Transport Rates of a Glutamate Transporter Homologue Are Influenced by the Lipid Bilayer

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Background: Structures of GltPh have been solved in the absence of a lipid bilayer.

Results: Transport rates of GltPh are influenced by lipid bilayer composition.

Conclusion: Transmembrane orientation and lipid-protein interactions influence transport in GltPh.

Significance: This provides a specific example of how interactions between lipid headgroups and membrane-bound proteins can influence function.

The aspartate transporter from *Pyrococcus horikoshii* (GltPh) is a model for the structure of the SLC1 family of amino acid transporters. Crystal structures of GltPh provide insight into mechanisms of ion coupling and substrate transport; however, structures have been solved in the absence of a lipid bilayer so they provide limited information regarding interactions that occur between the protein and lipids of the membrane. Here, we investigated the effect of the lipid environment on aspartate transport by reconstituting GltPh into liposomes of defined lipid composition where the primary lipid is phosphatidylethanolamine (PE) or its methyl derivatives. We showed that the rate of aspartate transport and the transmembrane orientation of GltPh were influenced by the primary lipid in the liposomes. In PE liposomes, we observed the highest transport rate and showed that 85% of the transporters were orientated right-side-out, whereas in trimethyl PE liposomes, 50% of transporters were right-side-out, and we observed a 4-fold reduction in transport rate. Differences in orientation can only partially explain the lipid composition effect on transport rate. Crystal structures of GltPh revealed a tyrosine residue (Tyr-33) that we propose interacts with lipid headgroups during the transport cycle. Based on site-directed mutagenesis, we propose that a cation-π interaction between Tyr-33 and the lipid headgroups can influence conformational flexibility of the trimerization domain and thus the rate of transport. These results provide a specific example of how interactions between membrane lipids and membrane-bound proteins can influence function and highlight the importance of the role of the membrane in transporter function.
branes containing His-GltPh were isolated and solubilized with grown and purified as described previously (13). Briefly, mem-
determined by swelling of vesicles measured by light scattering (Avanti Polar Lipids) and treated with Triton X-100 at a ratio by extrusion through 400-nm polycarbonate membranes 
nitrogen and thawed at least six times. Liposomes were formed resuspended in internal buffer (100 mM KCl, 20 mM HEPES-Tris, pH 7.5, 1 μM valinomycin, 100 nM L-[3H]aspartate) at 30 °C. At each time point, an aliquot was removed and diluted 10-fold into ice-cold quench buffer (100 mM LiCl, 20 mM HEPES-Tris, pH 7.5) fol-
the right-side out (RSO) orientation of GltPh, liposomes containing 1 mM MTSET were me-
gulated as described previously (13). Briefly, mem-
brane-containing His-GltPh were isolated and solubilized with n-dodecyl β-o-maltopyranoside (Anatrace), and protein was purified using nickel-nitrotriacetic acid resin (Qiagen). The histidine tag was subsequently removed by digestion with thrombin (10 units/mg of protein), and the protein further purified on a size exclusion column where the detergent was exchanged to MTSET that gave 10% inhibition (100 μM, which was immediate. The residual amount of MTSET was very quench buffer under vacuum; filters were combined with scin-
tillation fluid and assayed for radioactivity using a Trilux β counter (PerkinElmer Life Sciences).

The Na+ dependence of L-[3H]aspartate transport was deter-
by varying the extraliposomal Na+ concentration from 0.3 to 100 mM. Choline chloride was used to balance osmolarity. The K0.5 of aspartate transport was measured by varying the extraliposomal L-[3H]aspartate concentration from 3 to 1000 nM, maintaining NaCl concentrations at 100 mM. The influence of lipid bilayer composition on the transport-active limb of the GltPh, transport cycle was explored by loading liposomes with 100 mM NaCl, 20 mM HEPES-Tris, pH 7.5, 100 μM L-aspartate and diluting liposomes into uptake buffer as described previously (11). Initial rates of transport represent the amount of L-[3H]aspartate transport over the linear portion of the time course.

