 C to a density of  $5 \times 10^8$  cells per ml. CaCl<sub>2</sub> (5 × 10<sup>-3</sup> M) was then added, and incubation was continued for 5 min longer. Phage was added at a multiplicity of 1, and the suspension was incubated at 37 C without shaking for 20 min. The bacteria were then centrifuged in the cold and resuspended in sterile phosphate buffer (pH 7.0, 0.1 M). Samples (0.1 ml) of appropriate dilutions were spread on selective media. The recipient strain and the phage suspension were also spread separately on the same media to serve as controls. The plates were kept for 2 hr at room temperature and then incubated at the desired temperature as specified.

Isolation of recombinants and scoring for unselected markers. Recombinants were isolated and unselected markers were scored for as previously described (17).

Uptake of <sup>14</sup>C-glutamate. Glutamate uptake was studied by two different methods.

In the first method, incorporation of <sup>14</sup>C-L-glutamate (uniformly labeled, The Radiochemical Centre, Amersham, Bucks, England) into the proteins of logarithmically growing cultures was determined ac-

cording to the method of Ames (2) as described by

Halpern and Lupo (10), with some modifications. The

bacteria were grown to the logarithmic phase at 30 or 42 C. Cultures grown at 42 C were transferred to a shaking water bath at 30 C 90 sec before the addition of the radioactive glutamate. The incorporation mixtures were incubated at 30 C for 3 min, at which time trichloroacetic acid was added and the tubes were heated at 90 C for 10 min. The bacteria were filtered, washed, and dried; the radioactivity was determined as previously described (10). Rates were expressed as micromoles of glutamate incorporated per milligram (dry weight) of bacteria per hour.

The second method involved measuring the accumulation of 14C-glutamate by nongrowing cells. This was done according to the method of Kessel and Lubin (15) as earlier described (9, 10), with some modifications. The bacteria were grown to a density of 7 × 10° cells per ml in minimal medium containing 0.5% succinate (enriched as required) and were incubated in a shaking water bath at 30 or 42 C. Chloramphenicol was added to a final concentration of 200 µg/ml, and incubation was continued for 30 min longer. The uptake experiments were performed at 25 C. The cells were incubated in the presence of radioactive glutamate for 6 min. The incubation mixture was then filtered, washed, dried, and counted as described by Halpern and Even-Shoshan (9). The intracellular concentration of radioactive material was expressed as micromoles of glutamate per milliliter of intracellular water.

Determination of radioactive glutamate in the internal pool. Radioactive glutamate in the internal pool was determined according to the method of Halpern and Lupo (10). These authors found that about 75% of the radioactivity in the wild-type strain CS101 and in the glutamate-utilizing mutants tested was due to free glutamate. In the present work, we examined the radioactive pool in strain CS2<sup>TC</sup>, which grows on glutamate at 42 C but not at 30 C, and we also found that more than 75% of the radioactive material was free glutamate.

Determination of the rate of exit of <sup>14</sup>C-glutamate. The rate of exit of radioactive glutamate from preloaded chloramphenicol-treated cells was determined according to the method described by Halpern (8) with some modifications. The cultures were grown in a succinate (0.5%)-minimal medium enriched as required, at 30 or at 42 C. The cells were preloaded with <sup>14</sup>C-L-glutamate at 25 C for 6 min. The exit reaction was also carried out at 25 C. The bacterial densities in the uptake mixtures varied with the different strains, as follows [mg (dry weight) per ml]: CS101, 3.2; CS7, 0.8; CS7/50, 3.2; CS2<sup>TC</sup> grown at 42 C, 0.8.

Enzyme assays. Cell-free extracts were used for the determination of enzyme activities. The procedures used for the preparation of extracts and for measuring the activities of glutamate dehydrogenase and glutamate-oxalacetate transaminase were as described by Halpern and Lupo (10). Aspartase activity was determined from the amount of ammonia formed from aspartate in the following reaction mixture: L-aspartate, 0.02 M, adjusted to pH 7.5; MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.001 M; potassium phosphate buffer, pH 7.5, 0.05 M; bacterial extract equivalent to 1 to 4 mg of protein;

total volume, 2 ml. The reaction mixture was incubated for 10 to 30 min at 30 C. The reaction was stopped by the addition of trichloroacetic acid to a final concentration of 3.3%. The precipitate was removed by centrifugation, and ammonia was determined on samples of the clear supernatant liquid by nesslerization according to the method of Halpern and Umbarger (11). Enzyme activities were expressed in micromoles of substrate transformed per milligram of protein per hour.

