

UC San Diego

UC San Diego Previously Published Works

Title

Conserved movement of TMS11 between occluded conformations of LacY and XylE of the major facilitator superfamily suggests a similar hinge-like mechanism

Permalink

<https://escholarship.org/uc/item/4zn6f5jf>

Journal

Proteins Structure Function and Bioinformatics, 83(4)

ISSN

0887-3585

Authors

Västermark, Åke
Driker, Adelle
Li, Jiaqi
et al.

Publication Date

2015-04-01

DOI

10.1002/prot.24755

Peer reviewed

Conserved movement of TMS 11 between occluded conformations of LacY and XylE of the major facilitator superfamily suggests a similar hinge-like mechanism

Åke Västermark, Adelle Driker, Jiaqi Li, and Milton H. Saier Jr*

Department of Molecular Biology, University of California at San Diego, La Jolla, California 92093-0116

ABSTRACT

The Δ -distance maps can detect local remodeling that is difficult to accurately determine using superimpositions. Transmembrane segments (TMSs) 11 in both LacY and XylE of the major facilitator superfamily uniquely contribute the greatest amount of mobile surface area in the outward-occluded state and undergo analogous movements. The intracellular part of TMS11 moves away from the C-terminal domain and into the substrate cavity during the conformational change from the outward-occluded to the inward-occluded state. A difference was noted between LacY and XylE when they assumed the inward open state after releasing a substrate to the inside in which TMS11 of LacY moved further into the substrate release space, whereas in XylE, TMS11 slightly retracted into the C-terminal domain. Independent movement of the N-terminal half of TMS11 suggests that it is flexible in the middle. Repeat-swapped homology modeling was used to discover that a loop connecting TMSs 10 and 11 in LacY probably moves during the transition between the unavailable outward-open state and the outward-occluded state. TMSs 11 and the other elements displaying a notable domain-independent movement colocalize with the interdomain linker, suggesting that these elements could drive the alternating access movement between the domain halves. Preliminary evidence indicates that analogous movements occur in other members of the major facilitator superfamily.

Proteins 2015; 83:735–745.
© 2015 Wiley Periodicals, Inc.

Key words: TMS11; LacY; XylE; MFS; contact maps; occluded conformational states.

INTRODUCTION

The major facilitator superfamily (MFS) is the largest family of secondary carriers, containing 83 families within TC# 2.A.1 of the Transporter Classification Database, TCDB (www.tcdb.org)¹ and an additional seven families within the extended superfamily. Most MFS proteins have a 6 + 6 α -helical transmembrane segment (TMS) topology (but some have 6 + 2 + 6) where the two halves are connected by an interdomain linker that allows movement of the halves relative to each other during the alternating access transport cycle. The term “rocker switch” is specifically used to define a kinetic behavior of secondary carriers whose transport rate is temperature dependent, but the affinity is temperature independent,² and the term can also be used to describe the alternating access mechanism.³

The transport cycle involves at least four major states: outward open, outward and inward occluded, and inward open. The interdomain linker of LacY contains

three cytoplasmic helical segments that colocalize in space when the outward-open conformation is stabilized. Only the outward-open states of LacY and XylE have not been determined by X-ray crystallography. However, this state has been determined for an MFS homolog, the fucose porter, FucP.⁴ We attempt here to examine a model relying on the available X-ray data, a repeat-swapped homology model.⁵

Additional Supporting Information may be found in the online version of this article.

Grant sponsor: NIH; Grant numbers: GM077402 and GM094610 (to M.H.S.).

A.V. and M.H.S. conceived and designed the experiments; A.V., A.D., and J.L. performed the experiments and analyzed the data; A.V. and M.H.S. contributed reagents/materials/analysis tools; and A.V., J.L., and M.H.S. contributed to the writing of the manuscript.

The authors declare no competing financial interests.

*Correspondence to: Milton Saier, Department of Molecular Biology, University of California at San Diego, La Jolla, CA 92093-0116. E-mail: msaier@ucsd.edu
Received 6 November 2014; Revised 17 December 2014; Accepted 31 December 2014

Published online 13 January 2015 in Wiley Online Library (wileyonlinelibrary.com). DOI: 10.1002/prot.24755

Many MFS proteins use a proton gradient, typically created by primary active transporters, to allow intracellular accumulation of solutes such as lactose and xylose. The *Escherichia coli* lactose transporter, LacY, is one of the most intensely studied secondary carriers, having had the effects of point mutations on transport and counterflow investigated for each of its residues. The xylose transporter, XylE, provides a model system of an MFS belonging to the sugar porter family, which includes the human glucose transporters, the GLUTs, although some GLUTs have been structurally determined⁶, XylE was the first MFS transporter to have crystal structures determined for both inward- and outward-facing occluded states.^{7,8}

Δ -distance or difference distance mapping offers advantages over superimposition because it allows detection of local differences that superimposition of parts of homologous structures cannot reveal. We have previously published a Δ -distance map comparing the occluded states of XylE, showing that the majority of the interdomain distance remodeling is localized to the C-terminal bundle.⁹ Publication of the crystal structure of a sugar-bound state of LacY confirmed these observations.¹⁰ Note, however, that this finding does not contradict unfolding experiments,¹¹ because they measure other physical properties, not distance remodeling.

In this article we use an established technique, Δ -distance mapping, to compare structures in different conformational states. The concept of calculating Δ -distance maps is not new,^{12–14} but the recent availability of multiple conformational states for MFS homologs, LacY and XylE, has created a new application area. In fact, the most recent version of Chimera, release 1.10,¹⁴ contains a new function called RR distance maps. While this function can be used to compare two contact maps, it does not visualize whether the distances increase or decrease; it only shows distance remodeling as a color scale of standard deviations. Δ -distance mapping provides higher precision in localizing conformational changes compared to the superimposition-based methods. This approach reveals the main differences between LacY and XylE as they transition between their four main conformational states.

Since crystal structures of LacY and XylE are not yet available in the outward-open state, we used a repeat-swapped homology model to estimate the missing patterns and differences in internal distance remodeling patterns exhibited in these two transporters were identified. To complement the results from the Δ -distance mapping, a detailed description of the different conformational states and the changes occurring between them is provided. In particular, the movements of full-length and segmental TMSs displaced by the alternating access movement between domain halves were examined, and

correlated with the placement of these elements in relation to the interdomain linker.

