



Inducing nonlinear conductance and emergent memristance in open pores using blockers

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Memristance in biomembrane systems is an emerging area of research relevant to biocomputing. Work has so far focused on voltage-driven insertion of pore-forming peptides in droplet interface bilayer systems, design of asymmetric pores, and lipid membrane poration. However, only a few instances of always-open, pore-forming channels have been used to design and study the emergence of memristive behaviors in biomimetic membranes. In this work, we investigate the Outer Membrane Protein F (OmpF) as a model nanopore. Due to its nanoscale pore size (~1 nm diameter), OmpF can be transiently or partially blocked by a variety of molecules, including antibiotics, polyamines, and polypeptides. We reconstituted OmpF into droplet interface bilayers and examined electrical signatures of different blocking agents using cyclic voltammetry. We show that binding of some of these agents to pore-lining residues induces strongly nonlinear current–voltage (I – V) responses, which, in some instances, give rise to memristive behavior. We find that the interaction of ampicillin and spermine with OmpF at the tested conditions was insufficient to produce nonlinearity or memristance. However, interaction between arginine polypeptides and OmpF leads to length and charge-dependent nonlinear conductance with emergent memristance in the case of arginine nonapeptide. Molecular dynamics simulations showed that arginine pentapeptides (Arg-5) block the pore longitudinally as a stretched chain whereas bulkier and highly charged arginine nonapeptides (Arg-9) block the pore transversally by compaction of the peptide chain. Although both blocking conformations were sufficient to exhibit nonlinear dynamics, only the transversal blocking conformation led to

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emergent memristance, likely due to longer-lived interactions between binding sites and the Arg-9.

Introduction

In recent years, with the advancement of artificial intelligence, demand for energy required for computation has increased exponentially, and global power demand for computation is expected to surpass global power production by 2030.¹ Traditional computing devices follow the Von Neumann architecture, where the processing unit and memory unit are two separate entities.² Data needs to be continuously transferred from one unit to another.² This data transfer, which can cost up to 1000 times that of actual computation costs, is referred to as the Von Neumann bottleneck.³ The inefficiency caused by this bottleneck motivates research into devices where memory and processing units can be co-located.

Memristive devices or memristors^{4,5} are two-terminal devices in which memory and processing units are co-located.⁶ The conductance of these devices depends on their history of voltage and current.⁵

Leon Chua and co-workers showed that potassium and sodium ion channels in the Hodgkin–Huxley axon could be described as memristors.⁷ This work, along with others, has inspired work on biological memristors as a building block for brain-inspired computing hardware, including ion channel-doped biomembrane-based memristors.^{8–17}

Alamethicin-doped membranes have been extensively studied in droplet interface bilayer (DIB) systems.^{8,12,13,15} Alamethicin peptides form transmembrane ion channels following voltage-driven insertion into lipid membranes. In alamethicin-doped membranes, memristance emerges from the time difference between insertion and exit of the pore-forming peptides from lipid membranes as well as the time delay in area and thickness changes of the membrane due to voltage, which influences the total number of open channels that conduct ions.⁸ There are limited examples in the literature of non-voltage-gated protein channels that act as memristors. In one study, Koner *et al.* showed that a gramicidin-doped biomembrane serves as a memristor where memristance emerges from the dynamics of the reconfiguration of gramicidin.⁹ Paulo *et al.* demonstrated how hydrophobically gated nanopores can act as memristors.¹⁶ Very recently, Mayer *et al.* reported on open beta-barrel proteins (aerolysin, alpha-HL, and MspA) exhibiting memristance because of gated ion transport governed by lumen charge.¹⁷

OmpF is one of the major diffusion pores in the outer membrane of *E. coli*.¹⁸ OmpF is a homotrimer where each monomer is made up of 16-stranded anti-parallel beta-barrels.¹⁹ The L3 loop joins β -strands 5 and 6 of the barrel and forms the constriction zone in each pore.¹⁹ Literature reports suggest that OmpF itself exhibits gating behavior above certain voltage thresholds, but this behavior is highly dependent on experimental conditions²⁰ with varying voltage threshold values reported in the literature.^{20,21}

OmpF is also extensively studied for its antibiotic binding and transport properties as it is one of the pathways for antibiotic entry into the cell.^{18,22,23} There are conflicting reports on the antibiotic ampicillin blocking the OmpF pore.^{18,24,25} Earlier studies indicated ampicillin binding and blocking of OmpF pores,^{18,25} but



it was later shown that ampicillin shows negligible blocking of the OmpF pore. Rather, it was the degraded ampicillin product (penicilloic acid) that is actually responsible for OmpF blocking.²⁴

There are also studies to understand the interaction between OmpF pores and polyamines, where spermine (a type of polyamine) significantly reduces the activity of OmpF pores.^{26,27} Spermine reduces the activity of OmpF not by plugging the pore, but rather by stabilizing the closed conformation of the pore.²⁶

Lamichhane *et al.* studied the binding of arginine polypeptide chains with OmpF²⁸ and reported no translocation. However, another study using secondary blocker molecules attached to the periplasmic side of OmpF established that tri-arginine indeed translocates through OmpF whereas hepta-arginine is bulky enough to preclude translocation through OmpF pores.²⁹ We hypothesized that memristance could be induced in an open pore, such as OmpF, using voltage-responsive blocker molecules, inspired by voltage-gated ion channels, in which the probability of channel opening and closing produces memristance.⁷ This functionality is produced by blocker molecules that act as a pseudo-gate and the probability of blocking and unblocking of the molecule (dictated by its residence time on specific residues) will produce emergent memristance. In this work, we exploited the blocking of OmpF using various blocker molecules and attempted to induce memristance in an OmpF-doped DIB biomembrane system. We also conducted molecular dynamics simulations to understand the binding of blocker molecules with OmpF. To the best of our knowledge, this work is the first demonstration of the use of ion channel extrinsic blocking to induce memristance.

