Picomolar Fingerprinting of Nucleic Acid Nanoparticles Using Solid-State Nanopores

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Supporting Information

ABSTRACT: Nucleic acid nanoparticles (NANPs) are an emerging class of programmable structures with tunable shape and function. Their promise as tools for fundamental biophysics studies, molecular sensing, and therapeutic applications necessitates methods for their detection and characterization at the single-particle level. In this work, we study electrophoretic transport of individual ring-shaped and cube-shaped NANPs through solid-state nanopores. In the optimal nanopore size range, the particles must deform to pass through, which considerably increases their residence time within the pore. Such anomalously long residence times permit detection of picomolar amounts of NANPs when nanopore measurements are carried out at a high transmembrane bias. In the case of a NANP mixture, the type of individual particle passing through nanopores can be efficiently determined from analysis of a single electrical pulse. Molecular dynamics simulations provide insight into the mechanical barrier to transport of the NANPs and corroborate the difference in the signal amplitudes observed for the two types of particles. Our study serves as a basis for label-free analysis of soft programmable-shape nanoparticles.

KEYWORDS: nucleic acid nanoparticles, nanopore sensing, RNA and DNA nanotechnology, RNA ring, DNA cube

The advent of nanopores as a single-molecule platform has advanced studies of a wide range of species such as proteins, drugs, nucleotides, and polymers such as RNA, DNA, polypeptides, and polysaccharides.1−7 Voltage-driven passage of analytes through a nanopore results in transient occlusion of the pore, which temporarily reduces the ionic current. The depth, duration, and frequency of the current modulations contain information about size,8−10 charge,10,11 and structure12,13 of biomolecules and their interactions with the nanopore.3,14 Deciphering such ionic current signals enables biophysical characterization of single-molecule processes15,16 and contributes to development of single-molecule analytical methods16−19 such as nanopore DNA sequencing.20−23 Alongside the development of nanopore-based sensing techniques, advances in nucleic acid nanotechnology have enabled design and programming of complex molecular structures for applications in nanomedicine and nanobiotechnology.

Nucleic acid nanoparticles (NANPs) are self-assembled nanostructures composed of multiple RNA or DNA molecules.24 To date, a variety of distinct NANPs has been demonstrated differing from one another by their shape, size, internal connectivity, and physicochemical properties.54−47 Ongoing development of NANPs is motivated by their potential applications in medicine and biotechnology, including their uses as carriers for RNAi inducers,47−50 aptamers,48,51 fluorophores, and as customizable materials.52−54 Striking features of these nanoparticles are their precisely defined molecular structures, chemical stability, and tunable physicochemical properties. Moreover, due to their size uniformity and biocompatibility, NANPs are expected to outperform their metal and polymeric nanoparticle counterparts previously used for biosensing through antibody decoration, peptide coating, as well as DNA conjugation.55−58

In this work, we show that solid-state nanopores can be used to detect and characterize NANPs with high efficiency and precision. Akin to nanopore transport of deformable hydrogels,59−62 nanopore transport of hollow and flexible NANPs...
can be expected to report on their mechanical properties. Furthermore, distinguishing individual types of NANPs via a nanopore measurement can enable multiplexed detection of biomarkers using functional NANPs. Here, we characterize nanopore translocation of ring-shaped RNA and cube-shaped DNA nanoparticles and find that mechanical deformation of NANPs governs their passage through narrow pores. Next, we examine the detection limit of such nanopore measurements and show that individual particles from a binary mixture can be identified based on an ionic current pulse produced by a single nanopore passage of a nanoparticle.

