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Structure of a tetrameric MscL in an expanded intermediate state

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Abstract

The ability of cells to sense and respond to mechanical force underlies diverse processes such as touch and hearing in animals, gravitropism in plants, and bacterial osmoregulation 1, 2. In bacteria, mechanosensation is mediated by the mechanosensitive channels of large (MscL), small (MscS), potassium-dependent (MscK), and mini (MscM) conductances. These channels act as "emergency relief valves" protecting bacteria from lysis upon acute osmotic downshock3. Among them, MscL has been intensively studied since the original identification and characterization 15 years ago by Kung and co-workers4. MscL is reversibly and directly gated by changes in membrane tension. In the open state, MscL forms a nonselective 3 nS-conductance channel which gates at tensions close to the lytic limit of the bacterial membrane. An earlier crystal structure at 3.5 Å resolution of a pentameric MscL from Mycobacterium tuberculosis (TbMscL) represents a closed-state or nonconducting conformation5, 6. MscL has a complex gating behaviour; it exhibits several intermediates between the closed and open states, including one putative nonconductive expanded state and at least three sub-conducting states7. Although our understanding of the closed5, 6 and open8-10 states of MscL has been increasing, little is known about the structures of the intermediate states despite their importance in elucidating the complete gating process of MscL. Here we present the crystal structure of a truncation mutant (95-120) of MscL from Staphylococcus aureus (SaMscL-C 26) at 3.8 Å resolution. Strikingly, SaMscL-C 26 forms a tetrameric channel with both transmembrane (TM) helices tilted away from the membrane normal at angles close to that inferred for the open state9, likely corresponding to a nonconductive but partially expanded intermediate state.

The full-length SaMscL protein is 120 amino acid residues long and shares 40% and 51% homology with TbMscL and the *Escherichia coli* MscL (EcMscL), respectively. Initial

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Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

Author Contributions Z. F. L. designed and performed the experiments in molecular biology, biochemistry, crystallography and the structure analysis. Z.F.L. and C.S.G. conducted the downshock assays. C. S. G. was responsible for the protein reconstitution and electrophysiology. D. C. R. coordinated the project and contributed to the structure analysis. The manuscript was written by Z. F. L, C. S. G and D. C. R.

Author Information The atomic coordinates and structure factors have been deposited at the Protein Data Bank with accession numbers 3HZQ. Reprints and permissions information is available at www.nature.com/reprints.

electrophysiological characterization of SaMscL by Moe *et al*11 established that it exhibited characteristic membrane tension dependent MscL activity with significantly shorter open dwell times compared to EcMscL. Excised patches from azolectin liposomes containing SaMscL-C 26 show a stereotypical pressure dependent increase in channel activity, similar to patches containing wild-type EcMscL, with a single channel conductance comparable to EcMscL (Fig. 1). The open state of SaMscL-C 26 appears more stable than that of wild-type SaMscL (Fig. 1c and d), suggesting a role for the carboxy-terminal truncation in the gating behaviour change. Nevertheless, as shown in supplementary Figure 1, both full-length and C 26 SaMscL are capable of rescuing a mechanosensitive channel knockout strain of *E. coli* cells from osmotic downshock12, indicating that they are both functional *in vivo*. Consequently, truncation of 26 residues from the carboxy-terminus of SaMscL does not impair its function, consistent with previous reports for EcMscL13, 14.

SaMscL-C 26 forms a tetramer in the crystal with the molecular symmetry axis coincident with the crystallographic four-fold axis (supplementary Fig. 2), so that the asymmetric unit contains one subunit. The SaMscL-C 26 tetramer measures \sim 69 Å wide in diameter and \sim 37 Å thick (Fig. 2a, b), while the TbMscL pentamer, omitting carboxyl-terminal residues 102-151 that form a cytoplasmic helical bundle, measures \sim 52 Å wide and \sim 50 Å thick (Fig. 2c, d). Excluding the carboxy-terminal residues which have been deleted in SaMscL-C 26, the major differences between the amino acid sequences of SaMscL and TbMscL are localized to the periplasmic ends of TM1 and TM2 and their connecting loop. In vitro crosslinking experiments indicate that both C 26 and full-length SaMscL are tetrameric in detergent solution and that removal of the carboxy-terminal domain does not change the oligomeric state of SaMscL (supplementary Fig. 3). Although unexpected, it is not unprecedented that certain multimeric membrane proteins form distinct oligomers in different species. For example, the CorA Mg²⁺ transporter forms a pentamer in *Thermotoga* maritima and a tetramer in Bacillus subtilis and E. coli15, while the bacterial lightharvesting complex II may be either octameric or nonameric 16. The ring of c subunits of the membrane embedded F₀-ATPase also exhibits variable stoichiometry17.

