Flexible Gates Generate Occluded Intermediates in the Transport Cycle of LacY

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Abstract

The major facilitator superfamily (MFS) transporter lactose permease (LacY) alternates between cytoplasmic and periplasmic open conformations to co-transport a sugar molecule together with a proton across the plasma membrane. Indirect experimental evidence suggested the existence of an occluded transition intermediate of LacY, which would prevent leaking of the proton gradient. As no experimental structure is known, the conformational transition is not fully understood in atomic detail. We simulated transition events from a cytoplasmic open conformation to a periplasmic open conformation with the dynamic importance sampling molecular dynamics method and observed occluded intermediates. Analysis of water permeation pathways and the electrostatic free-energy landscape of a solvated proton indicated that the occluded state contains a solvated central cavity inaccessible from either side of the membrane. We propose a pair of geometric order parameters that capture the state of the pathway through the MFS transporters as shown by a survey of available crystal structures and models. We present a model for the occluded state of apo-LacY, which is similar to the occluded crystal structures of the MFS transporters EmrD, PepTSo, NarU, PiPT and XylE. Our simulations are consistent with experimental double electron spin–spin distance measurements that have been interpreted to show occluded conformations. During the simulations, a salt bridge that has been postulated to be involved in driving the conformational transition formed. Our results argue against a simple rigid-body domain motion as implied by a strict “rocker-switch mechanism” and instead hint at an intricate coupling between two flexible gates.

Introduction

Lactose permease (LacY) is the best-studied member of the major facilitator superfamily (MFS) of secondary transporters [1]. LacY of Escherichia coli is a symporter and uses the proton (H⁺) electrochemical gradient to transport galactosides, such as lactose, from the periplasm into the cytoplasm [2]. The structural elements that allow or prevent exchange of sugar and H⁺ between the central binding site and either the cytoplasm or the periplasm are often referred to as “gates” [3]. LacY transports substrate by switching from a conformation with an open cytoplasmic gate and a closed periplasmic gate to a periplasmic open conformation in which the states of the gates are reversed. This alternating access mechanism [4,5] has been detected [6] in site-directed alkylation [7] and cross-linking [8], tryptophan quenching [9], single-molecule Förster resonance energy transfer [10] and double electron–electron spin resonance (DEER) [11] experiments.

The transition between the cytoplasmic and periplasmic open states of LacY is not understood in full atomic detail. Partly, this reflects a lack of structural data. Only crystal structures for cytoplasmic open [12–15] LacY have been obtained. The crystal structure of the distantly related MFS transporter FucP from E. coli has been solved in a periplasmic open conformation [16]. Radestock and
Forrest have built two models of periplasmic open LacY: the first is based on its homology to the FucP structure and the second uses the internal inverted repeat symmetry of the transporter [17]. Biochemical studies [2] have identified key events in the transport. First, cytoplasmic open LacY releases the transported sugar to the cytosol. The subsequent deprotonation of E325 initiates the transition to the periplasmic open conformation [18]. Periplasmic open LacY is a minor conformer [2] in the absence of substrate. Protonation of H322 and substrate binding then stabilizes the periplasmic open state and initiates the return to the cytoplasmic open conformation, thereby carrying sugar and H⁺ across the membrane.

In addition to biochemical evidence, functional considerations provide some clues about the mechanism of the conformational transition. A transporter with both gates open would dissipate vital transmembrane gradients, such as the proton gradient in bacteria or the sodium gradient in higher eukaryotes to the detriment of the cell that has expended ~20% of its energy to establish the gradient [19]. The two gates of a transporter should therefore never be open at the same time [3] and it is thought that, to prevent this, the movement of the gates must occur in a precise order so that one gate closes before the other starts to open as in the operation of an airlock (Fig. 1a) [20]. Such an “airlock model” for transport predicts an occluded intermediate conformation that is closed to ion and substrate permeation from both cytoplasm and periplasm. The occluded conformation is thus fundamental to the transport mechanism. Preventing both gates from being open at the same time necessitates coupling between the movements of the two gates so that only the closure of one gate allows opening of the other gate. We describe the case where one gate must close fully before the other can begin to open as a *sequential motion*. In this case, the transporter passes through intermediates that are fully occluded.

It has been hypothesized that a rigid-body movement of the N- and C-terminal domains of LacY relative to one another [16] (Fig. 1b) would allow the cytoplasmic and periplasmic open states to interconvert. This model is called the “rocker-switch mechanism” [1,12]. The rigidity of the domains would provide a direct mechanical linkage between the two gates so that the closure of one gate would occur simultaneously with the opening of the other. We describe this as a *simultaneous motion*. A truly rigid motion of the two domains would probably not be able to form a fully occluded conformation. Allowing the domains some flexibility would allow the formation of an occluded conformation, ruling out the dissipation of transmembrane potentials through channel-like conformations. Hydrogen/deuterium exchange experiments have established that LacY is more dynamic than many other membrane proteins [21–23] and is therefore unlikely to contain large, rigid domains. The conformational flexibility of LacY has also been highlighted in NMR experiments [24] and in molecular dynamics (MD) simulations [24.25].

In the absence of direct evidence, we cannot be certain if LacY passes through an occluded intermediate and, if it does, what such an intermediate would look like. DEER (also known as PELDOR) measurements for a range of residue pairs on LacY have been interpreted to demonstrate the existence of an intermediate occluded state [11,26] and have recently been used to propose a model of an occluded conformation of LacY [27]. To date, occluded conformations of five MFS transporters at atomic resolution have been published. These are EmrD [28], NarU [29], XylE [30,31] (all from *E. coli*), PepTb [32] (from *Shewanella oneidensis*) and PiPT [33] (from the fungus *Piriformospora indica*). In addition, an occluded conformation of the MFS transporter OxlT from *Oxalobacter formigenes* was resolved by electron microscopy, albeit at a lower resolution of 6.5 Å [34].

Computer simulations can suggest intermediate conformations and plausible transition mechanisms [35] that may then be compared to experimental data. The crystal structures of LacY and GlpT transporters are in cytoplasmic open conformations. The cytoplasmic gate of both structures has been observed to undergo partial closure in equilibrium.

Fig. 1. Schematic models of transport resulting in alternating access. A transporter consisting of two domains (light and dark gray) may undergo conformational transitions that expose a central substrate binding site alternatively to the extracellular and the intracellular side. (a) In the *airlock model*, two gates open and close sequentially to form an outward open and inward open conformation. In order to prevent formation of a leak state, the gates must be strongly coupled so that they never open at the same time. (b) In the *rocker-switch model*, the two domains move simultaneously as rigid bodies relative to each other and thus form the outward open and inward open states. No additional coupling is required to prevent formation of a leaky transporter.
MD simulations, thereby forming partially occluded conformations [18,25,36]. The turnover number for LacY is $16-21 \text{ s}^{-1}$ [37], thus, on average, one transition every 50,000 $\mu$s would be expected. At present, standard atomic-resolution equilibrium MD simulations typically yield trajectories of the length of up to $\sim 1 \mu$s and hence we cannot readily characterize the kinetics of transport using such simulations. Reducing the level of detail modeled is one way to overcome this challenge and this approach enabled the transition between cytoplasmic and periplasmic open FucP to be studied using an elastic network model [38], which hinted at the existence of an occluded intermediate. Alternatively, fully atomistic approaches that bias the sampling, such as stratified umbrella sampling [39], dynamic importance sampling (DIMS) MD [40–42] or self-guided Langevin dynamics (SGLD) [43], can be used to study such conformational changes. Pendse et al. used SGLD to simulate the transition from cytoplasmic open to periplasmic open LacY [44]. Although transition intermediates obtained from the SGLD simulations contained both cytoplasmic and periplasmic constrictions, no detailed study of the likely functional relevance of these structures was presented.

