Structure of the mammalian TRPM7, a magnesium channel required during embryonic development

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The transient receptor potential ion channel subfamily M, member 7 (TRPM7), is a ubiquitously expressed protein that is required for mouse embryonic development. TRPM7 contains both an ion channel and an α-kinase. The channel domain comprises a non-selective cation channel with notable permeability to Mg2+ and Zn2+. Here, we report the closed state structures of the mouse TRPM7 channel domain in three different ionic conditions to overall resolutions of 3.3, 3.7, and 4.1 Å. The structures reveal key residues for an ion binding site in the selectivity filter, with proposed partially hydrated Mg2+ ions occupying the center of the conduction pore. In high [Mg2+], a prominent external disulfide bond is found in the pore helix, which is essential for ion channel function. Our results provide a structural framework for understanding the TRPM1/3/6/7 subfamily and extend the knowledge base upon which to study the diversity and evolution of TRP channels.

Among the more than 300 mammalian ion channels are two unique members, TRPM6 and TRPM7, that are ion channels and protein kinases (chanzymes) (1–3). TRPM6 and TRPM7 are ubiquitously expressed with prominent roles in early embryonic development (4–6). Tissue-specific knockout mice indicate that TRPM7 is required for normal organ development of the heart (7) and kidneys (4).

Similar to other TRPs, TRPM6 and TRPM7 comprise tetrameric ion channels with each subunit containing six-transmembrane (6-TM) segments. However, unlike most TRP channels, they are permeable to several divalent cations, including Zn2+, Mg2+, and Ca2+, as well as common monovalent cations such as Na+ and K+ (8–11). As chanzymes, the functions of TRPM6 and TRPM7 are correlated with proteolytic cleavage of the kinase domain, which is then translocated to the nucleus to phosphorylate histones and regulate gene expression (12–14). The majority of native TRPM6 and TRPM7 cellular protein is in unique intracellular vesicles (15–17). However, since these vesicles are inaccessible to patch clamp recording, all recordings are from channels in the plasma membrane. In addition to relative ion selectivities and biophysical properties, electrophysiological recordings have demonstrated that TRPM7 inactivation/deactivation is enhanced by phosphatidylinositol 4,5 bisphosphate [PI(4.5)P2] hydrolysis (18), intracellular Mg2+ (9), and intracellular acidification (19). Activators of a TRPM7 divalent cation current are unknown. However, removal of external divalent cations (9), acidification of the external environment (20), or addition of glutathione (17) potentiates TRPM7 currents by enhancing monovalent cation permeability.

Kuriyan and coworkers crystallized and solved the structure of TRPM7’s C-terminal isolated kinase domain (21). The kinase’s NT is similar to classical protein kinases, but its C-terminal lobe resembles ATP-grasp fold enzymes. The aspartic acid and glutamine residues important for catalysis are separated by one residue on a continuous kinked β-strand, with a critical lysine forming hydrogen bonds with the α-phosphate group and the adenine ring of ATP. A Zn2+ appears to be a resident stabilizing cation in the hydrophobic core of the C-terminal lobe. TRPM7’s isolated C-terminal coiled-coil domain was proposed to participate in tetrameric chazyme assembly by bundling the membrane-proximal C-terminal ~50 amino acids (22).

Based on sequence homology, the TRPMs can be divided into two subfamilies as TRPM2/4/5/8 and TRPM1/3/6/7 (23). This familial divide is significant because TRPM6/7 are permeable to divalent cations (including Zn2+ and Mg2+) whereas TRPM4/5 are monovalent cation-selective channels. Cryo-EM structures of human TRPM4, bird TRPM8, and sea anemone TRPM2 have been solved recently (24–27); however, none are available for the TRPM1/3/6/7 subfamily. In this study, we report cryo-EM structures of TRPM7 in EDTA (TRPM7-EDTA, to remove Ca2+–EDTA; TRPM7-Mg2+; and TRPM7-DVF have been deposited in the Electron Microscopy Data Bank under accession codes EMD-6795–EMD-7297–EMD-7298, respectively. Atomic coordinates of the mouse TRPM7-EDTA, TRPM7-Mg2+ and TRPM7-DVF have been deposited in the Protein Data Bank, www.pdb.org, under accession codes 5XZ5, 6BWF, and 6BWH, respectively.

