Modeling and Analysis of Intercalant Effects on Circular DNA Conformation

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Supporting Information

ABSTRACT: The large-scale conformation of DNA molecules plays a critical role in many basic elements of cellular functionality and viability. By targeting the structural properties of DNA, many cancer drugs, such as anthracyclines, effectively inhibit tumor growth but can also produce dangerous side effects. To enhance the development of innovative medications, rapid screening of structural changes to DNA can provide important insight into their mechanism of interaction. In this study, we report changes to circular DNA conformation from intercalation with ethidium bromide using all-atom molecular dynamics simulations and characterized experimentally by translocation through a silicon nitride solid-state nanopore. Our measurements reveal three distinct current blockade levels and a 6-fold increase in translocation times for ethidium bromide-treated circular DNA as compared to untreated circular DNA. We attribute these increases to changes in the supercoiled configuration hypothesized to be branched or looped structures formed in the circular DNA molecule. Further evidence of the conformational changes is demonstrated by qualitative atomic force microscopy analysis. These results expand the current methodology for predicting and characterizing DNA tertiary structure and advance nanopore technology as a platform for deciphering structural changes of other important biomolecules.

KEYWORDS: DNA conformation, nanopore, DNA translocation, ethidium bromide, intercalation, molecular dynamics

Over the course of its life cycle, a cell’s DNA undergoes many carefully orchestrated conformational changes that facilitate vital cellular processes such as replication and transcription. Consequently, unresolved conformational defects in the structure can interfere with critical interactions, resulting in genetic anomalies that culminate in cell death. As a result, numerous cancer treatments target DNA, seeking to induce structural changes to inhibit tumor growth. Medications such as the widely used anthracycline class of antitumor drugs interfere with cellular functionality by interrupting protein–DNA interactions. The specific molecular mechanisms are not completely understood, but the increased torsional stress on the DNA from the intercalation is thought to be a significant contributor. Although these important chemotherapy drugs have proven to be effective treatments for many forms of human cancers, their associated cardiotoxicity has pushed for the advancement of different analogues and formulations. The development of innovative cancer drugs can be greatly enhanced by high-throughput, low-cost methods to characterize their effects on DNA integrity and conformation. However, conventional investigations of conformational changes of DNA frequently require extensive preparation, labeling, or substrate adhesion, which may induce unexpected artificial conformational changes. Here, we show that the structural changes induced by the intercalation of ethidium bromide (EtBr) in circular DNA can be identified using a solid-state nanopore. Our experiments are supported by all-atom molecular dynamics simulations that provide fundamental insight into the
unwinding of the supercoiled DNA molecule, resulting in the formation of branched (looped) structures and distinctive signatures in the translocation events.

Although routinely considered as a next-generation platform to achieve rapid and inexpensive DNA sequencing, the most promising emerging application of solid-state nanopores is to provide a label-free molecular probe to investigate individual biomolecular structures in solution. This approach has been previously used to characterize linear DNA structures, protein binding to linear DNA, and other modifications to biomolecules. However, the evaluation of circular DNA has been limited to comparisons with linear conformations. Such comparisons are not adequate for predicting structural changes in DNA conformation in vivo, which requires a DNA molecule with a constrained configuration in the absence of cellular infrastructure. Circular covalently closed DNA has been shown to be a good model for representing the structure of DNA in vivo because it contains the constraints of closed domains found in DNA that give rise to its large-scale conformational properties. Being constrained, intercalation into circular DNA causes localized unwinding of the molecule, which subsequently causes a structural transformation according to the Calugareanu–Fuller–White model. EtBr is a very popular prototypical intercalating molecule that has been used for several decades to fluorescently label DNA and alter its mobility during electrophoresis. EtBr’s interaction with DNA has been well-characterized experimentally and serves as a good model for anthracyclines, such as daunorubicin and doxorubicin. Previous investigations have shown that, upon intercalation, EtBr reduces the negative charge of DNA while increasing the effective molecular weight. The intercalation also initiates a negative unwinding of the helix by reducing the helix angle by 26° per intercalated base pair, consequently lengthening the DNA molecule. However, while the intercalation of EtBr has been extensively studied, modeling the changes to conformation has been largely overlooked and no direct measurements of the resultant structural changes in a fluid environment have been reported.

