

Export of L-Isoleucine from *Corynebacterium glutamicum*: a Two-Gene-Encoded Member of a New Translocator Family

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Bacteria possess amino acid export systems, and *Corynebacterium glutamicum* excretes L-isoleucine in a process dependent on the proton motive force. In order to identify the system responsible for L-isoleucine export, we have used transposon mutagenesis to isolate mutants of *C. glutamicum* sensitive to the peptide isoleucyl-isoleucine. In one such mutant, strong peptide sensitivity resulted from insertion into a gene designated *brnF* encoding a hydrophobic protein predicted to possess seven transmembrane spanning helices. *brnE* is located downstream of *brnF* and encodes a second hydrophobic protein with four putative membrane-spanning helices. A mutant deleted of both genes no longer exports L-isoleucine, whereas an overexpressing strain exports this amino acid at an increased rate. BrnF and BrnE together are also required for the export of L-leucine and L-valine. BrnFE is thus a two-component export permease specific for aliphatic hydrophobic amino acids. Upstream of *brnFE* and transcribed divergently is an Lrp-like regulatory gene required for active export. Searches for homologues of BrnFE show that this type of exporter is widespread in prokaryotes but lacking in eukaryotes and that both gene products which together comprise the members of a novel family, the LIV-E family, generally map together within a single operon. Comparisons of the BrnF and BrnE phylogenetic trees show that gene duplication events in the early bacterial lineage gave rise to multiple paralogues that have been retained in α -proteobacteria but not in other prokaryotes analyzed.

Traditionally, bacterial export carriers were thought to confer resistance to antibiotics or remove waste products. However, it has become clear more recently that carriers also export primary metabolites such as sugars and amino acids. *Escherichia coli*, for instance, has three similar exporters encoded by the *setA*, *setB*, and *setC* genes, two of which (*setA* and *setB*) export lactose and glucose (21). A distant *E. coli* paralogue, YdeE, catalyzes arabinose export (4). Another homologue, the *Erwinia chrysanthemi* *sotB* gene product, probably exports lactose (6). The physiologically relevant functions of these exporters are poorly understood. For example, a triple *setABC* mutant has no observable phenotype under the conditions examined (21, 22). Possibly, sugar exporters are involved in the detoxification of nonmetabolizable sugars (22).

Two transporters in *E. coli* are known to catalyze amino acid export. One is encoded by the *rhtB* gene, whose overexpression results in increased extracellular accumulation of L-homoserine (42). The other is encoded by *ydeD*. Upon its overexpression, the concentration of extracellular L-cysteine metabolites is augmented (8). Whereas the primary functions of these *E. coli* exporters are not obvious, the basic amino acid exporter LysE of *Corynebacterium glutamicum* can clearly protect the cell against toxic levels of cytoplasmic cationic amino acids (3, 39). The absence of LysE results in growth arrest at elevated intracellular L-lysine or L-arginine concentrations

which occur during growth on complex medium or in the presence of basic amino acid-containing peptides. Expression of this exporter is strictly controlled by the intracellular concentration of the basic amino acids (3). Since LysE homologues are present in a large number of bacteria (40), one can anticipate that many bacteria will prove to exhibit this type of intracellular amino acid control. Indeed, amino acid export during growth of *Lactococcus lactis* on milk has been observed (16), although the exporters responsible for this activity have not yet been identified.

We have studied amino acid export in *C. glutamicum* and have identified the aforementioned basic amino acid exporter LysE (39) and the L-threonine exporter ThrE (34). Surprisingly, in both cases, new large families of translocator proteins of previously unknown function were identified (40, 41). Another amino acid of long-standing interest is L-isoleucine. This amino acid is excreted by *C. glutamicum* after deregulation of its biosynthetic pathway (10, 26). L-Isoleucine fluxes across the bacterial membrane are of particular interest from a mechanistic point of view since this amino acid is highly hydrophobic. In fact, net flux may result from diffusion and carrier-mediated export counteracting carrier-mediated import (44). The L-isoleucine importer, BrnQ, mediates Na⁺-dependent import of the three branched-chain amino acids (9, 35). The exporter, known to be driven by the proton motive force (14), has not been identified. We describe here the molecular identification of a translocator exporting L-isoleucine from the cell and show that this exporter represents the prototype of a novel two-gene-encoded translocator family widely distributed in bacteria and archaea.

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TABLE 1. Strains and plasmids used

Strain or plasmid	Relevant characteristics ^a	Reference or source ^b
Strains		
<i>E. coli</i>		
DH5 α MCR	<i>endA1 supE44 recA1 gyrA96 relA1 deoR U169</i> ϕ 80 <i>dlacZ</i> Δ M15 <i>mcrA</i> Δ (<i>mrr-hsdRMS-mcrBC</i>)	12
GM2929	<i>dam-13::Tn9 dcm-6 hsdR2 recF143 mcrA mcrB</i>	27
<i>C. glutamicum</i>		
ATCC 13032	Wild type	ATCC
ATCC 14752	Wild type	ATCC
13032 Δ <i>brnFE</i>	Wild type deleted of a 942-nt fragment of <i>brnFE</i>	This work
13032 Δ <i>lrp</i>	Wild type deleted of a 409-nt fragment of <i>lrp</i>	This work
13032:: <i>lrp</i>	Wild type with <i>lrp</i> disrupted by pK18mob <i>lrp</i> _{int}	This work
Plasmids		
pCGL0040	Donor of Tn5531 (IS1207, Km ^r), Ap ^r <i>oriV</i> _{Ec}	U53587
pET2	Promoter-probe vector, Cm ^r	38
pZ1	<i>E. coli-C. glutamicum</i> shuttle vector, Km ^r <i>oriV</i> _{Ec} <i>oriV</i> _{Cg}	23
pJC1	<i>E. coli-C. glutamicum</i> shuttle vector, Km ^r <i>oriV</i> _{Ec} <i>oriV</i> _{Cg}	7
pK18mob	Integration vector, Km ^r <i>oriV</i> _{Ec} <i>oriT</i>	33
pK19mobsacB	Integration vector, Km ^r <i>oriV</i> _{Ec} <i>oriT</i> <i>sacB</i>	33
pK19mobsacB Δ <i>brnFE</i>	pK19mobsacB with a 942-nt internal fragment deleted in <i>brnFE</i>	This work
pK19mobsacB Δ <i>brnF</i>	pK19mobsacB with a 642-nt internal fragment deleted in <i>brnF</i>	This work
pK19mobsacB Δ <i>lrp</i>	pK19mobsacB with a 409-nt internal fragment deleted in <i>lrp</i>	This work
pK18mob <i>lrp</i> _{int}	pK18mob with a 203-nt internal fragment of <i>lrp</i>	This work
pJC1 <i>brnFE</i>	pJC1 with a 1.3-kb PCR fragment containing <i>brnFE</i>	This work
pJC1 <i>brnF</i>	pJC1 with a 0.96-kb PCR fragment containing <i>brnF</i>	This work
pZ1 <i>brnE</i>	pZ1 with a 1.2-kb <i>EcoRI/XbaI</i> fragment of pUC18 <i>brnE</i> containing <i>brnE</i>	This work
pZ1 <i>lrp</i>	pZ1 with a 0.7-kb <i>EcoRI/XbaI</i> fragment of pUC18 <i>lrp</i> containing <i>lrp</i>	This work
pET2 <i>brnF-lrp</i>	pET2 with a 219-bp <i>BamHI</i> PCR fragment containing the promoter of <i>brnF</i> and <i>lrp</i>	This work
pET2 <i>brnE</i>	pET2 with a 239-bp <i>BamHI-KpnI</i> PCR fragment containing the promoter of <i>brnE</i>	This work
pUC18 <i>brnFE-lrp</i>	pUC18 with a 1.8-kb PCR fragment containing <i>brnFE-lrp</i>	This work
pUC18 <i>brnE</i>	pUC18 with a 1.2-kb PCR fragment containing the <i>brnE</i> gene	This work
pUC18 <i>lrp</i>	pUC18 with a 0.65-kb PCR fragment containing the <i>lrp</i> gene	This work

^a Km^r, kanamycin resistant; Ap^r, ampicillin resistant; Cm^r, chloramphenicol resistant. Subscripts: *Ec*, *E. coli*; *Cg*, *C. glutamicum*.

