Large-Conductance Transmembrane Porin Made from DNA Origami

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ABSTRACT: DNA nanotechnology allows for the creation of three-dimensional structures at nanometer scale. Here, we use DNA to build the largest synthetic pore in a lipid membrane to date, approaching the dimensions of the nuclear pore complex and increasing the pore-area and the conductance 10-fold compared to previous man-made channels. In our design, 19 cholesterol tags anchor a megadalton funnel-shaped DNA origami porin in a lipid bilayer membrane. Confocal imaging and ionic current recordings reveal spontaneous insertion of the DNA porin into the lipid membrane, creating a transmembrane pore of tens of nanosiemens conductance. All-atom molecular dynamics simulations characterize the conductance mechanism at the atomic level and independently confirm the DNA porins’ large ionic conductance.

KEYWORDS: DNA origami, lipid membrane, synthetic porin, ionic current recordings, molecular dynamics

Due to their diverse architectures, protein channels in natural lipid membranes are capable of fulfilling a variety of functions in living cells, from the recognition of substrates to the selective transport of ions or large biomolecules between cellular compartments.1 Synthetic channels have been proposed as components of drug-delivery systems, as antimicrobial agents, biosensors, filters, photosystems, catalysts,2 or as tools for synthetic biology;3 all undoubtedly requiring tailored architectures with a high level of customizability. Efforts to create synthetic channels started three decades ago with the first account by Tabushi et al.4 and are still ongoing.5 The architectural variability of biological membrane proteins, spanning 1 order of magnitude in channel diameter and 3 orders of magnitude in molecular weight and conductance, remains however widely unexplored and often inaccessible due to limitations of chemical synthesis methods especially for large synthetic channels.2 DNA has previously been used as an alternative highly stable and readily available chemical block for building transmembrane pores,6–11 but the pore’s architectures have so far been limited to the following three pore types: a bundle of six DNA duplexes with a nominal inner channel diameter of 2 nm,6–9,11 a bundle of four duplexes with a 0.8 nm channel,10 and a single membrane-spanning duplex which induces DNA-lipid channels at its circumference.12 Reported conductances of these DNA channels range from 0.112 to 1.6 nS.9,11

Here we expand the design space of synthetic lipid membrane pores beyond these limits by creating a significantly larger funnel-shaped porin from DNA origami.13 Made exclusively using off-the-shelf components, our DNA origami porin overcomes limitations of traditional chemical synthesis that finds creation of large channels challenging.2 The nominal cross section of the DNA porin is 6 nm, Figure 1B, which is wider than the cross section of large natural porins14 and comparable to the electrical diameter of the nuclear pore complex.15 Previously, funnel-shaped large-diameter DNA origami nanopores were inserted into SiN apertures,16 however insertion of large-diameter nanopores into lipid membranes has not been described until now.

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RESULTS AND DISCUSSION

Design and Characterization of the DNA Origami Porin. The design of our DNA origami porin derives from that of the very first DNA nanopore,16 which has been modified to accommodate 19 cholesterol anchors to facilitate insertion into a lipid membrane. The 5 MDa DNA origami porin was assembled using the 7249 base-long M13mp18 scaffold and 179 single-stranded DNA staples. For the DNA origami layout, positions of cholesterol anchors, and DNA sequences see Supporting Information, Figures S1–S4 and Tables S1–S6. Agarose gel electrophoresis yields a sharp band and confirms the stability of the structure in the measurement buffer (Supporting Information, Figure S5). Atomic force microscopy (AFM) measurements, Figure 1C, clearly resolve the funnel shape of the DNA origami nanopore, including its three segments with an average length of 20.8 ± 2.5, 23.6 ± 2.1, and 11.0 ± 1.7 nm (n = 10) for the wide, the middle, and the narrow sections of the porin structure, respectively, resulting in a total measured length of 55.4 nm. These values are in good agreement with the designed dimensions (20, 24, 10 nm; total length: 54 nm). For a detailed description of the image analysis see Supporting Information, Note S1, Figures S6, S7, and Tables S7, S8. Assuming geometrical packing and a 2 nm diameter of the DNA helix,17 the designed square segments have a width and height of 22 × 22, 18 × 18, and 10 × 10 nm². The measured dimensions (46.7 ± 2.0 × 2.67, 35.8 ± 2.0 × 5.95, and 18.9 ± 2.0 × 2.67 nm²) show the collapse of the hollow funnel when imaged in air. Consequently, the middle segment of the funnel (which is made of two layers of DNA) is roughly twice as high as the other segments (which are made from a single DNA layer).

