1 TITLE

2 TMEM16 and OSCA/TMEM63 proteins share a conserved potential to permeate ions and

- 3 phospholipids
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L4 ABSTRACT

The calcium-activated TMEM16 proteins and the mechanosensitive/osmolarity-activated L5 OSCA/TMEM63 proteins belong to the Transmembrane Channel/Scramblase (TCS) superfamily. ۱6 L7 Within the superfamily, OSCA/TMEM63 proteins, as well as TMEM16A and TMEM16B, are thought to function solely as ion channels. However, most TMEM16 members, including TMEM16F, maintain L8 an additional function as scramblases, rapidly exchanging phospholipids between leaflets of the ٤9 membrane. Although recent studies have advanced our understanding of TCS structure-function 20 relationships, the molecular determinants of TCS ion and lipid permeation remain unclear. Here we 21 show that single mutations along the transmembrane helix (TM) 4/6 interface allow non-scrambling 22 TCS members to permeate phospholipids. In particular, this study highlights the key role of TM 4 in 23 controlling TCS ion and lipid permeation and offers novel insights into the evolution of the TCS 24 superfamily, suggesting that, like TMEM16s, the OSCA/TMEM63 family maintains a conserved 25 potential to permeate ions and phospholipids. 26

INTRODUCTION

Ion channels and phospholipid scramblases catalyze the passive flux of ions and 28 phospholipids down their respective chemical gradients. Compared to ion channels, both our 29 understanding of scramblases and the evolutionary origins of phospholipid scrambling are 30 underdeveloped. Discoveries of two scramblase families-the TMEM16 calcium-activated 31 phospholipid scramblases (CaPLSases) and XKR caspase-dependent phospholipid scramblases-32 have revealed roles for scramblases in processes such as blood coagulation¹⁻³, angiogenesis⁴, cell 33 death signaling^{5,6}, phagocytosis⁶, cell migration^{7,8}, membrane repair^{9,10}, microparticle release³, cell-34 cell fusion^{11–13}, and viral infection^{14,15}. Among identified scramblases, TMEM16 CaPLSases are the 35 most extensively studied at the molecular level¹⁶. The TMEM16 family (Fig. S1a) was originally 36 classified as calcium-activated chloride channels (CaCCs) based on the first-discovered members. 37 TMEM16A and TMEM16B^{17–19}. Most remaining members, however, exhibit CaPLSase activity, with 38 TMEM16F representing the canonical CaPLSase²⁰ (Fig. 1a). Uniquely, CaPLSases also function as 39 non-selective ion channels along with their non-specific permeability to phospholipids^{2,21-25}. Substrate 10 11 permeation is facilitated by calcium-induced conformational changes causing the clamshell-like separation of TMs 4 and 6 to facilitate ion and phospholipid permeation^{26–28} (Fig. S1b-d). These 12 conformational changes further catalyzed phospholipid permeation by thinning the membrane near 13 the permeation pathway 29,30 . 14

The TMEM16 family together with the OSCA/TMEM63 family and TMC family^{31–34} collectively form the Transmembrane Channel/Scramblase (TCS) superfamily¹⁶. Despite the conserved 10-TM architectural core^{35–39}, phospholipid permeability has not been experimentally demonstrated in mechanosensitive OSCA/TMEM63 or TMC proteins, raising the intriguing question of how TCS proteins discriminate between ion and ion/phospholipid substrates. Interestingly, studies have shown that the TMEM16A CaCC can be genetically modified to enable phospholipid permeability, either by substituting TMEM16F domains as small as 15 amino acids²³, or through single mutations at the
 hydrophobic gate²⁶. These findings suggest a modest energetic barrier for scramblase activity and
 lead us to hypothesize that evolutionary relatives of the TMEM16 family may maintain a conserved
 potential for both ion and phospholipid permeation.

To test this hypothesis, we introduced single mutations along TM 4 of TMEM16F, TMEM16A, 55 OSCA1.2, and TMEM63A at sites near the corresponding hydrophobic gate of TMEM16F²⁶. 56 Mutations along the TM 4/6 interface in TMEM16F and TMEM16A resulted in constitutive 57 phospholipid scramblase activity. Strikingly, equivalent TM 4 mutations in OSCA1.2 and TMEM63A 58 also converted these ion channels into phospholipid scramblases, which were either constitutive or 59 activated by osmotic stimulation. Individual mutations also resulted in gain-of-function (GOF) ion 50 channel activity, suggesting that these mutations may disrupt a conserved activation gate in these 51 52 evolutionarily related proteins. Together, our findings i) advance the mechanistic understanding of gating and substrate permeation in the TMEM16 and OSCA/TMEM63 families, ii) underscore a key 53 design principle for phospholipid scramblases, and iii) support our hypothesis that the TCS 54 55 superfamily maintains a conserved potential for both ion and phospholipid permeation.

56 **RESULTS**

57 **TM 4 mutations result in a constitutively active TMEM16F scramblase**

⁵⁸Our previous study identified the activation gate of TMEM16F, consisting of hydrophobic ⁵⁹residues from TMs 4, 5, and 6, which collectively govern ion and phospholipid permeation²⁶. ⁷⁰Mutations to the activation gate result in GOF CaPLSases, some of which are constitutively activated ⁷¹without requiring calcium²⁶. The hydrophobic gate represents the most constricted point along the TM ⁷²4/6 interface in TMEM16F and mutations along this interface likely alter substrate permeation and ⁷³gating. Structural and computational studies on the fungal *Nh*TMEM16⁴⁰ and *Af*TMEM16²⁹ orthologs, ⁷⁴as well as human TMEM16K⁴¹ further support a clamshell-like gating model²⁶ whereby scrambling