^b ATCC, American Type Culture Collection.

MATERIALS AND METHODS

Bacteria, plasmids, and growth conditions. The strains and plasmids used are listed in Table 1. Strains were grown as described previously (34). The minimal medium for *C. glutamicum* was usually CGXII (26), but to quantify amino acid export, the minimal medium MMI was used (14), with (NH₄)₂SO₄ reduced to 0.5 g liter⁻¹. When appropriate, carbenicillin (50 μ g ml⁻¹), kanamycin (KAN; 15, 25, or 50 μ g ml⁻¹), or chloramphenicol (20 μ g ml⁻¹) was added to the medium.

Isolation of peptide-sensitive mutants and localization of transposon insertion sites. The transposon (Tn) delivery vector pCGL0040 was isolated from *E. coli* GM2929 grown in the presence of 50 μ g of KAN and 20 μ g of chloramphenicol ml⁻¹ (1). The plasmid was used to transform *C. glutamicum* ATCC 14752 to KAN resistance by using LBHIS plates with 15 μ g of KAN ml⁻¹ (19). The resulting Tn mutants were transferred to CGXII plates containing 25 μ g of KAN ml⁻¹ and either 3 mM of the dipeptide, Ile-Ile, or no peptide. The clones that exhibited retarded growth in the presence of Ile-Ile but grew normally in the absence of peptide were retrieved from the LBHIS master plate and retested in liquid CGXII medium. For that purpose, mutants were precultured on brain heart infusion medium (Difco) containing 25 μ g of KAN ml⁻¹. Minimal medium cultures (CGXII \pm 3 mM Ile-Ile, 25 μ g of KAN ml⁻¹) were inoculated at an initial optical density at 600 nm (OD₆₀₀) of 0.1 or 0.5. Mutants that confirmed the growth delay in the presence of Ile-Ile were saved for further studies. The cloning and sequencing of the Tn insertion site was performed as described previously (1, 34).

Construction of plasmids. Plasmids were constructed in *E. coli* DH5 α MCR from PCR-generated fragments (Expand High Fidelity PCR kit; Roche Diagnostics) by using *C. glutamicum* ATCC 13032 DNA as a template. In order to construct pJC1*brnFE*, the upstream primer 5'-GCTCTAGAACCCTTGTCAGC CAGTGCAGAGAT-3' and the downstream primer 5'-GCTCTAGAAAAATC CGCATCCCCTTCA-3' were used. The resulting fragment was *XbaI* digested and cloned into the *XbaI* site of pJC1. Using the same upstream primer as before

and primer 5'-GCTCTAGAGCCCGGAGCGCAAAAGTAAT-3', *brnF* was amplified. The fragment was cloned into the *XbaI* site of pJC1. *brnE* was amplified by using the primers 5'-ATTCATTCAGCCTGGAGGTGTCG-3' and 5'-AGCGCTGTCTGCTTAAGCCTTTTC-3'. The resulting fragment was blunted, cloned into the *SmaI* site of pUC18, reisolated as an *EcoRI-XbaI* fragment and, after blunting, ligated with *ScaI*-treated pZ1. Cloning of *lrp* was achieved by use of a 0.65-kb fragment obtained by PCR. The fragment was first ligated with pUC18, excised as an *EcoRI-XbaI* fragment, and then ligated with pZ1. For the cloning of the promoter regions of *brnF-lrp* and *brnE*, respectively, PCR fragments carrying the putative promoter regions were ligated with the promoter probe vector pET2.

To generate a chromosomal *brnFE* deletion, crossover PCR was applied (20). An 869-bp fragment carrying part of the 5' end of *brnF* and the 3' end of *brnE* with an internal 942-bp fragment deletion was generated (Fig. 2). The fragment was cloned into pK19mobsacB via its attached *EcoRI* and *XbaI* site. Similarly, pK19mobsacB derivatives were made with 642 bp of *brnF* deleted to allow an in-frame deletion of this gene, as was a derivative carrying a fragment with 409 bp of the *lrp* sequence deleted.

Construction of strains. *C. glutamicum* was transformed by electroporation (37), and the presence of replicative plasmids was verified by plasmid reisolation and restriction analysis. To obtain the *lrp* inactivation mutant, the nonreplicative plasmid pK18mob*lrp*_{int} was transferred to *C. glutamicum*. The correct integration of the vector into the chromosome was verified by PCR analysis. The *brnFE* deletion mutant and the *lrp* deletion mutant were made by using pK19mobsacB Δ *brnFE* and pK19mobsacB Δ *lrp*, respectively. Clones were selected for KAN resistance to establish integration of the plasmid in the chromosome. In a second round of positive selection by using sucrose resistance, clones were selected for deletion of the vector (33). The deletions in the chromosome were verified by PCR analysis or Southern blotting.

Primer extension. Total RNA was isolated from *C. glutamicum* by using extraction with hot acidic phenol (11). The transcriptional start sites were deter-

mined by primer extension by using SuperScript II reverse transcriptase (Gibco-BRL) with the primers labeled with [32 P]ATP. In parallel, the respective DNAs (pET2*pbrmFlp* and pET2*brmE*) were sequenced by using 32 P-labeled primers and the Thermosequenase kit. The sequencing reactions and primer extension products were heated at 95°C for 4 min, and 2- μ l samples were loaded onto a polyacrylamide gel.

Assay of amino acid export. To determine the amino acid export rates, cells pregrown in brain heart infusion medium (Difco) were washed once with ice-cold 0.9% NaCl and transferred into MMI (14) to give an initial OD₆₀₀ of 10, which corresponds to 3.0 mg (dry weight) ml⁻¹. After preincubation for 20 min at 30°C with rapid stirring (ca. 700 rpm on a magnetic stirrer), the assay of amino acid excretion was initiated by addition of dipeptide in final concentrations as indicated in Results. Processing of samples for separation of extra- and intracellular fluid was performed by using silicone oil centrifugation (17). In the resulting fractions, amino acids were quantified as their *o*-phthalaldehyde derivatives via high-pressure liquid chromatography. The intracellular volume used to calculate the internal amino acid concentration was 1.6 μ l mg (dry weight)⁻¹.

