Organization and Dynamics of the Red Blood Cell Band 3 Anion Exchanger SLC4A1: Insights From Molecular Dynamics Simulations

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Molecular dynamics (MD) simulations have provided new insights into the organization and dynamics of the red blood cell Band 3 anion exchanger (AE1, SLC4A1). Band 3, like many solute carriers, works by an alternating access mode of transport where the protein rapidly (10⁴ ions/s/Band 3) changes its conformation between outward and inward-facing states via a transient occluded anion-bound intermediate. While structural studies of membrane proteins usually reveal valuable structural information, these studies provide a static view often in the presence of detergents. Membrane transporters are embedded in a lipid bilayer and associated lipids play a role in their folding and function. In this review, we highlight MD simulations of Band 3 in realistic lipid bilayers that revealed specific lipid and protein interactions and were used to re-create a model of the Wright (Wr) blood group antigen complex of Band 3 and Glycophorin A. Current MD studies of Band 3 and related transporters are focused on describing the trajectory of substrate binding and translocation in real time. A structure of the intact Band 3 protein has yet to be achieved experimentally, but cryo-electron microscopy in combination with MD simulations holds promise to capture the conformational changes associated with anion transport in exquisite molecular detail.

Keywords: Band 3, anion transport, red blood cell, membrane transporters, molecular dynamics simulations

INTRODUCTION

Band 3, the red blood cell Cl⁻/HCO₃⁻ anion exchanger 1 (AE1), is the founding member of the SLC4 family of bicarbonate transporters and plays an essential role in respiration (Alper, 2009; Cordat and Reithmeier, 2014; Jennings, 2021). In the tissues, carbon dioxide enters the red blood cell primarily by diffusion where it is hydrated by carbonic anhydrase II to form bicarbonate and a proton that is buffered by hemoglobin. The bicarbonate is rapidly (10⁴ ions/s/Band 3) transported out of the red blood cell by the 1.2 million copies of Band 3 in exchange for chloride, increasing the blood’s capacity to transport carbon dioxide as plasma bicarbonate. In the lung, bicarbonate enters the red blood cell by Band 3 in exchange for chloride where it is dehydrated by carbonic anhydrase II to form carbon dioxide and water; the carbon dioxide diffusing out of the red blood cells to be expired.
was determined at neutral pH (Shnitsar et al., 2013). The isolated cytoplasmic domain of Band 3 is dimeric and undergoes dramatic pH-dependent conformational changes forming a more compact structure at low pH as revealed by the crystal structure (Zhou and Low, 2001).

Band 3 contains cysteine residues at residues 201 and 317 in the cytoplasmic domain (Figure 1B) that can be crosslinked to each other at neutral pH forming a covalent intermolecular dimer (Steck, 1972; Reithmeier and Rao, 1979). Interestingly, these two residues can be crosslinked intramolecularly to each other within each monomer in the isolated cytoplasmic domain. In the crystal structure of the isolated cytoplasmic domain, Cys 201 and Cys317 are ~29Å apart in each monomer and ~20Å apart across the dimer. This indicates that there is considerable conformational dynamics of the cytoplasmic domain. Perhaps, MD simulations can reveal the nature and scope of these conformational dynamics in the context of the isolated domain and the intact protein.

The cytoplasmic domain provides the site of interaction of Band 3 with hemoglobin, glycolytic enzymes, and the cytoskeleton (Low, 1986). Protein 4.2 stabilizes the interaction of Band 3 with ankyrin. The ankyrin binding site has been localized to a hairpin loop at residues 175–185 with contributions from residues 63–73 (Stefanovic et al., 2007). It may be possible to form a complex of the interacting portion of ankyrin with Band 3 for structural studies. Two Band 3 supramolecular complexes exist in red blood cells: the Band 3-ankyrin complex and the junctional complex (Bruce et al., 2003). The ankyrin complex not only includes Band 3, Glycophorin A, and protein 4.2 but also an association with the Rh complex (Rh-associated glycoprotein, Rh polypeptides, CD47, and LW), while the junctional complex includes Band 3, Glycophorin C, p55, protein 4.1, adducin, dematin, and GLUT1; both complexes provide linkages to the spectrin-actin cytoskeleton (Mankelow et al., 2012). Cryo-electron microscopy may be ideally suited to examine the structure of such supramolecular complexes in situ.