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most free divalent cations), in high external Mg$^{2+}$ (TRPM7-Mg$^{2+}$), and in buffer without added external Mg$^{2+}$ (TRPM7-divalent-free: DVF), to overall resolutions of 3.3, 3.7, and 4.1 Å, respectively.

**Overall Structure of the Mouse TRPM7 Tetramer Ion Channel**

We screened orthologs of TRPM7 and determined that mouse TRPM7 was a promising candidate for our structural studies. To improve the stability and homogeneity of mouse TRPM7, residues C-terminal to natural cleavage sites (12) (almost always 1281–1863) were removed, including the previously characterized kinase domain (21) (almost always 1580–1863). Maltose binding protein (MBP) was fused to the N terminus of TRPM7, and the fusion protein was expressed in the BacMam expression system (*Methods*). The MBP tag was not removed before structural determination, since its cleavage significantly reduced protein yield and stability. Purified and detergent-solubilized protein was exchanged with the amphiphil, PMAL-C8 for structure determination (*SI Appendix, Fig. S1 A and B*).

To evaluate the functional properties of our tagged and truncated TRPM7 construct without endogenous TRPM6/7 contamination [TRPM6 can heteromultimerize with TRPM7 (10)], we patch-clamped transiently transfected HEK293T cells in which TRPM6 and TRPM7 were both deleted (13). The resulting current-voltage (I–V) relationship elicited from a 200-mv ramp protocol (8, 9, 11, 28) displayed several TRPM7 characteristics, including a small and linear inward current, reversal near 0 mV, and a prominent outwardly rectifying current (*SI Appendix, Fig. S1 C and D*). Whole-cell currents also exhibited dialysis-induced growth (*SI Appendix, Fig. S1 E and F*), with a mean peak time (113 ± 21 s at +100 mV; n = 7) within the range of full-length TRPM7 (9). Moreover, as with full-length TRPM7 (8), currents activated rapidly and did not inactivate over a 500-ms test pulse; however, some current decay was observed at voltages positive to +80 mV (*SI Appendix, Fig. S1G*). Divalent cations are proposed to permeate TRPM7 through an anomalous mole fraction mechanism in which they also act as pore blockers (9). We therefore determined whether our truncated construct was blocked by high concentrations of external Mg$^{2+}$ and observed a substantial reduction of outward current with minimal change to inward current (*SI Appendix, Fig. S1 H and I*), analogous to full-length TRPM7 (9, 28). As our truncated construct is suitable for evaluating the Mg$^{2+}$ regulatory/permeability properties of TRPM7, we solved structures of TRPM7-EDTA, TRPM7-Mg$^{2+}$, and TRPM7-DVF using cryo-EM. Resolutions were estimated using the gold-standard Fourier shell correlation cutoff of 0.143 (*SI Appendix,* Fig. S2 and Table S1). We will primarily discuss TRPM7-Mg$^{2+}$, which represents the main features of all three structures.

The tetramer structure of TRPM7-Mg$^{2+}$ surrounds an ion channel pore which empties into a ~50 Å × 50 Å cytoplasmic space created by an ~100-Å-tall, ~120-Å-wide intracellular skirt (Fig. 1 A and B). Our density map was sufficient to enable tracing of the complete polypeptide chain, guided by strong densities for the bulky side chains in Arg, Lys, His, Trp, Tyr, and Phe (*SI Appendix, Fig. S3*). Despite <20% protein sequence identity between TRPM7 and other solved TRP channels [TRPVs (29, 30), polyeystin-2 (31), TRPLML1 (32–34) and TRPA1 (35)], the TRPM7 structure exhibits the fourfold symmetry and domain-swapping features of these and other 6- TM channels, such as the well-studied voltage-gated K$^+$ channels. The orientation and position of helices in the 6-TM bundles of TRPM7 are quite similar to other TRP channel structures (*SI Appendix, Fig. S4*). A short helix between S2 and S3 (“S2–S3 helix”) lies parallel to the intracellular lipid bilayer located close to the amphipathic TRP domain. This resembles that of other TRP structures, but is shorter by one helical turn compared with TRPM2 and TRPM4 (black boxes in Fig. 1C and *SI Appendix, Fig. S5 A and B*). We speculate that these partially membrane-embedded orthogonal helices may affect gating of the channel upon changes in bilayer composition and thickness.

