Regulating the Transport of DNA through Biofriendly Nanochannels in a Thin Solid Membrane

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Channels formed by membrane proteins regulate the transport of water, ions or nutrients that are essential to cells' metabolism. Recent advances in nanotechnology allow us to fabricate solid-state nanopores for transporting and analyzing biomolecules. However, uncontrollable surface properties of a fabricated nanopore cause irregular transport of biomolecules, limiting potential biomimetic applications. Here we show that a nanopore functionalized with a self-assembled monolayer (SAM) can potentially regulate the transport of a DNA molecule by changing functional groups of the SAM. We found that an enhanced interaction between DNA and a SAM-coated nanopore can slow down the translocation speed of DNA molecules and increase the DNA capture-rate. Our results demonstrate that the transport of DNA molecules inside nanopores could be modulated by coating a SAM on the pore surface. Our method to control the DNA motion inside a nanopore may find its applications in nanopore-based DNA sequencing devices.

Protein channels, embedded in a cell membrane, connect the cytoplasm of a cell with an extracellular fluid, playing an important role in transporting molecules in and out of a cell. Functional membrane channels include aquaporins that enhance the membrane permeability of water1, ion channels that selectively transport protons, potassium, sodium or calcium ions2, active transporters that bring in small nutrients such as vitamin B12, and so on. The high selectivity of each type of membrane channels, resulting from subtle and energetically favorable interactions between a transported solute and the channel's surface, ensures that other solutes are excluded. Inspired by transport mechanisms of membrane proteins, researchers have designed and built biomimetic channels or nanopores in a thin solid membrane. As a single-molecule technique, biomemetic channels can not only be used to transport ions or molecules but also be deployed to analyze transported solutes3–5. Due to its confined geometry, comparable to the size of a transported solute, a synthetic nanopore has been widely used as a sensor for structural analyses of a transported molecule, such as a DNA molecule with bound proteins6,7. However, because of the dynamic drilling process by a focused ion beam8 or a transmission electron beam9, each solid-state nanopore typically has its unique and uncontrolled surface features (e.g. charge distribution). Experimentally, it has been shown that interaction between DNA and the surface of a solid-state nanopore can affect the transport dynamics of the DNA. It is therefore essential to improve surface properties and avoid intermittent events of DNA sticking to the surface. Several methods have been developed, including assembling a fluidic lipid-bilayer on the surface of a solid-state nanopore10, coating DNA hairpins to the pore surface11, embedding a protein nanopore inside the solid-state one12, inserting a DNA origami pore inside the solid-state one13, chemically modifying the pore surface with organic molecules14–16, etc. Closely related to the last method, we suggest to coat a self-assembled monolayer (SAM) on the pore surface and modify functional groups (via chemical reaction) on the SAM surface to achieve improved transport of DNA.

By flushing chemical agents through a solid-state nanopore with a SAM containing 4-carboxyl benzyl phosphonic acids, we were able to in – situ switch the functionality of the SAM from being hydrophilic to being hydrophobic. Such change results in a reduced interaction between DNA and the coated pore. Theoretically, increasing the DNA-pore interaction can slow down the DNA translocation speed and increase the capture-rate at the same time. An optimum transport flux can be achieved for the intermediate-interaction region17. In this paper, we show that DNA interacts more strongly with the hydrophilic-SAM-coated pore than it does with the hydrophobic-SAM-coated one. Consistent with theoretical results, we found that in the hydrophilic-SAM-coated
pore the DNA’s translocation speed is slower but the capture rate is actually larger than in the hydrophobic-SAM-coated one. Our work highlights the possibility to fine-tune surface properties of a nanopore and to achieve desired transport rates of DNA molecules.

Results

Experiment of DNA translocation through SAM-coated nanopores. Figure 1a illustrates our experimental set-up. Two fluidic chambers (cis. and trans.) are separated by a 30-nm thick Si$_3$N$_4$ membrane and are connected by a nanopore drilled into the membrane. 4-carboxyl benzyl phosphonic acid molecules were used to coat the nanopore surface and the membrane surface. Figure 1b and Figure 1c respectively show TEM images of a nanopore before and after the coating. After the coating, a self-assembled monolayer (SAM) covers the surface of a solid-state nanopore and the radius of the pore is reduced. For each molecule in the SAM, the end containing a phosphonic acid group can covalently self-assemble to the pore surface. By means of organic syntheses on the other end of a coated molecule, different chemical groups can be positioned on the SAM surface, resulting in different functionalities of the SAM-coated nanopore. Here, we used trimethylsilyl diazomethane, as a methylating agent, to methylate carboxyl groups on the hydrophilic SAM surface, switching the hydrophilic SAM surface into the hydrophobic one (Figure 1d). Because of the solubility and the size of molecules in a desired SAM, we found that it was efficient to coat 4-carboxyl benzyl phosphonic acid molecules on the pore surface first and then to switch functional groups to target ones by esterification of carboxylic acid.