Methods

Materials

1,2-Diphytanoyl-*sn*-glycero-3-phosphocholine (DPhPC) lipids (25 mg ml⁻¹ in chloroform) were obtained from Avanti Research. Ampicillin sodium salt and spermine were purchased from Sigma-Aldrich, arginine peptides were custom synthesized by GenScript Biotech.

Preparation of vesicle solution

25 mg ml⁻¹ of DPhPC stock in chloroform was added to a round-bottom flask along with 2–3 ml of additional chloroform. A rotary evaporator was used to evaporate chloroform under reduced pressure and to prepare thin lipid films. The flask was then placed in a vacuum oven at room temperature for ≥ 2 hours to ensure complete removal of chloroform under reduced pressure. The residual lipid film was rehydrated using the appropriate volume of buffer (100 mM potassium chloride (KCl), 10 mM 3-(*N*-morpholino)propanesulfonic acid (MOPS), pH 7.0) and extruded (11 passes) using Avanti's mini extruder to form an aqueous vesicle solution at a target lipid concentration of 2 mg ml⁻¹. A 0.2 μ m PC membrane was used for extrusion.

Purification of OmpF

OmpF was expressed and purified according to procedures reported before.³⁰



Blockers tested

The blockers tested included ampicillin (net charge at pH 7(n): -0.75),²⁴ spermine ($n = +4$), tri-arginine peptide (Arg-3) ($n = +3$), penta-arginine peptide (Arg-5) ($n = +5$), hepta-arginine peptide (Arg-7) ($n = +7$), and nona-arginine peptide (Arg-9) ($n = +9$).

Preparation of electrodes

Rounded Ag/AgCl electrodes are needed to securely interact with the droplets within the droplet interface bilayer (DIB) system. The tips of the 0.127 mm diameter silver wire (Ag) electrodes were rounded by melting. Then, silver/silver chloride (Ag/AgCl) electrodes were prepared by dipping the ends of Ag electrodes into bleach for at least 20 minutes. The tips of these electrodes were then dipped in melted agarose to coat their ends and make them hydrophilic. The electrodes were connected to a patch clamp amplifier (HEKA, EPC 10). The electrode tips were then submerged in a hexadecane oil bath. The positions of the electrodes were manipulated by using micromanipulators.

Formation of droplet interface bilayers

400 nL of aqueous vesicle solution was pipetted onto each of the electrode tips in oil. After allowing 10–15 minutes for monolayer formation around the droplets, we used micromanipulators to bring the droplets into contact to form bilayers. Once the bilayer formed, we waited 3 minutes for it to stabilize before investigating its voltage-responsive properties.

Insertion of OmpF in droplet interface bilayer

Vesicles were premixed with OmpF (at a lipid to protein mass ratio (LPR) of 2000 : 1) and blocker molecules (different concentrations) before dispensing the droplet vesicles onto the electrodes. For inserting OmpF into DIB membranes, a droplet of prepared mixture of vesicles with OmpF and blocker molecules was added to the source electrode and a droplet of vesicles without any OmpF or blocker molecules was added to the grounded electrode (with the exception of the ampicillin case, where a droplet of vesicles with OmpF was added to the source electrode and a droplet of vesicles with blocker molecules was added to the ground electrode). Droplet interface bilayers were formed as described above. Then three DC voltage pulses of -100 mV each were applied for 120 s (with a wait time of 5 min in between).

For experiments of inducing memristance in the OmpF pore, pre-insertion of OmpF in the bilayer before memristance testing is desirable. To achieve this, the relative net charges of the protein and blocker molecules were utilized. OmpF is negatively charged and was added to the droplet attached to the source electrode, whereas blocker molecules were added to the source side or ground droplet, depending on whether they were positively charged or negatively charged, respectively (Table 1 and Fig. 1). We surmised that applying a negative potential at the source electrode would attract negatively charged molecules to the ground electrode, whereas positively charged molecules would move in the opposite direction. This procedure ensured that the initial few sweeps could be used to insert OmpF in bilayers, whereas the blocker molecule would move away from the bilayer, hence essentially separating the insertion of OmpF dynamics from the



Table 1 List of protein and blocker molecules along with charge, side added, and direction of force experienced

Protein/blocker molecules	Net charge at pH 7	Side in which protein/blocker molecules are added	Direction of force experienced during initial sweeps for OmpF insertion
OmpF	Negative	Source	Towards bilayer
Ampicillin	Negative	Ground	Away from bilayer
Spermine	Positive	Source	Away from bilayer
Arg-3	Positive	Source	Away from bilayer
Arg-5	Positive	Source	Away from bilayer
Arg-7	Positive	Source	Away from bilayer
Arg-9	Positive	Source	Away from bilayer

blocking dynamics of the blocker molecules. A similar approach to driving molecules towards bilayers has been used before in the literature²⁹

Electrophysiology

A HEKA EPC 10 amplifier was used to conduct electrophysiological studies, including current–voltage (I – V) sweeps. For the memristance tests in particular,