RESULTS AND DISCUSSION

To further our understanding of the changes to DNA conformation that occur due to the intercalation of ethidium between base pairs, all-atom molecular dynamics simulations were conducted to predict and identify unwinding of the DNA in the presence of EtBr. We began with a negatively supercoiled DNA ring of 180 base pairs and linking number (Lk) of 14 in 1 M KCl, as depicted in the schematic representation of the simulation system in Figure 1a. Three separate simulations were carried out on the DNA ring. We first simulated the DNA ring for 200 ns and calculated its writhe based on the change in the position of the center of mass of the base pairs. In the other two simulations, 6 and 12 ethidium ions were intercalated between the base pairs at approximately equidistant positions (Table S1, Supporting Information), and each system was simulated and allowed to evolve for 200 ns. A sample configuration of an ethidium ion located between two base pairs is shown in Figure 1b, and the time evolution of the writhe for the three simulations is shown in Figure 1c. In the first run, that is, the free DNA ring, the writhe becomes stable after about 20 ns of simulation and fluctuates around −0.65 ±
0.03 for the remainder of the 100 ns run. In the other two systems containing 6 and 12 ethidium ions, the calculated writhe increases during the first 50 ns of the simulation, reaching an average writhe of $-0.45 \pm 0.09$ and $-0.16 \pm 0.04$ over the final 100 ns of the simulation, respectively. The initial and final snapshots of these two systems are shown in Figure 1d,e. The figures show that the intercalated ethidium ions unwind the DNA ring, and the extent of unwinding correlates with the number of ethidiums in the system. Comparing the average writhe of the DNA ring with that of the DNA–ethidium complexes, it is possible to estimate the amount of unwinding per intercalated ethidium ion. The second simulation containing 6 ethidium ions yielded an unwinding of $12.0^\circ$ per intercalated ethidium, while the 12 ethidium simulation produced a $14.7^\circ$ unwinding per ethidium, smaller than the reported $26^\circ$ per ethidium.36,37 Experimentally, DNA intercalation has been demonstrated to require several seconds to reach a maximum unwinding,39 making it reasonable that a 200 ns long simulation is not sufficient to capture the complete conformational changes that correspond to a $26^\circ$ increase in writhe per ethidium. The degree of unwinding has also been reported to correlate with the extent of intercalation,39 so simulations of 6 and 12 ethidium ions per 180 base pairs may not be sufficient to achieve an average unwinding of $26^\circ$ per intercalated base pair.

Although the calculated unwinding is not as large as that in previous experimental reports, the simulations still reveal significant changes in the writhe over the 200 ns evolution period. Because of the intercalant-induced unwinding of the constrained molecule, the changes in writhe are expected to yield changes in the twist that we anticipate to manifest as branching observable with nanopore translocation. The predicted structural changes were assessed by supercoiled DNA translocation experiments through a solid-state nanopore sensor, which offers fast label-free detection and characterization of biomolecules in solution, as depicted in Figure 2a. The translocation of a molecule through a nanopore (Figure 2b) produces a characteristic drop in ionic current that is used to analyze the structural configuration. To characterize the effects caused by the intercalation of ethidium ions (Figure 2c), we first predicted changes to the translocation of DNA with further molecular dynamics simulations. The translocation of a 50 bp linear DNA molecule through a 3.8 nm diameter, 10.5 nm thick nanopore shown in Figure S1 (Supporting Information) reveals that the intercalation of ethidium ions does not have a significant impact on the current drop for the nanopore used in the simulation. However, with the reported increases in DNA mass and length as well as the decreased charge,27,30,33–35,40 an increase in the translocation time would be expected. From the simulations, it is anticipated that changes in the current drop will arise only from branching and not occur due to an increase in the intercalated DNA diameter.

Briefly, the nanopore experiments utilize a single $\sim 20$ nm pore drilled in a 20 nm thick Si$_3$N$_4$ membrane (Figure 2b) sandwiched between two flow cells containing electrolyte, thus forming the only electrical path between the reservoirs. An
applied voltage across the nanopore generates an ionic current that is interrupted by the translocation of DNA through the pore. The change in current, $\Delta I$, and the translocation time, $t_d$ (Figure 2d), of the translocation events are characteristic for the biomolecules. A detailed discussion of the experimental parameters is provided in the Methods section.

