Simulations of Electrophoretic RNA Transport Through Transmembrane Carbon Nanotubes

Urs Zimmerli and Petros Koumoutsakos
Computational Science and Engineering Laboratory, ETH Zürich, Switzerland

ABSTRACT The study of interactions between carbon nanotubes and cellular components, such as membranes and biomolecules, is fundamental for the rational design of nanodevices interfacing with biological systems. In this work, we use molecular dynamics simulations to study the electrophoretic transport of RNA through carbon nanotubes embedded in membranes. Decorated and naked carbon nanotubes are inserted into a dodecane membrane and a dimyristoylphosphatidylcholine lipid bilayer, and the system is subjected to electrostatic potential differences. The transport properties of this artificial pore are determined by the structural modifications of the membrane in the vicinity of the nanotube openings and they are quantified by the nonuniform electrostatic potential maps at the entrance and inside the nanotube. The pore is used to transport electrophoretically a short RNA segment and we find that the speed of translocation exhibits an exponential dependence on the applied potential differences. The RNA is transported while undergoing a repeated stacking and unstacking process, affected by steric interactions with the membrane headgroups and by hydrophobic interaction with the walls of the nanotube. The RNA is structurally reorganized inside the nanotube, with its backbone solvated by water molecules near the axis of the tube and its bases aligned with the nanotube walls. Upon exiting the pore, the RNA interacts with the membrane headgroups and remains attached to the dodecane membrane while it is expelled into the solvent in the case of the lipid bilayer. The results of the simulations detail processes of molecular transport into cellular compartments through manufactured nanopores and they are discussed in the context of applications in biotechnology and nanomedicine.

INTRODUCTION

Recent advances at the interface of nanotechnology and molecular biology present us with new opportunities for the development of devices such as nanoscale biosensors, molecular motors, and nanosyringes for application domains ranging from bioengineering to nanomedicine (1,2). Biological cells rely on nanoscale components such as ion channels and pumps that regulate key transport processes across cell membranes (3,4). Molecular transport across membranes may also be inflicted by actions of external agents such as viruses binding and puncturing the cell membrane to transfect it with their nucleic acids (5).

The critical role of transmembrane channels and pores in cell biology has prompted several efforts to develop synthetic structures (1,6), with characteristics such as ion and membrane selectivity, voltage and ligand gating, and blockage. Several techniques have been developed for the synthesis of artificial channels based on unimolecular macromolecules, micellar pores, and barrel-stave and barrel-hoop supramolecules (6). The requirements for these channels include structural stability, appropriate trans-membrane orientation, and regulated channel activity (1). These requirements are not always easy to achieve by self-organizing structures; at the same time, the activity regulation of these channels remains an ambitious goal. Alternatively, we may consider the implantation of artificial channels and pores in cell membranes by processes such as conjugation of biological molecules and synthetic polymers as well as by and in situ forming implants (7).

The requirements for structural reliability and low reactivity are met by silica and carbon nanotubes, which are envisioned as prototypes of artificial transmembrane channels and nanosyringes for targeted drug delivery (2). Transmembrane transport in natural ion channels, however, is largely controlled by the structure of membrane proteins whose structure and precise molecular actions may not be easily reproducible with today’s nanofabrication techniques. Simpler structures, such as silica-based nanopores (8), can be manufactured, which may provide us with enhanced capabilities for molecular sequencing and selective molecular transport. This possibility has been thoroughly investigated in a series of recent articles (9–13) that reported on the electrophoretically driven transport of DNA in nanometer-size silica pores. In Fyta et al. (14), the molecular dynamics (MD) simulations have been coupled with Lattice-Boltzmann models to study DNA translocation through a pore for extended timescales. The interface between biological systems and artificial nanopores was recently assessed by means of coarse-grained simulations (15–17). These simulations demonstrated that a generic, hydrophobic nanotube functionalized with hydrophilic ends can be spontaneously inserted into a lipid bilayer. In this work, we consider a specific carbon nanotube and we analyze the detailed molecular interactions of the membrane components with the nanotube using molecular dynamics.
Carbon nanotubes are consistent with the requirements for a model nanopore (17), as they are hydrophobic and can be decorated for insertion in membranes. Their low reactivity (18,19) suggests their use as nanometer-size syringes for drug delivery (20) and storage (21). In addition, CNTs can withstand a broad range of voltages and they have been suggested as artificial membranes for DNA sequencing and as stabilizing pores in biological membranes during electroporation. Several recent studies have assessed the interaction of small CNTs with water (22–26), ions (27,28), and nucleic acids (29–31). The regulated channel activity by nanotubes remains a challenge, however, because it requires the integration of suitable gating molecules during their manufacturing. At the same time, the relatively simple structure of the nanotubes, reminiscent of pores consisting of unimolecular macromolecules, can help us to assess the role of complex gating structures in biological channels.