Thiol modification was carried out by treating liposomes with 1 mM 2-(trimethylammonium)ethyl methanethiosulfon-
ate (MTSET) for 5 min at room temperature. This isolated the inside out orientation of GltPh. MTSET is membrane-imper-
meable, and inhibition is complete under these conditions. Higher concentrations of MTSET or longer reaction times did not result in more than a 50% reduction in transport (see Fig. 2B). The reducing agents dithiothreitol (DTT; 1 mM) and tris(2-carboxyethyl) phosphine (TCEP; 20 mM) were used to reduce GltPh thiol modification. To isolate the right-side out (RSO) orientation of GltPh, liposomes containing 1 mM MTSET were diluted into uptake buffer containing 20 mM TCEP, which is able to rescue the RSO orientation. The uptake was performed as soon as the liposomes were in the TCEP-containing buffer. The rate of transport was ~50% of that of untreated liposomes, which demonstrates that the reduction of A364C by excess TCEP is immediate. The residual amount of MTSET was very low (15 μM, which was ~6-fold lower than the concentration of MTSET that gave 10% inhibition (100 μM; see Fig. 2B).

Biochemical Evaluation of Transmembrane Orientation—
Liposomes were reconstituted with His-tagged GltPh (20 μg of protein/mg of lipid) and subjected to treatment with thrombin (10 units/mg of protein). Proteolysis was terminated by the addition of 10 mM EDTA and 1 mM AEBSF. Liposomes were...
centrifuged at 150,000 × g for 20 min, and pellets were resuspended in 12% SDS loading buffer and loaded on a 10–20% SDS-polyacrylamide gel.

Amido Black Quantification—A modified Amido Black assay (15), which is used to quantify microgram quantities of protein in the presence of milligram quantities of lipids, which can often interfere with more traditional visual colorimetric assays, was conducted. Liposome samples were dissolved with SDS, and protein was precipitated with trichloroacetic acid (Sigma) by vortexing. Samples were filtered under vacuum, and mixed cellulose ester membranes were stained with Amido Black stain. Protein spots were excised and soaked in elution solution, and the absorbance of the eluate was measured at 530 nm.

Data Analysis—Analysis of kinetic data was conducted using GraphPad Prism 5.0 (GraphPad Software, La Jolla, CA). All values presented are the mean ± S.E. of experiments performed in triplicate. One-way analysis of variance was performed with a Dunnett’s post hoc test. The p values < 0.05 were interpreted as statistically significant and are indicated in figures as * for p < 0.05, ** for p < 0.01, and *** for p < 0.001.

RESULTS

The Lipid Bilayer Influences Aspartate Transport Rates by GltP<sub>β</sub>—The aim of this study was to investigate the role of the lipid environment in GltP<sub>β</sub> function by varying the lipid composition of the liposomes in which GltP<sub>β</sub> is reconstituted. Aspartate transport via GltP<sub>β</sub> is coupled to the co-transport of three Na<sup>+</sup> ions, and in the presence of an inwardly directed Na<sup>+</sup> gradient, GltP<sub>β</sub> will transport aspartate into liposomes (Fig. 1, C and D). A common lipid condition used when studying GltP<sub>β</sub> is a 3:1 mixture of Escherichia coli polar lipid extract to trimethyl PE. Trimethyl PE is commonly referred to as 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine; however, we have chosen to use the N-methyl nomenclature here. The E. coli polar lipid extract is composed of ~10% CL, a bulky mitochondrial membrane lipid; ~20% PG; and ~70% PE (10, 12, 13, 16).

Purified GltP<sub>β</sub> protein was reconstituted in liposomes in which the primary phospholipid species was PE or its monomethyl, dimethyl, or trimethyl derivatives. F, initial rates of transport for wild-type GltP<sub>β</sub> reconstituted in liposomes as shown in E. Data represent the mean of experiments performed in triplicate, and error bars indicate S.E.