activity is promoted by separation of the TM 4/6 interface¹⁶ (Fig. S1b-d). The conformational transition 75 involves the N-terminal half of TM 4, which bends away from TM 6, and the C-terminal half of TM 6, 76 which collapses onto the calcium binding sites formed together with TMs 7 and 8¹⁶. Given that the N-77 termini of TM 4 are largely hydrophobic among TMEM16 CaPLSases (Fig. S1f) and mutations at 78 F518 lead to constitutive TMEM16F activity²⁶, we hypothesized that introducing a charged side chain 79 along the TM 4/6 interface might disrupt the hydrophobic gate and result in TMEM16 scramblases 30 with GOF scramblase activity. To test this hypothesis, we overexpressed eGFP-tagged mutant 31 constructs in TMEM16F knockout (KO) HEK293T cells and used an established scramblase 32 assay^{25,26,42} that detects phosphatidylserine (PS) exposure using fluorophore-tagged Annexin V 33 (AnV) as a reporter (Fig. 1b). In the absence of calcium stimulation, PS is predominantly in the inner 34 leaflet of the plasma membrane of TMEM16F wildtype (WT) expressing cells and is therefore not 35 labelled by extracellular AnV. Similar to TMEM16F F518K²⁶, overexpressing the single lysine 36 mutations TMEM16F I521K and M522K led to spontaneous, global exposure of PS on the plasma 37 membrane without requiring calcium stimulation (Fig. 1b, c). In contrast, T526K, which is one helical 38 39 turn below I521 and M522 (Fig. 1a), caused minimal spontaneous PS exposure (Fig. 1b, c). We further tested whether I521K, M522K, and T526K have GOF ion channel activity using whole-cell Э0 patch clamp. In the absence of calcium, I521K and M522K, but not WT TMEM16F or T526K, showed Э1 robust depolarization-activated outward rectifying currents (Fig. 1d-f). Our imaging and patch clamp Э2 experiments thus demonstrate that I521K and M522K are functionally expressed on the plasma ЭЗ membrane and their GOF phospholipid scramblase activity results in spontaneous, global PS Э4 exposure on the cell surface. Conversely, despite apparent plasma membrane localization, T526K Э5 exhibited strongly attenuated GOF ion channel and scramblase activities. The TM4/6 interaction is Э6 Э7 notably less prominent near T526 (Fig. 1a), two helical turns below the hydrophobic gate residue, F518²⁶, (Fig. 1a) and may represent a lower limit for this charge-induced effect. On the other hand, 98

I521 and M522 are approximately one helical turn below the hydrophobic gate, (Fig. 1a) supporting
 the idea that charged mutations along the TM 4/6 interface promote gate opening and substrate
 permeation (Fig. 1g).

Our previous study also showed that glutamate, but not alanine, substitution at F518 led to spontaneous scramblase activity in TMEM16F²⁶. To assess whether alternative side chains cause spontaneous scrambling below the hydrophobic gate in TMEM16F, we tested I521A and I521E. I521E led to spontaneous, global PS exposure in all transfected cells, whereas I521A failed to cause spontaneous PS exposure (Fig. S2a, b). This result is consistent with our previous finding, providing additional support that charged TM 4 mutations disrupt the TM 4/6 interface.

1611K on TM 6 results in a constitutively active TMEM16F scramblase

We next tested whether a TM 6 mutation along the TM 4/6 interface could promote scrambling.)9 Directly adjacent to I521 and M522 are a pair of glycine residues on TM 6 that are proposed to LO function as a hinge during calcium-dependent activation⁴³. We therefore chose to introduce a lysine Ι1 mutation one helical turn above this site at 1611, a residue adjacent to the pore-facing inner gate L2 residue I612²⁶ (Fig. 2a). Comparable to I521K (Fig. 1b, c), I521E (Fig. S2a, b), and M522K (Fig. 1b, L3 c), I611K resulted in spontaneous, global PS exposure (Fig. 2b, c). Likewise, I611K-expressing cells L4 exhibited robust, depolarization-activated outward rectifying currents in the absence of calcium (Fig. ۱5 2d-f). Together with our previous report that I612K is a GOF phospholipid scramblase and ion ۱6 channel²⁶. I611K further supports that charged mutations in TM 6 near the activation gate can also ι7 promote substrate permeation in TMEM16F (Fig. 2g). ٢8

19 TM 4 lysine mutations convert TMEM16A into a constitutively active scramblase

TMEM16A is a CaCC^{17–19} without scramblase activity^{23,26} (Fig. 3a-c). We previously reported that a single TM 4 lysine mutation, L543K, at the hydrophobic gate of TMEM16A (Fig. 3a) allows the CaCC to permeate phospholipids spontaneously, analogous to the F518K mutation in TMEM16F²⁶. <u>23</u> Thus, we reasoned that lysine mutations equivalent to TMEM16F I521K and M522K might also result in spontaneous phospholipid permeability. To test this hypothesis, we overexpressed eGFP-tagged 24 TMEM16A I546K and I547K mutants in TMEM16F KO HEK293T cells and assessed their ability to 25 expose PS on the plasma membrane with confocal microscopy. Similar to TMEM16A L543K²⁶, I546K-26 and I547K-expressing cells exhibit spontaneous, global PS exposure (Fig. 3b, c), analogous to the 27 equivalent TMEM16F mutations I521K and M522K (Fig. S1f, 1b, c). Similarly, whole-cell patch clamp 28 revealed GOF ion channel activity at depolarizing potentials, even in the absence of calcium <u>29</u> stimulation (Fig. 3d-f). In contrast, E551K, equivalent to T526K in TMEM16F, failed to exhibit PS 30 exposure, but showed modest ion channel activity at depolarizing potentials in the absence of 31 calcium, perhaps reflecting the gate is open sufficiently for ion but not phospholipid permeation. In 32 total, these results suggest that TM 4 mutations in this region destabilize interface to endow the 33 34 TMEM16A CaCC with GOF ion channel activity, and, in the case of I546K and I547K, constitutive phospholipid permeability (Fig. 3g). 35

