Role of Gly117 in the Cation/Melibiose Symport of MelB of Salmonella typhimurium

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Supporting Information

ABSTRACT: The melibiose permease of Salmonella typhimurium (MelB<sub>St</sub>) catalyzes symport of melibiose with Na<sup>+</sup>, Li<sup>+</sup>, or H<sup>+</sup>, and bioinformatics analysis indicates that a conserved Gly117 (helix IV) is part of the Na<sup>+</sup>-binding site. We mutated Gly117 to Ala, Pro, Trp, or Arg; the effects on melibiose transport and binding of cosubstrates depended on the physical—chemical properties of the side chain. Compared with WT MelB<sub>St</sub>, the Gly117→Ala mutant exhibited little difference in either cosubstrate binding or stimulation of melibiose transport by Na<sup>+</sup> or Li<sup>+</sup>, but all other mutations reduced melibiose active transport and efflux, and decreased the apparent affinity for Na<sup>+</sup>. The bulky Trp at position 117 caused the greatest inhibition of melibiose binding, and Gly117→Arg yielded less than a 4-fold decrease in the apparent affinity for melibiose at saturating Na<sup>+</sup> or Li<sup>+</sup> concentration. Remarkably, the mutant Gly117→Arg catalyzed melibiose exchange in the presence of Na<sup>+</sup> or Li<sup>+</sup>, but did not catalyze melibiose translocation involving net flux of the coupling cation, indicating that sugar is released prior to release of the coupling cation. Taken together, the findings are consistent with the notion that Gly117 plays an important role in cation binding and translocation.

The melibiose permease of Salmonella typhimurium (MelB<sub>St</sub>) catalyzes cotransport of galactoside with Na<sup>+</sup>, Li<sup>+</sup>, or H<sup>+</sup>. MelB<sub>St</sub> belongs to the glycoside/pentoside/hexuronide:cation family, a subgroup of the major facilitator superfamily (MFS) of membrane transport proteins. MelB of Escherichia coli (MelB<sub>Ec</sub>) is the best-studied member among all MelB orthologues. Recently, MelB homologues in human and mouse (called the major facilitator superfamily domain-containing proteins, MFS<sub>D</sub>) have been reported. Among them, MFS<sub>D2A</sub> protein, which is expressed in many cells, plays a role in adaptive thermogenesis; it has been also identified as them, MFSD2A protein, which is expressed in many cells, plays a role in adaptive thermogenesis; it has been also identified as the crystal structures of an MFS permease, the H+-coupled H+-binding site in LacY is located in the C-terminal helices.28 Located in the N-terminal helix bundle (Figure 1), whereas the H<sup>+</sup>-binding site in LacY is located in the C-terminal helices.28 On the basis of the location of the sugar-binding site in LacY, the melibiose-binding site is proposed to lie within the internal cavity (Figure 1). This model is consistent with numerous biochemical and biophysical results, as well as low-resolution EM structures of MelB<sub>Ec</sub>. The organization of the protein into two separate helix bundles, as well as the location of cosubstrates, are consistent with the alternating-access transport model, which has been recognized as a fundamental mechanism for many other secondary transporters.

MelB<sub>St</sub> shares 86% identity and 96% similarity of primary sequence with MelB<sub>Ec</sub> (Supporting Information, Figure S1). All nonconserved and 82% of the conserved variations occur in the C-terminal domain and the middle loop. Bioinformatics analysis suggests that the internal cavities including two cosubstrate-binding sites are well conserved between the two MelB permeases (Figure 1), implying similarity in function, which is supported by previous studies of melibiose transport and cation binding. The docked sugar is surrounded by potential H-bonding partners and aromatic residues (Figure 1), which share common features for sugar binding. A Na<sup>+</sup> has been proposed to bind between helices II and IV. A large body of experimental data, including those from mutagenesis, biochemistry, and FTIR spectroscopy, indicates that the carboxyl groups of conserved Asp55 and Asp59 (helix II) contribute to Na<sup>+</sup> binding to MelB<sub>Ec</sub>. Helix IV is in the center of a charge/H-bond network involved in the binding of the two cosubstrates (Figure 1).