In this paper, we will investigate if LacY switches conformation via an occluded state and what conformation the protein could plausibly adopt in such a state. These results will then allow us to analyze if the broad conformational changes involved in the transport mechanism are compatible with either the airlock or the rocker-switch mechanism. We will use both equilibrium and DIMS [40,41] MD simulations to study transitions between the

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**Fig. 2.** Atomistic simulations of LacY. The wild-type crystal structure of cytoplasmic open LacY (PDB code 28VN) (a) [14] and a FucP-based homology model for periplasmic open LacY (PMDB PM0077183) (b) [17] used in atomistic simulations are shown with the periplasmic face at the top and the cytoplasmic face at the bottom. The N-terminal and C-terminal domains, residues 7–185 and 220–399, are colored cyan and magenta, respectively. The linkers connecting the domains and flexible N- and C-terminal stretches are shown in black. (c) The membrane and solvent environment was represented implicitly as regions with differing dielectric constants $\varepsilon$ in electrostatics calculations (using the Poisson-Boltzmann approach [45]) and simulations of rare events (with the GBSW solvent model [46–48]). (d) In equilibrium MD simulations, both the protein and the membrane and solvent environment were represented in full atomistic detail. Cytoplasmic open LacY inserted into a POPC membrane, highlighted in gray, is shown here. The solvent molecules and ions are omitted for clarity. The structures were rendered in VMD [49].
cytoplasmic open and periplasmic open conformations of LacY. In particular, we will use the biased DIMS MD simulations in implicit solvent to accelerate sampling of conformational transitions and then study conformations in full detail with explicit solvent equilibrium MD (Fig. 2), similar to the approach taken by Pendse et al. [44]. The simulations reveal fully occluded intermediate structures. By introducing geometric order parameters that track the states of the two gates, we show that our results are consistent with crystal structures of related MFS transporters. Furthermore, our simulations also agree with experimental DEER measurements [11] of LacY and exhibit the formation of a salt bridge postulated to be involved in driving the conformational transition [15,18].

Results and Discussion

Equilibrium MD simulations of cytoplasmic open and periplasmic open conformations

We first simulated both end states of the conformational cycle in a 1-palmitoyl,2-oleoyl-sn-glycero-3-phosphocholine (POPC) membrane (Fig. 2) to probe the conformational flexibility (discussed in further detail in the supplementary information, especially in Figs. S1 and S2) of the transporter and establish a baseline for its dynamical behavior. The wild-type crystal structure of LacY [14] served as the cytoplasmic open conformation (see Table 1 for a list of all simulations). For the periplasmic open conformation, we used Radestock and Forrest's homology model [17], which is based on the periplasmic open FucP structure [16]. We simulated the apo conformation of both end states; that is, there was no sugar bound and E325 was not protonated. The apo conformations should be on the cusp of changing conformation [2] and thus simulation might reveal the initial stages of the transition. Since simple visual inspection of a static structure is not sufficiently accurate to determine if the central binding site is accessible to both substrate and protons, we will apply a range of computational methods to assess the functional state of the generated protein conformations (Fig. 3). We will first characterize the functional state of the transporter in the simulations (i.e., which gates are open or closed) by calculating the radius of the translocation pathway through the transporter. Then, we shall examine the density of water and the electrostatic free energy of a permeating H⁺. These are more sensitive and together will allow us to assign the likely functional state of any structure.

During equilibrium MD simulations of the model of periplasmic open LacY, the periplasmic gate remained open and the cytoplasmic gate stayed closed (see simulation POpenA in Fig. 3c and repeat simulation POpenB in the supplementary content). These simulations demonstrated that both the conformation and its functional state were maintained over the duration of the simulations.

In contrast, during simulations of cytoplasmic open LacY, the periplasmic gate remained closed but the cytoplasmic gate started to close. In simulation COpenA1, the minimum radius on the cytoplasmic side of the transporter decreased from ~4 Å in the crystal structure to ~1 Å (Fig. 3a). The same effect was seen in multiple simulations with different box dimensions and force fields (as evidenced by the pore radius profiles in Fig. S3). The observed closure did not therefore depend on the dimensions of the box or the force field used. The variation of pore radius along the membrane normal suggests that sugars would be sterically blocked from entering the central cavity from both cytoplasm and periplasm. Our results are consistent with earlier studies that observed a similar closure of the cytoplasmic gate in multiple equilibrium MD simulations [18,25] or a more limited closure after 10 ns of simulation [56]. Taken together, these results show that the cytoplasmic open apo structure of LacY, as resolved by X-ray crystallography, quickly relaxes to a more occluded structure that would prevent sugar

| Name of simulation | Starting structure | Number of POPC lipids | Length of simulation (ns) | Force field |
|--------------------|--------------------|-----------------------|--------------------------|------------|
| COpenA1            | 2V8N crystal structure | 126                   | 100                      | GROMOS96 43a1 |
| COpenA2            | COpenA1 end structure | 126                   | 100                      | GROMOS96 43a1 |
| COpenB1            | 2V8N crystal structure | 256                   | 100                      | GROMOS96 43a1 |
| COpenB2            | COpenB1 end structure | 256                   | 200                      | GROMOS96 43a1 |
| COpenC             | 2V8N crystal structure | 152                   | 100                      | CHARMM22 + CHARMM36 |
| POpenA             | FucP-based homology model | 254               | 100                      | GROMOS96 43a1 |
| POpenB             | FucP-based homology model | 254               | 100                      | CHARMM22 + CHARMM36 |
| Occ                | End structure DIMS cytoplasmic closure | 256             | 100                      | GROMOS96 43a1 |
| COpen-GBSW         | 2V8N crystal structure | Implicit membrane | 10 | CHARMM22 + GBSW |
| POpen-GBSW         | FucP-based homology model | Implicit membrane | -12 | CHARMM22 + GBWS |
| Occ-GBSW           | End structure of Occ | Implicit membrane | 10 | CHARMM22 + GBWS |
permeation and might be a functionally relevant intermediate [25].

We assessed if this intermediate is functionally occluded, that is, whether solvated protons could still permeate, despite the closure of the cytoplasmic gate. We first calculated water densities from the MD simulations to define aqueous pathways that could act as proton conduits. Since the desolvation of a charged species is energetically unfavorable (in the absence of compensating interactions), dehydration of a gate will prevent ion permeation [45,52]. The water density for the simulation of cytoplasmic open LacY (Fig. 3d and Fig. S4a and b) indicated that the constriction between the cytoplasm and the central

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**Fig. 3.** Functional characterization of LacY in MD simulations of multiple conformational states. Pore radii as determined by HOLE [50] are shown along the membrane normal z for the central cavity in simulations cytoplasmic open COpenA1 (a), occluded Occ (b) and periplasmic open POpenA (c). Shaded regions indicate 1 SD of the radius from the mean value. Water densities relative to bulk SPC water [51] (0.970 g/cm³) were calculated with gridcount [52,53] and visualized as 3D densities in UCSF Chimera [54,55]. Cuts through the densities for simulations COpenA1 (d), Occ (e) and POpenA (f) are shown. For COpenA1 and Occ, the last frame from 100 ns of equilibrium MD is depicted. For POpenB, the equilibrated starting structure for the 100-ns simulation is shown. The position-resolved electrostatic free energy $W_{elec}(r)$ for solvated H⁺ interacting with cytoplasmic open (g), fully occluded (h) and periplasmic open LacY (i), calculated with BornBrofiler [45] (see Materials and Methods) and visualized in UCSF Chimera [54,55] is shown.
cavity remained partially solvated even after the initial closure of the cytoplasmic gate, indicating that protons could still be expected to permeate and thus the conformation cannot be considered a functionally occluded state. No solvent-filled pathway connected the cavity and the periplasm in the simulation of cytoplasmic open LacY and thus the periplasmic gate remained completely shut. The cytoplasmic gate in the simulations of periplasmic open LacY remained dehydrated (Fig. 3f) whereas a wide aqueous pathway connected the central cavity to the periplasmic solution, suggesting the conformation represents a functional periplasmic open state.

