Molecular Dynamics Simulation of Melting of the DNA Duplex with Silver-Mediated Cytosine–Cytosine Base Pair

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Abstract: Metal-mediated base pairs in DNA double helix molecules open up broad opportunities for biosensors based on DNA clusters with silver due to their low toxicity and applicability in drug design. Despite intensive experimental and computational research, molecular mechanisms of stabilization of a double helix by silver-mediated base pairs are mainly unknown. We conducted all-atom molecular dynamics simulations of a dodecameric DNA double helix (sequence 5′-TAGGTCAATACT-3′-3′ATCCACTTATGA-5′) with either cytosine–cytosine or cytosine–Ag⁺–cytosine mismatch in the center of the duplex. We extended the previously proposed set of interaction parameters for a silver ion in the silver-mediated pair in order to allow for its dissociation. With this new potential, we studied how the addition of a silver ion could stabilize a DNA double helix containing a single cytosine–cytosine mismatch. In particular, we found out that the helix with cytosine–Ag⁺–cytosine mismatch has a greater melting temperature than the helix with cytosine–cytosine one. This stabilization effect of the silver ion is in qualitative agreement with experimental data. The central region of the duplex with cytosine–Ag⁺–cytosine mismatch (unlike with cytosine–cytosine mismatch) is stable enough to prevent bubble formation at moderate temperatures during melting. The results of this simulation can be used to devise novel metal-mediated DNA structures.

Keywords: metal-base pair; silver-mediated DNA; melting temperature; molecular dynamics

1. Introduction

DNA double helix is often used in nanotechnology for a wide range of applications. One promising example is the extensive search for alternatives for hydrogen bonds (H-bonds) between complementary nitrogenous bases. An attractive approach is to create a base pair with metal cations that have a high affinity for nitrogenous bases. This approach is promising for the creation of DNA-based nanomaterials with improved thermal, mechanical and chemical properties, as well as with new functionalities [1–4]. A DNA double helix in which nucleotide bases are paired by a metal ion instead of hydrogen bonds in Watson–Crick base pairs is called metalized DNA (M-DNA) [5]. Such systems are of interest due to their potential use as sensitive and specific metal sensors and biosensors [6,7]. As a rule, the inclusion of non-canonical base pairs in homochiral or heterochiral DNA leads to a decrease in the double helix stability [8]. Such decrease depends to a large extent on the specific structure of the non-canonical base pairs, their nature, and the sequence in which they are included. In 2009, Ono and Miyake [9] described two structures with metal-mediated pairs, in particular, cytosine–Ag⁺–cytosine (C:Ag:C). Their research has opened wide opportunities for DNA nanotechnology, nanoelectronics and DNA-origami due to the high specificity of silver binding [10,11]. Silver ions have attracted special attention due to their low toxicity and applicability in drug design [12,13]. As a result, numerous studies on artificial DNA containing non-canonical base pairs with silver ions were reported [14–17]. In natural DNA, silver-mediated pairing has been explored in canonical double helices with base pair mismatches [9] and in strands without canonical pairs [18–21]. High chemical
and thermal stability of the C:Ag:C base pair was demonstrated [22–26], and the role of pH in the stability of the C:Ag:C pair was studied [27]. The effect of silver on the melting temperature of DNA double helix with cytosine–cytosine (C:C) inclusions as a complementary pair has been experimentally observed [28,29]. The melting point of an oligonucleotide with C:C mismatch is increased in the presence of silver ions in solution.

To date, a number of theoretical models of base-pairing dynamics in DNA of various degrees of detailization have been developed. These models stimulate significant progress in the field of DNA melting [30,31]. The proposed models vary from the mesoscopic level (models based on the Peyrard–Bishop (PB) model [32,33]) to the coarse-grained model (CG) with several beads per nucleotide [34]. Mesoscopic models such as the PB model and its variations use a set of effective potentials to describe the main interactions in the DNA double helix, namely the hydrogen bonds and stacking interactions [35–38]. Using the mesoscopic models, the influence of single, double, and triple mismatches on duplex stability was investigated. It was found that many combinations of multiple mismatches are surprisingly stable [39]. Melting points of sequences containing metal-mediated C:C and thymine–thymine pairs were calculated [40]. The use of a CG model, where only monovalent and divalent ions are included, allows one to reproduce experimental melting temperatures for oligonucleotides [41]. In many mesoscopic and CG models, the effective non-linear potential, Lennard–Jones potential [42–44] or Morse potential [45–47] are used to mimic hydrogen bonding to describe the melting processes.

