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Krupnik, Tomas; Sobczak - Elbourne, Iwona; Lolkema, Juke S.

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Turnover and accessibility of a reentrant loop of the Na\textsuperscript{+}-glutamate transporter GltS are modulated by the central cytoplasmic loop

TOMAS KRUPNIK, IWONA SOBCZAK-ELBOURNE*, & JUKE S. LOLKEMA

Department of Microbiology, Groningen Biomolecular Sciences and Biotechnology Institute, University of Groningen, Groningen

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Abstract
GltS of Escherichia coli is a secondary transporter that catalyzes Na\textsuperscript{+}-glutamate symport. The structural model of GltS shows two homologous domains with inverted membrane topology that are connected by a central loop that resides in the cytoplasm. Each domain contains a reentrant loop structure. Accessibility of the Cys residues in two GltS mutants in which Pro351 and Asn356 in the reentrant loop in the C-terminal domain were replaced by Cys is demonstrated to be sensitive to the catalytic state supporting a role for the reentrant loops in catalysis. Saturating concentrations of the substrate L-glutamate protected both mutants against inactivation by thiol reagents, while the presence of the co-ion Na\textsuperscript{+} stimulated the inactivation of both mutants. Insertion of the 10 kDa biotin acceptor domain (BAD) of oxaloacetate decarboxylase of Klebsiella pneumoniae in the central cytoplasmic loop blocked the access pathway from the periplasmic side of the membrane to the cysteine residues in mutants P351C and N356C in the reentrant loop. Kinetically, insertion of BAD increased the maximal rate of uptake 2.7-fold while leaving the apparent affinity constants for L-glutamate and Na\textsuperscript{+} unaltered. The data suggests that insertion of BAD in the central loop results in conformational changes at the translocation site that lower the activation energy of the translocation step without affecting the access pathway from the periplasmic side for substrate and co-ions. It is concluded that changes in the central loop that connects the two domains may have a regulatory function on the activity of the transporter.

Keywords: Glutamate transport, domain structure, central loop, reentrant loop, biotin acceptor domain

Introduction
Escherichia coli K-12 grows on L-glutamate as a sole carbon and nitrogen source. Transport into the cell is mediated by three glutamate transport systems, an ABC transport system specific for glutamate and aspartate and two secondary transporters, the proton symporter GltP and the Na\textsuperscript{+} symporter GltS (Schellenberg and Furlong 1977, Deguchi et al. 1989, Wallace et al. 1990, Kalman et al. 1991). GltP is a member of the DAACS family (Dicarboxylate/Amino Acid:Cation Symporter; TC 2.A.23) (Saier 2000, Slotboom et al. 1999), while GltS belongs to the ESS family (Glutamate Sodium Symporter; TC 2.A.27) of secondary transporters. Members of the ESS family are exclusively found in the bacterial domain. GltS has high affinity for L-glutamate, but, with a lower affinity, also transports D-glutamate, α-methylglutamate, and homocysteic acid. Glutamate uptake activity by GltS was observed only in the presence of Na\textsuperscript{+}, irrespective of the pH, suggesting an obligatory coupling of glutamate and Na\textsuperscript{+} translocation (Tolner et al. 1995). The exact physiological function of each of the three transport systems is not known, but it is likely that they function under different conditions which requires some means of regulation of their activity either at the genetic or at the protein level.

Hydropathy profile analysis suggested that transporters of the ESS family share the same structure with transporters of the 2HCT family [2-HydroxyCarboxylate Transporter, TC 2.A.24 (Sobczak and Lolkema 2005a)], another family found only in bacteria and specific for 2-hydroxycarboxylates like citrate, malate and lactate (MemGen structural classification, class ST[3]) (Lolkema and Slotboom 1998, 2003). Transporters from the two families do not share significant sequence identity. Experimental evidence for the structural similarity has been presented (Dobrowolski et al. 2007, Dobrowolski and Lolkema...
2009, ter Horst and Lolkema, 2010) and the structural model of the transporters in class ST[3] is largely based on studies of GltS of E. coli and the Na\(^+\)-citrate symporter CitS of Klebsiella pneumoniae from the ESS and 2HCT families, respectively. The transporters also share a dimeric quaternary structure (Mościcka et al. 2009, Krupnik et al. 2011).