The secondary structure of the TM region of SaMscL-C 26 resembles that of TbMscL (Fig. 2e) with a few exceptions. TM1 of SaMscL-C 26 begins at the cytoplasmic surface and extends from Asn 13 to Phe 47, kinking near Pro 41 on the periplasmic side. The polypeptide chain continues through the periplasmic loop from Gly 48 to Gly 64 and crosses the membrane again through TM2 from Leu 65 to Leu 90. TM1 has the same length as in TbMscL, while the periplasmic loop and TM2 are 4 residues and 7 residues shorter respectively, than the corresponding parts of TbMscL. The largest changes in the two structures are the first 12 residues (which adopt an irregular structure in SaMscL-C 26 but a short α -helix in TbMscL) and the detailed conformations of the periplasmic loop. In addition, the intrasubunit crossing angle between TM1 and TM2 changes from 134° in TbMscL to 111° in SaMscL-C 26 (Fig. 3a, b). An overlay of TbMscL and SaMscL-C 26 monomers on TM1s suggests a counterclockwise pivoting of TM2 away from the membrane normal and a translation towards the periplasmic surface (Fig. 3a). In both structures, TM1 and TM2 from the adjacent subunit (TM2') form an antiparallel pair of helices with crossing angle = 169/185° in SaMscL-C 26/TbMscL. The TM1-TM2' pair of SaMscL-C 26 aligns

well with that of TbMscL (root mean square deviation of α -carbon atoms = 1.5 Å) indicating this pair might move as a rigid body during the conformational change (Fig. 3c and ref.18, 19). The extensive network of largely conserved hydrophobic residues at the TM1-TM2' interface presumably reflects the stability of this antiparallel helical pair (supplementary Fig. 4).

The SaMscL-C 26 tetramer is \sim 13 Å thinner along the membrane normal than the membrane spanning domain of TbMscL pentamer but up to 17 Å wider on the periplasmic surface. The periplasmic surface area of tetrameric SaMscL-C 26 is \sim 2200 Å², while that of pentameric TbMscL is ~2000 Å². As such, the membrane spanning region of SaMscL-C 26 tetramer is expanded within the membrane plane and compressed along the pore axis as compared to the TbMscL pentamer (Fig. 2 a-d). This expansion is also evident in the pore profiles of both channels (Fig. 3d). The constriction of SaMscL-C 26 at Val 21 is widened to ~ 6 Å diameter as compared to ~ 3 Å in TbMscL. Ala 25 and Leu 17, positioned to either side of Val 21 in the permeation pathway, move further away from the central axis, opening the pore to ~ 8 Å around those residues. The corresponding residues in TbMscL, Thr 25 and Leu 17, respectively, lie closer to the axis and hence the pore diameter in this structure narrows to ~ 4 Å in this region. Although the constriction in SaMscL-C 26 is wider than in TbMscL, theoretical studies indicate hydrophobic pores with diameters less than 9 Å and 13 Å are impermeable to water and ions20, respectively, implying that both structures are likely non-conducting since they each have narrow and hydrophobic pore constrictions. The expanded state of SaMscL-C 26 channel reflects the higher tilt angles of TM1 and TM2, relative to the membrane normal. In SaMscL-C 26, TM1 and TM2 are tilted 49° and 59° with respect to the pore axis, significantly larger than the 36-38° for the TM helices of TbMscL6, and close to the angles in an open-state model of EcMscL (~51° for TM1/ TM2)9. Consequently, the crossing angle between the axes of two adjacent TM1s in SaMscL-C 26 (-63°) changes by \sim -21° relative to -42° in TbMscL. The combination of the constricted pore and expanded conformation suggests SaMscL-C 26 represents an intermediate state during the transition from the resting closed state to the sub-conducting or open states.