#### RESULTS

Isolation of temperature-conditional glutamate-utilizing mutants. We suggested earlier (17) that glutamate permease synthesis in *E. coli* is regulated by a genetically controlled mechanism of repression. Permease synthesis is partly repressed in the wild-type strains, resulting in their inability to utilize glutamate as the sole source of carbon, and is derepressed in the glutamate-utilizing mutants. On this assumption, one should be able to obtain mutants with lesions in the regulator gene specifying the synthesis of the repressor, which would produce a thermolabile repressor. Such mutants could be readily detected by their ability to grow on glutamate at high, but not at low, temperatures.

Logarithmic cultures of strain CS101 were treated with UV and mutants were selected capable of growing in a glutamate-minimal medium enriched with methionine, at 42 C. Forty independent mutants were isolated and tested for their ability to grow on glutamate at 30 C. Three of these mutants, CS2<sup>TC</sup>, CS3<sup>TC</sup>, and CS4<sup>TC</sup>, were found to have acquired the Glut<sup>TC</sup> phenotype, i.e., the ability to utilize glutamate at 42 C but not at 30 C. The ability

of these mutants to grow on glucose or succinate at 30 C was unimpaired.

Determination of 14C-L-glutamate uptake and exit in strains CS101, CS7, CS8, and CS2<sup>TC</sup> grown at 30 C and at 42 C. The uptake and exit of <sup>14</sup>C-L-glutamate were measured as described in Materials and Methods. The results shown in Table 2 clearly demonstrate that the rate of exit was similar in the three strains examined. The exit of glutamate was about 2.5 times faster in bacteria grown at 42 C than in those grown at 30 C. The affinity of the glutamate permease for glutamate was determined by measuring glutamate incorporation into the proteins of exponentially growing cultures and accumulation by nongrowing cells. The two methods gave similar results. The  $K_m$  for glutamate in the wildtype CS101, in the Glut<sup>+</sup> mutant CS8, and in the Glut<sup>TC</sup> mutant CS2<sup>TC</sup> grown at either 30 or 42 C was 10<sup>-5</sup> M and differed only slightly in strain CS7 (5  $\times$  10<sup>-6</sup> M). However, the rate of uptake of glutamate and the maximal glutamateconcentrating capacity differed greatly in the four strains tested. The glutamate-utilizing mutants CS7 and CS8 showed a fivefold higher rate of glutamate uptake and accumulated four to six times more glutamate than did their Glutparent strain CS101, regardless of the temperature of growth. On the other hand, glutamate uptake in strain CS2<sup>TC</sup> exhibited a dramatic dependence on growth temperature. When grown at 42 C, CS2TC took up glutamate at a fast rate similar to those of the Glut+ mutants; however, CS2<sup>TC</sup> cultures grown at 30 C took up glutamate at a rate one-fifth of that of the Glut+ mutant,

Table 2. Determination of glutamate permease activity and glutamate exit in cultures grown at 30 C and at 42 C

		Uptake						
Strain	Growth temp	Glutamate incor	porated into proteins	Internal p	Exit half-life			
	·	V <sub>max</sub> <sup>a</sup>	K <sub>m</sub>	MCC <sub>p</sub>	K <sub>m</sub>	-		
	С		м	· · · · · · · · · · · · · · · · · · ·	м	min		
CS101	30	0.070	1 × 10 <sup>-5</sup>	4.0	1 × 10-5	21		
CS101	42	0.070	$1 \times 10^{-5}$	4.0	1 × 10 <sup>-5</sup>	8		
CS7	30	0.330	5 × 10 <sup>-6</sup>	16.0	5 × 10 <sup>-6</sup>	21		
CS7	42	0.320	5 × 10 <sup>-6</sup>	13.5	5 × 10-6	8		
CS8	30	_	1 × 10 <sup>-5</sup>	25.0	1 × 10-5	_		
CS8	42	_	1 × 10 <sup>-5</sup>	15.0	1 × 10 <sup>-5</sup>	_		
CS2TC	30	0.075	1 × 10 <sup>-5</sup>	4.5	1 × 10-1	21		
CS2TC	42	0.320	1 × 10 <sup>-5</sup>	14.6	1 × 10 <sup>-5</sup>	8		

