A hydrophobic funnel governs

monovalent cation selectivity in the ion channel TRPM5

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Data availability: MD

simulation inputs and analysis scripts used for this study are deposited in a public GitHub repository, available at: https://github.com/cmives/ Na_selectivity_mechanism_ of_TRPM_channels.

Funding: CMI was supported by the Medical Research Council [grant number MR/N013735/1], NJT and ATŞ were supported by the UKRI Biotechnology and Biological Sciences Research Council (BBSRC) [grant numbers BB/M010996/1 and BB/T00875X/1, respectively].

Competing interests: The authors declare no competing interests.

11 Abstract

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- A key capability of ion channels is the facilitation of selective permeation of certain ionic species
- across cellular membranes at high rates. Due to their physiological significance, ion channels are
- ¹⁴ of great pharmaceutical interest as drug targets. The polymodal signal-detecting Transient
- ¹⁵ Receptor Potential (TRP) superfamily of ion channels form a particularly promising group of drug
- targets. While most members of this family permeate a broad range of cations including Ca²⁺,
- 17 TRPM4 and TRPM5 are unique due to their strong monovalent-selectivity and impermeability for
- divalent cations. Here, we investigated the mechanistic basis for their unique
- ¹⁹ monovalent-selectivity by *in silico* electrophysiology simulations of TRPM5. Our simulations reveal
- ²⁰ an unusual mechanism of cation selectivity, which is underpinned by the function of the central
- ²¹ channel cavity rather than the selectivity filter. Our results suggest that a subtle hydrophobic
- ²² barrier at the cavity entrance ("hydrophobic funnel") enables monovalent, but not divalent
- ²³ cations to pass and occupy the cavity at physiologically relevant membrane voltages. Monovalent
- 24 cations then permeate efficiently by a co-operative, distant knock-on mechanism between two
- ²⁵ binding regions in the extracellular pore vestibule and the central cavity. By contrast, divalent
- cations do not enter or interact favourably with the channel cavity due to its raised
- ²⁷ hydrophobicity. Hydrophilic mutations in the transition zone between the selectivity filter and the
- ²⁸ central channel cavity abolish the barrier for divalent cations, enabling both monovalent and
- ²⁹ divalent cations to traverse TRPM5.

31 Introduction

- ³² The translocation of ions across cellular and organellar membranes via ion channels is essential to
- ensure cellular ionic homeostasis and provides a key pathway of intra- and intercellular communi-
- $_{34}$ cation. Ion channels catalyse the permeation of ions across the membrane up to an order of 10^8
- ³⁵ ions per second, while at the same time often displaying strict selectivity for particular ionic species
- 36 (Dudev and Lim, 2014; Roux, 2017; Kopec et al., 2018). The transient receptor potential (TRP) super-
- ³⁷ family of ion channels comprises a large group of cation-selective channels that are implicated in
- ³⁸ a wide range of physiological processes (*Ramsey et al., 2006; Khalil et al., 2018*). Due to their phys-
- ³⁹ iological importance, TRP channels are associated with a large number of pathological conditions
- 40 (Nilius, 2007), including in the aetiology of several rare, genetic conditions. Many members of the
- 41 TRP channel superfamily constitute major pharmaceutical target proteins (Moran, 2018; Koivisto
- 42 et al., 2022).
- Within the TRP channel superfamily, TRPM (transient receptor potential melastatin) channels
 form the largest subfamily, consisting of eight members (TRPM1-8) (*Nilius and Owsianik, 2011*;
 Samanta et al., 2018). TRPM channels assemble as homotetramers, in which each subunit provides
 six transmembrane helices (S1-S6), a cytosolic *N*-terminus domain composed of four melastatin
- homology regions, and a cytosolic C-terminus coiled-coil domain (*Hilton et al., 2019: van Goor*
- et al., 2020). In keeping with most members of the TRP superfamily, TRPM channels are described
- as being cation non-selective, that is, they conduct cations but do not differentiate substantially
- ⁵⁰ between cationic species. However in the TRPM subfamily, TRPM4 and TRPM5 are exceptions to
- this observation, since both channels are selective for monovalent cations and impermeable to
- ⁵² divalent cations (*Owsianik et al., 2006*). Thereby, TRPM4 and TRPM5 are the only members of the
- ⁵³ wider TRP superfamily to display selectivity for monovalent cations.
- Although TRPM4 and TRPM5 are close homologs, sharing both a high degree of sequence ho-
- mology and similar biophysical characteristics, there are some variations in their activation mecha-
- ⁵⁶ nisms. For example, while both channels are activated by raised intracellular Ca²⁺ concentrations,
- ⁵⁷ TRPM5 is approximately 20-fold more sensitive to Ca²⁺ than TRPM4 (Ullrich et al., 2005). Ion con-
- ⁵⁸ duction through TRPM5 has been implicated in the sensation of sweet, bitter, and umami tastes
- in type II taste bud cells (Pérez et al., 2002; Zhang et al., 2003), and in the secretion of insulin by
- ⁶⁰ pancreatic β-cells (*Brixel et al., 2010; Colsoul et al., 2010*). Consequently, TRPM5 is a potential drug
- target for a number of conditions, including metabolic conditions such as type II diabetes mellitus
- 62 (Vennekens et al., 2018). Several molecular structures of the TRPM4 and TRPM5 channels have
- been published to date, however an open-state structure has only been solved for TRPM5 (*Ruan et al.* 2021).
- In the present work, we set out to characterise the cation permeation mechanism of the TRPM5
- channel, focusing in particular on the basis for its monovalent cation selectivity, by conducting
- atomistic molecular dynamics (MD) simulations and *in silico* electrophysiology of the open-state
- structure of Danio rerio TRPM5 (Ruan et al., 2021) (PDB ID: 7MBS) in solutions of Na⁺, K⁺, and Ca²⁺
- ⁶⁹ ions. We recorded more than 700 individual ion permeation events from over 20 μ s of aggregated
- time from our *in silico* electrophysiology simulations.

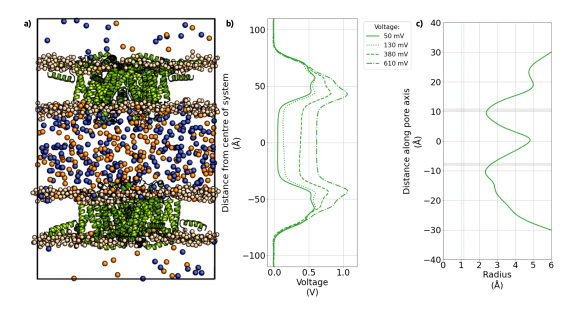


Figure 1. Structure and membrane voltage of CompEL simulations of TRPM5. **a**) Snapshot of the CompEL system showing the TRPM5 pore domain of *Danio rerio* used in this study inserted into a double bilayer simulation system in an anti-parallel fashion so that both proteins experience identical voltage polarity. Cations within the aqueous compartments are shown as spheres (orange: calcium; blue: sodium), highlighting the 9:1 ion concentration gradient between the compartments. **b**) The CompEL charge differences we applied across the aqueous compartments (Δq) resulted in transmembrane voltages of ~ -50 mV, -130 mV, -380 mV, and -610 mV in addition to the concentration gradient. **c**) Average pore radius of TRPM5 along the pore axis from MD simulations. The regions in grey shade represent the average positions of the major pore constrictions in TRPM5, formed by Q906 (*upper* gate) and 1966 (*lower* gate). The dashed line indicates the radius of a completely dehydrated Ca²⁺ ion for comparison.

Our findings reveal a new mechanism of ion selectivity, based on a hydrophobic barrier at the entrance to the central channel cavity, which shields the cavity from an influx of divalent cations. In this way, the central cavity forms a binding site for monovalent cations, but not for divalent cations. The conduction of monovalent ions thus becomes a synergistic process incorporating cooperativity between multiple binding sites.