To study the transport of biomolecules in differently functionalized nanopores, we loaded DNA molecules (20-nM) in a 1-M KCl electrolyte in the cis. chamber and applied a biasing electric field to drive DNA molecules through functionalized nanopores (Figure 1a). With the biasing electric field, an ionic current $i_{\text{ion}}$ through the pore can be measured. During the DNA translocation, such current is reduced due to the physical blockage by a DNA molecule ($\Delta I$ shown in Figure 1a). The duration of current blockage, or the residence time of DNA, can be used to calculate the translocation speed of DNA. Thus, when DNA transits a functionalized nanopore, the complex interplay of electric, hydrodynamic and tribological interactions on DNA could be revealed from fingerprints (such as $\tau$ and $\Delta I$) of an ion-current trace.

Because of the drilling process of nanopores, each pore has its own unique physical and geometrical properties that affect the ion conductivity. To pinpoint coating effects by different SAMs, we report experimental results obtained for the same pore (about 5 nm in diameter and about 4 nm in diameter after the coating). Similar experiments were repeated for different pores and results were consistent (supporting information). We observed DNA translocation events before the pore was fully wetted. Therefore, we only analyzed current data after the open-pore current reached its maximum value. Figure 2a and Figure 2b show 4 seconds of typical ion-current traces for DNA translocations through the same pore with hydrophilic-SAM-coated and hydrophobic-SAM coated surfaces, respectively. In each case, 2-Kbp DNA molecules were electrophoretically driven through the pore at a biasing voltage of 300 mV. Averaged over translocation events (number $N$), the dwell time of DNA in the hydrophilic-SAM-coated nanopore is about 0.4 ms ($N = 1451$, Figure 2c), about 2.7 times longer than the mean dwell time of 0.15 ms for DNA in the hydrophobic-SAM-coated pore ($N = 1942$, Figure 2d). This indicates that the pore surface is lubricated by the hydrophobic SAM. Because of reduced interaction (e.g. sticking) between DNA and the pore, DNA translocation traces tended to last longer for the hydrophobic-SAM-coated nanopore. The longest recorded trace of free DNA translocations corresponded to the pore coated with the hydrophobic SAM and lasted 160 s (Supplementary Information).

From measured open-pore currents (Fig. 2a and Fig. 2b), the average open-pore conductances were estimated to be 12.2 ± 0.5 nS for the hydrophilic pore and 14.5 ± 1 nS for the hydrophobic pore, which are of the right order of magnitude (6 nS) calculated$^{18}$ for the geometry of the pore considered. In a hydrophilic-SAM-coated nanopore, the SAM surface serves as a non-slip boundary for the confined electrolyte and velocities of ions and water approach zero near the surface. Therefore, the larger open-pore current in a hydrophobic-SAM-coated nanopore is due to finite mobilities of ions near such surface (serving as a slip-boundary).

The capture rates (i.e. the number of translocation events per unit time) are 38.4 Hz and 24.3 Hz for DNA transiting a hydrophilic-SAM-coated and a hydrophobic-SAM-coated pore (Figure 2a and

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**Figure 1** | Experimental set-up. (a) A schematic of DNA translocation (driven by a biasing electric field) through a SAM-coated nanopore. (b), (c) TEM images of a solid-state nanopore (12-nm-in-diameter) before and after coating with carboxyl benzyl phosphonic acid, respectively. Red circles highlight pore sizes. (d) Configurations of molecules in the hydrophilic and hydrophobic SAMs. The carboxyl benzyl phosphonic acid was used to coat a solid-state nanopore in a 30-nm-thick Si$_3$N$_4$ membrane. The other end can be changed from hydrophilic to hydrophobic status with trimethylsilyl diazomethane.