While this study is primarily limited to making comparisons between LacY and XylE, the only two proteins to have three determined conformational states, we attempt to generalize to other MFS proteins by creating homology models. This approach has identified conserved movements, also involving TMS11 for the fucose porter, FucP,⁴ and a member of the MFS peptide transporting POT/PTR family, NRT1.¹⁵ This study is the first of its kind, because it uses three recent developments: the repeat-swapped model of the outward-open state of LacY,⁵ the recently published crystal structure of the sugar-bound state of LacY,¹⁰ and the use of Δ -distance mapping to compare conformational states.⁹

METHODS

Δ -distance maps

We downloaded the LacY PDB files for the inward-open conformation, with an affinity inactivator (2Y5Y), and in the sugar-bound, outward-occluded state (4OAA). The range of residues was restricted to the greatest common range, from 4 (LEU) to 403 (PRO). We selected only “CA” (backbone α -carbon) atoms from the “A” chains. There was a two-line discrepancy in line count between the files, indicating missing atoms in 4OAA. Residues 191–206 were missing in 4OAA, and Residues 189–202 were missing in 2Y5Y. We deleted segments so that corresponding segments encompassing Residues 189–206 were missing in both files. As a result, both files had 382 lines, running from Residues 4 to 403 but missing 18 residues (189–206).

The CA atoms of XylE chain A of 4JA3 (8–465) and of 4GBY (5–479, no atoms missing between 8 and 465) were selected, yielding 426 and 475 lines, respectively. 4JA3 was missing 265–275, 304–309, 398–406, and 434–439 (inclusive). In total, 32 residues were missing in 4JA3 (8–465). To compensate for the missing atoms in 4JA3, we removed all atoms from 4GBY that were not present in 4JA3.

The data were analyzed using a Perl script that calculated all pairwise Euclidian distances in Angstroms, rounding the numbers to two decimal places. The resulting matrices were loaded as a tab-delimited spreadsheet in Microsoft Excel. The distance map of LacY (2Y5Y) was subtracted from 4OAA, generating a Δ -distance map for the comparison. Conditional formatting was applied, using the blue–white–red color scale. The quadrants of the plot representing the alternating access movements between the N- and C-terminal halves were not considered when applying conditional formatting, making the detection of small remodeling of helix distances within each half possible.

Repeat-swapped homology model of the outward-open states

A theoretical model of the outward-open conformation of LacY has previously been created using the repeat-swapping methodology.⁵ The model of the C-terminal half of the sequence was based on the N-terminal half of the inward-open structure and *vice versa*. In Modeller 9.7, LacY in the inward-facing state was used as the template. A sequence alignment from AlignMe¹⁶ was used to guide the homology modeling. The model was energy minimized using Charmm 3.4. The repeat-swapped model of the outward-open state of LacY is available in the Protein DataBank (PDB): PM0076824 (repeat-swapped model). Modeller was used to emulate the above-mentioned experimental procedure and build a theoretical model of the outward-open state of Xyle that was limited to the C-terminal 6 TMS bundle.

Use of RR distance maps to generalize to other members of the MFS

We used AlignMe PST¹⁶ to align TMSs 1–6 and TMSs 7–12 of FucP. The resulting alignment was used to guide Modeller to create a homology model of TMSs 7–12 (in an inward open-like state) using TMSs 1–6 in the outward-open state (3O7P) as the template. Thus, TMSs 7–12 in the outward-open state could be superimposed on TMSs 7–12 in the inward open-like state (the model). RR distance maps were used to detect differences between the two structures and identified a cross-shaped region located near TMSs 11, which coincides with elevated standard deviation measurements.

With the outward-occluded state of Xyle (4GC0) as the template, a homology model of NRT1, also in the outward-occluded state, was compared with the inward-open state of NRT1 of the POT/PTR family (4CL5; TCID 2.A.17.3.1) using the standard deviation measurement function of RR distance maps in Chimera 1.10. A region corresponding to a loop connecting TMSs 11 and 12 was found to display the greatest distance remodeling in the C-terminal domain, approximately 10–15 standard deviations of change.

Schematic illustration of the substrate translocation pathway

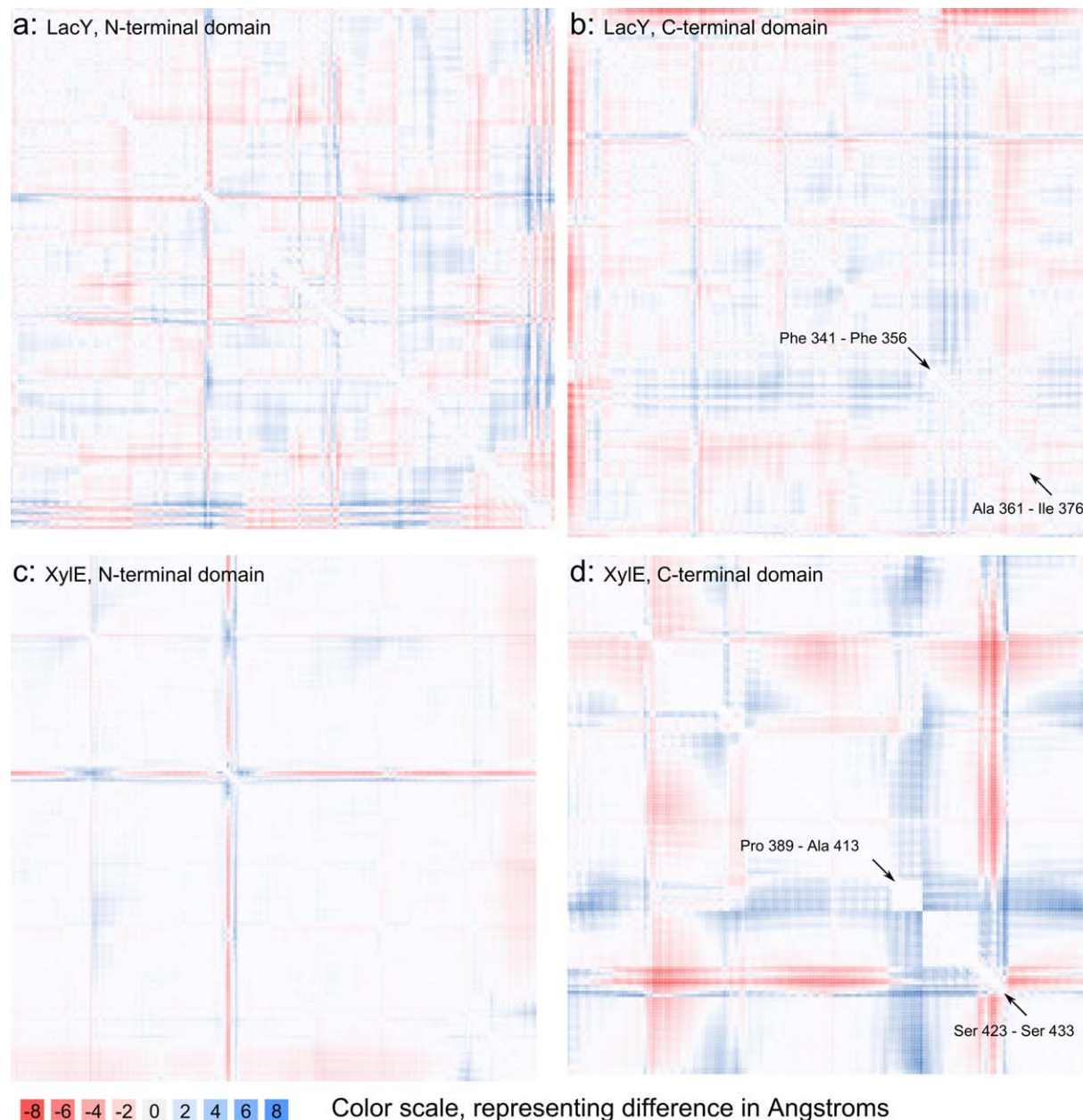
We used the model clipping function in Chimera to create images representing cross-sections of the protein at different membrane-parallel planes. We used the command “mclip all offset 1” to move the clipping plane by 1 Å. The cross-sections were exported to the Inkscape vector graphics drawing program where a vector path was placed around the edge of the channel in each cross-section. The images of the cross-sections were organized in a regular way such that the paths were also spaced regularly. The vector graphics file was parsed using a Perl

