High-speed atomic force microscopy reveals a three-state elevator mechanism in the citrate transporter CitS

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The secondary active transporter CitS shuttles citrate across the cytoplasmic membrane of gram-negative bacteria by coupling substrate translocation to the transport of two Na\(^+\) ions. Static crystal structures suggest an elevator type of transport mechanism with two states: up and down. However, no dynamic measurements have been performed to substantiate this assumption. Here, we use high-speed atomic force microscopy for real-time visualization of the transport cycle at the level of single transporters. Unexpectedly, instead of a bimodal height distribution for the up and down states, the experiments reveal movements between three distinguishable states, with protrusions of \(~1.0\) nm, \(~1.6\) nm, and \(~35\) nm above the membrane, respectively. Furthermore, the real-time measurements show that the individual protomers of the CitS dimer move up and down independently. A three-state elevator model of independently operating protomers resembles the mechanism proposed for the aspartate transporter GltPh. Since CitS and GltPh are structurally unrelated, we conclude that the three-state elevators have evolved independently.

Significance

As cellular membranes are impermeable to most molecules, transporter proteins are typically present in the membrane to transport small molecules in or out of the cell. Due to the small, nanometer size of these transporters, it is challenging to study their transport mechanism. Here, we use advanced microscopy approaches to study in real time and at the single-molecule level the mode of action of the dimeric CitS transporter. Using high-speed atomic force microscopy, we visualize the dynamic, elevator-like movement of the transporter, and we reveal that the two protomers move independently. We also discovered an intermediate state, reminiscent of another, unrelated transporter, indicating that independent evolutionary pathways have led to similar three-state elevator mechanisms.

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The authors declare no competing interest.

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and dynamic experiments have not been reported to test this assumption. In this respect, high-speed atomic force microscopy (HS-AFM) (12–15) offers the advantage of probing membrane proteins in liquid in their lipidic environment with high spatial and temporal resolution (16, 17). Uptake experiments performed on CitS report an apparent turnover rate of roughly once every second at saturating conditions (SI Appendix, Table S1) (18). This rate is within the scanning capabilities of HS-AFM, and here, we report HS-AFM experiments directly visualizing the dynamics of citrate transport, at the single-molecule level, to unveil the molecular mechanism behind transport.

**Results**

**Binding Order of Citrate and Sodium Ions to CitS.** Although it is well established that citrate transport mediated by CitS is strictly coupled to the cotransport of two sodium ions (19), the order in which substrate and coupling ions bind is unknown. For a proper interpretation of our HS-AFM data, such information is essential. Therefore, we set out to deduce the binding order from measurements of the initial rates of uptake of radio-labeled citrate into proteoliposomes reconstituted with purified CitS. We determined the rates as a function of a wide range of citrate and Na⁺ concentrations (Fig. 1A and SI Appendix, Table S1). These experiments provide a rich source of data on the kinetic mechanism, including the $K_M$ values for citrate at different fixed Na⁺ concentrations ($K_M^{C–Ci}$). In Fig. 1B, a double logarithmic plot is presented of these apparent affinity constants as function of the concentration of Na⁺. In such plots, linear relations are approached asymptotically in both the high- and low-Na⁺ concentration extremes, with slopes of either zero or negative integers (20–22). The negative value of the slopes in the low- and high-concentration regime indicate the number of Na⁺ ions that must bind before and after the citrate-binding event, respectively. For the discussion below (Discussion), the information on the number of sodium ions binding before citrate is most relevant. In Fig. 1B, the slope at low-Na⁺ concentrations approaches −1, and therefore, we conclude that citrate-binding is preceded by an ordered step involving one Na⁺ (20, 22).

