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Role of Intercalation on Electrical Properties of Nucleic Acids for use in Molecular Electronics

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Complete List of Authors:	Mohammad, Hashem; University of Washington, Electrical Engineering Demir, Busra; TOBB University of Economics and Technology, Department of Materials Science and Nanotechnology Engineering Akin, Caglanaz; TOBB University of Economics and Technology, Materials Science and Nanotechnology Engineering Luan, Binquan; IBM Research, Hihath, Joshua; University of California, Davis, Electrical and Computer Engineering Oren, Ersin Emre ; TOBB Economics and Technology University, Department of Biomedical Engineering; TOBB Economics and Technology University, Department of Materials Science and Nanotechnology Engineering Anantram, M.; University of Washington, Electrical Engineering	



"New Concepts" statement:

Facilitated by the advance of synthetic biology, designer DNA with predetermined sequences can be readily synthesized for various applications (e.g., biosensors, single molecule transistors and DNA storage) in the nascent field of molecular electronics. However, the lack of control on charge transport significantly limits its practice in nanoelectronics. While experiments have shown that the conductivity of DNA can be adjusted via intercalation with organic molecules, the mechanism is still not established. In this work, using rigorous calculations, we addressed this problem by studying the effect of intercalation on both the molecular structure of and charge transport along the nucleic acids. We estimated the location of Fermi energy using partial charge transfer analysis. This helped us show that the conductance of DNA can be altered depending on the redox state of the intercalator, induced energy levels, and the location of the Fermi energy. These results establish the theoretical basis for intercalation-regulated conductance and thus can guide the design of novel intercalator-nucleic acid systems for use in future molecular electronic devices.

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Role of Intercalation on Electrical Properties of Nucleic Acids for use in Molecular Electronics

Received 00th January 20xx, Accepted 00th January 20xx Hashem Mohammad^{a,#}, Busra Demir^{b,c,#}, Caglanaz Akin^{b,c}, Binquan Luan^d, Joshua Hihath^e, Ersin Emre Oren^{b,c,*}, M. P. Anantram^{a,*}

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Intercalating ds-DNA/RNA with small molecules can play an essential role in controlling the electron transmission probability for molecular electronics applications such as biosensors, singlemolecule transistors, and data storage. However, its applications are limited due to a lack of understanding the nature of intercalation and electron transport mechanisms. We addressed this long-standing problem by studying the effect of intercalation on both the molecular structure and charge transport along the nucleic acids using molecular dynamics simulations and firstprinciple calculations coupled with Green's function method, respectively. The study on anthraquinone and anthraquinoneneomycin conjugate intercalation into short nucleic acids reveals some universal features: 1) the intercalation affects the transmission by two mechanisms: a) inducing energy levels within the bandgap and b) shifting the location of the Fermi energy with respect to the molecular orbitals of the nucleic acid, 2) the effect of intercalation was found to be dependent on the redox state of the intercalator: while oxidized anthraguinone decreases, reduced anthraquinone increases the conductance, and 3) the sequence of intercalated nucleic acid further affects the transmission: lowering the AT-region length was found to enhance the electronic coupling of the intercalator with GC bases, hence yielding an increase of more than four times in conductance. We anticipate our study to inspire designing intercalator-nucleic acid complexes for potential use in molecular electronics via creating a multi-level gating effect.

Conceptual insights

Facilitated by the advance of synthetic biology, designer DNA with predetermined sequences can be readily synthesized for various applications (e.g., biosensors, single molecule transistors and DNA storage) in the nascent field of molecular electronics. However, the lack of control on charge transport significantly limits its practice in nanoelectronics. While experiments have shown that the conductivity of DNA can be adjusted via intercalation with organic molecules, the mechanism is still not established. In this work, using rigorous calculations, we addressed this problem by studying the effect of intercalation on both the molecular structure of and charge transport along the nucleic acids. We estimated the location of Fermi energy using partial charge transfer analysis. This helped us show that the conductance of DNA can be altered depending on the redox state of the intercalator, induced energy levels, and the location of the Fermi energy. These results establish the theoretical basis for intercalation-regulated conductance and thus can guide the design of novel intercalator-nucleic acid systems for use in future molecular electronic devices.

- ^a Department of Electrical Engineering, University of Washington, Seattle, WA, USA.
- ^b Bionanodesign Laboratory, Department of Biomedical Engineering, TOBB University of Economics and Technology, Ankara, Turkey.
- Operations of the second second
- ^d Computational Biological Center, IBM Thomas J. Watson Research, Yorktown Heights, NY 10598, USA.
- ^e Electrical and Computer Engineering Department, University of California Davis, Davis, CA, USA.
- # These authors contributed equally.
- * Corresponding authors; e-mails: eeoren@etu.edu.tr, anantmp@uw.edu

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Introduction

Electronic properties of DNA have attracted particular interest over the past 20 years¹ due to its long-range charge transport and self-assembly properties, making it a desirable nanoelectronics component. DNA consists of four main bases: guanine (G), cytosine (C), adenine (A), and thymine (T), and the stacking and pairing of these bases form a helical structure. The π - π interactions of the stacked bases lead to long-range charge transport. Simultaneously, due to its molecular recognition (exclusive base-pairing), sophisticated geometries can be precisely fabricated, overcoming the scaling challenges with solid-state/silicon-based electronics. Thus, the understanding of the electrical properties of DNA, either theoretically or

experimentally, becomes a key factor for future device developments.