script that converted the paths to coordinates viewable in molecular graphics programs. This PDB-like output could subsequently be opened in Chimera. By providing a reduced representation of the surface area, we were better able to view the shape of the channel.

Error-scaling of difference distance matrices

Difference distance matrices, which contain the absolute difference between Cartesian coordinate vectors of atoms, were first introduced in 1974¹⁷ and can be used to reconstruct the geometry of a molecule except its handedness. To be able to objectively compare protein structures, we have employed the normalization proposed in Thomas R. Schneider's error-scaled difference distance matrices.¹⁸ Noise can originate from coordinate errors, suggesting that difference distance matrices need to be normalized to their error levels. A heuristic approach to find the standard uncertainties has been developed in cases where the inversion of the full least-squares matrix is not available. These formulas can be used to take the B values (temperature factors) in PDB files and put them on a common scale so that their influence can be factored into difference distance matrices. The following parameters can be obtained from PDB files: (1) the number of fully occupied atomic sites, (2) the number of reflections, (3) the completeness, (4) the resolution, and (5) the R_{free} . The positional error is $\sigma_r^{\text{DPI}}(B_{\text{avg}}) = 3^{1/2} (N_i/n_{\text{obs}})^{1/2} C^{-1/3} R_{\text{free}} d_{\text{min}}$. For 2Y5Y, $N_i = 382$, $n_{\text{obs}} = 35,055$, $C = 97.7$ (completeness for range, %), $R_{\text{free}} = 0.30$ and $d_{\text{min}} = 3.38$. $\sigma_r^{\text{DPI}}(B_{\text{avg}})_{2Y5Y} = 0.040$. For 4OAA, $N_i = 382$, $n_{\text{obs}} = 20,954$, $C = 95.3$ (completeness for range, %), $R_{\text{free}} = 0.29$ and $d_{\text{min}} = 3.50$. $\sigma_r^{\text{DPI}}(B_{\text{avg}})_{4OAA} = 0.052$. For 4JA3, $N_i = 426$, $n_{\text{obs}} = 17,321$, $C = 82.3$ (completeness for range, %), $R_{\text{free}} = 0.31$ and $d_{\text{min}} = 3.80$. $\sigma_r^{\text{DPI}}(B_{\text{avg}})_{4JA3} = 0.074$. For 4GBY, $N_i = 426$, $n_{\text{obs}} = 19,517$, $C = 98.4$ (completeness for range, %), $R_{\text{free}} = 0.27$ and $d_{\text{min}} = 2.81$. $\sigma_r^{\text{DPI}}(B_{\text{avg}})_{4GBY} = 0.042$.

The estimated positional error is $\bar{\sigma}_{r,i} = \frac{\sigma_r^{\text{DPI}}(B_{\text{avg}})}{B_{\text{avg}}} B_i$. For 2Y5Y, $B_{\text{avg}} = 157.24$. For 4OAA, $B_{\text{avg}} = 157.54$. For 4JA3, $B_{\text{avg}} = 224.95$. For 4GBY, $B_{\text{avg}} = 75.72$. Note that, for Xyle, 4GBY has particularly high accuracy, whereas 4JA3 has poor accuracy [Fig. 1(c,d)]. The estimated standard deviation of an element of the difference distance matrix, $\sigma(\Delta_{ij}^{ab})$, can be estimated by calculating: $\sigma(\Delta_{ij}^{ab}) = \left[(\sigma_{r,i}^a)^2 + (\sigma_{r,j}^a)^2 + (\sigma_{r,i}^b)^2 + (\sigma_{r,j}^b)^2 \right]^{1/2}$. The elements of the error-scaled matrix can be obtained by calculating: $E_{ij}^{ab} = \Delta_{ij}^{ab} / \sigma(\Delta_{ij}^{ab})$. Note that four separate B values go into the normalization of each cell. While we noted a marked global difference in the average B values,

**Figure 1**

The Δ -distance maps of the occluded states of LacY and XylE. (a) and (b): Δ -distance map (LacY), calculated by subtracting the distance map of 4OAA (outward-occluded state) from that of 2Y5Y (inward affinity inactivator-occluded state); 2Y5Y-4OAA (“inward–outward” convention). Blue (positive difference) indicates that a distance between two backbone α -carbons (CA) increased as the protein transitioned to the inward occluded state, while red indicates a negative difference, meaning that the distance decreased. The first panel (a) shows quadrant 2 of the plot of the full structure and compares the distances in the N-terminal domain of LacY, and the second panel (b) shows quadrant 4, comparing distances within LacY’s C-terminal domain. (c) and (d) Δ -distance map of 4JA3-4GBY (XylE in the inward-occluded and outward-occluded states, respectively), comparing distances in the N-terminal (c) and C-terminal (d) domains. TMS10B and TMS11 are located between the arrows with associated residue ranges presented in panels (b) and (d).

especially between inward-occluded XylE (4JA3) and outward-occluded XylE (4GBY), this does not affect the TMS11 region. Plotting the raw B -values for the four files also revealed nothing that would suggest that the B -values would be comparatively high in the TMS11 region.