<sup>&</sup>lt;sup>a</sup> V<sub>max</sub> is expressed in micromoles of <sup>14</sup>C-L-glutamate per milligram (dry weight) of bacteria per hour. <sup>b</sup> MCC (maximal concentrating capacity) is expressed in micromoles of <sup>14</sup>C-L-glutamate per milliliter of intracellular water.

similar to the rate characteristic of the Glutparent strain CS101.

Growth of CS2<sup>TC</sup> on glutamate at different temperatures. The growth rate of CS2<sup>TC</sup> in a glutamate-minimal medium enriched with methionine, at 42 and 30 C, and the changes in the rate of growth after temperature shifts are given in Fig. 1. The results demonstrate a close correlation between the ability of CS2<sup>TC</sup> to grow on glutamate at 42 C, its inability to grow at 30 C, and the rates of glutamate uptake and the maximal glutamate-concentrating capacities at these temperatures (see Table 2).

Mapping of the gene determining the Glut<sup>TC</sup> phenotype in strain  $CS2^{TC}$  by interrupted mating. Preliminary experiments showed that it was possible to obtain  $Glut^{TC}$  recombinants with  $Hfr\ CS2^{TC}$  as the donor and  $F^-\ C600$  or  $F^-\ P678$  as the recipient. The results of an interrupted mating between  $Hfr\ CS2^{TC}$  and  $F^-\ C600$  are given in Fig. 2. These data clearly show that the mutation which enables growth on glutamate at a high temperature  $(gltR^{TL})$  does not map at and

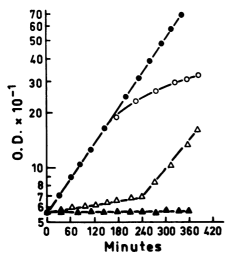


Fig. 1. Growth of strain  $CS2^{TC}$  in a glutamate-minimal medium at 30 and 42 C.  $CS2^{TC}$  was grown for 10 generations in a glutamate-minimal medium at 42 C to attain steady-state conditions. The culture was then divided into two flasks; one was left at 42 C and the other was transferred to 30 C. The culture at 42 C continued to grow exponentially. At 30 C, growth slowed down after 140 min and stopped entirely after three generations. Part of this culture was then transferred from 30 to 42 C, and growth was followed turbidimetrically as described in Materials and Methods. Symbols:  $\blacksquare$ , culture grown at 42 C;  $\bigcirc$ , culture transferred at zero time from 42 to 30 C;  $\triangle$ , culture grown at 30 C.

is not even linked to the *gltC* locus, which maps near the *mtl* gene (17). Similar results were obtained in experiments in which CS3<sup>TC</sup> and CS4<sup>TC</sup> served as donors. Several recombinants were tested for their ability to take up <sup>14</sup>C-L-glutamate after growth at 30 and at 42 C. All the recombinants tested exhibited a pattern similar to that of the donor strain CS2<sup>TC</sup>. The somewhat lower number of Glut<sup>TC</sup> recombinants as compared with the number of Thi<sup>+</sup> recombinants probably resulted from the greater interference with integration on the minimal selective medium for Glut<sup>TC</sup> recombinants (4, 17).

Mapping of the gltR<sup>TL</sup> locus by recombination analysis. The results of the interrupted mating indicated that the gltR<sup>TL</sup> locus entered before the thi locus. This observation was verified by an unselected-marker frequency analysis (Table 3), in which the difference in linkage within each pair of markers, depending on which of the two markers is selected for, is a consequence of the gradient of transmission (13).