**Imaging the CitS Transporter by HS-AFM.** To follow the conformational dynamics of CitS transporters by HS-AFM, we purified the dimeric protein and reconstituted it at high density in liposomes consisting of lipids extracted from _Escherichia coli_, the same lipid mixture in which CitS shows robust transport activity (Fig. 1). Upon adsorption onto a mica surface, the proteoliposomes formed a planar membrane bilayer with either the cytoplasmic side or the periplasmic side of the transporter facing up. The maximum protrusion of the transport domain when observed from the cytoplasmic side is expected to be <1.0 nm, while from the periplasmic side, this is >1.5 nm (23). Therefore, it is well distinguished, and we limited our observation to protein dynamics at the periplasmic side. Citrate transport by CitS is strictly coupled to the cotransport of Na⁺ ions (11). In order to understand the role of both Na⁺ and citrate in the transporting mechanism, we studied the membrane-embedded protein 1) in presence of saturated concentration of both Na⁺ (300 mM) and citrate (150 μM) (SI Appendix, Table S1), 2) in presence of Na⁺ but absence of citrate, and 3) in absence of both Na⁺ and citrate. A typical surface topography image of CitS in presence of both Na⁺ and citrate is depicted in Fig. 2A. Protrusions with a height of about 1.6 nm and an intradimer, peak-to-peak distance of about 6.8 nm were found. Comparison between the CitS features measured in presence of Na⁺ and citrate with the crystal structure of the protein (23) revealed a similar height and spacing of the protrusions, indicating that they correspond to the transport domains (Fig. 2B). In contrast, when the protein was observed in absence of both Na⁺ and citrate, the HS-AFM recording showed an increment in the intra-dimer spacing as can be seen in Fig. 2C–E. Interestingly, while in presence of Na⁺, almost no diffusivity of the dimers was observed, they became highly diffusive in the absence of Na⁺ (Movie S1). Sodium-free conditions are nonphysiological, but this change in behavior indicates that Na⁺ is playing a distinctive role in maintaining the stability of CitS in the membrane.

**Role of Na⁺ and Citrate in the Elevator Mechanism of CitS Transporting Domain.** Next, we focused on the axial movement of the transport domains. Based on the crystal structures, it was proposed that these domains have two distinguishable conformational states: outward-facing (elevated) and inward-facing (down) (SI Appendix, Fig. S1) (24). To record the appearance and disappearance of the protrusions, which correspond to movements of the transport domains between the elevated and down states, we recorded HS-AFM movies at 2.5 frames per second in the presence of Na⁺ and citrate. Snapshots of a recording are depicted in Fig. 3A and indeed reveal alternation between outward- and inward-facing states. Kymograph representations of eight distinguishable individual domains from

![Fig. 1.](https://doi.org/10.1073/pnas.2113927119) Kinetic studies of CitS. (A) Citrate transport rates reported in SI Appendix, Table S1 plotted as a function of citrate concentration. The lines represent the fits of the Michaelis–Menten equation to the data for uptake at Na⁺ concentrations of 0.5 mM (black), 1 mM (red), 2.5 mM (blue), 5 mM (cyan), 25 mM (magenta), 75 mM (yellow), 125 mM (dark blue), and 200 mM (brown). (B) Dependence of the apparent affinities ($K_a$) for citrate obtained from the fitted curves in A, plotted against the Na⁺ concentration in which they were obtained in a double logarithmic scale. The lines represent linear fits at low- and high-[Na⁺] regimes. The values obtained from the linear regression are $-0.73$ and $-0.04$ for the low- and high-[Na⁺] regime, respectively. The values will tend to an integer number at infinitely low and high [Na⁺]; therefore, we approximated such value to the closest integer number. The data points are the average of triplicate measurements, and the error bars represent the SEM.
four dimers (Fig. 3 B and C) show the transitions between the states over time.

For a more-quantitative analysis of the protruded height of the transport domains, we measured the height of the protrusion above the membrane frame by frame (SI Appendix, Figs. S2 and S3 and Methods). Fig. 4 A–C shows example traces of these measurements along with histograms of the height measurements of all analyzed CitS molecules. In the absence of Na\(^{+}\) and citrate (nonphysiological condition), the transporter exhibits a height of 1.0 ± 0.2 nm and does not show significant movement in the axial direction (Fig. 4A). Na\(^{+}\) alone is sufficient to trigger the elevator movement of the transport domain in CitS (Fig. 4B). Unexpectedly, instead of showing a bimodal height distribution (for up and down states) as deduced from the crystal structures (11, 23, 24), three distinguishable distributions at ∼0.5 nm, ∼1.0 nm, and ∼1.6 nm were observed in the presence of Na\(^{+}\) (Fig. 4C). After the addition of citrate, similar peaks as for the Na\(^{+}\)-alone situation were observed, but the distribution over the peaks was different. In order to characterize the data further, each data point was assigned to one of the three observed states: elevated, intermediate, or down (Fig. 4 A–C, Left). Fig. 4D and Movie S3 show AFM data of these three distinct states. The probability of finding the domains in one of these states is shown in Fig. 4E. The elevated state has the highest probability in both Na\(^{+}\)-containing conditions; however, the probability of finding the domain in the intermediate state is relatively higher when citrate is present. Interestingly, the domain has similar probability to be found in the down state both in absence and presence of citrate. This observation suggests that even when the absence of citrate does not allow the transporter to go through the entire alternating-access transport cycle, Na\(^{+}\) alone is sufficient to promote the conformational switch in the elevator mechanism.