Kinetic Analysis of Transport by GltP<sub>β</sub>—We sought to determine whether our observed influence of the lipid bilayer composition was due to changes in the apparent affinity of Na<sup>+</sup> and/or aspartate for GltP<sub>β</sub>. In the presence of saturating Na<sup>+</sup> concentrations, we determined that the half-maximal rates of aspartate transport were not significantly different comparing the mono-, di-, or trimethyl lipid compositions with PE (Table 1). The Na<sup>+</sup> dependence of transport was also determined by measuring the rate of transport with increasing concentrations of Na<sup>+</sup> in the presence of saturating L-[3H]aspartate, and simi-
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| Table 1 |
| --- |
| The effect of lipid bilayer composition on kinetics of transport by Glt$_{ph}$ |
| Shown are the initial rate of L-[3H]aspartate transport as a function of [L-[3H]aspartate] in the outside buffer in the presence of saturating Na$^+$ concentrations and the initial rate of L-[3H]aspartate transport as a function of [Na$^+$] in the outside buffer in the presence of saturating L-[3H]aspartate concentrations. Kinetic parameters for Glt$_{ph}$ reconstituted in varying lipid compositions were derived by fitting the L-aspartate concentration-response data to the Michaelis-Menten equation and fitting the Na$^+$ concentration-response data to the Hill equation. The values are averaged from experiments performed in triplicate, and errors indicate S.E. No significant difference was observed for $K_{m,O}$ values. |
| **Aspartate $K_{m,O}$ (nM)** | **Aspartate $V_{max}$ (nmol mg$^{-1}$ min$^{-1}$)** | **Na$^+$ $K_{m,O}$ (mM)** |
| **PE** | **Monomethyl PE** | **Dimethyl PE** | **Trimethyl PE** |
| 84.7 ± 8.6 | 57.9 ± 8.8 | 62.8 ± 5.9 | 67.5 ± 10.6 | 82.4 ± 14.6 |
| 10.9 ± 0.3 | 11.1 ± 0.4 | 6.1 ± 0.1 | 3.4 ± 0.1 | 1.6 ± 0.1 |
| 2.6 ± 0.2 | 3.2 ± 0.5 | 3.2 ± 0.6 | 3.4 ± 0.5 | 3.0 ± 0.2 |

In preliminary studies, we observed that the RSO and ISO populations were each contributing ~50% of the transport of untreated liposomes (Fig. 2C). When 1 mM MTSET was present on both sides of the liposome, transport was reduced to background levels, suggesting that both RSO and ISO transporters had been silenced by MTSET modification. Function could be rescued to pre-MTSET modification levels by treatment with DTT, which is a membrane-permeable reducing agent. Finally, we showed that transport by Cys-less Glt$_{ph}$ was not affected by application of MTSET (Fig. 2C). To confirm that the ~50% reduction in transport represents ~50% of the protein oriented in RSO or ISO orientation, we reconstituted His-tagged Glt$_{ph}$ (Glt$_{ph}$-His) into the 3:1 control lipid mixture and treated these liposomes with thrombin. Samples were separated on a 10–20% acrylamide gel, and two bands that correspond to Glt$_{ph}$-His and cleaved Glt$_{ph}$ were observed. We performed a time course of thrombin digestion. After 1 min, two bands appeared, the reaction was stable up to 6 h, and the bands were of equal density (Fig. 3A). This demonstrates that thrombin can only access around half of the protein to cleave the His tag and correlates with our uptake data where MTSET treatment reduced transport rates by ~50%. Taken together, these results demonstrate that A364C is only accessible from one side of the membrane, and through the use of cysteine-reactive methanesulfonate agents and reducing agents, we can selectively modify each population of transporters (RSO or ISO).

This thiol modification strategy allowed us to probe the sidedness of the transporters to determine the kinetic parameters of the RSO and ISO populations as well as understand the effect that lipid bilayer composition has on the transmembrane orientation of Glt$_{ph}$. We measured half-maximal rates of aspartate transport in the presence of 100 mM Na$^+$ and found that Glt$_{ph}$ has an aspartate $K_{m,O}$ of 39.3 ± 6.7 and 140.7 ± 26.2 nM for the RSO and ISO orientations, respectively (Table 2). This observation supports the role of Glt$_{ph}$ as a concentrative aspartate transporter that has a higher affinity for aspartate when the binding site is facing the external side of the cell. We also determined that the Na$^+$ dependence of transport was similar for both RSO and ISO orientations with Na$^+$ $K_{m,O}$ values of 2.2 ± 0.2 and 2.1 ± 0.3 mM, respectively (Table 2).

We then used this strategy to selectively inhibit the RSO transporters to investigate the effect of the different lipid compositions on the orientation of Glt$_{ph}$ in the liposomes. When A364C was reconstituted in PE liposomes, MTSET application reduced the initial rates of transport by 84.0 ± 3.3% (Fig. 3B), indicating that ~84% of the transporters were in the RSO orientation. When reconstituted in monomethyl and dimethyl PE liposomes, rates were reduced by 76.0 ± 1.9 and 64.0 ± 2.3%, respectively. Finally, L-[3H]aspartate transport in trimethyl PE liposomes was reduced by 51 ± 1% following treatment with MTSET (Fig. 3B). These results demonstrate that when Glt$_{ph}$ was reconstituted in lipids in which the headgroups are
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![Diagram A](image1.png)