Compared with other TRPM channels, TRPM7’s long N terminus is ~30% conserved by sequence homology; however, the overall architecture of this region is surprisingly similar, including the N-terminal domain (NT), the ankyrin-repeat domain, and the linker-helical domain (*SI Appendix, Fig. SSD*). A hydrophobic helix, formed by almost always 759–769 and anchored to the inner leaflet of the plasma membrane (red boxes of Fig. 1C and *SI Appendix, Fig. S5 A and C*), is similar to that observed in available TRPM structures, but is not present in other TRP channel structures. A long “connecting helix” (25 residues), present in TRPC3 (36), TRPC6 (37), TRPM2 (26), TRPM4 (24, 27), and TRPM8 (25), links the transmembrane domain and coiled-coil domain (Fig. 1C). Sequence alignments indicate that this connecting helix is conserved across TRPM family members (*SI Appendix, Figs. SSE and S6 A and B*).

**Ion Conduction Pore and Magnesium Binding Site**

TRPM7’s selectivity filter is formed by the backbone carbonyls of Phe1045/Gly1046 and Glu1047s side chain. The narrowest opening is 6.9 Å across the diagonally opposed Gly1046 residues (*SI Appendix, Fig. S7A*). Along the pore, His1093 (S6 cytoplasmic end) and Asn1097 (Cytoplasm+ TRPM domain) form another restriction site, defined as the lower gate. The most constricted point is Asn1097, where S6 helices from each subunit form a hydrophobic seal (*SI Appendix, Fig. S7A*). The distances between diagonal (4.1 Å) and neighboring (2.9 Å) side chains of Asn1097 are narrower than corresponding sites in the TRPA and TRPV structures. For comparison, the most restricted opening in TRPV1 is in the outer pore region, measuring 7.6 Å diagonally between opposed Gly643 residues (29). In TRPA1, the narrowest point (6.4 Å) is the lower gate (Val3961) (35).

Regardless of condition (with EDTA, DVF, or with high [Mg$^{2+}$]), the ion conduction pathway is essentially unchanged (Fig. 2A). The pore radius in all three structures differs by ~0.5 Å to 1 Å (Fig. 2B), which is most likely due to differences in resolution rather than structural modification.

A strong nonprotein density with a high signal-to-noise ratio is present in the ion permeation pathway of the TRPM7-Mg$^{2+}$ structure (Fig. 3 A and B). In contrast, peripheral densities are not found in the TRPM7-EDTA structure (Fig. 3C) at various contour levels (*SI Appendix, Fig. S8*). We interpret this density as a probable Mg$^{2+}$ ion located at the most restricted site of the selectivity filter (Fig. 3 D and E). Moreover, the distance between Mg$^{2+}$ and Glu1047s side chain (3.7 Å) and the backbone carbonyl of Gly1046 (4.0 Å; Fig. 3D) is large enough to accommodate a partially hydrated Mg$^{2+}$ ion.

**Assembly and Domain Interactions**

TRPM7’s “TRP domain” is the α-helical structure bending sharply toward the membrane immediately adjacent to the near-perpendicular S6 transmembrane helix (Fig. 1C). This TRP domain is a common feature of other TRP family members and is proposed to participate in subunit assembly or allosteric modulation of channel gating (38–41). In our structures, unlike those for TRPA1 and TRPV1, the pre-S1 helix does not interact with the TRP domain. Instead, Arg1115 and Lys1112 in the TRP domain (almost always 1099–1123) interact with the N-terminal Ser744 and Glu740, which are located on an elbow helix immediately after the pre-S1 helix (Fig. 4A and B). In addition, Trp1111 and Arg1115 form hydrogen bonds with Ala981 and Val982 on the S4–S5 linker, respectively (Fig. 4A and B). A π–π stacking interaction (Phe1118/Tyr1122) and a cation–π interaction (Arg1115/Trp1111) are also found on the TRP domain (Fig. 4A and B). These interactions are consistent with the TRP domain’s potential importance in gating. This hypothesis is supported by prior mutagenic studies in which...
Lys1112Gln and Arg1115Gln generated nonfunctional channels (42). Based on these and other structure–function studies of tetrameric ion channels (43–45), we hypothesize that the TRP domain and S4–S5 linker couple the movement of the “voltage sensor-like” S1 to S4 domains to the channel gate.