Gly117 is in the
**Biochemistry**

**Materials.** 1-[3H]Melibiose was custom synthesized by PerkinElmer (Boston, MT). The 2-(N-dansyl)aminoalkyl-1-thio-β-D-galactopyranoside (D25) was kindly provided by Drs. H. Ronald Kaback and Gérard Leblanc. Monensin and carbonyl cyanide m-chlorophenylhydrazone (CCCP) were purchased from Sigma-Aldrich. 2-(4′-Maleimidylanilino)-napthalene-6-sulfonic acid was purchased from Invitrogen. Oligodeoxynucleotides were synthesized by Integrated DNA Technologies. MacConkey agar media (lactose free) was from Difco. All other materials were reagent grade and obtained from commercial sources.

**Bacterial Strains and Plasmids.** *Escherichia coli* DW2 strain (melA, ΔmelB, ΔlacZ) obtained from Dr. Gérard Leblanc, was used for the functional characterization. *E. coli* XL1 Blue cells were used for DNA manipulations. The expression plasmid pK95 ΔAH/MelBΔ/His10,1,3, which encodes the full-length MelBΔ with L5→M and a His10-tag at the C-terminus (the wild type) was used as the template. Four individual mutants with residue Ala, Pro, Trp, or Arg at position Gly117 were constructed by a QuickChange Site-Directed Mutagenesis kit from Stratagene. All mutants have been confirmed by DNA sequencing.

**Growth of Cells and Protein Overexpression.** *E. coli* DW2 cells (melA, ΔmelB, ΔlacZ) containing a given plasmid were grown in Luria–Bertani (LB) broth with 100 mg/L ofampicillin in a 37 °C shaker overnight. The overnight cultures were diluted by 5% to LB broth supplemented with 0.5% glycerol and 100 mg/L of ampicillin, and constitutive overexpression was obtained by shaking at 30 °C for 5 h.

**Preparation of Right-Side-Out (RSO) Membrane Vesicles.** RSO membrane vesicles were prepared from *E. coli* DW2 cells by osmotic lysis,1,3,54 extensively washed, and resuspended in 100 mM KPi, pH 7.5 at a protein concentration of ∼25–30 mg/mL, frozen in liquid N2, and stored at −80 °C.

**Preparation of Crude Membranes.** The 5-h cultures with the expressed MelBΔ were washed with 20 mM Tris-HCl (pH 7.5) once, resuspended, and adjusted with the same buffer to A600 ~20. Approximately 100 μL of cells from each sample were sonicated in an ice-cold water bath (Branson 2510) for 5 min, three times, and centrifuged at 20816g for 15 min at 4 °C. The supernatant (80 μL) was subjected to ultracentrifugation at 384492g (TLA-100 rotor, Beckman Optima Max Ultra-centrifuge) for 20 min; the pellets were resuspended in 70 μL of 20 mM Tris-HCl (pH 7.5).

**SDS–12% PAGE and Western Blotting.** After protein assay using a Micro BCA Protein Assay kit (Pierce), ∼15 µg of total membranes were loaded onto each well of SDS–12% PAGE. The gel was transferred onto the PVDF membrane by the Trans-Blot Turbo transfer system (Bio-Rad) at 1.3 A, 25 V for 15 min. The blocked PVDF membrane was then reacted with the penta-His horseradish peroxidase conjugate and washed according to the protocols provided in the Penta-His HRP Conjugate kit (Qiagen). MelBΔ proteins were detected using the SuperSignal West Pico chemiluminescent substrate (Thermo Scientific) after exposing it to the X-ray film (Kodak BioMax XAR film).

**1-[3H]Melibiose Transport Assay.** *E. coli* DW2 cells expressing MelBΔ in the absence of melibiose were washed with 50 mL of 100 mM KPi, (pH 7.5, so-called Na+–free buffer) twice, followed by washing with 50 mL of 100 mM KPi, pH 7.5, 10 mM MgSO4. The cell pellets were resuspended with the latter buffer, adjusted to an A420 = 10 (~0.7 mg of protein/mL). Melibiose transport at a final concentration of 0.4 mM and a specific activity of 10 mCi/mm mol was assayed as described.1,3

**Melibiose Fermentation.** The DW2 cells (melA, ΔmelB, ΔlacZ) were transformed with a given plasmid, plated on MacConkey agar plates containing ~100 mM Na+ with supplementation of 1–30 mM melibiose (the sole carbohydrate source) and 100 mg/L of ampicillin, and incubated at 37 °C. After 18 h, the plates were viewed and photographed immediately.