In this study, we use MD simulations to characterize the behavior of decorated and naked CNTs embedded in membranes and assess the role of the CNT decoration for its stabilization in the membrane. We analyze the electrophoretic transport of RNA through these CNTs and quantify interactions of the membrane constituents with the carbon nanotubes. We find that the interplay between the lipid heads and the opening of the nanotube characterizes the electrostatic potential of the CNT, particularly during the electrophoretic transport of biomolecules. The article is structured as follows: In Methods, we describe the setup of the molecular dynamics simulations and present a novel method for inserting and stabilizing CNTs in membranes. In The Electrostatic Potential, we discuss the electrostatics of the CNT embedded in the dodecane membrane and the lipid bilayer, followed by RNA Translocation in Transmembrane CNTs with results from the simulations of the electrophoretic transport of RNA across a membrane-mimetic and a lipid bilayer. In the final section, we conclude with a summary of our findings and their implications for applications at the interface of biology and nanotechnology.

METHODS

Molecular dynamics

Molecular dynamics is a key methodology for the evaluation of new technologies and biological systems at the nanoscale (32). In this study, we employed the MD package FASTTUBE (33), developed in our group for the simulation of carbon nanotubes in aqueous environments, and we visualized the results using VMD (34). The electrostatic interactions were computed using the smooth particle-mesh Ewald method (35). A 32 × 32 × 48 grid was used and validation studies using a 48 × 48 × 64 grid did not show any significant differences in quantities such as the electrostatic potential maps and the structure and speed of translocation of the RNA. The van der Waals interactions and the real-space contribution to the Coulomb interaction were calculated using a cutoff of 1 nm. All systems were kept at a temperature of 323 K by applying a Berendsen thermostat and equilibrated to atmospheric conditions using a Berendsen barostat with a characteristic timescale of 0.1 ps. Using the Berendsen thermostat, the system was equilibrated by a Berendsen thermostat and a Berendsen barostat acting in the direction orthogonal to the membrane for 180 ps (36) each, with characteristic times of 0.1 ps. Using the Berendsen barostat, after equilibration, the system was relaxed at constant temperature for another 20 ps. The final extent of the computational domain was 6.1 nm × 6.1 nm × 8.214 nm. The coordinates for the dimeristoylphosphatidylcholine (DMPC) lipid bilayer were obtained from the Tieleman group (PDB file dmpc_npat.pdb from http://moose.bio.ucalgary.ca/index.php?page=Downloads). These coordinates are the result of a 1-ns-long simulation of 128 DMPC molecules solvated in 3655 water molecules.

The dodecane membrane mimetic was set up in a 6.1 nm × 6.1 nm × 10 nm computational box. A slab of 192 dodecane molecules in random orientation with a height of 2.5 nm were placed between two slabs of 3610 water molecules, each 3.1 nm thick. After energy minimization, this system was equilibrated by a Berendsen thermostat and a Berendsen barostat acting in the direction orthogonal to the membrane for 180 ps (36) each, with characteristic times of 0.1 ps. Using the Berendsen barostat, after equilibration, the system was relaxed at constant temperature for another 20 ps. The final extent of the computational domain was 6.1 nm × 6.1 nm × 8.214 nm. The coordinates for the dimeristoylphosphatidylcholine (DMPC) lipid bilayer were obtained from the Tieleman group (PDB file dmpc_npat.pdb from http://moose.bio.ucalgary.ca/index.php?page=Downloads). These coordinates are the result of a 1-ns-long simulation of 128 DMPC molecules solvated in 3655 water molecules.