Poisson–Boltzmann calculations of the electrostatic free energy of a solvated proton offer another way of assessing whether a gate is open since the heights of the energetic barriers between the cavity and bulk can be quantified [45]. The energies calculated for the cytoplasmic open (Fig. 3g) and periplasmic open (Fig. 3i) conformations suggest that the cytoplasmic and the periplasmic gates, respectively, offer no barrier to the movement of protons and thus would permit relatively free exchange of H+ between the bulk solvent and the central cavity. Electrostatic free energies in the cytoplasmic and periplasmic vestibules and the central cavity were only slightly less favorable than bulk (0–10 kJ/mol). The periplasmic gate of cytoplasmic open LacY is closed because H+ exchange between the central cavity and the periplasm would involve crossing a large volume with unfavorable electrostatic free energies (~100 kJ/mol; Fig. 3g). By the same reasoning, the cytoplasmic gate of the periplasmic open transporter is also closed to H+ permeation (Fig. 3i). The initial closure of the cytoplasmic gate seen in Fig. 3a led to only a limited increase in the electrostatic free-energy barrier in the space connecting the cavity to the cytoplasm, from 0–10 kJ/mol to 10–20 kJ/mol (Fig. S5). This modest increase in the electrostatic free energy for proton permeation together with the water density calculations suggests that the observed closure of the cytoplasmic gate resulted in a partially functionally occluded intermediate. The gate appears therefore impermeable to sugar but not to H+ exchange between the cytoplasm and the internal cavity.

**DIMS simulations of gate closure yield a fully occluded conformation**

Eight equilibrium MD simulations of cytoplasmic and periplasmic open LacY with an aggregated total time of 0.8 μs yielded neither a transition between the end states nor a functionally fully occluded intermediate. This is not surprising given that, on average, one transition would be expected to be observed every 50,000 μs for a typical turnover number of 16–21 s⁻¹ [37]. We therefore used DIMS MD [40] to investigate whether LacY passes through an occluded state as it changes from a cytoplasmic open conformation to a periplasmic open conformation. In DIMS MD, an importance sampling procedure is used to generate transitions between an initial structure and a target structure. DIMS MD has been successfully applied to study how other membrane proteins, such as a potassium ion channel [57] and a transporter [58], change conformation.

We first used DIMS MD to generate trajectories starting from the cytoplasmic open structure of LacY during which the cytoplasmic gate would close (these will be referred to as *DIMS-closure*). It was assumed that the cytoplasmic and periplasmic gates are located on the cytoplasmic and periplasmic sides of the transporter, respectively. Therefore, only the cytoplasmic half of periplasmic open LacY, that is, the one featuring the closed cytoplasmic gate, was used as the target structure. The DIMS importance sampling procedure then used this target structure to select pathways that led to cytoplasmic closure. The variation in the pore radius profile along the membrane normal of the final structure from the DIMS MD simulation showed constrictions on both the cytoplasmic and periplasmic sides (Fig. 3b), consistent with it being an occluded structure. A similarly constricted structure was generated from periplasmic open LacY using the periplasmic half of cytoplasmic open LacY, that is, the closed periplasmic gate, as the target structure (data not shown).

To help assess whether the DIMS method had produced a functionally fully occluded conformation, we simulated the final structure from DIMS MD for 100 ns (simulation Occ) using standard equilibrium MD. The putative occluded conformation was stable for the entire simulation as shown by the RMSD relative to the starting conformation (Fig. S1d) and the preservation of the secondary structure (Fig. S2c). The central water-filled internal cavity kept its shape. We then calculated how the pore radius varied along the membrane normal during the trajectory *Occ* (Fig. 3b). These pore profiles show that the minimum pore radii at the cytoplasmic and periplasmic constrictions remained smaller than the radius of a water molecule (~1.4 Å) during the entire simulation. The radius of gyration of lactose is 3.5 Å and therefore sugar substrates would not be able to access the central cavity. The pore profile is similar to the one observed in MD simulations of the occluded structure of the related MFS transporter EmrD [59].

The water density computed from the simulation of the occluded structure, *Occ*, was also consistent with both gates being closed. Slices through the three-dimensional (3D) water density show that regions with very low density separate the central cavity from both the cytoplasm and the periplasm. The internal cavity remained solvated (Fig. 3e) at between 0.7 and 1.4 times the bulk water density. Some water, however, remained in the interfacial regions. For example, on the cytoplasmic side, water
density at 70% of bulk water is apparent in the slice shown in Fig. 3e but only regions with low density connect it to the cytoplasm and the internal cavity. While ions are likely to be stable in the water-filled central cavity, there are no continuous permeation pathways joining either the periplasm or the cytoplasm to the central cavity.

The structures generated by the Occ simulation were clustered (see Materials and Methods). The last frame of the simulation was chosen as a representative structure (Fig. 4a) since it belonged to the dominant cluster. This structure is of comparable quality to the wild-type crystal structure of cytoplasmic open LacY (see the supplementary content). The electrostatic free energy of placing a proton in this structure was calculated as before. Energies in the internal cavity ranged from 0 kJ/mol to 20 kJ/mol (Fig. 3h), but importantly, very high electrostatic free energies (> 100 kJ/mol) were obtained in the spaces on the periplasmic and cytoplasmic sides of the internal cavity. Some regions with more favorable electrostatic free energies (20–50 kJ/mol) exist between the internal cavity and the cytoplasm and the periplasm, as can be seen on the periplasmic side of the slice through the 3D density. Given that such regions were surrounded by space very unfavorable to protons, it is reasonable to conclude that electrostatic barriers to ion permeation from both the cytoplasm and the periplasm exist. The electrostatics and water density calculations are thus consistent with this structure of the transporter being not only sterically closed to sugar substrates but also occluded to protons and thus preventing a proton leak across the membrane.

Moreover, the model for the occluded state (Fig. 4a) is also consistent with the proposed mechanism for the transport cycle [2]. The residues crucial for function E269, R302, H322 and E325 (highlighted in Fig. 4), as well as E126 and R144 (data not shown), surround the central cavity as expected. In particular, we found that E325 and R302 form a stable salt bridge in simulation Occ and in our occluded state model (Fig. 4c) but neither in the cytoplasmic (Fig. 4b) nor in the periplasmic open (Fig. 4d) structures that were the inputs for our study. The two residues approach each other during the DIMS-closure simulation but the salt bridge only fully forms during the Occ equilibrium simulations as shown in Fig. S6 and discussed in more detail in the supplementary content. Experiments on an R302C/E325C mutant using excimer fluorescence suggested that these residues could be in close proximity [61]. Functional assays of mutants indicated that the formation of a salt bridge between R302 and E325 enables the release of the transported proton [62] so that the apo protein could transition from a cytoplasmic open to a periplasmic open conformation [18]. Thus, our computer simulation reveals a necessary step in the transport cycle and captures details of the molecular interactions that are hard to discern from static structures alone.

Comparison of equilibrium MD simulations with DEER measurements

Having established that we have obtained a functionally, fully occluded state, we validated this model and the simulations by comparing them with

![Fig. 4. Occluded state model.](image-url)
distances derived from DEER experiments [11]. These have been previously interpreted as reporting the presence of cytoplasmic and periplasmic open states (revealed after the addition of the binding sugar, NPGal) and occluded conformations [27]. To better interpret these data, we have developed an approach (see Materials and Methods) that takes into account the flexibility of the spin label and thus predicts the electron spin–spin distance distribution that would be observed for a single structure or an entire MD trajectory. This is important as it allows us to compare like with like. We compared nine experimental distances to our simulations: three on the periplasmic side of the protein (residue pairs 105–310, 164–310 and 164–375; Fig. 5a) and six on the cytoplasmic side (residue pairs 73–401, 73–340, 136–340, 137–340 136–401 and 137–401; Fig. 5b). As the periplasmic open homology model [17] lacks some C-terminal residues, spin–spin distances 73–401, 136–401 and 137–401 can only be compared for the simulations of cytoplasmic open LacY.

The distance distributions calculated from the simulations of cytoplasmic open LacY agree well with the DEER experiments on LacY in the presence of NPGlc. This non-binding sugar ensures that the protein is found predominately in cytoplasmic open conformations [11]. In general, there is substantial overlap between the distributions calculated from simulations and the positions of the major peaks from the DEER experiments (Fig. 5, red lines versus gray histograms; Fig. S7a in the supplementary content). The absolute heights and widths of the experimental peaks are not expected to quantitatively agree with the simulations because the three independent simulations of individual conformational states do not capture the relative distribution of different states observed in a single experiment.

The distance distributions estimated from the simulation of periplasmic open LacY (blue lines in Fig. 5c) show the pattern expected for a periplasmic open state, that is, the spin label distances on the periplasmic side are shifted to larger values and the distances on the cytosolic side are shifted to smaller values compared to the equivalent distances estimated from the simulations of the cytoplasmic open transporter. One exception is distance 164–375 where the end of helix 5 with residue 164 moved closer to residue 375 on helix 11 at the beginning of the simulation and thus inverted the expected pattern. The distributions from the simulations generally overlap with the positions of the major experimental peaks for predominately periplasmic open LacY (Fig. 5c, blue lines; Fig. S7b).