RESULTS AND DISCUSSION
In this work, we explore nanopore transport of NANPs using two representative NA structures that are further referred to as DNA cubes and RNA rings. Each structure is composed entirely of either DNA or RNA strands, self-assembled according to the prescribed connectivity rules. The DNA cubes are assembled through intermolecular canonical Watson–Crick base pairing, whereas the RNA rings are formed via RNA–RNA tertiary interactions known as kissing loops. To initiate the magnesium-dependent kissing loop interactions, individual monomers of RNA rings must be...
prefolded prior to assembly. In contrast, monomers entering the composition of DNA cubes were designed to avoid any internal secondary structures. To assess the physical characteristics of these NANPs, we used atomic force microscopy (AFM) (Figure 1a,b) and gel electrophoresis (Figure 1c), both of which show the formation of monodisperse and highly regular structures. Our nanopore setup is schematically shown in Figure 1d. The nanopore forms a solitary electrolyte contact between the cis and trans side of the membrane. Applying an electric field across the pore produces a steady-state ion current as well as a localized electric field that captures and drives charged nanoparticles through the pore. Each time a nanoparticle interacts with the pore, the ion flux through the pore is reduced transiently, producing an electrical spike. 

Previously, increasing the electric field was found to facilitate nanopore capture of biomolecules from the bulk solution and increase the signal-to-noise ratio of the produced electrical spikes. A major downside, however, was the shorter duration of the nanopore translocation process, which compromised detection efficiency. In the case of NANPs, however, we find that it is possible to increase the capture rate without compromising the detection efficiency. Employing pores with diameters in the range of 9−10 nm, just smaller than the size of a NANP, forces NANPs to reside at the nanopore entrance for a prolonged time before being squeezed through the nanopore by the electric field. As a result, every particle is detected with high signal-to-noise ratio using high-bandwidth electronics.

Sample current traces for RNA rings through a 9 nm pore at different applied voltages are shown in Figure 2a. The current is normalized by its open-pore value to highlight changes in the fractional blockade. We find that the fractional current blockade increases with applied bias. Further, we observe a seemingly counterintuitive result of longer dwell times at 400 mV than at 200 mV bias, which defies the previously observed trends in translocation of DNA and proteins through nanopores. Increasing the applied bias beyond 400 mV reveals the trend expected with translocation, namely, faster dwell times with increasing voltage. We rationalize these results as follows: below a certain voltage threshold, the RNA rings only collide with the pore and do not translocate because the electromotive force that drives the NANPs is insufficient for squeezing them through. As a result, the NANPs dwell in the pore entrance for durations that span orders of magnitude. Beyond a certain threshold defined by the electric field strength, the NANPs traverse the pore. A similar mechanism was previously suggested to govern transport of double-stranded DNA through very narrow pores in silicon nitride membranes, as well as the docking translocation of DNA origami nanoplates in 5−30 nm nanopores. The deformation-controlled mechanism of NANP translocation is also borne out by the observation of increasing fractional blockades with voltage (Figure 2c), which reports on how deep the NANPs enter into the pore constriction. The transition from collisions to translocations manifests itself in Figure 2b as a peak in the dwell times at ~400 mV, although we find the exact location of the peak to depend on the pore and particle geometry. Another example of

Figure 3. Concentration limit of NANP detection. (a–h) Current traces recorded from samples containing RNA rings at different concentrations. All measurements were carried out using a 9 nm pore at 1 V applied bias; the data were low-pass filtered at 500 kHz. Shaded regions in panels a–c highlight occurrence of shallow events. (i) Capture rate versus concentration of RNA rings. Red markers: capture rate based on total number of events. Black markers: capture rate after omission of the shallow events with fractional blockade less than 15%. Experiments performed with 400 mM KCl (10 mM Tris, 2 mM MgCl2, pH 7.9).
similar fabrication protocol. Thus far, our results suggest that the molecules of similar mass are not expected to produce signals of disrupt the structural integrity of the rings, as linear RNA may be accounted for by the nanopore-shape asymmetry.