The carbon nanotubes are based on a (14,14) armchair-CNT with a length of 2.35 nm and a 1.87-nm carbon-to-carbon diameter corresponding to a pore size of ~1.5 nm (29). We considered two variations of this CNT: The “naked” CNT (with unfilled valences at the ends) and a decorated/doped CNT, where alternating hydroxyl and hydrogen termini were added at its edges (H/OH CNT). The CNT diameter is comparable with the constriction...
region of the α-hemolysin channel (9,29,43,) and the CNT length is selected so that it minimizes the hydrophobic mismatch with the DMPC membrane (44,45). Due to the limited size of the computational domain, the membrane could not compensate for the effects of a hydrophobic mismatch, which led to destabilization of the lipid bilayer structure.

In these simulations, we considered the transport of single-stranded RNA consisting of 20 adenosine nucleotides. The RNA is hydroxylated at both ends and the charge of the whole system is neutralized by randomly adding potassium counterions to the solvent.

Membrane-CNT setup and equilibration

We present a snapshot (Fig. 1) from the simulations of the insertion of the RNA into the CNT to define the geometry of the configuration. Using a cylindrical coordinate system (r, θ, z), the nanotube axis is nominally aligned along the z axis while the membrane surface is largely parallel to the (r, θ) plane.

Embedding the CNT in a membrane

The embedding of a CNT in a membrane is hindered by the irregular, unsteady distribution of the lipids. To address this difficulty, Faraldo-Gomez et al. (46) presented a two-stage method, first generating a cavity in a lipid bilayer and then using steered molecular dynamics to insert it in a protein structure.

We introduce a novel technique for the insertion of CNTs into pre-equilibrated membranes. As a first step, the solvent around the membrane is removed and the bilayer structure is adjusted to accommodate the CNT. In the case of the DMPC bilayer, 14 lipid molecules were removed from the central area of the bilayer corresponding to the nominal volume that would be occupied by a (14,14) CNT. No molecules were removed from the dodecane membrane. Subsequently, a scaling technique was used to insert the CNT: the CNT was initially contracted to a single line of carbon atoms, threaded into the cavity opened in the DPMC bilayer or into the unaltered dodecane membrane. The length-scale parameters σ of all the Lennard-Jones interactions of the CNT molecules were initially set to zero and then linearly scaled to their original value during a period of 2 ps. In the next 28 ps, the radius of the CNT was grown back to its original size. During this process, the positions of the CNT atoms were adjusted so that they correspond to a linear scaling of the CNT radius. Furthermore, the motion of the membrane atoms was restricted to the two in-plane dimensions and the respective velocity components of the membrane atoms were capped at a maximal value of 0.3 nm ps⁻¹. The system was equilibrated for 1 ns and subsequently samples were taken every 1 ps during the course of 1 ns.

We note that this approach differs from the insertion technique presented in Faraldo-Gomez et al. (46) in that it does not require any surface triangulation and it relies on quantities that are computed routinely in MD simulations.

Solvation

After the insertion of CNT into the membrane, the system is solvated by two slabs of ~1 molar potassium chloride solution. In both cases, 150 potassium and 150 chlorine ions were added to 8127 (for the dodecane membrane) and 8211 (for the DMPC bilayer) water molecules.

For the DMPC bilayer, during the first 40 ps of solvation all phosphorous atoms are kept fixed and the water is relaxed to wet the membrane. This is followed by 4 ps of dynamics where the phosphorous atoms are restricted to move only within the plane of the membrane. After further relaxation and a total simulation time of 60 ps, the system is relaxed using a Berendsen barostat during 200 ps. This is followed by a further relaxation at constant volume for 40 ps.

In the case of the dodecane membrane, its molecules were kept fixed during the first 40 ps of simulation and the water was relaxed to wet the membrane. Subsequently, the velocity of the dodecane molecules was capped at 0.3 nm ps⁻¹ for 4 ps. After a total simulation time of 80 ps, the system was coupled to a Berendsen barostat for 120 ps. The system was further equilibrated at constant volume for another 100 ps to obtain a total of 300 ps for solvation, relaxation, and equilibration.