RESULTS

Δ -distance mapping

We calculated Δ -distance maps comparing the distance remodeling between outward- and inward-occluded states in the N- and C-terminal domains of LacY and

Table I

Features of 2Y5Y and 4OAA (inward affinity inactivator-bound and outward-occluded states of LacY, respectively)

Name	Residues in sequence	Location in structure
<i>N</i> -terminal cross 1	Leu 70–Leu 72	Border between TMSs 2 and 3
<i>N</i> -terminal cross 2	Tyr 113–Gly 115	Mid-TMS4
Edge of <i>N</i> -terminal quadrant	Leu 178–Lys 188	<i>C</i> -terminal half of TMS6
Edge of <i>C</i> -terminal quadrant	Ala 207–Ser 209	Middle of interdomain linker; end of small helical segment protruding from TMS7 ^a
Big blue cross (<i>C</i> -terminal quadrant)	Phe 341–Phe 356	<i>N</i> -terminal half of TMS11
Big red cross (<i>C</i> -terminal quadrant)	Ala 361–Ile 376	<i>C</i> -terminal half of TMS11

The shift in the numbering between the location in the diagram (Fig. 1) and the actual residue number in the protein is due to missing residues between the structures.

^aThere are two additional helical units that are close together when the inward face of the transporter is closed.

Xyle (Fig. 1; Tables I and II). This outward- to inward-occluded state transition, as discussed later, represents the largest conformational change that occurs between any of the four known states of these transporters. We also included Δ -distance maps comparing inward-occluded and inward-open states of LacY and Xyle (Supporting Information) and analogous comparisons using theoretical models of outward-open states (see “Repeat-swapped homology modeling” section). These are treated separately for two reasons: (1) they represent more limited conformational changes, and (2) they represent usage of theoretical data and are more speculative. Together, these maps cover all four known conformational states that these MFS transporters can assume.

To clarify, the measurements taken, and the differences calculated between corresponding measurements, were not transformed before the values were pasted into an Excel spreadsheet, but normalization was carried out separately using ESCET 0.7. The color scale was normalized based on the current value distribution. We have indicated this fact by including a scale bar in Figure 1, translating values to Angstroms.

Repeat-swapped homology model of the outward-open states

When we compared the theoretical models of the outward-open states of both LacY and Xyle with their X-ray crystallography-determined outward-occluded coun-

terparts, we noted that the differences in RMSD values (5.8 and 6.4 Å for LacY and Xyle, respectively) were larger than in the comparison between the inward-occluded and the inward-open states of Xyle (1.8 Å). This suggests that the biological differences that exist between the modeled outward-open state and the outward-occluded state are partly masked by differences that the methodology introduces.

By using Δ -distance maps, we compared the outward-occluded state of LacY with the repeat-swapped model of the outward-open state.⁵ A cross-shaped pattern in the fourth quadrant of the plot represents internal movement in the *C*-terminal 6 TMS bundle. The position of this cross corresponds to a short loop connecting TMSs 10 and 11 (Supporting Information Fig. S1). The red color of the cross indicates that this part of the structure moves closer to the rest of the *C*-terminal domain when shifting from the outward-open to the outward-occluded state. We performed an analogous contact map comparison for the *C*-terminal domain of Xyle but were unable to determine corresponding characteristic patterns.

Pairwise comparison of states

“4OAA” (outward-occluded LacY)

Six TMSs contribute to the internal surface of the upper layers of the substrate translocation pathway: TMSs 1 and 7, TMSs 2 and 8, and TMSs 11 and 5.

Table II

Features of 4JA3 and 4GBY (inward- and outward-occluded states of Xyle, respectively)

Name	Residues in sequence	Location in structure
<i>N</i> -terminal cross	Ile 112–Thr 117	Long loop between TMSs 3 and 4 ^a
Edge of <i>N</i> -terminal quadrant	Thr 247–Gly 263	Corresponds to third interdomain helical segment
Big blue cross (<i>C</i> -terminal quadrant)	Pro 389–Ala 413	<i>C</i> -terminal 1/3 of TMS10 (TMS10C), <i>N</i> -terminal 1/3 of TMS11 (TMS11N) ^{b,c}
Big red cross (<i>C</i> -terminal quadrant)	Ser 423–Ser 433	<i>C</i> -terminal half of TMS11

^aThis extended loop is not found in LacY.

^bThe TMS10B region contains the E(X₆)R motif located at 397–404, conserved across GLUTs 1–4.³ Arg 160–Glu 397 and Glu 153–Arg 404 form interdomain salt bridges that are broken in the inward open state. A third bridge, formed between Glu 397 and Arg 341, is missing in the inward-occluded structure but present in the outward-occluded and inward-open states. Trp 392 forms part of the cytoplasmic lid of the substrate cavity in the outward-occluded state.³

^cIncorrectly identified as being restricted to TMS10B in Quistgaard *et al.* The differences identified by superimposition are the regions that could be identified from averaged superimposition solutions and can differ from the more accurate localization of movement observed by using Δ -distance maps.

TMSs 1 and 7 make up the sides of the elongated cavity, whereas TMSs 8 and 5 and TMSs 2 and 11 connect the sides.

At the altitude of ~ 15 Å from the base of the membrane (it is generally accepted that the fully dehydrated part of the bilayer is 15–25 Å thick,¹⁹ although the full thickness of the membrane is 70–80 Å, including the head groups), the channel area from TMSs 2 and 8 shrinks, while the area from TMSs 4 and 10 emerges to contribute substantially. TMSs 5 and 11 continue to comprise significant surface area. Although TMSs 1 and 7 also contribute channel surface area, they will cease to contribute 10–20 Å further down.

The floor of the cavity is mainly composed of TMSs 10 and 11, but also of TMSs 4 and 5, which are slightly further up (Fig. 2).

“2Y5Y” (inward, affinity inactivator-occluded LacY)

Six TMSs contribute to the internal surface of the lower cytoplasmic open end of the substrate translocation pathway: TMSs 10 and 4, TMSs 8 and 2, and TMSs 5 and 11. TMSs 10 and 4 make up the sides of the elongated cavity, whereas TMSs 8 and 5 and TMSs 11 and 2 connect the sides.

Toward the inner opening of the substrate translocation pathway, the surfaces of TMSs 8 and 2 are on opposite sides of the channel, as are those of TMSs 10 and 4 and TMSs 11 and 5. TMSs on the outskirts of the structure, such as TMSs 1, 6, and 7, do not seem to contribute much surface area to the channel. Toward the middle of the membrane, TMSs 1 and 7 steadily become more apparent and begin to converge with TMSs 2, 4, 8, and 10 as the opening gradually becomes more visible.

The TMS7 surface first emerges between TMSs 10 and 11, and the TMS1 surface emerges between TMSs 4 and 5, although TMSs 4, 5, and 11 are not found deep in the cavity. TMSs 1 and 7 contribute the largest surface area to the floor of the cavity, while TMSs 2 and 7 form narrow stripes on one side, and TMSs 8 and 10 do the same on the other side.