The distance in time units between *gltR* and the *thi* locus as determined by the interrupted mating experiment was about 3 min (Fig. 2). This places the *gltR* gene on the *E. coli* K-12 chromosome map (19) near the *metA* locus (Fig. 3). A three-factor-cross analysis was therefore performed in which a prototroph derivative of CS2<sup>TC</sup> served as the donor and F- DF1933

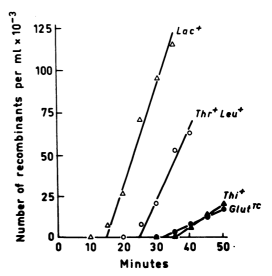


Fig. 2. Mapping of the gltR gene by interrupted mating. A temperature-sensitive glutamate-utilizing mutant,  $Hfr\ CS2^{TC}$ , served as the donor and strain F-C600 was used as the recipient. Glut $^{TC}$  recombinants were selected on a glutamate-minimal medium enriched with methionine at 42 C.

Table 3. Unselected-marker frequency analysis by conjugation of the linkage relationships between the thi and gltR loci in E. coli K-12a

Selected	No. of re-	Unselected	No. of re-	Linkage
marker	combinants	marker	combinants	
gltR <sup>TL</sup> thi <sup>+</sup>	125 111	thi <sup>+</sup> gltR <sup>TL</sup>	45 106	% 36.0 95.5

<sup>a</sup> Hfr CS2<sup>TC</sup> and F<sup>-</sup> C600 served as donor and recipient, respectively.

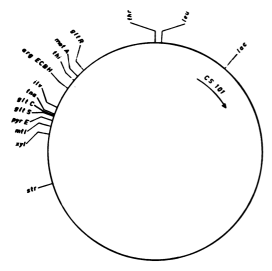


Fig. 3. Escherichia coli K-12 chromosome drawn to scale according to Taylor and Trotter (19) showing the positions of gltC, gltS, and gltR.

(arg<sup>-</sup>, met<sup>-</sup>) served as the recipient. Arg<sup>+</sup> recombinants were selected, and Met<sup>+</sup> and Glut<sup>TC</sup> recombinants were scored by the replica method. The results shown in Table 4 establish the following order of genes: argECBH, metA, gltR.

Linkage analysis by transduction between gltR<sup>TL</sup>, argECBH, metA, and thi. Phage Plkc was grown on strain CS2<sup>TC</sup>. The recipient strains were DF1933 and P678. The results shown in Table 5 confirm the order of genes established by conjugation.

Thermolability of the gltR<sup>TL</sup> product. The results described so far provide tentative support for our assumption that the glutamate permease synthesis is controlled by a mechanism of repression (17), in which gltR plays the role of the repressor-forming gene. In wild-type strains which do not grow on glutamate, the synthesis of the permease is partly repressed. The behavior of the Glut<sup>TC</sup> mutants would then be due to the production of a thermolabile repressor, which

is destroyed at high growth temperatures, resulting in derepression of glutamate permease synthesis and in the acquisition of the ability to grow on glutamate at 42 C.

Evidence pointing to the gltR product as a thermolabile repressor was obtained in the following experiment.  $CS2^{TC}$  and CS101 were grown in a succinate-minimal medium enriched with methionine, in a shaking water bath at 30 C. After 10 doublings (18), the cultures were diluted into fresh medium to a density of 1.5  $\times$ 107 cells per ml and further incubated for 90 min. The two cultures were then divided each in two equal portions and the four flasks were immediately placed in a shaking water bath at 30 C. Samples (8 ml) were withdrawn from each flask at the times indicated in Fig. 4. The rate of glutamate uptake was measured by mixing each sample with 2 ml of <sup>14</sup>C-L-glutamate (1.5 µc per  $\mu$ mole) to a final concentration of 5  $\times$  10<sup>-6</sup> M and incubating the mixture for 3 min at 30 C. Trichloroacetic acid was then added to a final concentration of 5%, and the amount of radioactive material incorporated into protein was determined. After two samples were taken from each of the four flasks, the remaining cultures were filtered on a membrane filter (47 mm in diameter,  $0.45-\mu m$  pore size), and the filtered growth media were returned to the respective flasks. The cells were washed five times with 10 ml of 0.06 м phosphate buffer (pH 7.0), prewarmed to 30 C, and resuspended in 3 ml of buffer. One suspension of each strain was incubated at 44 C and the other at 30 C for 10 min and transferred back to its growth medium at 30 C (18). The cultures were incubated, sampled, and tested for glutamate uptake for a further 80 min. As shown in Fig. 4, heating of CS2<sup>TC</sup> for 10 min at 44 C in the absence of growth increased the differential rate of glutamate per-