Insertion of RNA into CNTs

The simulations of RNA transport through the CNT were initialized by placing the RNA in the solvent so that the first phosphorous atom from the 3'-terminus is situated 0.8 nm outside the CNT and on the tube axis. The whole RNA was initially restrained for 46 ps and subsequently only the first phosphorous atom from the 3'-terminus was restrained, while the rest of the RNA was allowed to move freely. Both, the RNA and the membrane were solvated simultaneously.

After solvation and relaxation, the RNA was steered into the CNT (47). A harmonic guiding potential with a force constant of 30,000 kJ nm⁻² mol⁻¹ was applied to the first phosphorous atom from the 3'-terminus. The RNA was threaded into the CNT by moving the reference point of the steering potential along the CNT axis during 280 ps. After the steering, the phosphorous atom was restrained at the opening of the CNT to allow for relaxation of structural strain imposed during the steering.

THE ELECTROSTATIC POTENTIAL

The electrostatic potential maps are used to characterize the transport in transmembrane pores (48). We study naked and H/OH-decorated carbon nanotubes, embedded in a dodecane
membrane and a lipid bilayer and subjected to an electrostatic potential difference. The random distribution of dodecane molecules and the ordered structure of the lipid molecules are reflected on the electrostatic potential of the system. In the interior of the membrane, the lipid tails of the DMPC have a composition similar to that of the molecules in the dodecane membrane, but their lipid headgroups carry partial charges. The headgroups interact with the decorated rims of the CNT and alter the electrostatic potential in the vicinity of the membrane and near the openings of the transmembrane pore, as detailed later.

The electrostatic potential is assessed following the approach of Aksimentiev et al. (9). We evaluate the electrostatic potential using the method of SPME (33,35). The instantaneous electrostatic potential is radially averaged over 1 ns and presented as an axisymmetric electrostatic potential map. In the related figures, the electrostatic potential map is complemented with the contour lines of the number density for characteristic atoms of the respective setup. These density profiles serve to illustrate the location and mobility of the membrane and the carbon nanotubes.

**Electrostatics of a CNT in a membrane**

We first present the electrostatic characteristics of a CNT embedded in a membrane without the application of an external potential. The distribution of the electrostatic potential helps us to assess the characteristics of the CNT as a transmembrane channel.

**CNT in a dodecane membrane**

The electrostatic potential map of an H/OH-CNT in an \( \sim 2.5 \) nm thick dodecane membrane is presented in Fig. 2 along with radially averaged density profiles of their constituent atoms. The hydroxyl and hydrogen termini stabilize the CNT in its transmembrane position and the hydrophilic interaction between the water solution and the termini prevents the dodecane molecules from creeping around the tube rim. The CNT interior forms a channel of low electrostatic potential across the membrane. The electrostatic potential map exhibits distinct zones of high and low potential that are highly correlated with the location of the membrane and the CNT. We measure a potential difference of 0.65 V between the bulk of the ionic solution and the interior of the dodecane membrane, in close agreement with the potential difference found between bulk water and the interior of a lipid bilayer (38).

**CNT in DMPC bilayer**

The electrostatic potential maps for the decorated H/OH-CNT and for the naked CNT in a DMPC lipid bilayer are shown in Fig. 3. The H/OH-CNT exhibits a channel of low electrostatic potential connecting both sides of the membrane. On both ends of the membrane, zones of high electrostatic potential protrude into the nanotube and constrict the effective channel opening by 50\%, to \( \sim 1 \) nm in diameter. For the naked CNT these effects are more pronounced, and we observe zones of high electrostatic potential extending across the whole channel and forming barriers (Fig. 3).

The CNT-DMPC electrostatic potential maps are significantly different from the ones reported for the CNT in the dodecane membrane. The differences are attributed to the DMPC lipid heads that extend further into the solution to access water molecules to satisfy their hydrophilicity. The lipid molecules are arranged into an hourglass shape around the CNT, widening above the area of constriction, while the lipid headgroups are fanning out over the tube rims partially covering the pore openings. This effect is well illustrated by the radially averaged density profiles shown in Fig. 3.