**FIGURE 2.** Thiol modification strategies to observe sided properties of Glt<sub>ph</sub>. A, A364C was only accessible from one side of the membrane. Following treatment of liposomes with 1 mM MTSET (red triangles) A364C is modified resulting in inhibition of the RSO transporters while the ISO transporters are still functional as A364C is unmodified (represented by SH). The strategy for modifying ISO transporters involved loading liposomes with MTSET (MTSET<sub>ISO</sub>) (100 mM KCl, 20 mM HEPES-Tris, 1 mM MTSET) followed by performing uptake in buffer containing TCEP (orange circles) (100 mM NaCl, 20 mM HEPES-Tris, pH 7.5, 1 μM valinomycin, 20 mM TCEP, 100 mM L-[3H]aspartate) to selectively rescue RSO transporters. B, concentration-response data for the effect of increasing incubation concentrations of MTSET on A364C in liposomes. Data were fit to an inhibitor concentration-response curve for display purposes. 1 mM MTSET was incubated with liposomes containing A364C for 10 min. MCAT was performed in buffer containing 100 mM NaCl, 20 mM HEPES-Tris, pH 7.5. Concentration-response data for the effect of MTSET treatment on A364C in liposomes. Data were fit to an inhibitor concentration-response curve for display purposes. C, concentration-response data for the effect of MTSET treatment on A364C in liposomes. Data were fit to an inhibitor concentration-response curve for display purposes. C, concentration-response data for the effect of MTSET treatment on A364C in liposomes. Data were fit to an inhibitor concentration-response curve for display purposes.

![Diagram B](image2.png)

**FIGURE 3.** Inhibition of RSO transporters reveals the distribution of transmembrane orientation. A, strategy for thrombin cleavage of His-tagged Glt<sub>ph</sub>, reconstituted in 3:1 E. coli polar lipid extract:trimethyl PE. The hexahistidine tag is susceptible to thrombin cleavage (scissors). Thrombin (10 units/mg of protein) was added to liposomes containing His-Glt<sub>ph</sub> (20 μg/mg of lipid). At the indicated time points, the digest was stopped with 1 mM AEBSF and 10 mM EDTA. Samples were run on a 10–20% SDS-polyacrylamide gel. B, percentage of transport remaining after 1 mM MTSET (in 100 mM KCl, 20 mM HEPES-Tris, pH 7.5) incubation of A364C reconstituted in liposomes. Data were normalized to transport of untreated liposomes in each lipid composition. Data represent the mean of experiments performed in triplicate, and error bars indicate S.E.

**TABLE 2**

| Lipid Composition | ISO | RSO |
|-------------------|-----|-----|
| Aspartate K<sub>0.5</sub> (nm) | 140.7 ± 26.2 | 39.3 ± 6.7 |
| Na<sup>+</sup> K<sub>0.5</sub> (mm) | 2.1 ± 0.3 | 2.2 ± 0.2 |

unmethylated (such as PE) the transporters are more correctly oriented. In addition, the outwardly directed substrate binding site of Glt<sub>ph</sub> had a 3.5-fold higher affinity for aspartate than the inwardly directed binding site (Table 2). In combination, these results only account for an ~18% difference in the rate of transport we observed and cannot fully account for the ~75% difference in transport rate observed between PE- and trimethyl PE liposomes. Therefore, other factors to explain these differences must exist.