The transmembrane domains of TRPM7 are overlapped with other known core TRP structures in SI Appendix, Fig. S4, exhibiting remarkably similarity. The largest differences between TRP channels lie in the N and C termini. TRPV1’s ankyrin repeats 3 and 4 interact with the C-terminal domain via three-stranded antiparallel β-sheets (46). In TRPA1, a β-strand links a short, poorly resolved α-helix, buried in the inner leaflet of the membrane, to a TRP-like domain/C-terminal coiled-coil domain connecting loop (35). In contrast, TRPM7’s N/C-terminal interaction is through a well-resolved connecting helix, encoded by sequences that are relatively conserved among TRPM subfamily members (SI Appendix, Fig. S6 A and B). The center of these interactions is Tyr1181 and Phe1182, which form a π–π interaction and further interact with residues Trp642 and Glu692 on the NT (Fig. 4 A and C). A hydrogen bond and salt bridge involving Glu1184 and Glu1174 in the connecting helix with Arg699 and Try711 in the NT stabilize these domain interactions (Fig. 4 A and C).

**Major Structural Differences Between TRPM7 and TRPM4**

We next compared the structure of TRPM7-Mg$^{2+}$ with the structure of TRPM4, a monovalent cation-selective channel and member of TRPM2/4/5/8 subfamily (26). First, the conserved disulfide bond near the pore loop in TRPM7 (formed by Cys1056 and Cys1066; SI Appendix, Fig. S6 C and D) is closer to the S5–S6 helices than that of TRPM4 (Fig. 5 A and B). Second, the TRPM7 (1045FGGE1047) and TRPM4 (975FGQ977) selectivity
filters differ by one amino acid (Fig. 5C). This disparity is essential to TRPM7 divalent cation permeation, as mutating Glu1047 to Gln (the equivalent TRPM4 residue) abolished Ca\(^{2+}\) and Mg\(^{2+}\) permeability in full-length TRPM7 (28). We tested this mutation (Glu1047Gln) in our truncated TRPM7 protein and observed a similar result, with inward currents becoming inwardly rectifying in the mutant (6.5-fold greater at \(−100\) mV; Fig. 5D), while outward rectification was unchanged above 0 mV. Moreover, in the *Nematostella vectensis* (NvTRPM2) structure, the corresponding outermost selectivity filter residue is also a glutamate. However, in *Homo sapiens* (HsTRPM2), the equivalent residue is a glutamine (by sequence analysis). Electrophysiologically, this difference was correlated with an \(~70\)-fold greater \(P_{\text{Ca}}/P_{\text{Na}}\) in NvTRPM2 compared with HsTRPM2 (26). Thus, a negatively charged glutamate in the outermost selectivity filter position likely facilitates divalent cation permeability in TRPM channels (*SI Appendix*, Fig. S7 B and C). Third, the residues of the lower gate, which create the most constricted region of both pores, differ between TRPM7 (Asn1097) and TRPM4 (Ser1044). Here, asparagine–asparagine side chain separations in TRPM7 are smaller than those between serines in TRPM4 (Fig. 5C and *SI Appendix*, Fig. S7 B and C). However, the isoleucine and asparagine in the lower gate are well conserved between TRPM7, NvTRPM2, and hTRPM2 (*SI Appendix*, Fig. S7 B and C). Finally, the cytosolic domains are distinctly organized. In TRPM7, hydrophobic loops and two helices belonging to the NT are anchored to the inner leaflet of the plasma membrane (black boxes, Fig. 5E). This is more complex than the loop and short helix in TRPM4. The movement of this region may produce additional conformational changes in the cytosolic domains, perhaps leading to large vertical and rotational motions of the NT of the channel (Fig. 5E).