Ani9, a selective TMEM16A inhibitor, attenuates I546K ion and phospholipid permeation

We then tested whether Ani9, a selective, extracellular TMEM16A inhibitor⁴⁴, could attenuate phospholipid and ion permeation through I546K. Similar to our previous observation of TMEM16A L543K²⁶, chronic incubation of Ani9 at 10 µM also prevented spontaneous PS exposure for I546K (Fig. S3d). Interestingly, 10 µM Ani9 reversibly inhibited voltage-elicited TMEM16A I546K currents by over 50% (Fig. S3a-c). These results demonstrate that although the TM 4 GOF mutations alter TMEM16A substrate permeation, they retain their sensitivity to Ani9 inhibition (Fig. S3e). This further supports that the TM 4 GOF mutations of TMEM16A mediate the spontaneous, global PS exposure.

L438K on TM 4 converts OSCA1.2 into a constitutively active scramblase

We next applied our approach to a TMEM16 relative from the OSCA/TMEM63 family (Fig.
 S1a), which was first discovered in plants as a family of mechanosensitive and osmolarity-activated

cation non-selective ion channels⁴⁵ (Fig. 4a). We hypothesized that analogous lysine mutations on 17 TM 4 along the TM 4/6 interface of OSCA/TMEM63 proteins would result in GOF channels that may 18 also permeate phospholipids. Within the family, we chose OSCA1.2 from Arabidopsis thaliana due to 19 previous structural³⁵ and biophysical⁴⁶ characterization demonstrating that the channel is activated 50 directly by membrane tension. Overexpressing eGFP-tagged OSCA1.2 WT in TMEM16F KO 51 HEK293T cells did not induce PS exposure, demonstrating that OSCA1.2 WT lacks spontaneous 52 phospholipid scramblase activity (Fig. 4b). We then introduced TM 4 lysine mutations at analogous 53 sites, as identified by structural (Fig. 4a) and sequence alignment (Fig. S1f). Strikingly, a single point 54 mutation, L438K, causes cells overexpressing the OSCA1.2 mutant to exhibit spontaneous and 55 alobal PS exposure (Fig. 4b, c). This mirrors our results with both TMEM16F (Fig. 1) and TMEM16A 56 (Fig. 3), suggesting that the L438K mutation allows the OSCA1.2 channel to permeate phospholipids. 57 58 Next, we used inside-out patch clamp to examine if L438K also enhances OSCA1.2 channel activity. We found that the mutant significantly left-shifts the conductance-voltage (G-V) relationship (Fig. 4e, 59 f) and accelerates channel activation kinetics (Fig. 4g) compared to WT. Under -50 mmHg of 50 pressure, L438K has a half-maximal voltage ($V_{0.5}$) of 66.7±3.7 mV, while the WT $V_{0.5}$ is nearly 51 108.7±5.6 mV (Fig. 4f), underscoring that this mutation also promotes channel gating. Together, 52 these experiments show that, like TMEM16A, a single lysine mutation near the putative gate allows 53 the mechanosensitive and osmolarity-activated OSCA1.2 channel to become spontaneously 54 permeable to phospholipids (Fig. 4h). 55

To help visualize how L438K alters the TM 4/6 interface of OSCA1.2, we created a homology model of the mutant using the Swiss-PDB server and inserted it into a phospholipid membrane with the CHARMM-GUI webserver⁴⁷. After equilibration, we ran 900 ns atomistic molecular dynamics (MD) simulations for both WT and L438K in GROMACS, using a combination of conventional and metadynamic simulations. Remarkably, the L438K trajectory showed considerably increased

hydration of the pore region compared to WT (Fig. S4b). This enhanced hydration is attributed to a
 modest expansion of the TM 4/6 interface throughout the simulation (Fig. S4a). Together with our
 functional results, these MD simulations reinforce our hypothesis that disruption of the TM 4/6
 interface of OSCA1.2 promotes GOF ion and phospholipid permeation.

One helical turn above OSCA1.2 L438, at the corresponding site to the activation gate residue 75 F518 in TMEM16F, is an endogenous lysine at position 435 (Fig. S5a). This positioning places two 76 positively charged residues in close proximity within TM 4 of the OSCA1.2 L438K mutant. Previous 77 studies on model transmembrane helical peptides have demonstrated that hydrophilic side chains 78 within the hydrophobic core of the membrane promote trans-bilayer phospholipid transport and this 79 effect is further enhanced when these hydrophilic residues are stacked within the helix⁴⁸. To explore 30 the role of the endogenous K435 in facilitating L438K-mediated spontaneous phospholipid 31 32 permeability, we assessed PS exposure in the K435L and K435L/L438K mutants. Remarkably, the double mutant K435L/L438K, which shifts the lysine to position 438, resulted in a modest level of 33 spontaneous PS exposure (31%, Fig. S5b, c). This level is intermediate between the WT (K435) with 34 35 no PS exposure and the L438K mutant (K435/K438) with 63% spontaneous PS exposure (Fig. 4c/Fig. S5b, c). In contrast, K435L (L435/L438) did not induce significant spontaneous PS exposure 36 (3%, Fig. S5b, c). These findings, consistent with previous studies on model transmembrane 37 peptides⁴⁸, highlight the role of multiple charged resides in TM 4 in promoting phospholipid 38 scrambling. Moreover, our results further support the hypothesis that the instability of the TM 4/6 39 interface in OSCA1.2 is crucial for controlling phospholipid and ion permeation. Э0