Calculations. Observed efflux rates of branched-chain amino acids are due to diffusion, active export, and active import (44). Therefore, the carrier-mediated export rate is calculated as: $V_{ex} = V_{eff} - K_d \cdot [aa]_{in} + V_{in}$, where V_{eff} is the observed efflux rate as provided in the graphs. $K_d \cdot [aa]_{in}$ is the efflux due to diffusion, which is dependent on the determined internal amino acid concentration ($[aa]_{in}$) in nanomoles per microliter. Due to the low external concentrations of the branched-chain amino acids accumulating in the experiments, these were neglected in the calculations. The known diffusion constants (K_d values) are 0.13 for L-Ile and L-Leu and 0.09 for L-Val in μ l mg (dry weight)⁻¹ min⁻¹ (18, 24). V_{in} is the rate of uptake of the branched-chain amino acids. Uptake was invariably observed under all conditions assayed (9, 14, 35, 44). The uptake rates measured were 1.1, 0.94, and 1.3 nmol min⁻¹ mg (dry weight)⁻¹ for L-Ile, L-Leu, and L-Val, respectively. The active export rates resulting from this approximation are expressed in nanomoles per minute per milligram (dry weight).

Computer methods. The programs and computer methods used here were as described previously (41).

Nucleotide sequence accession number. The sequence data have been submitted to the GenBank database under accession numbers AF454053 (*brnFE-lrp*), AF454055 (*chl1*), and AF454054 (*htaA*).

RESULTS

Isolation of peptide-sensitive mutants and characterization of Tn insertion loci. After the observation that the growth of a *lysE*-null mutant of *C. glutamicum* unable to export L-lysine is sensitive to the presence of lysine-containing peptides (39) and, similarly, that the growth of a *thrE* mutant is sensitive to the presence of threonine-containing peptides (34), we screened 1,800 Tn mutants of *C. glutamicum* ATCC 14752 for sensitivity to 3 mM of the isoleucine dipeptide, Ile-Ile, on plates. This strategy was based on the knowledge that *C. glutamicum* can hydrolyze cytoplasmic Ile-Ile and that internally accumulated L-isoleucine is toxic (44). The screening procedure yielded three clones that were studied in liquid culture (Fig. 1A). Clone 1-75 showed weak growth retardation upon addition of Ile-Ile that was comparable to that of clone 1-25 (not shown), whereas clone 1-8 exhibited strong growth retardation. Cloning and analysis of the Tn adjacent sequences revealed that in clone 1-25 a polypeptide encoding a putative heme transporter, *htaA*, was disrupted, and that in clone 1-75 the Tn was inserted into the promoter region of a putative magnesium chelatase subunit, *chl1*. Although the only currently known function of this latter gene is to incorporate Mg²⁺ into the protoporphyrin skeleton of photosynthetic organisms, attempted inactivation in *C. glutamicum* was not possible. From this observation we suggest that Mg²⁺ incorporation into an unidentified cellular component is essential for the growth of *C. glutamicum*.

In clone 1-8, with the strongest peptide sensitivity, the Tn

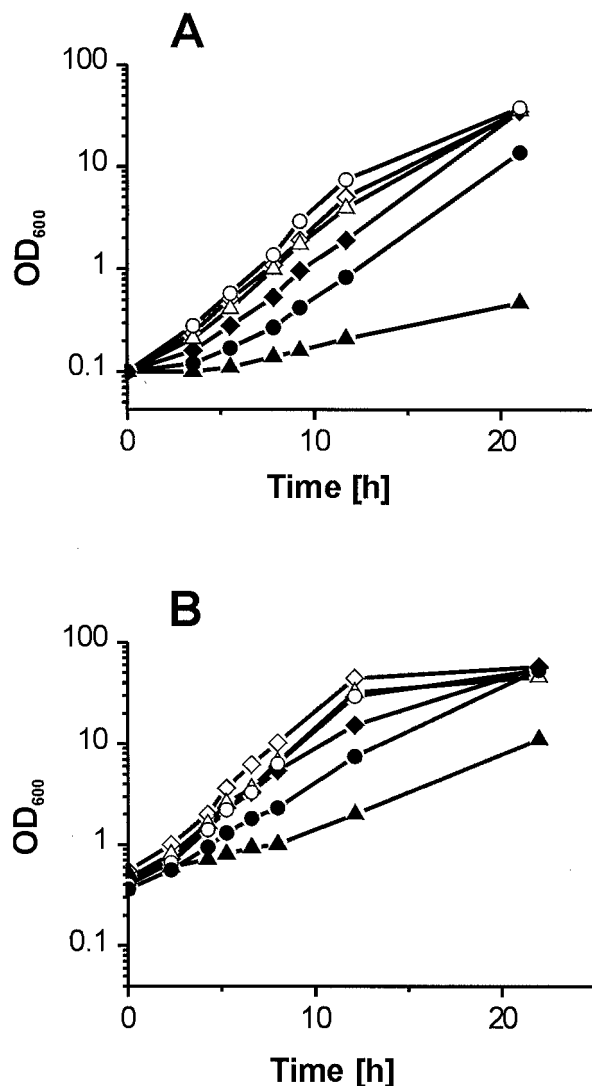


FIG. 1. Peptide-sensitive growth of Tn mutants. (A) Growth of Tn mutants 1-75 (circles) and 1-8 (triangles) compared to the control strain, *C. glutamicum* ATCC 14752/pZ1 (diamonds), in the presence (solid symbols) or absence (open symbols) of 3 mM Ile-Ile. (B) Growth of the deletion mutant 13032 Δ *brnFE* with control vector pJC1 (triangles) or pJC1*brnFE* (circles) compared to the wild-type 13032/pZ1 (diamonds) in the presence (solid symbols) or absence (open symbols) of 5 mM Ile-Ile.

was inserted into a gene whose homologue in *Bacillus subtilis* is *azlC* and which when deleted together with *azlD* renders *B. subtilis* more susceptible to azaleucine (2). The Tn-inactivated gene in *C. glutamicum* ATCC 13032 is 756 nucleotides (nt) in size and encodes a polypeptide with a molecular weight of 27,300 (Fig. 2). It overlaps by 4 bp the downstream gene of 327 nt that is predicted to encode a polypeptide of 11,500 Da. In accordance with the subsequent functional analyses, the two corresponding genes in *C. glutamicum* were termed *brnF* and *brnE* for branched-chain amino acid export. Both polypeptides exhibit pronounced local hydrophobicities (see below). Divergently transcribed from *brnFE* is a putative regulatory gene