The last decade has witnessed clear demonstrations of the sensitivity of short oligonucleotides' electrical conductance (5 nm range) to chemical modification of nucleotide,^{2–4} conformational change,^{5–8} and single base mismatch.⁹ The ability to tune and control these factors has wide implications in electronics, sensors,^{10–19} and sequencing.^{20–22}

Intercalation is an insertion of small molecules between bases of a nucleic acid structure, triggering structural changes such as local unwinding and elongation.²³ Small molecules called intercalators are often used as anticancer drugs to inhibit nucleic acid replication²⁴ or as fluorescent markers to visualize the structure.²⁵ The interaction between intercalators and nucleic acids has been studied both experimentally and computationally.^{26–31} Consequently, there has been a growing interest in controlling nucleic acid conductance by adding intercalators.

Recently, a graphene quantum dot intercalator in a 59-basepair long abasic-DNA was found to enhance the charge transfer rate.³² An 11-basepair DNA intercalated with coralyne showed three times larger conductance than the native DNA at negative biases.³³ However, other studies have shown a contrasting impact of intercalation on DNA conductance. Harashima et al.34 demonstrated that intercalation using ethidium bromide increased conductance by four times; they also showed that using Hoechst-33258 as a minor groove binder did not induce an increase in conductance on the same ds-DNA having 8 basepairs. On the other hand, an earlier experiment conducted by Wang et al.35 demonstrated that the same intercalator (ethidium bromide) decreased the conductance of 20 basepairs long ds-DNA. Recently, Xiang et al.36 reported that an anthraguinone (Ag) molecule directly attached to the end of the backbone of ds-DNA via an uracil (U) base could modify the conductance of a ds-DNA based on its redox state. In their experiment, they showed that the Aq moiety intercalated the complementary strand between the guanine bases, and the reduced state Aq yielded higher conductance, while the oxidized state Aq yielded lower conductance than the unmodified DNA. The experimental papers do not resolve the underlying mechanisms in a unified manner as exemplified from the literature. A recent modelling paper by Aggarwal et al.37 reported the effect of ethidium and daunomycin intercalators on the conductance of 8 and 12 base-pair long DNA strands. They used ab initio calculations and showed that upon intercalation, the conductance increases by one order of magnitude mainly due to reducing the twist angle between the intercalated base pairs, enhancing their structural alignment.

In this paper, we explore the fundamental pathways to alter the conductance of DNA upon intercalation, which is relevant to molecular electronics. We employ a combination of molecular dynamics (MD) simulations, density functional theory (DFT), and Green's function-based charge transport calculations. We focus on two different nucleic acids, the ds-DNA and the DNA:RNA hybrid, both intercalated with Aq. We study the impact of redox state, location, and quantity of intercalators in altering ds-DNA's electronic properties considering different

measurement scenarios. We report a new approach to analyze the effect of intercalations on the Fermi energy using partial charge transfer concept. Subsequently, we demonstrate that Aq intercalation in ds-DNA can increase or decrease the conductance, depending on the oxidation state, due to the following two reasons: 1) Intercalation decreases the bandgap energy by inducing new energy levels within the bandgap region of the bare ds-DNA, either close to the HOMO region (reduced Aq case) or in the middle of the bandgap (oxidized Aq case), and 2) intercalation shifts the Fermi energy with respect to the HOMO of the intercalated ds-DNA. We also report that the effects of Aq intercalation on both ds-DNA and DNA:RNA hybrids are alike.

We additionally model intercalation by the anthraquinoneneomycin conjugate (AqNEO) in DNA:RNA hybrids as the binding stability experimentally studied by Degtyareva *et al.*³⁸ Neomycin has been shown to increase the thermal stability of RNA duplexes³⁹ and also the binding affinity of the conjugated intercalator to DNA:RNA duplexes⁴⁰. We show that the neomycin part of the conjugate, acting as a groove binder, has a small effect on the conductance, which agrees with the literature results.^{34,35} We report that this is due to the indistinctive changes on molecular orbitals around HOMO region of DNA:RNA hybrid.