A439K on TM4 converts OSCA1.2 into an osmolarity-sensing scramblase

Interestingly, mutating the neighboring amino acid (Fig. 4a), A439, to lysine resulted in minimal
 spontaneous PS exposure (10%, Fig. 4b, c). We thus reasoned that A439K scramblase activity may
 require additional stimulation to trigger scramblase activity. Given that WT OSCA1.2 ion channel

Э5 activity can be induced by hypotonic treatment (Fig. S6), we acutely treated WT- and A439Kexpressing cells with a hypotonic solution (120 mOsm/kg) and assessed scramblase activity using Э6 time-lapse imaging. Indeed, PS exposure was robustly induced for the A439K mutant (Fig. 5b, c) but Э7 not WT (Fig. 5a, c) in response to hypotonic stimulation. Inside-out patch clamp further demonstrated 98 that A439K enhances OSCA1.2 ion channel activity as evidenced by the accelerated activation)9 kinetics (Fig. 4g) and left-shifted *G-V* relationship (Fig. 4f). Our experiments thus indicate that A439K)0 modestly disrupts OSCA1.2 gating and converts the osmolarity-activated ion channel (Fig. S6) into an)1 osmolarity-sensing phospholipid scramblase (Fig. 5d).)2

TM 4 lysine mutations convert TMEM63A into a constitutively active scramblase

Finally, we turned our attention to TMEM63A to further investigate the evolutionary)4 conservation of our observation that TM 4 lysine mutations convert TMEM16 and OSCA members)5)6 into scramblases. TMEM63s represent the mammalian members of the OSCA/TMEM63 family with three members present in humans (TMEM63A-C). Recent structural and functional characterizations)7 indicate that TMEM63s function as mechanosensitive ion channels gated by high-threshold)8)9 membrane tension and, in notable contrast to all other structurally resolved TCS superfamily members, they likely function as monomers^{49–51}. Given their structural homology to TMEM16s and Γ0 OSCA1.2⁴⁹, we hypothesized that TM 4 mutations in TMEM63A would also result in GOF activity. We Ι1 again identified residues near the putative gate by structural (Fig. 6a) and sequence alignment (Fig. 12 S1f), selecting W472 (equivalent to F518 in TMEM16F, L543 in TMEM16A, and K435 in OSCA1.2), L3 S475 (equivalent to I521 in TMEM16F, I546 in TMEM16A, and L438 in OSCA1.2), and A476 L4 (equivalent to M522 in TMEM16F, I547 in TMEM16A, and A439 in OSCA1.2). Indeed, ۱5 overexpressing eGFP-tagged mouse TMEM63A with single lysine mutations at either W472 or S475 ۱6 ι7 led to spontaneous PS exposure (Fig. 6b, c), though the AnV staining revealed punctate rather than global patterns of PS exposure induced by their OSCA1.2 counterparts. Notably, A476K failed to ٢8

٢9 show obvious PS exposure (Fig. 6b, c), even after 120 mOsm/kg hypotonic treatment (Fig. S7b, c), perhaps reflecting differences in mechanical pressure threshold between OSCA1.2 and TMEM63A⁴⁹. 20 To confirm membrane localization and further probe mutant effects, we exploited TMEM63A ion 21 channel function using the cell-attached patch clamp configuration. As TMEM63A exhibits voltage-22 dependent activity under high pressure, we compared mutant and WT I-V relationships at -80 mmHg. <u>23</u> W472K, S475K, and A476K caused marked reductions in V_{0.5} from 122.3±3.5 mV for WT to 95.9±6.3 24 mV, 92.1±8.6 mV, and 96.8±4.2 mV respectively (Fig. 6d-f). By comparison, mock-transfected 25 controls failed to elicit current (Fig. 6d, e). Together, these results are consistent with our 26 observations in TMEM16F, TMEM16A, and OSCA1.2, indicating that single lysine mutations along 27 TM 4 of TMEM63A can facilitate both ion and phospholipid permeation (Fig. 6g). 28

29 DISCUSSION

Mechanistically, our study improves models of TMEM16 substrate permeation and gating. We 30 identified multiple mutations in TMEM16F and TMEM16A that promote phospholipid permeation and 31 cause commensurate changes in ion channel activities. Our findings complement functional 32 33 characterizations of TMEM16F gating mutants where gate destabilization is inversely correlated with side chain hydropathy^{26,52}. Lysine is above only arginine on the hydropathy index and thus likely 34 explains why it readily destabilizes the gate. For instance, F518K exhibits spontaneous PS exposure, 35 even when the calcium binding site is destroyed²⁶, whereas F518H does not exhibit spontaneous PS 36 exposure⁵². Interestingly, the recent TMEM16F F518H structure (Fig. S1e) shows local membrane 37 thinning due in part to unexpected conformational changes in TM 3⁵². Future structural studies are 38 needed to assess whether TMEM16A or OSCA/TMEM63 mutant scramblases also promote 39 membrane thinning and/or conformational rearrangements in TM 3. More broadly, our results 10 highlight an increasingly appreciated design principle of scramblases where polar and charged 11 residues often line a membrane-spanning groove. This observation has been noted for 12

TMEM16^{53,41,54}, Xkr^{55,56}, and opsin⁵⁷ scramblases and should be a key criterion for identifying and
 characterizing new scramblases.