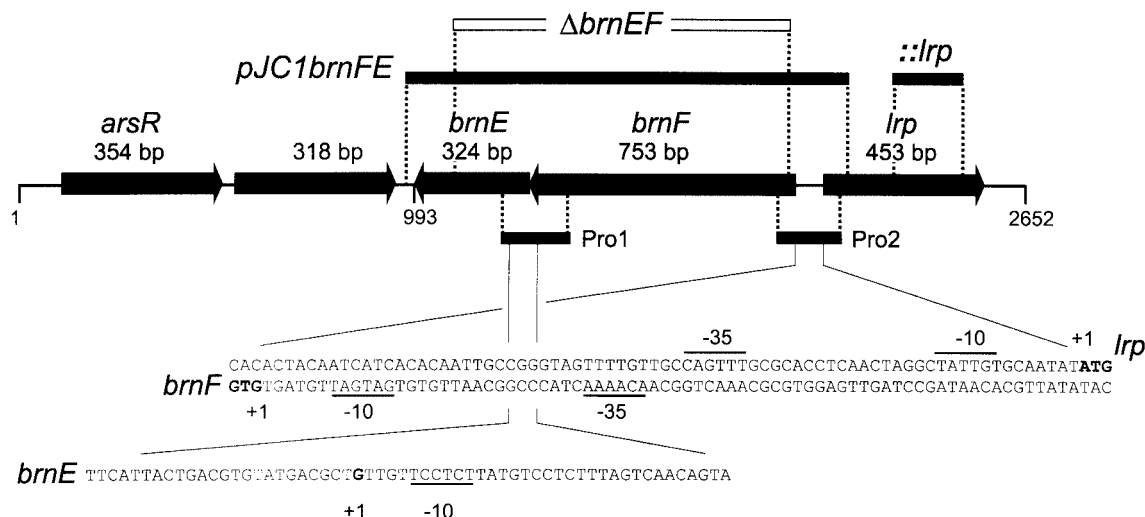


FIG. 2. Overview on the *brnE-brnF-lrp* locus of *C. glutamicum* and regions with identified promoter activity. In the upper part of the figure, the thick arrows represent the genes with their sizes in base pairs. Above them, two fragments used for plasmid construction and *lrp* inactivation, respectively, are given, as well as the part of the genome deleted in the *brnFE* deletion mutant. In the lower part of the figure, the two fragments, Pro1 and Pro2, used for primer extension are depicted, together with the sequence of the promoters identified. The transcriptional start sites are marked “+1.” These are underlined, as are the -10 and -35 promoter regions. The initiation codons for *lrp* and *brnF* are indicated in boldface. The separate *brnE*-transcript initiation might not contribute for polypeptide formation (see the text).

that encodes a protein exhibiting 37% identity to Lrp of *E. coli* (13).

Deletion of *brnFE* and growth characteristics. With the objective of deleting the chromosomal *brnFE* locus, the mobilizable replacement vector pK19mobsacB $\Delta brnFE$ was constructed carrying sequences flanking *brnFE* (Fig. 2). It was transferred into *C. glutamicum* ATCC 13032, and two rounds of positive selection were applied to integrate and subsequently remove the vector (33). PCR analysis of 10 clones revealed that in 3 of the clones *brnFE* was in fact deleted, whereas in 7 clones the wild-type locus was reconstituted. These numbers indicate that these genes may not play a major housekeeping function for cellular metabolism or growth of *C. glutamicum*. Growth of the deletion mutant in liquid culture was indistinguishable from that of the wild type (Fig. 1B), thus confirming this conclusion. Only when the Ile-Ile dipeptide was present in the medium was growth reduced. When the deletion mutant was transformed with pJCb*rnFE*, the strain regained its peptide-resistant growth phenotype (Fig. 1B). This effect was not observed when *brnF* or *brnE* was individually overexpressed, providing evidence that both genes together promote resistance to the peptide (not shown). In view of the fact that deletion of the *brnFE* homologues, *azlCD*, abolishes azaleucine resistance in *B. subtilis* (2), we also assayed growth of the recombinant *C. glutamicum* strains in response to azaleucine. Although the growth rate of the wild-type strain was reduced from 0.34 to 0.15 h⁻¹ in the presence of 0.5 mM azaleucine, deletion or overexpression of *brnFE* did not influence growth either with or without azaleucine (not shown).

Determination of the transcriptional start sites of *brnF*, *brnE*, and *lrp*. To define the genes, transcript initiation sites were measured. For this purpose a 219-bp *Bam*HI fragment carrying the divergent promoter regions of *lrp* and *brnF* (Pro2 in Fig. 2) was cloned in the two possible orientations into the

promoter probe vector pET2 (38). Similarly, a 239-bp *Bam*HI-*Kpn*I fragment (Pro1) carrying a possible start site of *brnE* was cloned. The resulting plasmids rendered *C. glutamicum* resistant to chloramphenicol. The MICs caused by the *brnF* and *lrp* promoters were 60 and 50 $\mu\text{g ml}^{-1}$, respectively; that of the *brnE* promoter was 25 $\mu\text{g ml}^{-1}$.

The results of the primer extension experiments with the sequencing reactions carried out in parallel are shown in Fig. 3. In the case of *brnF* and *lrp*, the transcript initiation sites are identical to those of their deduced polypeptides (Fig. 2). Such leaderless genes are a frequent feature of *C. glutamicum* and related organisms (25). The close proximity of the genes separated by just 74 nt, as well as their overlapping promoters, indicates that *brnF* expression might be a target of the *lrp*-encoded regulator (see below). By using fragment Pro1, an additional transcript initiation site was observed within *brnE* (Fig. 2). However, the corresponding “ -10 ” consensus sequence is located 2 nt closer to the determined transcript initiation site than in most other *C. glutamicum* promoters (28). This fact and the low MIC make it unlikely that this site contributes to transcript formation. Instead, we assume that there is a single message for both *brnF* and *brnE*.

Increased L-isoleucine export is due to *brnFE* overexpression. In order to perform functional analyses of *brnFE*, we measured export rates in the presence of Ile-Ile. Addition of this peptide leads to increased intracellular L-Ile steady-state levels (44). The actual cytoplasmic concentrations are determined by the rates of import and hydrolysis of the peptide, as well as the L-Ile efflux rate. Upon addition of 3 mM Ile-Ile, the internal L-Ile concentration increased up to 23 mM in strain 13032/pJCb*rnFE*, up to 45 mM in strain 13032/pZ1, and up to 87 mM in strain 13032 $\Delta brnFE$ /pJC1 (Fig. 4A).

The rates of L-isoleucine efflux from these strains are fairly comparable, which may at first seem surprising. However, this

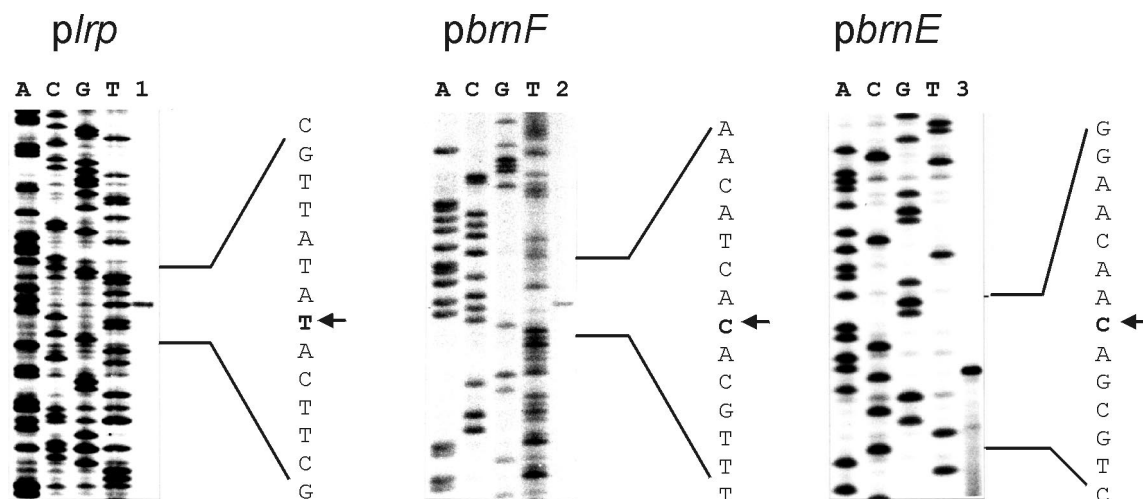


FIG. 3. Mapping of the transcriptional start sites of *lp* (*plrp*), *brnF* (*pbrnF*), and *brnE* (*pbrnE*) by primer extension analysis. In each experiment, the primer extension product was run in the numbered lane. The sequencing ladder (labeled A, C, G, and T) of the coding strand was generated by using the same primer that was used for primer extension. The respective transcriptional start site is indicated by the arrow.