Next, we looked into the potential cooperativity in movement of the two protomers of the CitS dimer (SI Appendix, Fig. S4). Comparing the idealized traces of individual protomers suggests that the protomers act independently (SI Appendix, Figs. S4 and S5A and Fig. 4 B and C). However, it is often observed that one of the protomers is more active than the other within the period of recording (SI Appendix, Fig. S4). We estimated that a fast-protomer shows four times more jumps to the down state than the slower protomer (SI Appendix, Fig. S4B). Quantifying the relative frequencies of all six conformation combinations of all measured dimers (SI Appendix, Fig. S5B) and the position of one protomer with respect to the other (SI Appendix, Fig. S5C), no clear signs of cooperativity are observed. It can be concluded that the protomers move up and down independently of each other. Therefore, we can analyze their dynamics separately. To scrutinize these protomer dynamics further, the relative transition frequencies from one state to another were determined (Fig. 4F). While the transporter shows elevator movements in both the presence and absence of citrate (with Na\(^{+}\) present in both cases), the six possible state transitions revealed major differences in the two conditions. In particular, in absence of citrate, the most-frequent transition is between the elevated and down states. In contrast, in presence of citrate transitions between elevated and intermediate states are the most-frequently occurring ones. The transition between intermediate and down states exhibits relatively low frequencies for both conditions. Our observations reveal that while Na\(^{+}\) is playing a pivotal role in switching the transporting domain from the elevated to down state and vice versa, the presence of citrate makes this domain to fluctuate more in between elevated and intermediate states.

**Dynamics of Transporting Domain in CitS Transporter.** In order to validate our interpretation of the existence of three distinguishable states we used an infinite Hidden Markov Model (25) (iHMM; Methods), which allows analyzing the experimental time traces without any constraints in the number of states. A small majority of the HS-AFM height-time traces analyzed by the iHMM successfully yielded three states (SI Appendix, Fig. S6A), but for certain traces, the least-populated state could not be recognized by the iHMM (SI Appendix, Fig. S6B). These concerned the down state in presence of Na\(^{+}\) and citrate and the intermediate state in presence of Na\(^{+}\) alone. The analysis of such traces, showing a very limited number of visits to the third state, yielded two-state trajectories instead (SI Appendix, Fig. S6B). A comparison of tracking three states using iHMM and idealized traces has been depicted in SI Appendix, Fig. S7 for both Na\(^{+}\)-alone as well as Na\(^{+}\) and citrate conditions. Next, we have determined the dwell-time distribution for individual states in presence and
absence of citrate (both containing Na\textsuperscript{+}; SI Appendix, Fig. S8) by using idealized traces. The dwell-time distributions were well represented by a single exponential function, as expected for a Markovian system, and yielded the stochastic rate constants for leaving a specific state. This rate appears to be similar in absence and presence of citrate except for the intermediate states. To relate the dynamics of the transport domain with its function, we determined the rate constants for transitions between states by calculating the transition probability matrix of the idealized traces (Methods). Rate constants were also obtained from the iHMM analysis, and the comparisons are depicted in SI Appendix, Table S2 (in presence of Na\textsuperscript{+} only) and SI Appendix, Table S3 (in presence of Na\textsuperscript{+} and citrate). A similar trend for both approaches is found. Using this data, we evaluated the free energy differences between the corresponding states (Methods and SI Appendix, Fig. S9). In absence of citrate, there is 1.6 k\textsubscript{B}T of free energy difference for a transition from the elevated to down state ($\Delta G_{\text{ele-down}}^{0}$); 2.5 k\textsubscript{B}T of free energy difference for a transition from the elevated to intermediate state ($\Delta G_{\text{ele-inter}}^{0}$); and $-0.3$ k\textsubscript{B}T of free energy difference for a transition from the intermediate to down state ($\Delta G_{\text{inter-down}}^{0}$). When citrate is present, the estimated free energy differences are 1.4 k\textsubscript{B}T, $-0.2$ k\textsubscript{B}T, and 1.3 k\textsubscript{B}T, respectively (SI Appendix, Fig. S9). This reveals that independent of the presence or absence of citrate, a similar value of free energy difference between elevated and down state is observed, while for transitions involving the intermediate state, significant differences arise.