**Figure 2** | Translocation of DNA through the same nanopore coated with different SAMs at 300 mV. (a) hydrophilic-SAM coating; (b) hydrophobic-SAM coating, switched from the hydrophilic-SAM coating after adding trimethylsilyl diazomethane. (c), (d) typical electric signals of DNA translocation events.
Gaussian-like, similar to previous experimental results for DNA and hydrophobic-SAM-coated pores, respectively. Events with long dwell times in Figure 3a (e.g. from 2 ms to 100 ms) correspond to translocation of DNA through the hydrophilic-SAM-coated nanopore. The capture rate increased when the biasing voltage changed from 300 mV to 400 mV. At an even higher biasing voltage (500 mV), the capture rate was reduced likely because of the stronger interaction (in a higher electric field) between the DNA and the SAM on the top surface of the membrane (Fig. 1a).

Figure 3a shows the scatter plots of conductance changes versus dwell times for both cases. Two well-defined clusters can be identified, corresponding to two distinctive dynamics of DNA transport in two-SAM-coated nanopores. Besides different residence times of DNA in SAM-coated pores, translocation-induced current blockades are also different. At a biasing voltage of 300 mV, average blockade currents, with standard deviations of the mean, were 1,200 ± 28 pA and 1,529 ± 37 pA for hydrophilic-SAM-coated and hydrophobic-SAM-coated pores, respectively.

The distributions of blockade currents, shown in Figure 3b are Gaussian-like, similar to previous experimental results for DNA translocation in a bare solid-state nanopore. The wider distribution of the conductance change for the hydrophobic pore may result from larger fluctuation at the water-SAM (hydrophobic) interface.

For comparison, we normalized the mean current blockages with their respective open-pore currents. The normalized values are 32.4% and 35.6% for hydrophilic-SAM-coated and hydrophobic-SAM-coated nanopores. Note that the normalized value for the hydrophobic-SAM-coated nanopore is about 10% larger than that for the hydrophilic-SAM-coated one. Theoretically, for a high-ion-concentration electrolyte (such as 1 M used in our experiment), normalized blockade currents can be simply estimated by $S_{DNA}/S_{pore}$, where $S_{DNA}$ and $S_{pore}$ are the cross-section areas of DNA and the coated pore. As shown in the supporting materials, because of the hydrophobic-SAM surface, the effective radius of the pore (accessible by water) is 1.9 nm instead of 2 nm for the hydrophilic-SAM-coated one. Thus, the resulting cross-section area for the hydrophobic-SAM-coated nanopore is about 10% less than that for the hydrophilic-SAM-coated one, which accounts for the difference between normalized values of current blockades for two SAM-coated pores. For the hydrophilic-SAM-coated pore, if the radius of DNA is 1.1 nm, the predicted normalized value is 30.3%, consistent with the experimentally obtained value.

Figure 3c and Figure 3d show the trends regarding the average dwell time and the average conductance change, as a function of biasing voltages, for hydrophilic-SAM-coated and hydrophobic-SAM-coated nanopores. The dwell time decreases as the voltage changes for both cases when the voltage increases and, correspondingly, the conductance changes are nearly constant (Figure 3d). Conductance changes measured for the hydrophilic-SAM-coated pore are about 80% of those for the hydrophobic-SAM-coated one. Figure S1 in the Supplementary Information shows 1.5 seconds of typical ion-current traces at 300, 400, and 500 mV for DNA translocations in the pores with hydrophilic and hydrophobic SAMs.

Note that we repeated the same experiment for different nanopores with the same SAM coating. We found that results are very consistent. Details are presented in the Supplementary Information (section 6: Fig. S5–S8 and Table AT1).

**Table 1 | Frequencies (Hz) f of DNA translocation events in two SAM-coated nanopores at different biasing voltages (mV)**

| Voltage (mV) | 300 mV | 400 mV | 500 mV |
|-------------|--------|--------|--------|
| hydrophobic-SAM $f_1$ | 24.3   | 39.6   | 31.2   |
| hydrophilic-SAM $f_2$ | 38.4   | 56.9   | 53.9   |
| Ratio $f_1/f_2$ | 63%    | 70%    | 58%    |