Figure 4. Discrimination of RNA rings from DNA cubes. (a) Current traces measured from a binary mixture of DNA cubes and RNA rings (1 nM ring; 1.4 nM cube). The translocation experiments were performed using a 9.5 nm diameter pore, with a 500 mV transmembrane voltage; data were low-pass filtered at 500 kHz. (b) Close view of the current trace indicates two distinct blockade levels associated with each nucleic acid nanoparticle. (c) Scatter plot of the fractional blockade versus dwell time measured at 500 mV applied bias displays two distinct populations corresponding to the DNA cubes (n = 422) and the RNA rings (n = 376). (d) Scatter plot similar to (c) obtained at an applied bias of 800 mV. In this panel, data were processed using a 1 MHz low-pass filter (n_cube = 573, n_ring = 498). Experiments performed with 400 mM KCl (10 mM Tris, 2 mM MgCl_2, pH 7.9).

dwell time dependence is shown in Supporting Information (SI) Figure S1, where the peak voltage is at ∼700 mV. Note that the nanopore shape drawn in Figure 1d is derived from the prior tomography-based studies of pores produced using a similar fabrication protocol.69,70 Thus far, our results suggest that nanopore translocation of NANPs involves processes that occur at two separate time scales: the dwell time scale associated with NANPs residing at the nanopore entrance and the passage time scale associated with actual nanopore translocation of NANPs. Whereas the dwell times of the RNA rings span several orders of magnitude (see SI Figure S1 and Figure 4c,d), the passage times are expected to be much shorter than the dwell times, given that the nanopore translocation can only happen after deformation and/or buckling of the stalled nanoparticles. Thus, although we would expect to see deeper blockades when NANPs pass through the pores, we do not detect them presumably due to the passage time scales shorter than those detectable by our electronics (>1 μs detection limits).

In order to verify successful translocation of NANPs through the nanopore, we performed nanopore recapture experiments (see Figure 2d).72 In these experiments, a transition in the pore current level from the NANP-occupied state back to the open state triggers the hardware to reverse the bias polarity after a set delay, which we have chosen to be 200 μs. Such rapid switching of bias polarity results in recapture of the same NANP that just passed through the nanopore. Recapture of the rings, which occurs with a probability of ∼10%, validates our interpretation of long-lasting blockades as NANP capture terminated by nanopore translocation. Furthermore, ionic current blockades produced by both translocation and recapture are very similar (more example events are shown in SI Figure S2), with slight variations in the exact signal amplitude and durations (<20%) that may be accounted for by the nanopore-shape asymmetry. This finding suggests that nanopore translocation does not disrupt the structural integrity of the rings, as linear RNA molecules of similar mass are not expected to produce signals of similar amplitudes and durations given that the translocation kinetics of a NANP is conditioned by both the size of the ring and its mechanical compliance.

In a typical nanopore experiment, the voltage applied across the pore serves to facilitate the capture of analytes as well as to translocate them through the pore. Therefore, capture and translocation are coupled, which makes detection of trace amounts of analyte challenging. Indeed, although detection limits for a particular analyte can be increased by increasing the applied bias, any increase in applied bias would also speed up the translocation process, which usually compromises detection. However, in contrast to free translocation of particles through nanopores that are larger than the analytes, in the regime of deformation-controlled transport, capture and translocation can be decoupled. In Figure 3a–h, we show example 60 s current traces for RNA rings at 1 V applied bias for concentrations ranging from 10 pM to 1.25 nM. The experiment was carried out by serial dilution (2×) of a 5 nM solution of RNA rings, followed by recording of ∼150 s of data at each concentration for mean capture rate calculation. Figure 3i displays a log–log plot of the mean capture rate of the RNA rings as a function of their concentration. Red markers were obtained by calculating the capture rate based on the total number of recorded events, whereas the black markers represent capture rates for only events with amplitudes that are larger than 15% of the open-pore current (mean blockades of rings ∼30%). With the shallow events omitted, a power law fit to the log–log data yields a power of 1.12, which closely matches the expected linearity (1.0 power), the discrepancy either stemming from volume estimation error or from pipet miscalibration. However, it is noteworthy that omitting the random (and rare) shallow events that result either from pore surface contamination or from buffer contamination allows us to push detection limits to the pM levels. Finally, we have calculated the capture radius of the rings from the observed capture rates, yielding values of ∼100 nm; the diffusion coefficient of an RNA ring was estimated to be 40 μm²/s using
We note that for a previous study of double-stranded DNA capture into small (4 nm diameter) nanopores,72 the capture radius was found to be commensurate with the DNA radius of gyration. In contrast, the higher voltages used in our experiment, coupled to high efficiency detection that results from deformation-controlled transport, yields a capture radius for the deformable structure that is an order of magnitude larger than its radius of gyration.