Confocal Fluorescent Imaging of Membrane Attachment. To probe effective attachment of the DNA origami porin to lipid membranes, we carried out confocal fluorescent imaging experiments. For this purpose, three Cy3-labeled fluorescent DNA oligomers were incorporated during assembly of the funnel-shaped DNA porin at its wider end. The same construct was prepared without the cholesterol membrane anchors as a negative control. After addition to giant unilamellar vesicles (GUVs), bright rings appeared around the vesicles in the confocal plane for the cholesterol-tagged sample, Figure 2A, whereas no membrane adhesion was observed for the control sample, Figure 2B.

Ionic Current Recordings. We subsequently carried out ionic current recordings in solvent-containing membranes18 to prove the membrane-insertion capabilities of the synthetic DNA porin and to determine its ionic conductance. As shown in Figure 3A, multiple high-conductance insertion steps were observed. A stepwise decrease in conductance, Figure 3B, could be caused by DNA porins escaping from the membrane. Additional insertion and closure traces are presented in the Supporting Information, Figures S8 and S9. The log-scale histogram obtained from such conductance steps, Figure 3C, is rather broad. The mean stepwise current increase associated with a DNA porin insertion lies at 30 nS in 1 M KCl, 10 mM 2-(N-morpholino)ethanesulfonic acid (MES), pH 6.0. A stepwise decrease in conductance, Figure 3B, could be caused by DNA porins escaping from the membrane. Additional insertion and closure traces are presented in the Supporting Information, Figures S8 and S9. The log-scale histogram obtained from such conductance steps, Figure 3C, is rather broad. The mean stepwise current increase associated with a DNA porin insertion lies at 30 nS in 1 M KCl, 10 mM 2-(N-morpholino)ethanesulfonic acid (MES), pH 6.0. The widespread of conductance values is likely to be caused by...
multiple factors. Previous MD simulations have suggested that the conductance of DNA pores can be influenced by membrane pressure.\(^{19}\) Hence, a certain amount of variability is expected among experiments which involve breaking and reforming the membrane. Agarose gel electrophoresis (Supporting Information, Figure S5) indicates the presence of dimerized DNA origami porins (\(-15\%\)), which could account for higher conductances if they are capable of inserting jointly. Deviations from perpendicular insertion orientation with respect to the lipid bilayer plane is likely to account for further variability.

To determine the current−voltage (I−V) characteristics of the DNA porin, only insertions which were stable across the voltage range of \(\pm 100\) mV and for the duration of the I−V recording were taken into account. I−V characteristics, Figure 3D, were found to be largely ohmic between \(-50\) and \(+50\) mV. The dashed line represents a linear fit with a gradient of \(20\) nS for this subset of stable insertions. A corresponding trace is shown in the Supporting Information, Figure S10. Deviations from the linear behavior are often observed above \(\pm 50\) mV: The DNA origami porins can switch to lower conductance states or disappear entirely as the voltage is increased, in line with previous observations for smaller DNA-based membrane pores.\(^{9,10}\) Rarely, the conductance increases at higher voltages while the current exhibits large fluctuations. Although some insertions could be stable for tens of minutes (Supporting Information, Figures S8E, S10), transient insertions or attempts prevail in ionic current recordings (Supporting Information, Figure S9D), contributing to the width of the reported conductance histogram.