**Molecular dynamics simulation.** To interpret experimental results, we carried out all-atom MD simulations of DNA transiting a functionalized nanopore on the IBM Bluegene supercomputer. Figure 4a illustrates the simulation system that mimics the experimental set-up. The 6-nm-diameter channel was made from the amorphous SiO$_2$ solid. Octanol or octane molecules were covalently linked to Si atoms that were evenly distributed on the channel surface, which respectively results in a hydrophilic or hydrophobic SAM on the channel surface. Note that, limited by the availability of force fields, the composition of a simulated SAM is different from that in experiment. However, both the pore radii after the coating and functional chemical groups on a SAM-coated surface, are the same in simulation and in experiment. A dsDNA molecule was solvated in a 100 mM KCl electrolyte. The complex of the DNA and the electrolyte was confined inside the functionalized nanopore. Periodic boundary conditions were applied in all three directions in simulation. The length of the dsDNA molecule is the same as the channel length. End effects (such as slow entry into and fast exit away from a nanopore) of dsDNA were not present because of the covalent linking of the 5’-end and the 3’-end (in each DNA strand) across the periodic boundary. A biasing electric field,
3.1 mV/nm (about 100 mV biasing in experiment), was applied in the z direction to drive the dsDNA molecule through the channel. Details on the simulation set-up are provided in the “method” section. The translocation of DNA in each SAM-coated nanopore was simulated for 600 ns. Figure 4c shows different trajectories of DNA opposite the electric field direction in two SAM-coated nanopores. In a hydrophobic-SAM-coated nanopore, the mean translocation velocity of DNA is about 0.21 nm/ns. At short time scales (e.g. 10 ns), the velocity of DNA fluctuates and even the reversal motion (in the direction of the electric field) of DNA was observed. In the weak electric field applied in simulation, it is possible that in a short time interval the electroosmotic flow near the DNA surface exerts a larger hydrodynamic friction force than the electric driving force. When DNA is in the hydrophilic-SAM-coated nanopore, the motion of DNA adopts a stick-slip fashion. It is conceivable that DNA can have a stronger interaction with a hydrophilic SAM than with a hydrophobic SAM (see below). Generally, when DNA is contacting the SAM (e.g. around 200 ns in Figure 4c), DNA either stops or moves slower (than near the channel center). The mean velocity of DNA’s driven motion includes both the hydrodynamic friction and the contact friction between DNA and the SAM. Therefore, as shown in Figure 4c, DNA transits faster in the hydrophobic-SAM-coated pore than in the hydrophilic-SAM-coated one.

Figure 5a shows the time-dependent radial motion of DNA during the translocation (also see movies in Supplementary Information). When DNA was in a hydrophobic-SAM-coated nanopore, DNA was always near the center of the channel and was at most a few angstroms away from the center. From an energy point of view, the solvation-energy barrier prevents the hydrophilic DNA molecule from contacting a hydrophobic SAM. Thus, the electric driving force is balanced only by the hydrodynamic friction. When DNA was in a hydrophilic-SAM-coated nanopore, DNA can move away from the center and temporarily stay at the SAM surface. The force resisting DNA’s driven motion includes both the hydrodynamic friction and the contact friction between DNA and the SAM. Therefore, as shown in Figure 4c, DNA transits faster in the hydrophobic-SAM-coated pore than in the hydrophilic-SAM-coated one.

Figure 5b and 5c show density functions, $\rho(r)$, illustrating radial distributions of DNA in hydrophilic-SAM-coated and hydrophobic-SAM-coated nanopores. The probability to find DNA, at a distance of $r$ measured from the central axis of DNA to the central axis of a coated

Figure 5 | Radial motion of DNA in SAM-coated nanopores. a) Time-dependent positions of DNA, in the radial direction. (b,c) Radial distribution of DNA in a hydrophobic-SAM-coated (b) and a hydrophilic-SAM-coated (c) nanopore. Dashed lines show Gaussian distributions. (d) Illustration of water-mediated interaction between DNA and a hydrophilic SAM. Water molecules are shown as green sticks. (e) Illustration of the direct interaction between DNA and a hydrophilic SAM. In (d) and (e), DNA is at the lower left corner; part of the SiO$_2$ solid surface is at the upper right corner; the hydrophilic SAM is in the “stick” representation. Hydrogen bonds are presented as gray lines.
pore, \( r(2\pi r dr) \). When DNA is in the hydrophilic-SAM-coated nanopore, the radius-dependent density function satisfies a single-Gaussian distribution centered at the origin (Figure 5b). Thus, DNA is effectively confined near the center of a pore.