After having characterized the nanopore transport of RNA rings, we have carried out similar measurement using DNA cubes (see Figure 1 and SI Figure S3). Overall, DNA cubes exhibit very similar transport behavior, including very broad dwell time distributions, increased fractional blockade with increasing voltage, and nonmonotonous behavior of dwell times with increasing voltage values. A simple theoretical estimate of the fractional blockade produced by a DNA cube is in good agreement with the results of our measurements (see SI, section III). Owing to their 3D structure, DNA cubes can block a larger volume of the pore in its high electric field zone, hence causing larger current blockades. This permits the use of the same pore to discriminate DNA cubes from RNA rings. Figure 4a directly shows the possibility of such shape differentiation. Clearly, two distinct blockade levels are present in the trace, as further exemplified in the close-up of Figure 4b. Experiments conducted with individual NANPs as well as their binary mixtures (1 nM ring; 1.4 nM cube) confirmed that the higher-level blockade belongs to the cubes (see Figure S4). The scatter plot of the fractional blockade versus dwell time clearly demonstrates two mutually exclusive populations, indicating that cubes and rings can be differentiated based on single pulses with ≥99% efficiency (Figure 4c). In other words, each event in the scatter plot can be mapped to either a ring or a cube, based on its fractional blockade amplitude. Given that both particles deform as they enter the pore constriction, increasing the applied bias to 800 mV results in increased fractional current blockades for both NANPs while also reducing their corresponding mean dwell times (Figure 4d). These results were reproduced using 15 different nanopores in the 9.5 ± 1 nm range at different applied voltages despite small variabilities in the fractional current blockades and translocation threshold voltage that are related to the exact nanopore geometry. It is important to note that in the gel electrophoresis experiment performed using a ring–cube mixture (Figure 1c), the two types of NANPs, despite being different in shape and size, migrate very similarly and hence are difficult to distinguish from one another. In contrast, the deformation-controlled transport through nanopores enables accurate differentiation of the two NANP species. An even more accurate quantification of a composition of a NANP mixture, in addition to measuring the capture rate, would require accounting for the relative capture probability for each nanoparticle, which in practice would reduce to a calibration measurement for each particle type.

To elucidate the microscopic mechanism of NANP permeation through solid-state nanopores and to obtain an independent assessment of the ionic current blockade levels, we constructed several all-atom models of a nanopore system containing either a DNA cube or an RNA ring nanoparticle.
(Figure 5a). To increase the computational efficiency of molecular dynamics (MD) simulations, the nanopore shape was chosen to reproduce the cis half of the experimental nanopore system, with the trans side cut off at the membrane midplane. Prior to simulations of the nanopore transport, the nanoparticles were equilibrated for at least 60 ns in bulk 400 mM KCl solution until their root-mean-square deviation (rmsd) from the idealized geometry had reached steady values of 1.2 and 0.6 nm for the DNA cube and the RNA ring, respectively (SI Figure S8). Such rather large equilibrium rmsd values reflect considerable structural fluctuations that nevertheless do not affect integrity of the particles. Indeed, more than 90% of all base pairs remained intact during the free equilibration simulation (SI Figure S8).