Table 4. Three-factor-cross analysis of the gltR<sup>TL</sup> gene in E. coli K-12 by conjugation<sup>a</sup>

Selected marker	Unselected markers <sup>b</sup>	No. of recom- binants	Percentage of recom- binants
argECBH+	metA+, gltRTL	89	68.5
argECBH+	metA+, gltR+	15	11.5
argECBH+	metA-, gltR+	26	20.0
argECBH+	metA-, gltRTL	2	1.5
J	, ,		1

<sup>a</sup> Hfr CS2<sup>TC</sup> and F- DF1933 served as donor and recipient, respectively.
 <sup>b</sup> The gltR<sup>TL</sup> recombinants were selected at 38

<sup>b</sup> The gltR<sup>TL</sup> recombinants were selected at 38 C because DF1933 did not grow well at 42 C. At 38 C, growth of Glut<sup>TC</sup> mutants on glutamate is slower than at 42 C and recombinant colonies appeared only after 4 days of incubation.

TABLE 5. Determination of	linkage between	gltR, metA,	and argECBH by transduction
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Expt no.	Donor strain	Recipient strain	Selected marker	No. of recombinants	Unselected markers	No. of recombinants	Linkage
1	CS2TC	DF1933	argECBH+	156	metA <sup>+</sup>	25	16.0
2	CS2TC	DF1933	argECBH+	156	gltR <sup>TL</sup>	0	<0.6
3	CS2TC	DF1933	metA+	469	argECBH <sup>+</sup>	140	29.8
4	CS2TC	DF1933	metA+	469	gltR <sup>TL</sup>	12	2.6
5	CS2TC	DF1933	metA <sup>+</sup>	469	argECBH+, gltRTL	0 0	<0.2
6	CS2TC	P678	thi <sup>+</sup>	150	gltRTL		<0.6
7	CS2TC	P678	gltR <sup>TL</sup>	150	thi+		<0.6

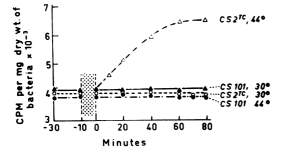


Fig. 4. Thermolability of the glutamate permease repressor in the temperature-sensitive glutamate-utilizing mutant  $CS2^{TC}$ .

mease synthesis during subsequent growth at 30 C. This was manifested by the continuous increase, for about 60 min, in the specific activity of the permease in CS2TC heated at 44 C. In contrast, a CS2<sup>TC</sup> culture heated at 30 C and CS101 cultures heated at either 30 or 44 C exhibited low specific activities which remained constant throughout the entire experiment. It is of interest to point out that the highest specific activity of glutamate permease in CS2TC grown at 30 C after a brief treatment at 44 C was only about 1.5-fold higher than that of the control cultures, whereas CS2TC cells grown for 10 generations at 44 C exhibited a rate of glutamate uptake four times as high as that of cells grown at 30 C (see Table 2). These results can be best explained by the production of a thermolabile repressor by the gltRTL allele of the repressorforming gltR gene. Destruction of the repressor by brief heating results in a burst of permease synthesis at a high, derepressed rate. Accumulation of new repressor molecules following the resumption of growth at the lower temperature brings about a gradual reestablishment of repression.