**Discussion**

Using HS-AFM, we studied the functional mechanism of CitS transporters, and our real-time imaging approach revealed that the proposed elevator mechanism (6, 11, 23) is real. Surprisingly, while the crystal structures suggested a two-state elevator mechanism, it turns out that the dynamic CitS elevator mechanism goes through three discrete states, termed here elevated, intermediate, and down. The three states are distinguished solely by the observed elevation of the transport domains in the AFM measurements and do not directly relate to their functional states (apo, bound, nonoccluded, occluded, etc.). Indeed, multiple functional states could be associated with the same elevation.
Unexpectedly, this three-state elevator mechanism resembles what has been found for the aspartate transporters GltPh and GltTk, in which crystal or cryogenic electron microscopy (cryo-EM) structures have resolved the three positions of the elevator (10, 26, 27). In the case of GltPh, HS-AFM studies showed that the three states were indeed visited under near-physiological conditions (28). Also, the approximate seconds time scale of the movements up and down in CitS and the lack of cooperativity in the elevator-like movements of the individual protomers in the homodimeric CitS assembly resemble the functional properties of the homotrimeric aspartate transporter GltPh and GltTk (10, 16, 29). These similarities are surprising, because the aspartate transporters are not related in sequence and structural fold to CitS. Therefore, the use of three-state elevator
mechanisms, lack of cooperativity, and seconds time scale of the movements appear to have evolved in parallel. The observation that two structurally unrelated proteins operate using a three-state mechanism may suggest that there is a constraint that precludes a two-state mechanism. One speculative possibility is that it is more difficult to prevent ion leaks in a two-state mechanism, but more work is needed to substantiate such speculation. A number of other transporters also use elevator-type mechanisms, and it is worth testing in future whether they also visit more structural states than currently elucidated at high resolution (30–32). Finally, increased temporal resolution using line-scan HS-AFM could improve the kinetic analysis (28), including a better view on the energy landscape of the different states.

The elevator movements of CitS were observed in the presence of Na\(^+\) and in the presence of both Na\(^+\) and citrate, while in absence of both ligands, the dimeric protein adopts a conformation in which the transport domains of both protomers are immobile and are separated further than in the conditions that show dynamic behavior. Complete absence of Na\(^+\) is nonphysiological, and therefore, we do not take the observed immobility into account for the proposed mechanism of transport (see below). We speculate that the structural integrity of CitS is compromised in the absence of Na\(^+\), resulting not only in wider separation of the protomers but also in increased diffusion through the membrane.

In the conditions in which dynamic elevator movements are observed, the elevated position at 1.7 nm is dominant in presence of Na\(^+\), while in the presence of Na\(^+\) and citrate, the intermediate position at about 1 nm exhibits similar likelihood as the elevated state. Fig. 5 shows the energy landscape of the transporter in absence and presence of citrate, visualizing the free energy differences between the three states. Next to the citrate-dependent difference in rates from and to the intermediate state, it can also be seen that the presence of citrate slightly flattens the energy landscape, easing transitions. In the presence of citrate and Na\(^+\), the transporter visits the three elevator states. It is likely that the carrier in the elevated state is fully loaded, because of the saturating concentrations of substrate and coion. A fully loaded elevated state is also consistent with the observations from the crystal structure in which both Na\(^+\) and citrate are bound to the outward-facing transport domain (11). In physiological conditions, it is expected that the binding of ligands from the environment occurs in this state (11, 23). The down state likely corresponds to the inward state observed in the crystal structure. In physiological conditions, citrate and Na\(^+\) are released into the cytoplasm from this state. However, because, in both HS-AFM and crystallization condition, the concentrations of substrate and coion were saturating and no membrane gradient was present, the down state must represent the inward-loaded state as observed in the crystal structures (11, 23). Movements between the up and down states in the presence of citrate and Na\(^+\) occur directly or via the intermediate state. The possibility to bypass the intermediate state when moving between the down and elevated states resembles what was observed for the aspartate transporter GltPh (16). It is not clear whether binding and unbinding of transported cargo can also take place in the intermediate state or that the state takes an occluded conformation. Further structural characterization is needed to answer this question.