To simulate nanopore transport, equilibrated NANPs were placed at the opening of the nanopore (Figure 5a) and, following short equilibration, were simulated under a transmembrane bias of either 200 or 500 mV. Several independent simulations were performed at each bias; Table S1 provides a summary of all MD runs. In some simulations performed at a 500 mV bias, the particles were observed to pass through the nanopore constrict, rapidly exiting on the trans side; one such trajectory is featured in Figure 5b. Interestingly, the DNA cube maintained its integrity after the translocation (SI Figure S9). In the majority of the simulations, however, the particles became wedged at the cis entrance of the nanopore. Two such stable conformations are shown in Figure 5c,d; SI animations 1–5 represent simulation trajectories. Once the structures had reached an equilibrium conformation and their movement effectively ceased, see SI Figure S7 and Table S1, we calculated the ionic current for each of the structures directly from the motion of the ions through the nanopore. For comparison, we also simulated the ionic current through the nanopore containing no nanoparticles. Figure 5e plots the current blockade percentage 100 \((I_0 - I)/I_0\) where \(I\) is the ionic current passing through the nanopore and \(I_0\) is the open-pore current. The DNA cube blocked a higher percentage of the current than the RNA ring in our simulations, at both biases, containing no nanoparticles. Figure 5e plots the current blockade percentage 100 \((I_0 - I)/I_0\) where \(I\) is the ionic current passing through the nanopore and \(I_0\) is the open-pore current. The DNA cube blocked a higher percentage of the current than the RNA ring in our simulations, at both biases, containing no nanoparticles.

CONCLUSIONS

NANPs have shown promise as enablers of advanced biophysical studies and as probes for molecular sensing approaches. Here, we have used measurements of ionic current to characterize the process of NANPs’ transport through solid-state nanopores, identifying conditions where deformability of the nanoparticles determines their transport characteristics. Specifically, we find that elastic deformation of NANPs in response to the applied electric field permits them to pass through nanopores smaller than the particle size but only after the force on the particles in the nanopore exceeds a threshold value. Such deformation results in longer dwell times, which considerably improves NANP detection limits even at a high applied bias. Furthermore, nanoparticles of different types (i.e., RNA rings and DNA cubes) produce distinct populations of ionic current blockade, and the type of individual particle transiting a nanopore from a particle mixture can be determined from a single current blockade. Such a tremendous distinguishability, combined with picomolar sensitivity, poises nanopore-based sensing of NANPs as an attractive candidate for development of a multiplexed detection platform for general biomarker sensing. Finally, our work introduces nanopores as sensitive tools for investigation of the physical properties of multistrand nanoassemblies, which are challenging to characterize otherwise, especially when several nanoparticle types of comparable sizes, charges, and molecular weights are simultaneously present in the mixture, even at pM concentrations.

METHODS

Nanoparticle Synthesis and Characterization. All NANPs were assembled by combining individual monomer components at equimolar concentrations. The oligonucleotides encoding the composition of NANPs are listed in the Supporting Information. DNA oligonucleotides were purchased from IDT (idtdna.com), and RNA strands were produced by in vitro runoff transcription using PCR-amplified DNA templates. For that, synthetic DNAs coding for the sequence of the designed RNA were amplified by PCR using primers containing the promoters for T7 RNA polymerase. PCR-amplified and purified (DNA Clean&Concentrator-5, Zymo Research) DNA templates (0.2 μM) were transcribed with “home-made” T7 RNA polymerase in 80 mM HEPES-KOH, pH 7.5; 2.5 mM spermidine; 50 mM DTT; 25 mM MgCl₂; 5 mM NTPs. Transcription was stopped with RQ1 DNase. Transcribed RNAs were purified with 8 M urea denaturing gel electrophoresis (PAGE, 15% acrylamide). The RNAs were eluted from gel slices overnight at 4 °C into 1× TBE buffer containing 300 mM NaCl and then precipitated in 2.5 volumes of 100% ethanol. Samples were then rinsed with 90% ethanol, vacuum-dried, and dissolved in double-deionized water.

For assembly of DNA cubes, corresponding oligonucleotides (purchased from IDT) were mixed in doubly deionized water, heated to 95 °C for 2 min, snap-cooled to 25 °C, and incubated for 20 min. For assembly of RNA rings, mixtures of RNAs (individually transcribed and purified) were heated to 95 °C for 2 min, followed by snap-cooling on ice for 2 min, and incubation at 30 °C for 30 min. An assembly buffer (1× concentration: 89 mM tris-borate (pH 8.3), 2 mM MgCl₂, 50 mM KCl) was added following the heating step to all assemblies.