**Pore Loop Cysteines**

The movement and organization of the extracellular pore loop may be important to TRPM7 channel function. Conformational changes of the neighboring pore helices, S5 and S6, usually affect key channel properties. In the TRPM7-Mg\(^{2+}\) structure, we observed...
a pore loop disulfide bond between Cys1056 and Cys1066 (Fig. 6 A and B), which is conserved across the TRPM family (SI Appendix, Fig. S6 C and D). If this configuration is not due to protection from radiation damage (47), we hypothesize that divalent cations, such as Mg$^{2+}$, may stabilize this disulfide bond, since it is only observed in the TRPM7-Mg$^{2+}$ structure. Moreover, external application of the reducing agent glutathione potentiated TRPM7 currents by enhancing monovalent cation permeability (17), implicating an externally located disulfide bond. We tested whether the Cys1056/Cys1066 disulfide bond is essential for divalent permeability by mutagenesis and patch-clamp electrophysiology. When either or both cysteines are mutated, the resulting currents (I-V relationships) are more linear (Fig. 6 C and D), in contrast to the inward rectification noted in the prior glutathione results (17). Thus, mutation of these cysteines globally disrupts TRPM7 function by mechanisms which will require more investigation.

**Discussion**

Mg$^{2+}$ is the second most abundant cellular divalent cation. However, relatively little is known about the molecular components and mechanisms that regulate Mg$^{2+}$ compared with Ca$^{2+}$ homeostasis. In this study, we presented cryo-EM structures of the Mg$^{2+}$-permeable TRPM7 in three ionic conditions: TRPM7-DVF, TRPM7-Mg$^{2+}$, and TRPM7-EDTA. There are three conclusions from the work. First, in the TRPM7-Mg$^{2+}$ structure, a nonprotein density we presume is Mg$^{2+}$ is found within the selectivity filter. Mg$^{2+}$ has well-defined first and second hydration shells; the dehydration energy of the first hydration shell is 476 kcal/mol (48), at least 4 times that of Na$^+$ and K$^+$, making the passage of fully dehydrated Mg$^{2+}$ energetically unfavorable. In contrast, the second hydration shell of Mg$^{2+}$ is similar to bulk water (49). Thus, we propose that the backbone carbonyl of Gly1046 and the carboxyl group of Glu1047 replace the waters of...
the second hydration shell and compensate for the energetic cost of dehydration when hydrated Mg$^{2+}$ enters the pore. Importantly, the key negatively charged selectivity filter Glu1047 differs from the orthogonal Gln in TRPM4/5’s monovalent cation selective filter. Second, the channel’s lower gate has two constriction sites (Ile1093 and Asn1097), which is identical to that of NvTRPM2. Interestingly, Asn1097 occupies a similar position in TRPM1, TRPM3, TRPM6, and TRPM7, but not in the monovalent-selective TRPM4/5. As the only polar amino acid in the lower gate, the side chains of Asn1097 from each monomer form a polar ring at the cytoplasmic pore entrance. In addition, asparagine may also bind the first hydration shell of Mg$^{2+}$ (50). Third and finally, the pore helix’s disulfide bond is required for proper channel assembly/function and is a general feature of the TRPM family.

In summary, TRPM7’s structure sheds light on many questions raised by previous electrophysiological and physiological studies. The next steps will be to define open, desensitized, and blocked states of the TRPM7-EDTA, TRPM7-Mg$^{2+}$, and TRPM7-EDTA structures, 3.5 μM of purified TRPM7 in PMAL-C8 at 4 mg/mL to 5 mg/mL for electron cryomicroscopy.

For the high Mg$^{2+}$ structure, 300 mM MgCl$_2$ was added to the DVF buffer for every step of extraction and purification. For the TRPM7-EDTA structure, 2 mM EDTA (pH 8.2) was added to the protein extraction DVF buffer, and 5 mM EDTA was added to purified protein before freezing. In our DVF buffer, we determined that NaCl (Fisher Scientific) contributed the most contaminating Ca$^{2+}$ (0.002%, 3 μM) and Mg$^{2+}$ (<0.001%, <1.5 μM). Using MaxChelator software (https://web.stanford.edu/~patton/webmaxcs.htm), the free Ca$^{2+}$ and Mg$^{2+}$ in the purified TRPM7-EDTA protein were estimated as 3.8 pM and 895 pM, respectively. We note that these values may change under cryoconditions, as MaxChelator estimates are limited to a minimum temperature of 0°C.