**MD Simulations.** To independently evaluate the ionic conductance of our synthetic DNA-based porin, we built an all-atom model including all components of the experimental system: the DNA origami porin itself, the DphPC lipid bilayer, ions, and water molecules, Figure 4A, resulting in a system of 7,963,516 atoms. In the first 48 ns of the equilibration simulation, the DNA porin was restrained to its initial coordinates, allowing the membrane and the solvent to adopt an equilibrium configuration. The restraints were gradually removed over 14.4 ns. The system was subsequently simulated in the absence of restraints for another 19.2 ns. The DNA porin structure after free equilibration simulation is shown in the Supporting Information, Figure S12. During equilibration simulations, lipid molecules rearranged around the transmembrane part of the DNA porin, forming a water-filled passage along the DNA porins’ outer surface, Figure 4B. The formation of such water-filled passages has previously been reported for a transmembrane-spanning porphyrin-tagged DNA duplex.\(^{12}\) The process of lipid rearrangement is shown as a time series in the Supporting Information, Figure S11. Approximately 900 water molecules, or 50 per 180° segment of each DNA duplex, surrounded the DNA porin within a 1 nm thick rectangular slab centered at the middle plane of the membrane. The interior volume of the channel was filled with solution as well. This gives rise to an effective electrical diameter of approximately 11 nm for the DNA porin, making it larger than the electrical diameter proposed for the nuclear pore complex (6 nm).\(^{15}\) The equilibrium ion concentration outside the DNA porin was close to the target bulk values (1.1 M KCl and 20 mM MgCl\(_2\)); the concentration of Mg\(^{2+}\) was found to vary considerably within the system, increasing to 0.6 M within the DNA porin’s walls, Supporting Information, Figure S12.

To evaluate its ionic conductance, the DNA porin was simulated at +100, +30, −30, and −100 mV transmembrane biases, reproducing the experimental voltage range. The duration of each MD simulation was 19.2 ns at \(\pm 100\) mV and 48 ns at \(\pm 30\) mV, which was sufficient to observe statistically significant displacement of ions within the MD trajectories.\(^{19−21}\) Supporting Information, Movie S1, described biases, reproducing the experimental voltage range. The duration of each MD simulation was 19.2 ns at \(\pm 100\) mV and 48 ns at \(\pm 30\) mV, which was sufficient to observe statistically significant displacement of ions within the MD trajectories.\(^{19−21}\) Supporting Information, Movie S1, described biases, reproducing the experimental voltage range. The duration of each MD simulation was 19.2 ns at \(\pm 100\) mV and 48 ns at \(\pm 30\) mV, which was sufficient to observe statistically significant displacement of ions within the MD trajectories.\(^{19−21}\) Supporting Information, Movie S1, described.
in Note S2, illustrates the MD trajectory of the system. The cumulative charge transmitted across the lipid bilayer over time is plotted in Figure 4C. Around 80% of ions flow through the central pore, while ion flow along the outside of the channel contributes 20% to the total conductance. Figure 4D shows the histogram of the simulated conductance with a mean of 46.6 nS (dashed line, for additional conductance histograms see Supporting Information, Figure S13).

Although the width of the simulated conductance histogram is similar to that obtained from our experiments, the time scale of our MD simulation precludes us from making a definitive statement about the possible origin of broad conductance histograms. At the time scale of 50 ns, a considerable fraction of the ionic current noise is produced by thermal fluctuations. Another factor is the deformation of the funnel structure and fluctuations of the toroidal lipid pore, which can be appreciated from the Supporting Information, Movie S1, and snapshots of the equilibrated structure, Supporting Information, Figure S1A, B. 32% of the experimentally recorded insertion steps fall outside the conductance range obtained from MD simulations. Assuming that the simulated range is correct, this could represent the fraction of pores that inserted in an orientation different from that considered in MD simulations.

In absolute numbers, 23 of the experimentally obtained insertion steps had lower conductances, potentially due to angled insertion, whereas 3 exhibited higher conductances, potentially due to the insertion of dimers. The I–V curve obtained from simulations, Figure 4E, is ohmic and yields an average conductance of 46.6 nS. The simulated conductance is thus in very good agreement with the experimental data, taking into account the difference in the access resistance conditions and systematic overestimation of bulk electrolyte conductance in the simulation.19