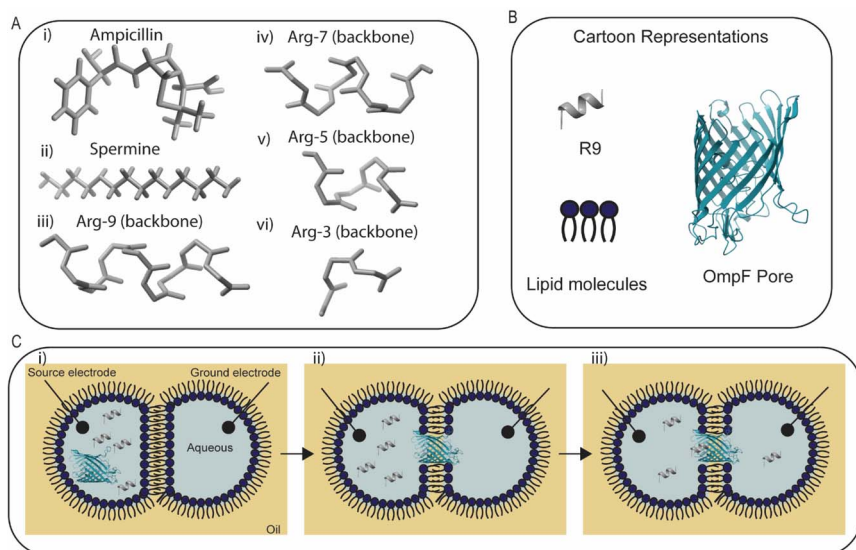


Fig. 1 Chemical structures of protein and peptide molecules, along with the schematic of the process of open pore blocking. (A) Chemical structures of (i) ampicillin, (ii) spermine, and backbone structure of arginine polypeptides as (iii) Arg-9, (iv) Arg-7, (v) Arg-5, (vi) Arg-3. (B) Representations of the molecules used in the schematic in part (C). (C) Schematic of the process of OmpF blocking using chained arginine molecules: (i) 2 mg per ml DPhPC DIBs containing buffer (100 mM KCl, 10 mM MOPS, pH 7) as well as OmpF and blocker molecules are added on the source electrode and 2 mg per ml DPhPC DIBs containing only the same buffer are added to the ground electrode. (ii) Initially, only a negative DC potential is applied so that only OmpF channels get inserted into the bilayer without affecting the blocker molecules. (iii) Binding dynamics of the blocker molecules were investigated by applying triangular voltage sweeps. UCSF ChimeraX was used to show the structure of protein and blocker molecules.



zero-mean continuous triangular voltage sweeps were applied on the droplet interface bilayer system with an amplitude of ± 150 mV and varying sweep rates. A 1 kHz Bessel filter was used during the measurement. A 100 Hz display filter was applied for post-processing of the data.

Molecular dynamics simulations

The initial coordinates for the OmpF porin (PDB ID: 2OMF) embedded in a 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC) bilayer were generated using the CHARMM-GUI Membrane Builder.³¹ Three distinct systems were constructed: (i) OmpF in POPC (control), (ii) OmpF/POPC with an Arg5 peptide (Arg-5), and (iii) OmpF/POPC with an Arg9 peptide (Arg-9). Peptides were placed in the extracellular vestibule, and overlapping water molecules were removed using a 2 Å distance cutoff. The systems were solvated with TIP3P water molecules and neutralized with NaCl to mimic experimental ionic strength conditions.

Simulations were performed using NAMD 3.0 with the CHARMM36m force field. Additional CHARMM-based equilibration was performed, plus 10 ns of equilibration before introducing the peptides. Long-range electrostatic interactions were computed using the Particle Mesh Ewald (PME) method, and van der Waals interactions were treated with a force-based switching function between 10 and 12 Å.

A timestep of 2 fs was used, and a constant temperature of 303.15 K was maintained using Langevin dynamics. Pressure was regulated at 1.01325 bar using the Nosé–Hoover Langevin piston method. A constant external electric field of -0.5 kcal (mol Å e)⁻¹ was applied along the *z*-axis, corresponding to a trans-membrane potential of approximately 2.5 V.³² While this potential exceeds experimental voltages, it is a standard approach in porin simulations to accelerate sampling and overcome the free energy barriers of ion/peptide transport within computationally accessible timescales.^{32–35}

For each of the three systems (control, Arg5, and Arg9), three independent simulation replicas were executed for 15 ns each, starting from different initial velocity distributions to ensure statistical robustness.

Results and discussion

A bilayer without any protein acts as a capacitor and allows very minimal ion transport, as illustrated in Fig. 2A, and the current–voltage (*I*–*V*) curve shows an almost flat horizontal line showing minimal transport and large capacitance. We conducted preliminary screening experiments with six distinct blocker molecules (representing three classes: antibiotics, polyamines, and polypeptides) to study OmpF blocking and their ability to induce nonlinear behaviors and memristance in OmpF-based DPhPC lipid membranes using the DIB method (Fig. 1). These blockers included ampicillin, spermine, tri-arginine peptides (Arg-3), penta-arginine peptides (Arg-5), hepta-arginine peptides (Arg-7), and nona-arginine peptides (Arg-9). As seen in Fig. 2, *I*–*V* sweeps for preliminary evaluation of memristive behavior showed no signs of emergent memristance or distinctive nonlinear behaviors for the OmpF–membrane system in the presence of ampicillin (0.13 μM) or spermine (0.3 mM) (blocker concentrations cover the range of