**Lipid Bilayer Composition Influences Substrate-bound Isomerization of Glt<sub>ph</sub>**—The transport cycle of Glt<sub>ph</sub> begins as an empty transporter with binding sites for Na<sup>+</sup> and aspartate that are outward facing (Fig. 4A). Once Na<sup>+</sup> and aspartate have bound (step i), the loaded transporter isomerizes from outward facing to inward facing (step ii). The strong electrochemical gradient for Na<sup>+</sup> prompts unbinding of the coupled ions and aspartate (step iii) to leave an empty transporter with binding sites for Na<sup>+</sup> and aspartate facing inward that then relocates...
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FIGURE 4. Lipid bilayer composition influences substrate-loaded isomerization of Glt$_{\text{ph}}$. A, schematic of Glt$_{\text{ph}}$ transport cycle where the empty transporter (Glt$_{\text{ph}}$ (out)) is in the outward facing state and may bind Na$^+$ and aspartate (step i) to form the substrate-loaded complex (Glt$_{\text{ph}}$, 3 Na$^+$, Asp$^-$). This complex can isomerize from the outward facing state to the inward facing state (step ii). The substrate and co-transported ions dissociate (step iii), leaving an empty transporter in the inward facing state (Glt$_{\text{ph}}$, (in)) to relocate to the outward facing state (step iv). Inset, schematic of experimental conditions for counterflow experiments where Na$^+$- and aspartate-loaded liposomes were used to isolate the transport-active limb of the transport cycle (shaded). B, time course of L-[$^3$H]aspartate accumulation in liposomes. Counterflow assays were performed with internal buffer containing 100 mM NaCl, 20 mM HEPES-Tris pH 7.5, 100 $\mu$M unlabeled L-aspartate and an external buffer containing 100 mM NaCl, 20 mM HEPES-Tris, 100 $\mu$M L-[$^3$H]aspartate. Symbols represent PE liposomes (open circles), dimethyl PE liposomes (open triangles), and trimethyl PE liposomes (inverted open triangles). Background is shown as open diamonds. Data represent the mean of experiments performed in triplicate, and error bars indicate S.E.

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Rationale for Site-specific Interactions—The structure of the intermediate outward facing state (Protein Data Bank code 3V8G) of Glt$_{\text{ph}}$ proposes that although the trimerization domain is primarily rigid there are small movements into the surrounding lipid bilayer (9). The ~7-Å flexing movement of TM1 (Fig. 5A) in particular is believed to accommodate the rotation of the transport domain as it completes its isomerization between the outward facing state and inward facing state. We asked whether a residue on TM1 could affect the favorability of this movement into the surrounding lipid bilayer, for example through direct protein-lipid interactions or steric bulk, and could account for the sensitivity of Glt$_{\text{ph}}$ to the lipid environment. In Glt$_{\text{ph}}$, the upper portion of TM1 contains a tyrosine (Tyr-33) residue that may be involved in site-specific interactions (Fig. 5A) consistent with x-ray crystallographic structures of bovine PE-binding protein and murine PE-binding protein bound to zwitterionic lipids (21, 22).

The involvement of Tyr-33 in a lipid-protein interaction was investigated through mutagenesis. The phenol side chain of tyrosine imparts both an aromatic nature and hydrogen-bonding capacity. To dissect these properties, Y33F was generated to remove the hydrogen-bonding capacity of the phenol hydroxyl, and Y33S was generated to maintain hydrogen-bonding capacity of the side chain but remove the aromaticity and side chain bulk. mutant transporters were purified to homogeneity and shown to be functional when reconstituted into control lipid conditions with no significant change in the $K_{0.5}$ for aspartate compared with wild-type Glt$_{\text{ph}}$ (Fig. 5B and Table 3). The mutant transporters were then reconstituted into liposomes composed of PE and trimethyl PE, and rates of transport were examined.

When reconstituted into PE liposomes, Y33F showed initial rates of transport comparable with wild-type Glt$_{\text{ph}}$ (WT), whereas the rate of transport of Y33S was 1.9 ± 0.1-fold higher than that of WT (Fig. 5C). These effects were even more marked when Y33S was reconstituted into trimethyl PE liposomes, which displayed a 3.3 ± 0.1-fold increase in initial rate (Fig. 5D). These results are consistent with a cation–π interaction occurring between Tyr-33 and the zwitterionic lipid headgroups. By maintaining this π interaction in the Y33F mutant, we observed rates comparable with WT. When we removed the potential of cation–π interactions entirely via Y33S, rates increased significantly. Y33W was also generated but was found to function ~0.64-fold less efficiently in PE liposomes and 0.81-fold less efficiently in trimethyl PE liposomes (Fig. 5, C and D). Cysteine residues at the edges of transmembrane domains typically serve to anchor transmembrane helices to the lipid bilayer (23), and the Y33W mutation may reduce the ability of the transporter to undergo important conformational changes within the lipid bilayer. To address the site specificity of this putative
cation–π interaction, we created similar mutations at a nearby tyrosine residue (Tyr-35). Although changes were observed when Y35 was mutated to Phe and Ser, the nature of the residue at position 35 was not affected by lipid composition to the same extent as the residue at position Tyr-33 (Fig. 5, C and D).