is due to the significant L-Ile efflux component caused by diffusion (44). This carrier-independent flux component is determined by the concentration gradient across the plasma membrane. Based on the known diffusion constant, at the high internal concentration of 87 mM in the deletion mutant, efflux due to diffusion is $6.4 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$. At the lower concentration of 23 mM in the overexpressing strain, it is only $2.6 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$. The L-isoleucine uptake rate of $1.1 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$ detracts from the apparent L-isoleucine efflux rate (9, 35). Accordingly, the active export rate is the difference of the total efflux minus diffusion plus uptake. When we calculated the active export rates (see Materials and Methods), values of $7.8 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$ for the overexpressing strain, $0.4 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$ for the deletion mutant, and $4.2 \text{ nmol min}^{-1} \text{ mg (dry weight)}^{-1}$ for the wild type were derived (Table 2). Upon overexpression of either *brnF* or *brnE*, active export was not increased (Table 2), in accordance with the failure of each of these genes singly to relieve peptide-sensitive growth. Also, with a *brnF* in-frame deletion, no export was obtained (Table 2).

In separate experiments, 10 mM Ile-Ile was used with the overexpressing strain and 0.5 mM Ile-Ile was used with the deletion mutant in order to achieve more comparable internal concentrations of L-Ile (Fig. 4B). Total efflux proved to be highest with *pJC1brnFE* and lowest with the deletion mutant. Nevertheless, the calculated export rates were almost identical to those determined in the experiment described earlier (Table 2). These experiments (i) confirm that L-isoleucine export is due to *brnFE* expression, (ii) suggest that both genes are required for export, and (iii) make it unlikely that an additional carrier contributes appreciably to L-isoleucine export. In *C. glutamicum*, under identical conditions but without peptide, the internal concentrations of the branched-chain amino acids do not exceed 2 mM (not shown).

L-Val and L-Leu export. As previously reported, loading *C. glutamicum* from the external side with L-Leu or L-Val resulted in reduced L-Ile efflux by up to 25% (14). We used the iden-

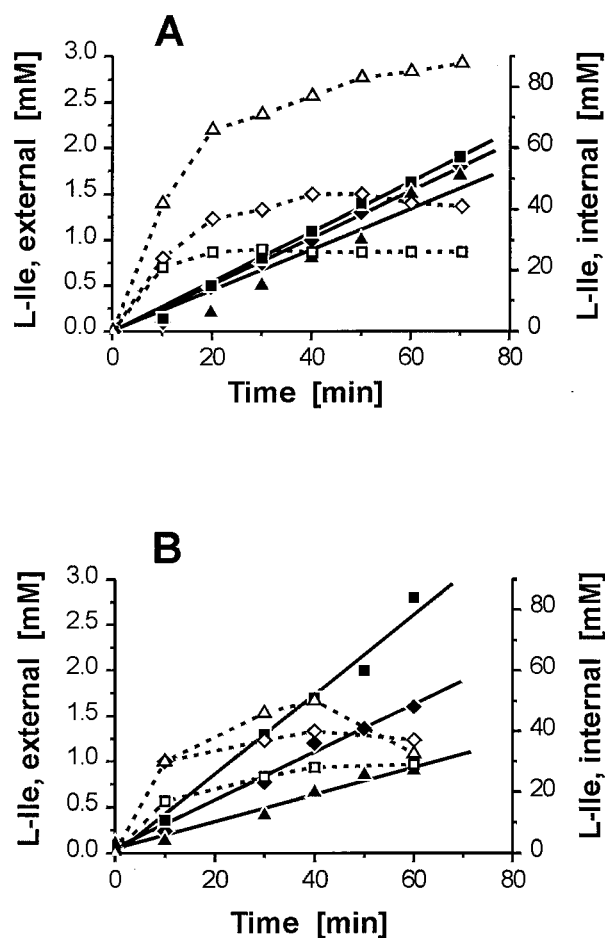


FIG. 4. Internal L-isoleucine concentrations and isoleucine efflux after *brnFE* expression. (A) Efflux (solid symbols) and internal L-isoleucine concentrations (open symbols) upon addition of 3 mM Ile-Ile to 13032/pJC1*brnFE* (squares) and 13032Δ*brnFE*/pJC1 (triangles) compared to the wild-type strain 13032/pZ1 (diamonds). (B) Same as panel A, except that 10 mM Ile-Ile was used with strain 13032/pJC1*brnFE* and 0.5 mM Ile-Ile was used with strain 13032Δ*brnFE*/pJC1.

TABLE 2. Export rates due to *brnF*, *brnE*, and *lrp* expression

<i>C. glutamicum</i> strain	Export rate (nmol min ⁻¹ mg [dry wt] ⁻¹) ^a of:		
	L-Isoleucine	L-Valine	L-Leucine
13032/pJC <i>brnFE</i>	7.8 ± 1.3 ^a	3.3, 3.2	6.3
13032Δ <i>brnFE</i> /pJC1	0.4 ± 0.2	-0.1 ± 0.9	0.6 ± 0.4
13032Δ <i>brnF</i> /pJC1	-0.9 ± 0.6	ND	ND
13032/pZ1	4.2 ± 0.4	2.4, 2.0	4.0
13032Δ <i>brnFE</i> /pJC <i>brnFE</i>	6.9	4.5	8.1
13032/pJC1 <i>brnF</i>	4.6	ND	ND
13032/pJC1 <i>brnE</i>	3.7	ND	ND
13032Δ <i>brnFE</i> /pJC1 <i>brnE</i>	0.5, 0.3	ND	ND
13032Δ <i>brnFE</i> /pJC1 <i>brnF</i>	0.4, 0.2	ND	ND
13032/pZ1 <i>lrp</i>	3.6	ND	ND
13032Δ <i>lrp</i>	0.0, -0.7	ND	ND
13032 <i>lrp</i> ::pK18mob <i>lrp</i> _{int}	0.7	ND	ND

^a Values for the calculation of standard deviations (provided where applicable) were derived from three independent measurements. ND, not done.

tified exporter and peptides to assay more directly whether BrnFE accepts the other branched-chain amino acids. With 5 mM Ala-Val in the deletion mutant, up to 82 mM L-Val accumulated, and in the deletion mutant with pJC1*brnFE*, ca. 30 mM L-Val accumulated (Fig. 5A). The total L-Val efflux rate by the deletion mutant was substantially lower than that by the overexpressing strain. This may be due to the known low membrane diffusibility of L-Val (24), as becomes evident from comparison of the L-Ile and L-Val efflux rates in the deletion mutant under otherwise identical conditions (compare Fig. 4A with Fig. 5A). Using a diffusion rate constant of 0.09 μl mg (dry weight)⁻¹ min⁻¹ (18, 24), the calculated export rates for L-Val are -0.1 nmol min⁻¹ mg (dry weight)⁻¹ with the deletion mutant and 4.5 nmol min⁻¹ mg (dry weight)⁻¹ with the same strain but overexpressing *brnFE* (Table 2). In addition, with the wild-type strain carrying pJC1*brnFE*, the L-Val export rate proved to be 3.3 nmol min⁻¹ mg (dry weight)⁻¹. This confirms the conclusion that L-Val export is mediated by the *brnFE*-encoded exporter. This rate is determined to be ca. 60% of that of L-Ile export.