Finally, we can validate the model of the occluded state against the DEER data. The spin–spin distance distributions calculated from the simulation Occ of the occluded model (Fig. 5c, magenta lines) are consistent with the experimental data. On the cytosolic side of the protein, the predicted spin–spin distances were shorter than those for cytoplasmic open LacY. The distributions for 73–340 and 136–340 coincided with minor peaks in the DEER experiments with NPGlc. These peaks are indicative of cytoplasmic closure as they overlap with the spin–spin distance distribution from the simulation of periplasmic open LacY where the cytoplasmic gate is closed. Peaks for 73–340, 137–340 and 136–340 for Occ also overlap with peaks from experiments.

**Fig. 5.** Experimental and simulated spin–spin distance distributions. (a) Location of spin labels on residues (black spheres, Cα atoms) and experimentally measured distances (black lines) on the periplasmic face of LacY (periplasmic open conformation). (b) Spin–spin distances on the cytoplasmic face (cytoplasmic open conformation). (c) Comparison of experimental DEER spin–spin distributions [11] (gray histograms, “DEER data”) to computed distributions from equilibrium MD simulations of LacY in three different conformational states: periplasmic open (blue lines, “Popen”), cytoplasmic open (red lines, “Copen”) and occluded (magenta lines, “Occ”). DEER experiments were performed in the presence of the non-binding sugar NPGlc, which shifts the conformational equilibrium toward the cytoplasmic open state [11].
recorded in the presence of NPGal that coincide with peaks for the simulation of the periplasmic open transporter (Fig. S7b in the supplementary content).

On the periplasmic side, the distances closely matched the distances for cytoplasmic open LacY (Fig. 5c, red lines), which is not surprising given that the periplasmic gate is closed. The peak for 164–310 for \( \text{Occ} \) closely follows the distribution from the simulations of cytoplasmic open LacY where the periplasmic gate is closed. For residues 164–310, the experimental DEER peaks at 31 Å and 37 Å in presence of binding and non-binding sugar have been attributed to conformations with both gates closed [27]. Encouragingly, the two peaks intersect the spin–spin distribution calculated from \( \text{Occ} \).

Order parameters allow comparison with crystal structures of MFS proteins

In order to compare our model of fully occluded LacY to known occluded MFS transporter structures such as PepT\( \text{So} \) [32], XylE [30,31], EmrD [28], NarU [29] and PiPT [33], we propose a pair of geometric order parameters. Putative gates were identified by comparing the experimental structures of cytoplasmic open LacY and the models of periplasmic open LacY [17]. The key feature of the closed periplasmic gate is the tight packing of helices 1 and 7, which seals off the central cavity to the periplasm. A similarly close packing of helices 4 and 10 is the crucial element of the cytoplasmic gate. Although other helices [32] have also been implicated in the gating of MFS transporters, we have focused on helices 1 and 7 and on helices 4 and 10 to capture the essence of the mechanism. The order parameter for the periplasmic gate is defined as the distance between the pair of \( \alpha \)–\( \alpha \) atoms in helices 1 and 7 with the shortest inter-helical distance in the wild-type crystal structure of LacY: this is I32–N245. The order parameter for the cytoplasmic gate is defined in a similar way, except that we now determine the pair of residues on helices 4 and 10 that shows the smallest \( \alpha \)–\( \alpha \) distance in the homology model of periplasmic open LacY. The cytoplasmic order parameter is therefore defined as the distance between the \( \alpha \)–\( \alpha \) atoms of residues E126 and C333. This choice of order parameters is consistent with previous studies of gating (see the supplementary content).

When the periplasmic and cytoplasmic gates are closed, the transmembrane helices 1 and 7 as well as helices 4 and 10 are kinked, enabling them to pack tightly against each other, sealing off any pathways across the membrane. Sequence motifs that facilitate helix bending by disrupting the transmembrane helices (such as prolines) are conserved across the MFS family [17,63]. This observation suggests that key features of the gating mechanism could be shared between MFS transporters. By comparing the conformations of the gates in both the available crystal structures and our simulations, we tested the hypothesis that the overall conformational changes in the transport cycle are conserved across the MFS superfamily [17] (Table 2).

The order parameters not only tracked the state of the gates in the simulations but also consistently ordered all the available MFS transporter structures (Fig. 6a). For cytoplasmic and periplasmic open LacY, the order parameters agreed with both the pore profile and the water density analysis. The equilibrium simulation of the occluded structure, \( \text{Occ} \), sampled a region of order parameter space (Fig. 6a) intermediate between cytoplasmic and periplasmic open LacY. The cytoplasmic order parameter remained predominately \(<13\) Å while the periplasmic order parameter was \(<8\) Å. Both gates were shut, which is consistent with the calculations of the pore radius (Fig. 3b).

The order parameters intuitively classify all the available MFS transporter crystal structures (Fig. 6a) even though the pairwise sequence identify between members of the MFS is typically low, for example, no more than 12% between LacY and any of the other transporters studied here [64] up to a maximum of 46% for the closely related peptide transporters PepT\( \text{St} \) and GkPOT [65]. The cytoplasmic open crystal structures for LacY, GlpT, PepT\( \text{St} \) and GkPOT and the inward open structure of XylE mapped onto the region of order parameter space sampled by the simulations of cytoplasmic open LacY. The two virtually identical FucP crystal structures were close to the region of order parameter space explored by the simulations of periplasmic open LacY, while the recently solved periplasmic open structure of YajR was located slightly farther outside. The crystal structures of NarK and PepT\( \text{So} \) border on the region of order parameter space sampled by the simulations of cytoplasmic open LacY during which it partially closed, while NarU B is only slightly removed, all of which is consistent with the partially cytoplasmic closed structure of LacY constituting a transition intermediate [25]. Importantly, the occluded crystal

**Table 2.** Order parameters for MFS transporters.

| Transporter | Cytoplasmic gate distance (\( \alpha \)–\( \alpha \)) TM4–TM10 | Periplasmic gate distance (\( \alpha \)–\( \alpha \)) TM1–TM7 |
|-------------|-------------------------------------------------|-------------------------------------------------|
| EmrD        | Thr119 and Phe311                               | Ala32 and Cys229                               |
| FucP        | Thr143 and Ser370                               | Gin51 and Ile282                               |
| GlpT        | Trp138 and Ile368                               | Pro53 and Thr278                               |
| LacY        | Glu126 and Cys333                               | Ile32 and Asn245                               |
| NarU        | Met1149 and Phe367                              | Ser54 and Ala273                               |
| NarK        | Met1515 and Phe370                              | Ser56 and Ala275                               |
| PepT\( \text{So} \) | Ser131 and Leu427                           | Thr36 and Ser320                               |
| PepT\( \text{St} \) | Ser130 and Gly407                         | Ala34 and Ser303                               |
| PiPT        | Ala154 and Ala439                               | Asn53 and Leu332                               |
| POT         | Asn138 and Gly420                               | Val47 and Thr314                               |
| XylE        | Met1149 and Ser396                              | Ser22 and Tyr298                               |
| YajR        | Ser125–Ser328                                  | Thr37 and Pro236                               |
structures of PepTSo, EmrD, NarU (A and B) and NarK and cytoplasmic partially occluded structure of XyIE mapped to the intermediate region accessed by Occ, with the occluded structure of PIPT close by. Classifying the crystal structure of periplasmic partially occluded XyIE [30] using our order parameters puts it outside the region sampled by Occ. While the periplasmic order parameter for XyIE was similar to EmrD (~8.0 Å), the cytoplasmic order parameter for XyIE was very low, ~5.1 Å. This is much lower than the cytoplasmic order parameters seen in our simulations or any other MFS transporter structure and probably reflects the presence of the additional intracellular helices that pack close to the cytoplasmic gate and that are not present in the other MFS transporter structures.