The differences between the naked and the H/OH-CNT are attributed to the solvation of the ester groups of the lipids. The ester groups are stabilized by forming hydrogen bonds with the hydrogen and the hydroxyl groups at the decorated tube rim (Fig. 4). In the case of the naked CNT, this interaction is lacking and the lipids slide along the CNT and toward the water phase, so that the ester groups come into contact with the water at the tube rim (Fig. 4). This configuration is energetically more favorable, allowing for solvation of the carbonyl groups, while at the same time the lipid headgroups extend further over the tube rim, allowing the ammonium groups to leap into the area of the tube opening (Figs. 3 and 4).

The fanning of the lipid heads into the tube opening is reflected in the distribution of the standard deviation of the
electrostatic potential (Fig. 5) having large values in the zone occupied by the lipid headgroups. The charge separation of the choline group promotes significant fluctuations (of \( \sim 5\,\text{ns}^{-1} \)) of the electrostatic potential, which are more pronounced for the naked CNT than for the H/OH-CNT. These oscillations are attributed to the strongly correlated reorientation of the lipid headgroups (Fig. 6).

We note that the hydrophilicity of membrane heads and the corresponding CNT rim decorations are critical in determining the access to the nanotube entrance. This finding hints at possible regulation and gating mechanisms that may be accomplished by specifying the molecules for the CNT rims so as to control their interactions with the corresponding lipid heads.

Application of a transmembrane voltage difference

A transmembrane voltage difference (8,9,48) is applied to drive a single-stranded RNA segment of 20 adenosine nucleotides through the CNT pore embedded in the membrane.

We first demonstrate that the electrostatic voltage differences induce an asymmetry on the lipid head arrangement on the entrance and on the end-side of the CNT, and affect the overall electrostatic potential maps. We then discuss the electrophoretic transport of the RNA through the CNT and identify a stacking process that depends on steric interactions with lipid heads at the tube entrance and hydrophobic interactions with the walls of the nanotube.

CNT in a dodecane and a DMPC membrane

An electrostatic potential of 0.95 V is applied across the system of the H/OH-CNT embedded in the dodecane membrane. The electrostatic potential (Fig. 7) is largely uniform on either side of the membrane, although it changes sharply within the transmembrane nanopore. The potential maps are qualitatively similar to the corresponding maps of solid-state nanopores presented by Heng et al. (10).

In the case of the DMPC membrane, an electrostatic potential difference of 1.05 V is applied, resulting in the electrostatic potential maps shown in Fig. 8 for both naked and H/OH CNTs. We observe a constant electrostatic potential for the H/OH-CNT on both sides of the membrane accompanied by a high barrier on the side of high electrostatic potential just outside the opening of the CNT (Fig. 8). On the side of low electrostatic potential, a similar barrier is absent for the H/OH-CNT, leading to an opening reflected in a bay of low electrostatic potential extending into the pore. This distribution is due to the absence of lipid headgroups in the area of that tube opening, which is also confirmed by the radially averaged density profiles (Fig. 8).

A high barrier on the side of the high potential is also observed in the case of the naked CNT. In contrast, however, with the H/OH-CNT, this barrier is inside the CNT opening. The asymmetry of the electrostatic potential map of the H/OH-CNT is attributed to a reorientation of the membrane, subject to the electrostatic potential differences in the vicinity of the tube entrance. The positively charged ammonium groups of the lipids (38) on the side of high electrostatic potential are pushed by \( \sim 0.7\,\text{nm} \) into the area of the H/OH-CNT opening (compare Fig. 9). For the naked CNT, we observe a similar effect with an ammonium density peak just
outside the tube opening that is pushed ~0.7 nm into the tube (Figs. 3 and 8) On the membrane side with low electrostatic potential, the ammonium groups are subject to electrostatic forces which, in turn, pull them away from their equilibrium position, leading to their upright position close to the tube entrance (Fig. 9).

In the case of the H/OH-CNT, the lipid headgroups are hardly leaping into the tube area even in the absence of an electrostatic potential difference (Fig. 4). As a consequence, the electrostatic potential difference has a negligible effect on their density distribution on the side of low electrostatic potential (Figs. 3 and 8). Contrastingly, for the naked CNT in the absence of an electrostatic potential difference, the lipid headgroups are leaping into the area of the tube opening (3). As the electrostatic potential difference is applied, the ammonium density retracts by ~0.5 nm away from the tube axis. The electrostatic barrier reported in the absence of an electrostatic potential difference across the membrane is weaker but still persists in the presence of a potential difference.