Figure 1 represents the structural model of the core of the transporters in structural class ST[3] which is the same as the model for GltS of E. coli (Sobczak and Lolkema 2005a). Transporters in other families may have additional transmembrane segments (TMS) at the N- or C-terminus. The core structure consists of two domains sharing the same fold and consisting of 5 TMS each (Lolkema et al. 2005, Lolkema 2006). The two domains have opposite orientations in the membrane, an architecture that is frequently observed in membrane proteins (Sobczak and Lolkema 2005b, Forrest et al. 2011). The N-termini of the N- and C-domain are in the periplasm and cytoplasm, respectively, and the two are connected by a large cytoplasmic loop. Strong evidence for the two domain structure was obtained recently by swapping the order of the N- and C-domain encoding parts of the gltS gene. The engineered gene encoded a transporter GltS\(^{\text{swap}}\) with similar activity as GltS (Dobrowolski and Lolkema 2010). In addition, transport activity was recovered by expressing the two domains as separate proteins from an artificial operon (GltS\(^{\text{split}}\)). In the model, the domains contain a reentrant loop structure positioned in between the 4th and 5th TMS that are believed to be involved in catalysis (Dobrowolski and Lolkema 2009, Sobczak and Lolkema 2003, 2004). A mutational study of the GGXG motif found in the reentrant loops of GltS and CitS showed the functional importance of the loops (Dobrowolski and Lolkema 2009). When replaced by a Cys residue, only a single residue in the reentrant loops in the C-domains of GltS and CitS (Pro351 and S405, respectively) was accessible from the periplasmic side by a bulky membrane impermeable thiol reagent (Dobrowolski and Lolkema 2009). In the 3D-structure, the two domains are believed to fold upon each other with the two reentrant loops forming the translocation pathway (Dobrowolski et al. 2010).

Translocation of substrate and co-ions by this class of transporters is believed to follow an alternate access mechanism in which movement of the two domains relative to one another results in alternate access of the binding sites positioned at the domain interface, similarly as observed for other classes of transporters (Forrest et al. 2011). Here, a potential role of the cytoplasmic loop that connects the two domains in regulation of the activity of the transporters is suggested by demonstrating that an insertion of a 10 kDa protein in the loop increases the transport rate and effects the accessibility of Pro351 at the vertex of the reentrant loop and of the nearby residue Asn356 that in the model is deeper inside the translocation pore.

**Methods**

**Growth conditions and membrane preparation**

GltS and GltS derived transporters were expressed in Escherichia coli strain DH5\(\alpha\) under control of the arabinose promoter using pBAD24 (Invitrogen) derived plasmids. Freshly transformed bacteria were used to inoculate an overnight preculture. A total of 50 ml of preculture was added to 1 l of Luria Broth (LB) medium containing 50 \(\mu\)g/ml ampicillin in a 5 l flask at 37°C that was under continuous shaking at 200 rpm. At an optical density of OD\(_{660}\) = 0.6 0.1% w/v arabinose (Sigma-Aldrich GmbH, Steinheim, Germany) was added to induce expression of the transporters after which the culture was allowed to grow for an additional hour. Cells were harvested by spinning at 8000 rpm at 4°C for 10 min, washed with 50 ml 50 mM KPi, pH 7 at 4°C and resuspended in the same buffer.

Cytoplasmic membranes were isolated by breaking cells resuspended at high density in 100 ml and on ice with a Soniprep 150 sonicator operated at an amplitude of 10 \(\mu\)m and using 15 cycles of 15 s ON and 15 s OFF. Cell debris and unbroken cells were removed by
spinning for 5 min at 13,000 rpm at 4°C. Membranes were collected from the supernatant by ultracentrifugation for 25 min at 80,000 rpm at 4°C.