The density plot and the accompanying histograms of the translocation data from circular DNA (Figure 2e) revealed two distinct regions of current drop, which suggests two populations of different DNA molecular conformations. The distribution of the maximum $\Delta I$ is shown in the inset histogram in Figure 2e. The majority of the translocations occur in Region I, corresponding to longer translocation time and smaller current drop. The small temporal distribution of the events indicates a relatively uniform population of DNA, and a current drop of $\sim 200$ pA suggests the simultaneous translocation of two double strands ($N = 2$) consistent with DNA in the circular covalently closed conformation. We further investigated this by atomic force microscopy (AFM) and found a generally uniform circular conformation, as demonstrated by the representative AFM height image in Figure 2f and Figure S3a (Supporting Information). The inclusion of circular nicked DNA in the population would also exhibit a similar uniform structural conformation.41,42 However, its presence would be clearly observable in the translocation plots.25,26 The second region of events occurred with a shorter translocation time, but with twice the current drop ($\sim 400$ pA). This suggests that four double strands ($N = 4$) are occupying the nanopore, which consequently decreases the effective length of the DNA and leads to a decreased translocation time. Linear DNA translocation is known to exhibit a similar behavior of decreased translocation time for a doubling of the current drop, which is attributed to a folded DNA conformation.25,26,43,44 Although we cannot entirely exclude the possibility of folded conformations in Region II, folded events seem unlikely with supercoiled DNA due to the stiffness of the molecule. Previous work with nicked circular DNA, which is a more relaxed conformation than supercoiled, did not reveal the occurrence of folded events.25,26

To identify the configuration of the DNA during translocation, we analyzed the translocation events that comprise the two regions on the density plot. Region I in Figure 2e was thought to be a circular closed configuration. The translocation events shown in Figure 3a confirm a uniform conformation, indicating a structure represented by the illustration in Figure 3b. The events comprising Region II indicate different structural features that give rise to the multilevel current drops shown in Figure 3c. These typical events feature two discrete current drops, each with a magnitude of 200 pA. The second current drop occurring in the middle of the translocation event suggests the presence of a branched (looped) structure similar to the conformation illustrated in Figure 3d and demonstrated in the representative AFM image in Figure 2f. These data also support the hypothesis that the secondary current drops are not caused by folded events. We hypothesized that the addition of an intercalant to unwind the circular DNA would initiate further branching of the circular DNA due to the unwinding predicted by the simulations and previous experimental studies.36–38

Under identical translocation conditions, we added EtBr to the DNA to initiate structural changes. As shown by the density plot in Figure 4a, the translocation time of the events increased by at least 85%, and there were three regions of current drops as shown by the inset histogram. Comparing the density plot in Figure 2e with Figure 4a, we observed that the majority of events have shifted from having a current drop of 200 pA to events with a 400 pA current drop. To determine if the changes in translocation could be due to changes in conformation, we used AFM to identify relative changes in the DNA structure after intercalation. The AFM height image in Figure 4b and Figure S3b (Supporting Information) shows a significant increase in branched structures relative to those seen in Figure 2f, which we attribute to the interaction with EtBr.

Again, we analyzed the translocation events to determine the structural conformation of the DNA. For uniform supercoiled DNA intercalated with ethidium ions, the simulations suggest the translocation events will remain at 200 pA, but with a longer translocation time. The events in Region 1, shown in Figure 5a, were uniform with a current drop of 200 pA, which suggests that the DNA remains in the circular closed configuration. However, the translocation time increased by $\sim 265\%$, which we attribute to the increased length, increased mass, and decreased charge arising from ethidium ion intercalation. An illustration of the increased length is shown in Figure 5b and is consistent with the results of the simulation.