The paper is organized as follows: first, we introduce the cases under study and discuss the structural impact of intercalation (MD results). We follow it by energy levels analysis (quantum mechanics calculations). Then, we focus on the energy levels induced by the intercalators and their impact on transmission. Afterward, we study the effect of having two intercalators on altering electron transmission through DNA. After this focused analysis on the intercalator induced levels, we shift our attention to the overall conductance. Thus, we analyze the partial charge transfer rates to predict the Fermi energy for conductance comparisons. Finally, we study an additional case of a structure with a shorter AT region and illustrate key differences.

Results and Discussion

We study a family of three structures: 1) Aq (both oxidized and reduced states) intercalation into a 15 base-pair ds-DNA with the sequence 3'-G3A9G3-5'. 2) Aq and AqNEO intercalation into a DNA:RNA hybrid having the same sequence, with the RNA strand formed by the purines and the DNA formed by the pyrimidines. 3) Aq (both oxidized and reduced states) intercalation into a ds-DNA with a shorter AT region, having the sequence 3'-G3A3G3-5'. We first focus on 1 and 2 (Figure 1a) to understand the intrinsic effects of the intercalation on the strands. The 15 base-pair structures are expected to be comparatively resistive due to the high number of AT basepairs, creating a barrier for the charge to traverse the structure to the terminal guanines. Thus, as for the 3rd structure, we lowered the AT-region length from 9 to 3 base-pairs. As will be shown, the effect of intercalation on both ds-DNA and DNA:RNA hybrid is similar, hence moving forward in the transport calculations, the discussion will focus on the ds-DNA system.



Figure 1: Schematic representation of the system: a, Simulated intercalation cases for the 15 base-pair long structures. b, Oxidized and reduced states of the Aq molecule. c, Decoherence model for charge transport calculations.

The exact location of intercalation is difficult to identify experimentally as Aq and its derivatives can reside between any consecutive base-pairs depending on the molecular interactions between the intercalator molecule and the nucleic acids.^{41,42} To account for this, we consider intercalation between a) AT basepairs at 7 and 8 ($I_{7,8}$) and b) GC base-pairs at 13 and 14 ($I_{13,14}$) among other possibilities (Figure 1). We simulate ds-DNA with and without the intercalator using NAMD⁴³ with CHARMM 36 force field⁴⁴. We choose representative structures after more than 80 ns MD simulations (see SI methods). In both cases, oxidized and reduced state Aq are studied, called hereafter as Ox-Aq and Re-Aq for brevity. As for the DNA:RNA system, the hybrid structure is built using Amber NAB Tool,^{45,46} and then Ox-Aq is also intercalated at $I_{7,8}$ and $I_{13,14}$ (see SI methods). For this system, we additionally consider Aq conjugated with neomycin acting as a groove binder (AqNEO) to study its effect on charge transport. We use AMBER with ff99OL347,48 and Bsc149 force fields to simulate the DNA:RNA system and used the clustering method to choose a representative structure from 50 ns MD simulations. The DFT calculations are then carried out using Gaussian 16⁵⁰ software package with B3LYP/6-31G(d,p) basis set together with the polarizable continuum model. After obtaining the Hamiltonian from the DFT results, the charge transport calculations are performed for a system consisting of a contact-DNA-contact configuration (Figure 1c). The model is based on the Green's function method with probes to include decoherence (see SI for further details).

Structural Analysis

We choose representative structures from each MD simulation via the RMSD-based clustering analysis of the trajectories for the quantum mechanical calculations (Figures S1-S4). The structural changes in these representative structures upon intercalation are studied using the root mean square deviation (RMSD) calculations and structure dendrograms generated by RMSD matrices and the Agnes function (Figure S5), a hierarchical clustering algorithm.⁵¹ We find that the intercalation at $I_{13,14}$ position causes a smaller structural change compared to $I_{7,8}$ position. We also observe intercalation into the middle of the structure causes base flip-out and base sliding in DNA:RNA for both Ox-Aq- $I_{7,8}$ and AqNEO- $I_{7,8}$ cases (Figure S6). This is consistent with our expectation that intercalation-induced stress closer to the ends can be relaxed more easily. While it changes the total charge of the system, the π - π interaction of the DNA:RNA bases are modulated to a small extent compared to intercalation. Our simulations demonstrate an increase in the binding affinity of intercalator when conjugated with neomycin (Table S1). The structural comparison of Aq- and AqNEO- intercalated DNA:RNA hybrids show that the duplex structures are similar when intercalated at $I_{13,14}$ position (RMSD: 0.87 Å) however when intercalated at $I_{7,8}$ position (RMSD: 2.32 Å) the conjugated tail part causes relatively higher structural differences (Figure S5b).

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Figure 2: LUMO and HOMO plots both without and with Aq (ISO value is 0.02). a and b, LUMO and HOMO distribution of ds-DNA. c and d, LUMO and HOMO distribution of ds-DNA intercalated with Ox-Aq between 7th and 8th base pairs. e and f, LUMO and HOMO distribution of ds-DNA intercalated with Ox-Aq between 13th and 14th base pairs. g and h, LUMO and HOMO distribution of ds-DNA with same intercalation position with c and d using Re-Aq. i and j, LUMO and HOMO distribution of ds-DNA with same intercalation position with e and f using Re-Aq.