Comparison of the LacY outward-occluded (4OAA) and putative inward-occluded (2Y5Y) states: the tilting movement of TMS11

TMSs 1 and 7, the floor of the 2Y5Y transport pathway, form the sides of the open end of 4OAA. Conversely, the floor of 4OAA, mainly TMSs 10 and 11, but also TMSs 4 and 5, are the major contributors to the occluded “channel” surface at the open end of 2Y5Y.

Δ -distance maps have determined movement of TMS11 when switching from the outward-occluded to the inward-occluded state, in which the *N*-terminal half of TMS11 moves away from the other parts of the

C-terminal domain, and the *C*-terminal half of the TMS11 approaches the rest of the *C*-terminal domain.

Differences between outward-occluded LacY (4OAA) and outward-occluded Xyle (4GBY)

Close to the extracellular sides, both structures appear similar; the same six TMSs contribute surface area lining of the “channel”: TMS2, TMS1, TMS5, TMS8, TMS7, and TMS11, in clockwise order. However, LacY has a better-defined cavity opening, whereas the cavity opening in Xyle is a shallow valley.

Progressing further down the Xyle channel, it turns toward the TMS2/TMS4 side and narrows until its surface area is composed almost exclusively of TMSs 2 and 4. At an intermediate point before the digression, TMSs 5 and 8 cease to contribute to the channel that leads toward the floor of the main cavity. In addition, TMSs 1 and 7 start to form a divider between two subsections of the channel: the shallower TMS5/TMS8 pair and the deeper TMS2/TMS4 pair subsections.

The channel space of Xyle takes on a more complicated shape, potentially containing a “bubble” at the bottom that appears closed off from both ends, but the molecular dynamics of the structure likely permit passage of water. The TMSs contributing surface area to the bottom of the “bubble” are equivalent to the floor TMSs of LacY with the sole exception of TMS4.

Differences between inward inactivator-bound (2Y5Y) LacY and inward-occluded (4JA3) Xyle

At the closed extracellular side, LacY and Xyle both contain surface area from the same TMSs: TMSs 1, 7, 8, and 10. However, in Xyle, there is surface area derived from TMSs 5 and 11 as well.

When observing the Xyle channel opening from the cytoplasmic side, it is important to note that two entrances, divided by a thin column, where TMSs 10 and 11 connect, merge into one slightly wider channel approximately 6 Å down. Here, only four TMSs contribute surface area: TMSs 1, 4, 10, and 11. While both entrances are open to the cytoplasmic side of the molecule, only one is accessible from the hydrophobic wall of the membrane. Two to 3 Å further down, the trough that opens to the side closes, and the channel forms a narrower bottleneck made of TMSs 1, 5, 10, and 11. It widens into a chamber approximately in the middle of the molecule. The floor at this level is mainly composed of TMS7, and the walls derive from TMSs 1, 5, 7, 8, 10, and 11.

DISCUSSION

TMS11 undergoes an analogous movement in LacY and Xyle (Fig. 1, Tables I and II), in which the *N*-terminal halves of the TMSs become more distant from the

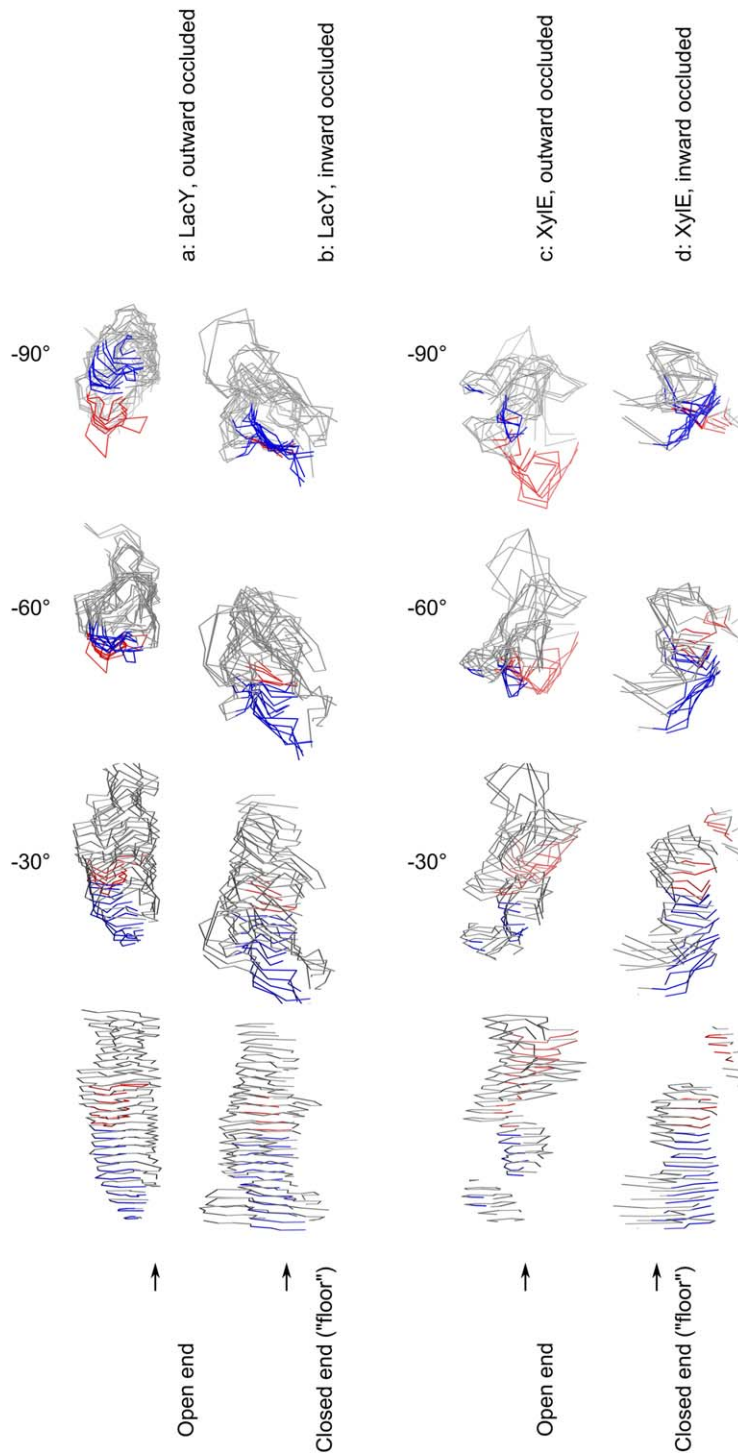


Figure 2

Topographical display of the substrate translocation pathways. The pathway (in the central display, at 0° rotation), contained within the gray, blue, and red outlines, is oriented with the extracellular side at the top of the diagram and the cytoplasmic side at the bottom. It is also shown at -90° , -60° , and -30° rotations: (a) outward-occluded LacY (4OAA); (b) inward affinity inactivator-occluded LacY (2Y5Y); (c) outward-occluded XylE (4GBY); (d) inward-occluded XylE (4JA3). The red and blue traces indicate where TMS11 contributes surface area, in which red signifies that the distance to the C-terminal domain has decreased and blue that it has increased (when transitioning from the outward-occluded to the inward-occluded state; same meaning of the color scheme as in Fig. 1). TMS11 contributes the greatest amount of mobile surface area and displays the greatest individual movement in relation to its domain half in both proteins. Each figure is made from 25 Å steps, starting from the bottom of each channel. The molecular graphics file used to create this display is available on request.

rest of the structure in the inward state while the C-terminal halves of the TMSs move closer to the C-terminal six TMS bundle in the same state. It similarly appears that two other distantly related MFS permeases, FucP and NRT1, exhibit similar TMS11-associated movements.