**Melibiose Efflux and Exchange.** RSO membrane vesicles containing MelBΔ in 100 mM KPi, pH 7.5, 10 mM MgSO4, and...
were concentrated to 25–30 mg/mL, and pre-equilibrated overnight on ice with melibiose (20 mM, 10 mCi/mmol), 0.75 μM monensin, and 10 μM CCCP, in the absence or presence of 20 mM NaCl or LiCl. Aliquots (2 μL) were diluted 200-fold into a given buffer in the absence (efflux) or presence (exchange) of 20 mM unlabeled melibiose, and reactions were terminated by dilution and rapid filtration at a given time.

**FRET (Trp → D2G).** Fluorescence was measured with an AMINCO-Bowman Series 2 spectrometer. Steady-state measurements were performed in a 3-mm quartz cuvette (Starna Cells, Inc.) with RSO membrane vesicles at a protein concentration of ~0.5 mg/mL in 100 mM KPi, pH 7.5. With excitation wavelength at 290 nm, the FRET intensity was recorded for 60 s at 500 nm with slit widths of 2 mm for both excitation and emission.

**Determination of Apparent Cation-Stimulation Constants (K_0.5^{Na^+} and K_0.5^{Li^+}) of the D2G FRET.** NaCl or LiCl was consecutively added into a 3-mm quartz cuvette containing RSO vesicles after an addition of 10 μM D2G (a K_0 value for the WT). Water at an identical volume was used for the control. The FRET signals were recorded for 60 s at each condition and mean values were used for further calculation. The increase in intensity (ΔF_{water}/I_0), the difference before (I_0) and after successive additions of cation solution (ΔF_{Na^+/Li^+}/I_0), was expressed as the percentage of the I_0 (ΔF_{Na^+/Li^+}/I_0). FRET stimulation at each cation concentration was corrected by subtracting corresponding ΔF_{water}/I_0 value, and then plotted as a function of Na^+ or Li^+ concentration. The K_0.5^{Na^+} or Li^+ values are determined by fitting the data to a hyperbolic function.

**Determination of the Melibiose Concentration Corresponding to Half-Maximal Displacement of Bound D2G (IC50).** Melibiose was added stepwise to a 3-mm quartz cuvette containing RSO vesicles after the additions of D2G (10 μM) and NaCl or LiCl (20 mM or 200 mM) until no change occurred in fluorescence emission at 500 nm. An identical volume of water was added at each point as a negative control. FRET signals were recorded for 60 s after each addition, and mean values were calculated. The decrease in intensity after each addition of melibiose (ΔF_{Mel}) was corrected by subtracting the emission change obtained with water (ΔF_{water}) and plotted as a function of melibiose concentration. The IC50 values were determined by fitting the data to a hyperbolic function.

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**RESULTS**

**Construction and Protein Expression of Gly117 Mutants.** Gly at position 117 in MelB_{St} was changed by site-directed mutagenesis to four different residues: Ala (small volume); Pro (a helix destabilizer); Trp (aromatic ring); or Arg (positive charge). Membrane proteins were separated by SDS-PAGE, and the proteins were detected by both silver nitrate and Western blot using anti-His antibody (Supporting Information, Figure S2). Compared to the WT, most of the mutants had similar levels of membrane expression, with the exception of G117R, which had a reduced expression.

**Melibiose Active Transport in Intact Cells.** In a nominally Na^+-free buffer, the WT catalyzes melibiose transport at an almost linear rate for 15 s to 80 nmol/mg protein at 2 min (Figure 2), which was significantly stimulated by Na^+ or Li^+ as shown previously. Addition of the protonophore CCCP abolished transport, indicating that the primary driving force for transport is the electrochemical H^+ gradient. Furthermore, when unlabeled melibiose was added at each time point during transport, with a 10-min incubation prior to the filtration, the entire internal pool of melibiose was exchanged with extracellular melibiose (Supporting Information, Figure S3). Thus, there is little or no hydrolysis or chemical alteration of the accumulated intracellular melibiose.