Quantum Mechanical Modeling



Figure 3: Energy levels and bandgap values for different intercalation cases of ds-DNA and DNA:RNA. a, Energy levels for ds-DNA system. Ox-Aq induces a LUMO into the bandgap of DNA, while Re-Aq induces both HOMO and LUMO which are close to energy levels of the DNA, respectively. b, Energy levels for DNA:RNA system with Ox-Aq and AqNEO cases. Both Ox-Aq and AqNEO induces LUMO into the bandgap of DNA:RNA.

An essential effect of Aq intercalation on the DNA is in the HOMO and LUMO spatial distribution. Without intercalation, the LUMO is localized at a single Thymine (Figure 2a), whereas its HOMO is delocalized on two guanines (Figure 2b). However, when Ox-Aq is introduced into the system, its effect is seen as adding a new unoccupied level (LUMO) localized at the Aq. This result is true for both the $I_{7,8}$ and $I_{13,14}$ cases (Figure 2c,e). As for the HOMO, it is still localized at the guanines (Figure 2d,f). However, in the Re-Aq case, both HOMO and LUMO are localized at the Aq (Figure 2g-j). Therefore, it adds two energy levels into the bandgap as opposed to Ox-Aq. The added levels for Re-Aq (Ox-Aq) near HOMO (LUMO) are fully occupied (empty). Hence, this is not like the conventional doping of semiconductors where p-doping (n-doping) has unoccupied (occupied) levels induced near the HOMO (LUMO) band. The bandgap value changes as the intercalator adds new energy levels into the DNA. In the Ox-Aq case, the LUMO induced by Aq is seen in the middle of the bandgap closer to the LUMO region of ds-DNA (Figure 3a). The bandgap is reduced by more than 1.2 eV, from 3.67 eV of bare DNA to 2.12 eV for $I_{7,8}$ and 2.45 eV for $I_{13,14}$. The Re-Aq induces two new levels in the bandgap region, a HOMO and a LUMO, which also decrease the bandgap compared with the bare DNA, to 2.97 eV and 2.86 eV for $I_{7,8}$ and $I_{13,14}$ intercalations, respectively. Similar trends are observed for DNA:RNA system (Figure S7). We observe the LUMO localizing on the intercalating molecule in both Aq and AqNEO cases. As for the HOMO, it remains localized on the guanines. The bandgap is reduced by 1.48 eV for the Aq case and by 1.66 eV with AqNEO (Figure 3b). We also notice that the incorporation of neomycin as a groove binding molecule does not significantly alter the molecular orbitals of the system in the HOMO region (AqNEO cases in Figure 3b). However, the LUMO region is shifted to lower energies for the AqNEO cases. The average shift in the induced energy between Aq and AqNEO cases is 175 meV. Thus, we find that the groove binder neomycin does not cause a significant effect on the molecular orbitals of the system or the intercalator induced energy levels. In the following section, we explore the transport calculation results for the different cases, specifically at the induced energy levels.

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transmission peak is ~14%. As for the Re-Aq, it induces an occupied level close to the HOMO vicinity. Here, the Re- $I_{7,8}$ has



Figure 4: Transmission plots for a, ds-DNA system in the HOMO region (stars: Re-Aq induced peaks). b, ds-DNA system in the Aq-induced region (stars: Ox-Aq induced peaks). c, DNA:RNA system in the Aq-induced region (stars: Aq and AqNEO induced peaks). The x-axis is energy because these results do not depend on the occupancy or Fermi factor of the electrons. Conduction plots for d, ds-DNA in the HOMO region, e, ds-DNA in Aq-induced region and f, DNA:RNA in Aq-induced region.

The direct comparisons between the original states of ds-DNA and DNA:RNA is discussed in prior references.^{52,53} In this work, we focus on the intercalator's effect on charge transport in the ds-DNA and DNA:RNA systems (Figure S8). The transmission is plotted as a function of energy ranging from -6 to 0 eV in Figures S9, S10 and S11. Figure 4 shows the transmission near the intercalator induced energy levels. The transmission increases at the Aq-induced energy levels by 16-34% compared to bare DNA (see transmission at the starred peaks in Figure 4). The Re-Aq cases ($I_{7,8}$ and $I_{13,14}$ in Figure 4a) show a ~16% increase, whereas $Ox-I_{7,8}$ and $Ox-I_{13,14}$ increase the transmission by 23% and 34%, respectively (Figure 4b). Similarly, the DNA:RNA system with AqNEO (Aq) exhibits the same trend (Figure 4c): the intercalator-induced peaks increase the transmission at their respective energies compared to the bare system. It is essential to mention that the induced peaks for Aq (or AqNEO) in its oxidized state are deep in the bandgap region of the no intercalated cases, -2.8 eV < Energy < -2.4 eV (Figure 4b,c and Figures S9 and S10). This range is more than 1.5 eV above the anticipated Fermi energy (E_f) location, which is assumed to be in the HOMO vicinity.