We examined the Δ -distance maps representing the subsequent transitions between the inward-occluded and the inward-open states of both LacY and XylE and discovered an important difference. We had previously reported⁹ that, in XylE, the N-terminal portion of TMS11 “reverses” by moving back, closer toward the C-terminal six TMS bundle (Supporting Information Fig. S3). However, we found that, in LacY (comparing the affinity activator-bound state with the inward-open state), the N-terminal half of TMS11 in fact “strengthens” the pattern observed when going from the outward-occluded to the inward-occluded state. Thus, it moves even further away from the C-terminal six TMS bundle (Supporting Information Fig. S2). This subtle but notable difference in movement does not mean that the N-terminal half of TMS11 in LacY moves inward to close the cytoplasmic opening. The N-terminal half of TMS11 of XylE only bends slightly toward the rest of the C-terminal bundle, thus qualitatively constituting a reversal of the movement when going between the two occluded states without affecting the width of the substrate pathway. Although this difference may be substrate specificity related, it, nevertheless, shows that the middle of TMS11 must be flexible to physically squeeze the substrate translocation pathway.

A conceptual difference introduced by Solcan *et al.*²⁰ is that while there could be a “hinge” that can bend in the middle of TMSs 10 and 11, this movement could also be “teeter-like”, containing a pivot. These concepts are not mutually exclusive.

While Δ -distance maps are accurate in detecting which part of a sequence moves, detecting the difference between a “pivot point” and a hinge, where bending can occur, is difficult because it necessitates accurate detection of how the angle has changed. In an earlier publication, it had been suggested that the entire TMS 10–11 hairpin opened the gate to the inside in the PepT transporter of the POT family.²¹ Solcan *et al.* reported that some bending/pivoting occurred in the middle of TMSs 10 and 11 at Gly407 and Trp427, respectively. Solcan *et al.* tentatively extrapolated to other members of the MFS (GlpT, FucP, EmrD, and LacY), showing that they also contain a bending point in TMS10 localized to a conserved glycyl residue. These authors investigate the movement of the extracellular end of TMS11 (which moves away from its domain half in Fig. 1), and the authors report how it is packed closer to both TMS2 and TMS7. Solcan *et al.* discuss the transition from the occluded form to the inward-open form without making a clear distinction as to how the transition between the two occluded forms differ from the transition between

the inward-occluded and inward-open forms. In addition, Quistgaard *et al.*^{8,22} previously discussed the movement of TMS10B, and while we had initially thought that this was a mislocalization of the TMS11 movement, based on an incorrect superimposition, reexamination of the residue ranges highlighted in Figure 1 showed that there is evidence of TMS10B movement in XylE. This movement includes Pro389–Ser433 but is not observed in LacY where the movement involves Phe341–Ile376. This could indicate that the conserved glycyl residue identified by Solcan *et al.* may not be an actual bending point.

Careful examination of the structures also showed that while XylE has a discontinuity in TMS10, LacY has this type of discontinuity in TMS11. According to our view that the TMS11-associated movement is caused by movement of the interdomain linker, the movement of TMS11B might consequently be seen as more “indirect.” The conserved tryptophanyl residue in TMS11 has been shown to be substitutable by phenylalanine without loss of function. The question is whether or not TMS11 bends in the middle or is just a pivot point. Tryptophanyl residues are not as flexible as glycyl residues, and we have demonstrated the coupled movement of the extracellular end of TMS11, suggesting that it might be more similar to a pivot than a hinge. We had previously suggested that there is a subtle difference between LacY and XylE in how they behave in the transition from the inward-occluded state to the inward-open conformation (see above). In Supporting Information Figures S2 and S3, we can see evidence of bending, inasmuch as we detect independent movement of the intracellular halves of TMS11 (TMS11A) without coupled movement of TMS11B.

We would like to comment on the use of the structure 2Y5Y as a representation of the “inward-occluded” conformation of LacY. This structure represents LacY bound to a substrate-like affinity inactivator, not a substrate, in the inward conformation, and this conformation is indeed similar to the inward-open conformation of LacY (2V8N), displaying a global RMSD of 1.6 Å between the two structures. In contrast, the inward-occluded and inward-open states of XylE display a global RMSD of 1.9 Å. Strictly speaking, we think that 2Y5Y is an approximation of the inward-occluded conformation but cannot be considered the actual inward-occluded state because of the presence of the inactivator. The subtle difference in biological movement between LacY and XylE as it goes from the inward-occluded (or inward affinity inactivator-bound) state to the inward-open state is probably not attributable to 2Y5Y not representing a genuine “inward-occluded state.” Thus, 2Y5Y may not be identical to the substrate-bound form, as noted earlier, but it may be closer to that form than it is to the inward-open state.

It is a potential misconception of Δ -distance maps that the values found are related to values obtained by

superimposition. A superimposition of two structures will generally produce a globally averaged best solution, but the values in Δ -distance maps show individual distances independent of the rest of the structure. For example, if we superimpose the outward-occluded (4GBY) and inward-occluded (4JA3) conformations of XylE, we obtain a distance measurement of 3.26 Å between the C α atoms of Ala409 in the two conformations. However, the corresponding position in Figure 1(d) is white, indicating a distance difference of zero (0). While this may seem contradictory, it is not, because all Δ -distance maps contain a diagonal of self-comparison distances. While the values in a Δ -distance map represent movement in relation to all other C α atoms, the movement of a certain atom in relation to itself is always, going to be zero. The distance of 3.26 Å found, using a superimposition-based approach, does not represent and cannot be interpreted as a movement because it only represents a displacement that has resulted from the iterative process of finding the best global superimposition average. This is precisely why the use of Δ -distance maps should be strongly encouraged over superimposition-based methodologies. It also provides a likely explanation for why Quistgaard *et al.*^{8,22} only identified movement associated with TMS10B and not TMS11.