Our findings also advance our understanding of evolutionary relatives of the TCS superfamily¹⁶ 15 and help uncover their roles in human diseases. Although OSCA/TMEM63 proteins are not known to 16 scramble phospholipids and we did not detect obvious hypotonicity-induced PS exposure in cells 17 overexpressing WT OSCA1.2 (Fig. 5a) or TMEM63A (Fig. S7a), we show that single mutations in TM 18 4 of OSCA1.2 and TMEM63A convert these osmolarity-activated and/or mechanosensitive ion 19 50 channels into phospholipid scramblases, similar to our findings with TMEM16A mutants. We thus speculate that the conserved structural architecture in the transmembrane region endows TCS 51 proteins with a potential to scramble phospholipids, though this capability may have been lost by 52 some members during evolution. This hypothesis is especially intriguing given the recent OSCA open 53 54 state structures, which detail dramatic conformational rearrangements of TMs 3-6 leading to a phospholipid-lined (or proteolipidic) pore near TMs 4 and 6⁵⁸. It will be interesting to test whether 55 equivalent mutations can convert transmembrane channel-like (TMC) proteins—the third TCS relative 56 of TMEM16 and OSCA/TMEM63¹⁶—into phospholipid scramblases. TMC1 is best known for its role 57 in auditory sensation, and thus far has mostly been characterized in vivo due to expression difficulties 58 in heterologous systems⁵⁹. However, recent *in vivo* characterization of mouse TMC1 M412K, known 59 as the *Beethoven* mutation, provided an important insight⁶⁰. The deafness-associated mutation is 50 located in TM 4 (Fig. S1f; equivalent of M522 in TMEM16F, I547 in TMEM16A, A439 in OSCA1.2, 51 and A476 in TMEM63A) and results in constitutive PS exposure when expressed in the hair cell 52 membranes of both heterozygous and homozygous mice⁶⁰. This raises the intriguing possibility that 53 the *Beethoven* mutation may enable TMC1 to spontaneously permeate phospholipids, leading to a 54 loss of membrane homeostasis and ultimately, auditory sensation⁶¹. The possibility of converting 55 TMC proteins into phospholipid scramblases should be thoroughly investigated. Additionally, disease-56

- associated mutations in TMEM63 proteins are present along the TM 4/6 interface, such as TMEM63B
- ⁵⁸ T481N⁴⁹. We speculate that introducing more hydropathic side chains along this interface may lead to
- 59 spontaneous ion and or phospholipid permeability, perhaps contributing to underlying
- 70 pathophysiology.
- 71 METHODS

72 Cloning and mutagenesis

- All constructs used a peGFP-N1 vector backbone. GFP mock controls used the empty peGFP-N1
- vector. Wild type sequence and mutation numbers correspond to NCBI: NP_780553.2 (*Mus musculus*
- 75 TMEM16F) with a three amino acid (MQM) N-terminal truncation, NCBI: NP_001229278 (Mus
- 76 musculus TMEM16A), GenBank: AIU34614.1 (Arabidopsis thaliana OSCA1.2), and NCBI:
- NP_001404481.1 (*Mus musculus* TMEM63A). *At*OSCA1.2 and *Mm*TMEM63A cDNAs were
- ⁷⁸ subcloned using In-Fusion Snap Assembly (Takara, #638947). Point mutants were generated by
- 79 PCR site-directed mutagenesis with primers from IDT DNA Technologies. Sequences were confirmed
- 30 by Sanger sequencing (Azenta).

31 Bioinformatics

- 32 The following sequences were obtained from UniProt and aligned using Clustal Omega: Q8BHY3-2
- 33 (*Mm*TMEM16A), Q9NQ90 (*Hs*TMEM16B), Q9BYT9 (*Hs*TMEM16C), Q32M45 (*Hs*TMEM16D),
- 34 Q75V66 (HsTMEM16E), Q6P9J9 (MmTMEM16F), Q6IWH7 (HsTMEM16G), Q9HCE9
- 35 (*Hs*TMEM16H), A1A5B4 (*Hs*TMEM16J), Q9NW15 (*Hs*TMEM16K), C7Z7K1 (*Nh*TMEM16), Q4WA18
- 36 (AfTMEM16), Q9XEA1 (AtOSCA1.1), Q5XEZ5 (AtOSC1.2), B5TYT3 (AtOSCA1.3), A0A097NUQ0
- 37 (AtOSCA1.4), A0A097NUS0 (AtOSCA1.5), A0A097NUP1 (AtOSCA1.6), A0A097NUP8 (AtOSCA1.7),
- 38 A0A097NUQ2 (AtOSCA1.8), A0A097NUQ5 (AtOSCA2.1), A0A097NUS5 (AtOSCA2.2),
- 39 A0A097NUP6 (*At*OSCA2.3), A0A097NUQ3 (*At*OSCA2.4), A0A097NUQ7 (*At*OSCA2.5), Q9C8G5
- 30 (AtOSCA3.1), A0A097NUT0 (AtOSCA4.1), Q91YT8 (MmTMEM63A), Q5T3F8 (HsTMEM63B), and

- 31 Q9P1W3 (*Hs*TMEM63C), Q8R4P5 (*Mm*TMC1), Q8R4P4 (*Mm*TMC2), Q7TQ69 (*Mm*TMC3), Q7TQ65
- 32 (*Mm*TMC4), Q32NZ6 (*Mm*TMC5), Q7TN60 (*Mm*TMC6), Q8C428 (*Mm*TMC7), and Q7TN58
- 33 (*Mm*TMC8). A subset of the alignment was selected for Fig. S1. Structural models were obtained
- ³⁴ from the PDB, aligned, and visualized using Pymol (Schrödinger).
- **Cell culture**
- ³⁶ The HEK293T cell line was authenticated by the Duke Cell Culture Facility. The TMEM16F KO
- HEK293T cell line was generated by the Duke Functional Genomics Core and characterized in
- previous studies^{25,26}. All cells were cultured with DMEM (Gibco, #11995-065) supplemented with 10%
- ³⁹ fetal bovine serum (FBS) (Sigma-Aldrich, # F2442) and 1% penicillin/streptomycin (Gibco, #15-140-
- 122) at 37°C in 5% CO₂-95% air.