The mutant G117A catalyzed Na^+- or Li^+-dependent melibiose transport similar to the WT (Figure 2) with somewhat lower H^+-coupled transport activity (i.e., in the absence of Na^+ or Li^+). All other mutants (P, W, or R) exhibited H^+-coupled melibiose transport at a level indistinguishable from CCCP-treated WT or nontransformed DW2 cells, and showed no response to addition of Na^+ or Li^+.

**Melibiose Fermentation.** On MacConkey agar plates containing 1–30 mM melibiose, untransformed DW-2 cells formed pale/white colonies with a translucent background, denoting little or no melibiose translocation across the membranes. At >10 mM melibiose, DW2 cells, containing the chromosome-encoded α-galactosidase and recombinantly overexpressing WT MelB_{St} formed magenta colonies on a hazy background indicating melibiose fermentation and therefore downhill melibiose influx catalyzed by MelB_{St} (Figure 3). At 5 mM melibiose, few colonies were pink, whereas at 2.5 mM or lower, the colonies were pale yellow/brown. Mutant G117A exhibited color similar to WT. Mutant G117P exhibited red or pink colonies only at high concentrations of melibiose, indicating limited melibiose influx and fermentation. Mutants G117R and G117W formed colonies that were indistinguishable from untransformed cells even at 30 mM melibiose, indicating no fermentation.

**Melibiose Efflux and Exchange.** In the absence of Na^+ or Li^+, RSO membrane vesicles containing WT MelB_{St} exhibited a slow rate of melibiose efflux down a concentration gradient and a slightly faster rate of equilibrium exchange (Figure 4). Whereas the efflux rate was stimulated by Na^+ or Li^+, the exchange rate was enhanced by Na^+ but inhibited by Li^+. In the presence of Na^+, 96% of the intravesicular melibiose exchanged with the extracellular unlabeled melibiose within 5 min (data not shown). Consistent with the results in intact cells, little or no melibiose hydrolysis occurred within the RSO vesicles.
The G117W mutant did not catalyze either melibiose efflux or exchange in the presence of H+, Na+, or Li+. The G117P mutant catalyzed significantly reduced rates of efflux and exchange. Strikingly, mutant G117R exhibited little or no efflux but catalyzed a relatively fast rate of exchange in the presence of Na+ or Li+, but not H+.

$K_{0.5}^{Na^+}$ or $K_{0.5}^{Li^+}$ for D2G FRET. FRET from endogenous Trp residues to the dansyl moiety of bound D2G has been demonstrated with MelB$_{Ec}$ and MelB$_{St}$. Thus, WT MelB$_{St}$ exhibits FRET upon the addition of D2G, which is stimulated by addition of Na+ or Li+ and reversed by melibiose. The difference in intensity observed before and after melibiose addition reflects the D2G bound specifically to MelB$_{St}$ at saturating Na+ or Li+ concentrations. The Na+ or Li+ enhancement of FRET (difference in intensity before and after cation addition, $\Delta I_{Na^+ or Li^+}$) results from combined effects of an increase of D2G binding per se and a conformational change (distance and/or environment) induced by Na+ or Li+ binding. Mutations may affect Na+- or Li+-induced intensities by (a) decreasing the binding affinity for either or both cosubstrates, and/or (b) modifying the microenvironment surrounding donor Trp residue(s), and/or (c) restricting the conformational change. Thus, the same concentrations of Na+- or Li+-bound ternary complex from different mutants may not yield the same $\Delta I/I_0$ value.