In order to assess the robustness of our conclusions, we introduced a fairly drastic change into our equilibrium simulations by changing the underlying force field from GROMOS96 to CHARMM22. As discussed in the supplementary content, both sets of simulations behaved very similarly and, in particular, cover similar regions of order parameter space (Fig. S8b). Thus, the order parameter analysis and the simulations are robust against changes in the simulation parameters and are therefore likely to reflect upon a real property of the LacY protein, namely, the flexibility of its cytosolic and periplasmic gates.

Taken together, our analysis suggests that key features of the gating mechanism are conserved across the MFS transporters and a meaningful structural comparison is possible even in cases such as XyIE where additional structural elements interact with the gates. In particular, our model for the occluded conformation of LacY is consistent with the known structural evidence for occluded conformations in the MFS superfamily, as revealed by our order parameter analysis.

In a recent study, Madej et al. modeled the occluded state of LacY by homology to the crystal structure of partially occluded PepTSo [27]. Even though their model and ours were derived by completely different methods, they are generally quite similar with a Cα RMSD of 4.4 Å computed with PyMOL (Schrödinger, LLC); the agreement is higher for the C-terminal domain (3.2 Å) than for the N-terminal domain (4.8 Å). The similarity is also emphasized by the order parameters for the gates of the occluded homology model. The cytoplasmic and the periplasmic gates are closed in the homology model and the model...
maps onto the region of the two-dimensional order parameter plot (Fig. 6a) populated by occluded structures from our simulations, both from the equilibrium simulation of the occluded state, Occ, and from the simulation of the transition between cytoplasmic and periplasmic open LacY (DIMS-c.open-p.open) described in the next section.

**DIMS transitions between cytoplasmic and periplasmic open LacY**

In order to test the hypothesis that the conformational change proceeds via an intermediate occluded state, we generated an ensemble of 80 transitions between the end states. These DIMS MD simulations (DIMS-c.open-p.open) started with the cytoplasmic open wild-type crystal structure of LacY and used as the target structure the entire N- and C-terminal domains (residues 7–185 and 220–399) of the homology model of periplasmic open LacY. During the DIMS MD simulations, the structure of the transporter changed from a cytoplasmic open/periplasmic closed state to a cytoplasmic closed/periplasmic open conformation as shown by analysis of pore radius profiles (Fig. S11 in the supplementary content). These transitions connect the broad wells sampled by the equilibrium MD simulations of cytoplasmic and periplasmic open LacY in our order parameter space (Fig. 6b). Intermediate structures in the DIMS transitions between cytoplasmic and periplasmic open LacY were occluded according to both analysis of the pore radius (Fig. S11) and the order parameters (Fig. 6b). These structures occupy a region in order parameter space that is sampled by the Occ equilibrium MD simulation (Fig. 6a and b). Thus, two different DIMS MD simulations produced structures with both their gates impermeable to H⁺ and sugar; a full transition between cytoplasmic and periplasmic open LacY (DIMS-c.open-p.open) and a transition using partial structural information to close the cytoplasmic half of cytoplasmic open LacY (DIMS-closure) together with subsequent equilibrium MD.

**Coupling of the flexible gates**

The results of our simulations lie somewhere between a strict rocker-switch mechanism, where the gates open simultaneously [1,16], and a strict airlock mechanism [20], where the gates open simultaneously (Fig. 1b). We would expect a single, diagonal straight line to connect the cytoplasmic and periplasmic open states in the order parameter plot (Fig. 6) in the case of the former and an “L” shape for the latter. The simulated transitions between the cytoplasmic open and periplasmic open conformations using DIMS (DIMS-c.open-p.open) did not fall on either set of hypothetical lines. We were able, however, to close one gate independently of the other by only applying a bias to the open gate (DIMS-closure; Fig. 6b), which highlights the structural flexibility in LacY and is consistent with a gating motion more similar to an airlock mechanism. A simple structural comparison also argues against the rigid-body motion implied by a literal rocker-switch mechanism. Significant intra-domain arrangements (all with Cα RMSD > 2.9 Å) are apparent when comparing the N- and C-terminal domains of representative structures for the three key states—cytoplasmic open, periplasmic open and occluded (Fig. S12).

The conformational flexibility detected by previous studies also argues against a literal rocker-switch mechanism and is consistent with the results from our simulations. The conformational flexibility of LacY has been previously highlighted in hydrogen/deuteron exchange experiments [21–23] and in intra-domain cross-linking experiments [66,67], which argue for high intra-domain flexibility. Furthermore, equilibrium MD simulations of LacY have suggested that its individual helices are flexible [18,25] during the initial stages of the conformational switch. SGLD simulations [44] of LacY have shown intra-domain re-arrangements of the helices, as well as global motions during the transition itself, consistent with our conclusions.

In summary, our simulations provide multiple lines of evidence for the existence of a functionally occluded state in LacY, as hinted at by a range of experimental studies. A model for the intermediate occluded state was derived using molecular simulation that only utilized the crystal structure of the cytoplasmic open conformation and a homology model of the periplasmic conformation. It agrees with the available structural information on occluded states of MFS transporters and experimental DEER data. Residues known to be involved in the transport cycle occupy positions predicted in other studies. Calculation of geometric order parameters enabled comparison of all MFS transporter X-ray crystal structures, an approach that will become increasingly fruitful as more structures become available. The movements of flexible gates were seen to be coupled and thus lead to the formation of occluded transition intermediates. While global motions of the N- and C-terminal domains relative to one another do play a role, we rule out a strict “rocker-switch” mechanism involving rigid domain movements. We have shown that it is productive to describe LacY (and by extension, the other MFS transporters) as multi-gated pores [68] by (1) assessing the barriers to substrate permeation for different states in the transport cycle and (2) quantifying the changes in the gates by order parameters. A similar approach has already proven successful for the LeuT-like transporters [58,69,70] and could therefore act as a general paradigm to computationally analyze secondary transporter function.
Materials and Methods

Membrane protein insertion

The LacY wild-type crystal structure (PDB code 2V8N [14]) was downloaded from the Protein Data Bank. The homology model of Radestock et al. [17] for periplasmic open LacY (P0077183), which is based on the periplasmic open structure of FucP, was obtained from the protein model database [17,71]. These structures, as well as a model for a fully occluded structure of LacY obtained from DIMS simulations, were inserted into POPC bilayers by an established protocol [51,72,73]. In all structures, His322 was neutral as suggested by the proposed mechanism [2] and modeled as the imidazole N resonance except for simulations COpenB1 and COpenB2 in which the N2 nitrogens were added, resulting in a physiological free NaCl concentration of 100 mM.

Atomistic equilibrium simulations

Atomistic simulations were run in GROMACS 4.5 [74], either with the GROMOS96 43a1 [75] force field with Berger’s lipid parameters [76] (“GROMOS”) or with the CHARMM22 force field [77], including the CMAP correction [77,78] and CHARMM36 lipid parameters [79] (“CHARMM”). All atomistic simulations were performed at 315 K and 1 bar. The Parrinello–Rahman barostat was used for semi-isotropic pressure coupling [90]. Temperature coupling was achieved by velocity rescaling with a time constant of 0.1 ps [81]. The P-LINCS algorithm was used to constrain bonds involving hydrogen atoms [82]. The SPC [53] and the CHARMM TIP3P [83] water models were used for atomistic simulations with GROMOS and CHARMM, respectively. The radius for neighbor list searches was set to 10 Å and 12 Å, respectively, for simulations with GROMOS and CHARMM force fields. Long-range electrostatics were simulated with the particle mesh Ewald method [84], either with a real-space cutoff of 10 Å (GROMOS) or a switching function between 8 Å and 10 Å (CHARMM). Van der Waals forces were cut off at 10 Å (GROMOS) or switched off between 8 Å and 10 Å (CHARMM). Production runs were preceded by energy minimization and 5 ns equilibration using position restraints with a harmonic force constant of 1000 kJ/mol nm² on the protein heavy atoms.

Cluster analysis

The simulation of the occluded conformer of LacY was analyzed using the algorithm by Daura et al. [85] in order to pick relevant frames for further analysis. All protein atoms were considered in the geometric clustering and a distance cutoff values of 0.25 nm was employed. The structure used for electrostatics calculations—the final structure of the trajectory Occ—was part of the dominant cluster with a cutoff of 0.25 nm.