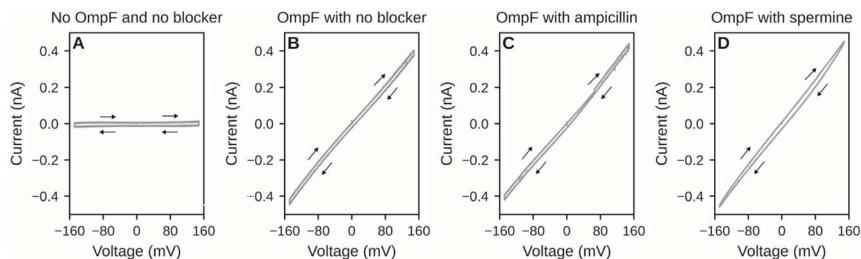


Fig. 2 I - V sweeps for zero mean triangular voltage input show no emergent nonlinear behavior or emergent memristance for ampicillin or spermine molecules driven through OmpF pores. (A) The I - V sweep for the DIB with no OmpF and no blocker molecules showing minimal transport. (B) The I - V sweep for the DIB with OmpF pore, but no blocker molecules, produces a straight line. (C) The I - V sweep for the DIB containing the OmpF pore, along with ampicillin molecules. Here, we also observe almost linear I - V curves representing no/negligible memristance as evidenced by a lack of a pinched hysteresis loop. (D) The I - V sweep for the system containing the OmpF pore with spermine molecules. Here, we do observe a slightly different curve, but no memristive behavior. The sweep rate applied to generate A was 10 mV s^{-1} , and for the other three plots, 20 mV s^{-1} . The area between the curves is because of the large capacitance of the bilayer, as explained elsewhere³⁶ (typical values of capacitance for these experiments were 400 – 800 pF , except for the spermine case, where that value was 50 – 60 pF).

blocker to pore molar ratio of ~ 5 to $11\,000$). Nonlinear I - V behaviors and emergent memristance were observed with arginine peptides of increasing length (Fig. 3 & 4).

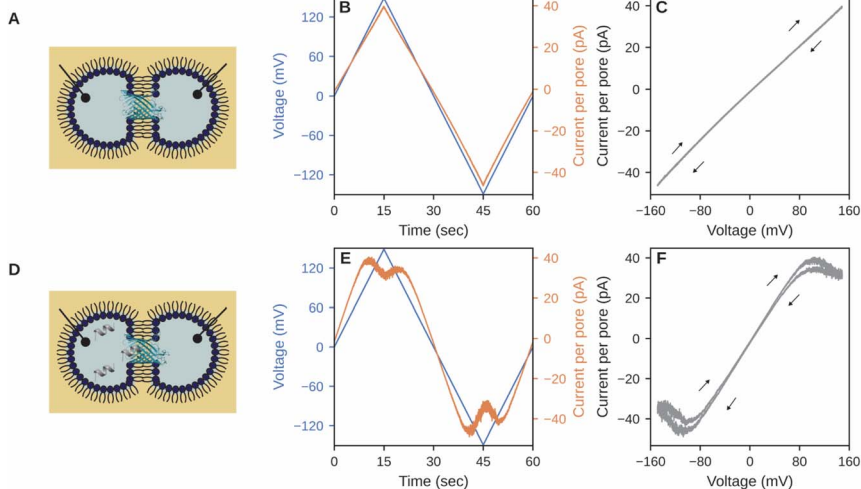


Fig. 3 Current response for the triangular sweep voltage applied to the DIB with and without Arg-9 molecules. A–C represent the experiments where the droplet attached to the source electrode contains OmpF protein channels (LPR 2000 : 1) but doesn't contain any blocker molecules. D–F represent experiments in which the droplet attached to the source electrode contains both OmpF protein channels (LPR 2000) as well as blocker molecules (Arg-9, 0.003 mM concentration). The buffer used for all the cases is 100 mM KCl , 10 mM MOPS , $\text{pH } 7.0$. Sweep rate is 10 mV s^{-1} . Current per pore was calculated by dividing the total current response by the number of pores inserted (Fig. S2). Sweep-wise variation is shown in Fig. S3.



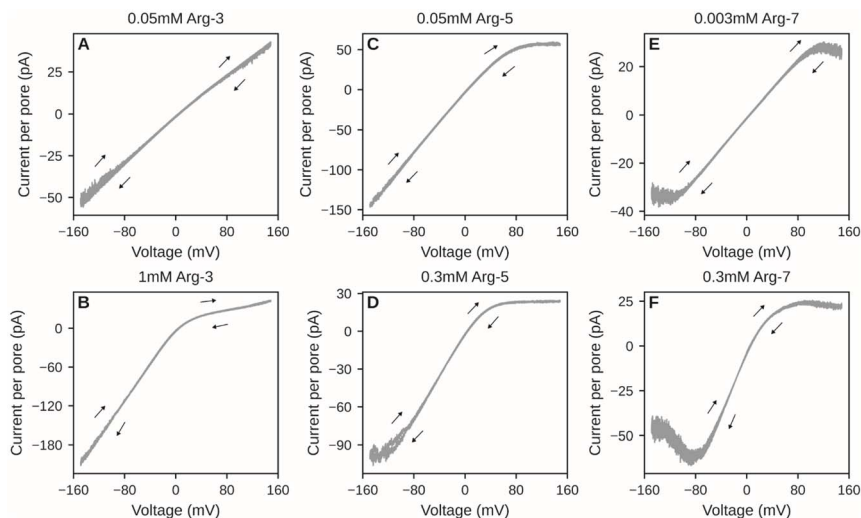


Fig. 4 Current per pore vs. voltage curves with different arginine chain lengths used as blockers of OmpF pore with input of zero mean triangular voltage sweeps showed increasing nonlinearity with longer chain lengths and concentrations of arginine polypeptides. (A) I - V curve with 0.05 mM Arg-3 concentration showed little blocking. (B) I - V curve with 1 mM Arg-3 concentration showed blocking during the positive sweep but no memristance. (C) I - V curve with 0.05 mM of Arg-5 concentration showed blocking during the positive sweep but no memristance. (D) I - V curve with 0.3 mM of Arg-5 concentration showing blocking during the positive sweep. (E) I - V curve with 0.003 mM of Arg-7 concentration showing blocking in both directions. (F) I - V curve with 0.3 mM of Arg-7 also showed blocking in both directions. Current per pore was calculated as mentioned above, and the number of pores was estimated in Fig. S4.