We expect that the closer the Aq-induced levels get to the high density of states (DOS) region, the higher the transmission becomes. This is due to the enhanced coupling of Aq to neighboring bases. The location of intercalation has a small effect on the induced energy levels as shown in Figure 3. For instance, changing the Ox-Aq location from $I_{7,8}$ to $I_{13,14}$ induces an energy shift of 245 meV, and the corresponding increase in

its HOMO energy level closer to the HOMO of Re- $I_{13,14}$ by 163 meV, and results in a ~25% increase in transmission peak (Figure 4). As for the DNA:RNA system with Aq, Figure 4c shows that by changing the location of intercalation from $I_{7,8}$ to $I_{13,14}$, the induced level shifts by 91 meV closer to the LUMO region, only increasing the transmission peak by 7%. On the other hand, in the neomycin conjugate (AqNEO) case, the induced level shifts by 89 meV away from LUMO. However, this opposite shift yields a change in transmission peak of less than 1%. We find that transmission is not affected by the addition of the groove-binder, which is in qualitative agreement with experiments^{34,35} that finds groove-binding molecules have an insignificant impact on DNA conductance.

Although we observe energy shifts and transmission peaks can quantitatively vary, the percentage of transmission increase to the no-intercalation cases is not significant. These results indicate that the spatial location of intercalation -alone- cannot be the reason behind the intercalation induced conductance differences reported in the literature. We also considered a hopping model, which supports this conclusion (Figures S12, S13 and S14). Thus, in the next section, we will look at the effect of having double intercalations. We also like to note that since the effect of intercalation yields similar trends regardless of the system (ds-DNA or DNA:RNA), the following sections will focus on the ds-DNA system.

Double Intercalation

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Multiple intercalations may occur in experiments depending on the intercalator molecule concentration and the length of the DNA strand.^{54,55} We intercalate two anthraquinone molecules into pre-stretched target positions corresponding to $I_{7,8}$ and $I_{13,14}$ positions. Following this, we minimize and equilibrate the system with MD simulations.

terminal case (Figure 6b), the Fermi energy is swept between the HOMO and LUMO energies using a gate electrode (Figure 6b). In the two-terminal case (Figure 8b) we estimate the Fermi energy location. We assumed that the electrodes are coupled to the DNA ends via thiol linker groups in both experiments, which is the common approach in STM-Break Junction



Figure 5: Double-Intercalation effect on Transport properties of the ds-DNA. a, the 2D DOS plot of double Re-Aq, b, the 2D DOS plot of double Ox-Aq, showing the energy levels distributions along the strand.

The transmission results show that in the Re-Aq case, a single transmission peak occurs near the HOMO region for double intercalators (TwoRe), which is ~83% higher than the single intercalation cases (Figure 4a). However, in the Ox-Aq case, the double-intercalation (TwoOx) yields two transmission peaks close in amplitude to the single intercalation cases (Figure 4b). To understand this, we look at the 2D DOS plot. The plot shows the distribution of electrons as a function of both space and energy. Figure 5a shows that the two intercalators induce two nearly degenerate levels near the HOMO region of the DNA for the Re-Aq case. Hence, having higher coupling with the neighboring bases of the DNA, they yield a large transmission peak. However, in the Ox-Aq case, the energy levels are deep in the bandgap region (dark blue area in Figure 5b). Therefore, they are residing in an almost isolated region with low coupling to the DNA bases. Hence, increasing the number of Re-Aq intercalators can yield higher transmission probability at the induced levels.

So far, we have focused our discussion on the Aq-induced levels. The calculations show that the induced energy levels inside the bandgap cause a transmission increase of less than 45% for this 15 base-pair long strand. At the same time, a higher percentage increase is reached with double-intercalators. Compared with literature, in which more than three times increase was reported, we deduce that the induced levels cannot be the only factor in increasing the DNA conductance. To probe other mechanisms that may influence the conductance, in the next section, we look at the overall conductance profile of the ds-DNA cases at different energy regions.

Conductance and Fermi Energy Location

In this section, we present results for two types of modelling scenarios: ds-DNA is connected in between two electrodes with and without a gate electrode system, respectively named as the three-terminal and the two-terminal scenarios. In the threesetups.^{56,57} This assumption provides us with the information that for small biases we operate in the linear response regime^{4,58,59} (where most of the potential drop occurs across the electrode-thiol linker junction). However, we have not explicitly included the thiol linkers or the electrodes due to the enormous computational complexity. More details are available in the SI about the transport calculations.