76 Methods & Materials

77 TRPM5 system construction

- 78 A truncated TRPM5 simulation system consisting of the membrane-domain of the channel was
- ro constructed by using residues 698-1020, including the resolved N-acetyl- β -D-glucosamine of the
- ⁸⁰ glycosylated N921 residue, of the Danio rerio TRPM5 structure (Ruan et al., 2021) (PDB ID: 7MBS).
- $_{\text{e1}}$ We also modelled the bound Ca²⁺ cations occupying the Ca_{TMD} binding sites at E768 and D797 in
- $_{*2}$ each subunit, which have been proposed to be implicated in Ca²⁺-dependent activation of TRPM5.
- ⁸³ The system was built using the CHARMM-GUI server (*Jo et al., 2008*). The charged N- and C-terminal
- residues were neutralised by capping with acetyl (ACE) and N-methylamide (CT3) groups, respec-
- tively. All missing non-terminal residues were modelled using CHARMM-GUI (*Jo et al., 2014*).
- ⁸⁶ The structure was aligned in the membrane using the PPM server (*Lomize et al., 2012*), inserted

- into a 1-palmitoyl-2-oleoyl-sn-glycerol-3-phosphocholine (POPC) bilayer of 160 x 160 Å size with
- the CHARMM-GUI membrane builder (*Jo et al., 2007; Wu et al., 2014*), and then solvated. lons
- were added with GROMACS 2020.2 (Abraham et al., 2015; Lindahl et al., 2020) to neutralise any
- ⁹⁰ system charges and add ions to a concentration of either 150 mM NaCl, 150 mM KCl, 150 mM CaCl₂
- ⁹¹ (referred to as mono-cationic solutions), or a mixture of 75 mM NaCl and 75 mM CaCl₂ (referred
- to as di-cationic solutions). In the case of simulations containing Ca²⁺, the standard CHARMM36m
- parameters for Ca^{2+} ions were then replaced with the multi-site Ca^{2+} of Zhang *et al.*, (*Zhang et al.*,
- **2020**). This multi-site model has used been used to investigate Ca^{2+} permeation in a number of
- ⁹⁵ channels, including including the type-1 ryanodine receptor *Zhang et al.* (2020); Liu et al. (2021),
- MMPA receptors (Schackert et al., 2022), the E protein of SARS-CoV-2 (Antonides et al., 2022), and
- TRPV channels (Liu and Song, 2022; Ives et al., 2023).

Molecular dynamics simulation details

All simulations were performed using GROMACS 2020.2 (Abraham et al., 2015: Lindahl et al., 2020) 99 or GROMACS 2022 (Bauer et al., 2022), together with the CHARMM36m force field for the protein. 100 lipids, and ions (except for Ca²⁺ as noted above) (*Huang et al., 2016*). The TIP3P water model was 101 used to model solvent molecules (*Jorgensen et al., 1983*). The system was minimised and equi-102 librated using the suggested equilibration input scripts from CHARMM-GUI (Lee et al., 2016). In 103 brief, the system was equilibrated using the NPT ensemble for a total time of 1.85 ns with the 104 force constraints on the system components being gradually released over six equilibration steps. 105 The systems were then further equilibrated by performing a 15 ns simulation with no electric field 106 applied. To prevent closing of the lower hydrophobic gate of the pore, harmonic restraints were ap-107 plied to maintain the distance between the α -carbon atoms of the lower gate residue 1966 of each 108 respective chain. The temperature was maintained at T = 310 K using the Nosé-Hoover thermostat 109 (Evans and Holian, 1985), and the pressure was maintained semi-isotropically at 1 bar using the 110 Parrinello-Rahman barostat (Parrinello and Rahman, 1981), Periodic boundary conditions were 111 used throughout the simulations. Long-range electrostatic interactions were modelled using the 112 particle-mesh Ewald method (Darden et al., 1993) with a cut-off of 12 Å. The LINCS algorithm (Hess 113 et al., 1997) was used to constrain bond lengths involving bonds with hydrogen atoms. Hydrogen mass re-partitioning (HMR) of the system was used to allow the use of 4 fs integration time steps in 115 simulations of NaCl solutions. The multi-site Ca²⁺ model used for simulations of CaCl₂ however is 116 incompatible with a 4-fs time step, and therefore any simulations involving Ca²⁺ cations were per-117 formed with HMR but at a time step of 2-fs. A summary of all simulations performed is presented 118 in Table 1, and in more detail in Tables S1 and S2.

120 CompEL simulations

- We employed the computational electrophysiology (CompEL) protocol (Kutzner et al., 2011, 2016)
- of GROMACS to create a transmembrane voltage and drive ion permeation in an anti-parallel dou-
- ¹²³ ble membrane system, such that both channels experienced the same voltage polarity with neg-
- ative polarity in the intracellular region. Simulations were performed in a di-cationic solution of
- ¹²⁵ 75 mM NaCl and 75 mM CaCl₂ with a range of ionic imbalances (Δ q), resulting in membrane volt-

Protein	Transmembrane voltage methodology	Voltage	lon solution	Simulation duration	
		-50 mV	75 mM NaCl +	3 x 500 ns	
	CompEL		75 mM CaCl ₂	3 X 300 113	
	(anti-parallel)	-160 mV	75 mM NaCl +	3 x 500 ns	
	(and parallely		75 mM CaCl ₂	3 X 300 H3	
		- 380 mV	75 mM NaCl +	3 x 500 ns	
TRPM5			75 mM CaCl ₂	5 X 300 H3	
		-610 mV	75 mM NaCl +	3 x 500 ns	
			75 mM CaCl ₂	3 × 300 113	
	External applied field	-50 mV	150 mM NaCl	3 x 250 ns	
			150 mM CaCl ₂	3 x 250 ns	
		-200 mV	150 mM NaCl	3 x 250 ns	
			150 mM CaCl ₂	3 x 250 ns	
		-340 mV	150 mM NaCl	3 x 250 ns	
			150 mM KCl	3 x 250 ns	
			150 mM CaCl ₂	3 x 250 ns	
		-130 mV	150 mM NaCl	3 x 250 ns	
TRPM5			150 mM CaCl ₂	3 x 250 ns	
F904T		-200 mV	150 mM NaCl	3 x 250 ns	
			150 mM CaCl ₂	3 x 250 ns	

Table 1. Summary of simulations performed in this study.

ages of \sim -50 mV, -130 mV, -380 mV, and -610 mV. To further drive cation permeation, we also generated a neutral ion concentration gradient of 9:1 between the extracellular and intracellular solutions (Figure 1). All CompEL simulations were 500 ns long and repeated three times for each

- $_{129}$ system, resulting in an aggregated simulation time of 3 μ s per membrane voltage due to the double
- 130 channel nature of these simulations.

131 External applied field simulations

- 132 In addition to CompEL simulations, we also performed simulations in mono-cationic solutions of
- 133 150 mM NaCl, 150 mM KCl, and 150 mM CaCl₂, using an applied electric field to produce membrane
- voltage Aksimentiev and Schulten (2005). Fields of -0.03, -0.0175, or -0.0044 V nm⁻¹ were applied,
- resulting in transmembrane voltage of ~-340 mV, 2-00 mV, or -50 mV, respectively, with negative
- polarity in the intracellular region. All applied field simulations were 250 ns long and repeated
- 137 three times for each system.

¹³⁸ Simulation analysis

- Analysis of MD trajectory data was performed using in-house written Python scripts, utilising GRO-
- MACS modules (Abraham et al., 2015; Lindahl et al., 2020), the SciPy library of tools (Oliphant, 2007;

- 141 Pérez and Granger, 2007; Millman and Aivazis, 2011; Van Der Walt et al., 2011), and MDAnalysis
- (Michaud-Agrawal et al., 2011; Gowers et al., 2016). Analysis of the pore architecture was per-
- formed using CHAP (Rao et al., 2019). All plots were generated in Python using Matplotlib (Hunter,
- 2007) and Seaborn (Waskom et al., 2018). All MD input and analysis scripts used for this study
- are deposited in a public GitHub repository, available at: https://github.com/cmives/Na_selectivity_
- 146 mechanism_of_TRPM_channels.

¹⁴⁷ Calculating conductance and selectivity from *in silico* electrophysiology experiments

- The conductance of the channels (C_{ion}) was calculated according to Equation 1, where N_p is the
- number of permeation events, Q_{ion} is the charge of the permeating ion in Coulomb, t_{traj} is the length
- $_{150}$ of the trajectory, and V_{tm} is the transmembrane voltage. The mean conductance and standard error
- were calculated from overlapping 50 ns windows of the trajectory.

$$C_{ion} = \frac{N_p \times Q_{ion}}{t_{iraj} \times V_{im}}$$
(1)

The selectivity (P_{Na}/P_{Ca}) from the di-cationic CompEL simulations was calculated as the ratio between the total sum of Na⁺ permeation events and the total sum of Ca²⁺ permeation events across all simulations in a certain voltage or concentration regime.