)1 Transfection

- ¹² Plasmids were transiently transfected into TMEM16F KO HEK 293T cells by using X-tremeGENE9
- 13 (MilliporeSigma), X-tremeGENE360 (MilliporeSigma), or Lipofectamine 2000 (Thermo Fisher). Media
- vas replaced with calcium free DMEM (Gibco, 21068-028) 3-4 hours after transfection. The
- ¹⁵ transfected cells were imaged or patched 18-24 or 18-48 hours after transfection, respectively.

Fluorescence imaging of scramblase-mediated PS Exposure

A Zeiss 780 inverted confocal microscope was used to monitor scramblase activity in live cells using)7 the methods described in previous publications^{12,25,26}. 18-24 hours after transfection, the cells were)8 incubated in AnV buffer (1:175 dilution of the fluorescently tagged AnV (Biotium) in Hank's balanced)9 salt solution) immediately before and throughout the duration of the imaging experiment without a Γ0 formal incubation period. Spontaneous PS positive cells were readily labeled by fluorescently tagged Ι1 AnV. Results were quantified as a percentage of PS positive cells among all GFP positive cells. For Γ2 osmolarity activation, 2 mM CaCl₂ in ddH₂O was added to the AnV buffer at a 2:1 ratio. The final L3 osmolarity was ~120 mOsm/kg as measured by a micro-osmometer (Advanced Instrument). Cells L4

overexpressing WT or mutant were treated with low osmolarity AnV buffer, and the scramblase
 activity was measured by recording fluorescent AnV surface accumulation at 5 (Fig. 5) or 7 (Fig. S7)
 second intervals. A custom MATLAB code was used to quantify AnV signal and is available at Github

18 (https://github.com/yanghuanghe/scrambling_activity).

19 Electrophysiology

All electrophysiology recordings were conducted using an Axopatch 200B amplifier with the signal digitally sampled at 10 kHz using an Axon Digidata 1550A (Molecular Devices, Inc.). All

electrophysiology recordings were carried out at room temperature 18-48 hours after transfection.

23 Glass pipettes were pulled from borosilicate capillaries (Sutter Instruments) and fire-polished using a

24 microforge (Narishige). Pipettes had resistances of $2-4 \Box M\Omega$ in physiological bath solution.

Inside-out patch clamp recordings. The pipette solution (external) contained 140 mM NaCl. 25 10 mM HEPES, 2 mM MgCl₂, adjusted to pH 7.3 (NaOH), and the bath solution (internal) contained 26 140 mM NaCl, 10 mM HEPES, 5 mM EGTA, adjusted to pH 7.3 (NaOH). OSCA1.2 WT and 27 mutations were held at constant pressure administrated using a syringe calibrated with a manometer. 28 similar to a previous study⁶². In our experience, the constant pressure and variable voltage protocol <u>29</u> achieved more consistent measurements. Patches were held at a membrane potential of -60 mV 30 and at the indicated pressure, then stimulated using the indicated voltage protocol, taking advantage 31 of the larger OSCA1.2 currents elicited by depolarizing potentials. 32

Cell-attached patch clamp recordings. For TMEM63 recordings, cell-attach mode was used
 instead of inside-out to avoid breaking the giga-ohm seal under higher applied pressure and voltage.
 The bath solution contained (in mM): 140 KCI, 10 HEPES, 2 MgCl₂, 10 glucose, pH 7.3 adjusted with
 KOH. The pipette solution contained (in mM): 130 NaCl, 5 KCl, 10 HEPES, 10 TEA-Cl, 1 CaCl₂, 1
 MgCl₂, pH 7.3 (with NaOH). The mechano-activated current was evoked with a 200 ms pressure
 pulse at -80 mmHg using a high-speed pressure clamp system (HSPC-1, ALA Scientific Instruments,

Farmingdale, NY). The membrane potential inside the patch was held at -60 mV. The voltage pulse alone was run first followed by voltage pulse with pressure. The mechanosensitive current was obtained by subtracting the voltage pulse from the voltage pulse with pressure.

Whole cell TMEM16 patch clamp recordings. The pipette solution (internal) contained 140
mM□CsCl, 10 mM□HEPES, 5 mM□EGTA, adjusted to pH 7.3 (CsOH), and the bath solution
(external) contained 140 mM□NaCl, 10 mM HEPES, 5 mM□EGTA, adjusted to pH 7.3 (NaOH).
Patches were held at a membrane potential of -60□mV, then stimulated using the indicated voltage
protocol.

Whole cell hypotonic patch clamp recordings. The pipette solution (internal) contained 140 17 mM Na gluconate, 10 mM HEPES, 1 mM MgCl₂, and 0.2 mM EGTA, adjusted to pH 7.3 (NaOH). 18 Cells seeded on a small section of cover glass were placed in bath solution (external) containing 140 19 mM Na gluconate, 10 mM HEPES, 1 mM MgCl₂, and 0.2 mM EGTA adjusted to pH 7.3 (NaOH). 50 Patches were held at a membrane potential of -60 mV, then stimulated using the indicated voltage 51 protocol using a 2-second sweep protocol. After 5 sweeps, ddH₂O was slowly and gently added to the 52 53 bath at a ratio of 2:1 using a hypodermic needle to induce hypotonic cell swelling. The sweep protocol continued for a minimum of 4 minutes after hypotonic stimulation. 54

55 Data analysis for electrophysiology

59

All data analysis was performed using Clampfit (Molecular Devices), Excel (Microsoft), MATLAB
 (MathWorks), and Prism softwares (GraphPad). Individual *G-V* curves were fitted with a Boltzmann
 function,

$$G = \frac{G_{max}}{1 + e^{\frac{-ZF(V - V_{0.5})}{RT}}}$$

where G_{max} denotes the fitted value for maximal conductance, $V_{0.5}$ denotes the voltage of half maximal activation of conductance, *Z* denotes the net charge moved across the membrane during the transition from the closed to the open state, and *F* denotes the Faraday constant.