In the presence of Na\(^+\) but in the absence of citrate, CitS can bind at least one sodium ion (Fig. 1), as determined in the kinetic analysis. With only Na\(^+\) bound, cycling between inward- and outward-exposed conformations would lead to uncoupled leakage of Na\(^+\), and therefore, these transitions should not occur. Surprisingly, the HS-AFM data revealed that movements between elevated and down states take place in the presence of only Na\(^+\). There are two possible explanations for this observation. The first is that full movement between elevated and down states can take place with sodium ion(s) bound but that the sodium ion(s) cannot be released once the opposite side is reached. Although theoretically possible, this explanation is not consistent with the known crystal structures that revealed exposed binding sites in the outward and inward states (11, 23). The alternative explanation is that the sodium ion concentration is not fully saturating in the absence of citrate. This is a well-known phenomenon in other sodium-coupled transporters such as GltPh (10), in which sodium ions bind weakly and cooperatively to aspartate increases the apparent affinity for Na\(^+\). It is noteworthy that the equilibrium binding constant of Na\(^+\) in the absence of citrate (defined by the K\(_D\)) is difficult to determine but likely is of much-lower affinity than the apparent K\(_M\) values for Na\(^+\) observed under transport (nonequilibrium) conditions (11, 22, 33). Thus, in the presence of only Na\(^+\), a fraction of the transporters is likely unoccupied and adopts an apo conformation, which has not been resolved in the available crystal structures. This conformational movement between outward- and inward states is possible, as reorientation of the empty carrier is part of the transport cycle of sodium-coupled symporters. The HS-AFM data show that the intermediate state may be visited on the way, although it is less favored than when citrate is also bound. In the sodium-bound conformation, the transporters would not be able to move between elevated and down states, but it is possible that they can still move between outward and intermediate or inward and intermediate states without being able to complete the full elevator movement. The inability to fully move when only Na\(^+\) is bound can be explained from the crystal structures. In the Na\(^+\)-only state, two arginine residues in the binding-side region (R402 and R428) likely prevent the elevator movement by exposing the sidechains at the interface between the scaffold and transport domains. In the fully loaded state (citrate plus Na\(^+\)-bound), these arginine side chains are more buried, because they interact directly with carboxylate groups of the substrate. In the apo state, the side chains may also be buried as they can now occupy the empty Na\(^+\)-binding sites. In both cases, the elevator movement would not be blocked.

By studying the dynamics of the CitS transporter, we were not only able to reveal the independent movement of the individual protomers but also that the elevator movement displays three different states. These unexpected insights raise the question of how other transmembrane transporters function, and the reported HS-AFM approach paves the way to address this question.