When DNA is in the hydrophilic-SAM-coated nanopore, DNA can diffuse away from the pore center and temporarily stay at the coated surface. Intuitively, two Gaussian peaks should be present in the density function, representing higher probability-densities at the center and at the pore surface. Surprisingly, between those two peaks mentioned above, there is an extra Gaussian peak, centered at a distance of 0.36 nm from the pore center. From simulation trajectories when DNA motion stalls, we found that DNA could interact with the hydrophilic SAM in two different ways. Figure 5d shows that the interaction between DNA and the hydrophilic SAM is mediated by a thin (3–4 Å) layer of water molecules. When confined between two hydrophilic surfaces close to each other, water molecules exhibit dramatically slow dynamics because of hydrogen bonding to both surfaces (i.e. surface-induced solidification of water). Such water-mediated adhesion between DNA and the SAM surface could temporarily stall DNA motion. Another type of interaction is that DNA contacts the hydrophilic SAM-surface directly. As shown in Figure 5e, phosphate groups of DNA form hydrogen bonds with hydroxyl groups on the SAM surface, interlocking two surfaces. Because the radius of DNA is about 1.2 nm and the radius of a SAM-coated nanopore is about 2 nm, DNA is about 0.7 nm away from the pore center when the contact (defined when the distance between two surfaces is about 0.1 nm) occurs. This is consistent with the Gaussian peak shown at \( r = 0.7 \) nm in Figure 5c.

The above two mechanisms of interfacial locking rely on the formation of hydrogen bonds (between two surfaces) that normally require a typical time \( t \). If DNA moves fast, \( \sigma/\tau < t \), where \( \sigma \) is a typical shearing distance of one surface relative to the other one before the interfacial locking occurs. Hydrogen bonds between DNA and the SAM-coated surface are always disrupted by the fast shearing process. Therefore DNA cannot be stalled if the interfacial adhesion of DNA is too fast. This is confirmed in an independent set of simulations when the electric field strength is 10 times larger (Figure 4b). Results, that velocities of DNA in both SAM-coated nanopores were close and DNA did not stall during the time of simulation, indicate no interfacial locking in a high biasing electric field because the electric driving force is much larger than the friction force on the SAM-surface.

Discussion

We have shown experimentally and theoretically that a functionalized nanopore in a thin solid membrane, mimicking a cell membrane, can be used to control the transport of other biomolecules, such as proteins. In a bio-mimicking fashion, a functionalized solid-state nanopore could even be selective, becoming a sodium/potassium channel or a nutrient transporter. This may require a more complicated coating mechanism, such as differential coating or patterned coating of different types of molecules inside a nanopore drilled through a sandwich-like membrane.

Methods

Synthesis of coating molecules. We synthesized the organic coating compound 4-carboxyl benzylphosphonic acids. (1) 4-Bromomethylbenzonic acid (2.15 g, 10 mM) was suspended in 10 mL of triethyl phosphate and the mixture was heated at 90°C under nitrogen for 5 hours. After cooling to room temperature, excess triethyl phosphate was removed under vacuum and the solid residue was re-crystallized from toluene to give diethyl 4-carboxybenzylphosphonate (2.21 g, 81%) as white needles. NMR (DMSO-d6): ppm 1.6 t = 6.2 Hz, 6H; 3.16 d, J = 22 Hz, 2H; 7.38 d, 6 = 8 Hz, 2H; 7.86, d, J = 8 Hz, 2H. (2) Diethyl 4-carboxybenzylphosphonate (1.5 mL) was suspended in 30 mL of 20% hydrochloric acid and the mixture was heated to reflux under nitrogen for 6 hours. After cooling to room temperature, the solvent was removed under reduced pressure and the residual solid was re-crystallized from isopropyl alcohol to give 4-carboxybenzylphosphonic acid (650 mg, 61%) as white crystals, NMR (CD3OD-d4), ppm, 3.24 d, J = 22 Hz; 7.43, d, J = 22 Hz, 2H and 7.97, d, J = 8 Hz, 2H.

The coating of a nanopore (drilled into a SiN membrane) is achieved by immersion of the membrane into a solution containing 2 mM 4-carboxybenzylphosphonic acid in ethanol for 12 hours and rinsing it with ethanol afterwards for about 1 to 2 hours. The phosphonic acid group covalently and exclusively bonds to the SiN membrane surface and the carboxylic acid group is exposed on the SAM surface. At pH 7.0, the resulting SAM surface is neutral and hydrophilic. To switch surface functionality, a substrate coated with monolayer of 4-carboxybenzylphosphonic acid was immersed in a solution of trimethylsilyl diazomethane (2 M solution in hexane, 2 mL) in 50 mL of anhydrous tetrahydrofuran. To this solution was added 1 mL of methanol and kept at room temperature for 4 hours. The substrate was removed from the solution and rinsed in methanol and dried under stream of nitrogen to give methyl ester of the monolayer.