Right-side-out (RSO) membranes were prepared from an E. coli culture by the osmotic shock procedure as described (Kaback 1974). RSO membranes were resuspended in 50 mM KPi pH 7.0 and aliquoted in 150 μl portions that were stored at −80°C. Membrane protein concentration in the samples was determined by DC protein assay kit (BioRad Laboratories, Hercules, CA, USA).

**Construction of plasmids**

All genetic manipulations were done in E. coli DH5α using standard techniques. Plasmids pGltS, pGltS P351C and pGltS N356C encode the wild type GltS protein and the GltS mutants P351C and N356C, respectively. The constructs that encode GltS variants extended with an N-terminal His₆-tag were described before (Dobrowolski et al. 2007, Dobrowolski and Lolkema 2009). Plasmid pGltS-BAD206 encodes the GltS protein with the Biotin Acceptor Domain (BAD) of oxaloacetate decarboxylase of K. pneumoniae inserted at position 206 of wild type GltS as described before (Krupnik et al. 2011).

Plasmid pGltS-BAD206 was mutated to introduce mutations P351C and N356C in the BAD insertion variant of GltS yielding plasmids pGltS-BAD206 (BAD-P351) and pGltS-BAD206 N356C (BAD206-N356C). The mutants were constructed by PCR using the QuickChange site-directed mutagenesis kit (Stratagene, La Jolla, CA, USA). The presence of the mutations was confirmed by DNA sequencing (Service SX, Leiden, The Netherlands).

**Expression levels of GltS and GltS derivatives**

Expression levels of GltS and GltS derivatives were estimated from the staining intensity of the partial purified proteins after SDS-PAGE. Samples of RSO membranes or cytoplasmic membranes obtained by sonication and containing between 0.5 and 1 mg of membrane protein were solubilized in 1 ml of 50 mM KPi pH 7 buffer containing 400 mM NaCl, 10% glycerol and 1% Triton X-100 for 1 h at 4°C while shaking gently. Solubilized proteins were recovered by spinning for 25 min at 4°C in a Beckman TLA 100.4 rotor at 80,000 rpm. The supernatant was applied to a 50 μl bed volume Ni-NTA resin equilibrated with 50 mM KPi pH 8 buffer containing 600 mM NaCl, 10% glycerol, 0.1% Triton X100, and 30 mM imidazole and left overnight at 4°C while shaking. Next, the resin was spun down in a table centrifuge for 1 min at 3,500 rpm and the supernatant was removed. Subsequently the resin was washed twice with equilibration buffer and once with the same buffer but containing 40 mM imidazole. Proteins were eluted with 50 μl of 50 mM KPi pH 7 buffer containing 600 mM NaCl, 10% glycerol, 0.1% TritonX, 500 mM imidazole and analyzed by SDS-PAGE. Staining intensity of the appropriate bands was visualized and quantified using a Fujifilm LAS-4000 image analyzer (Fuji).

**Transport studies**

RSO membrane vesicles were diluted to 1 mg/ml in 50 mM KPi pH 6 containing 10 or 100 mM NaCl as indicated. Aliquots of 100 μl were incubated for 2 min at 30°C with 10 mM K-ascorbate and 100 μM phenazine methosulfate (PMS) under a flow of water-saturated air and continuous stirring with a magnetic bar. At t = 0, [14C]-proline or L-[14C]glutamate was added to final concentrations of 2.2 and 2.9 μM, respectively. Uptake was stopped at 5, 10, 20, 40 s by the addition of 2 ml of ice-cold 0.1 M LiCl followed by immediate filtering over cellulose nitrate filters (0.45 μm pore size) and washing of the filter with an additional 2 ml of the LiCl solution. Filters were collected and their radioactivity was measured in a liquid scintillation counter. Background activities due to the chromosomal copy of the gltS gene were determined in parallel experiments by the uptake activity of membranes treated with an excess of NEM (GltS-P351, GltS-N356C and BAD206-N356C) or by the uptake of membrane vesicles derived from cells containing the empty vector (BAD206-P351).