Coarse-Grained Simulations. To determine if the gain in free energy produced by incorporation of cholesterol anchors into a lipid bilayer membrane can indeed compensate the free energy penalty associated with the formation of a lipid pore, we estimated the free energy of pore formation using the MARTINI coarse-grained model,23 which is known to semiquantitatively reproduce the mechanical properties of lipid bilayers.24,25 Starting from an equilibrated lipid bilayer system, a pore of radius $R_p$ was created by applying a cylindrical half-harmonic wall potential to lipid molecules.25 Figure 5A. Figure 5B plots the average pressure exerted by the lipid molecules on the potential wall as a function of the pore radius $R_p$. The pressure increases initially reaching 300 bar for $R_p = 0.5$ nm, then decreases monotonically as $R_p$ increases, leveling off at ~20 bar. The work required to form a pore of radius $R_p$ can be determined by integrating the pressure–volume $(pV)$ work, see Methods section. Previous MD studies estimated the insertion free energy of a single cholesterol molecule into a lipid bilayer at $\Delta G_{CHL} = -75$ kJ/mol.26 For the DNA porin insertion to be favorable, the sum of the pore formation work, $W_f(R_p)$, and the insertion free energy of N cholesterol anchors, $\Delta G_{CHL}(N)$, must be negative. Figure 5C plots the value of the sum as a function of the pore radius and the number of cholesterol anchors. For the DNA porin system reported in this work, $N = 19$ and $R_p \sim 5.5$ nm (determined from the lipid headgroup density, Supporting Information, Figure S11), so $W_f(R_p) + \Delta G_{CHL}(N)$ is roughly $-400$ kJ/mol, a value favoring insertion of the DNA porin into a lipid bilayer.

CONCLUSION

In the present study, we have demonstrated a synthetic DNA membrane porin with the largest conductance known to date. Electrophysiological experiments and MD simulations have shown that the conductance of our DNA origami porin is over an order of magnitude larger compared to all previous DNA membrane pores. In nature, such high conductances are rare and have only been measured for few toxins27,28 or suggested for the nuclear pore complex. Our work demonstrates that cholesterol-based membrane anchoring is a viable method for creating larger DNA pores with diameters of several nanometers. From a methodological point of view, we introduce MD-guided design of synthetic pores by demonstrating excellent agreement between simulation and experiment.

The choice of an appropriate nanopore architecture will ultimately depend on the target application. Large scaffold-based DNA nanopores like our DNA origami porin are ideally suited for functionalization and can provide space for the passage of larger biomolecules. Such DNA pores are prime candidates for single-molecule sensing or as a research tool to study transport at molecular level if the occurrence of stable insertions can be enhanced and match the longevity of protein pores. In biological environments, our large synthetic DNA porins could mimic the behavior of toxins or more complex systems like the nuclear pore complex. With this combination...
of experiments and MD simulations expanding the design space of synthetic membrane pores, we envision custom-made pores to become a versatile toolbox for cross-disciplinary applications; even if today, we may still be humble apprentices of nature.

**METHODS**

**Design of the Cholesterol-Tagged DNA Origami Porin.** The funnel-shaped DNA origami was designed on a square lattice adopted from an earlier design\(^\text{15}\) using the open source plugin in caDNAno.\(^\text{12}\) Custom DNA oligomers (for sequences see Supporting Information, Tables S1–S6) were purchased from Integrated DNA Technologies, Inc.

**Molecular Self-Assembly.** Ten nM of the M13mp18 scaffold (New England Biolabs) was mixed with 100 nM of the 179 staples and annealed for 23 h in 40 mM Tris-HCl, 45 mM boric acid, 1 mM EDTA, 14 mM MgCl\(_2\), pH 8.2 using a protocol described previously.\(^\text{27}\)

**Purification.** Structures were purified from excess staples via spin-filtration with 100 kDa MWCO filters (Amicon) in 40 mM Tris-HCl, 45 mM boric acid, 1 mM EDTA, 2 mM MgCl\(_2\), pH 8.2. The MgCl\(_2\) concentration was reduced to prevent sticking to the filter membrane and readjusted to 14 mM after filtration. This protocol yielded approximately 65 ng/μL assembled structures as determined by UV–vis spectroscopy (NanoDrop 2000, Fisher Scientific). The concentration was then adjusted to 10 nM.

**Attachment of Cholesterol Anchors.** The cholesterol-modified strands (3’ cholesterol-TEG from Integrated DNA Technologies, 5’ cholesterol-C6 from Biomers, for sequences see Supporting Information, Table S6, positions are indicated in Figure S4) were heated to 50 °C for 10 min to dissolve aggregates and incubated with the DNA origami for 10 min at room temperature in 5X excess and 40 mM Tris-HCl, 45 mM boric acid, 1 mM EDTA, 14 mM MgCl\(_2\) directly before each experiment.