Emergence of nonlinear I - V curves and memristance by OmpF blocking by Arg-9

OmpFs inserted into the bilayer allow conductive transport of ions, resulting in a linear I - V profile (in this case with minimal contribution of capacitance) (Fig. 3A-C). Arginine is a positively charged amino acid and can interact with negatively charged residues on the L3 loop within the OmpF pore (Fig. S1). The interaction of Arg-9 with OmpF (Fig. 3D) induced nonlinearity and emergent memristance in the system. When the voltage was increased from 0 to 150 mV, initially the current also increased linearly, indicating nearly constant conductance (Fig. 3E). But above a certain threshold voltage, the conductance decreased with increasing applied voltage. This trend indicates a tendency for increased blocking with increasing voltage, presumably because of the blocker molecules being driven to and then interacting strongly with particular charged residues within the pore. In the reverse cycle from 150 mV to 0 mV, the conductance increased (indicated by the increase in slope of the I - V curve) till it reached the value reached in the 0 mV to 150 mV direction. This reversible change in conductance indicates voltage-dependent blockage of the OmpF pore. It also indicates that a persistent potential is required to keep the pore blocked, and reducing the potential initiates the unblocking process. The I - V curve of the protein pore blocked with Arg-9 molecules at a concentration of 0.003 mM



showed a hysteresis loop (Fig. 3F) and a symmetric, pinched I - V hysteresis loop, meeting one of the main criteria for classifying systems as possessing emergent memristance. To confirm whether the hysteresis is memristive rather than a distortion caused by capacitive currents, we varied the sweep rate of the triangle voltage waveform to test for hysteresis at various frequencies. Fig. S5 shows the results that capacitive I - V characteristics are only observable at very high frequencies (>100 mV s $^{-1}$). When we adjusted for these by subtracting the capacitive current associated with a DIB device, we observed hysteresis effects at higher frequencies. We hypothesize that Arg-9, because of its higher charge and larger size, binds in a manner that the blocking and unblocking kinetics become sufficiently differentiated, leading to memristance.

A similar pattern is observed during the negative voltage cycle (0 to -150 mV and back). It is worth noting that we did not add any blocker to the other side of the membrane (*i.e.*, the opposite droplet). Further investigation is needed to understand the reasoning behind this observed phenomena.

Emergence of nonlinear dynamics by Arg-3, Arg-5, and Arg-7

OmpF exhibited nonlinear I - V behaviors in the presence of Arg-3, Arg-5, and Arg-7 at certain concentrations, although they did not result in significant I - V hysteresis (Fig. 4). Arg-3 at a concentration of 0.05 mM showed some transient blocking, but this blocking was not sufficient to significantly alter the linearity of the I - V curve (Fig. 4A). However, at 1 mM, Arg-3 induced a nonlinear response in the OmpF I - V trace at positive voltages, but no memristance (Fig. 4B). Arg-5 and Arg-7 also induced nonlinear I - V behavior, although they also did not exhibit memristance (Fig. 4C-F). We observed minimal blocking in the case of Arg-3 and Arg-5 during the negative voltage cycle (0 to -150 mV and back), whereas we observed blocking during the negative voltage cycle for Arg-7 (similar to Arg-9 as discussed above). According to literature, Arg-3 can translocate through the pore, whereas Arg-7 does not.²⁹ Our MD simulation results also show that Arg-5 could translocate across the pore, whereas Arg-9 did not under the conditions simulated. Literature results combined with our MD simulations suggest that although Arg-3 and Arg-5 can translocate through the pore, the concentration on the other side of the pore does not reach a level during I - V sweeps to result in nonlinear IV relationships at negative voltages. Similar to the case of Arg-9, the reason for blocking observed during the negative cycle for Arg-7 is unclear and further investigations are necessary to provide a convincing explanation for this observation.

Theoretical model

We developed a probabilistic model inspired by the Hodgkin-Huxley model³⁷ to describe ion channel dynamics underlying OmpF blocking and the resulting IV behavior. In this model, current can be simply modeled by single channel conductance, the number of channels, and their open probability:

$$I = G_u A N_a (1 - p) V$$

G_u is conductance per channel (monomer), A is membrane area, N_a is areal channel density, and p is probability of channel being closed at any point. The



probability, p , of blocking the OmpF pore in a DPhPC membrane can be modeled with a first-order linear differential equation as

$$\frac{dp}{dt} = \beta(p_{ss}(V) - p)$$

where β is the rate constant of pore blocking, and p_{ss} is the steady state probability of pore blockage at a particular voltage. p_{ss} is a static function of the transmembrane voltage, which we define as

$$p_{ss} = \begin{cases} 0, & V < V_T \\ \alpha(|V| - V_T), & V_T \leq V \leq V_M \\ 1, & V > V_M \end{cases}$$

where α is the proportionality constant representing the dependency of pore blocking probability on voltage. V_T is the threshold voltage after which pore blocking starts. V_M is the voltage at which probability of pore blockage becomes 1. The blocking of pores is responsible for the nonlinearity of the curve (hence it is dependent on α), whereas the memristance is dependent on how quickly the probability reaches its steady state and hence is dependent on β .