(2)

53 *Model building and MD simulations*

The initial model of OSCA1.2 for atomistic MD simulations and homology modeling was constructed 54 based on PDB: 6MGV. Missing segments were modeled using the Swiss-PDB server 55 (https://swissmodel.expasy.org/). The OSCA1.2 L438K mutation structure was modeled using 6MGV 56 as the template, and the RMSD between the mutation and the original structure of 6MGV was less 57 than 0.5 Å. To mimic the state of transmembrane proteins in the plasma membrane, the model was 58 inserted into a lipid membrane consisting of POPC (exoplasmic leaflet) and 1:1 POPS/POPE 59 (cytoplasmic leaflet) using the CHARMM-GUI web server⁴⁷. To neutralize charge levels in the 70 simulation system, 150 mM KCl solution was added. The final simulation boxes contained ~600 lipid 71 molecules, with ~290.000 atoms. During the simulation, the CHARMM36m all-atom force field was 72 used. The temperature was maintained at 310 K to maintain the fluidity of lipids. During the atomistic 73 simulation, long-range electrostatic interactions were described by the Particle Mesh Ewald (PME) 74 algorithm with a cutoff of 12 Å. Van der Waals interactions were cut off at 12 Å and the MD time step 75 was set at 2 fs. All atomistic MD simulations were conducted using GROMACS-2022.3 software 76 77 package, and enhanced sampling MD simulations were conducted using GROMACS-2022.3 with the COLVARS module. Before the production process, the system performed 5,000 steps of energy 78 minimization. Subsequently, all systems were equilibrated using the default equilibration parameters 79 provided by CHARMM-GUI. After equilibration, the velocities of all atoms were randomly generated, 30 and the product simulation system consisted of a 2-stage MD simulation. In step 1, the system used a 31 conventional MD simulation run for 200 ns. In step 2, the system used a metadynamics simulation run 32 for ~700 ns. During enhanced sampling MD simulations, the sampling interval of the collective 33 variable was adjusted to adapt the lipid movement trajectory. VMD and PYMOL were used to analyze 34 35 simulation trajectories and water occupancy.

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22 AUTHOR CONTRIBUTIONS

- H.Y. and Y.Z. conceived and H.Y. supervised the project. Y.Z. and A.J.L. imaging. P.L. and A.J.L.
- electrophysiology. A.J.L. sequence and structure alignments. A.J.L. and Y.C.S.W. cloning and
- ²⁵ mutagenesis. Z.P. plasmids. Y.Z. MATLAB codes. M.S. MD simulations, supervised by Y.Z. A.J.L.
- and H.Y. drafted the manuscript with input from all authors.

27 **COMPETING INTERESTS**

²⁸ The authors declare no competing interests.

FIGURES







35	HEK293T cells expressing eGFP-tagged TMEM16F wildtype (WT), I521K, M522K, and T526K
36	(center column). CF 594-conjugated Annexin V (AnV, right column) labelled PS exposing cells. BF
37	denotes bright field images (left column). (c) Quantification of the percentage of cells with AnV
38	labelling for TMEM16F WT (n=5), I521K (n=5), M522K (n=5), and T526K (n=5) transfected cells.
39	Values were derived from images of biological replicates, with error bars representing standard error
10	of the mean (SEM). Statistical comparisons to T526K were done using unpaired t-tests with Welch's
11	correction (ns: p>0.05, ****: p<0.0001). (d) Representative current recordings and (e) current-voltage
12	(I-V) relationships of whole cell patches from TMEM16F KO HEK293T cells expressing eGFP-tagged
43	TMEM16F WT (n=6), I521K (n=5), M522K (n=5), and T526K (n=7). Currents were elicited by the
14	voltage protocol shown with the pipette solution containing 5 mM EGTA. Dotted line denotes zero
15	current. (f) Quantification of current at +160 mV. Currents in (e) and (f) were normalized to cell
16	capacitance with the mean \pm SEM calculated from independent patches. Statistical comparisons to
17	T526K were done using unpaired t-tests with Welch's correction (*: p<0.05, **: p<0.01). (g) Lysine
18	mutations along TM 4 in TMEM16F enable spontaneous phospholipid permeation in the absence of
19	calcium.



50

Figure 2: I611K on TM 6 enables TMEM16F channel and scramblase activities in the absence
of calcium stimulation. (a) I611 highlighted on the TMEM16F CaPLSase structure with side chains
shown as yellow sticks (PDB 6QPB). (b) Representative images of TMEM16F knockout (KO)
HEK293T cells expressing eGFP-tagged TMEM16F wildtype (WT) and I611K (center column). CF
594-conjugated Annexin V (AnV, right column) labelled PS exposing cells. BF denotes bright field
images (left column). (c) Quantification of the percentage of cells with AnV labelling for TMEM16F WT
(n=5) and I611K (n=5) transfected cells. Values were derived from images of biological replicates,

58	with error bars representing standard error of the mean (SEM). Statistical comparison was done using
59	an unpaired t-test with Welch's correction (****: p<0.0001). (d) Representative current recordings and
50	(e) current-voltage (I-V) relationships of whole cell patches from TMEM16F KO HEK293T cells
51	expressing eGFP-tagged TMEM16F WT (n=6) and I611K (n=5). Currents were elicited by the voltage
52	protocol shown with the pipette solution containing 5 mM EGTA. Dotted line denotes zero current. (f)
53	Quantification of current at +160 mV. Currents in (e) and (f) were normalized to cell capacitance with
54	the mean \pm SEM calculated from independent patches. Statistical comparison was done using an
55	unpaired t-test with Welch's correction (***: p<0.001). (g) A lysine mutation on TM 6 in TMEM16F
56	enables spontaneous phospholipid permeation in the absence of calcium.