**Methods**

**Protein Expression and Purification of MBP-TRPM7.** The mouse TRPM7 construct (amino acids 1 to 1280) was cloned into the pEG BacMam vector (52). An MBP tag was added to TRPM7’s N terminus via the linker, N555NNNNNNNNNNLQEVQLFEQPQ, containing the HRV 3C cleavage site (underlined). P3 baculovirus was produced in the Bac-to-Bac Baculovirus Expression System (Invitrogen) using Spodoptera frugipera (SPF) cells transduced at a density of 2 to 3 × 10$^6$ cells/mL. For protein expression, HEK293S GnTI− cells (lacking N-acetylglucosaminyltransferase I) were grown in Freestyle 293 Expression medium (Life Technologies) supplemented with 2% FBS and 1% penicillin-streptomycin (37°C, 8% CO$_2$), and transduced with P3 baculovirus once cells reached a density of 2 to 3 × 10$^6$ cells/mL. After 12 h to 24 h, 10 mM sodium butyrate was added, and the temperature was reduced to 30°C. Cells were harvested 72 h after transduction and frozen at −80°C. Before solubilization, cells were resuspended in 30 min in a DVF buffer containing 30 mM Hapes, 150 mM NaCl, 1 mM dithiothreitol (DTT), pH 7.5 with EDTA-free protease inhibitor mixture (Roche). MBP-TRPM7 membranes were mechanically homogenized and solubilized for 2 h to 3 h in a buffer containing 1.0% (w/v/o) N-dodecyl-beta-D-maltopyranoside (DDM; Affymetrix), 0.1% (w/v/o) cholesteryl hemi-succinate (CHS; Sigma), 25 mM Hapes, 150 mM NaCl, 1 mM DTT, pH 7.5 with EDTA-free protease inhibitor mixture (Roche). The supernatant was isolated by centrifugation at 100,000 × g for 60 min, followed by incubation in amyllose resin (New England BioLabs) at 4°C overnight. The resin was washed with 20 column volumes of 25 mM Hapes, 150 mM NaCl, 0.05% (w/v/o) DDM, 0.01% (w/v/o) CHS, 1 mM DTT, pH 7.5 with EDTA-free protease inhibitor mixture (Roche). The protein was eluted with four column volumes of 25 mM Hapes, 150 mM NaCl, 0.05% (w/v/o) DDM, 0.01% (w/v/o) CHS, 1 mM DTT, pH 7.5 with EDTA-free protease inhibitor mixture (Roche) and 40 mM maltose. The protein was then concentrated and mixed with poly (maleyl anhydride-alt-1-decene) substituted with 3-(dimethylamino)propylamine (PMAL-C8, anatrace) at 1:4 (wt/wt) with gentle agitation for 4 h. Bio-Beads SM-2 (50 mg/mL reconstitution mixture; Bio-Rad) were added to remove additional detergent, and a disposable Poly-Prep column was used to remove Bio-Beads. The protein was then concentrated to 0.5 mL with a 100-kDa molecular weight cutoff concentrator (Millipore) before further purification on a Superose 6 column in a buffer composed of 25 mM Hapes, 150 mM NaCl, 1 mM DTT, pH 7.5. The peak, corresponding to tetrameric C-terminally truncated TRPM7, was collected and concentrated to 4 mg/mL for electron cryomicroscopy.

**Electron Microscopy Data Collection.** For the TRPM7-EDVF, TRPM7-Mg$^{2+}$ and TRPM7-EDTA structures, 3.5 μL of purified TRPM7 in PMAL-C8 at 4 mg/mL to 4.5 mg/mL was applied onto a glow-discharged, 400 mesh copper grid (Quantifoil). Grids were blotted for 7 s at 100% humidity and flash frozen in liquid nitrogen-cooled liquid ethane using an FEI Vitrobot Mark I (FEI). The grid was loaded onto an FEI TF30 Polara electron microscope operating at 300 kV accelerating voltage. TRPM7 image