Instruments and data analyses. The platform comprises a 30-nm-thick SiN membrane with a 5-nm-diameter nanopore. The membrane is mounted in a fluidic cell, separating the cis. and trans. reservoirs. The cis. reservoir is filled with an aqueous electrolyte/DNA solution. The 1 M KCl electrolyte (measured PH = 6.9, no buffer) was made using DI water (millipore, MA). The 2-K-base-pairs dsDNA was purchased from Fermentas (Thermo Scientific, PA, US). DNA test solutions were mixed using fresh DI water and their pH values were confirmed by measurements, right before translocation experiments. Fluidic cells were flushed with fresh solutions before every experiment. Two Ag/AgCl electrodes are immersed into two reservoirs, electronically connecting the flow cell to an ionic current measurement set-up. Thus, DNA molecules in the cis. compartment were electrically driven to the trans. compartment through the nanopore. The set-up includes a computer-controlled patch clamp amplifier (Axon Axopatch 200B, Molecular Devices), and a DAQ card (Digidata 1440A, Molecular Devices). All data were acquired at 250 KHz sampling and were analyzed using a 100 KHz low-pass filter. We extracted the mean dwell time from the histogram of dwell times by an exponential fitting.

Nanopore fabrication. The fabrication of the nanopore membrane starts from a conventional 750-µm-thick 200-mm-diameter (100) Si wafer. Firstly, low pressure chemical vapor deposition (LPCVD) was used to deposit low-stress 30-nm-thick SiN on both sides of the Si wafer. Then 250 nm SiN was deposited on the backside of the Si wafer using plasma enhanced chemical vapor deposition (PECVD). 200-nm-thick PECVD SiO2 and 200-nm-thick PECVD SiN were deposited sequentially on the front side of the Si wafer. Then a square hole was etched through the 280 nm-thick SiN (250 nm PECVD SiN plus 30 nm LPCVD SiN) on the backside of the wafer into the Si substrate by reactive ion etching (RIE). The Si wafer was then put into 80°C, 25% (w/w) TMAl solution to etch away the backside Si material. The etching was anisotropic, with theetch rate in (110) direction being much larger than the one in (111) direction. Hence, an inverted pyramid is etched into Si material before the etch stops on the 30-nm-thick PECVD SiN, forming a 100 µm X 100 µm square-shaped free-standing membrane made of 30-nm-SiN/200-nm-SiO2/200-nm-SiN (bottom from top). A 5-µm-diameter hole was made at the center of the free-standing membrane, by etching through the top two layers (200 nm-SiN/200 nm-SiO2) and creating a free-standing membrane area (made of 30-nm-thick SiN). A nanopore (typical diameters ranging from 3 nm to 10 nm) was then drilled through the free-standing membrane using a focused electron beam in a transmission electron microscope (JEOL 3000F).
Molecular dynamics simulation. All-atom molecular dynamics (MD) simulations were carried out on the IBM Bluegene supercomputer, using the software package NAMD. The CHARMM27 force-field [1] was used to model DNA and organosilane molecules. We applied the TIP3P model [2] for water molecules, silica force-field [3] for a SiO2 solid, and standard force fields [4] for ions. All simulations were carried out in the NVT (T = 300 K) ensemble. To maintain the system at constant temperature T, Langevin dynamics was applied to atoms in the SiO2 solid. These atoms were additionally restrained harmonically to occupy amorphous lattice sites. Each simulation system measures 96 × 96 × 32 Å³. With periodic boundary conditions in all three dimensions, long-range coulomb interactions were computed using particle-mesh Ewald (PME) full electrostatics. The van der Waals interaction between atoms was calculated using a smooth (10–12 Å) cutoff.

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Author contributions

All discussed and designed the project. D.W. and S.H. performed experiments of nanopore coating and DNA translocation. B.L. did MD simulations. H.P. fabricated the SiNx sandwich for DNA position control. The authors gratefully acknowledge the microelectronics Research Laboratory (MRL) and specifically Phil Waggoner and Stephen Rossnagel at the IBM T. J. Watson Research Center for assistance with fabricating membrane devices. We express our gratitude towards Kathy Reuter for pore drilling and imaging support, and Stanislav Polonsky as well as Young Kwark for their help in calibrating the electronic measurement setup. Ajay Royyuru provided valuable advice. The work presented was supported by a grant (Number: R01HG005110) from the National Human Genome Research Institute.

Additional information

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