In order to quantify cation binding to MelB$_{St}$ mutants, $\Delta I_{Na^+ or Li^+}/I_0$ was measured as a function of Na+ or Li+ concentration (Figure 4, upper panels). The stimulation constants ($K_{0.5}^{Na^+}$ and $K_{0.5}^{Li^+}$) for D2G FRET in WT MelB$_{St}$ were 0.9 mM and 1.5 mM, respectively, and the mutant G117A yielded similar values (Table 1). $K_{0.5}^{Na^+}$ and $K_{0.5}^{Li^+}$ values for G117P were 2- and 6-fold higher than those for the WT, respectively. With mutant G117R, both $K_{0.5}^{Na^+}$ values increased by 12-fold. The $K_{0.5}^{Li^+}$ for G117W was only 2.5-fold higher than obtained for WT, but no Na+ stimulation was observed even at 500 mM (data not shown). In all three mutants, the maximum stimulations by Na+ or Li+ were largely reduced implying that either the microenvironment and/or distance between D2G and donors are changed, or that the number of bound D2G molecules is reduced by lower sugar affinity. In either case, the constant ($K_{0.5}^{Na^+ or Li^+}$) reflects the cation-binding affinity.

IC$_{50}$ for Melibiose Displacement of Bound D2G. In order to quantify melibiose binding to MelB$_{St}$, 10 mM D2G, Na+ or Li+, and then the titrate melibiose were consecutively added to RSO vesicles. IC$_{50}$ values for WT MelB$_{St}$ were 3.8 mM or 1.7 mM at 20 mM Na+ or Li+, respectively, and 2.2 mM or 1.8 mM at 200 mM Na+ or Li+, respectively (Figure 4, bottom panels; Table 2). The IC$_{50}$ for mutant G117A was about 2-fold higher than that of the WT at 20 mM Na+ or Li+. In the presence of 20 mM Na+ or 200 mM Li+, IC$_{50}$ values for mutant G117W were 50 mM or 14.1 mM, respectively, and for mutant G117P were 11.3 mM or 7.5 mM, respectively. Mutant G117R exhibited IC$_{50}$ of 5.6 mM or 4.7 mM in the presence of 200 mM Na+ or Li+. FRET intensities for mutants G117W or P, with Na+ as the cosubstrate, were too weak to allow the determination of IC$_{50}$ value.
**DISCUSSION**

An ordered-binding model for cation/melibiose symport in MelBEc has been suggested. An extended model for the melibiose efflux mechanism is proposed for MelBSt here (Figure 6). In this simplified scheme, melibiose efflux down a sugar concentration gradient consists of eight steps (black solid arrows): 

- **Step-1**, Na⁺-bound inward-facing state, or PI:Na⁺ state
- **Step-2**, Na⁺- and melibiose-bound inward-facing state, or PI:Na⁺:Mel state
- **Step-3**, occluded Na⁺- and melibiose-bound state, or P:(Na⁺:Mel) state
- **Step-4**, Na⁺- and melibiose-bound outward-facing state, or PO:Na⁺:Mel state
- **Step-5**, Na⁺- and melibiose-bound outward-facing state, or PO:Na⁺ state
- **Step-6**, empty outward-facing state, or PO:empty state
- **Step-7**, occluded empty state, or P:(empty) state
- **Step-8**, empty inward-facing state, or PI:empty state

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**Table 1. Cation Stimulation Constants for D²G FRET (K_{0.5} Na⁺ or Li⁺, mM)**

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
<th>G117A</th>
<th>G117W</th>
<th>G117P</th>
<th>G117R</th>
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</thead>
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<tr>
<td>Na⁺</td>
<td>0.9</td>
<td>1.7</td>
<td>a</td>
<td>23.2</td>
<td>11.2</td>
</tr>
<tr>
<td>Li⁺</td>
<td>1.5</td>
<td>0.97</td>
<td>3.6</td>
<td>9.5</td>
<td>17.2</td>
</tr>
</tbody>
</table>

“Not determined due to weak signals.”

**Table 2. IC_{50} for Melibiose Displacement of Bound D²G (mM)**

<table>
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<tr>
<th></th>
<th>WT</th>
<th>G117A</th>
<th>G117W</th>
<th>G117P</th>
<th>G117R</th>
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<tr>
<td>Na⁺</td>
<td>20 mM</td>
<td>3.8</td>
<td>8.1</td>
<td>a</td>
<td>a</td>
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<tr>
<td></td>
<td>200 mM</td>
<td>2.2</td>
<td>b</td>
<td>a</td>
<td>a</td>
</tr>
<tr>
<td>Li⁺</td>
<td>20 mM</td>
<td>1.7</td>
<td>3.4</td>
<td>50.0</td>
<td>11.3</td>
</tr>
<tr>
<td></td>
<td>200 mM</td>
<td>1.8</td>
<td>b</td>
<td>14.1</td>
<td>7.5</td>
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</tbody>
</table>

“Not determined due to weak signals.”