Band 3 is a mixture of dimers and tetramers in the membrane and when solubilized in mild detergent solutions (Casey and Reithmeier, 1991). In contrast, the isolated membrane domain (residues 361–911) is exclusively a dimer (Reithmeier, 1979). The crystal structure of the dimeric membrane domain in the outward-facing conformation (Arakawa et al., 2015; Reithmeier et al., 2016; Figure 2) reveals a complex folding pattern with 14 transmembrane (TM) segments in a 7+7 TM inverted repeat common to members of the SLC4 family, the related SLC26 anion transporter family, and the unrelated SLC23 family of nucleobase transporters (Chang and Geerttsma, 2017). Each subunit consists of two sub-domains, the gate domain (TM 5, 6, 7, 12, 13, and 14) at the dimer interface and the core domain (TM 1, 2, 3, 4, 8, 9, 10, and 11) that contains the substrate binding site (Figure 3). The dimer is held together by interactions involving the extracellular ends of TM5 and 6. The Tanner lab (Groves et al., 1998) showed that complementary TM fragments of Band 3 (e.g. TM 1–2 + TM 3–4) representing the entire sequence could reassemble into functional units when expressed in Xenopus oocytes, including the unexpected finding that non-complementary TM1-5 and
TM8–14 entirely missing TM6 and 7 was functional. The anion binding site is centrally located between the amino-termini of the short TM3 and TM10 segments with a positive charge contributed by Arg730 located in TM10. The short carboxyl-terminal tail of Band 3 interacts with carbonic anhydrase II forming the basis for a bicarbonate transport metabolon (Sterling et al., 2001).

**BAND 3 KINETICS**

Extensive kinetic studies have been carried out on anion transport in red blood cells (Knauf, 1979). Firstly, the affinity for anions is in the mM range, allowing a very rapid off-rate. Secondly, there is a single central anion binding site. Thirdly, Band 3 works by an alternating access mode of transport. Fourthly, the anion binding site is accessed by water-filled cavities. Fifthly, the empty carrier cannot undergo the transition enabling tight 1:1 exchange. Sixthly, anion binding reduces the energy barrier for the transition. Finally, anion exchange is very rapid (~10^4 ions/s). The molecular details that allow for such rapid transport remain to be fully described.

Kinetic studies have also shown that SLC4A2 (AE2) operates by a ping-pong mechanism (Falke and Chan, 1985). This suggests that the nature of the dimer interface may play a role in the mode of transport and its allosteric regulation in the SLC4 family of transporters.

The structural and functional studies of Band 3 described above have provided new insights into the mechanism of action of this transport protein. Yet, a complete molecular description of the conformational change associated with transport remains elusive. Furthermore, there are a number of inherited conditions, such as hereditary spherocytosis (HS), Southeast-Asian ovalocytosis (SAO), and distal renal tubular acidosis (dRTA) caused by mutations that affect Band 3 folding, trafficking, and function. A full understanding of the effect of these mutations on the structure and dynamics of Band 3 awaits further exploration. MD simulations are ideally suited to explore these issues since this powerful computational method provides a direct link between the structure and the conformational landscape accessible to the protein molecule. These dynamic insights will reveal how Band 3 works at the molecular level and how mutations associated with disease affect their ability to fold and function properly.

**MOLECULAR DYNAMICS SIMULATIONS**

Molecular dynamics (MD) simulations of membrane proteins in complex lipid bilayers, both at the coarse-grained and atomistic levels, have provided new insights into the organization and dynamics of these proteins (Marrink et al., 2019). Detergents are commonly used to solubilize, purify, and crystallize membrane proteins. Native membrane proteins however function within complex lipid bilayers. Some crystal structures of membrane proteins contain tightly-bound lipids, including cholesterol (Hunte and Richers, 2008). In the case of Band 3, removing these lipids results in destabilization of the protein and aggregation (Maneri and Low, 1989; Vince et al., 1997). Lipids play a key role in oligomerization of membrane transporters (Gupta et al., 2017).

Membrane transport proteins, like Band 3, are commonly found as oligomers, mostly as dimers. The oligomeric state allows for allosteric interactions between the subunits. In addition, the proper assembly of oligomers is a requirement for exit from the ER and may be part of protein quality control. As we will discuss, the dimeric structure of membrane transport proteins can provide a rigid scaffold to allow the relative movement of mobile elements associated with transport. In the case of Band 3 and transport proteins with a similar 7+7 inverted repeat structure, it is the gate domain that forms a rigid dimer interface and the core domain that binds substrate and undergoes movement to provide alternating access to the substrate binding site.