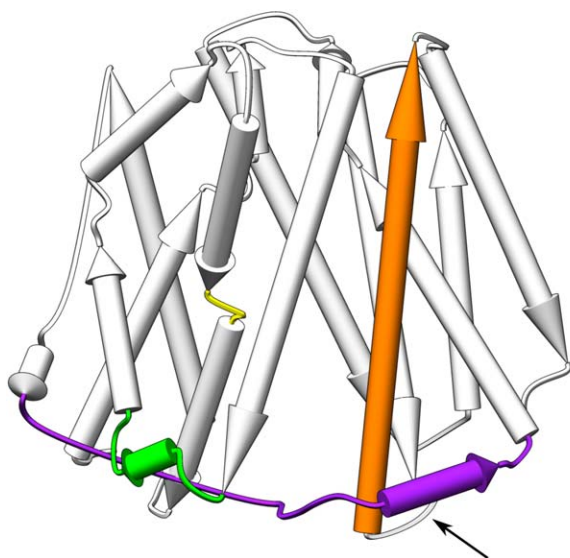
We have relied on theoretical models of the outward-open states of LacY and XylE, created using repeat-swapped homology modeling. Only subtle differences

exist between the outward-open and outward-occluded states because the major conformation changes occur between the two occluded states. However, we did note that a loop connecting TMSs 10 and 11 in LacY (but not in XylE) displayed a marked movement between the outward-open and outward-occluded states.

These findings illustrate the value of Δ -distance maps to facilitate interpretation of multistate transport structures. This becomes particularly evident when noting that the movement of TMS10B, reported earlier,^{7,8} in fact, also includes the *N*-terminal third of TMS11.

In the outward-occluded state of LacY, TMS11 lines the entire cavity, and although it is in principle equivalent to TMS5 in the first half of the protein, TMS11 uniquely contributes to both the side of the outward opening and the floor of the cavity, sealing the cavity from the cytoplasm. In the inward-occluded state, however, TMS11 only contributes to the surface at the cytoplasmic opening of the cavity and does not line the side or floor of the cavity (Fig. 2). The *N*-terminal part of TMS11 forms the floor in the outward-occluded state but an edge of the cytoplasmic open side in the inward-occluded state. The *C*-terminal part of TMS11 forms a side of the extracellular opening in the outward-occluded state (thus being tilted) but an outer scaffolding to the closed end in the inward-occluded state. TMSs 2 and 7 do not contribute to the floor in the outward-occluded state, but they may play a role in facilitating the tilting

a: LacY, inward occluded



b: LacY, outward occluded

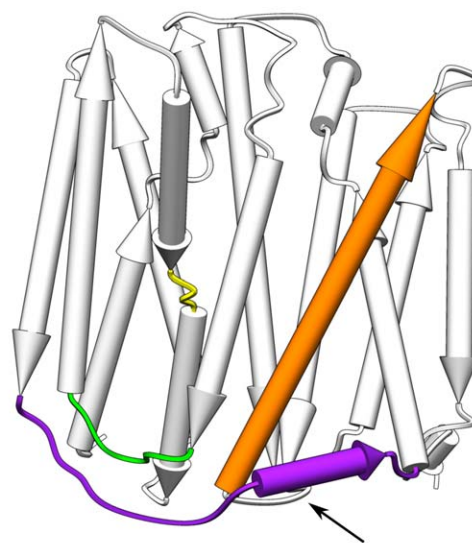


Figure 3

Overview of colocalization of the interdomain linker with structural elements: (a) the inward-occluded state and (b) the outward-occluded state of LacY. The interdomain linker, the loop between TMSs 2 and 3, the mid-TMS4 linker, and TMS11 are highlighted in purple, green, yellow, and orange, respectively. All of these elements generate strong signals on the Δ -distance maps (Fig. 1; Tables I and II) and are colocalized with the interdomain linker. In general, the same pattern holds true for XylE, except that there is a long loop between TMSs 3 and 4, which is located on the extracellular side of the protein that displays independent movement (Table II).

mechanism of TMS11, whose *N*-terminal part is located between these two TMSs, under the interdomain linker.

The tilting movement of TMS11 is related to the location of TMS11 under helices belonging to the interdomain linker. The displacement of TMS11 could either be an effect or a cause of the alternating access movement. TMS11 contains a charge pair whose polarity can be reversed without loss of activity and which is important for membrane insertion of LacY.²³ We have also noted that other portions of the structures displaying individual movement in relation to their domain halves, including the linker between TMSs 2 and 3 and the mid-TMS4, were in close contact with the interdomain linker, providing further evidence for this pattern (Fig. 3).

To generalize the conclusion that a significant TMS11-associated movement is a common feature of MFS members beyond LacY and XylE, we have illustrated using RR distance maps (Supporting Information Fig. S4) that, when the fucose porter, FucP, switches from the outward to the inward state, a significant TMS11-associated distance remodeling occurs similar to that noted for LacY and XylE.

A similar effect was observed within a member of the POT/PTR peptide transporter family, where a comparison between the inward-open and outward-occluded states of NRT1 revealed a significantly elevated standard deviation measurement for a loop located between TMSs 11 and 12. These findings, revealed by examination of recently published MFS structures, show that TMS11-associated movement is not restricted to XylE and LacY. Because these additional findings are based on a homology modeling approach, any further generalizations would be unreliable. However, these initial homology modeling results give us increased confidence that the model presented for either LacY or XylE may be extended to the MFS in general.

There are four principal reasons to believe that the identified movement of TMS11 is a genuine biological effect and not an artifact (which could be the case when comparing X-ray structures of relatively low resolution): (1) the observed effect is on the order of magnitude of 4–6 Å, greater than the approximate resolution limit used of 3 Å, (2) the pattern of TMS11-associated movement is recurrent between multiple comparisons, (3) the similar movement (although incorrectly characterized as a movement limited to TMS10B) has been documented before for XylE,^{8,22} FucP,²⁰ and POT, and (4) these observations are resilient to error-scaling¹⁸ (Supporting Information Fig. S5). The PDB files for LacY and XylE used to draw the principal conclusions of this study were: 4GBY (2.81), 2Y5Y (3.38), 4OAA (3.50), 2V8N (3.60), 4JA3 (3.80), and 4JA4 (4.20), in order of decreasing accuracy, measured in Angstrom resolution, ranging from 2.81 to 4.20 (Table III). However, to make fully validated comparisons, the percentile ranks of R_{free} , clashscore, Ramachandran outliers, side-chain outliers,

Table III

Summary Data of Six Principal Structural Models of Conformational States of LacY and XylE Used in This Study

PDB ID	Protein	Conformation	Resolution ^a
4OAA	LacY	Outward occluded	3.50
2Y5Y	LacY	Inward, affinity inactivator occluded	3.38
2V8N	LacY	Inward open	3.60
4GBY	XylE	Outward occluded	2.81
4JA3	XylE	Inward occluded	3.80
4JA4	XylE	Inward open	4.20

^aThe magnitude of the TMS11 movement differs in panels (b) and (d) of Figure 1, measuring 4 Å for LacY and 6–8 Å for XylE. On average, resolution limits are better for LacY, as they never reach the 4 Å threshold, but not for XylE in the inward occluded state (resolution only 4.2 Å).

and RSRZ outliers have to be considered. We, thus, consider that the differences detected in the Δ -distance maps represent true statistical differences and do not result from inaccuracies in the modeling procedure. Our conclusions are adequately supported by the data, because, with the exception of the repeat-swapped model, the distances we detected are greater than the low-resolution noise, and the same patterns reemerge between multiple comparisons.