DIMS MD

DIMS MD [40,41] was used to sample transitions between cytoplasmic and periplasmic open LacY. Simulations were run with the CHARMM 36a1 program using the CHARMM22 all-atom force field [77,78]. A soft ratcheting bias \( \Delta \phi = 10^{-4} \) Å was used in DIMS-c-open, \( \Delta \phi = 10^{-5} \) Å was used in DIMS-c-periplasmic-open, and \( \Delta \phi = 10^{-6} \) Å was used in DIMS-closure. Langevin dynamics were run at 300 K with a timestep of 2 fs and a friction coefficient of 25.0 ps⁻¹. For efficiency, we modeled the protein environment with the GBSW implicit solvation model [46,47], which has previously been used to obtain DIMS transitions for the KcsA ion channel in a membrane environment [57]. The CMAP dihedral correction and atomic radii optimized for GBSW were used [87]. The GBSW model approximates the bilayer as an exclusion zone from the aqueous phase (\( \varepsilon = 80 \)) [46]; here, we employed a GBSW membrane thickness of \( h_{\text{memb}} = 40 \) Å and a smoothing length of \( w_{\text{memb}} = 6.5 \) Å (Fig. 2c) to approximate the POPC membrane geometry seen in the explicit solvent simulations (Fig. 2d). The dielectric changed smoothly from \( \varepsilon = 1 \) to \( \varepsilon = 80 \) in the head group region. The protein was positioned in the implicit membrane by structural alignment to a fully atomistic system generated by the membrane insertion protocol described above using MDAnalysis [90–92] and energy minimized. The half-smoothing length \( w_{\text{memb}} \) for the dielectric boundary was set to the default of 0.3 Å and the non-polar solvation contribution \( s_y \) was 0.04 kcal/(mol Å²). The default non-bonded options for GBSW simulations were employed, with an inner cutoff of 16 Å and a cutoff for the energy calculation of 16 Å. A cutoff for non-bonded list generation of 20 Å was used. The RMSD of the non-hydrogen backbone atoms for the N-terminal domain (residues 7–185) and the C-terminal domain (residues 220–399) to the target conformation was used as a progress variable in DIMS. The target structure was reoriented relative to the current structure every 10 MD steps. Non-hydrogen backbone atoms of the N- and C-terminal domains were used for rotational and translational fit. To close the cytoplasmic or periplasmic gate, we used only atoms with \( z < 0 \) Å or \( z > 0 \) Å (where \( z = 0 \) Å marks the midplane of the membrane) for the RMSD calculation assessing the change in progress variable that drives the soft ratcheting bias. Eighty simulations were performed with differing seeds for the random number generator that drives the stochastic term of the Langevin equation so that each simulation sampled a different microscopic pathway.

HOLE analysis

Centered and fitted trajectories were used for pore radii analysis of equilibrium MD trajectories. Conformations were written out every 0.1 ns and their radial pore profiles were calculated with the program HOLE [50]. NumPy [88] was used to histogram the pore profiles, which allowed the calculation of average pore profiles and standard
deviations, and figures were plotted with Matplotlib [89]. All frames from the 80 DIMS trajectories between cytoplasmic and periplasmic open LacY were analyzed by HOLE. For each structure, the backbone RMSD distance of the N-terminal domain (residues 7–185) and the C-terminal domain (residues 220–399) to the periplasmic open target was calculated with MDAnalysis [90–92]. The HOLE profiles and the RMSD distances were gridded and interpolated with Matplotlib to be shown as contour plots. Python code for the pore radius analysis is available as part of the HOLE analysis module of the MDAnalysis library [90].

Water density analysis

Water densities were calculated for centered and RMS-fitted trajectories using the program gridcount1 [52]. The densities were represented using the volume viewer in UCSF Chimera [54,55] and measured relative to the density of SPC bulk water at standard conditions, 0.970 g/cm³ [93].

DEER distance analysis

DEER is an EPR technique for measuring distances between two spin labels that have been covalently attached to a protein. Two cysteine residues are introduced into the protein and subsequently labeled. The positions are chosen to report on the expected conformational change. The study by Smirnova et al. used (1-oxyl-2,2,5,5-tetramethylpyrroline-3-methyl)-methanethiosulfonate (MTSL) spin labels [11]. MTSL has a linker with five rotatable bonds and is therefore very flexible. The distance distributions between the two spin labels measured by the experiment for LacY are typically broad and often multi-modal. The distributions are therefore a convolution of the flexibility of the MTSL spin label and the conformational spread of the proteins in the sample. To ensure that we compared like with like, we developed a method that (1) maps rotamer libraries of the MTSL spin label onto each position, (2) discards those rotamers that sterically clash with the protein (typically distances >2 Å) and (3) calculates all (weighted) distance pairs between the remaining rotamers and (4) thereby estimates a distance distribution for that structure. The code was written in Python using the MDAnalysis library [90] and incorporates a published rotamer library for MTSL [94]. It is available for download from the MDAnalysis website². Our approach improves upon an existing method [94] by increasing computational efficiency and implementing, via the MDAnalysis library [90], analysis of ensembles of hundreds of structures, which allowed us to estimate distance distributions for entire simulation trajectories. For spin–spin distance pairs 73–340, 136–340 and 137–340, a clash distance cutoff of 1.5 Å was used. The estimated distance–distance distributions were then compared to the previously reported DEER measurements of LacY [11].

Order parameter analysis

Two order parameters that report on whether a MFS transporter structure is cytoplasmic or periplasmic open were defined for EmrD (PDB code 2GFP) [28], FucP (3O7Q, 3O7P) [16], GltP (1PW4) [95], cytoplasmic open LacY (1PV6, 1PVT, 2CFQ, 2CFP, 2V8N, 2V5Y) [12–15] or periplasmic open LacY (homology model: PM0077183, repeat swapped model: PM0076824) [17], NarU (4IUB) [29], NarK (4JR9) [96], PepTSo (2XUT) [32], PepTSo (4APS) [97], Pipt (4J05) [33], Pot (4IKV) [65], XylE (4GBY, 4JA3, 4JA4) [30,31] and YajR (3WDO) [98] are summarized in Table 2. Distance matrices were calculated for Cα atoms of the pairs of transmembrane helices 1 and 7 for the closed periplasmic gates of LacY, GltP, EmrD, NarK, NarU, PepTSo, Pipt, Pot and XylE using MDAnalysis [90] (see Table S2 for definitions of the transmembrane helices). The closest contact defined the Cα–Cα distance that serves as the order parameter. Similarly, the cytoplasmic order parameter was calculated for the closed cytoplasmic gates of FucP, EmrD, NarU, PepTSo, Pipt, XylE and YajR, as well as the homology and the repeat swapped model for periplasmic open LacY. No structure featuring the closed cytoplasmic exists for GltP, NarK, PepTSo and Pot. Similarly, the structures of the closed periplasmic gates of FucP and YajR are unknown. The order parameters were defined via structural alignments. GltP, NarK, PepTSo and Pot were aligned with cytosolic open LacY by the DALI pairwise alignment server [99], which was also used to align YajR to the periplasmic open homology model of LacY. For FucP, the refined structural alignment with cytoplasmic open LacY obtained by Radestock et al. [17] that underlies the homology model for periplasmic open LacY was used.

Poisson-Boltzmann calculations

In order to assess the energetics of proton permeation through aqueous pathways in a semiquantitative manner, we computed the position-resolved electrostatic free energy \( W_{\text{elec}}(r) \) for a solvated proton on a 3D grid of ion positions \( r \). Proton transport in aqueous solution involves a complicated “fluxional complex” in which the Eigen \( (H_{2}O^{+}) \) and Zundel cation \( (H_{2}O_{2}^{+}) \) hydration structures of the proton rapidly interconvert [100]. In order to quantify the energetics of solvated protons inside a membrane protein, we adopted the much simpler Born description, which treats the proton as a cavity in a continuum solvent with an effective radius \( a \) and a charge of +1. The use of such a simple model is motivated by the insight that the energetics are likely dominated by the electrostatics of the charge in the low dielectric membrane compared to smaller differences due to the detailed proton–water interactions. The effective Born radius of the solvated proton was estimated as \( a = 1.4848 \ \text{Å} \) by solving the Born equation \( a = \sqrt{166 \ \text{Å}^{2}\text{Kcal mol}^{-1}} q^{2}/\Delta G_{\text{solv}}(1-1/80) \) with the experimental hydration free energy of the oxonium ion \( (H_{2}O^{+}) \) at the 1 M standard state, \( \Delta G_{\text{solv}} = -110.4 \pm 0.7 \ \text{Kcal/mol} \) [101].