**THE TM1 SIGNAL ANCHOR**

The first TM segment of Band 3 (Figure 3) acts as a signal anchor to target nascent Band 3 to the endoplasmic reticulum.
(ER) membrane (Popov et al., 1997). The first TM segment translocates the following TM2/3 region into the ER lumen with TM4 acting as a stop-transfer sequence (Cheung and Reithmeier, 2005). TM1 moves laterally from the translocon into the lipid bilayer, likely along with TM4. TM2 and 3 subsequently fold into the nascent structure. TM1-4 makes up part of the core domain. MD simulations showed that isolated TM1 can assume a native helical conformation similar to that found in the native protein (Fowler et al., 2017).

A nine-amino acid deletion (Ala400–Ala408) at the cytoplasmic interface region found in SAO Band 3 prevents the shortened TM segment from interacting properly with the lipid bilayer and forming a stable TM segment (Cheung et al., 2005). This feature accounts for the negative effect of the deletion on the folding and trafficking of SAO Band 3 (Moriyama et al., 1992; Sarabia et al., 1993). The SAO deletion removes Pro403 within a bend that connects the short H1 helix to the TM1 helix (Figure 3), which may result in a change in the relative orientation of the cytoplasmic and membrane domains. The more distal Pro419 places a flexible hinge in the middle of TM1 due to loss of backbone hydrogen bonds to residues 3 and 4 proximal, leaving the carbonyl groups at Ala415 and Ala416 free for other hydrogen-bonding interactions. Early NMR studies (Chambers et al., 1999) of synthetic peptides corresponding to TM1 revealed the role of proline residues in disrupting its helical structure and the effect of the SAO deletion. Despite these studies, the structural effect of the SAO mutation on the membrane domain of Band 3 has not been fully described. Current studies using MD simulations to construct a molecular model of SAO Band 3 will allow us to examine these details at the atomistic level.

**BAND 3-GLYCOPHORIN A INTERACTIONS**

Band 3 interacts with Glycophorin A forming the Wright (Wr) blood group antigen. Central to this interaction is a salt bridge between Glu658 in Band 3 and Arg61 in Glycophorin A (Bruce et al., 1995). Wr(a+b+) heterozygotes also have Lys at residue 658. Glycophorin A interacts with Band 3 in the ER facilitating its trafficking to the cell surface (Groves and Tanner, 1992; Pang and Reithmeier, 2009). MD simulations of the membrane domain of Band 3 and the dimeric TM portion of Glycophorin A were carried out in a complex lipid bilayer (Kalli and Reithmeier, 2018). Glycophorin A was positioned in the greatest distance from Band 3 while maintaining the Glu658-Arg61 ionic interaction. Glycophorin A moved closer to Band 3 suggesting that in addition to the interaction between Glu658 and Arg61 there are also interactions between the helical TM segments and extracellular regions of Glycophorin A and similar regions in Band 3 (Figure 4A). Figure 4B shows the residues involved in the interaction of Band 3 and Glycophorin A. Changing Glu658 to Lys resulted in loss of the interaction with Arg61 but the TM region of Glycophorin A still interacted.
with Band 3. This suggests that the Wright blood group antigen is probably due to a local conformational change in the epitope rather than complete dissociation of the Glycophorin A/Band 3 complex. Large-scale simulations showed that the Glycophorin A dimer can bridge Band 3 dimers resulting in the dynamic formation of a supramolecular complex of long strands of alternating Band 3 and Glycophorin A dimers. It would be of considerable interest to use MD simulations to examine the effect of Band 3 mutations (e.g., G701D) on the structure of Band 3 and its interaction with Glycophorin A.