Methods

Expression and Purification of CitS. The expression vector pBAD.CitS (4) was transformed in E. coli/ MC1061 competent cells, and the recombinant strain was then cultivated in Luria-Bertani broth (LB) medium supplemented with 100 \(\mu\)g/mL ampicillin. The protein expression was induced with 0.05%
L-arabinose when the culture reached an optical density of 0.8 at 600 nm. Three hours after induction, cells were harvested (6,500 RPM, 4 °C, 15 min, Beckman JLA 10.100 rotor) and resuspended in 20 mM Tris HCl (pH 8) and resuspended to a concentration of 100 mg/mL, and 10 μM phenylmethylsulfonyl fluoride (PMSF), and 1 mM MgSO4 before being passed through a Con- stant System cell disruptor precooled at 4 °C and operated at 20 KPSI (Constant Systems Ltd., Daventry, United Kingdom). The remaining unbroken cells and cell debris were separated by centrifugation (7,500 RPM, 4 °C, 20 min, Beck- man 50.2 Ti, 30,000 g) and the crude cytoplasmic membrane in the superna- tant were then pelleted (40,000 RPM, 4 °C, 15 min, Beckman 45 Ti) and finally, resuspended in 20 mM Tris HCl (pH 8) for storing at −80 °C. The crude mem- branes containing CitS were solubilized in solubilization buffer (50 mM Tris HCl [pH 8], 300 mM KCl, 15 mM imidazole [pH 8], and 1% n-dodecyl-β-D-maltoside (DDM)) on a rocking platform at 4 °C for 45 min, and the solution was then centrifuged (55,000 RPM, 4 °C, 30 min, Beckman MLA 55 rotor), and the supernatant was incubated for 1 h with 500 μL of pre-equilibrated (50 mM Tris HCl [pH 8], 300 mM KCl) Ni-Sepharose resin. The mixture was then loaded on a Poly-Prep column (Bio-Rad), the unbound protein solution was let flow through, and the nonspecifically bound proteins were washed with 20 CV of cold washing buffer (50 mM Tris HCl [pH 8], 300 mM KCl, 60 mM imidazole [pH 8], and 0.15% DDM). The protein was eluted using consecutive additive of 300, 750, and 40 μL of cold elution buffer (50 mM Tris HCl [pH 8], 300 mM KCl, 500 mM imidazole [pH 8], and 0.15% DDM), the second fraction was then further purified by size-exclusion chromatography (SEC) on a Superdex 200 10/300 GL column and operated at 20 °C. The protein was added to the membrane on the side of a test tube containing 200 μL of ice-cold quenching buffer (10 mM potassium phosphate buffer [pH 7] and 200 mM KCl). The content of the tube was then poured onto a BA 45 nitrocellulose filter which was then washed with 2 mL of quenching buffer. The filters finally dissolved in scintillation mixture Ultima Gold (Perkin-Elmer), and the β-decay from the radiolabeled substrate was counted.

Citis Reconstitution in Liposomes for Kinetic Analysis. The lipids used to reconstitute CitS contained a 3:1 mixture of E. coli/PC (Avanti). Liposomes were homogenized by extruding 11 times through a 400-nm pore size filter and subsequently diluted to 5 mg/mL in 50 mM potassium phosphate buffer, pH 7. To allow the insertion of the protein into the bilayer, the lipids were destabi- lized by step-wise addition of 10% Triton X-100, while scattering was followed at a wavelength of 540 nm. The titration was stopped once the absorption sig- nal decreased to about 60% the maximum value reached. Purified protein was added at a protein/lipid ratio (wt/wt) of 1:1800. The protein lipid mixture was incubated for 30 min at room temperature, and then the detergent was removed by addition of BioBeads in three steps: First, 15 mg/mL BioBeads was added followed by incubation for 60 min at 4 °C, then 19 mg/mL BioBeads was added followed by overnight incubation at 4 °C. Finally, 29 mg/mL Bio- beads was added followed by 120 min incubation at 4 °C. BioBeads were then removed, and the proteoliposomes were pelleted (80,000 RPM, 25 min, 4 °C, Beckman MLAB0) and resuspended in 50 mM potassium phosphate buffer (pH 7) to a final lipid concentration of 20 mg/mL. The proteoliposomes were subjected to three cycles of freeze-thawing using liquid nitrogen and stored in liquid nitrogen until use.

Citis Steady-State Rates Measurement. The luminal buffer in each proteolipo- somal preparation was changed to 10 mM potassium phosphate buffer (pH 7) and 200 mM KCl. The proteoliposomes were first pelleted (80,000 RPM, 25 min, 4 °C, MLA80) and then resuspended in the desired luminal buffer. Three freeze-thaw cycles were performed in order to allow the exchange of buffer in the luminal area of the proteoliposomes. The solution was then extruded 11 times through a filter with 400-nm pore size in order to obtain homoge- neously sized unilamellar vesicles, which were pelleted (80,000 RPM, 25 min, 4 °C, TLA100.3) and resuspended to a final lipid concentration of 100 μg/mL. Steady-state rate cuve were measured at different concentrations of substrate and coions. An array of conditions was prepared as described in SI Appendix, Table S1. The blank measurement for each condition was measured by pipet- ing the liposome on the side of a test tube containing 200 μL of reaction buffer and subsequently flushing them in the reaction buffer with 2 mL of ice- cold quenching buffer (10 mM potassium phosphate buffer [pH 7] and 200 mM KCl). The content of the tube was then poured onto a BA 45 nitrocellulose filter which was then washed with 2 mL of quenching buffer. The filters finally dissolved in scintillation mixture Ultima Gold (Perkin-Elmer), and the β-decay from the radiolabeled substrate was counted.