¹⁵⁵ Identification of cation binding sites from MD simulations of TRPV channels

Cation binding sites were identified by plotting timeseries of each permeating ion with respect to their position along the pore axis. To further validate these positions, a 3D density mesh was generated for cations within 10 Å of the protein. This analysis was performed on a trajectory of concatenated, three-fold replicated 500 ns simulations in mono-cationic solutions with a voltage of \sim -50 mV produced by the CompEL method.

- ¹⁶¹ Characterising permeation cooperativity through mutual information with SSI from PENSA
- ¹⁶² To characterise the level of co-operativity of the ion permeation mechanisms within the TRPM5
- channel, we used PENSA to calculate the state-specific information (SSI) shared between discrete
- state transitions in the occupancy distributions of both of the pore binding sites **Thomson et al.**
- (2021); Vögele et al. (2022). The methodology has been described in greater detail in our previous
 work *lves et al.* (2023).

¹⁶⁷ In brief, a timeseries distribution with a timestep of 20 ps for each binding site was obtained. ¹⁶⁸ For each frame, the ion's atom ID number was recorded if an ion occupied the binding site in this

- ¹⁶⁹ frame (occupied state). By contrast, if the binding site was unoccupied (vacant state), an ID of -1
- was recorded. We then quantified by mutual-information whether ion transitions from occupied
- to vacant, or vice versa, at one site were coupled to similar ion transitions at the second ion binding
- site. To account for statistical noise that can arise from distributions even if they are uncorrelated
- with one another due to small-batch effects (McClendon et al., 2009; Pethel and Hahs, 2014), we
- calculated a statistical noise threshold. This threshold level was subtracted from the measured SSI
- values to yield the excess mutual information, or excess SSI (exSSI) above noise.

176 Results

¹⁷⁷ Cation conductance of the TRPM5 channel in di-cationic solutions

We performed in silico simulations of open state Danio rerio TRPM5 (Ruan et al., 2021) embed-178 ded in a dual POPC lipid bilaver system, with a di-cationic solution of 135 mM NaCl and 135 mM 170 CaCl₂ in the central dense aqueous compartment, and 15 mM NaCl and 15 mM CaCl₂ in the outer diluted aqueous compartments, respectively (Figure 1). An anti-parallel CompEL double bilayer 181 setup (Kutzner et al., 2011) was used to yield a bio-mimetic transmembrane voltage of ~ -50 mV 182 across both embedded channels, as well as higher voltages of -130 mV, -380 mV and -610 mV to 183 increase the number of permeation events and improve the statistics of our analyses (Figure 1). 184 The 9:1 ion concentration gradient between the middle and the outside bulk compartment acted synergistically with the membrane voltage to drive ion permeation. 186 Our simulations showed a continuous flow of permeating ions, resulting in a total of 374 per-187 meation events across all investigated simulation conditions performed with the CompEL setup. 188 Even though the ion gradient provided an additional driving force for permeation alongside the 189 voltages, the calculated conductances from our *in silico* electrophysiology simulations, in a range

¹⁹¹ between 7 and 38 pS (Table 2), were generally in good agreement with the published conductance

values of 23–25 pS from *in vitro* electrophysiology experiments on TRPM5 in NaCl based solutions

193 (Hofmann et al., 2003; Prawitt et al., 2003).

Low-voltage simulations in di-cationic solutions show exclusive permeation of Na⁺ through TRPM5

At the lowest simulated voltages of ~ -50 mV and -130 mV, we observed complete Na⁺-selectivity in mixed Ca²⁺/Na⁺ solutions, with no recorded Ca²⁺ permeation during an accumulated simulation time of 1.5 μ s. During the same time span, 15 (-50 mV) and 18 Na⁺ ions (-130 mV) traversed the TRPM5 pore, respectively, in accordance with its general conductance level (Table 2).

Analysis of the pore architecture of TRPM5 showed no major conformational changes throughout the course of the simulations. The TRPM5 pore possesses two main constrictions: an upper constriction formed by the sidechains of Q906 and by G905 of the three-residue selectivity filter (SF), and a lower constriction formed by the sidechains of 1966 of the lower gate (Figure 1). A minor constriction can also be observed ~13 Å above the SF, in the extracellular pore vestibule (EPV) (Figure 2). This constriction is formed by the turret loop between the pore helix (PH) and the S6 helix.

In our simulations at -50 mV and -130 mV, Na⁺ cations first entered the EPV region of the TRPM5 pore, where they showed a broad association with the protein matrix. Permeating Na⁺ cations then traversed the SF rapidly, and entered the pore cavity. They spent a substantial amount of time occupying the cavity before passing through the lower gate and exiting the pore at the intracellular face.

As opposed to monovalent Na⁺, Ca²⁺ ions did not readily enter the inner pore of TRPM5 during the course of the simulations. Ca²⁺ cations chiefly occupied the EPV region at the extracellular entrance (see Figure 2). 3D density maps of Na⁺ and Ca²⁺ ions further confirmed this observation

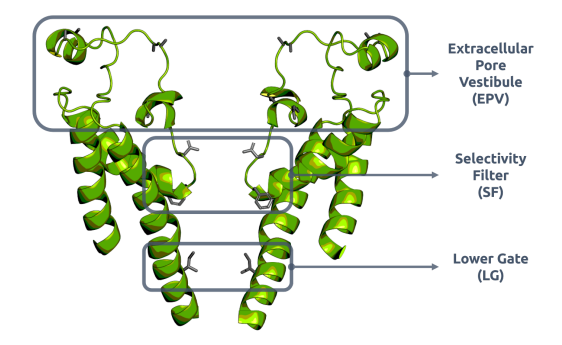


Figure 2. Overview of the structure of the TRPM5 channel of *Danio rerio* used in this work (two of the four subunits are omitted for clarity). TRPM5 has a short, three-residue selectivity filter (SF) consisting of Q906, G905, and F904. The hydrophobic lower gate (LG) of TRPM5 is formed by I966. In this study, the pore is defined as the region between the the two constrictions of the channel, namely Q906 of the SF and I966 of the lower gate. Above the pore is the extracellular pore vestibule, which contains a number of acidic residues, such as E910, E911, D919, D920, D925, and E928. All residues mentioned by name are displayed as grey sticks.

(Figure 3). The maps show substantial Ca²⁺ density in the EPV, particularly near the acidic residues
on the loop between the PH and the S6 helix, namely: E910, E911, D919, D920, D925, and E928. We
observed that Ca²⁺ ions occasionally migrated from the EPV toward the pore, however they were
blocked from entering the cavity at the SF, particularly at the constriction formed around G905 and
F904 from each subunit (Figure 3).

²²⁰ Voltage dependence of simulated TRPM5 ion selectivity

As the membrane voltage was increased in our CompEL simulations, we observed the Na⁺ selectivity (P_{Na}/P_{Ca}) to be diminished (Table 2). At a voltage of both ~ -50 mV and ~ -130 mV, we recorded complete Na⁺-selectivity, with no Ca²⁺ permeation events in any of the simulations. At a voltage of ~ -380 mV, the *in silico* electrophysiology simulations continued to display slightly Na⁺-selective permeation; however, when the voltage was further increased to ~ -610 mV, the Na⁺-selectivity was lost. Furthermore, higher-voltage simulations also yielded a small number of Cl⁻ permeation events, with anions permeating through to the extracellular solution. Our findings suggest relatively weak cation binding sites within the pore domain, in line with

²²⁸ Our findings suggest relatively weak cation binding sites within the pore domain, in line with ²²⁹ the absence of negatively charged residues lining the SF and inner cavity, due to the substantial

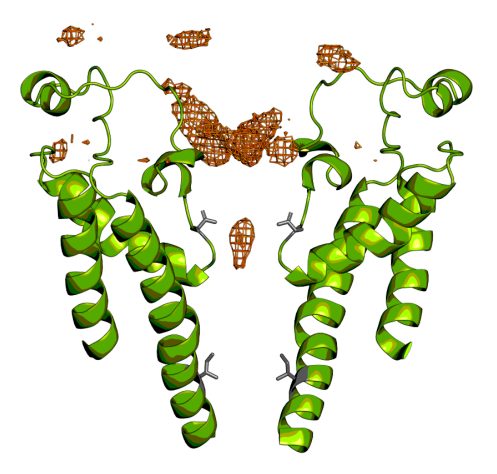


Figure 3. 3D density map of Ca^{2+} cations around the EPV and SF of TRPM5. The density of Ca^{2+} ions was calculated from concatenated trajectories of TRPM5 in a di-cationic solution under a transmembrane voltage of ~ -50 mV generated by the CompEL method. A major density maximum is seen within the EPV, where Ca^{2+} associates. Occasionally, Ca^{2+} ions migrated into the SF (minor density maximum at Q906); however, they were not able to traverse past the SF at bio-mimetic voltages. The sidechains of Q906 of the SF and I966 of the lower gate are shown as sticks in grey.