**BAND 3-LIPID INTERACTIONS**

Band 3 is embedded in a complex lipid bilayer and has extensive interactions with both phospholipids and cholesterol. MD simulations of the membrane domain Band 3 have revealed specific binding sites for acidic lipids and cholesterol (Kalli and Reithmeier, 2018). Cholesterol was found at the dimer interface where it may play a structural role in stabilizing the interaction between the gate domains of the two subunits (Figure 5B). In addition, the MD simulation studies have also suggested that there is enough space in the dimer interface for one phospholipid or sphingomyelin molecule. Cholesterol is known to inhibit anion transport and so may also play a regulatory role (Gregg and Reithmeier, 1983). Although Band 3 interacted with both 1-palmitoyl-2-oleoyl-phosphatidylserine (POPS) and phosphatidylglycerol 4,5-bisphosphate (PIP2) lipids, MD simulations have shown a strong preference for PIP2 lipids. PIP2 provides binding sites for cytoplasmic proteins such as Protein 4.1. Band 3 is therefore a central organization center in the red blood cell membrane by direct interaction with proteins and associated lipids like PIP2. Thus, the interaction of proteins with Band 3 may be regulated via modification of associated lipids.

Molecular dynamics simulations and modeling of intact Band 3 were used to provide an atomistic view of the interaction mass spectrometry (MS) and proteomics of the cytoplasmic domain (missing the first 54 disorderd residues), the membrane domain, and the connecting linker in a model of AE1 in a complex lipid bilayer (de Vecchis et al., 2019). During the simulations, the cytoplasmic domain moved closer to the membrane domain forming a more compact structure. The flexible linker was involved in the interaction of the membrane and cytoplasmic region but its removal after Band 3 adopted a more compact structure did not change the stability of the structure. Analysis of the possible orientations of the cytoplasmic region relative to the membrane region of Band 3 based on available experimental data indicated that a structure with the C-terminal dimerization arms (residues 314–347) of the cytoplasmic region facing the transmembrane region of Band 3 was most likely (Figure 5). The two twisted models in which there was a domain swap in the orientation of the cytoplasmic domain allowed closer packing of the complex compared to the other orientations. Structures of SLC26 transporters, which also feature a 14 TM 7+7 inverted repeat topology (Baranovski et al., 2020), revealed a domain swap of the C-terminal STAS domain with major interactions between the STAS domain and the membrane domain being responsible for dimer stability (Chang et al., 2019; Walter et al., 2019). There are interactions of the cytoplasmic domain of Band 3 with the membrane domain in a cross-over structure that may stabilize the dimer; however, both domains in isolation remain dimeric. This structural and functional independence of the two domains in Band 3 may not be a feature of other members of the SLC4 family, whose transport activity is regulated by intracellular pH and other factors as opposed to Band 3 that works near maximum velocity (Zhang et al., 1996).

The red blood cell membrane contains lipid nanodomains (“rafts”) enriched in cholesterol and sphingomyelin (Simons and Ikonen, 1997). Atomistic MD simulations with the membrane region of Band 3 inserted in membranes that contained 1,2-dipalmitoyl-phosphatidylcholine (DPPC), 1,2-dilinoleoyl-phosphatidylcholine (DLiPC), and cholesterol (4:3:3) showed that there was an asymmetric interaction of Band 3 with the phase-separated lipid nano domains in the two leaflets in this bilayer (Jin et al., 2020). At the end of these simulations, the contacts between Band 3 and DLiPC and DPPC were ~4.1 in the outer leaflet and ~3.2 in the inner leaflet. This suggests that in the outer leaflet the protein contacts mainly low-Tm Ld domains, whilst in the inner leaflet Band 3 contacts both Ld and high-Tm lipid and cholesterol Lo domains. Simulations of Band 3 in a DPPC, DLiPC with 30 or 50mol % cholesterol showed that a higher cholesterol concentration results in the deformation of the bilayer.
some helices in Band 3. Simulation with a bilayer that contained 1,2-distearoyl-phosphatidylserine (DSPS), DLiPC, and cholesterol revealed strong electrostatic interactions between phosphatidylserine (PS) lipids and Band 3. As a result, Band 3 interacts with Lo domains at specific sites in both leaflets; this is not the case in the simulations with DPPC, DLiPC, and cholesterol.

The bacterial purine transport UraA has a similar 7 + 7 inverted topology as Band 3 (Lu et al., 2011). MD simulations of the UraA monomer in a complex lipid bilayer revealed specific interactions with lipids, particularly cardiolipin (Kalli et al., 2015). MD simulations of the fungal proton-driven purine transporter UapA support the essential role of lipids in stabilizing its functional dimeric structure and an elevator model for transport (Alguel et al., 2016). MD simulations of a mutant form of UapA (G411V) trapped in the inward-facing state identified an essential arginine-rich lipid-binding site at the dimer interface (Pyle et al., 2018). The fumarate transporter SLC26Dg crystallized as a monomer but is reported to be a dimer in a lipid bilayer (Geertsma et al., 2015). All of these studies point to an essential role of lipids in stabilizing the dimeric state of these transport proteins.