**Labeling studies**

RSO vesicles were incubated for 5 min at room temperature in 50 mM KPi pH 7 buffer with or without the indicated concentrations of L-glutamate (K⁺ salt) and Na⁺ (chloride salt). The samples were treated with 1 mM of NEM or 0.25 mM of AmdoS for 10 min unless otherwise stated. The thiol reagent solutions were prepared freshly. The treatment was stopped by addition of 10 mM DTT. Subsequently, the membranes were collected by spinning at 7,000 rpm for 5 min at 4°C, followed by resuspension and three times washing in 50 mM KPi pH 7 buffer. RSO membranes were resuspended in 50 mM KPi pH 6 containing 100 mM NaCl. Control samples were included that were not treated with the thiol reagents but with and without L-glutamate. Rates of uptake in these samples were comparable, indicating efficient removal of L-glutamate by the washing procedure.
Cells expressing the GltS variants were treated in the same way, followed by a second treatment with 0.1 mM fluorescent maleimide (FM) for 1 min at 20°C. The reaction was stopped with 1 mM DTT and the cells were washed twice with 50 mM KPi pH 7 buffer. Direct labeling of the proteins by FM was performed in the same way but the first labeling step with NEM or AmdS was omitted. Following purification and SDS-PAGE, the fluorescence intensity of the bands was visualized using a Fujifilm LAS-4000 image analyzer (Fuji).

Materials

Ampicillin was obtained from Roche Diagnostic GmbH, Mannheim, Germany, arabinose from Sigma-Aldrich GmbH, Steinheim, Germany, and [1,5-14C]-citrate and L-[14C]-glutamate from Amer sham Pharmacia, Roosendaal, The Netherlands.

Results

Kinetics of L-glutamate uptake by RSO membranes containing GltS and GltS-BAD206

Before (Krupnik et al. 2011), the biotin acceptor domain (BAD) of the oxaloacetate decarboxylase of Klebsiella pneumoniae, a 10 kDa protein, was inserted at position 206 in the central loop of the Na+-dependent glutamate transporter GltS (see Figure 1) for cross-linking studies. Transport activity of GltS-BAD206 and the wild type transporter was measured by the rate of uptake of radiolabeled L-glutamate by right-side-out (RSO) membrane vesicles derived from E. coli DH5α producing the proteins with six histidine residues added at the N-terminus (His6-tag). RSO membrane vesicles rather than whole cells were used to prevent further metabolism of the substrate following uptake. L-glutamate accumulation was driven by a proton motive force that was generated using the artificial electron donor system ascorbate/PMS (Konings et al. 1971). Membrane vesicles prepared from the host cells contain a low level of Na⁺-dependent L-glutamate transport activity due to expression of the gltS gene encoded on the chromosome. Background activity was approximately 20–25% of membranes containing in addition plasmid encoded GltS (Figure 2A) (Dobrowolski et al. 2007, Dobrowolski and Lolkema 2009).

RSO membranes containing GltS-BAD206 took up L-glutamate with an initial rate that was within experimental error the same as the rate observed with membranes containing wild-type GltS (Figure 2A).