- ²³⁰ effect of supra-physiological membrane voltages. To further explore the ion permeation dynamics
- in TRPM5 and their underlying determinants, we thus aimed to enhance the sampling of both
- ²³² Na⁺ and Ca²⁺ permeation, while at the same time remain within the Na⁺-selective voltage regime.
- $_{233}$ We selected an intermediate voltage of \sim -340 mV for investigating permeation in mono-cationic
- $_{234}$ solutions to ensure a sufficient number of traversals of both Ca²⁺ and Na⁺ ions in the monovalent-
- 235 selective regime.

²³⁶ Mechanistic insights into ion permeation in TRPM5 from mono-cationic solutions

- ²³⁷ We conducted *in silico* electrophysiology simulations with an applied electric field, generating a ²³⁸ membrane voltage of \sim -340 mV, to investigate the permeation mechanism of Na⁺, K⁺ and Ca²⁺
- ions in mono-cationic solutions at sufficient sampling efficiency (Table 3). As shown in Figure 4, we
- observed a clear difference between the behaviour of monovalent Na⁺ and K⁺ ions in the channel
- and the divalent Ca^{2+} ions, especially near and in the central cavity.

Table 2. Calculated conductances and selectivities from CompEL simulations of ion permeation in the TRPM5 channel. Mean inward conductances and standard error of the mean (SEM) were calculated from overlapping 50 ns windows from three-fold replicated 500 ns simulations of an anti-parallel double bilayer system. The number of permeation events associated with the conductance for each cation is displayed in brackets below the respective conductance values. Mean selectivity ratios of Na⁺ and Ca²⁺ permeation events and SEM were calculated from three-fold replicated 500 ns simulations.

Conc.	Voltage					
gradient	(mV)		P_{Na}/P_{Ca}			
gruurent		Na⁺	Ca ²⁺	Cl ⁻	Overall	
	-50	16 ± 3.1	0 ± 0.0	0 ± 0.0	16 ± 3.1 7 ± 1.3	∞
135 mM :	-50	(15)	(0)	(0)		
155 mM . 15 mM	-130	7 ± 1.3	0 ± 0.0	0 ± 0.0		∞
	-150	(18)	(0)	(0)		
	-380	4 ± 0.5	5 <u>+</u> 0.8	0 <u>+</u> 0.1	9 ± 1.0	1.9 ± 0.41
	-300	(32)	(19)	(1)		
	-610	10 ± 0.9	29 ± 2.5	-1 ± 0.2	38 ± 3.1	0.8 ± 0.13
	-010	(115)	(168)	(6)	50 <u>±</u> 5.1	

As can be seen, whereas Na⁺ and K⁺ ions occupied the central cavity of the channel for most 242 of the simulated time, permeating Ca^{2+} ions traversed the inner cavity rapidly, not exhibiting any 243 apparent immobilisation within the cavity. Despite occupying the cavity for extended periods of 244 time, Na⁺ and K⁺ ions did not seem to bind to a particular binding position or residue within the 245 cavity, but instead explored nearly the entire cavity volume before they permeated through the 246 lower gate to the intracellular side. In this way, the cavity serves to store a monovalent ion rather 247 than providing specific binding sites for it. A similar behaviour has recently been described for 248 simulations of Na⁺ ions in the cavity of the homo-dimeric, endo-lysosomal Na⁺-selective cation 249 channel TPC2 (Milenkovic et al., 2021). 250

Looking at the density of ions along the pore axis, and using the negative logarithmic density 251 as an estimate for the underlying free energy profile at the examined non-equilibrium permeation 252 conditions under a membrane voltage of -340 mV, it can be observed that the cavity region formed 253 only a shallow, broad energy minimum for the permeating monovalent cations, whereas in con-25/ trast, permeating Ca^{2+} ions experienced a small apparent energy barrier in the same region (Fig-255 ure 5). Both monovalent and divalent ions showed further binding to a relatively shallow binding 256 site at the EPV. In addition, all ion types experienced a slight energy barrier to translocation near 257 the intracellular channel exit (hydrophobic lower gate). Notably, the ions did not show major inter-258 actions with the SF. This observation, again, is in accordance with observations made in simulations 259 of TPC2 (Milenkovic et al., 2021). 260

We conducted additional simulations using applied external electric fields of differing magnitudes. Similar to the low-voltage CompEL simulations, whereas the main features of the ion den-

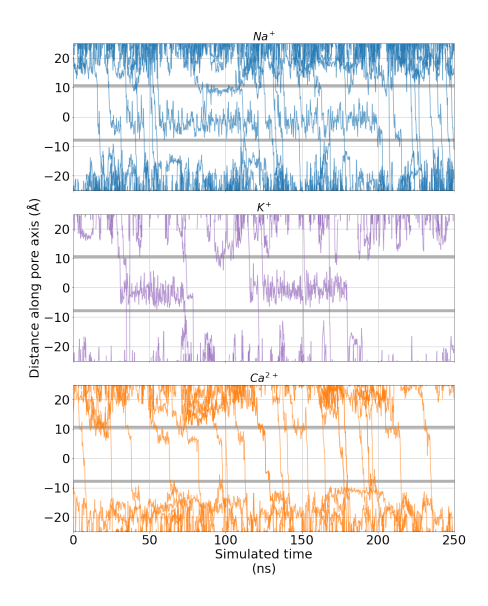


Figure 4. Exemplar permeation traces of the *z*-coordinate of permeating Na⁺ (*blue, top*), K⁺ (*purple, middle*), and Ca²⁺ (*orange, bottom*) over time, plotted from simulations performed in a mono-cationic solution with an applied electric field (-340 mV). The shaded grey regions represent the average position of the pore constrictions formed by Q906 in the SF (*upper*) and I966 of the hydrophobic gate (*lower*). Please note, only cations that fully permeate through the pore within the 250 ns simulations are shown in the plot.

- ²⁶³ sity and free energy estimates occurred across all tested voltages, Ca²⁺ was increasingly excluded
- ²⁶⁴ from the cavity at these lower voltages, and no longer able to enter into the cavity at the lowest
- voltage of -50 mV during the time span of our simulations (Figure S1). This again showed that the
- ion selectivity of TRPM5 was voltage-dependent in the simulations. We therefore next aimed to elu-
- cidate the molecular foundations of this behaviour and, importantly, the ion selectivity of TRPM5
- ²⁶⁸ in general.

> **Table 3.** Calculated conductances from applied field simulations of ion permeation in the TRPM5 channel. Mean inward conductances and standard error of the mean (SEM) were calculated from overlapping 50 ns windows from three-fold replicated 500 ns simulations of a single bilayer system. The number of permeation events observed for each cation is displayed in brackets below the respective conductance values.