**Atomic Force Microscopy.** Five μL of the DNA origami porin (10 nM) in 10 mM Tris-HCl, 1 mM EDTA, 20 mM MgCl\(_2\), pH 8.0 was deposited on a freshly cleaved mica surface (Agar Scientific) and incubated for 90 s. Subsequently, the surface was rinsed 3X with 1 mL of Milli-Q water (Merck Millipore) to remove excess sample and blow-dried with nitrogen. Imaging was carried out using a Cypher S AFM (Oxford Instruments) in amplitude modulation in air and at room temperature using AC240TS cantilevers (Olympus) with a nominal spring constant of 2 N/m. The set-point to free amplitude ratio was generally kept around 70% with a free oscillation amplitude of 20 nm.

**General MD Methods.** All MD simulations were performed using the program NAMD2,\(^\text{53}\) periodic boundary conditions, the CHARMM36 parameter set for water, ions, and nucleic acids,\(^\text{54}\) CHARMM parameters for the DphPC lipid bilayer,\(^\text{55}\) custom parametrization of ion–DNA and ion–ion interactions.\(^\text{56}\) All simulations employed a 2–2–6–6 multiple timestepping, SETTLE algorithm to keep water molecules rigid,\(^\text{57}\) RATTLE algorithm to keep all other covalent bonds involving hydrogen atoms rigid,\(^\text{38}\) a 8–10–12 Å cutoff for van der Waals and short-range electrostatic forces. Long-range electrostatic interactions were computed using the particle mesh Ewald (PME) method\(^\text{59}\) over a 1.2 Å resolution grid.\(^\text{60}\)

**Equilibration of the All-Atom Model.** To equilibrate the DNA origami porin, we first cut away a 11 nm slab of the initial all-atom model containing the lipid membrane (5 nm thick) and the adjacent 6 nm-thick cross-section of the solvated DNA origami. The resulting system was energy-minimized for 1200 steps and equilibrated for 48 ns, allowing the lipid bilayer and the solution to adopt equilibrium conformation around the structure; the DNA atoms were restrained to maintain their initial coordinates during this equilibration simulation (with the spring constant \(k_{ramp} = 1 \text{kcal/(mol Å}^2\)). Following that, the equilibrated lipid bilayer and the surrounding solvent were combined with the full-length DNA origami porin. The resulting system was equilibrated under a network of elastic restraints that maintained distances between atomic pairs at their initial values; such elastic restraints excluded hydrogen atoms, phosphate groups, atoms in the same nucleotide, and pairs separated by more than 8 Å. The system was simulated under such elastic restraints for 144 ns, the spring constants of the restraints were decreased from 0.5 to 0.1 and then to 0.01 kcal/(mol Å\(^2\)) every 4.8 ns. All equilibration simulations were performed under the NPT condition, where the number of atoms (\(N\)), pressure (\(P\)), and temperature (\(T\)) were kept constant. The pressure was set to 1 atm using the Nosé–Hoover Langevin piston method.\(^\text{43,44}\) The temperature was maintained at 295 K using a Langevin thermostat.\(^\text{45}\) The ratios of the system’s dimensions along the \(x\) and \(y\) axis were constrained, while the \(z\) axis was decoupled. Following that, the system was simulated in the absence of any restraints for 19.2 ns. During all MD simulations, the system’s coordinates were recorded every 48 ps.

**MD Simulation of Ionic Current.** All simulations of the ionic current were performed in the constant number of atom, volume, and temperature ensemble. A voltage drop, \(V\), across the system was produced by applying an external electric field \(E\) such that \(V = -EL\),...
where \( L \) was the length of the simulation system in the direction of the applied field.\(^{46}\)

**Ionic Current Calculations.** Prior to calculations of the ionic current, frames of the MD trajectory were aligned\(^{45}\) using a two-step process. First, we shifted the \( x \) and \( y \) coordinates of all atoms in the simulation system by the same amount to maintain the center of mass coordinate of the DNA origami porin constant within the plane of the lipid bilayer. Next, the \( z \)-coordinates of all atoms in the system were shifted by the same amount to maintain the \( z \)-coordinate of the lipid bilayer center of mass constant. To reduce thermal noise originating from stochastic displacements of ions in the bulk solution, the ionic current calculations were carried out within the \(-l/2 \leq z \leq l/2\) region of the system, where \( l = 30 \) Å. The instantaneous current was computed as