Plots in Fig. 5 are results from the model with assumed values of α and β . A value of single channel conductance, $G_u = 0.28$ nS,³⁸ was approximated from literature (shown in SI) and used, and the number of inserted channels per bilayer ($A \times N_a$) was assumed as 100, as the number of pores inserted varied mostly in the range of 20–200 for our experiments, as shown in Fig. S2 and S4. In Fig. 5A, the curves resulting from a low value of α indicate that the probability of blocking at steady state is lower compared to the higher value of α , resulting in curves seen in Fig. 5B and C. A higher value of β indicates that dp/dt , the rate of change of probability of blocking, is high, and thus the probability of blocking reaches its steady state relatively rapidly. Therefore, no memristance is observed with the higher value of β compared to the lower value of β , where we do observe memristance (indicated by the pinched hysteresis loop).

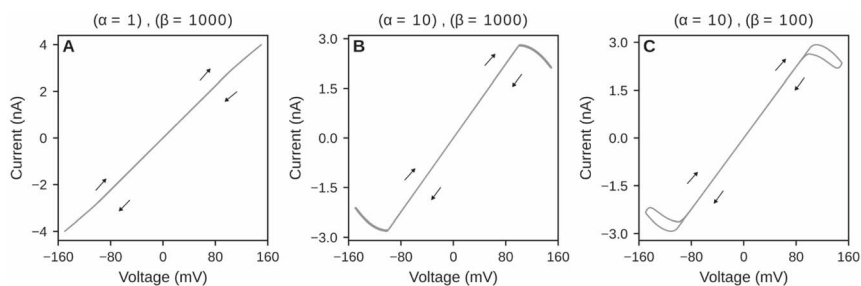


Fig. 5 Our model reproduces the behavior seen with OmpF pores and arginine blockers using chosen values of model parameters. I – V curves with different parameter values of α and β . (A) Low value of α and high value of β results in a linear profile, indicating no blocking and no memristance. (B) A higher value of α with a high value of β results in blocking but no memristance. (C) If we decrease the β value, keeping the α value the same, we observe the emergence of memristance.



Molecular dynamics simulation

Molecular dynamics simulations of a single OmpF pore reveal a pronounced reduction in ionic flux across the pores blocked with arginine peptide variants relative to the control. MD inherently samples short timescales, limiting the number of ion conduction events that can be observed (a conductance of 300 pS corresponds to ~ 2 crossings per 10 ns window). To obtain better statistics within the accessible simulation timeline, we applied a transmembrane voltage equivalent to 2.8 V. The use of unphysiologically high voltages in our MD simulations is consistent with prior studies.^{32,33,39–42} We lightly restrained lipid phosphorus atoms and C α atoms of the equilibrated protein, a stabilization strategy that preserves lipid bilayer, and the protein's secondary structure while allowing side-chain mobility.^{43,44} Under the simulation conditions, the system without any peptides sustains a steady ionic current of 880 ± 21 pA, whereas the OmpF pore in the presence Arg-5 and Arg-9 peptides exhibited substantial current suppression, with mean currents of 240 ± 15 pA and 208 ± 15 pA, respectively (Fig. 6). Notably, during the initial ~ 2 ns, the currents observed in both Arg-5 and Arg-9 trajectories closely follow the same trend as the control simulation. However, this transient conductive phase is quickly followed by a rapid transition to a predominantly blocked state, marked by extended periods of reduced flux punctuated by rare ion-translocation events.

Visualizations of the trajectories showed that while both peptides, Arg-5 and Arg-9, reside within the beta-barrel, they exhibit distinct conformational behaviors within the protein pore. Arg-9 exhibits a compact, globular structure that stably occupies the central pore axis, generating a large steric and electrostatic barrier to entering the constriction created by loop L3. In contrast for Arg-5, the peptide unfolds and a subset of arginine residues (Arg1, Arg2, and Arg3) bind in

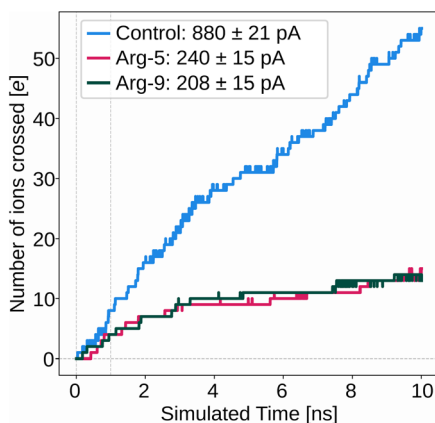


Fig. 6 MD simulations can reproduce current blockades under different conditions. The curves show the total number of ions that successfully crossed the membrane over the simulated time. The blue line represents the control system (no peptides present), and exhibits a steady ion flux corresponding to an average current of 880 ± 21 pA. In contrast, the addition of peptides Arg-5 (magenta) and Arg-9 (dark green) significantly attenuates ion transport, reducing the current to 240 ± 15 pA and 208 ± 15 pA, respectively.



the pore lumen, and constrict the pore along the pore length rather than just at the entrance (Fig. 7).