As indicated by Figure 4a, upon addition of EtBr, the majority of events shifted to Region 2, indicating that intercalation caused branching in the DNA conformation. Although the current blockage remained at 400 pA, the translocation time of branched DNA increased by $\sim 286\%$, which is consistent with changes to molecular length, mass, and net charge. The branching structures can be identified by the discrete current drops in Figure 5c, suggesting a conformation like that illustrated in Figure 5d.

The translocation events comprising Region 3 have multiple discrete current drops, shown in Figure 5e, with a maximum current drop of 600 pA. Again, the current drops occur in multiples of 200 pA, consistent with the translocation of multiply branched structures, as illustrated in Figure 5f. Further analysis of the translocation events reveals that approximately 5% of the total events correspond to a current drop of over 700 pA, shown in Figure 5g, that were not represented in the histogram. Additional primary branches or secondary branches (branched branches), as illustrated in Figure 5h, would be expected to produce an 800 pA current, which is observed for about one-half of the events. We attribute the discrepancy in current drop to signal filtering since the translocation of these individual features is so fast.
The analysis of the event morphology suggests branched structures translocating the nanopore, implying that multiple pairs of helices occupy the nanopore simultaneously. Although the hydrodynamic radius of this particular plasmid used has not been reported, previous work using smaller nanopores in similar high salt concentrations (1 M KCl) demonstrated the simultaneous translocation of multiple helices. Circular DNA translocation using 10 nm diameter pores report branched conformation events \( (N=4) \), and up to five helices of linear DNA have been shown to concurrently translocate using a \(~10\) nm pore. The nanopores used in our investigation had approximately 4 times the cross sectional area and up to 8 times greater volume than the nanopores used in the previous reports, so it is reasonable that multiple pairs of helices \( (N=4, N=6) \) can translocate the nanopore simultaneously. Furthermore, comparing the translocation of the closed circular conformation with and without intercalated EtBr, we found that translocation of intercalated DNA does not change the current drop (within the experimental error). EtBr is known to also bind external to the DNA helix, and high concentrations have been reported to increase the hydrodynamic radius, however, our results suggest that the concentration of EtBr used was low enough not to greatly change the hydrodynamic radius of the circular DNA.

### CONCLUSIONS

In summary, we predict and evaluate changes to DNA conformation due to the intercalation of EtBr into circular DNA. All-atom molecular dynamics simulations predict that the intercalation will induce unwinding of the helix. Although not observed on the time scales used in simulation (200 ns), we hypothesize that the unwinding gives rise to branches in the DNA molecule that we experimentally characterized using solid-state nanopore translocation and further identified with AFM. This work represents a significant advancement in simulating changes to circular DNA conformation and is supported by experimental results. Our findings provide fundamental insight into characterizing DNA structural conformations, which we envision to have important implications for anticancer drug treatment and design.
METHODS