The value for each uptake rate reported in Si Appendix, Table S1 repre- sents the average and SE of three independent biological replicates, each con- stituted by two or three technical replicates. The substrate-dependent uptake rates obtained at fixed concentration of Na+ were plotted as a function of cit- rate (app) related to the MM equation (SEE on a Superdex 200 10/300 GL column) increase 10/300 GL (GE Healthcare) column using SEC buffer (10 mM Heps-KOH [pH 8], 100 mM KCl, and 0.15% DDM) as running buffer, and the temperature was kept at 4 °C for the whole procedure. Uptake assays were performed to assess that CitS retained its activity after purification.

High-Density Reconstitution of CitS on Membrane. To achieve a densely packed membrane patch, we followed the procedure by Keble and col- leagues (17). This procedure densely packed liposomes were mixed at a lipid to protein ratio of 1:800. The protein lipid mixture was incubated for 30 min at room temperature, and then the detergent was removed by addition of BioBeads in three steps: 15 mg/mL of BioBeads was added and the solution was incubated for 30 min at 4 °C, then 19 mg/mL BioBeads was added followed by O/N incubation at 4 °C. Finally, 29 mg/mL Bio- beads was added followed by 120 min incubation at 4 °C. BioBeads were then removed, and the proteoliposomes were pelleted (80,000 RPM, 25 min, 4 °C, Beckman MLAB0) and resuspended in 10 mM potassium phosphate buffer (pH 7), to a final lipid concentration of 20 mg/mL. The proteoliposomes were subjected to three cycles of freeze-thawing using liquid nitrogen and extruded 11 times through a 400-nm pore size polycarbonate filter and pel- leted again (80,000 RPM, 25 min, 4 °C, Beckman MLAB0). The liposomes were resuspended to a concentration of 100 mg/mL, and 10 μL finally was added to 1 mL of solution containing 10 μM citrate (of which 0.5 μM was labeled with 14C), 200 mM NaCl, and 10 mM potassium phosphate buffer (pH 7). The reaction was started by adding 200 μL of reaction buffer in 2 mL of ice-cold quenching buffer containing 10 mM potassium phosphate buffer (pH 7) and 200 mM KCl after 10, 20, 30, 60, and 90 s from the start of the reaction. The quenched reaction was filtered through a BA 45 nitrocellulose filter and then washed with 2 mL of quenching buffer. The filters finally dissolved in scintillation mixture Ultima Gold (Perkin-Elmer), and the β-decay from the radiolabeled substrate was counted.

AFM Experiments. The AFM images were acquired in amplitude modulation tapping mode in liquid, using high-speed atomic force microscopes (both a home-built system of T.A. and a commercial system from Research Institute of Biomolecule Metrology, Japan) (35, 36). The HS-AFM imaging was performed using small silicon nitride cantilevers (BL-AC10-D5, Olympus, Japan) with a nominal spring constant of ~0.1 N/m and a resonance frequency of ~400 to 500 kHz in buffer. The probe tip was ground on top of the cantilever by elec- tron beam deposition (37) using an electron beam lithography system (EL5- 7500UK, Elionix, Japan). The tips were then sharpened using plasma cleaning for 30 s. The applied-imaging forces were controlled by using a cantilever set- point amplitude of 0.8 nm with 1-nm free oscillation amplitude. During incu- bation and imaging, a buffer containing 50 mM potassium phosphate, pH 7, was used, supplemented with 300 mM NaCl or 300 mM NaCl and 150 μM cit- rate, pH 7, when indicated. Prior to HS-AFM imaging, the CitS-reconstituted liposomes were incubated on a freshly cleaved mica for 15 to 20 min. After incubation, the surface was cleaned thoroughly using the pH 7 phosphate buffer (when appropriately supplemented with Na+ or Na+ and citrate). The buffer conditions were changed (adding Na+ or adding Na+ and citrate) by using a syringe pump connected to a 55-μL measurement chamber. Buffer flow was kept constant at 60 μL/min, and a pumping time of about 5 min to assure a complete buffer exchange. Measurements were started after buffer exchange had finished.