**CONFORMATIONAL CHANGES ASSOCIATED WITH TRANSPORT**

Band 3, like many facilitated transporters, works by an alternating access model. The crystal structure of the membrane domain has revealed key features of this mode of transport (Reithmeier et al., 2016). First, there is a single anion binding site located between the N-termini of two short α-helices (TM3 and 10) with participation of the side chain of Arg730, consistent with earlier kinetic studies. This site is occupied by one of the sulfonyl groups of the inhibitor Diisothiocyanostilbene-disulfonate (DIDS), which locked the protein into the outward-facing state. A second feature of Band 3 is the presence of two domains, a so-called gate domain at the dimer interface and a core domain that has a predominate interaction with the lipid bilayer. The structure resembles a mobile bone and socket with slippery hydrophobic interacting surfaces. The interface between these two domains is dynamic allowing alternating substrate access from the outside and inside.

Molecular dynamics simulations combined with experimental data have shown two substrate binding sites in the outward-facing state of Band 3 and the related human electrogenic sodium bicarbonate cotransporter (HNBCe1, SLC4A4; Zhekova et al., 2021). One site is located at the entry of the outward-facing cavity and the other one in a more central position toward the middle of the protein. Band 3 residue R730 was shown to be critical for anion binding for both the entry and central binding sites. The cryo-electron microscopic structure of a sodium-dependent chloride/bicarbonate exchanger (SLC4A8) in the outward-facing state had well-defined densities for both the Na⁺ close to D800 and bicarbonate in the central substrate binding site (Wang et al., 2021). Interestingly, the dimeric structure was further stabilized by a compact extracellular N-glycosylated domain between TM5 and 6 that contained two disulfide bonds. Dimerization would confer stability between the two interacting gate domains in the plane of the membrane, suggesting that it is the core domain that moves during the transport cycle.

**ROCKER AND ELEVATOR TRANSPORT MODELS**

There are two basic structural mechanisms for alternating access model of membrane transport: the rocker switch or rocking bundle model and the elevator model (Diallinas, 2021). In both models, there is a central substrate binding site located...
The 7 + 7 TM inverted repeats consisting of TM1-7 and TM8-14 in each Band 3 monomer have a similar fold. The 7 + 7 inverted repeat structure is consistent with an alternating access model of transport. The repeats are however not homologous and differ greatly in amino acid sequence resulting in an asymmetric structure. In addition, it is not the repeats that move relative to one another but rather the core and gate domains.

A model (Figure 7) of the inward-facing state of Band 3 was constructed using repeat-swap homology modeling whereby the second seven TM repeat assumes the conformation of the first seven TM repeat and vice versa (Ficici et al., 2017). The modeling suggested that the transport mechanism of Band 3 involves an elevator-like translation of about 8 Å to the bilayer normal (~11 Å in total) and rotation of ~17 degrees of the substrate-binding core domain relative to the nearly stationary dimerization gate domain. In the outward facing state, the water-filled passage is lined by TM1 and 3 from the core domain and TM5 and 13 from the gate domain, while in the inward-facing state the 7 + 7 symmetry-related TM8 and 10 and TM6 and 12 are predicted to line the passage. Early studies showed that crosslinking the two monomers via residues now known to be within the gate domain in the TM5-6 loop (Lys551–Lys562) did not inhibit transport indicating that the dimer interface does not undergo a major conformational change during transport (Jennings and Nicknish, 1985).