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**Fig. 4.** Assembly and domain interactions. (A) One TRPM7 subunit from the plane of the membrane with regions of interest depicted by boxes. (B) Expanded view of A showing interactions between the N terminus (pink) and the TRP domain (blue). (C) Expanded view of A showing interactions between N (pink) and C (green) termini. Residues at domain interfaces are labeled, with potential hydrogen bonds and electrostatic interactions shown as dashed lines.
Fig. 5. Structural comparison of TRPM7 and TRPM4. (A) Overlaid monomer of TRPM7 (blue, pink, and cyan) and TRPM4 (gray). (B) Enlarged view of key pore loop disulfide bond between Cys1056 and Cys1066. (C) Aligned TRPM7 (blue) and TRPM4 (gray) dimer pores. Residues of interest are labeled and shown as sticks. (D) Whole-cell I-V relationships of HEK293T TRPM6/7 double KO cells expressing MBP-tagged truncated mouse TRPM7 (blue), with 4 mM Mg\(^{2+}\) (gold), mTRPM7 in 4 mM Mg\(^{2+}\), or truncated mouse E1047Q-TRPM7 (black). Bar charts represent the mean currents recorded at +100 mV and −100 mV; E1047Q (n = 18, 103 ± 10 pA/pF and −82 ± 1 pA/pF), mTRPM7 (n = 10, 111 ± 10 pA/pF and −12 ± 0.3 pA/pF), and mTRPM7+4 mM Mg\(^{2+}\) (mTRPM7+Mg\(^{2+}\), n = 10, 5 ± 1 pA/pF and −2 ± 0.1 pA/pF). (E) Side and bottom views of superimposed cytosolic domains of TRPM7 (blue and pink) and TRPM4 (gray); boxes highlight extracellular regions containing cysteines.
Stacks were recorded on Gatan K2 Summit (Gatan) direct detector set in superresolution counting mode using SerialEM (53), with a defocus range between 1.5 μm and 3.0 μm. The electron dose was set to 8 e− per physical pixel per second; subframe time of 200 ms and total exposure time of 10 s resulted in 50 subframes per image stack. The total electron dose was 52.8 e− per square angstrom (~1.1 e− per square angstrom per subframe).

**Image Processing and 3D Reconstruction.** Similar strategies were used for the image processing and 3D reconstruction of TRPM7-DVF, TRPM7-Mg2+, and TRPM7-EDTA structures. Image stacks were gain-normalized and binned by 2× to a pixel size of 1.23 Å before drift and local movement correction using motionCor2 (54). The images from the sum of all frames with dose weighting were subjected to visual inspection, and poor images were removed before particle picking. Particle picking and subsequent bad particle elimination through 2D classification was performed using Python scripts/programs developed by Maofu Liao (55) with minor modifications in the 8 coins method. The refined atomic model was further visualized in COOT. A few residues with side chains moving out of the density during the refinement were subsequently achieved based primarily on the clearly defined side chain densities of bulky residues such as Phe, Tyr, and Trp, as well as some Arg and Lys residues. Resolution of the first part of NT (1 to 150) were insufficient for backbone tracing, and the polyalanine model was used for that region. The refined atomic model was further visualized in COOT. A few residues with side chains moving out of the density during the refinement were fixed manually, followed by further refinement. The TRPM7 model was then subjected to global refinement and minimization in real space using the PHENIX (60) module “phenix.real_space_refine” (61), and geometries of the model were assessed using MolProbity (62) in the comprehensive model validation section of PHENIX. The final model of TRPM7-EDTA structure exhibited good geometry as indicated by the Ramachandran plot (preferred region, 97.8%; allowed region, 2.1%; outliers, 0%)