Voltage (mV)	lon solution	Conductance (pS)		
	150 mM NaCl	19 <u>+</u> 6.7		
-50		(4)		
	150 mM CaCl ₂	0 ± 0.0		
		(0)		
	150 mM NaCl	17 ± 3.1		
-200		(15)		
	150 mM CaCl ₂	9 <u>+</u> 3.8		
		(4)		
	150 mM NaCl	52 <u>+</u> 5.6		
-340		(83)		
-540	150 mM KCl	21 ± 4.2		
		(34)		
	150 mM CaCl ₂	85 <u>+</u> 5.0		
		(54)		

269 Solvation profiles of cations during channel permeation

To probe if cation desolvation played a part in selective ion permeation, we calculated the num-270 ber of water oxygen atoms within a 3 Å radius around the ions, representing their first solvation 271 shell (Figure 5). Amongst other mechanisms (Ives et al., 2023; Zhang et al., 2023), the desolvation 272 of permeating ions has previously been reported to represent an important potential selectivity 273 mechanism in ion channels (Noskov and Roux, 2007; Kopec et al., 2018). Differences in the des-274 olvation energies of permeating ions provide a thermodynamic penalty which can underpin the 275 more favourable permeation of an ionic species over another. Here, the free energy required to 276 desolvate Ca²⁺ strongly exceeds that for Na⁺ and K⁺ (*Marcus, 1991*), such that this difference could 277 give rise to monovalent-selectivity in TRPM5. 278

In the bulk solution of the simulated systems, Na⁺, K⁺, and Ca²⁺ ions showed the expected water coordination number of their solvation shells. As both Na⁺ and K⁺ ions entered the pore of TRPM5, they became partially desolvated by Q906, with its side chain displacing 1–2 water molecules from the first solvation shell of the ions. After traversing the constriction at Q906, the monovalent ions were then resolvated in the pore cavity, before again being partially desolvated at the hydropho-

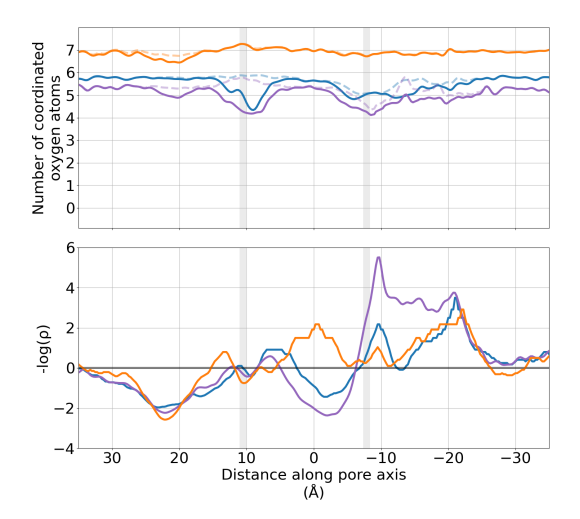


Figure 5. Solvation and log-density profiles of Na⁺ (*blue*), K⁺ (*purple*) and Ca²⁺ (*orange*) cations in the TRPM5 pore from simulations with a mono-cationic solution, and an applied transmembrane voltage of ~ -340 mV. (a) The mean number of oxygen atoms of water molecules (*solid line*) and of protein oxygen atoms (*dashed line*) within 3 Å of each permeating cation is shown. (b) Negative logarithmic density profiles of permeating cations as estimates of the non-equilibrium energy surface the ions experience in the pore (energy unit: k_B T). Minima reflect binding sites, while maxima indicate barriers between the binding sites. The location of the pore constrictions formed by Q906 (*upper*) and I966 (*lower*) are shown as grey regions. The curves in both plots have been smoothed using a Gaussian filter with a sigma value of 2.

- ²⁸⁴ bic lower gate formed by I966. By contrast, the rapidly permeating Ca^{2+} ions did not show any ²⁸⁵ significant desolvation when they crossed the SF, cavity, or hydrophobic lower gate of TRPM5.
- The solvation profiles of permeating cations displayed an additional region of differing desol-
- vation within the EPV region, highlighted previously (Figure 2). In this region, Ca^{2+} and K^+ ions
- were partially desolvated, indicating closer interactions with the acidic residues in the EPV region.
- ²⁸⁹ By contrast, Na⁺ cations did not show any desolvation in this location. We observed similar sol-
- vation profiles for permeating cations in both our simulations using an external applied electric
- ²⁹¹ field in mono-cationic solutions (Figure S2) and in the CompEL simulations in di-cationic solutions

- ²⁹² (Figure S3), across a range of voltage magnitudes.
- ²⁹³ Summarising, these findings suggest that ion desolvation in the SF or inner pore is not a major
- ²⁹⁴ factor in achieving selectivity for monovalent cations. Since both monovalent and divalent cations
- ²⁹⁵ occupied the EPV, filtering for monovalent ions must occur later in the permeation pathway. How-
- ²⁹⁶ ever, Ca²⁺ ions were not desolvated when they traversed the inner cavity. As the energetic penalty
- for desolvating Ca^{2+} is far larger than for Na^+ or K^+ (*Marcus, 1991*), the observed desolvation pro-
- files therefore suggest that desolvation does not underpin the deselection of Ca^{2+} ions in TRPM5.

²⁹⁹ Why does the central cavity form an attractive site for monovalent cations but a ³⁰⁰ repulsive site for divalent cations?

The presence of a water-filled internal cavity is a conserved feature amongst cation-selective chan-301 nels. The cavity serves to maintain a high degree of ion hydration despite locating to the centre 302 of the hydrophobic lipid bilaver, and to focus the membrane voltage difference onto the SF (Dovle 303 et al., 1998). Like in other cation channels, we observed in TRPM5 that the major permeating 304 species, Na⁺ and K⁺ ions, were re-hydrated and transiently captured in the cavity, following their 305 permeation through the SF (Milenkovic et al., 2021). At higher voltages, Ca²⁺ ions were able to enter 306 and traverse the cavity, but did not alter their hydration number during this process. This suggests 307 that they did not interact favourably with any of the cavity-lining residues or the cavity overall. At 308 lower voltages, by contrast. Ca^{2+} ions were completely excluded from entering the cavity. 300

We therefore investigated the difference between the pore and cavity properties of the highly 310 Ca²⁺-selective TRP channel, TRPV5, and the monovalent-selective TRPM5. Contrary to TRPM5, the 311 cavity of TRPV5 shows a high occupancy with Ca^{2+} ions (*lves et al., 2023*). As displayed in Figure 6, 312 the general features of the pore are preserved with a constriction at the extracellular SF. a wider 313 internal cavity region, and a second constriction at the intracellular gate. TRPM5 has a markedly 314 shorter SE, while its cavity is wider than that of TRPV5. However, there is a substantial difference 315 in the pore lining of the two TRP channels. Whereas the TRPV5 SF is a strongly hydrophilic region. 316 TRPM5 does not display increased hydrophilicity within its SF. The transition from the SF to the 317 cavity is slightly hydrophobic in TRPM5, while this is a hydrophilic region in TRPV5. There are no 318 differences between the hydrophobicity of the two channels at the intracellular gates. 310

The differing properties of the inner pore (cavity and SF) suggest that the absence of a favourable 320 interaction of TRPM5 with Ca^{2+} in this region arises due to the raised hydrophobicity of its SE and 321 upper portion of its inner cavity (hydrophobic funnel). In particular, the transition zone between 322 the SF and the cavity in TRPM5 is lined by large hydrophobic residues at the bottom of the SF. 323 especially F904 and I903. This sequence is shared with TRPM4, which is also a monovalent-cation 324 selective channel (Fig. 7A). On pore-forming helix S6 of TRPM5, the additional hydrophobic residues 325 V959 and 1962 line the cavity towards the hydrophobic lower gate at 1966, whereas only two polar side chains, N958 and N962, are involved. The conservation level of the large hydrophobic residues 327 lining the cavity is generally high (dark and light pink surface colour in Fig. 7B). 328

The energetic cost of placing monovalent cations into a hydrophobic environment is smaller than that for divalent cations, even when they retain their hydration shells. The Born energy for divalent cations, for example, quantifying this electrostatic energy penalty in continuum models, is

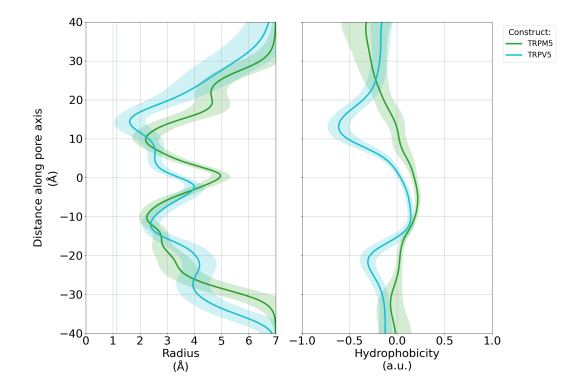


Figure 6. Pore architecture of the monovalent-selective TRPM5 channel (*green*) and the Ca²⁺-selective TRPV5 channel (*cyan*) from MD simulations. The average pore radius (**a**) and hydrophobic profile (**b**) for each channel was calculated using CHAP (**Rao et al., 2019**). The standard deviation is shown as shaded regions. The profile of the TRPV5 was generated from simulation data previously published (*lves et al., 2023*). The shaded grey regions represent the average position of the pore constrictions formed by Q906 in the SF (*upper*) and I966 of the hydrophobic gate (*lower*) in TRPM5.