AFM Data Analysis. The acquired AFM images were analyzed using home- built software of Kanazawa University (Kodec), Igor Pro, and ImageJ with additional home-written plug-ins. Height measurements were performed on
raw images after tilt correction. The evaluation of real heights of a single transporting domain was performed frame by frame by calculating the height of the domain and subtracting the height of the membrane in close vicinity (SI Appendix, Fig. S3 A and B). The height of a single transporting domain was calculated by tracking individual protomers and measuring the maximum height in the region of interest (15, 38). To exclude the possibility of surface fluctuations interfering with the height measurements, we estimated the distribution of roughness of the membrane surface over time. This resulted in a single Gaussian distribution around 0.02 ± 0.3 nm (SI Appendix, Fig. S3C). Given an SD of 0.3 nm, the multiplex fits in Fig. 4 A–C reveal clearly distinguishable peaks. For the data in the right panels of Fig. 4 A–C, the height dynamics of, respectively, 5, 8, and 14 dimers were analyzed for over 3,000 s for each condition. All stated errors in the text are SDs unless otherwise mentioned. The HS-AFM height versus time traces were idealized in Fig. 4 for each condition. All stated errors in the text are SDs unless otherwise mentioned.

Discussion of the Kinetic Analysis Using Idealized and/or iHMM. We have used both the idealized traces and iHMM to describe the three-states mechanism of citrate transporter CitS. While iHMM allowed us to identify three states without defining a prerequisite number of states as input parameter, the method fails to identify three states for some short traces, in which transition to one of the states is present but not for a significant amount of time. Examples are shown in SI Appendix, Fig. 56. It is observable that the trace SI Appendix, Fig. S6A shows a reasonable transition between the states (also observable in the histogram); as a result, iHMM could successfully detect three states. However, the trace in SI Appendix, Fig. S6B has only few visits to the down state (also observable in the histogram) within the duration of recording, causing iHMM not being able to detect three states. In order to include all possible transitions and to calculate the transition frequencies, we have used idealized traces. Comparative trajectories by fitting iHMM and idealized traces are shown in SI Appendix, Fig. S7. The kinetic analysis was done using both idealized traces and using iHMM and are summarized in SI Appendix, Tables S2 and S3.

Data Availability. All study data are included in the article and/or supporting information.

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Kinetic Analysis of Idealized Traces. The idealized traces of individual protomers obtained by thresholding were used to obtain the transition probability matrix and dwell-time distributions (SI Appendix, Fig. S5), as described in ref. 39. A custom-written MATLAB script was used to compute the counting matrix (M) from the individual-state trajectories, in which every matrix element (Mij) represents the number of transitions from state i to state j after a certain time t (frame rate). The matrix was row normalized to obtain the transition probability matrix, with its individual terms (Pij) giving the frequency of the protomer transition from state i to state j. The expression kij = −ln(1 − Pij)/t was used to estimate the rate constants (kij) governing the individual-state transitions (SI Appendix, Tables S1 and S2).

Hidden Markov Analysis. The iHMM (25) was applied to analyze the HS-AFM height versus time traces of individual protomers. This generalization of the HMM, based on Bayesian nonparametrics, allows analyzing the experimental time traces without a priori setting of a specified number of states. All traces were analyzed by the iHMM for an initial number of states of 10, which were reduced by the algorithm to two or three states before the first 500 iterations. All traces are shown in SI Appendix, Fig. S5b. After these burn-in periods of 500 iterations, additional 1,500 to 2,000 iterations were performed to compute the transition probability matrix for every frame. Transition-concentration (c) and base-concentration (c) parameters were varied between 1 and 103 and 10−1 and 10−3, respectively, keeping the rest of the parameters as default (25). The rate constants derived from the iHMM analysis are presented in SI Appendix, Tables S2 and S3. It is important to note that given the temporal resolution (300 to 400 ms per frame), the characterization of short-lived states will give rise to some uncertainties, and very short-lived states will not properly be identified by our approach. Unless otherwise mentioned, all presented errors are in terms of SD.

Estimation of Free Energy Difference. The estimation of free energy difference ∆G° between state j and i is calculated as:

$$\Delta G^0_{i,j} = -\ln(k_{i,j}/k_{j,i}) \cdot k_B T,$$

where kij is the estimated rate constant for the transition from i to j and kji is the estimated rate constant for the transition from j to i. k_B is the Boltzmann constant and T absolute temperature.

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