Support for the elevator model for Band 3 comes from structural studies of the related plant (Thurtle-Schmidt and Stroud, 2016) and yeast SLC4 borate transporter BorA (Coudray et al., 2017). The Arapidopsis protein crystallized as a dimer with each subunit having the same 14 TM fold as Band 3. Although no substrate could be identified at the 4.1 Å resolution, the structure was in an occluded state in which the core domains are rotated inward toward the gate domain with the extracellular ends of TM1, 3 and 8 moving inward by about 8 Å to close off the extracellular side of the protein. The structure of the yeast BorA protein was determined from helical membrane crystals at 6 Å resolution. Homology modeling (~23% sequence identity) and MD simulations using Band 3 as a template and water accessibility revealed that the BorA was in the inward-facing conformation with a rigid body rotation of the core domain by ~10 degrees against the immobile gate domain relative to Band 3. Comparison of the inward-facing structure with an outward-facing model of BorA did not reveal any translation of the core domain, leading the author to suggest that BorA does not operate by an elevator mechanism. It is important to recognize that the structure represents a substrate-bound

**Rocker transport model:**

**Elevator transport model:**

![Figure 6](https://www.frontiersin.org)
occluded form of BorA and not the empty carrier in its full inward-facing state. The same can be said for the structure of Band 3 that was determined in the presence of the bound inhibitor DIDS and does not represent the empty outward-facing state. Thus, the conformational changes associated with the full transport cycle may be more profound that observed in substrate or inhibitor-bound forms.

The 7+7 inverted repeat topology of the SLC4 family is also shared with members of the SLC23 family of nucleobase transporters. The crystal structure of the proton-driven uracil symporter, UraA in an inward-facing state was the first to reveal the 14 TM 7+7 inverted repeat topology (Lu et al., 2011). The UraA structure was originally reported as a monomer; however, the crystal lattice indicated that the protein is dimer with interactions between the gate domains. The structure of UraA with bound uracil in an occluded state in dimeric form was subsequently determined (Yu et al., 2017). The core domains of the inward-facing and occluded state could be superimposed, while there were conformational changes within the gate domain. There are structural changes in TM5 and its inverted partner TM12. Both TM5 and 12 rotate around an axis perpendicular to the interface between the core and gate domains. TM5 and 12 are both straight α-helices in the inward-facing state but both are kinked in the occluded state. The periplasmic end of TM5 moves toward the core domain, while the cytoplasmic end of TM12 moves away creating an inward-facing opening. In addition, the periplasmic loop connecting TM13 and 14 formed a rigid β-hairpin “paddle” in the occluded state. The binding of uracil in the core domain remains the same in both states. There is a translocation of ~5 Å of the substrate in the core domain between the occluded state of UraA and inward-facing state of UapA, consistent with an elevator model of transport. There were also significant conformational changes observed in the gate domain of UraA. The higher resolution of the occluded state at 2.5 Å identified water molecules bridging uracil to essential water-binding residues (Wat 1 to Ser72 and Glu241 and Wat 2 to Glu241 and His245). In the previous inward-facing state, a detergent molecule occupied the position of the Wat2 water molecule. MD simulations of UraA in a lipid bilayer identified a proton translocation pathway involving loss of water binding to Glu241 and interaction with His245.

UraA exists as a functional dimer in the membrane and is equilibrium between monomer and dimer in detergent (Yu et al., 2017). This points once again to the essential role that lipids play in the stabilization of the oligomeric state of membrane transport proteins. The dimer is held together by hydrophobic interactions between the gate domains with TM13 in one subunit packed into TM5 and 12 in the other subunit.

Our MD simulations of monomeric UraA in the absence of uracil in a lipid bilayer mimicking the bacterial inner membrane found that cardiolipin interacted with three specific sites, one on the outer surface (Arg321) and two on the inner surface (Arg299/Arg4 and Arg265/Lys109; Kalli et al., 2015). Mutation of the interacting arginine or lysine residues at these sites resulted loss of lipid binding. Cardiolipin may stabilize the structure of UraA and promote dimer formation. The cardiolipin on the outer surface of UraA is close to the transport pathway and may act as a “proton trap” to channel protons to the transporter. Cardiolipin on the inner surface may channel protons from the transporter. Cardiolipin may play a similar functional role in proton translocation in bacterial and mitochondrial proton-driven symporters that are embedded in membrane rich in cardiolipin. In the absence of uracil, the MD simulations of the UraA monomer showed that the inward-facing transporter assumed a more closed state, due to relative movement of the gate and core domains. This closed state in the absence of substrate may represent an intermediate in the empty carrier changing from the inward-facing to outward-facing state to complete the transport cycle. The transport cycle involves sequential
binding of the proton and uracil to the outward-facing state, a conformational change through an occluded state to the inward facing state, sequential release of the proton and uracil, and finally a conformational change of the now empty carrier to return to the original outward-facing state. This transport cycle may explain why symporters (co-transporters) tend to be crystallized in the inward-facing state in the presence of substrate. In contrast, antiporters (exchangers) undergo the conformational change in the presence of substrate and must be trapped in one conformational state with the use of inhibitors.