³³² four times larger than the penalty associated with monovalent cations (*Born, 1920*). As observed in

the simulations under increased voltage, the protein matrix does not form favourable interactions

- $_{334}$ with Ca²⁺ ions in its inner pore. The hydrophobicity of the TRPM5 pore gradually increases along
- the pore axis, with only few hydrophilic sites within the SF or central cavity.

Our results thus suggest that Ca²⁺ ions are unable to enter the increasingly hydrophobic environment of this region at physiological voltages, and only penetrate past the SF under high-voltage conditions, above the physiologically relevant level.

³³⁹ Selectivity for monovalent cations is linked to permeation co-operativity between

340 two binding regions

- ³⁴¹ We hypothesised that, due to the presence of an additional binding or 'storage' region for mono-
- valent cations in the internal cavity compared to Ca²⁺, the permeation mechanism for monovalent
- ions may be more efficient than for Ca²⁺. In previous work, we developed a mutual-information
- based quantification method of the level of cooperativity when ions permeate across multiple ion
- channel binding sites, termed state-specific information (SSI; (Thomson et al., 2021; Vögele et al.,

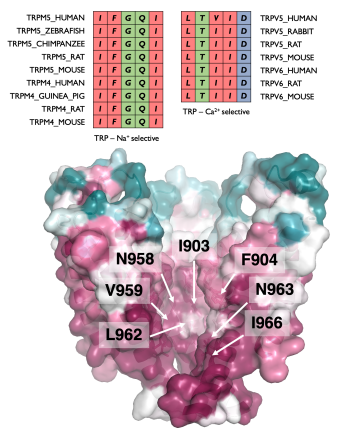


Figure 7. (Top) Sequence conservation of the SF and transition zone to the inner cavity in the monovalent-selective channels TRPM5 and TRPM4 compared to the Ca²⁺-selective channels TRPV5 and TRPV6. Colours according to Clustal Omega convention (*Sievers et al., 2011*). (Bottom) Evolutionary conservation of the pore cavity in TRPM5 channels. Evolutionary conservation scores were calculated using ConSurf (*Yariv et al., 2023*). The evolutionary conservation scores were projected onto the structure of TRPM5 from *D. rerio*, with one subunit removed for clarity. Figure made with Pymol (*DeLano, 2002*).

- 2022; Ives et al., 2023)). In brief, SSI quantifies the probability that a state change of one binding
- ³⁴⁷ site, such as a change from binding an ion to becoming vacant upon ion permeation, is correlated
- to a similar state change in a second binding site, in which case the unbinding events are coupled
- to one another for instance by a knock-on mechanism.
- We applied the SSI approach to ion conduction in TRPM5, focusing on the pair of ion binding regions at the EPV and the channel cavity (see Figure 2). These two binding areas are shallow and relatively distant to one another, but locate directly to the main pore axis. In addition, they show moderate to high occupancy with monovalent cations, respectively (Figure 8).
- By using SSI, we found that both Na⁺ and K⁺ ions displayed a high level of correlation between binding and unbinding at the two successive sites, whereas the permeation of Ca²⁺ ions showed only a low degree of correlation slightly above the noise level (Figure 8). This suggests that a distant
- ³⁵⁷ knock-on mechanism is in operation between incoming monovalent ions, which, when unbinding
- ³⁵⁸ from the EPV bind transiently at the SF, as well as over substantial time spans within the cavity. In
- other words, the cavity serves as a reservoir, more likely to release a Na⁺ or K⁺ ion to the cytoplasm
- when a further monovalent cation approaches and inserts into the TRPM5 pore.

- ³⁶¹ By contrast, Ca²⁺ ions permeated on their own. Their traversal is likely to be driven solely by the
- ³⁶² supra-physiological transmembrane electric field. Accordingly, lower-voltage simulations did not
- ³⁶³ show any permeating Ca²⁺. This selectivity for monovalent ions was abolished when the driving
- ³⁶⁴ force for Ca²⁺ permeation exceeded a certain threshold. Permeation at higher voltages could thus
- be described as a 'pull-through' of Ca^{2+} ions across the otherwise unfavourable environment of the
- inner cavity for divalent ions. This voltage-driven 'pull-through' occurs due to the higher charge of
- Ca²⁺, doubling the Coulombic driving force compared to the monovalent ions, but is unlikely to be
 physiological.
- Our results suggest that the occurrence of two binding regions for monovalent ions, one of which is a reservoir site in the central cavity, enhance permeation efficiency via a distant knock-on mechanism. By contrast, Ca²⁺ ions are prevented from entering the pore cavity at physiologically relevant membrane voltages by a hydrophobic gate region, abolishing the reservoir binding within and thus disrupting any sizeable permeation cooperativity.

The transition zone from the selectivity filter to the central cavity is key for monovalent cation selectivity in TRPM5

To test the hypothesis that the hydrophobicity of the transition zone connecting the SF to the 376 central cavity provides the primary barrier to the permeation of divalent cations, we performed 377 applied-field simulations of TRPM5 with a F904T mutation, which increases the hydrophilicity of 378 this area, at two voltage magnitudes. In the Ca²⁺-selective TRPV channels TRPV5 and TRPV6, a thre-379 onine residue is located at the structurally equivalent position. Since they also contain a charged 380 glutamate in their selectivity filter, which shows a high affinity for Ca^{2+} but is absent in TRPM5, we 381 did not expect the TRPM5 F904T mutant to show a similarly high degree of Ca²⁺-selectivity. Rather, 382 we hypothesised that the added hydrophilicity of the mutant may reduce the height of the hy-383 drophobic barrier for both monovalent and divalent cations, that is, increase the flow of Na⁺ while 384 allowing the passage of Ca^{2+} in physiologically relevant voltage ranges. 385

Three-fold replicated simulations of the F904T mutant in 150 mM CaCl₂ at -130 mV showed 13 completed Ca²⁺ permeation events within a total time of 750 ns. In the same time span, 27 Na⁺ ions traversed the mutant channel in 150 mM NaCl solution (Table S3). The permeation numbers correspond to a selectivity of P_{Na}/P_{Ca} of ~2, otherwise not observed below a voltage of -380 mV. Additional control simulations at -200 mV in mono-cationic solution displayed 15 Na⁺ and 4 Ca²⁺ permeation events in the WT (Table S2), whereas the mutant conducted 88 Na⁺ ions and 28 Ca²⁺ ions within the same accumulated time span (Table S3).

These results show that the F904T mutation indeed facilitated the permeation of Ca^{2+} , while at the same time increasing the Na⁺ flux. We conclude that the hydrophobic transition zone between the SF and the central cavity plays the major role in governing the selectivity of TRPM5 for monovalent cations at physiological voltages.

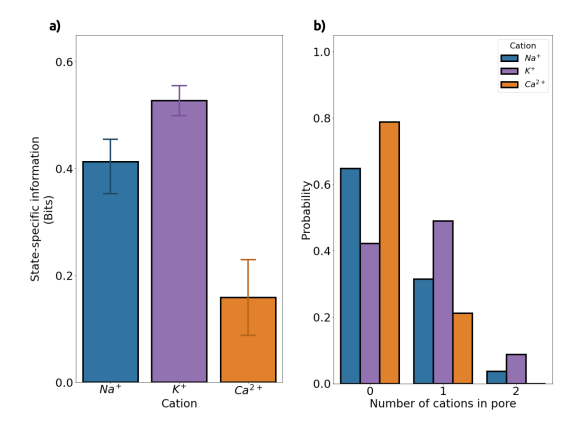


Figure 8. State-specific information (SSI) of cation transitions between binding sites and the average number of each cation within the pore of TRPM5. **(a)** Excess state-specific information (*exSSI*) between the EPV and pore cavity ion binding sites, quantifying the degree of co-operativity in the permeation mechanisms of Na⁺ (*blue*), K⁺ (*purple*), and Ca²⁺ (*orange*) ions. The mean exSSI and SEM between transitions from the two binding sites were calculated from simulations performed in mono-cationic solutions with an externally applied voltage of ~ -340 mV. **(b)** Mean probability for the number of each cationic species within the inner TRPM5 pore, calculated from non-overlapping 50 ns windows from three-fold replicated 250 ns simulations. We defined the inner pore as the region between the constrictions formed by Q906 in the SF and 1966 at the lower hydrophobic gate.