Within the framework of Poisson-Boltzmann theory, \( W_{\text{elec}}(r) \) is the electrostatic free energy of the ion bound to the protein compared to the ion and protein in bulk solution separately:

\[
\Delta W_{\text{elec}}(r) = G_{\text{protein + ion}}(r) - G_{\text{protein}} - G_{\text{ion}}
\]

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1 available from https://github.com/orbeckst/gridcount
2 http://mdanalysis.googlecode.com
In the case of membrane proteins, the reference energy $G_{\text{protein}}$ is calculated for the protein embedded in a membrane-like environment as described below. Such an approach has been used previously to calculate the electrostatic free-energy profile (“Born profile”) for ions in channels and pores along the permeation pathway [45]. Here this concept is extended to generate the 3D free-energy landscape of an ion in the heterogeneous environment of the LacY transporter. The linearized Poisson-Boltzmann equation was solved with APBS [102] at an ionic strength of 100 mM NaCl and a temperature of 298.15 K for ion positions $r$ generated with a customized version of HOLLOW [103] at a point spacing of 1 Å. A dielectric constant of $\varepsilon = 80$ was assigned to bulk water and a dielectric constant of $\varepsilon = 10$ was assigned to the protein. An implicit membrane was modeled after the membrane observed in the MD simulations as a low dielectric slab with a hydrophobic core of thickness 33.5 Å with $\varepsilon = 2$ and a head group region of thickness 6.5 Å with $\varepsilon = 20$ (Fig. 2c). The water-accessible interior of the transporter was modeled as a high dielectric ($\varepsilon = 80$). Manipulation of the dielectric maps was carried out with software based on the draw_membrane tool in APBSmem [104] and an updated version of the BornProfiler analysis package [45]. The calculations employed a manual focusing strategy in three steps on a computational grid of $129 \times 129 \times 129$, starting from a cubic bounding box of length 250 Å, via a 100-Å box, to a final dimension of 50 Å, centered on the position of the ion. Increasing the computational grid to $161 \times 161 \times 161$ did not change the energies and hence the results can be considered converged with respect to the grid. Starting structures for simulations $C\text{Open}A1$ and $P\text{Open}$—which had been equilibrated in atomistic membranes and are fully cytoplasmic and periplasmic open respectively—as well as final structures from MD simulations of $C\text{Open}A1$, which showed partial closure, and fully closed $O\text{cc}$ were used for Poisson-Boltzmann calculations. Partial charges and radii from the CHARMM force field were assigned with pdb2pqr [105]. All ionizable residues were simulated in their default charge state as either suggested by PROPKA [106] or because the default protonation state better described a transporter in that part of the transport cycle that is believed to occur without involvement of substrate or proton transport [2].

Accession number

A representative structure of the occluded conformation (the last frame of the simulation $O\text{cc}$) has been deposited in the protein model database [71] under accession code PM0079313 and is also available in the supplementary content.

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Appendix A. Supplementary Data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jmb.2013.10.024.

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Abbreviations used:

DEER, double electron–electron spin resonance; DIMS, dynamic importance sampling; MD, molecular dynamics; MFS, major facilitator superfamily; POPC, 1-palmitoyl,2-oleoyl-sn-glycero-3-phosphocholine; SGLD, self-guided Langevin dynamics; 3D, three-dimensional.

References

[1] Forrest LR, Krämer R, Ziegler C. The structural basis of secondary active transport mechanisms. Biochim Biophys Acta 2011;1807:167–88.
[2] Guan L, Kaback HR. Lessons from lactose permease. Annu Rev Biophys Biomol Struct 2006;35:67–91.
[3] Gadsby DC. Ion channels versus ion pumps: the principal difference, in principle. Nat Rev Mol Cell Biol 2009;10:344–52.
[4] Jardetzky O. Simple allosteric model for membrane pumps. Nature 1966;211:969–70.
Occluded intermediate conformations of LacY

[5] Mitchell P, editor. Translocations through natural membranes. Adv Enzymol Related Areas Mol Biol. John Wiley & Sons, Inc.; 1967 [Edited by Nord, F. F.].

[6] Kaback HR, Smirnova I, Kasho V, Nie Y, Zhou Y. The alternating access transport mechanism in LacY. J Membr Biol 2011;239:85–93.

[7] Kaback HR, Dunten R, Frillingos S, Venkatesan P, Kwaw I, Zhang W, et al. Site-directed alkylation and the alternating access model for LacY. Proc Natl Acad Sci U S A 2007;104:491–4.

[8] Zhou Y, Guan L, Freites JA, Kaback HR. Opening and closing of the periplasmic gate in lactose permease. Proc Natl Acad Sci U S A 2008;105:3774–8.

[9] Smirnova I, Kasho V, Sugihara J, Kaback HR. Probing of the osmosensor ProP. Biochim Biophys Acta Biomembr 2011;1808:1108–15.

[10] Kaback HR, Smirnova I, Kasho V, Nie Y, Zhou Y. The alternating access transport mechanism in LacY. Proc Natl Acad Sci U S A 2009;106:21561–6.

[11] Madej MG, Soro SN, Kaback HR. Apo-intermediate in the transport cycle of lactose permease (LacY). Proc Natl Acad Sci U S A 2012;109:E2970–8.

[12] Hofyoake J, Sansom MSP. Conformational change in an MFS protein: MD simulations of LacY. Structure 2007;15:873–84.

[13] Mchaurab HS, Steed PR, Kazmier K. Toward the fourth dimension of membrane protein structure: insight into dynamics from spin-labeling EPR spectroscopy. Structure 2011;19:1549–61.

[14] Madej MG, Soro SN, Kaback HR. Apo-intermediate in the transport cycle of lactose permease (LacY). Proc Natl Acad Sci U S A 2012;109:E2970–8.

[15] Yin Y, He X, Szewczyk P, Nguyen T, Chang G. Structure of the multidrug transporter EmrD from Escherichia coli. Science 2006;312:741–4.

[16] Yan H, Huang W, Yan C, Gong X, Jiang S, Zhao Y, et al. Structure and mechanism of a nitrate transporter. Cell Rep 2013;7:617–26.

[17] Sun L, Zeng X, Yan C, Sun X, Gong X, Rao Y, et al. Crystal structure of a bacterial homologue of glucose transporters GLUT1–4. Nature 2012;490:361–6.

[18] Quistgaard EM, Löw C, Moberg P, Träsaugues L, Nordlund P. Structural basis for substrate transport in the GLUT-homology family of monosaccharide transporters. Nat Struct Mol Biol 2013;20:766–8.

[19] Newstead S, Drew D, Cameron AD, Postis VLG, Xia X, Fowler PW, et al. Crystal structure of a prokaryotic homologue of the mammalian oligopeptide-proton symporters, PepT1 and PepT2. EMBO J 2010;30:417–26.

[20] Pedersen BP, Kumar H, Waight AB, Risenmay AJ, Roe-Zurr Z, Chau BH, et al. Crystal structure of a eukaryotic phosphate transporter. Nature 2013;496:533–6.

[21] Hirai T, Subramaniam S. Structure and transport mechanism of the bacterial oxalate transporter OxIT. Biophys J 2004;87:3600–7.

[22] Stansfeld PJ, Sansom MSP. Molecular simulation approaches to membrane proteins. Structure 2011;19:1562–72.

[23] Enkavi G, Tajkhorshed E. Simulation of spontaneous substrate binding revealing the binding pathway and mechanism and initial conformational response of OipT. Biophys Chem 2010;149:1105–14.

[24] Villanen P, Garcia ML, Kaback HR. Purified reconstituted lac carrier protein from Escherichia coli is fully functional. Proc Natl Acad Sci U S A 1984;81:1629–33.

[25] Chang S, Li K-S, Hu J-P, Jiao X, Tian X-H. Allosteric and transport behavior analyses of a fucose transporter with network models. Soft Matter 2011;7:4661.

[26] Fowler PW, Sansom MSP. The pore of voltage-gated ion channels becomes strained when closed. Nat Commun 2013;4:1872.