Isolation of Glut<sup>-</sup> mutants from the Glut<sup>+</sup> strain CS7 (gltC7). The gltC locus had been mapped by interrupted mating with HfrC Glut<sup>+</sup> mutants serving as donors and the P678 wild-type

strain as the recipient (17). Since no Glut+ recombinants appeared until 6 to 7 min after the time of entry of the gltR locus, it would seem that the gltC mutants examined carried a normal wild-type gltR allele. Moreover, as shown in Table 9, mutant CS7 (gltC°) as the donor in a transduction experiment yielded Glut<sup>+</sup> transductants at a high frequency  $(1.6 \times 10^{-6})$ . Since gltR and gltC are too far apart to be cotransduced, these Glut+ transductants must have inherited the gltR allele of the recipient, which was wild type. The fact that the gltC mutants and recombinants are derepressed in spite of the presence of a normal gltR gene suggests that the gltC mutations may be of the operator-constitutive type (14). That the gltC mutants carry mutations in a regulatory rather than in the structural gene of glutamate permease is also supported by the finding that the affinity of the permease of these strains for glutamate is the same as in the wild-type parent. One should be able to find structural gene mutants among the apparent Glut revertants of any Glut mutant. If indeed gltC is the operator gene, then the mutations in the structural gene of glutamate permease should be closely linked to the gltC

Cultures of E. coli K-12 CS7 were treated with nitrosoguanidine. Mutants that lost the ability to grow on glutamate, but still grew normally on glucose and succinate, were isolated from the mutagenized cultures after two cycles of penicillin treatment. The mutants were tested for the activities of aspartase, glutamate dehydrogenase, and glutamate-oxalacetate transaminase, for the rate of 14C-L-glutamate uptake, for the affinity of the glutamate permease for glutamate, and for the rate of glutamate exit from preloaded cells. Several mutants of independent origin were found to have an altered glutamate permease, as manifested by a decrease in its affinity for glutamate and by the lower rate of glutamate uptake. One such mutant, CS7/50, is described in Table 6. One can see that the activity of the glutamate permease in this mutant is about one-

1125

			En						
Strain	Growth temp			Accumulation of glutamate <sup>b</sup>		Exit, half-life	GDH¢	GOT⁵	Aspartase <sup>d</sup>
		$V_{\mathrm{max}}$	K <sub>m</sub>	мсс	K <sub>m</sub>				
	С		М		м	min			
CS7	30	0.33	5 × 10 <sup>-6</sup>	16.5	5 × 10 <sup>-6</sup>	21	25.0	38.0	2.8
CS7	42	0.32	5 × 10 <sup>-6</sup>	13.5	5 × 10 <sup>-6</sup>	8	25.0	39.0	3.0
CS7/50	30	0.11	1 × 10 <sup>-4</sup>	5.5	1 × 10-4	25	26.0	40.0	2.7
CS7/50	42	0.12	1 × 10 <sup>-4</sup>	5.0	1 × 10-4	8	25.0	40.0	3.2

<sup>&</sup>lt;sup>a</sup> The rate of glutamate uptake was expressed in micromoles of <sup>14</sup>C-L-glutamate per milligram (dry weight) of bacteria per hour.

third of that of the parent strain and the  $K_{\rm m}$  for glutamate of the mutant permease is 20-fold higher than that of the parent CS7 strain. The altered affinity of the permease suggests that the mutation in CS7/50 occurred in the structural gene of the glutamate permease.

Mapping of the CS7/50 mutation by transduction. The donor strains were prototroph (Met<sup>+</sup>) derivatives of CS101, CS2<sup>TC</sup>, and CS7. The recipient was CS7/50. Glut<sup>+</sup> and Met<sup>+</sup> transductants were selected. The *met* locus served as an outside marker for comparison of transduction frequencies. If gltC and gltS were closely linked genes, CS7 as the donor should give Glut<sup>+</sup> transductants at a much higher frequency than would CS101 or CS2<sup>TC</sup>. On the other hand, if there were no linkage between the two genes, all three of the transductions should give similar results. The results presented in Table 7 show that gltC and gltS are very closely linked.

Further mapping of gltC and gltS by an unselected-marker frequency analysis by transduction, involving known adjacent markers, was performed. CS7 served as the donor, and P678, GR103, AB1450, and KL141 were the recipients. The results shown in Table 8 are in agreement with those of our previous mapping experiments (Table 7; see also reference 17). On the basis of the known linkage of the tna, pyrE, and ilv loci (6, 19) and the linkage of tna, gltC, gltS, and pyrE obtained here, we concluded that gltC and gltS were located between pyrE and tna, very close to the tna locus.