All NANPs were characterized by the electrophoretic mobility shift assays carried out on 8% nondenaturing native PAGE (37.5:1, 2 mM MgCl₂) and visualized with a Bio-Rad ChemiDoc MP system using total staining with ethidium bromide (EtBr). All gels were run for 30 min at 4 °C, 300 V. NANPs were further visualized by AFM. For that, 5 μL of 50 nM NANPs was deposited on APS-modified mica, incubated for ~2 min, and air-dried, as described previously. AFM visualization was carried out in tapping mode on a Multimode AFM Nanoscope IV system (Bruker Instruments, Santa Barbara, CA). The images were recorded with a 1.5 Hz scanning rate using a TESPA-300 probe from Bruker with a resonance frequency of 320 kHz and spring constant of about 40 N/m. Images were processed by the FemtoScan software package (Advanced Technologies Center, Moscow, Russia).73,74

Nanopore Fabrication and Measurement. The nanopore devices are 5 × 5 mm² chips with a 50 nm thick freestanding silicon nitride membranes at the center. Nanopores were drilled using a JEOL 2010F transmission electron microscope operating at 200 kV. After fabrication, nanopore chips were cleaned using hot Piranha (sulfuric acid/hydrogen peroxide, 2:1), rinsed first by hot DI water and then by a stream of DI water, and finally dried under a gentle stream of nitrogen. Next, the chips were mounted in a custom PTFE cell, and a stream of DI water, and nitrogen. Next, the chips were mounted in a custom PTFE cell, and a stream of DI water, and nitrogen. Next, the chips were mounted in a custom PTFE cell, and a stream of DI water, and nitrogen. Next, the chips were mounted in a custom PTFE cell, and a stream of DI water, and nitrogen. Next, the chips were mounted in a custom PTFE cell, and a stream of DI water, and nitrogen. After drying, the exposed areas of the chips as close as possible to the silicon substrate and the silicon nitride layer were incubated for 20 min in 2% SDS/0.5 M NaOH solution at 45 °C, followed by snap-cooling on ice for 2 min, and incubation at 30 °C for 30 min. An assembly buffer (1× concentration: 89 mM tris-borate (pH 8.3), 2 mM MgCl₂, 50 mM KCl) was added following the heating step to all assemblies.

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nanopore. Experiments were performed with 400 mM KCl (10 mM Tris, 2 mM MgCl₂, pH 7.9). This low concentration of Mg²⁺ ions was added to the buffer to help with preserving the nanoparticles. The silver/silver chloride electrodes interface the fluids with the electronics. The ionic current was recorded using a Chimera VC100 amplifier (Chimera Instruments LLC), digitized at 4.17 M sample/s and digitally low-pass filtered.

Before each experiment, the conductance of the nanopores was measured and compared against the theoretical values to confirm the dimensions of the nanopores. Next, the nanoparticles were added to the cis chamber (grounded), and a positive bias was applied to the trans chamber. Upon applying the bias, the charged nanoparticles were electrophoretically driven through the nanopore and the translocation events were observed in the form of spikes in the DC current. The height and width of these events, that is, current blockade by a nanoparticle and the dwell time of a nanoparticle in the nanopore contain information about the structure of the nanoparticles and their interactions with the nanopore. Additionally, the interevent time distribution (the time interval between two successive events) was used to determine the capture rate, which was then correlated with the concentration of nanoparticles. PyhIon, a nanopore data analysis package, was used to extract such data from the recordings. Further analysis and distribution fittings were performed with MATLAB R2014.