The SaMscL-C 26 structure is clearly expanded in the crystal, but why was this conformation trapped? It is possible that removal of 26 residues from the carboxy-terminus lowers the energy barrier for the transition from the closed to expanded state and crystal packing facilitates this specific expanded conformation. There are several lines of evidence consistent with this idea. Examination of the TbMscL structure suggests that the carboxy-terminal bundle acts as a plug or pre-filter to the permeation pathway which may dissociate to permit the passage of large solutes, including small proteins21-23, through the channel. This is consistent with recent electron microscopy work, which showed that a pore is present at the center of a gain-of-function mutant (G22N) of EcMscL, whereas the cytoplasmic protrusion is absent24. We suggest that the carboxy-terminal helix bundle stabilizes the channel in the closed state and restrains the TMs from tilting into the expanded conformation. The dissociation of this bundle triggered by the mechanical force transmitted from the lipid bilayer removes the restraint, allowing the TMs to tilt and transit to the expanded state. Indeed, SaMscL-C 26 reconstituted in liposomes is more stable in the open state than the wild-type channel, which displays fast flickering from the closed to open

state11 (Fig. 1c and d), suggesting that the truncation mutant may sample the expanded state more readily under crystallization conditions. The relatively small detergent micelle of n-dodecyl-N,N-dimethylamine-N-oxide (LDAO)25 might also favor the expanded state by exerting curvature stress on the channel.

To model the gating process of MscL, a series of structural models of tetrameric SaMscL from closed to open state were constructed using relationships between the helix tilting angles (η), interhelical-crossing angles (α) and the minimum pore radius (R) of the oligometric proteins 26. We use equations $\cos \alpha = \cos^2 \eta + \sin^2 \eta \cos \theta$ and $R = d(\tan \eta \cot(\alpha/2))$ (-1)/2, where $\theta = 360^{\circ}/N$ with N = 4 or 5 for a tetramer or pentamer and d is the helix diameter 26, as the basis for modelling a series of tetrameric structures with the α angles determined from the pentameric structures at different states. Here we propose a two step helix-pivoting model of SaMscL gating. In this model, it is assumed that the TM1-TM2' pair rotates and shifts as a rigid body18, 19. The first pivoting step occurs during the transition from the closed state to a state near the expanded intermediate (Fig. 4a, b). TM1 pivots clockwise about Val 21 by $\sim 22^{\circ}$ and slides along the surface of TM2 (of the same subunit) from Gln 68 down to Ile 75. This induces TM2 to rotate counterclockwise about the pore axis and incline toward the membrane plane along with TM1. Increasing the TM1 tilt angle requires that the helices move farther apart to maintain a symmetric oligomer. As TM2 is in close interaction with TM1 from the next subunit, these movements are synchronized throughout the tetramer. The crossing angle between two adjacent TM1s increases from -42° to -68° at this stage. This angle reaches -67° in a previous open-state model8, 18 and -55° in an alternative model9, bracketing the value (-63°) observed in the expanded state structure, suggesting that it is likely near the edge of the open state. At the end of this first step, the periplasmic loop becomes fully stretched between intrasubunit TM1 and TM2, and if tilting continues, the channel would become thinner than the membrane.

The constraint of the membrane thickness leads to a proposed second pivoting step, defined as a counterclockwise rotation of TM1-TM2' pair about an axis running normal to the membrane and through Gly 48 near the carboxy-terminal end of TM1. The helices rotate by 16° (Fig. 4a and c). The highly conserved Gly 48 (Fig. 2e) appears well suited to serve as the pivot for the outward swinging of the TM1-TM2' pair. This second-step action expands the pore diameter to 22 Å, compatible with the observed large conductance. In MscS, the pore-lining helix also pivots around a glycine residue and moves outwards from the central axis to expand the pore constriction27. The periplasmic loop of MscL provides the flexibility for the substantial relative movements of the TM1 and TM2 observed within the same subunit, but also defines the steric limit of channel expansion in the fully open state. Previous functional studies on the periplasmic loop suggested it may act as a spring, influencing channel kinetics and mechanosensitivity13, 28-30. During the first step of this transition, the periplasmic surface of the SaMscL expands significantly while the channel constriction remains essentially closed until the second step takes place (Fig. 4a, d).

In summary, we have determined the crystal structure of a carboxy-terminal truncation mutant of *S. aureus* MscL and demonstrated that SaMscL forms a tetrameric pressure gated channel. This structure represents a likely expanded intermediate conformation of MscL

between the resting closed state and the open state. A two step helix-pivoting model of the gating mechanism has been proposed.

Methods Summary

The SaMscL gene was inserted into a pET15b vector with a stop codon at Glu 95 introduced by site-directed mutagenesis to generate the SaMscL-C 26 mutant. The channel was overexpressed in BL21-Gold cells and purified in the detergent LDAO by immobilized metal affinity and gel filtration chromatographies. Crystals were grown using the hanging-drop vapor diffusion method. The structure was solved by the method of multiple isomorphous replacement with anomalous scattering.