Fig. 6. Comparison of TRPM7-Mg2+ and TRPM7-EDTA structures. (A) Superimposition of the transmembrane module from TRPM7-Mg2+ (blue) and TRPM7-EDTA structures (pink). (B) Site of the broken disulfide bond between Cys1056 and Cys1066 in the pore domain. Key discernible residues, Cys1056 and Cys1066, are labeled. Shown is superimposition of the disulfide bond between S5 and S6 in the TRPM7-Mg2+ (blue) and TRPM7-EDTA (pink) structures. (C) Whole-cell I-V relationship of HEK293T TRPM6/7 double KO cells expressing MBP-tagged truncated mouse TRPM7 (mTRPM7; orange), C1056S (red), C1065S (green), and C1056S/C1066S (violet). (D) Mean currents of mTRPM7 and mTRPM7 cysteine(s) mutant measured at ±100 mV and ±100 mV; nontransfected (n = 10, 2.3 ± 0.5 and –1.6 ± 0.1), mTRPM7 (Time-zero) (n = 12, 5.1 ± 0.1 and –2.1 ± 0.1 pA/pF), C1056S/C1066S (n = 12, 55 ± 3 and –22 ± 3 pA/pF), C1056S (n = 12, 124 ± 5 and –56 ± 2 pA/pF), C1065S (n = 10, 90 ± 10 and –25 ± 4 pA/pF), and mTRPM7 (n = 10, 111 ± 10 and –12 ± 1 pA/pF).
The pore radius was calculated using HOLE (63). For the TRPM7-DVF and TRPM7-Mg structures, the structure of TRPM7-EDTA at 3.3 Å was docked into the cryo-EM maps, followed by manual adjustment in COOT. The models, then subjected to the same global refinement and minimization process described for TRPM7-EDTA, and geometries of the models were also assessed using MolProbity (62). The final models of TRPM7-DVF and TRPM7-Mg structures exhibited good geometry as indicated by the Ramachandran plot.

Electrophysiology. All electrophysiological experiments were performed in HEK293T TRPM7-GFP double KO cells to eliminate endogenous TRPM6 and TRPM7 currents (13). The cells were cultured in DMEM (ATCC, 30-2002) with 10% FBS, 1% Pen/Strep, 10 mM MgCl₂, 3 µg/mL of blasticidin and 0.5 µg/mL of puromycin. Cells were transiently transfected (48 h) with the MBP-tagged truncated mouse TRPM7 construct used for structural investigation or various mutants (C1056S, C1066S, C1056S/C1066S). E1047Q mouse TRPM7 was kindly provided by Lixia Yue, University of Connecticut Health Center, Farmington, and we truncated this construct to mimic the mouse TRPM7 channel used for structural investigation. Cells were cotransfected with EGFP at 10% of total DNA for visualization and cell selection.

Cells were perfused at a rate of 1 mL/min to 2 mL/min in a modified Tyrode's solution containing (millimolars): 140 NaCl, 5 KCl, 2 CaCl₂, 10 glucose, and 10 Heps; pH 7.4. Where indicated, cells were perfused with a high Mg²⁺ solution containing (millimolars): 110 MgCl₂, 10 Heps; pH 7.4. Patch pipettes of 2 MΩ to 5 MΩ contained (millimolars): 120 CsCH₃SO₄, 8 NaCl, 10 CsBAPTA, 2 CaCl₂, 2 NaATP, 10 Heps; pH 7.4. For experiments utilizing the high Mg²⁺ external solution, a modified pipette solution contained (millimolars): 120 CsCH₃SO₄, 8 NaCl, 10 mM HEDTA, 5 CsBAPTA, 1 CaCl₂, 2 NaATP, 10 Heps; pH 7. All solutions were stored at 4°C.

Whole-cell currents were recorded at room temperature using an Axopatch 200B patch-clamp amplifier controlled via a Digidata 1440A (Molecular Devices) and pClamp 10.3 software. Data were digitized at 10 kHz and low-pass-filtered at 2 kHz. In most recordings, cells were held at 0 mV, and 200-ms ramps from −100 mV to 100 mV were applied at 0.5 Hz. Where indicated, cells were held at 0 mV and stepped from −100 mV to +100 mV in +10-mV increments (2 Hz per step). Before recordings, currents were corrected for pipette (fast) capacitance, whole-cell capacitance, and series resistance compensated to 80%. Currents were offline-corrected for liquid junction potential using previously described methods (64).

Data and Software Availability. Cryo-EM electron density maps of the mouse TRPM7-EDTA, TRPM7-Mg²⁺, and TRPM7-DVF and have been deposited in the Electron Microscopy Data Bank under accession code EMD-6975, EMD-7297 and EMD-7298, respectively. Atomic coordinates of the mouse TRPM7-EDTA, TRPM7-Mg²⁺ and TRPM7-DVF have been deposited in the Protein Data Bank under accession code 6ZXS, 6BWD and 6BWF, respectively.

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