Interestingly, with Leu-containing peptides (Leu-Leu or Ala-Leu) in the *brnFE*-overexpressing strain, the internal L-Leu concentration never exceeded 15 mM. Nevertheless, with 5 mM Leu-Leu, a high efflux rate (8.6 nmol min⁻¹ mg [dry weight]⁻¹) resulted with 13032Δ*brnFE*/pJC1*brnFE* (Fig. 5B). Thus, from the calculated export rate (Table 2), we conclude that L-Leu and L-Ile are exported at comparable rates. For further confirmation, we conducted a competition experiment in which both amino acids were simultaneously present as a substrate of the exporter. The dipeptide, Leu-Ile, was used for this purpose (Fig. 6). In spite of a comparatively low concentration of ca. 10 mM for both amino acids, which could be due to limited peptide uptake, comparable export rates were observed for L-Ile and L-Leu of 2.5 and 1.8 nmol min⁻¹ mg (dry weight)⁻¹, respectively. This proved not to be true for the deletion mutant (data not shown). This finding shows that the exporter does not discriminate between the two hydrophobic amino acids. In contrast to the experiment shown in Fig. 4, a low L-isoleucine export rate suggests that the carrier is not saturated at the low internal concentrations present. This is in accordance with the observed low affinity of the system (ob-

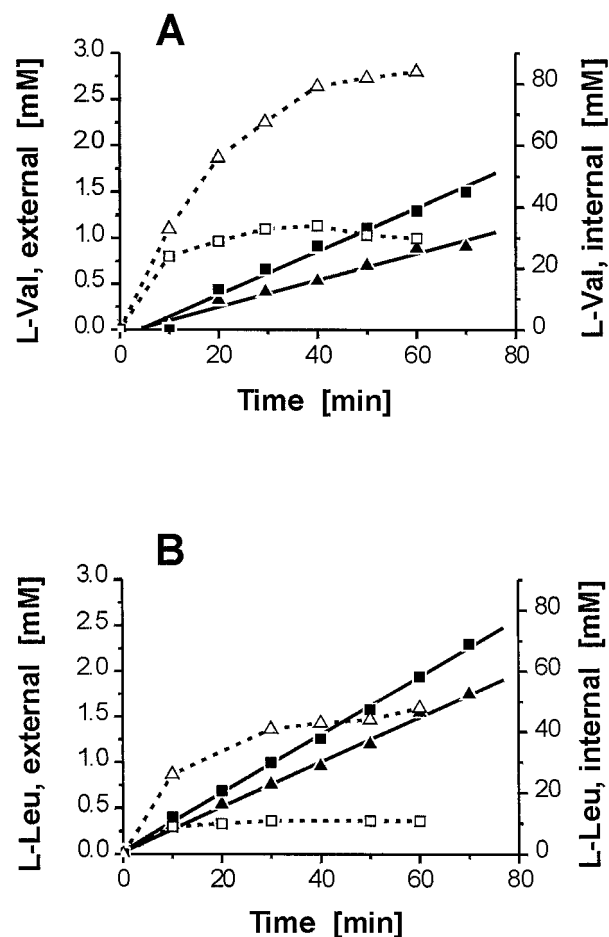


FIG. 5. Efflux (solid symbols) and internal concentrations (open symbols) of L-valine and L-leucine by *C. glutamicum* ATCC 13032Δ*brnFE*/pJC1 (triangles) and ATCC 13032Δ*brnFE*/pJC1*brnFE* (squares). (A) Response in the presence of 5 mM Ala-Val. (B) Response in the presence of 5 mM Leu-Leu.

servable K_m , 21 mM [44]). It suggests a comparable affinity of the system for these two amino acids.

Involvement of *lrp* in export. *lrp* maps upstream of *brnFE* (Fig. 2). To reveal whether *lrp* is involved in regulation of the exporter, we constructed an *lrp* disruption mutant. Strain 13032*lrp*::pK18mob*lrp*_{int} no longer proved to export L-isoleucine (Table 2). An additionally used *lrp* deletion mutant confirmed this result, indicating that Lrp is essential for the export of the branched-chain amino acids, probably by acting as an activator of *brnFE* expression.

Homologues of BrnFE in other organisms. Table 3 lists homologues of BrnF (left) and BrnE (right) retrieved from the databases. These proteins are derived exclusively from prokaryotic organisms, with gram-negative and gram-positive bacteria as well as archaea being represented. Most organisms with fully sequenced genomes that are represented in Table 3 have just one pair of these proteins. However, the closely related α-proteobacteria *Agrobacterium tumefaciens*, *Sinorhizobium meliloti*, and *Mesorhizobium loti* have multiple paralogues. The two *Helicobacter pylori* pairs of homologues are orthologues derived from two different strains of this organism.

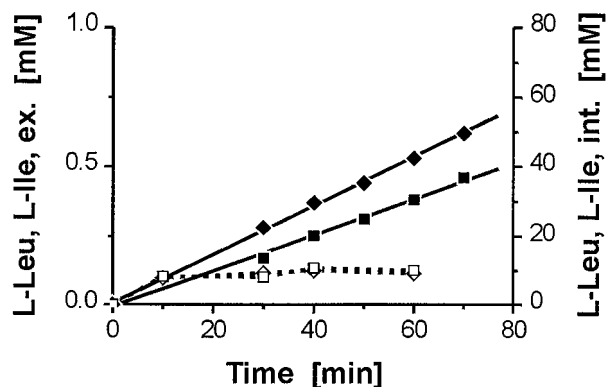


FIG. 6. Efflux (solid symbols) and internal concentrations (open symbols) of L-leucine (squares) and L-isoleucine (diamonds) with *C. glutamicum* ATCC 13032 after the addition of 10 mM Leu-Ile.

It can be seen from the data summarized in Table 3 that most BrnF homologues are of 218 to 256 amino acids (aa) long, whereas all BrnE homologues are 98 to 118 aa. However, the *Methanococcus jannaschii* BrnF homologue, as well as the *Neisseria meningitidis* BrnF homologue, is truncated, and no

BrnE homologue could be identified in the two strains of the latter organism that have been sequenced.