1) Three-Terminal Scenario (Fermi Energy Sweep)

Here, we assume a fixed small bias applied to the molecule while we sweep the Fermi energy *via* a third gate electrode and calculate the conductance. The conductance plot is shown in Figure 6a as a function of E_f , where we shifted the HOMOs of all molecules to match. If the Fermi energy is at the HOMO for each case (or HOMO plus a few hundred meV), it is apparent that the conductance is largest for oxidized Aq (Ox- $I_{13,14}$), followed by bare DNA and the two remaining oxidized cases. In contrast, the Re-Aq cases have the lowest conductance values. The reason behind this is that the HOMO of the Re-Aq cases is localized at the Aq, whereas it is localized at the DNA strand for Ox-Aq (see Figure 2). Therefore, we predict that the three-terminal experiments should result in this conductance trend when E_f is close to the HOMO of each structure.





In contrast to a sweeping the Fermi energy, in the next section we explore a broader approach by estimating the location of Fermi energy for each case using the principle of partial charge transfer.

2) Two-Terminal Scenario (Fermi Energy Location)

The second type of modelling scenario involves two-terminal setup (without a gate). Here, we do not know where the contact's Fermi energy lies a priori, but it is determined via the partial charge transfer method. In molecular junctions, it is known that the nature of the interface (gold electrode / ds-DNA) affects the Fermi energy of the contact. The adsorption of the molecule to the contacts causes both broadening of the molecular energy levels and partial charge transfer. Both the geometry and the orientation of the gold contacts may affect the amount of broadening and charge transfer within the molecular system under study. Including these effects in the calculations will, in principle, allow the evaluation of the Fermi energy position. This approach has been utilized previously on small molecules resulting in a wide range of Fermi energy locations extending from HOMO to LUMO.⁶⁰ Given this and the difficulty of including gold contacts for studying a molecule of more than a thousand atoms,⁶¹ we take an alternate approach proposed by Zahid and collaborators.⁶⁰ They suggested taking the Fermi energy rather as a fitting parameter for the calculations of molecular shifts due to partial charge transfer.

In the DNA-contact system, we expect a shift in the molecular energy levels with respect to the Fermi energy of the gold contacts. Partial charge transfer occurs from the contact to the DNA, shifting the MOs to higher energies. The E_f is in general closer to HOMO than to LUMO. Our approach here is to estimate E_f using the rate of change in energy as we vary the number of electrons in the system within the bandgap region. The higher the rate is, the higher the shift in the MOs, and the closer the HOMO gets to E_f . To achieve this, we first estimate the number of electrons in the bandgap region (Figure S15) by integrating the local density of states (LDOS) using the Green's function as:

$$N(E_i) = \iint LDOS(x, E) f(E - E_i) dx dE$$

= $\frac{1}{\pi} \iint - \operatorname{Im} \left[diag(G^r) \right] f(E - E_i) dx dE$ (1)

where *x* represents the location of the atoms in the system, and $f(E - E_i)$ is the Fermi-function, which describes the probability of electron occupancy at energy *E* when $E_f = E_i$. After obtaining *N* versus E_i , we calculate the derivative (dE_i/dN) , which gives us the average change in energy dE_i for a small change in electron number dN (Figure S16). The average change per one electron for each case is listed in Table 1. The values show that Ox-Aq cases have the lowest rates, by an average difference of 0.81 eV/electron from bare DNA. Re-Aq cases have a higher rate than the bare DNA by an average of 0.47 eV/electron. This trend can explain how partial charge transfer affects each case: when the same amount of partial charge is transferred from gold contact to DNA; 1) Ox-Aq has the lowest shift of molecular orbitals, which translates to E_f being the

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farthest from HOMO region. 2) Re-Aq, has the largest shift, making E_f the closest to HOMO. 3) Bare DNA is between Ox-Aq and Re-Aq cases.

Therefore, we can estimate the location of E_f by incrementing the amount of partial charge transfer (dN), using the following equation:

$$E_f = E_{f_0} - \left(\frac{dE_i}{dN}\right) \times dN \tag{2}$$

where E_{f_0} resembles Fermi energy of the electrode before making the contact-molecule junction, dE_i/dN is taken from Table 1, dN is a sweeping parameter, and E_f is the location of the Fermi energy after the MOs shift due to the partial charge transfer. We assumed different starting points for E_{f_0} , and extracted the conductance of each case at their respective E_f . Next, we compared the conductance values with the bare DNA case, and the ratios are plotted in Figure 7. We plotted the conductance ratio for each case as a function of partial charge transfer (dN) for E_{f_0} = -3.5 eV in Figure 7a (Figures S17 and S18 for -3.0 eV, -3.2 eV, -3.5 eV, and -4.0 eV). Notice how at dN > 0, conductance ratios for TwoRe and Re- $I_{13,14}$ become greater than 1, while Re- $I_{7,8}$ needs dN > 0.2 electron to reach ratios greater than 1. As for the Ox-Aq cases, the ratios are below 1 for dN > 0.