397 Discussion

The TRP channel superfamily encompasses a broad range of cation-selective ion channels of great 398 physiological and biomedical importance (Ramsey et al., 2006; Khalil et al., 2018). While most 300 members of the superfamily translocate both monovalent cations and divalent Ca²⁺ at similar per-400 meabilities, TRPV5 and TRPV6 are strongly Ca²⁺-selective (Ives et al., 2023) and TRPM4 and TRPM5 401 are selective for monovalent cations (Owsianik et al., 2006). In contrast to most other Na⁺-selective 402 channels (Dudev and Lim, 2014), the SFs of TRPM4, TRPM5, and the endo-lysosomal TPC2 do not 403 contain charged residues, while they retain a relatively high abundance of hydrophobic residues 404 (Ruan et al., 2021; Milenkovic et al., 2021). 405 Ion selectivity is usually linked to the presence of specific binding sites in the channels' SF and in-406 ner pore (Hille, 2001; Zhou et al., 2001), cooperativity between the permeation kinetics of multiple 407

- such binding sites (*Hille, 2001; Köpfer et al., 2014; Ives et al., 2023; Derebe et al., 2011*), ion desol-

vation (*Noskov and Roux, 2007; Kopec et al., 2018*), or size exclusion effects (*Hille, 2001; Dudev and Lim, 2014*). Our simulations showed, however, that no high-affinity binding sites for cations exist
within the pore of TRPM5, that only minor ion desolvation effects occur, and that size exclusion does not play a role in selectivity. The channel can conduct both monovalent and divalent cations at slightly increased membrane voltages. We observed two shallow, broad ion binding regions for monovalent cations in the TRPM5 channel; one within the EPV above the SF, and a second within the central cavity, whereas divalent cations did not interact favourably within the cavity.

Our findings suggest a new mechanism of monovalent cation selectivity, in which the combination of an uncharged, relatively hydrophobic SF and the presence of large hydrophobic side chains at the entrance to the central channel cavity determine the monovalent-selectivity of TRPM5. Since the traversal of a divalent cation through this hydrophobic funnel incurs a larger energy penalty as compared to a monovalent cation, this hydrophobic region creates a higher energy barrier for the permeation of divalent cations (*Born, 1920*). The hydrophobic energy barrier difference generated in this way is of moderate magnitude, and therefore can be overcome by divalent cations in the supra-physiological voltage range.

Under physiologically relevant voltages, this barrier and the generally largely hydrophobic char-424 acter of the central cavity prevent divalent cations from entering and residing within the central cav-425 ity, whereas monovalent cations readily enter and occupy the cavity for substantial time spans. Nu-426 merous closed-state structures of TRPM4, a close monovalent-selective homolog of TRPM5, have 427 been published within the PDB. Several of these structures include Na⁺ cations modelled within 428 the pore cavity (Guo et al., 2017: Duan et al., 2018), Consequently, these structures (PDB IDs 6BC). 429 6BCL, and 6BWI) suggest that the presence of a monovalent cation binding region in the inner 430 cavity is a conserved feature amongst the monovalent-selective TRPM4 and TRPM5 channels. 431

⁴³² Due to this additional binding region, a distant knock-on mechanism is established between ⁴³³ an incoming monovalent cation and the monovalent cation stored within the cavity, which greatly ⁴³⁴ increases permeation efficiency. By contrast, under supra-physiological voltage conditions, diva-⁴³⁵ lent cations are simply 'pulled through' the hydrophobic cavity on their own, exhibiting no inter-⁴³⁶ actions with the cavity matrix or other cations. Our application of the mutual-information based ⁴³⁷ SSI method (*Ives et al., 2023; Vögele et al., 2021*) to ions permeating through TRPM5 showed only ⁴³⁸ negligible cooperativity for Ca²⁺ permeation.

Finally, we examined whether a hydrophilic mutation at the entrance to the central cavity facilitated the flux of divalent cations through this region. According to the findings discussed above, this was indeed the case in the F904T mutant of TRPM5, with substantial Ca^{2+} permeation in a voltage range that did not allow for Ca^{2+} permeation in the WT channel. In line with a hydrophobic barrier that exists for both monovalent and divalent cations, but is larger for divalent cations, the flow of Na⁺ also increased in the mutant, raising its overall conductance level.

445 Acknowledgments

We thank the University of Dundee I.T. services for maintenance of the School of Life Sciences

high-performance computing (HPC) cluster which was utilised in this research.

448 Author contributions

- CMI and UZ conceived the idea and designed the computational study, CMI conducted and anal-
- 450 ysed the WT simulation data, ATŞ conducted and analysed the F904T mutant simulations, NJT anal-
- 451 ysed the SSI data, UZ supervised the work, CMI and UZ wrote the manuscript with contributions
- 452 from NJT, and all authors edited and reviewed the manuscript.

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626 Supplementary

627 Summary of MD simulations used within this study

Table S1. Summary of CompEL simulation details of the TRPM5 channel. All simulations were conducted in a di-cationic solution of 75 mM NaCl and 75 mM CaCl₂. In all simulations, the Ca^{2+} cations occupying the Ca_{TMD} were modelled, and remained bound for the duration of the simulations.

Protein	TRPM5						
	7MBS						
Structure	(698-1020)						
Force field			1M36m				
Water		TIF	23P				
In silico electrophysiology		Com	npEL				
methodology	(anti-pa	rallel with a 9:1		radient)			
			• 75mm CaCl ₂				
			IARMM36m)				
lon	274 Ca^{2+} (Zhang et al.)						
		814 Cl ⁻ (CH)	ARMM36m)				
Independent simulations	3	3	3	3			
Total simulation time (µs)	1.5	1.5	1.5	1.5			
Total aggregated simulation	3	3	3	3			
time (µs)							
lonic ratios between	239 : 27 Na⁺	239 : 27 Na⁺	239 : 27 Na⁺	239 : 27 Na⁺			
compartments	239 : 35 Ca ²⁺	239 : 35 Ca ²⁺					
(Extracellular : Intracellular)	681 : 133 Cl ⁻	680 : 134 Cl ⁻	677 : 137 Cl ⁻	673 : 141 Cl ⁻			
Estimated voltage (mV)	-50	-130	-380	-610			
	15 Na⁺	18 Na⁺	32 Na⁺	115 Na⁺			
Permeation events	0 Ca ²⁺	0 Ca ²⁺	19 Ca ²⁺	168 Ca ²⁺			
	0 Cl ⁻	0 Cl ⁻	1 Cl ⁻	6 Cl ⁻			
Total number of permeation	15	18	52	289			
events							

Table S2. Summary of external applied field simulation details of the TRPM5 channel. All simulations were conducted in a mono-cationic solution of either 150 mM NaCl, 150 mM KCl, or 150 mM CaCl₂. In all simulations, the Ca²⁺ cations occupying the Ca_{TMD} were modelled, and remained bound for the duration of the simulations.

Protein	TRPM5								
Structure		7MBS							
		(698-1020)							
Force field		CHARMM36m							
Water					TIP3P				
In silico electrophysiol	ogy								
methodology		External applied field							
		150 mM NaCl		150 mM KCl					
Le		267 Na+ (CHA	RMM36m)	267 K+ (CHARMM36m)			150 mM CaCl ₂		
lon		275 Cl [.] (CHAF	RMM36m)		(CHARMM36m)		1 Ca ²⁺ (Zhang <i>et al</i> .)		
		4 Ca ²⁺ (CHAF		4 Ca ²⁺ (CHARMM36m)		542 Cl ⁻ (CHARMM36m)			
Independent simulation	ons	3			3		3		
Total simulation time		0.75		0.75		0.75			
Estimated voltage (m			-340		-340		-340		
		83 N	a ⁺	34 K+		54 Ca ²⁺			
Permeation events		0 CI	-	0 Cl ⁻		0 Cl ⁻			
Total number of permea	ation								
events		83		34		54			
Protein			TRPM5						
Structure		7MBS							
			(698-1020)						
Force field				CHARMM36m					
Water				TIF	23P				
In silico electrophysiology				External a	oplied field				
methodology				450 1411 0					
		50 mM NaCl	150 mM	CaCl ₂	150 mM NaCl		150 mM CaCl ₂		
lon		a ⁺ (CHARMM36m)	271 Ca ²⁺ (Zhang e	ang et al.)	267 Na ⁺ (CHARMM3		271 Ca ²⁺ (Zhang et al.)		
		l ⁻ (CHARMM36m)	542 Cl ⁻ (CHAI	RMM36m)	275 Cl ⁻ (CHARMM36m)		542 Cl ⁻ (CHARMM36m)		
Independent simulations	4 Cd-	2+ (CHARMM36m)			4 Ca ²⁺ (CHARMM36m)		3		
Total simulation time (µs)	0.75		3 0.75		0.75		0.75		
Estimated voltage (mV)	-50		-50		-200		-200		
	-50 4 Na*		-50 0 Ca ²⁺		15 Na ⁺		-200 4 Ca ²⁺		
Permeation events		4 118 0 Cl ⁻	0 Cl ⁻		0 Cl ⁻		0 Cl ⁻		
Total number of permeation									
events		4 0		15			4		

Table S3. Summary of external applied field simulation details of the TRPM5 F904T channel. All simulations were conducted in a mono-cationic solution of either 150 mM NaCl, or 150 mM CaCl₂. In all simulations, the Ca^{2+} cations occupying the Ca_{TMD} were modelled, and remained bound for the duration of the simulations.