Crosslinking was performed using N,N-dicyclohexylcarbodiimide at room temperature for 30 min-1 h and the products were analyzed by SDS-polyacrylamide gel electrophoresis.

The MJF465 (mscl⁻, mscs⁻, msck⁻) (DE3) strain was used for the osmotic downshock assay. Cells hosting EcMscL and an empty vector were included as positive and negative control sets, respectively. For the osmotic shock, the culture was diluted 1:500 into sterile doubledistilled H₂O. For the mock shock, LB media with 500 mM NaCl was used instead of H₂O.

MscL protein was reconstituted into giant azolectin liposomes via the dehydration/ rehydration method and assayed by patch clamp electrophysiology. Patches containing at least 1-10 channels were excised and currents measured at various pressures. Recording and pipette solutions both contained 200 mM KCl, 40 mM MgCl₂, 5 mM HEPES, pH 7.2.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. SaMscL-C 26 is a pressure sensitive channel

a, Pressure sensitive response of EcMscL and SaMscL-C 26 proteins reconstituted into azolectin. Excised patches were held at +60 mV and pressure clamped from 0 to -40 mm Hg (EcMscL) or -75 mm Hg (SaMscL-C 26). In both cases, channel activity steadily increases with greater negative pressure and decreases when the pressure is removed. **b-d**, Single channel recordings of EcMscL (b), SaMscL-C 26 (c), and wild-type SaMscL (d). EcMscL and SaMscL-C 26 channels open and close with discreet jumps in current. Calculated single channel conductances for EcMscL and SaMscL-C 26 are 2.9 and 2.6 nS respectively. In contrast, single channel SaMscL openings flicker rapidly and are too brief to calculate a conductance, as observed previously11. C denotes the closed state, while O1 and O2 indicate the opening of one and two channels, respectively.

Figure 2. Structures of SaMscL-C 26 and TbMscL

a and **b**, The crystal structure of tetrameric SaMscL-C 26 viewed along the membrane plane and the pore axis (from the periplasmic side), respectively. **c** and **d**, The membrane spanning domain of the TbMscL structure (residues 1-101 of PDB set 2OAR) viewed as in a and b. Val 21 is presented as a stick model while the polypeptide chains are shown as ribbons. **e**, Structure-based sequence alignment of SaMscL, EcMscL and TbMscL. The red arrow indicates the truncation point (Glu 95) of the SaMscL-C 26 mutant.

a, **b**, Intrasubunit conformational change visualized by aligning the two monomeric structures on TM1 and TM2, respectively. **c**, The superposition of intersubunit TM1-TM2' pair from the two structures. **d**, The pore profiles of SaMscL-C 26 and TbMscL along the central axes. The top two surface presentations are the sectional views of SaMscL-C 26 (left) and TbMscL (right) with the atoms around the constriction colored in yellow, red and blue for carbon, oxygen and nitrogen respectively. In the bottom profiles, the regions

corresponding to SaMscL-C 26 and TbMscL are colored yellow and blue, respectively. The residues lining the constriction are labeled in the same color codes.

Figure 4. The two-step helix-pivoting model of SaMscL gating

a, The relationship among TM1 tilting angle η , TM1-TM1' crossing angle α , and the pore radius R at Val 21 derived from the gating models. The * symbol indicates the position of the SaMscL-C 26 structure on the curve. The yellow, cyan and magenta spheres indicate the positions of the resting closed, turning-point intermediate and open states on the curve. S₁-S_n denotes the multiple subconducting states between the expanded and open states. At step 1, η and α are greatly changed due to the helix tilting motion, while R is only slightly increased. As the helices swing away from the pore axis at the step 2, R increases rapidly,

while η and α remain constant. b, The first pivoting step of TM1-TM2' occurs about Val 21 (red point). The pair pivots as a rigid body rotating 22° from the resting closed (yellow, based on TbMscL structure5, 6) to the turning-point intermediate state (cyan, near the SaMscL-C 26 structure). c, The second pivoting step occurs about the red-line axis through Gly 48, from the intermediate to open state (magenta, based on the Sukharev and Guy model8). d, The relationship among η , α and the periplasmic surface area of the models. At step 1, the periplasmic surface area expands more drastically than at step 2.