Phylogenetic tree of the LIV-E family. The phylogenetic trees for the BrnF and BrnE homologues are shown in Fig. 7A and B, respectively. Several features are worthy of note. First, there are two sets of orthologous α -proteobacterial proteins (Atu1, Sme1, and Mlo1; Atu2, Sme2, and Mlo2) which cluster loosely together with the *Desulfovibrio gigas* protein. The third set of the Atu3 and Sme3 paralogues cluster together, but they are clearly distant from the other orthologous proteins. It seems clear that the gene duplication events giving rise to these paralogues occurred early, before divergence of the species. Second, most of the gram-positive bacterial homologues cluster loosely together. In Fig. 7A, they fall into three clusters: one including the high G+C gram-positive bacterial proteins, Sco and Cgl; one including the low G+C gram-positive bacterial proteins, Spn, Lla, and Sau; and a third bearing only the *Bacillus halodurans* (Bha) protein. The single *B. subtilis* (Bsu) homologue does not cluster with the *B. halodurans* protein but instead clusters with the γ -proteobacterial proteins, Hin and Pmu, on both trees. The Bsu protein may have been acquired by horizontal transfer. Analyses of the genes encoding the *B.*

TABLE 3. Sequenced proteins of the branched-chain amino acid exporter (LIV-E) family^a

Protein abbreviation	Organism	BrnF homologues		BrnE homologues	
		Size (aa)	GI no.	Size (aa)	GI no.
Atu1	<i>Agrobacterium tumefaciens</i>	240	15157197	104	15157198
Atu2	<i>Agrobacterium tumefaciens</i>	234	15156113	100	15156114
Atu3	<i>Agrobacterium tumefaciens</i>	275	15156429	114	15156430
Afu	<i>Archaeoglobus fulgidus</i>	219	11499344	100	11499343
Bha	<i>Bacillus halodurans</i>	237	10175532	102	10175533
Bsu	<i>Bacillus subtilis</i>	254	3121821	110	3121820
Cgl	<i>Corynebacterium glutamicum</i>	251	14273894	108	14273895
Dra	<i>Deinococcus radiodurans</i>	235	7471172	98	7473007
Dgi	<i>Desulfovibrio gigas</i> ^b	235	12802727	103	
Eco	<i>Escherichia coli</i>	245	3123142	111	2506710
Hin	<i>Haemophilus influenzae</i>	244	1176150	109	1176149
Hpy-A	<i>Helicobacter pylori</i> (J99)	228	12230844	118	11387375
Hpy-B	<i>Helicobacter pylori</i>	228	3123102	115	3123101
Lla	<i>Lactococcus lactis</i>	235	12724620	108	12724619
Mlo1	<i>Mesorhizobium loti</i>	242	13471536	98	13471535
Mlo2	<i>Mesorhizobium loti</i>	244	13472209	99	13472208
Mja	<i>Methanococcus jannaschii</i> ^c	73	3219930	110	3219926
Nme	<i>Neisseria meningitidis</i> ^d	160	11353336		
Pmu	<i>Pasteurella multocida</i>	240	12720674	109	12720675
Pch	<i>Pectobacterium chrysanthemi</i>	247	14970542	113	14970543
Pae	<i>Pseudomonas aeruginosa</i>	252	11349405	104	11349404
Rsp	<i>Rhodobacter sphaeroides</i> ^e	235	7657933		
Sme1	<i>Sinorhizobium meliloti</i>	238	15075182	102	15075181
Sme2	<i>Sinorhizobium meliloti</i>	256	15074061	100	15074062
Sme3	<i>Sinorhizobium meliloti</i>	248	15074421	111	15074422
Sau	<i>Staphylococcus aureus</i>	231	13699927	109	13699928
Spn	<i>Streptococcus pneumoniae</i>	218	14971611	107	14971612
Sco	<i>Streptomyces coelicolor</i>	256	7479559	102	7479560
Vch	<i>Vibrio cholerae</i>	239	11354432	105	11355554

^a GI, GenInfo Identifier (GenBank).

^b The open reading frame for this organism is not included in the current databases and was derived from the nucleic acid sequence.

^c The small size of the *M. jannaschii* BrnF homologue is due to an apparent genomic deletion. If this is a natural deletion rather than a sequencing error, the *M. jannaschii* homologues would be expected to be nonfunctional.

^d Two strains of *N. meningitidis*, serogroup A strain Z2491 and serogroup B strain MC58, exhibit apparently truncated BrnF homologues but no identifiable BrnE homologue. If this is due to a natural genomic deletion that occurred prior to the divergence of these two strains, these homologues would be expected to be nonfunctional pseudogenes.

^e No BrnE homologue could be identified for *R. sphaeroides*, although a full-length BrnF homologue was identified. However, the fully sequenced genome for this organism was not available at the time these studies were conducted.

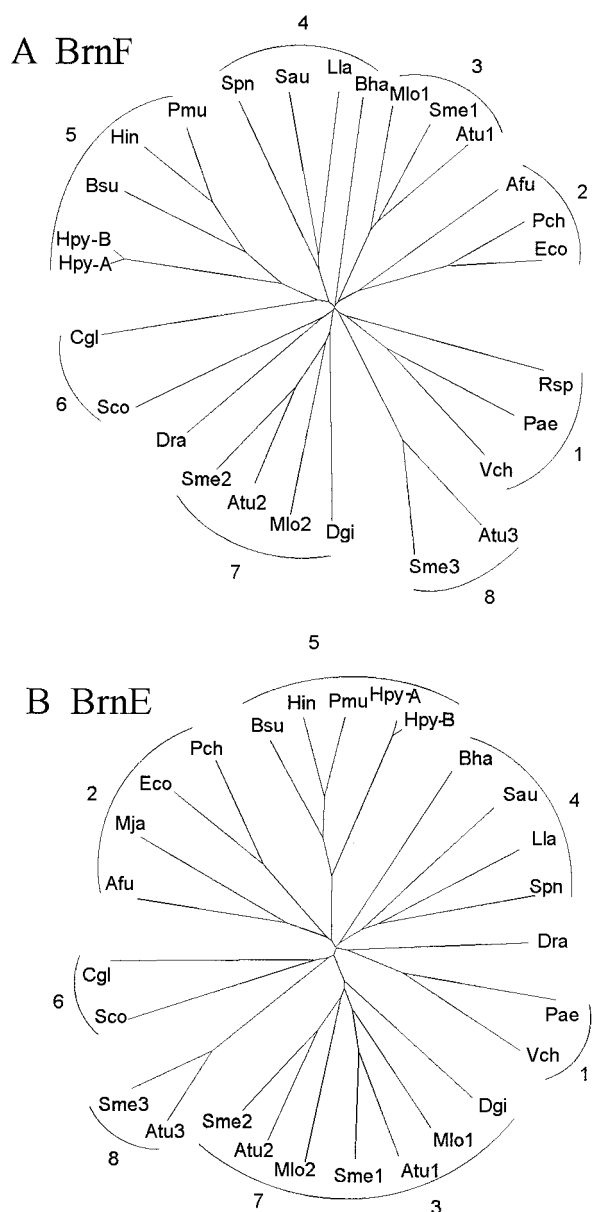


FIG. 7. Phylogenetic trees for the BrnF family (A) and the BrnE family (B). Protein abbreviations are as defined in Table 3. The trees were based on multiple alignments generated by using the CLUSTAL X program (36).

subtilis homologues revealed significant (4%) differences in G+C content, and codon frequencies relative to these values in *B. subtilis* genes as a whole were also noticeably different. Finally, the γ -proteobacterial homologues (Pmu, Hin, Pca, Pae, Eco, and Vch) fall into three distant clusters on both trees, suggesting that at least three early gene duplication events occurred during evolution of the γ -proteobacterial lineage, as was evidently true for the α -proteobacteria. However, in contrast to the α -proteobacteria, with which all three paralogues were retained, at least two of the three were lost from each γ -proteobacterium.