To determine the specific activity of the insertion mutant relative to wild-type GltS, the expression levels and quality of the membranes were determined. The latter was inferred from pmf driven uptake of radiolabeled proline by an endogenous transporter using the same batch of membranes under the same conditions (Figure 2B). Initial rates of uptake of proline by the membranes containing the plasmid encoded GltS and GltS-BAD206 transporters were not significantly different, while the control membranes took up proline at a 3-fold higher rate. It follows that the contribution of the endogenous GltS transporter to the L-glutamate uptake rates (Figure 2B) by the former two types of membranes is largely overestimated and amounts to less than 10%. The latter result is in agreement with experiments where the background rate was estimated by inactivating a mutant GltS (Krupnik et al. 2011). More importantly, it follows that membranes derived from cells overexpressing GltS or GltS-BAD206 are of comparable quality allowing for a direct comparison of the activity of the transporters in the two types of membranes. The His-tagged GltS proteins were partially purified by Ni²⁺-NTA affinity chromatography from the same batch of RSO membranes used in the uptake experiments. Densitometric analysis of the intensity of the Coomassie Brilliant Blue stained bands after SDS-PAGE revealed an intensity ratio of GltS over GltS-BAD206 of 2.2 (Figure 2C). Assuming the same staining efficiency of GltS and the BAD domain and taking into account the different molecular masses of GltS and GltS-BAD206 (43.5 and 52.9 kDa, respectively), the expression level of the insertion mutant in the membranes would be a factor of 2.7 lower. The relation between uptake activity and amount of transporter protein in the membranes was determined by preparing RSO membranes from cells harboring the appropriate plasmid that were induced with 0, 0.001 and 0.1% arabinose in the medium (Figure 2D). Induction with 0.1% arabinose corresponds to the standard growth condition. No protein was detected with the uninduced cells. For both GltS and GltS-BAD206, linear relations were obtained that extrapolated back to an initial rate of 0.25 nmol/min/mg, which corresponds to the uptake activity of the uninduced cells. It follows that for both types of membranes, uptake activity was proportional to the amount of plasmid encoded protein in the membrane. In conclusion, the specific activity of GltS-BAD206 was 2.7 times higher than observed for wild type GltS at concentrations of L-glutamate and Na⁺ of 2.9 μM and 10 mM, respectively.

In the presence of 10 mM Na⁺, the total initial rate of L-glutamate uptake at increasing L-glutamate concentrations in the range between 2.9 and 52.9 μM followed a saturation curve with an affinity constant K_M of 13.5 ± 3.5 μM and a maximal rate of 15.0 ±
1.6 nmol/min.mg of membrane protein for membranes containing wild type GltS (Figure 3, insert). The affinity constant was a factor of two lower than the value reported before (Tolner et al. 1995).

At a constant L-glutamate concentration of 2.9 µM, initial rates increased sigmoidal with increasing concentrations of Na⁺ covering a range between 0.1 and 100 mM (Figure 3). The rate was half the maximal rate at a Na⁺ concentration of 5.8 ± 0.5 mM. The sigmoidal curve indicates that more than one Na⁺ ions are symported with the L-glutamate anion, in agreement with the electrogenic nature of transport reported before (Tolner et al. 1995). The same analysis with membranes containing the GltS-BAD206 insertion mutant yielded an affinity constant for L-glutamate that was not significantly different than observed for the wild type, \( K_M = 15.0 ± 3.0 \) µM (Table I). Also, the Na⁺ concentration giving half the maximum rate was within the error observed for the wild type, \( K_{0.5} = 5.6 ± 0.3 \) mM. For both analyses, no correction for the background activity was applied because the difference between the parameters was not significant. Since the kinetic parameters of wild type GltS and the insertion mutant GltS-BAD206 are similar within experimental error, it follows that insertion of the BAD domain in the cytoplasmic central loop of GltS results in a 2.7-fold increase of the maximal turnover rate per GltS molecule (Table I).

**Inactivation of reentrant loop mutants P351C and N356C by thiol reagents**

Residues Pro351 and Asn356 of GltS were replaced with Cys residues before (Dobrowolski and Lolkema 2009) as part of a cysteine scanning mutagenesis study of the putative cytoplasmic reentrant loop in the C-terminal domain of GltS (see Figure 1). In a stretch of 18 residues, mutation of residues
Pro351 and Asn356 to Cys rendered the transporter sensitive to inactivation by the membrane permeable thiol reagent N-ethylmaleimide (NEM). In addition, P351C was also inactivated by membrane impermeable 4-acetamido-4′-maleimidylstilbene-2,2′-disulfonic acid (AmdiS) indicating accessibility of the thiol from the periplasmic side of the membrane. Pretreatment of RSO membranes containing the mutants in 50 mM KPi buffer pH 7 for 10 min at increasing concentrations of the thiol reagents followed by measurement of the residual activity revealed a much higher sensitivity of the P351C mutant than the N356C mutant for inactivation by NEM (Figure 4A, 4B, open bars).