\[
I(t + \Delta t/2) = \frac{1}{\Delta t} \sum_i q_i (\zeta_i(t + \Delta t) - \zeta_i(t))
\]

where

\[
\zeta_i(t) = \begin{cases} \zeta_{i,f}(t), & (i = K^+, \text{Cl}^-, \text{Mg}^{2+}; j = x, y, z) \\
\zeta_{i,l}(t), & (i = K^+, \text{Cl}^-, \text{Mg}^{2+}; j = x, y, z) \\
\zeta_{i,f}(t), & (i = K^+, \text{Cl}^-, \text{Mg}^{2+}; j = x, y, z) \\
\zeta_{i,l}(t), & (i = K^+, \text{Cl}^-, \text{Mg}^{2+}; j = x, y, z)
\end{cases}
\]

the sum over \( i \) indicates a sum over all ions, \( \Delta t \) is the time interval between two consecutive frames of the MD trajectory, and \( q_i \) is the charge of ion \( i \).\(^{47}\) The average current of a trajectory was computed by summing up all instantaneous currents and dividing by the number of coordinate frames of the trajectory. To estimate the error, the ionic current trace was first block averaged with a block size of 2.88 ns. The reported standard errors of the mean were calculated from the block-averaged current traces.

**Calculations of the Local Density and Local Ionic Current Flux.** The local density and the local ionic current flux were computed as described previously.\(^{21}\) We divided the simulation system into 5 × 5 × 5 Å grids and calculated the average density of the selected atom species in each grid using a sampling frequency of 240 ps. The local current in each grid in a given direction \((x, y, z)\) was calculated by

\[
I_j = \sum_i q_i \times f_{ij}, \quad (i = K^+, \text{Cl}^-, \text{Mg}^{2+}; j = x, y, z)
\]

where \( i \) is the ion species \((K^+, \text{Cl}^-, \text{Mg}^{2+})\), \( q_i \) is the charge of the ion, and \( f_{ij} \) is the flux of the ion in the given direction. We averaged the three-dimensional (3D) density and flux data in the cylindrical coordinate overview the azimuthal angle to obtain the mean density and mean flux on the \( r - z \) plane as described previously.\(^{19}\) Following that, the 2D density and flux were made symmetric about the \( z \) axis by making a mirror image \((r \rightarrow -r)\). Finally, we used the contour and streamplot function in the python matplotlib package to generate the local density and flux plots, which were then assembled into the final figures.

**Coarse-Grained Simulation of Lipid Pore Formation.** The MARTINI simulations were performed using the Gromacs 5.0.4 package with a 20 fs time step and 12 Å cutoffs for nonbonded forces.\(^{18}\) The half-harmonic potential was implemented using the MRDUN program of the Gromacs package.\(^{25}\) First, we equilibrated a lipid bilayer membrane containing 8192 dioleoyl-phosphatidylcholine (DOPC) lipid molecules and 96,000 water beads in a 50 × 50 × 9 nm³ simulation box at zero surface tension under periodic boundary condition. We chose to simulate DOPC lipids over DPhPC because the current MARTINI force field does not provide parametrization for DPhPC. DOPC is analogous to DPhPC within the MARTINI’s coarse-graining framework that maps four hydrocarbon atoms to one MARTINI bead. Following that, multiple systems, each containing a single pore of prescribed radius, were created by applying a half-harmonic cylindrical potential, \( V_r(r) \), to all DOPC beads: \( V_r(r) = 0.5k(r - R_p)^2 \) for \( r < R_p \) and 0 otherwise, where \( R_p \) is the pore radius and \( r \) is the distance from the pore axis. Each system was simulated for 300 ns; the average pressure was computed using the last 200 ns of simulation. The error bars were estimated as the standard error of 10 ns block averages.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.6b03759.

Layout, positioning of cholesterol anchors, and DNA sequences of the DNA origami porin, agarose gel electrophoresis, analysis of AFM imaging, additional ionic current traces. Results of MD simulations including arrangement of lipid head groups at lipid–DNA interface, local concentration of ions near the DNA origami porin, histograms of the simulated ionic current, and caption describing the Movie S1 (PDF) Movie S1 (AVI)

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Notes

The authors declare no competing financial interest.

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