In silico topological analyses of BrnF and BrnE homo-

logues. Figure 8 shows average hydropathy, amphipathicity, and similarity plots for the full-length BrnF and BrnE homologues listed in Table 3. The programs predict the presence of seven TMSs in BrnF homologues and four transmembrane spanners (TMSs) in BrnE homologues. The larger BrnF homologues are clearly better conserved than the smaller BrnE homologues.

The similar spacing of the last four BrnF peaks and the BrnE peaks suggested that these sequence dissimilar proteins arose from a common ancestor. We therefore examined BrnF homologues for regions of sequence similarity with BrnE. The last four putative TMSs in BrnF Mlo2, for example, gave a comparison score of four standard deviations with the corresponding BrnE protein, corresponding to a probability of ca. 10^{-5} that the sequence similarity observed occurred by chance. While suggestive of homology, this value is insufficient to establish common origin (32). We believe that the BrnFE system may have arisen by a gene triplication event from a four-TMS precursor, followed by loss of a single N-terminal TMS and extensive sequence divergence.

DISCUSSION

LysE was the first amino acid exporter characterized from *C. glutamicum* (39), and the use of peptides has recently resulted in the identification of the second such exporter, ThrE (34). *C. glutamicum* BrnFE, described here, represents the third amino acid exporter to be identified. In each of these cases, the discovery of a new large family of bacterial proteins resulted (39, 41). We suggest that all members of these novel families catalyze amino acid export, a function of general relevance that has previously been neglected.

BrnFE exports the three branched-chain amino acids, L-isoleucine, L-leucine, and L-valine, at comparable rates, although it appears that L-valine is exported somewhat less efficiently than is L-leucine or L-isoleucine. One other exporter of the LIV-E family, for which limited functional data are available, is from *B. subtilis* (2). In this case, inactivation of the BrnE homologue (AzID) abolishes resistance to azaleucine. It can be surmised that AzID functions as an azaleucine exporter. Based on the available functional data, as well as on the similar structures of all family members, it can be suggested that export of hydrophobic amino acids is a characteristic of all members of the LIV-E family. However, it is not clear whether the export of amino acids observed with BrnFE is its primary physiological function or, alternatively, if there is an unrecognized function for this translocator. An argument in favor of the latter idea is that the *B. subtilis* genes encoding the BrnFE homologues are cotranscribed with *brnQ*, encoding the uptake system for branched-chain amino acids. Coexpression of these genes would be expected to create an energy-consuming futile cycle unless strictly regulated. Nevertheless, for production of L-isoleucine when the biosynthetic pathway is deregulated (10), BrnFE might significantly contribute to the efflux of this amino acid.

Based on the export properties of the *C. glutamicum* *brnFE* mutants, together with the phylogenetic data reported, we suggest that the two proteins of the system function together as an obligatory pair. The LIV-E family apparently consists of many two-component permeases where one component is predicted

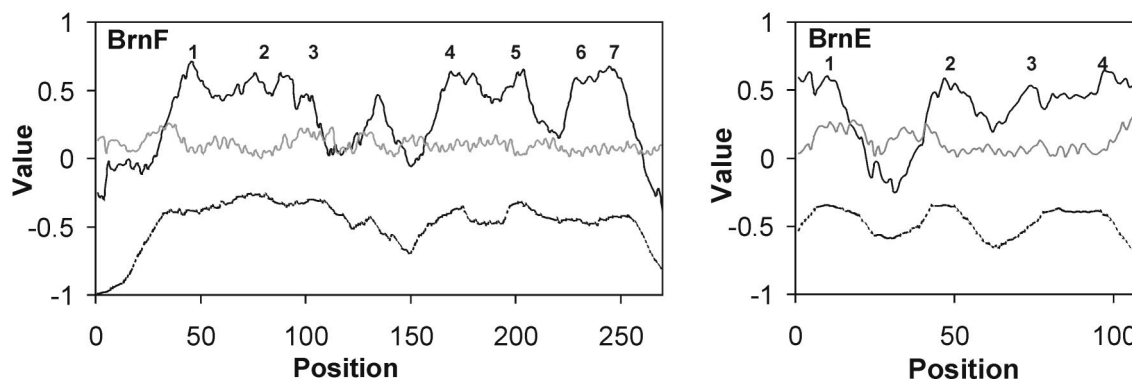


FIG. 8. Average hydropathy (solid line), amphipathicity (shaded line), and similarity (dashed line) for the BrnF and BrnE families. The plots were generated by using the AveHAS program (43), based on multiple alignments generated with the CLUSTAL X program.

to have four TMSs (BrnE) and the other is predicted to have seven TMSs (BrnF), with the latter being better conserved than the former (Fig. 8). Almost all secondary carriers consist of systems with 10 to 14 TMSs, and these can exist at either one or two gene products (15, 29). In the latter case, a homo- or hetero-oligomeric system is presumed to be the active species (5). The BrnFE exporters appear to be no exception when heterodimeric structures are likely to be the active species. This uniform topological characteristic of secondary carriers may reflect the evolutionary process whereby simple channel-type proteins gave rise to carriers, often via intragenic duplication events (31). The variations on this theme, however, are remarkable (30, 32), and BrnFE, with a uniform four plus seven TMSs and two-component heterodimeric topology, represents a unique example. This suggests that the LIV-E family may exhibit unusual mechanistic features among secondary carriers.

The phylogenetic data showed that, except for α -proteobacteria, no prokaryotic organism examined has more than one member of the LIV-E family. However, the results showed that these proteins in the different bacteria are not always orthologues. Instead, early gene duplication events evidently gave rise to multiple paralogues in a primordial bacterium. These paralogues were apparently transmitted faithfully to present-day organisms only in the case of *A. tumefaciens* and *S. meliloti*, for which three paralogous systems remain. In *Mesorhizobium loti*, two paralogues are still present. Many other prokaryotes with completely sequenced genomes lack homologues. In a few cases, only the BrnF homologue or a truncated version of this protein was identified (see Table 3). We suggest that in these organisms these systems are inactive and may be in the process of being genetically deleted from the genomes.

In at least one organism, *B. subtilis*, it appears that a LIV-E family transporter was acquired by a fairly recent horizontal gene transfer event. The *B. subtilis* homologues cluster with those of the α -proteobacteria instead of with the low G+C gram-positive organisms, including *B. halodurans* (Fig. 7). Further indication of horizontal transfer of *brnFE* homologues results from the clustering of the encoding genes with other resistance genes in several organisms (unpublished observations). Thus, a chloramphenicol transporter maps adjacent to the *E. coli brnFE* genes; erythromycin resistance genes are near

the *brnFE* genes in both *Yersinia pestis* and *E. coli*; a cadmium/zinc/cobalt resistance gene is adjacent to such genes in *H. pylori*, and an arsenate resistance gene is found adjacent to *brnFE* in *C. glutamicum* (Fig. 2).

Since the loss of paralogues in most proteobacteria appears to have been random (see Fig. 7), we suggest that the paralogous proteins have retained their ancestral function with little variation in substrate specificity. The differences between paralogues in the α -proteobacteria may reflect dissimilar regulatory characteristics. The fact that the phylogeny of LIV-E family members does not correlate with that for the organismal 16S rRNAs suggests that these carriers do not play roles as essential "housekeeping" proteins such as biosynthesis of essential cellular constituents or degradation of energy sources. In agreement with this suggestion is the fact that the LIV-E family carriers in both *C. glutamicum* and *B. subtilis* are apparently nonessential (2).

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