These distinctions are further highlighted by plotting the radius of gyration (R_g) and end-to-end distance (R_{ee}) distributions (Fig. 8). For Arg-9, the trajectory shows a clear transition toward a collapsed, globular state, where the R_{ee} drops significantly over time (from ~ 18 Å to < 10 Å) while R_g remains tightly clustered around 6.8 Å. This quantitative shift mirrors the visual transition into a stable, globular plug. Conversely, the Arg-5 plots reveals a very gradual increase in the end-to-end distance, indicating the “unfolding” behavior observed in the

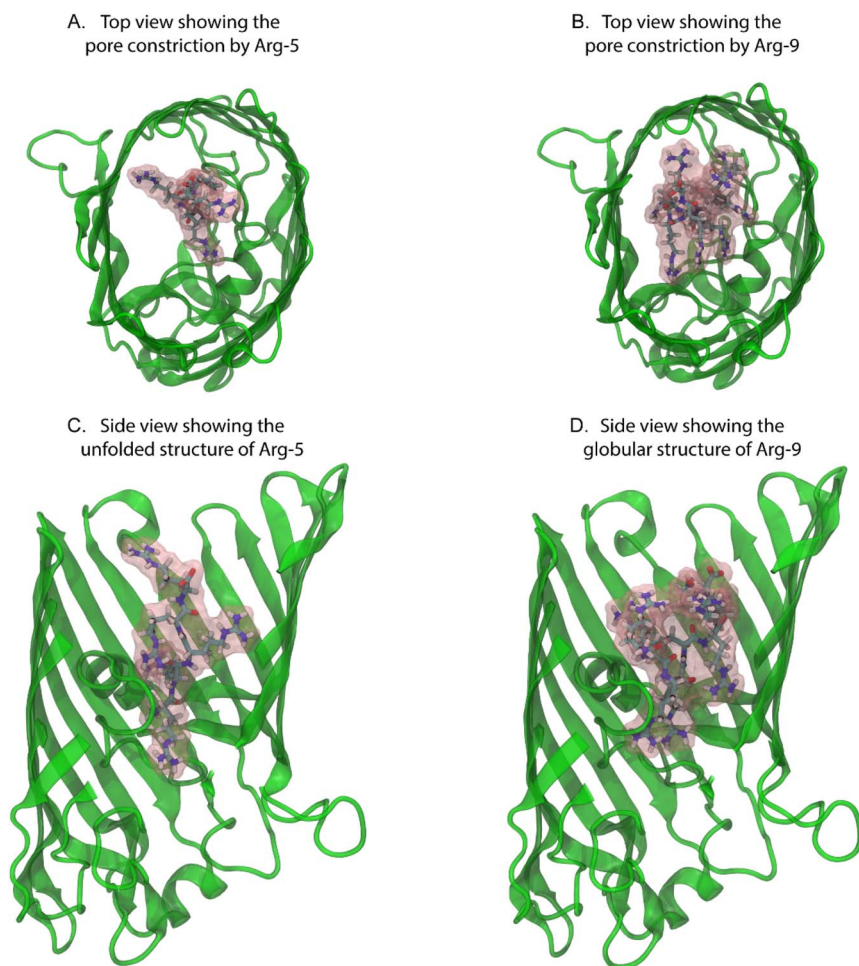


Fig. 7 Structural basis for pore constriction by Arg-5 and Arg-9 peptides inside the OmpF lumen. (A and B) Top-down views of the pore lumen showing the physical occlusion caused by the presence of (A) Arg-5 and (B) Arg-9. The peptides (pink surface/licorice) reside within the channel interior, significantly reducing the effective pore radius available for ion permeation. (C and D) Longitudinal side views illustrating the distinct conformational states of the peptides. (C) Arg-5 adopts a more elongated, unfolded structure that spans a larger vertical section of the pore, whereas (D) Arg-9 maintains a more compact, globular structure.



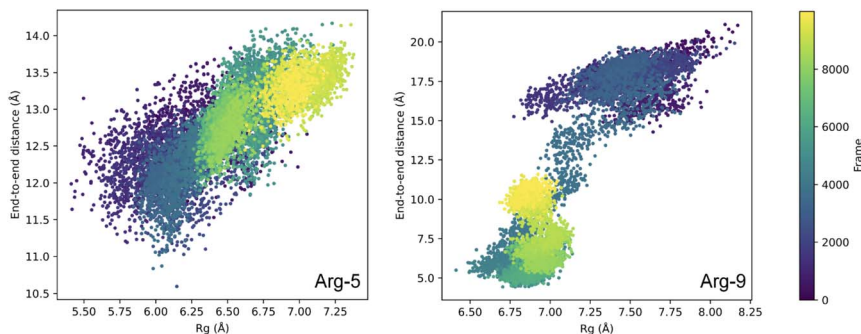


Fig. 8 Conformational analysis of Arg-5 and Arg-9 peptides during pore occlusion. Scatter plots illustrate the relationship between the radius of gyration (R_g) and end-to-end distance (R_{ee}) over the simulation trajectory. Arg-9 shows a transition toward a collapsed, globular state characterized by a significant drop in R_{ee} from 18 Å to <10 Å. In contrast, Arg-5 exhibits a gradual increase in R_{ee} , indicating an “unfolding” behavior where the peptide stretches its backbone while remaining within the barrel lumen. These distinct structural metrics confirm that Arg-9 blocks the pore via a dense steric bulk, while Arg-5 achieves occlusion through multi-point constriction along the pore axis.

visualizations. The peptide remains physically small enough to fit within the barrel but stretches its backbone to allow side chains of Arg1–3 to protrude into the lumen. Together, these metrics confirm that Arg-9 blocks the pore through a single, dense steric bulk, whereas Arg-5 achieves occlusion through a more elongated, multi-point constriction along the pore axis.