 Table 1: Average rate of change in energy with respect to number of electrons for ds-DNA

Molecule	dE_i/dN (eV/electron)	Change
TwoOx	2.61	-0.93
Ox-I _{7,8}	2.61	-0.93
Ox- <i>I</i> _{13,14}	2.97	-0.57
ds-DNA	3.54	0
TwoRe	3.75	0.21
Re- <i>I</i> _{7,8}	4.13	0.59
Re- <i>I</i> _{13,14}	4.16	0.62

Figure 7b summarizes the results for the different cases. The results are consistent for the different E_{f_0} cases and show a maximum increase in conductance of ~1.6 times for Re-Aq, with an average ratio higher than bare DNA. The Ox-Aq cases have average ratios of less than 1. These results are at least qualitatively consistent with the conductance measurements reported in the literature,³⁶ which show that $G(E_f^{\text{Re-Aq}}) > G(E_f^{\text{DNA}}) > G(E_f^{\text{Ox-Aq}})$; however, the structure in the reference is for a different Aq-based strand. An example graph is plotted in



Figure 7: Conductance ratio with respect to ds-DNA for the different intercalation cases. a, Conductance ratio as a function of partial charge transfer for E_{f_0} = -3.5 eV. The conductance is extracted at each E_f value based on Eq. 2 and compared with the conductance of ds-DNA without intercalator. The cut-off seen in some curves near 0.25 electron (Re-Aq cases) are because E_f has reached their respective HOMO which is set as the cut-off for incrementing *dN*. b, The maximum, minimum, and average values are taken for *dN* range going from 0.001 electron to 0.4 electron.

Figure 8a for dN= 0.2 eV and E_{f_0} = -3.5 eV values showing $G(E_f \text{ Re-Aq}) > G(E_f \text{DNA}) > G(E_f \text{Ox-Aq})$.

To summarize, the location of E_f for the DNA does impact the **a** $_{f} \times 10^{-11}$ **b**



Figure 8: a, Example of conductance plot as a function of Fermi energy for ds-DNA intercalation cases for a chosen dN = 0.2 eV and $E_{f_0} = -3.5$ eV values, where E_f is calculated using Eq. 2. b, two-terminal scenario.

conductance ratios between bare DNA and DNA + Aq. The present analysis shows that E_f of Re-Aq intercalation cases are closer to the HOMO region than E_f of bare DNA. As a result, the conductance can be as high as 1.6 times for DNA + TwoRe case (Figure 7b). In contrast, we find E_f to be farther away from HOMO in Ox-Aq cases, and the conductance are lower than bare DNA. These results indicates that there are two mechanisms are in play in increasing the conductance of Re-Aq intercalated DNA: 1) a Fermi energy shift, and 2) induced energy levels above the HOMO to help traverse the AT region.

Effect of AT Region Length

In this section, we study a strand with a shorter AT-region to investigate the effect of AT region length on the Aq-induced energy levels and the conductance. Here, we simulate the 3'-GGGAAAGGG-5' strand, with Aq intercalating at $I_{5,6}$, or $I_{7,8}$, and both (two-Aq). The main result is that the Re-Aq induced levels are now closer to the HOMO region of the ds-DNA, lowering the energy separation from 500 meV in 3'-G3A9G3-5' (see Figure 3a) to less than 70 meV in 3'-G3A3G3-5' (Figure S19). We attribute this result to the increase in coupling between the two GC regions as well as the intercalated Re-Aq (Figures S20, S21, S22 and S23).

Transport calculations in Figure S24 show that the increase in transmission at the Re-Aq induced levels (-4.75 eV < Energy < -4.8 eV) is now higher compared to the longer AT case (Figure 4). Re- $I_{5,6}$, has two transmission peaks with more than 3x increase and the two-Aq case having a peak of more than 5x higher transmission than the bare-DNA. The energy of the induced level shifts by 50 meV for the Re-Aq when going from Re- $I_{5,6}$, to Re- $I_{7,8}$, and the transmission changes by 30%. Whereas the variation in the Ox-Aq cases is 120 meV and the transmission changes by less than 2.5%.

We also carried out the Fermi energy location analysis: 1) Threeterminal scenario where E_f is changed with a gate, and 2) Two-Terminal scenario where we estimate E_f location based on partial charge transfer. For the first scenario, Figure S25 shows that the conductance for E_f at HOMO (or at a few hundred meV higher energies), the Ox-Aq cases are higher than bare DNA. Furthermore, Re- $I_{5,6}$ is only marginally greater than the bare DNA at HOMO while it decreases at higher energies, with the remaining two Re-Aq cases displaying lower conductance in general. Again, these findings are due to the localization of the HOMO at the Re-Aq, while HOMO of the oxidized cases is primarily localized on the DNA bases.