Protein	TRPM5 F904T								
Structure	7MBS (698-1020)								
Force field		CHARMM36m							
Water		TIF	23P						
In silico electrophysiology methodology	External applied field								
lon	150 mM NaCl 267 Na+ (CHARMM36m) 275 Cl- (CHARMM36m) 4 Ca2+ (CHARMM36m)	150 mM CaCl2 271 Ca2+ (Zhang et al.) 542 Cl- (CHARMM36m)	150 mM NaCl 267 Na+ (CHARMM36m) 275 Cl- (CHARMM36m) 4 Ca2+ (CHARMM36m)	150 mM CaCl2 271 Ca2+ (Zhang et al.) 542 Cl- (CHARMM36m)					
Independent simulations	3	3	3	3					
Total simulation time (μs)	0.75	0.75	0.75	0.75					
Estimated voltage (mV)	-130	-130	-200	-200					
Permeation events	27 Na+ 0 Cl-	13 Ca2+ 0 Cl-	88 Na+ 0 Cl-	28 Ca2+ 0 Cl-					
Total number of permeation events	27	13	88	28					

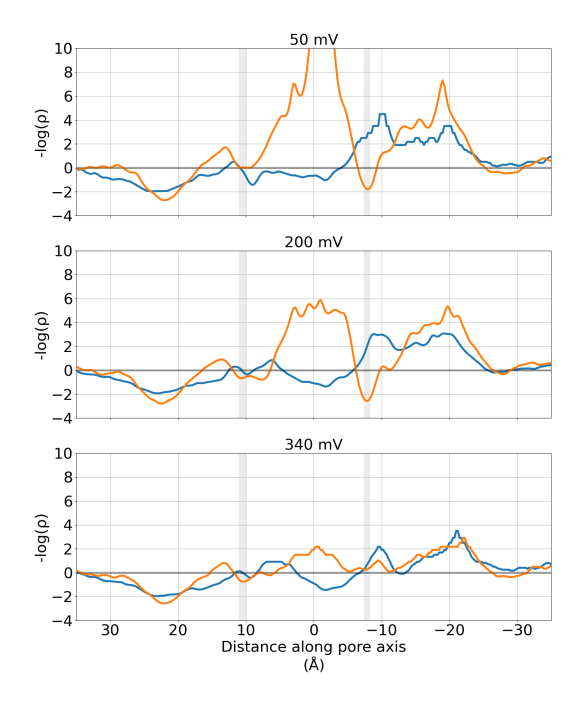


Figure S1. Negative logarithmic density profiles of permeating cations along the pore of the TRPM5 at different voltages. These simulations were performed in a mono-cationic solution, with an external applied electric field used to produce transmembrane voltages of ~ -50 mV (*top*), ~ -200 mV (*centre*), and ~ -340 mV (*bottom*). The logarithmic ion densities represent quasi-free energies (with a nominal unit of kT). The location of the pore constrictions formed by Q906 (*upper*) and I966 (*lower*) are represented as grey regions. Both plots have been smoothed using a Gaussian filter with a sigma value of 2.

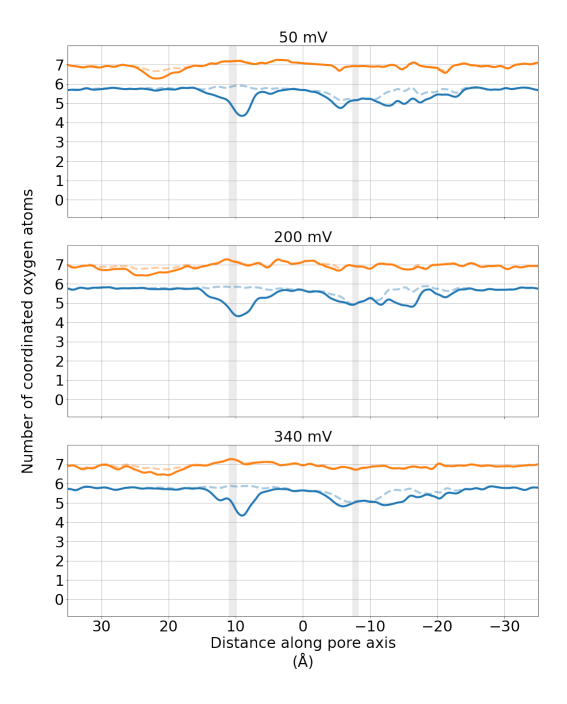


Figure S2. Solvation profiles of Na⁺ (*blue*) and Ca²⁺ (*orange*) cations through the TRPM5 pore. These simulations were performed in a mono-cationic solution, with an external applied electric field used to produce transmembrane voltages of ~ -50 mV (*top*), ~ -200 mV (*centre*), and ~ -340 mV (*bottom*). The mean number of oxygen atoms of water molecules (*solid line*) and of any oxygen atoms of any molecule (*dashed line*) within 3 Å of each permeating cation is plotted. The location of the pore constrictions formed by Q906 (*upper*) and I966 (*lower*) are represented as grey regions. All plots have been smoothed using a Gaussian filter with a sigma value of 2

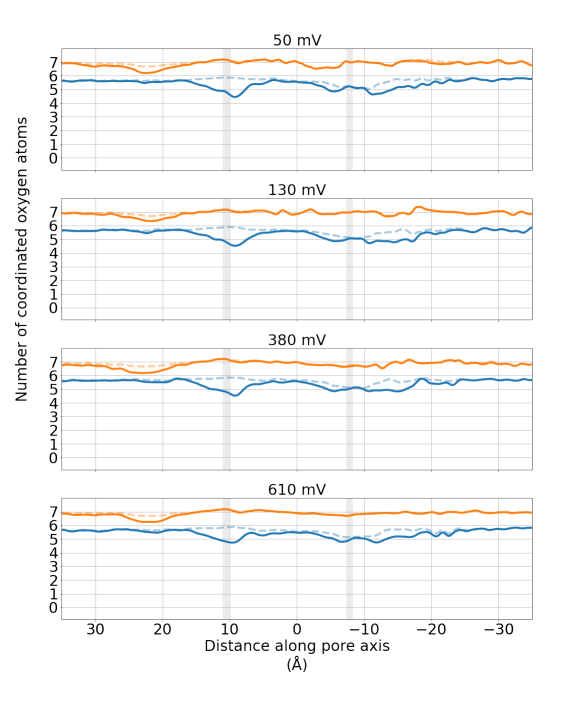


Figure S3. Solvation profiles of Na⁺ (*blue*) and Ca²⁺ (*orange*) cations through the TRPM5 pore. These simulations were performed in a di-cationic solution, with the CompEL methodology used to produce transmembrane voltages of ~ -50 mV (*top*), ~ -130 mV (*second from top*), ~ -380 mV (*second from bottom*), and ~ -610 mV (*bottom*). The mean number of oxygen atoms of water molecules (*solid line*) and of any oxygen atoms of any molecule (*dashed line*) within 3 Å of each permeating cation is plotted. The location of the pore constrictions formed by Q906 (*upper*) and I966 (*lower*) are represented as grey regions. All plots have been smoothed using a Gaussian filter with a sigma value of 2.