The order of gltC and gltS with reference to the tna locus was established by transduction between CS7/50 (gltS-, gltCc) as the donor and GR103 (tna<sup>-</sup>) as the recipient. A two-pointcross experiment with CS7 (gltS+, gltCc) as the donor and GR103 as the recipient was performed for comparison of the frequencies of Glut+ transductants. The results presented in Table 9 show that the order is gltS, gltC, tna, since this arrangement requires a double crossing-over in order to obtain Tna- Glut+ transductants in experiment 1, whereas the alternative order, gltC, gltS, tna, would necessitate a quadruple crossing-over event, which is by far too rare to account for the high percentage of cotransduction observed between gltC and tna (72.2%). Another point of significance is the 40-fold higher frequency of Glut+ transductants obtained with strain CS7 as the donor, as compared with the cross in which CS7/50 served as the donor. The lower frequency observed in the latter case was obviously due to the forced crossing-over between the very close gltS and gltC loci.

## **DISCUSSION**

Wild-type strains of E. coli K-12 are unable to grow on glutamate as the sole source of carbon. However, glutamate-utilizing mutants of this organism have been readily isolated, after treatment with different mutagenic agents (10, 17). Earlier studies from this laboratory showed that the transport of glutamate into the cell was the limiting factor in the utilization of this compound as an energy and carbon source for growth. Both wild-type and glutamate-utilizing E. coli strains were shown to be capable of actively transporting

<sup>&</sup>lt;sup>b</sup> Maximal glutamate-concentrating capacity (MCC) was expressed in micromoles of <sup>14</sup>C-L-glutamate per milliliter of intracellular water.

<sup>&</sup>lt;sup>c</sup> GDH and GOT were determined on extracts prepared from cultures grown in a glucose-minimal medium enriched as required.

<sup>&</sup>lt;sup>d</sup> Aspartase activity was determined on extracts prepared from cultures grown in a succinate-minimal medium enriched as required.

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Table 8. Unselected-marker frequency analysis by transduction of the linkage between mtl, xyl, gltC, gltS, tna, and ilva

Donor strain	Recipient strain	Selected marker	No. of transductants	Unselected marker	No. of transductants	Linkage
						%
CS7	P678	xyl+	150	gltC7	0	< 0.7
CS7	P678	mtl+	150	gltC7	0	< 0.7
CS7	GR103	gltC7	140	tna+	100	71.4
CS7	AB1450	gltC7	100	ilv+	0	<1.0
CS7	KL141	PyrE+	175	gltS+	31	17.5

<sup>&</sup>lt;sup>a</sup> All of the Glut<sup>+</sup> transductants tested showed a high glutamate permease activity and a high permease affinity for glutamate.

TABLE 9. Ordering of the gltC, gltS, and tna loci by transduction

Expt. no.	Donor strain	Recipient strain	Frequency of transduction to Glut <sup>+2</sup>	No. of ina <sup>+</sup> /no. of transductants examined (cotransduction of ina <sup>+</sup> )	gltC- tna linkage
1 2	CS7/50 CS7	GR103 GR103	$4 \times 10^{-7} (180)$ $1.6 \times 10^{-6} (500)$	130/180 100/140	% 72.2 71.4

<sup>&</sup>lt;sup>a</sup> The frequency of transduction was determined by relating the number of transductants to the number of infective particles in the transduction mixture. The numbers in parentheses indicate the number of transductants counted. All of the Glut<sup>+</sup> transductants tested showed a high glutamate permease activity and a high permease affinity for glutamate.

the amino acid across the cell membrane. However, the rate of uptake and the capacity for accumulating glutamate were several-fold higher in the mutants as compared with wild-type strains which were unable to grow on glutamate (9, 10, 17).

Interrupted-mating experiments performed on four glutamate-utilizing mutants independently isolated from *E. coli* K-12 CS101 and one mutant of the *E. coli* K-12 Hayes strain showed that in all these cases the mutation mapped near the *mtl* locus (17). The wild-type allele of Hfr CS101 (Glut<sup>-</sup>) was also mapped near the mannitol gene.

Although these studies did not provide any direct evidence on the nature of the glutamate

permeation gene affected by these mutations, we tentatively assumed it to be a control gene rather than the structural gene of glutamate permease. Our assumption was based on the observation that the affinity of the permease for glutamate in all of the mutants examined was practically the same as in the wild-type parent strains.