**Molecular Dynamics Simulations.** All MD simulations of the NANPs/nanopore systems were carried out using NAMD², periodic boundary conditions, the CHARMM36 force field, and a custom force field for silica, and the CUFIX corrections for ions. Multiple timesteps was used: local interactions were computed every 2 fs, whereas long-range interactions were computed every 6 fs. All short-range nonbonded interactions were cut off starting at 0.8 nm and completely cut off by 1.0 nm. Long-range electrostatic interactions were evaluated using the particle-mesh Ewald method computed over a 0.1 nm spaced grid. SETTLE and RATTLE algorithms were applied to water and nucleic acid hydrogen atoms, respectively. A Langevin thermostat of 1.0 ps⁻¹ damping constant was coupled to silica atoms to maintain constant temperature. Aoms of silica membrane were also harmonically restrained to their initial coordinates; the spring constant of the restraints was 200 kcal mol⁻¹ Å⁻². Constant pressure simulations employed a Nosé–Hoover Langevin piston.

The atomic-scale model of a solid-state nanopore was obtained by annealing high-temperature (7000 K) silica melt in the presence of a grid-based potential that defined the shape of the nanopore. During the annealing simulation, the temperature of the system was set to 7000 K for 500, 2000, and 300 K for 40,000, 400,000, 100,000, and 100,000 steps, respectively. These simulations were performed in vacuum using the BKS force field. The nanopore shape was chosen to match the nanopore geometry realized in the experiment, the nanopore had an hourglass shape with the middle section approximated by a 9 nm diameter cylinder and the nanopore entrances approximated by 30° angle cones. To reduce the cost of MD simulations, the nanopore shape was cut in half along the midplane of the membrane, resulting in an asymmetric system shown in Figure 4A. After being annealed, atoms located farther than 1 nm away from the nanopore surface were removed, further reducing the computational cost of the simulations. Initial models of NA particles were built by arranging DNA and RNA strands according to the particle’s idealized geometry. The cube and the ring nanoparticles were submerged in a cubic volume of 400 mM KCl electrolyte 14 and 18 nm on each side, respectively. The systems were equilibrated in the constant number of particles, pressure, and temperature (NPT) ensemble for over 60 ns each (P = 1 atm; T = 295 K). The equilibrated systems were then merged with the all-atom model of the nanopore, placing each NANP at the entrance of the nanopore with its center of mass initially located at the nanopore axis. The ring particle was initially oriented normal to the silica membrane. Three versions of the cube/nanopore system were built differing by the orientation of the cube with respect to the nanopore (SI Figure S6). Water and ions that clashed with the membrane were removed; additional volumes of 400 mM KCl solution were added to fully wet the nanopore and to form water-filled compartments on either side of the membrane. The final systems measured 16 × 16 × 23 nm³ and contained approximately 470,000 atoms. The systems were equilibrated in the NPT ensemble for approximately 5 ns. Following the equilibration, the simulations were run in the constant number of particles, volume, and temperature (NVT) ensemble. An external electric field, E = −V/Lₑ, was applied along the nanopore axis to produce the target drop of the electric potential, Vₑ over the system’s dimension in the direction of the applied field, Lₑ. In all simulations, a short-range repulsive potential was applied to atoms of NANPs to prevent their permanent binding to the nanopore surface.

**ASSOCIATED CONTENT**

Supporting Information

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Compositions of the nanoparticles; translocation of the DNA cubes and RNA rings through a nanopore at different voltages; recapture of RNA rings at different voltages; estimation of the fractional current blockade; initial orientation of the NANPs in MD simulations; simulated displacement of NANPs through a nanopore; structural fluctuations of NANPs in bulk solution; structural integrity of NANPs during simulated nanopore translocation; animations illustrating MD trajectories of a DNA cube translocation through a solid-state nanopore with different initial orientations under a 200 mV bias; an MD trajectory of a DNA cube translocation under a 500 mV bias, an MD trajectory of an RNA ring translocation under a 200 mV bias translocation (PDF)

Animation M1 (MPG)

Animation M2 (MPG)

Animation M3 (MPG)

Animation M4 (MPG)

Animation M5 (MPG)

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**Notes**  
The authors declare no competing financial interest.

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Assembly of Sequence-Defined Polymers on DNA Cages.
W.; Joshi, A. A.; Rusling, J. F. Resistive-Pulse Measurements with Nanopipettes: Detection of Vascular Endothelial Growth Factor C