We note that the electrostatic potential maps persist for potential differences between 0.2 V and 1.5 V and they are significantly different from the maps obtained for a CNT in a dodecane membrane (Fig. 7) and for solid-state nanopores (11). The lipid headgroups reorient when they are subjected to an electrostatic potential difference leading to different barriers of electrostatic potential for H/OH and naked CNTs. These differences are an essential feature of the steric interactions between an artificial nanopore and a lipid bilayer, and they need to be taken into consideration when designing the insertion of synthetic nanopores (2) in biological membranes. The sensitivity of the lipid head arrangement in naked and decorated CNTs can provide guidance for the design of suitable unsteady voltage differences across the CNT that may regulate access to its entrance. The interplay of nanotube-based pores and the lipid heads is further exemplified in the following section presenting the translocation of RNA through a CNT.

RNA TRANSLLOCATION IN TRANSMEMBRANE CNTS

The translocation of ions, DNA, and RNA through biological and synthetic pores has been investigated recently by large-scale MD simulations (10,11,27,32,49) to provide guidance for the design of innovative sequencing devices. In this article, we present MD simulations of RNA transport through a CNT embedded in a dodecane membrane and a DMPC bilayer to study the interplay of the lipid bilayers and the CNT.
in the translocation of RNA molecules. These simulations provide insight for applications ranging from drug delivery (50) to molecular machines (1,51), where artificial pores are implanted in lipid bilayers.

RNA across a dodecane membrane

A single-stranded piece of RNA consisting of 20 adenosine nucleotides is transported through a naked and H/OH-CNT embedded in a dodecane membrane. The transport is driven by transmembrane potential differences of 1.50–2.10 V in steps of 0.1 V. The potential differences employed in this study are significantly lower than potential differences employed for DNA translocation across synthetic pores (27,49) as the bilayer membrane is ruptured for potential differences of 2.4 V and beyond.

During the translocation, the entrance and exit of the RNA bases in the nanotube is affected by steric interactions of the CNT with the heads of the lipid bilayers, with the RNA bases folding backward with regard to the direction of translocation (52). The transport inside the pore is characterized by hydrophobic interactions between the nucleotide bases (29,30) and the walls of the carbon nanotube (Fig. 10).

Inside the tube, the RNA bases maximize contact with the CNT, sliding across the pore with their plane largely parallel to the tube walls (Fig. 11). Single and stacked pairs of nucleotide bases are observed, unstacking at the entrance of the CNT, while small groups of RNA bases are observed to re-stack at their exit from the CNT. During the translocation, the RNA backbone is exposed toward the tube center (Fig. 12), solvating the phosphate and sugar groups of the backbone by the water inside the CNT. This arrangement of the RNA is persistent throughout the simulations and suggests the possibility of using functionalized nanotubes to identify and interact with the RNA bases.

We quantify the translocation of the RNA by relating the speed of translocation with the potential differences applied across the membrane. The speed of translocation is defined by the time between the exit of nucleotide 3 from the tube and the exit of nucleotide 16. This definition was chosen to exclude contributions by end-effects such as nucleotides being expelled from the tube entrance and reduced hydrophobic interactions at the end of the simulation when the RNA is not extended across the whole CNT. For potential differences of 1.20 V and beyond, the RNA was driven through the CNT in relatively short time spans of \( \sim 10 \) ns. For potential differences between 1.20 and 2.15 V we observe a correlation with the speed of translocation measured in nucleotides per nanosecond (Fig. 11). The measurements were fitted with an exponential function, common in kinetic theories and in drift-diffusion models of molecular translocations in pores (29). The parameters of this exponential were determined by a least-squares fit resulting in an empirical relation between the activation voltage difference (\( \Delta V \)) and the translocation speed (\( v \)), 

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v = 0.087 (\text{Nuc/ns}) \exp(-2.57\Delta V)
\]

The speed of translocation exhibits large fluctuations (Fig. 12) and a
staircase pattern for potential differences \(<1.55\) V with the longest stop of translocation lasting for \(\sim1.2\) ns. These events are attributed to repeated conformational changes of the RNA during its transport through the CNT. The RNA bases form stacks as they approach the entrance of the CNT, as their hydrophobic interactions intensify. This layering of the RNA chain is broken when a base enters the CNT and the interfacial area is hydrated. In turn, a secondary hydrophobic interaction takes place between the base and the walls of the CNT. The base approaches the CNT wall and aligns itself parallel to it, while at the same time, water molecules are being expelled from that region. The RNA bases are folding backward with regard to the direction of transport, leading to short temporal trappings. This is manifested in the steplike behavior of the trajectories as the RNA bases overcome the energy barrier and unstack.