The structure of the proton-driven nucleobase xanthine symporter, UapA from Aspergillus was determined in the substrate-bound inward-facing state (Alguel et al., 2016). UapA had the same 7+7 inverted repeat fold as UraA with corresponding core and gate domains. The substrate xanthine was held within the core domain between the two TM3 and 10 half helices in a similar position as uracil in UraA. UapA crystallized as a dimer with extensive interactions involving TMs12-14 in the gate domain. The dimeric structure of UapA was found to be essential for trafficking to the cell surface and transport function (Kourkoulou et al., 2019). Mutations affecting transport and substrate specificity were found in the substrate-binding core domain as expected (e.g., Phe 406) but also in the dimerization gate domain in TM 12 and 14. A disulfide bond in the extracellular loop connecting TM 3 and 4 was essential for proper folding and trafficking. Mutation of Arg481 affects substrate specificity. This residue is located close to the substrate binding site of its partner subunit. Co-expression of inactive mutants with the normal protein resulted in a dominant-negative effect on transport consistent with the dimer being the functional transport unit. A model of the outward-facing state of UapA was built using Band 3 as a template. In UapA, the core domain is displaced downward toward by ~10Å in the inward-facing state, while the gate domain remains in fixed position in the membrane. Tightly-bound lipids (PI and PE) were found associated with UapA and stabilized the dimeric form of the protein (Pyle et al., 2018). MD simulations localized PI to specific basic residues on the cytoplasmic surface near the dimer interface. Mutation of specific arginine residues involved in PI binding resulted in destabilization of the dimer and loss of transport activity.

There are likely subtle but important differences in the details of the conformational changes in the 14 TM 7+7 inverted repeat transporters. For example, in the UraA proton-uracil co-transporter (symporter), the empty carrier must be able to move from the inward-facing to the outward-facing state to complete the transport cycle. In exchangers (antiporters) like Band 3, the empty carrier cannot undergo this conformational change to the outward state without bound substrate.

The crystal structure of the membrane domain of Band 3 is locked in the outward-facing state and we do not yet have a structure of the inward-facing conformation. We are currently using the inward-facing and occluded states of UraA to build equivalent models of Band 3. It is unlikely that transport occurs by simple rigid body movement of the two domains. MD simulations will allow us to examine the full extent of the structural changes associated with the transition from outward to inward-facing state of Band 3 with a focus on the relative movement of the gate and core domains.

**CONCLUSION**

Finally, an excellent example of the conformational dynamics of Band 3 is the senescence antigen (Badior and Casey, 2021). Early work showed that the Band 3 sequence encompassing residues 812–830 plays a role in binding endogenous senescence antibodies to aged red blood cells. In the crystal structure of the membrane domain, this region is on the cytoplasmic side of the membrane and consists of two short α-helices joined by a reverse turn linking TM12 and 13 (Figure 3). Using cysteine scanning mutagenesis of Band 3 and the membrane domain expressed in HEK cells, it was demonstrated that this region can assume an extracellular disposition. TM13 and 14 are shorter than typical TM α-helices and may not have a very stable interaction with the lipid bilayer (Deber et al., 2002). The interface region of the bilayer is about 15Å thick allowing interacting TM segments of membrane proteins some flexibility in their position relative to the plane of the membrane (Reithmeier, 1995). The ~30Å thick hydrophobic zone is likely a greasy slide, again allowing some movement of mobile domains. Cys843 has been identified as a site of palmitoylation in Band 3, yet is located in the middle of TM13. Fatty acylated residues are typically located at the membrane interface with the fatty acid positioned along with other fatty acyl groups in the lipid bilayer (Cheung and Reithmeier, 2004). This indicates that TM13 has a dynamic aspect and that the non-enzymatic palmitoylation of Cys843 may occur as red blood cells age and be part of the senescence clock.
of Band 3 expressed in yeast (Sekler et al., 1995) or transfected cells (Okawa et al., 2014).

AUTHOR CONTRIBUTIONS
RAFR and ACK contributed equally to the writing of this review. References were complied by RAFR, and figures were prepared by ACK. All authors contributed to the article and approved the submitted version.

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