“Not determined.”

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**Figure 5.** Apparent affinity. RSO vesicles (0.5 mg/mL, 100 mM KPi, pH 7.5) containing WT or a MelBSt mutant were used for D²G FRET assay at excitation and emission wavelengths of 290 and 500 nm, respectively. (Top panels): Determination of K_{0.5} Na⁺ and K_{0.5} Li⁺ for the D²G FRET. With 10 μM D²G, Na⁺ or Li⁺ was added at increasing concentrations. The normalized emissions were plotted as a function of Na⁺ (circles) or Li⁺ (triangles) concentration. K_{0.5} Na⁺ or K_{0.5} Li⁺ value was determined by fitting a hyperbolic function to the data. (Bottom panels): Determination of IC_{50} for melibiose displacement of bound D²G. With D²G (10 μM) and Na⁺ or Li⁺ at a given concentration, melibiose was added stepwise until the displacement of the bound D²G was complete. After correction for the water dilution effect, the change in intensity (ΔF) was plotted as a function of melibiose concentration. The IC_{50} value was determined by fitting a hyperbolic function to the data obtained in the presence of 20 mM Na⁺ (circles with dashed lines), 200 mM Na⁺ (circles with solid lines), 20 mM Li⁺ (triangles with dashed lines), or 200 mM Li⁺ (triangles with solid lines).

**Figure 6.** An ordered-binding model for melibiose/Na⁺ efflux catalyzed by MelBSt (see text). The N- and C-terminal domains of MelBSt molecule are colored in green and blue, respectively. Sugar and Na⁺ are shown as black hexagons and red spheres, respectively. PI or PO represents the permease at an inward-facing or outward-facing conformation, respectively. P:(Na⁺:Mel) or P:(empty), cosubstrates-bound or unloaded permease at the occluded state, respectively.
Opening of the cytoplasmic cavity. The melibiose efflux involves the whole cycle, and melibiose exchange reaction involves only steps 1–5.

Previously, we have experimentally determined that the extracellular release of cation is the rate-limiting step for melibiose efflux catalyzed by MelBSt and melibiose is released from MelBS extracellularly prior to the release of Na+. These results are consistent with that observed in MelBEc.22 Here, we provide further evidence to support this notion. Cells expressing mutant G117R neither ferment melibiose nor catalyze melibiose active transport or efflux. Remarkably, however, this mutant catalyzes Na+- or Li+- but not H+-coupled equilibrium exchange. Thus, in the presence of 20 mM melibiose and Na+ or Li+ on both sides of the membrane, the mutant can undergo the global conformational change involved in an alternating access-type mechanism but is defective in releasing Na+ or Li+ (Figure 6, step 6), implying that Gly117 plays an important role in the cation translocation mechanism. This behavior of mutant G117R, similar to that of E325A LacY,58,59 strongly supports the ordered-binding mechanism as postulated (Figure 6).

The rate of exchange is believed to correlate with the binding affinity. Accordingly, at saturating Na+ or Li+ concentration, the IC50 for melibiose displacement of bound D2G exhibits only a 2.5-fold increase (Table 2). If we assume that the mutation has a similar effect on D2G and melibiose, then it can be estimated that the apparent binding constant (Kd) for melibiose is increased by less than 4-fold, implying that sugar binding is only slightly affected by the Arg at position 117, and therefore Gly117 may not coordinate melibiose directly.