Molecular Dynamics Simulations. Molecular dynamics simulations were carried out using the program NAMD.37 The AMBER force field parmbsc037 was used to describe interaction of the atoms. A circular covalently closed DNA molecule composed of 180 base pairs with four separate regions of sequences has been used. The sequence of the simulated DNA ring is d(GpC)45d(GpG)45d(GpC)45d(GpG)45d, which consists of four regions to make it easier to get a “figure 8” shape for the ring. We avoided A–T base pairs in the sequence to reduce the probability of bubble formation and local denaturation. The initial configuration of the DNA ring is generated by the 3DNA software,39 assuming a perfect circular geometry for the ring with base pair step parameters corresponding to a linking number of 14 as an input. Such a DNA ring is an underwound conformation compared to the equilibrium values of the intrinsic twist of the base pairs.35–37 To equilibrate this circular DNA, we applied generalized Born implicit solvent implemented in NAMD.38 We start with 10 000 steps of minimization and then carry out 20 ns simulations at a constant temperature of 298 K and an ion concentration of 1 M. After this stage, the DNA configuration changes to a “figure 8” shape and can be used as input for an explicit all-atom MD simulation. We used the final configuration and solvated it with TIPS water molecules in a periodic box of 120 × 150 × 290 Å3 and added K+ and Cl− ions to achieve a salt concentration of 1 M. Such a system contains approximately 520 000 atoms. For simulations including ethidium bromide molecules, we increased the size of the water box to 140 × 160 × 350 Å3 to be able to capture conformational changes of the DNA ring while keeping enough distance between the DNA and its periodic images. In this case, the simulation box consists of about 730 000 atoms. The particle mesh Ewald summation method50 has been employed for electrostatic interactions with the time step chosen to be 1 fs. The system has been minimized for 10 000 steps and then simulated for 5 ns at a constant temperature of 298 K and constant pressure of 1 atm while harmonic constraints were imposed on all DNA atoms. The constraints were then released, and the system was equilibrated for 25 ns at constant temperature. At this stage, we take the final equilibrated configuration and start three separate simulations. The first simulation system consists of the DNA ring in the water box. In the second and third systems, we manually insert six and 12 ethidium ions between the DNA base pairs, uniformly distributed along the length of the DNA ring. The interatomic parameters and partial charges of the ethidium ions were taken from a distributed along the length of the DNA ring. The interatomic parameters and partial charges of the ethidium ions were taken from a previous investigation,59 and the initial positions of the ethidiums were manually adjusted to make it as close as possible to the available crystallographic information.61 Accordingly, six and 12 Cl− ions were randomly selected and removed from the two simulation systems to maintain charge neutrality of the system. Each system was minimized for 20 000 steps and equilibrated at constant temperature for 2 ns with harmonic constraints applied to both DNA and ethidiums. The constraints were released, and the system was minimized again for 10 000 steps. After this initial minimization and equilibration, each simulation is continued at constant temperature for 200 ns to observe the effect of the ethidiums on conformational changes of the DNA ring. As a control simulation, the DNA ring has also been simulated in the absence of ethidium bromide for 100 ns. The simulation trajectories are sampled every 1 ps for further analysis. Writh is calculated numerically by discretization of the Gauss double integral along a polygon of 180 segments.62

DNA Molecules. The circular plasmid pTYB21 (7514 bp) in 10 mM Tris and 1 mM EDTA was purchased from New England Biolabs and used for translocation and AFM investigations without further purification.

AFM Imaging. Sample Preparation. For non-EtBr-treated samples, 20 μL of 1x TAE buffer (40 mM Tris base, 20 mM acetic acid, and 1 mM EDTA, pH 8.0) containing 20 mM MgCl2 and 5 μL of 4 nM circular DNA (pTYB21) in 1x TAE buffer were incubated together on a freshly cleaved mica surface for 5 min. The divalent magnesium cations served to screen the negatively charged DNA from the negatively charged mica surface, thereby enhancing adsorption of DNA on the mica. The samples were then rinsed with 1 mL aliquot of ultrapure (18.2 MΩ-cm) water to remove nonadsorbed DNA before adding 20–40 μL of 12 mM NiCl2 in 1x TAE buffer.

For EtBr-treated samples, 20 μL of 1x TAE buffer containing 20 mM MgCl2 and 5 μL of 1x TAE buffer containing 4 nM circular DNA (pTYB21) were incubated together on a freshly cleaved mica surface for 5 min. The samples were then rinsed with a 1 mL aliquot of ultrapure (18.2 MΩ-cm) water before adding 20 μL of 15 mM MgCl2 in 1x TAE buffer. Two microliters of 1.26 μM EtBr was sequentially added to the sample, mixed using a pipet, and allowed to incubate for 5–10 min. Twenty microliters of solution was removed from the sample and immediately replaced by 20 μL of 12 mM NiCl2 in TAE buffer before imaging with the AFM.

Imaging Conditions. All images were acquired in fluid (15 mM MgCl2, in 1x TAE or 12 mM NiCl2, in 1x TAE) using a Bruker MultiMode 8 AFM operated in PeakForce tapping mode (20–40 mV PeakForce set point, corresponding to ≤1 nN) with a Bruker ScanAsyst Fluid+ probe (k = 0.7 N/m, f0 = 120–180 kHz). Additional imaging buffer was added as necessary during imaging to counteract evaporation.

Image Editing Procedure. All image processing was carried out using NanoScope Analysis version 1.50 or WSxM 5.0.63 All images were cropped and then the first-order plane fit to remove sample tilt. If necessary, a first-order line-by-line flatten was subsequently applied to account for slight line-to-line fast axis piezo offsets.