DISCUSSION

In this study, we investigated the effect of the lipid environment on the function of the aspartate transporter Glt\textsubscript{ph}, a prokaryotic homologue of the SLC1 family of membrane transporters. We demonstrated a functional effect of the headgroup chemistry of the lipids in which Glt\textsubscript{ph} was reconstituted that was due to a combination of transmembrane orientation and constraining cation–π interactions between the lipid headgroups and a tyrosine residue in TM1.

A common lipid composition used when studying reconstituted membrane proteins is a 3:1 mixture of \textit{E. coli} polar lipid extract to trimethyl PE. In this system, Glt\textsubscript{ph} is orientated ~50% RSO and 50% ISO (13). We have developed a novel thiol modification system using a single cysteine mutant (Glt\textsubscript{ph} A364C) and methanesulfonate reagents that allows us to study the sided kinetics of Glt\textsubscript{ph}. Using this method, we showed that in liposomes in which the predominant lipid is PE the ratio of RSO to ISO changed to 85% RSO and 15% ISO. This suggests that the nature of lipid headgroups is an important factor in determining transporter orientation that presumably is due to specific lipid headgroup-protein interactions. The effect of lipid composition orienting membrane proteins has also been observed with proteorhodopsin in lipid bilayers (24). This method allowed us to determine the $K_{0.5}$ for aspartate and Na\textsuperscript{+} for each face of the transporter. Although we saw no difference in the affinity of Na\textsuperscript{+} for either side of the transporter, we observed that the affinity for aspartate was ~3.5-fold higher for the RSO compared with ISO transporters. This is in contrast to work conducted by Reyes \textit{et al.} (25) who observed similar affinities for aspartate for both sides of the transporter when measuring binding to detergent-solubilized protein. In contrast, our transport assays represent the ensemble of binding, unbinding, and substrate translocation ($K_{0.5}$) of Glt\textsubscript{ph} reconstituted into a lipid bilayer and are in agreement with studies on the human and salamander glutamate transporters also showing a higher affinity for glutamate for the extracellular facing binding site (26–

![Image of Figure 5](image-url)

**FIGURE 5. Functional properties of tyrosine 33 mutants.** A, superimposition of the trimerization domain of the outward facing (green; Protein Data Bank code 2NWX), intermediate outward facing (pink; Protein Data Bank code 3V8G), and inward facing (blue; Protein Data Bank code 3KBC) crystal structures. Structures were aligned to TM4. The transport domain was omitted for clarity. Tyr-33 is shown as spheres. The figure was made using the program PyMOL (38). B, size exclusion column profile for wild-type Glt\textsubscript{ph} (black), Y33F (red), Y33S (orange), Y33W (yellow), Y35F (green), and Y35S (blue). Inset, SDS-PAGE of purified wild-type Glt\textsubscript{ph} (lane 2), Y33F (lane 3), Y33S (lane 4), Y33W (lane 5), Y35F (lane 6), and Y35S (lane 7). Lane 1 contains ladder. C, fold change in L-\textsuperscript{3H}aspartate transport rates for wild-type and mutant Glt\textsubscript{ph} reconstituted in PE liposomes. D, fold change in L-\textsuperscript{3H}aspartate transport rates for wild-type and mutant Glt\textsubscript{ph} reconstituted in trimethyl PE liposomes. Data represent the mean of experiments performed in triplicate, and error bars indicate S.E. with $p < 0.05$ (*), $p < 0.01$ (**), and $p < 0.001$ (***). Compared with WT in each lipid species.

**TABLE 3**

The effect of mutagenesis of Tyr-33 on kinetics of transport by Glt\textsubscript{ph}

Shown are kinetic parameters for mutant Glt\textsubscript{ph} reconstituted in liposomes composed of a 3:1 mixture of \textit{E. coli} polar lipid extract:trimethyl PE. Data were derived by fitting the L-aspartate concentration-response data to the Michaelis-Menten equation. The values are averaged from experiments performed in triplicate, and errors indicate S.E. No significant difference was observed for $K_{0.5}$ values.

|         | WT   | Y33F | Y33S | Y33W | Y35F | Y35S |
|---------|------|------|------|------|------|------|
| Aspartate $K_{0.5}$ (nM) | 84.7 ± 8.6 | 86.3 ± 13.0 | 116.0 ± 19.8 | 61.0 ± 9.0 | 57.5 ± 9.5 | 57.7 ± 10.6 |

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28). The difference in $K_{d,5}$ for the RSO compared with ISO transporters that we observed can partially explain the differences observed in the PE liposomes compared with the trimethyl PE liposomes, but other factors are required to fully explain the observed effects (Fig. 6A).