CONCLUSIONS

By using superimposition-based approaches, the movement of TMS11 in XylE had not been characterized correctly,^{7,8} because movement of the *N*-terminal half of this TMS appeared to be restricted to the *C*-terminal part of TMS10 (TMS10B), and a similar error in localization was noted for the *N*-terminal part of TMS11 in LacY. In this article, we have shown that more accurate positioning can be achieved using Δ -distance mapping. This established method^{12–14} could have wide applicability as more conformational state data become available. The majority of the TMS segments, including TMS11, which display intradomain movement, independent of the two domains' alternating access movement, were found in close proximity to the interdomain linker helices. This commonality is interpreted as either a consequence of either the alternating access movement or a driving force causing it.

TMS11 in both LacY and XylE undergoes a shared and crucial movement during the shift between outward- and inward-occluded states in which the *N*-terminal (intracellular) halves of TMS11 move away from the *C*-terminal (extracellular) six TMS bundles, while the *C*-terminal halves of TMS11 approach them. TMSs 11 in both porters form the floor of the channel in the outward-occluded state and contribute more mobile internal surface area than any of the other TMSs, indicating that TMS11 plays a unique role in opening the substrate translocation pathway to the cytoplasmic side.

REFERENCES

- Saier MH Jr, Reddy VS, Tamang DG, Västermark A. The transporter classification database. *Nucleic Acids Res* 2014;42(Database issue):D251–D258.
- Law CJ, Yang Q, Soudant C, Maloney PC, Wang DN. Kinetic evidence is consistent with the rocker-switch mechanism of membrane transport by GlpT. *Biochemistry* 2007;46:12190–12197.
- Forrest LR, Krämer R, Ziegler C. The structural basis of secondary active transport mechanisms. *Biochim Biophys Acta* 2011;1807:167–188.
- Dang S, Sun L, Huang Y, Lu F, Liu Y, Gong H, Wang J, Yan N. Structure of a fucose transporter in an outward-open conformation. *Nature* 2010;467:734–738.
- Radestock S, Forrest LR. The alternating-access mechanism of MFS transporters arises from inverted-topology repeats. *J Mol Biol* 2011;407:698–715.
- Deng D, Xu C, Sun P, Wu J, Yan C, Hu M, Yan N. Crystal structure of the human glucose transporter GLUT1. *Nature* 2014;510:121–125.
- Sun L, Zeng X, Yan C, Sun X, Gong X, Rao Y, Yan N. Crystal structure of a bacterial homologue of glucose transporters GLUT1-4. *Nature* 2012;490:361–366.
- Quistgaard EM, Löw C, Moberg P, Trésaugues L, Nordlund P. Structural basis for substrate transport in the GLUT-homology family of monosaccharide transporters. *Nat Struct Mol Biol* 2013;20:766–768.
- Västermark A, Lunt B, Saier M. Major facilitator superfamily porters, LacY, FucP and XylE of *Escherichia coli* appear to have evolved positionally dissimilar catalytic residues without rearrangement of 3-TMS repeat units. *J Mol Microbiol Biotechnol* 2014;24:82–90.
- Kumar H, Kasho V, Smirnova I, Finer-Moore JS, Kaback HR, Stroud RM. Structure of sugar-bound LacY. *Proc Natl Acad Sci U S A* 2014;111:1784–1788.
- Serdiuk T, Madej MG, Sugihara J, Kawamura S, Mari SA, Kaback HR, Müller DJ. Substrate-induced changes in the structural properties of LacY. *Proc Natl Acad Sci U S A* 2014;111:E1571–E1580.
- Schneider M, Brock O. Combining physicochemical and evolutionary information for protein contact prediction. *PLoS One* 2014;9:e108438.
- Chen H, Tian R, Ni Z, Zhang Z, Guo Q, Saier MH Jr. Conformational transition pathway in the inhibitor binding process of human monoacylglycerol lipase. *Protein J* 2014;33:503–511.
- Huang CC, Meng EC, Morris JH, Pettersen EF, Ferrin TE. Enhancing UCSF chimera through web services. *Nucleic Acids Res* 2014;42(Web Server issue):W478–W484.
- Parker JL, Newstead S. Molecular basis of nitrate uptake by the plant nitrate transporter NRT1.1. *Nature* 2014;507:68–72.
- Stamm M, Staritzbichler R, Khafizov K, Forrest LR. AlignMe—a membrane protein sequence alignment web server. *Nucleic Acids Res* 2014;42(Web Server issue):W246–W251.
- Nishikawa K, Ooi T. Comparison of homologous tertiary structures of proteins. *J Theor Biol* 1974;43:351–374.
- Schneider TR. Domain identification by iterative analysis of error-scaled difference distance matrices. *Acta Crystallogr D Biol Crystallogr* 2004;60(Pt 12 Pt 1):2269–2275.
- Gorter E, Grendel F. On bimolecular layers of lipoids on the chromocytes of the blood. *J Exp Med* 1925;41:439–443.
- Solcan N, Kwok J, Fowler PW, Cameron AD, Drew D, Iwata S, Newstead S. Alternating access mechanism in the POT family of oligopeptide transporters. *EMBO J* 2012;31:3411–3421.
- Newstead S, Drew D, Cameron AD, Postis VL, Xia X, Fowler PW, Ingram JC, Carpenter EP, Sansom MS, McPherson MJ, Baldwin SA, Iwata S. Crystal structure of a prokaryotic homologue of the mammalian oligopeptide-proton symporters, PepT1 and PepT2. *EMBO J* 2011;30:417–426.
- Quistgaard EM, Löw C, Moberg P, Nordlund P. Metal-mediated crystallization of the xylose transporter XylE from *Escherichia coli* in three different crystal forms. *J Struct Biol* 2013;184:375–378.
- Dunten RL, Sahin-Tóth M, Kaback HR. Role of the charge pair aspartic acid-237-lysine-358 in the lactose permease of *Escherichia coli*. *Biochemistry* 1993;32:3139–3145.