Nanopore Translocation. Silicon nitride nanopores were fabricated in 20 nm thick suspended membrane TEM grids (Norcada) using a JEOL 2010F field emission gun transmission electron microscope. All investigations were conducted using nanopores with diameters of a least ~20 nm to ensure adequate capacity for simultaneous translocation of a molecule with multiple branches. Each nanopore was fixed between two fluid reservoirs (~100 μL each) filled with a buffered ionic solution (1 M KCl, 10 mM Tris, 1 mM EDTA). Two Ag/AgCl electrodes were immersed in the ionic solution and connected to the headstage of an Axopatch 200B amplifier (Molecular Devices), biased at 100 mV. The amplifier was linked to a Digidata 1440 (Molecular Devices) data acquisition system for recording and analysis. Circular covalently closed DNA was added to the cis reservoir, and the recorded signal was filtered with the 10 kHz low-pass hardware filter from the Axopatch amplifier. Intercalation experiments were performed in identical conditions with the addition of 50 μM EtBr added to the cis chamber (0.5 μM final concentration). Attempts to increase the EtBr concentration resulted in longer translocation events with a decreased current drop resembling linear DNA translocation (data not shown) that we hypothesize to be intercalant-induced DNA breaking.8,64 Further characterization of these events is beyond the scope the work presented here. Data analysis was performed with Clampfit (Molecular Devices) and Origin8.5 (OriginLab) software packages and custom routines developed with MATLAB (MathWorks). To prevent collisions from being considered as translocation events, we defined our analysis routine to exclude events with tΔ less than 70 μs and required Δt to be greater than 140 ps.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.nano.6b04876.

Table of ethidium positions for MD simulations; MD simulation of linear DNA translocation; large-area AFM height images of DNA on mica and AFM height profiles (PDF)

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Supporting Information

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Table S1. Positions of the Ethidium molecules in DNA ring simulations.

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Molecular dynamics simulations of translocation of linear DNA through a nanopore.

To study the effect of ethidium bromide molecules on ionic current and translocation time of the DNA, we carried out simulations of a linear double stranded DNA of 50 bp passing
through a silicon nitride (Si$_3$N$_4$) nanopore. The DNA sequence was chosen to be (ApA) except for the first three terminal base pairs which were (CpC) to avoid denaturation at both ends.

The hourglass-shaped silicon nitride pore had a 5.4 nm diameter at the entrance and an inner diameter of 3.8 nm. The partial charges and van der Waals parameters of the pore atoms were taken from a previous investigation. The DNA configuration is created by the 3DNA software using average values of the base pair and base step parameters. We added TIP3P water molecules and K$^+$ and Cl$^-$ ions to achieve a 1 M ionic concentration. Hexagonal periodic boundary conditions were applied. The total simulation box consisted of about 263,000 atoms.

The DNA was initially positioned in such a way that the first two terminal base pairs were inside the pore. During the simulation, harmonic constraints were applied to the Si$_3$N$_4$ atoms to maintain pore geometry during the translocations. The epsilon values in the Lennard-Jones potentials between the DNA atoms and the pore atoms were reduced to one third of their original values while keeping the same sigma values to decrease the friction between the DNA and the pore to make translocation events possible in the current simulation times.

Two sets of simulations were carried out. In the first set, translocation of a linear DNA through the pore was simulated by applying a constant electric field. In the second set of simulations, three ethidium ions were inserted between the base-pairs at positions 12, 25 and 39, and the translocation of the intercalated DNA through the pore was simulated in the presence of a constant electric field.

Each system was equilibrated in a multi-step process. After 10,000 steps of minimization, the system was simulated for 5 ns at a constant temperature of 298 K and constant pressure of 1 atm while the first two DNA base-pairs were harmonically constrained. In the next step, we applied a constant electric field corresponding to a voltage of 1 V across the pore and released
the harmonic constraints on the DNA while simulating the system at constant volume and a constant temperature of 298 K. For those simulations that involve ethidium ions, we inserted the ions in such a way as to be close to the available crystallographic data of the local orientation of the intercalated ethidiums and carried out an extra 10,000 steps of minimization while having constraints on the DNA.