The outward occluded and inward occluded crystal structures of GltPh suggest that the transport domain undergoes a large movement (~18 Å) relative to the stable trimerization domain during transport (8). However, subtle movements of the trimerization domain have been observed, and it was proposed that these movements are required to accommodate the large movement of the transport domain (9). We propose that the likelihood of residues engaging a cation−π interaction varies with different lipid headgroups, which in turn affect the mobility of TM1 and thus the rate of translocation. When we consider the chemistry of these lipid headgroup moieties, continuing along the series from trimethyl PE to PE, the positive charge of the headgroup becomes more labile and is less able to engage a cation−π interaction. Removing this putative cation−π interaction by altering the lipids into which GltPh is reconstituted allows the requisite movement of TM1 and thus translocation to be more dynamic.

Our results also demonstrate lipid-specific effects that can be augmented by the nature of a residue at the extracellular edge of TM1 that is predicted to face into the lipid membrane. When the Y33S mutant was reconstituted in the strong cation−π partner lipid trimethyl PE, it displayed an approximately 3-fold increase in initial rates compared with WT reconstituted in the same lipid. We suggest that this is because the constraining cation−π interaction is now absent due to the serine mutation. Conversely, the Y33W mutant reduced transport rates compared with wild type, which may be explained by the side chain forming stronger cation−π interactions, anchoring TM1 of GltPh to the membrane, slowing important structural movements required for the isomerization of the substrate-loaded transporter. These results further support our proposal that movements of TM1 into the surrounding lipid bilayer are determined by both the lipid bilayer composition and the chemical nature of residues on TM1 (Fig. 6B). Although our results provide specific examples of how the lipid bilayer can influence transport rates in GltPh, they do not preclude the influence of diffuse, nonspecific features of the bilayer such as intrinsic curvature or steric bulk on transport rates.

Other studies investigating the dynamics of GltPh using a variety of methods have also observed an effect of lipids in determining transport properties. Electron paramagnetic resonance studies by Hänelt et al. (29) have shown a pronounced role of the lipid bilayer to favor intermediate conformations (like that of the intermediate outward facing state), and single molecule fluorescence resonance energy transfer studies suggest that the intermediate conformations of GltPh are rate-determining (30, 31). Finally, a molecular dynamics study using GltPh found that inclusion of constraints imposed by the lipid bilayer was required to permit the conformational changes necessary to allow alternating access of the transporter (32). Our data reveal an effect of the lipid bilayer on isomerization of the Na$^+$ and substrate-loaded transporter (Fig. 4) and further support the growing evidence for a specific role of the lipid bilayer in GltPh function.

The influence of the lipid bilayer composition appears to be a feature shared across a variety of membrane-bound proteins (33–37). Hakizimana et al. (17) and Gustot et al. (18) have observed an influence of lipid bilayer composition on the multidrug transporters LmrP and HorA, respectively. A conserved aspartate residue in LmrP is believed to participate in a hydrogen bond with the lipid headgroups of PE, monomethyl PE, and dimethyl PE. However, transport is abrogated when LmrP is reconstituted in trimethyl PE because this requisite hydrogen bonding is no longer possible (17). In HorA, the tilt angle of the α helices vary by ~10° when PE was compared with trimethyl PE, resulting in an uncoupling of ATP hydrolysis and substrate transport (18).

In addition to influences of transmembrane orientation, we have identified a site-specific, lipid-specific interaction between GltPh and the lipid bilayer that we propose controls the substrate-loaded isomerization step. GltPh cannot progress through the transport cycle without entering intermediate conformations, and lipid bilayer composition, reported through
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Tyr-33, influences the favorability of Glt$_{\text{Ph}}$ to enter these rate-determining intermediate conformations. Our results offer a putative mechanism for Glt$_{\text{Ph}}$ to be coupled to the chemical composition of the membrane.

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Note Added in Proof—The following review of the effects of the lipid environment on membrane protein topology and function has been added to those cited in the version of this article that was published on February 20, 2015 as a Paper in Press: Bogdanov, M., Dowhan, W., and Vitrac, H. (2014) Lipids and topological rules governing membrane protein assembly. Biochim. Biophys. Acta 1843, 1475–1488.

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