Concentrations of NEM as low as 0.1 mM completely inactivated the former while a 30-fold higher concentration was needed for complete inactivation of the latter. Uptake activities by the membranes were corrected for the background activity of the mutants was measured by treating the membranes with the thiol reagents (Dobrowolski et al. 2007). The sensitivity of the P351C mutant was slightly higher than the N356C mutant for inactivation by concentrations of AmdiS that were 10 times higher than the concentration required to inactivate the wild type version (Figure 4C). It follows that insertion of the BAD domain in the cytoplasmic central loop of GltS makes the residue at the vertex of the putative reentrant loop positioned at the periplasmic side (Figure 1) virtually inaccessible for thiol reagents NEM and AmdiS.

**Effect of substrate L-glutamate and co-ion Na⁺ on inactivation of the P351C and N356C mutants**

The presence of a saturating concentration of 1 mM L-glutamate ($K_M = 13.5 \mu M$; see Figure 3, insert) protected both the P351C and N356C mutants against inactivation by the thiol reagents. Treatment of RSO membranes containing the P351C mutant with 0.25 mM of AmdiS for 10 min resulted in 78% of inactivation of uptake activity, while the presence of 1 mM of L-glutamate during the inactivation essentially completely protected the transporter (Figure 5A).

Full protection was also observed when NEM instead of AmdiS was used as the thiol reagent (not shown). NEM inhibited the N356C mutant catalyzed uptake activity by 62% when used at a concentration of 1 mM of NEM for 10 min, while no inactivation was observed in the presence of 1 mM L-glutamate (Figure 5B). In the BAD inserted variant of the N356C mutant a similar protection was observed with a concentration of 3 mM NEM (not shown).

The effect of the co-ion Na⁺ on the inactivation of the mutants was measured by treating the membranes with the thiol reagents in the presence of 10 mM Na⁺.
and a saturating concentration of 100 mM of Na\(^+\) (K\(_{0.5}\) = 5.8 mM; Figure 3). For both mutants, the presence of Na\(^+\) had the opposite effect than the presence of L-glutamate discussed above. Under the conditions of the experiment and within experimental error, AmdiS completely inhibited the P351C mutant in the presence of Na\(^+\) and the same was observed for the inhibition of the N356C mutant by NEM treatment (Figure 6, open bars).

Apparently, the presence of Na\(^+\) increased the sensitivity to the thiol reagents. Insertion of the BAD domain in the P351C mutant rendered the mutant insensitive to inactivation, also in the presence of Na\(^+\). The residual activity was down to 80–90% under all conditions tested (Figure 6A, shaded bars). This level of inactivation is unspecific and due to a small inactivating effect on the pmf generating system by the treatment (Dobrowolski et al. 2007, Dobrowolski and Lolkema 2010). Treatment of the RSO membranes containing the BAD206-N356C mutant with 3 mM NEM for 20 min reduced the initial rate of uptake down to 26%. In contrast to what was observed with the N356C mutant, the presence of Na\(^+\) protected the insertion mutant against inactivation resulting in 48% and 92% residual activity in the presence of 10 and 100 mM of Na\(^+\), respectively (Figure 6B, shaded bars). It follows that where the presence of Na\(^+\) increases the reactivity of the thiol in the putative reentrant loop in the N356C mutant, the opposite effect was observed when the BAD domain is inserted in the central loop.