On this assumption, the low level of permease activity in the wild-type strains would be due to partial repression of glutamate permease synthesis. Mutations in the control gene (designated as gltC) result in derepression of glutamate permease synthesis and thus enable the cell to take up glutamate at a rate sufficient to support growth.

If this interpretation is correct, one should be

<sup>&</sup>lt;sup>a</sup> The frequency of transduction was determined by relating the number of transductants obtained to the number of infective particles in the transduction mixture. The numbers in parentheses indicate the total number of transductant colonies found.

<sup>&</sup>lt;sup>b</sup> A number of Glut<sup>+</sup> transductants were isolated and tested for glutamate uptake and permease affinity for glutamate. All of the transductants tested had a high affinity for glutamate and exhibited a high rate of uptake, similar to that of strain CS7.

able to find mutations in the repressor-forming gene that lead to temperature-sensitive regulation of glutamate permease synthesis. Mutants producing a thermolabile repressor, or in which the synthesis of active repressor is prevented at high temperatures (18), should be able to grow on glutamate only at high temperatures. One would also expect to find Glut—mutants in which the mutation occurred in the structural gene of glutamate permease. In at least some of these mutants, a change in the affinity of the permease for glutamate would be expected.

The two types of mutants predicted by the repression hypothesis were indeed found, as described in the Results section (Tables 2 and 6, and Fig. 1). Three temperature-sensitive glutamate-utilizing mutants were independently isolated from the wild-type CS101 strain. As predicted, these mutants grew readily on glutamate at 42 C but could not utilize glutamate for growth at 30 C. That the mutation indeed affected glutamate uptake was clearly demonstrated by the fact that cultures of the temperature-sensitive (CS2<sup>TC</sup>) mutant grown at 42 C exhibited a high rate of glutamate uptake and a high capacity for glutamate accumulation, similar to those of the temperature-insensitive Glut+ mutants discussed above. However, when grown at 30 C, CS2TC showed low glutamate permease activity similar to that of the wild-type parent, which does not grow on glutamate at all. It is significant that the affinity of the CS2TC permease for glutamate remained the same as in the parent strain, and was not affected by the growth temperature. The rates of glutamate exit from preloaded cells were the same in the temperaturesensitive mutants as in the wild type and in the temperature-insensitive mutants (Table 2).

Experiments in which CS2<sup>TC</sup> cells grown at 30 C were exposed to brief heat treatment in the absence of growth, and then were allowed to resume growth at 30 C and were tested periodically for glutamate permease activity (Fig. 4), showed that the mutation resulted in the formation of a thermolabile repressor, rather than in a temperature-sensitive synthesis of repressor (18). The possibility that the heating inactivated a permease inhibitor rather than a repressor is extremely unlikely in view of the gradual increase in the specific activity of the permease following the heat treatment. If an inhibitor of permease activity were removed, one would expect an immediate relief from inhibition, not followed by any further increase in permease activity during subsequent growth at 30 C. We therefore conclude that the mutation occurred in the repressor-forming gene (designated as gltR), resulting in the synthesis of a thermolabile repressor and thus leading to temperature-sensitive regulation of glutamate permease synthesis.

Genetic analysis of the  $gltR^{TL}$  mutation unequivocally places the gltR gene close to the metA locus, that is, at a distance of about 6 to 7 min [in time of chromosome transfer according to Taylor and Trotter (19)] from the gltC locus (see Fig. 3).

In view of these findings, it seems reasonable to assume that the control gene gltC may be the operator of the glutamate permease operon. This assumption is supported by the isolation of Glut mutants with a lesion in the structural gene of glutamate permease, in which both the transport activity and the affinity for glutamate are greatly reduced (see Table 6). Mapping experiments revealed that the structural gene, gltS, is very closely linked to the gltC locus (see Tables 7 and 9). Cis-trans tests are now in progress to establish the dominance relationships among the different genes of this system. Preliminary results indicate that the gltCc allele, when in a cis position to gltS+, is dominant over gltC+, as indeed would be expected if gltC were the operator gene.

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