[27] Perilla JR, Beckstein O, Denning EJ, Wolff TB. Computing ensembles of transitions from stable states: dynamic importance sampling. J Comput Chem 2011;32:196–209.

[28] Wolff TB. Path corrected functionals of stochastic trajectories: towards relative free energy and reaction coordinate calculations. Chem Phys Lett 1998;365:411–7.

[29] Beckstein O, Denning EJ, Perilla JR, Wolff TB. Zipping and unzipping of adenylate kinase: atomistic insights into the ensemble of open ↔ closed transitions. J Mol Biol 2009;394:160–76.

[30] Wu X, Brooks BR. Self-guided Langevin dynamics simulation method. Chem Phys Lett 2003;381:512–8.
[44] Pendse PY, Brooks BR, Klauda JB. Probing the periplasmic-open state of lactose permease in response to sugar binding and proton translocation. J Mol Biol 2010;404:506–21.

[45] Beckstein O, Tai K, Sansom MSP. Not ions alone: barriers to ion permeation in nanopores and channels. J Am Chem Soc 2004;126:14694–5.

[46] Im W, Feig M, Brooks CL. An implicit membrane generalized bond theory for the study of structure, stability, and interactions of membrane proteins. Biophys J 2003;85:2900–18.

[47] Im W, Lee MS, Brooks CL. Generalized Born model with a simple smoothing function. J Comput Chem 2003;24:1691–702.

[48] Feig M, Im W, Brooks CL. Implicit solvation based on generalized Born theory in different dielectric environments. J Chem Phys 2004;120:903–11.

[49] Humphrey W, Dalke A, Schulten K. VMD: visual molecular dynamics. J Mol Graphics 1996;14:33–8.

[50] Smart OS, Goodfellow MJ, Wallace BA. The pore dimensions of gramicanid A. Biophys J 1993;65:2455–60.

[51] Stansfeld PJ, Sansom MSP. From coarse grained to atomistic: a serial multiscale approach to membrane protein simulations. J Chem Theory Comput 2011;7:1157–66.

[52] Beckstein O, Sansom MSP. Liquid-vapor oscillations of water in hydrophobic nanopores. Proc Natl Acad Sci U S A 2003;100:7063–8.

[53] Hermans J, Berendsen HJC, van Gunsteren WF, Postma JPM. A consistent empirical potential for water–protein interactions. Biopolymers 1984;23:1513–8.

[54] Petterson EF, Goddard TD, Huang CC, Couch GS, Greenblatt DM, Menc EC, et al. UCSF Chimera—a visualization system for exploratory research and analysis. J Comput Chem 2004;25:1605–12.

[55] Goddard TD, Huang CC, Ferrin TE. Visualizing density maps with UCSF Chimera. J Struct Biol 2007;157:281–7.

[56] Yin Y, Jensen MO, Tajkhorshid E, Schulten K. Sugar binding and protein conformational changes in lactose permease. Biophys J 2006;91:3972–85.

[57] Denning EJ, Woolf TB. Cooperative nature of gating transitions in K(+) channels as seen from dynamic importance sampling calculations. Proteins 2010;78:1105–19.

[58] Shimamura T, Weyand S, Beckstein O, Rutherford NG, Hadden JM, Sharples D, et al. Molecular basis of alternating access membrane transport by the Rhodobacter sphaeroides transhydrogenase transporter Mhp1. Science 2010;328:470–3.

[59] Baker J, Wright SH, Tama F. Simulations of substrate transport in the multidrug transporter EmrD. Proteins 2012;80:1620–32.

[60] Dahl ACE, Chavent M, Sansom MSP. Bendix: intuitive helix geometry analysis and abstraction. Bioinformatics 2012;28:2193–4.

[61] Jung K, Jung H, Wu J, Privé GG, Kaback HR. Use of site-directed fluorescence labeling to study proximity relationships in the lactose permease of Escherichia coli. Biochem Biophys Acta 1993;32:12273–8.

[62] Sahin-Toth M, Kaback HR. Arg-302 facilitates deprotonation of Glu-325 in the transport mechanism of the lactose permease from Escherichia coli. Proc Natl Acad Sci U S A 2001;98:6068–73.

[63] Vardy E, Steiner-Mordoch S, Schuldiner S. Characterization of bacterial drug antiporters homologous to mammalian neurotransmitter transporters. J Bacteriol 2005;187:7518–25.

[64] Prlic A, Bliven S, Rose PW, Bluhm WF, Bizon C, Godzik A, et al. Pre-calculated protein structure alignments at the RCSB PDB website. Bioinformatics 2010;26:2983–5.

[65] Doki S, Kato HE, Solcan N, Iwaki M, Koyama M, Hattori M, et al. Structural basis for dynamic mechanism of proton-coupled symport by the peptide transporter POT. Proc Natl Acad Sci 2013;110:11343–8.

[66] Zhang W, Guan L, Kaback HR. Helices VII and X in the lactose permease of Escherichia coli: proximity and ligand-induced distance changes. J Mol Biol 2002;315:53–62.

[67] Zhou Y, Madej MG, Guan L, Nie Y, Kaback HR. An early event in the transport mechanism of LacY protein: interaction between helices V and I. J Biol Chem 2011;286:30415–22.

[68] Läuger P. Kinetic properties of ion carriers and channels. J Membr Biol 1980;57:163–78.

[69] Abramson J, Wright EM. Structure and function of Na⁺-symporters with inverted repeats. Curr Opin Struct Biol 2009;19:425–32.

[70] Krishnamurthy H, Piscitelli CL, Gouaux E. Unlocking the molecular secrets of sodium-coupled transporters. Nature 2009;459:347–55.

[71] Castrignanò T, De Meo PDO, Cozzetto D, Talamo IG, Tramontano A. The PMDB Protein Model Database. Nucleic Acids Res 2006;34:D306–9.

[72] Bond PJ, Sansom MSP. Insertion and assembly of membrane proteins via simulation. J Am Chem Soc 2006;128:2697–704.

[73] Scott KA, Bond PJ, Ivetac A, Chetwynd AP, Khalid S, Sansom MSP. Coarse-grained MD simulations of membrane protein-bilayer self-assembly. Structure 2006;16:621–30.

[74] Hess B, Kutzner C, van der Spoel D, Lindahl E. GROMACS 4: algorithms for highly efficient, load-balanced, and scalable molecular simulation. J Chem Theory Comput 2008;4:435–47.

[75] van Gunsteren WF, Billette SR, Eising AA, Hünenberger PH, Krüger P, Mark AE, et al. Biomolecular simulation: the GROMOS96 manual and user guide, VdW Hochschulverlag AG an der ETH Zürich. Switzerland: Zürich; 1996.

[76] Berger O, Edholm O, Jähning F. Molecular dynamics simulations of a fluid bilayer of dipalmitoylphosphatidylcholine at full hydration, constant pressure, and constant temperature. Biophys J 1997;72:2002–13.

[77] Mackerell AD, Bashford D, Dunbrack RL, Evanseck JD, Field MJ, Fischer S, et al. All-atom empirical potential for molecular modeling and dynamics studies of proteins. J Phys Chem B 1998;102:3586–616.

[78] Mackerell AD, Feig M, Brooks CL. Extending the treatment of backbone energetics in protein force fields: limitations of gas-phase quantum mechanics in reproducing protein conformational distributions in molecular dynamics simulations. J Comput Chem 2004;25:1400–15.

[79] Klauda JB, Venable RM, Freites JA, O’Connor JW, Tobias DJ, Mondragon-Ramirez C, et al. Update of the CHARMM all-atom additive force field for lipids: validation on six lipid mixtures. J Phys Chem B 2010;114:7830–43.

[80] Parrinello M, Rahman A. Polymeric transitions in single crystals: a new molecular dynamics method. J Appl Phys 1981;52:7182–90.

[81] Bussi G, Donadio D, Parrinello M. Canonical sampling through velocity rescaling. J Chem Phys 2007;126:14101.

[82] Hess B. P-LINCS: a parallel linear constraint solver for molecular simulation. J Chem Theory Comput 2008;4:116–22.

[83] Jorgensen WL, Chandrasekhar J, Madura JD, Impey RW, Klein ML. Comparison of simple potential functions for simulating liquid water. J Chem Phys 1983;79:926–35.
Occluded intermediate conformations of LacY