**Accessibility of reentrant loop mutants P351C and N356C in whole cells**

*E. coli* DH5\(\alpha\) cells producing GltS and the reentrant loop mutants P351C and N356C were treated with the fluorescent thiol reagent fluorescein maleimide to detect thiols accessible at the outer cell surface. Following treatment with the label for 10 min in KP\(\text{i}\) pH 7 buffer, the proteins were purified from the cells and labeling was analyzed by fluorescence imaging of the gel after SDS-PAGE. The three GltS alleles showed more or less the same intensity after protein staining (Figure 7A).
Wild type GltS showed no fluorescence indicating that under the conditions of the experiment none of the endogenous Cys residues were accessible for fluorescein maleimide added to the outside of the cells (ter Horst and Lolkema 2010). The same was observed for the N356C mutant showing that the Cys residue at the position of Asn356 was not accessible as well by this reagent. In contrast, the P351C protein was clearly fluorescent indicating that the Cys residue at the position of Pro351 was exposed at the external cell surface. Pretreatment of cells containing the P351C mutant with membrane impermeable AmdiS prevented subsequent labeling with the fluorescent label demonstrating that position 351 was also accessible for the bulky and charged AmdiS in whole cells (Figure 7B). The presence of 1 mM of L-glutamate during the treatment with AmdiS followed by washing of the cells to remove L-glutamate did result in fluorescent labeling indicating that the substrate protected against AmdiS labeling.

The results show the same behavior of GltS and the mutants P351C and N356C in whole cells and RSO membranes. Thiols of the wild type and N356C mutant are not accessible from the outside, in contrast to the thiol at position 351 of the P351C mutant. The latter is protected from labeling in the presence of glutamate.

Discussion

The two domains of the transporters in the different families of structural class ST[3] in the MemGen classification are connected by a large cytoplasmic loop that shows up as a typical dip in the hydropathy profile of the proteins as is seen for two-domain transporters in other structural classes as well. The GltS and CitS proteins as well as the H+/lactose symporter LacY, a major facilitator superfamily (MFS) transporter (class ST[1]), have been produced as active ‘split’ permeases consisting of the two domains as separate proteins which shows that a covalent connection between the domains is not essential (Bibi and Kaback 1990, Dobrowolski et al. 2010, Dobrowolski and Lolkema 2010). Nevertheless, in nature, such constructs are not observed, suggesting that a loop linking the two
domains did have some advantage during evolution. This is in contrast to, for instance, ABC transporters or phosphoenolpyruvate dependent phosphotransferase system transporters where the functional units consist of either a single polypeptide containing multiple domains or a complex built from different subunits. In some families of class ST[3] secondary transporters, the connecting loop is extended and probably folds into a cytoplasmic domain, suggesting a possible function of the loop in the activity of the transporter (Lolkema et al. 2005, Barabote et al. 2006, Nanatani et al. 2007). In a previous study we noticed that insertion of the 10 kDa biotin acceptor domain of K. pneumoniae in the central cytoplasmic loop of GltS, while not significantly affecting the uptake activity in RSO membranes under the conditions employed, seemed to result in less of the hybrid protein in the membranes (Krupnik et al. 2011). A detailed analysis presented here, taking into account different effects that might contribute to the observed uptake rates showed that the maximal rate catalyzed by the GltS-BAD206 transporter protein was a factor of 2.7 higher than observed for GltS while the kinetic affinities for L-glutamate and the Na\(^+\)-ions were unaltered. This suggests that the wild type transporter protein may not be in the “optimal” conformation when embedded in the membrane of isolated RSO membrane vesicles. It also suggests that changes in conformation of the central loop may be used to modulate the activity of the transporter thereby providing a post-translational regulatory mechanism for glutamate uptake activity to the cell. Possible effectors might be cytoplasmic pH affecting the conformation of the central loop or a cellular metabolite that binds to the loop.