57

Figure 3: Lysine mutations along TM 4 enable TMEM16A channel and scramblase activities in
the absence of calcium stimulation. (a) Top: TMEM16A is a calcium-activated chloride channel
Bottom: TM 4 mutant locations mapped on TMEM16A CaCC structure with side chains shown as
yellow sticks (PDB 5OYG). (b) Representative images of TMEM16F knockout (KO) HEK293T cells
expressing eGFP-tagged TMEM16A wildtype (WT), I546K, I547K, and E551K (center column). CF

73	594-conjugated annexin V (AnV, right column) labelled PS exposing cells. BF denotes bright field
74	images (left column). (c) Quantification of the percentage of cells with AnV labelling for TMEM16A WT
75	(n=4), I546K (n=4), I547K (n=4), and E551K (n=5) transfected cells. Values were derived from images
76	of biological replicates, with error bars representing the standard error of the mean (SEM). Statistical
77	comparisons were done using unpaired t-tests with Welch's correction (*: p<0.05, **: p<0.01). (d)
78	Representative whole-cell current recordings and (e) current-voltage (I-V) relationships of whole cell
79	patches from TMEM16F KO HEK293T cells expressing eGFP-tagged TMEM16A WT (n=14), I546K
30	(n=5), I547K (n=5), and E551K (n=6). Currents were elicited by the voltage protocol shown with the
31	pipette containing 5 mM EGTA. Dotted line denotes zero current. (f) Quantification of current at +160
32	mV. Currents in (e) and (f) were normalized to cell capacitance with the mean \pm SEM calculated from
33	independent patches. Statistical comparisons were done using unpaired t-tests with Welch's
34	correction (*: p<0.05, ****: p<0.0001). (g) Lysine mutations along TM 4 in TMEM16A enable
35	spontaneous phospholipid permeation in the absence of calcium.

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ЭЗ	(right column) labelled PS exposing cells. BF denotes bright field images (left column). Asterisk
€	highlights a PS positive cell for the A439K mutant. (c) Quantification of the percentage of cells with
) 5	AnV labelling for OSCA1.2 WT (n=4), L438K (n=7), and A439K-transfected cells (n=6). Statistical
) 6	comparisons were conducted with unpaired t-tests with Welch's correction (*: p<0.05, ****: p<0.0001).
) 7	(d) Representative current recordings and (e) normalized conductance-voltage (G-V) relationships of
98	inside-out patches from TMEM16F KO HEK293T cells expressing eGFP-tagged OSCA1.2 WT (n=8),
) 9	L438K (n=8), and A439K (n=6). Currents were elicited by the voltage protocol shown next to the listed
)0	pressures. Dotted lines denote zero current. (f) Quantification of half-maximal voltage at -50 mmHg
)1	for WT (109 mV), L438K (67 mV), and A439K (63 mV). Error bars represent standard error of the
)2	mean (SEM) calculated from independent patches. Statistical comparison was conducted with
)3	unpaired t-tests with Welch's correction (***: p<0.001, ****: p<0.0001). (g) Quantification of activation
)4	$ au_{ m on}$ at -50 mmHg and 160 mV for WT (41 ms), L438K (13 ms), and A439K (16 ms). Error bars
)5	represent standard error of the mean (SEM) calculated from independent patches. Statistical
)6	comparison was conducted with unpaired t-tests with Welch's correction (***: p<0.001, ****:
)7	p<0.0001). (h) A lysine mutation along TM 4 converts the OSCA1.2 channel into a phospholipid
)8	scramblase with spontaneous phospholipid permeability.









18 Figure 6: Lysine mutations along TM 4 enable TMEM63A channel and scramblase activities.

(a)Top: TMEM63A is an ion channel gated by high threshold membrane tension. Bottom: the TM 4/6

- 20 interface of HsTMEM63A (PDB 8GRS) with key residues shown as yellow sticks using amino acid
- numbering corresponding to the mouse ortholog. (b) Representative images of TMEM16F KO
- 12 HEK293T cells expressing eGFP-tagged TMEM63A WT, W472K, S475K, and A476K (center column).
- ²³ CF 594-conjugated AnV (right column) labelled PS exposing cells. BF denotes bright field images (left

24	column). (c) Quantification of the percentage of cells with AnV labelling for TMEM63A WT (n=4),
25	W472K (n=4), S475K (n =4), and A476K-transfected cells (n=4). Statistical comparisons were
26	conducted with unpaired t-tests with Welch's correction (**: p<0.01, ****: p<0.0001). (d)
<u>2</u> 7	Representative current recordings and (e) normalized conductance-voltage (I-V) relationships of cell
28	attached patches from TMEM16F KO HEK293T cells expressing either eGFP mock-transfected (n=7)
<u>29</u>	or eGFP-tagged TMEM63A WT (n=7), W472K (n=7), S475K (n=5), or A476K (n=7). Currents
30	represent the subtraction of voltage alone from currents elicited by the voltage and pressure protocols
31	shown. Dotted line denotes zero current. Note that the mock control was normalized to the mean
32	maximal current elicited from WT-transfected cells. (f) Quantification of half-maximal voltage at -80
33	mmHg for WT (122 mV), W472K (96 mV), S475K (92 mV), and A476K (97 mV). Error bars represent
34	standard error of the mean (SEM) calculated from independent patches. Statistical comparison was
35	conducted with unpaired t-tests with Welch's correction (*: p<0.05, **: p<0.01, ***: p<0.001). (g)
36	Lysine mutations along TM 4 in TMEM63A enable spontaneous phospholipid permeability.