In WT MelBSt, we have been observed previously1 that Li+ stimulates melibiose efflux and an outwardly directed Li+ gradient increases exchange; however, at equal Li+ concentrations on both sides of the membrane, the melibiose exchange rate is slower than the efflux rate. In MelBSt, Li+ also inhibits melibiose exchange, but not methyl-β-D-thiogalactoside exchange. Although the mechanism of inhibition is still not clear, it is possible that MelB/Li+/melibiose complexes favor an occluded state (Figure 6, step 3) in which there is no exchange between free and bound sugars.7 In the presence of Li+, the Gly117 → R mutation inhibits melibiose efflux but stimulates exchange at a rate even faster than that obtained with WT MelBSt. It is likely that the mutation may decrease the stability of the MelB/Li+/melibiose complexes at step 3, thus facilitating melibiose exchange at a rate similar to that obtained in the presence of Na+.

Six of eight endogenous Trp residues are located in the N-terminal helix bundle. Trp64 (helix II) and Trp299 (IX) of MelBSt are responsible for 80% of the D2G FRET, and apparently Trp116 does not contribute significantly to the FRET.25 Thus, it seems unlikely that the inserted Trp at position 117 could serve as an efficient FRET donor or significantly quench the Trp64 and Trp299 emission directly. Mutant G117W specifically blocked Na+-induced D2G FRET (ΔI), but the apparent affinity for Li+ did not change significantly (<3-fold increase in KdLi+, Table 1). Thus, the lack of Na+-induced D2G FRET (ΔI) may be mainly due to a large decrease in Na+ binding.

While in most Gly117 mutants there are reductions in binding of Na+, Li+, and melibiose, the effect on Na+ binding is more severe than those on Li+ and melibiose. The MelB orthologue of Klebsiella pneumoniae (MelBkp) couples melibiose transport with H+ and Li+ but not Na+.61 It has been demonstrated that Asn58, which is adjacent to Asp59, is important for Na+ recognition, and MelBSkp has Ala at position S8.31 It seems likely that selective elimination of Na+ binding in MelB could be achieved by a minor modification around the cation-binding site. In addition, the effective inhibition of melibiose binding in mutant G117W (Table 2) may be due to a steric effect of the bulky side chain, which may also partially account for the effects of other mutations on sugar binding.

Gly is known to destabilize α-helices, and the replacement with Pro, another α-helix destabilizer that may alter the tilt in helix IV, partially inhibits both efflux and exchange reactions, as well as impairs affinity for the cosubstrates to a similar extent. Thus, the data support the notion that the melibiose/cation symport mechanism requires precise positioning of helix IV.

If the carbonyl oxygen at position 117 is a part of the cation-binding site, then a small neutral replacement should not have a significant effect. Accordingly, placement of Ala at position 117 has little or no effect on Na+- or Li+-coupled melibiose transport (Figure 2) or the apparent affinity for Na+, Li+, or melibiose (Tables 1 and 2). Taken as a whole, our findings show that the placement of Arg or Trp at position 117 abolishes melibiose uphill transport, efflux, and fermentation; G117R and P greatly decrease apparent affinities for Na+ and Li+. Remarkably, the mutant G117R retains significant sugar affinity and catalyzes melibiose exchange, a partial reaction, with bound Na+ or Li+, but does not catalyze melibiose translocation involving net flux of the coupling cation. In conclusion, Gly117 plays an important role in cation binding and translocation of MelBSt, which is consistent with our threading structure.13

ASSOCIATED CONTENT

Supporting Information

Conservation between MelBSt and MelBEc, SDS–PAGE analysis and Western blotting, as well as melibiose exchange with intact cells are available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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MelB, melibiose permease of Salmonella typhimurium; MelB\textsubscript{Es}.
MelB of Escherichia coli; LCy, lactose permease of E. coli; melA, gene encoding α-galactosidase; LB broth, Luria–Bertani broth; D\textsuperscript{2}G, 2′-(N-dansyl)aminoalkyl-1-thio-β-D-galactopyranoside; CCCP, carbonylcyanide-m-chlorophenylhydrazone; \(K_{\text{m}}\)\textsuperscript{Na+ or Li+}, the Na+ or Li+ stimulation constant for D\textsuperscript{2}G FRET; IC\textsubscript{50}, the response to tunicamycin. Proc. Natl. Acad. Sci. U. S. A. 108, 11756–11765.


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The crystal structure of a sodium galactose transporter reveals mechanistic insights.