As for the second scenario, the calculated dE_i/dN values yield the same trend as the 3'-G3A9G3-5' structure (Table S3). By performing the same conductance ratio analysis, we get a maximum conductance increase of 2.2 for TwoRe, 4.0 for Re- $I_{5,6}$, and 3.0 for Re- $I_{7,8}$, with an overall average ratio greater than 1 (Figure 9). The ratio drops below 1.0 for different dN values depending on the intercalation location for the oxidized cases. For example, Figure 9a shows the conductance ratios for the E_{f_0} = -3.5 eV case (for other cases Figure S26 and S27). Here, TwoOx displays ratio less than 1.0 at dN > 0, Ox- $I_{7,8}$ drops below 1.0 at dN > 0.1 electron, and $Ox I_{5,6}$ drops below 1 at dN > 0.2electron. These results confirm that for the shorter AT-region strand, location of intercalation, number of intercalations, and partial charge transferred have an important effect on the conductance of the DNA. The general trend, however, is still G($E_f^{\text{Re-Aq}}$ > G(E_f^{DNA}) > G($E_f^{\text{Ox-Aq}}$).

In summary, the reduced state of intercalator increases the conductance more significantly as the AT region length decreases. In the shorter case, the intercalator-induced energy levels have lower energy separation from the HOMO level. Therefore, they help increase the coupling between the GCs on the two sides of the DNA. This finding is consistent for any amount of partial charge transfer. The Ox-Aq causes the DNA conductance to decrease, and the decrease is more sensitive to the amount of charge transfer and the intercalation location compared to the Re-Aq.



Figure 9: Conductance ratio with respect to ds-DNA for the different intercalation cases (3'-G3A3G3-5'). a, Conductance ratio as a function of partial charge transfer for $E_{f^{\circ}}=-3.5$ eV. The conductance is extracted at each E_f value based on Eq. 2 and compared with the conductance of ds-DNA without intercalator. The cut-off seen in some curves near 0.2 electron (Re-Aq cases) are because E_f has reached their respective HOMO which is set as the cutoff for incrementing dN. b, The maximum, minimum, and average values are taken for dN range going from 0.001 electron to 0.31 electron (see SI for more E_{f_0} cases).

Conclusions

In this paper, we investigated the effect of Aq-intercalation on the conductance of nucleic acids, ds-DNA and DNA:RNA hybrid, through a combination of molecular dynamics, *ab-initio* simulations, and charge transport calculations. We find the intercalation to yield similar effects on the DNA:RNA hybrid

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system. The DNA:RNA + AqNEO analysis shows that the groove binding molecule increases the stability of the molecular structure while having a small effect on the molecular energy levels and conductance compared to the intercalators. Thus, an important finding of this work that one can use a groove binder conjugated to the intercalator to increase the system's overall structural stability^{39,40} without affecting the electronic properties induced by the intercalator, a crucial feature for the thermal stability of future DNA based molecular devices. We showed that intercalation can modulate the conductance of ds-DNA. One can control this by using parameters such as the redox state of the intercalator, the location of Fermi energy, and the length of the strand.

Our results show that although Aq induces energy levels in the bandgap of the ds-DNA, this is not like the conventional doping in semiconductors. The added levels for Re-Aq (Ox-Aq) near HOMO (LUMO) are occupied (empty). Therefore, the electrons must travel through these induced energy levels to traverse the structure, as they do not increase the number of charge carriers in the system. In addition to the induced energy levels in the bandgap, the contact's Fermi energy with respect to the molecule modifies the conductance. The Fermi energy analysis shows that Aq can either increase or decrease DNA conductance based on its redox state. The Re-Aq was assessed to have the Fermi energy closest to the HOMO region, with Ox-Ag having it the farthest away from HOMO. Hence, yielding the relation $G(E_f^{Re-Aq}) > G(E_f^{DNA}) > G(E_f^{Ox-Aq})$ with maximum conductance 1.6 times higher (TwoRe) than the bare DNA. This relation agrees with the experiment conducted,³⁶ where they have shown that the DNA conductance can increase or decrease based on the state of the anthraquinone. Furthermore, lowering the AT-region length from 9 base pairs to 3 increases the impact of the Re-Aq, yielding a maximum conductance rise of 2.5 times for two intercalators and more than 4.0 times increase for a single intercalator. This means, increasing the number of intercalators does not always yield an enhanced conductance, especially for shorter strands. Our results also show that the experimental setup may alter the measured conductance. The three-terminal case, where the gate voltage is swept, the Ox-Aq intercalation yields higher conductance than the Re-Aq intercalation case, when the relative Fermi energy locations with respect to the HOMO levels are the same. Overall, we demonstrated that one can modulate the conductance, by altering the number of intercalators, the ATregions length, and/or the experimental setup. A possible application for intercalation in addition to modulating nucleic acid conductance arises from utilizing multiple intercalators with different redox potentials (such as Ethidium Bromide), which in principle can help to create a multi-level gating effect for the same strand.

Conflicts of interest

There are no conflicts to declare.

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