The strength of the electrostatic potential difference controls the stacking and unstacking of the bases, and the switch between their hydrophobic interaction outside the entrance of the CNT and their hydrophobic alignment with the walls inside the CNT. At electrostatic potential differences of...
0.955 V, the RNA gets trapped after 1 ns, as the first two nucleotides are expelled from the tube. In the next eight nanoseconds, no noticeable motion was observed, although the RNA inside the CNT exhibits spatial fluctuations of ~0.2 nm along the tube axis. The RNA outside the entrance of the CNT moves similarly to free RNA in solution, until, after another seven nanoseconds, one of the terminal attaches to the membrane and remains attached to the membrane throughout the rest of the simulation.

We note that, even at higher electrostatic potential, the RNA bases do not leave the CNT spontaneously, but are eventually pushed away by the remaining RNA bases sliding down within the CNT. One of the first nucleotides, however, usually reattaches quickly to the outside of the tube or the dodecane membrane. The RNA leaving the tube, buckles and forms a loop, which eventually turns into a coiled-up structure. The expelled RNA remains coiled-up at the tube exit during these limited simulation times of 5–10 ns. The RNA was not detached completely from the CNT in any of the simulations conducted in this study that had one or two of the terminal bases always hydrophobically attached to the CNT. (This is similar to the results reported in (8) for synthetic pores.)

For larger potential differences, we observe similar trapping and attachments of the RNA with intermittent periods of fast translocation with speeds of four nucleotides per nanosecond for an electrostatic difference of 1.2 V. For potential differences larger that 1.55 V, we observe trapping of the RNA, but we did not observe any events where individual nucleotides leaving a stacked geometry and entering the tube could be identified by a steplike trajectory as found for smaller potential differences (Fig. 12). The strong driving forces at large potential differences mask the signature of the sequential breaking of the stacking.

**RNA across a DMPC bilayer**

Simulations of RNA transport are performed for a tube embedded in a DMPC bilayer, to study the effects induced by the structure of the lipid bilayer membrane. The same single-stranded piece of RNA consisting of 20 adenosine nucleotides is transported through a naked CNT and an H/OH-CNT embedded in a DMPC bilayer, which was subject to a potential difference of 1.6 V.

The lipid headgroups of the DMPC membrane extend over the CNT and interact with the RNA (Fig. 13), slowing down its translocation in the tube. In addition, the motion of the RNA is sterically hindered as the lipid heads obstruct the pore opening, similar to the effect described by de Groot et al. (53) and comparable to the entropic barriers reported in Lim et al. (54).

The RNA structure within the CNT in the DMPC lipid bilayer does not significantly differ from the structure in the case of the dodecane membrane. The RNA bases are folded backward with regard to the direction of transport, and they are hydrophobically attracted to the walls of the CNT exposing the bases toward the center of the tube. The translocation (Fig. 14) of the RNA through the tube takes ~12 ns with an average speed of translocation of 1.5 nucleotides/ns. This speed is significantly lower than the one observed in the case of the dodecane membrane, for similar voltage differences. This effect is attributed to the increased steric interactions of the lipid heads with the RNA. And contrary to the case of dodecane membrane, the RNA does not reattach to CNT as the lipid molecules shield the exterior of the CNT. The RNA leaves the exit area and protrudes into solution while after ~10 ns the foremost nucleotides get into contact with the lipid bilayer. The RNA, however, is not adsorbed on the lipid bilayer surface but readily diffuses back into solution.