Each system was simulated for 60 ns, during which the entire DNA molecule was translocated through the pore. To achieve good statistical information on the translocation events, each simulation was carried out three times starting with different initial velocities. For every simulation, the number of translocated base pairs and the ionic current versus time was recorded. In addition to the above simulations, we performed another one on a system that did not contain a DNA molecule to obtain the open pore current of the system at a voltage of 1 V.

Figure S1a shows the number of base pairs that are passed through the center of the pore versus time. The presence of ethidium ions decreases the translocation time due to their positive charge and extra mass added to the DNA string.

To observe whether intercalated ethidiums affect the current blockade, we calculated the ionic current from:

\[ I(t) = \frac{1}{\Delta t L_z} \sum_{i=1}^{N} q_i \left( z_i(t + \Delta t) - z_i(t) \right) \]  

(1)

where \( I(t) \) is the current at time \( t \), \( q_i \) is the partial charge of atom \( i \), \( \Delta t = 1 \) ps, \( L_z \) is the length of the simulation box along the pore, and the summation is over all the atoms carrying partial charges in the box. The atomic coordinates are recorded every 1 ps and the current is blocked averaged over every 500 data points.

The average current over the three realizations for each simulation is plotted in Figure S1b. Snapshots of the simulation trajectory of the DNA carrying ethidiums are shown in Figure S3.
It can be seen that in both cases (with and without ethidiums) DNA translocation blocks the ionic current of the pore. The results show that although they increase the blockade time, intercalated ethidium ions do not have a significant effect on the current drop.

Figure S2 shows the current drop as a function of time for each of the three translocation events shown in Figure S1. Despite thermal fluctuations, the three curves show the same overall pattern as the averaged curve (black line) indicating a well-defined behavior.

Since the length of the DNA in our simulations is comparable to the persistence length of the double stranded DNA, the results of the simulations may not be directly comparable to experiments. As the DNA length becomes comparable to or larger than the persistence length of the DNA, entropic effects become more important in determining the translocation time of the DNA through nanopores, thus, making the comparison between our simulations and experimental data unrealistic. However, as the DNA passes through the pore, the drop in the ionic current due to blockage of the pore depends mainly on the hydration diameter of the DNA rather than its length (as long as the length of DNA is longer than the pore length), and as long as ethidium intercalation does not change the persistence length of the DNA, one can compare the change in the ionic current before and after intercalation while neglecting the entropic effect. Our simulations suggest that even at pore diameters comparable to the effective diameter of the DNA, ethidium does not change the amount of the current drop indicating that ethidium intercalation does not change the effective diameter of the DNA.
Figure S1. A 260,000 atom all-atom molecular dynamics simulation of the translocation of a 50 bp linear DNA intercalated with 3 ethidium ions through a solid-state nanopore. (a) Number of base pairs translocated through the center of the pore averaged over three separate simulations for a linear DNA double strand (blue) and the same DNA carrying three intercalated ethidium ions (red). (b) Current versus time for linear DNA (blue) and DNA with ethidium (red). The average open pore current is shown with a green line. (c) Snapshots of the translocation with 1 V transmembrane voltage and 1 M ion concentration.
Figure S2. The current drop for each of the three translocation events of the double-stranded DNA with bound ethidiums. Green, blue, and red curves show the currents calculated for each trajectory, and the black curve show the average current over the three runs.
Figure S3. Large area AFM height images of (a) circular DNA on mica with no EtBr and (b) circular DNA on mica with EtBr. Without EtBr, the circular DNA appears as large loops or bunched up piles of loops. Even in the loop piles, the circular DNA appears to exhibit very little branching. With EtBr, the circular DNA exhibits a very different structure with a large degree of branching. Images were processed using WSxM 5.0.\textsuperscript{8}
Figure S4. Three AFM height profiles across circular DNA with EtBr on mica. The color of the profile corresponds to the color of the profile line in the AFM image. The blue profile crosses an apparent single double helical section and yields a height of ~1.5 nm. The magenta and green profiles cross sections in which double helices appear to twist around each other and yield heights of ~2.8 nm, almost double the height of the single double helix section. Image was processed using WSxM 5.0.⁸
References