The functional consequence of these structural differences is most evident under high applied voltage (equivalent to 3.5 V), where a complete translocation event was observed for Arg-5 (Fig. 9). In this trajectory, the longitudinal orientation of the Arg-5 peptide allowed it to thread through the constriction zone and eventually escape into the bulk solvent on the *trans* side. This exit resulted in a full recovery of the ionic current to levels indistinguishable from the control (pore-only) simulations. Arg-9, on the other hand, remained securely lodged within

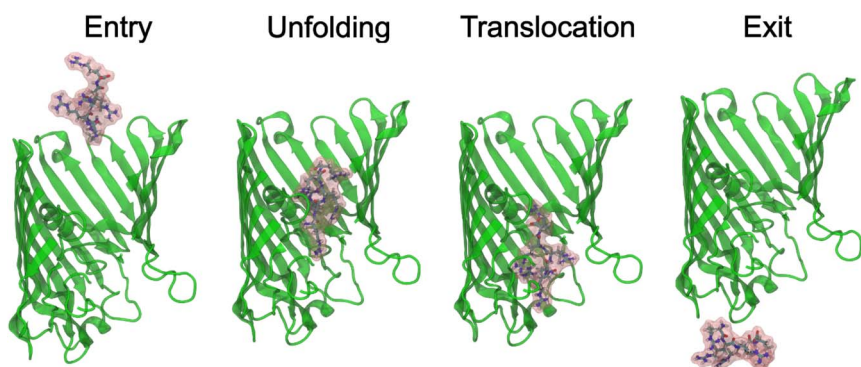


Fig. 9 Translocation mechanism of Arg-5 under higher applied voltages. The time-series snapshots illustrate the sequential stages of the Arg-5 peptide passing through the membrane channel under higher applied voltage in the MD simulations (3.5 V).



the barrel throughout the same simulation timescale. These results indicate that while Arg-5 can effectively constrict the pore at lower potentials, its 'unfolded' binding mode is more susceptible to voltage-driven translocation than the compact 'plug' formed by Arg-9.

Overall, MD simulations reveal a difference in the configuration of Arg-5 vs. Arg-9 when blocking the OmpF pore. Both molecules clearly demonstrated a drop in conductance when they block the pore. However, as shown in the R_g vs. R_{ee} plot (Fig. 8) Arg-5 polypeptide adopts an elongated, unfolded conformation that occupies the pore longitudinally, whereas Arg-9 collapses into a compact globular conformation, occluding the pore transversely. Fig. 7 illustrates this difference in conformation taken by Arg-5 vs. Arg-9 when blocking the pore. These results suggest that both Arg-5 and Arg-9 can block the OmpF pore and we believe that this blocking leads to nonlinearity in the I - V curves. However, the binding mode for smaller peptides like Arg-5 is strikingly different from the binding mode of Arg-9. Transversal blocking of the pore by Arg-9 potentially introduces a lag in unbinding kinetics and results in emergent memristance. On the other hand, longitudinal blocking of the pore by smaller peptides like Arg-5 does not create any lag in the unbinding process, and hence we did not observe emergent memristance. Further work is needed to explain how this difference in binding mode may result in emergent memristance.

Conclusions

Three different classes of blocker molecules were investigated for their potential to induce nonlinearity and pinched hysteresis (a signature of emergent memristance) in OmpF-doped biomembranes. Whereas ampicillin and spermine were unable to produce nonlinearity and memristance in the system, bulkier and highly charged arginine polypeptides (Arg-3, Arg-5, Arg-7, and Arg-9) were able to induce nonlinearity in the OmpF doped biomembrane. Some emergent memristance was also observed in the case of Arg-9. MD simulations indicate that both Arg-5 and Arg-9 can block the OmpF pore and reduce the conductance. However, simulations also indicated a difference in blocking conformation for Arg-5 and Arg-9. Where Arg-5 unfolded and blocked the OmpF pore longitudinally, Arg-9 collapsed into a globular structure and blocked the pore transversally. Both blocking conformations have resulted in nonlinear dynamics, but emergent memristance in Arg-9 is likely the result of transversal and likely long-lived blocking of the pore, whereas longitudinal blocking in the smaller peptide (Arg-5) was unable to lead to such emergent memristance. Further investigation is required to understand how this difference in binding conformation translates to emergent memristance. A theoretical probabilistic model was also built and used to show the transformation of linear dynamics to nonlinear dynamics by varying the α (proportionality constant) parameter and the transformation of nonlinear dynamics to memristance by varying the β (rate constant of pore blocking) parameter. This study shows that open pores can exhibit memristance dynamics in the presence of blocker molecules.

Looking forward, the results of this work contribute to the wider application of artificial membrane systems mimicking biological neurons, which rely on ion channel dynamics, both voltage gating and molecule-binding mediated, to produce signalling. The device presented in this work utilizes a blocking



mechanism that is among a growing library of literature examples of ion channel-based dynamics that seek to emulate neuronal and synaptic signalling. Furthermore, like other biologically derived computing systems, our device operates in the nanoampere and millivolt range, resulting in significantly lower power consumption than silicon-based devices. The field of biomimetic nanofluidics for neuromorphic applications seeks to leverage the low-power signalling of ion channel-based devices to achieve the computational efficiency of the brain.

Author contributions

R. D. and M. K. conceptualized the project. R. D., L. L., K. G. and M. R. performed the experiments. P. R. G. performed the molecular dynamics simulations. N. X. A. developed the theoretical model. R. J. V., K. G. and D. B. purified the protein. A. F., H. B. and S. S. helped with experimental design and manuscript editing. R. D., P. R. G. and M. K. performed the analysis and wrote the manuscript with the help of other co-authors.

Conflicts of interest

There are no conflicts to declare.

Data availability

All the data used in this manuscript have been uploaded to Texas Data Repository and can be accessed using the following link: <https://doi.org/10.18738/T8/MTA7H4>.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d6fd00008h>.

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