It is important to recall here that polarizability effects have not been accounted for in these simulations, largely due to their computational expense. We estimate, however, based on previous findings (41,42), that the inclusion of polarizability effects in this configuration would have increased the affinity of the RNA with the walls of the CNT, thus further enhancing hydrophobic effects and further delaying the transport of the RNA through the CNT. The study of polarizability effects in the electrophoretic translocation of RNA in pores and channels, and the embedding of the MD simulations in a multiscale framework where solvation effects are accounted for by these mean-field theories, is a subject of our current investigations.

**SUMMARY AND CONCLUSIONS**

We use MD simulations to study the electrophoretic transport of a short, single-stranded RNA segment through naked and decorated carbon nanotubes embedded in lipid bilayer and dodecane membranes. These simulations rely on a novel method to insert nanotubes in bilayers. The results enable observations with molecular detail for the interactions of the nanotube with the membrane components. Simulations with dodecane membranes and DMPC lipid bilayers demonstrated that lipid heads, and their coordination with the molecules at the rim of the nanotube, determine to a large extent the
transport properties of the nanotube. These properties are further quantified by presenting electrostatic potential maps in the vicinity of the nanotube with and without electrostatic potential differences. We note in particular that the lipid heads of the DMPC bilayer extend above the CNT rims to be solvated by water, thus blocking the entrance of the nanotube and forming an electrostatic potential barrier. This effect may help explain the structure of biological transmembrane pores that exited into the solution.

The transport properties of the CNTs were further quantified by measuring the speed of translocation of a short, single-stranded RNA segment. During the electrophoretic transport of RNA, the lipid heads in the DMPC membrane were found to interact sterically with the transported RNA bases and reduce by half its speed of translocation when compared to the case where the CNT is embedded in a dodecane membrane. In addition, the RNA was found to stay attached to the dodecane membrane, after its exit from the CNT, similar to a behavior that has been observed for other artificial pores. In contrast, it appears that the DMPC lipid heads help the RNA to overcome an entropic barrier so that it enters the solvent after its exit from the CNT. We note, however, that even at strong transmembrane potential differences a few of the terminal nucleotide bases remain hydrophobically attached to the CNT. Chemical modifications to the CNT interior and the tube rims that weaken the hydrophobic interaction of RNA bases and the CNT could promote the detachment of the RNA from the CNT. Possible applications may, however, benefit from the fact that RNA does not detach from the CNT, since the sustained attachment may allow the retraction of the RNA after exposing it to the cell interior. The transport of RNA across a carbon nanotube transmembrane pore compares well with results reported on synthetic silica nanopores (10) and the effects of constriction are similar to results on α-hemolysin (11). While transport velocities in transmembrane CNTs seem to exceed those of an α-hemolysin, they seem to be slower than those observed in synthetic silica pores (10).

Inside the nanotube, the CNT exhibits a strong hydrophobic interaction with the RNA bases, which align themselves to the walls of the nanotube, leading to a stacking and unstacking process during the transport of the RNA. The preferential orientation of the RNA bases inside the nanotube may form the basis of rational functionalization of the CNT interior for rapid sequencing protocols and the development of novel molecular assembly scaffolds.

The results of these simulations help to demonstrate that embedding carbon nanotubes inside membranes requires a careful consideration of their functionalization and their interactions with the surrounding membrane components. The interaction of suitably decorated CNT rims and membrane components may be exploited to regulate the transport properties of the nanotube, an important feature for biomedical applications such as artificial nanosyringes, controlled release drugs (50), and vectors for transfection (55).

Finally, we note that the design of devices based on the interface of biological molecules with carbon nanotubes has to take into account the toxicity of the nanotubes and identify processes by which the administered CNTs (56) and fullerenes (57,58) will leave the body. Recent studies have shown that water-soluble fullerenes do not enter membranes. The results of these simulations indicate that decorated CNTs are
well accommodated in lipid bilayers, resulting in well-defined transmembrane channels that, at the same time, may lead to increased toxicity and cell lysis.

Current work is geared toward modeling and assessing polarizability effects in membrane-CNT systems and the embedding of these MD simulations in a multiscaling framework. As imaging technologies are progressing toward capturing the motion of single molecules in motion inside CNTs (59), we are entering an era where the results of MD simulations of translocating molecules may be validated by experimental works.

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REFERENCES