The structural model of GltS shows putative reentrant loops in the N- and C-terminal domains (Figure 1) that are believed to form (part of) the translocation site in the 3D structure. Close vicinity of the two reentrant loops was demonstrated by cross-linking studies (Dobrowolski et al. 2010). The present data demonstrates that this part of the protein is likely to be involved in the substrate and co-ion binding pocket and supports the involvement of the reentrant loops in catalysis. Cysteine scanning mutagenesis of the putative reentrant loop region in the C-domain of GltS identified only position 351 in the P351C mutant being accessible to membrane impermeable AmdiS suggesting that the residue at this position would be the one most exposed to the periplasm and, therefore, be at the vertex of the reentrant loop (Dobrowolski et al. 2007). Consequently, Asn356 would be in the ingoing stretch, inside the pore (see Figure 1). The relative reactivity of the P351C and N356C mutants with thiol reagents reported here are consistent with this view. The reactivity of the P351C mutant with NEM was found to be at least one order of magnitude higher than observed for the N356C mutant, while the latter does not react with AmdiS at all. The observations are in line with a lower accessibility of the cysteine residue at position 356. The accessibility of the cysteines at the position of Pro351 and Asn356 was strongly reduced in the presence of a saturating concentration of L-glutamate and enhanced in the presence of the co-ion. The effect of the presence of L-glutamate and Na\(^+\) on the reactivity of the two mutants was similar suggesting that for both membrane impermeable AmdiS and membrane permeable NEM the access pathway to the cysteine residues was from the periplasmic side of the membrane. Binding of L-glutamate or Na\(^+\) added at the periplasmic side recruits the transporter molecules to the outward-facing conformation. Protection by L-glutamate most likely arises from steric hindrance in the access pathway or from direct involvement of the target residue in substrate binding. In this respect it may be noted that both Pro351 and Asn356 are highly conserved residues in the ESS family. Enhancement of the reactivity in the presence of Na\(^+\) would be a direct effect of locking the transporters in the outward-facing conformation without the need of recruiting (part of) the proteins from the inward-facing conformation during inactivation. Because of its small size, the Na\(^+\) ions are unlikely to restrict the access pathway.

The change in turnover rate of GltS by the insertion of the BAD domain in the loop that connects the two domains parallels changes in the accessibility of residues in the cytoplasmic reentrant loop in the C-domain that may shed some light on the mechanism of the activation. Insertion of the domain in the central loop reduces the accessibility of positions 351 and 356 at the periplasmic side of the membrane. The P351C insertion mutant appears to be insensitive to AmdiS as well as NEM, irrespective of the catalytic state of the transporter (Figures 4 and 6). The N356 insertion mutant showed a reduced reactivity with NEM, but at higher concentrations of the thiol reagent, inactivation was clearly observed (Figure 6B). Importantly, binding of Na\(^+\) protected the N356C-BAD206 mutant against NEM, while it enhanced the inactivation of N356C strongly suggesting different access pathways. If the enhanced reactivity of N356C is explained by recruitment in the outward facing state (see above), it follows that the N356C-BAD206 mutant reacts in the inward facing conformation with membrane permeable NEM coming from the cytoplasm. Summarizing, the dominant effect of the insertion of the BAD domain is a strong restriction of the access pathway to positions 351 and 356 in the reentrant loop for thiol reagents from the periplasmic side of the membrane. At the same time, position 356, but not position 351, appears
to become more accessible from the cytoplasmic side. The equilibrium of the transition between the inward and outward facing conformations appears to be shifted to former conformation.

Conclusion

The Na+-glutamate transporter GltS is believed to consist of two homologous domains that are connected by a large hydrophilic cytoplasmic loop. The translocation site is believed to be formed at the interface of the two domains where two reentrant loops interact. Cysteine residues engineered in the reentrant loop in the C-terminal domain that enters the membrane from the cytoplasmic face of the membrane are accessible from the periplasm and their reactivity is modulated by the presence of substrate and co-ion. Insertion of the 10 kDa BAD polypeptide in the loop that connects the two domains in the cytoplasm blocks the accessibility of the cysteine residues in the reentrant loop from the periplasm. Kinetically, the insertion results in an increase of the maximal rate of transport by a factor of 2.7. It follows that changes in the connecting loop affect turnover and may have a function in regulation of transporter activity.

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Declaration of interest: The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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