DNA melting has also been studied by means of all-atom molecular dynamics (MD) simulation [48–50]. However, the number of all-atom MD simulation studies of DNA with metal-mediated pairs is very limited. The shortest possible Ag⁺-mediated cytosine homobase duplex, a tetramer formed by Ag⁺ bridges between two DNA strands, has previously been studied [16]. The Lopez-Acevedo group reported a study of Ag⁺-paired cytosine homobase DNA strands using a combination of quantum and classical MD calculations [51]. They showed that a parallel double helix has high stability, whereas an antiparallel double helix untwists and loses its helical structure. Cytosines are paired in a transoid conformation in a parallel double helix, while in an antiparallel double helix, they are in a cisoid conformation (see Figure 1a,b). The bond in the C:Ag:C pair in the cisoid conformation is weaker than in the transoid one due to an additional hydrogen bond formed between the oxygen of one cytosine and the amino group of the complementary cytosine [51–53]. The approach developed in the works of Lopez-Acevedo et al. [51] was successfully used in the MD simulation of silver-mediated base pairing in parallel-stranded DNA [29]. However, the MD simulation of the destruction of the C:Ag:C pair is challenging to date. The potentials are usually used to describe the bond of a silver ion with the nitrogen of cytosine as a covalent bond. MD’s description of a covalent bond is based on a harmonic potential that does not provide a possibility for bond breaking. Therefore, a different potential has to be used to describe the destruction of the C:Ag:C pair. One possibility is to use the Morse potential, which explicitly includes the effects of bond breaking and the existence of unbound states.

In this article, we propose to use the Morse potential to describe the bond of a silver ion with cytosine by analogy with a hydrogen bond in some mesoscopic models. We simulated an antiparallel double-stranded DNA helix with C:Ag:C mismatch. We fitted parameters of the Morse potential that reproduced the process of melting a pair mediated by silver. We obtained melting curves and estimated melting temperatures. We believe that melting temperature studies of M-DNA will certainly enhance our knowledge about DNA double helix properties associated with metal-mediated base pairs.
2. Materials and Methods

Double-stranded DNA helix (12 base pairs) with the sequence 5'-TAGTCAATAC-3' - 3'-ATCCACTATGA-5' with a mismatch of complementarity in the sixth position, as C:C was considered. The initial structure was generated using the Avogadro program [54].

Initially, the DNA double helix was placed in a cubic box so that the minimum distance between the helix and the edges of the box exceeded 1 nm. Then, water molecules were added to the system. We used the TIP3P water model. To neutralize the charge of the system, 22 Na$^+$ ions were added to the calculation cell. Calculations were carried out in a periodic cell using stochastic dynamics as a thermostat. Equations of motion were integrated using a leap-frog version of the Verlet algorithm. The integration step was 2 fs. Constant temperature was maintained using a stochastic thermostat, and constant pressure was maintained using a Parrinello–Rahman barostat with a time constant of 2 ps. Bonds with hydrogen atoms were constrained using the LINCS algorithm. The cutoff radii for Coulomb and van der Waals interactions were taken to be 15 Å. PME method was used to consider long-range Coulomb interactions. For MD simulation, the Gromacs 2023.1 software package [55] was used.

The parameters of potentials are a combination of the Amber99-parmbsc1 force field [56] and original parameters for Ag$^+$ and cytosine interactions. Parameters of interaction between cytosine and a silver ion were taken from the literature, with some modifications as follows. Partial charges of atoms on cytosine interacting with silver ions have been corrected according to the paper [51]. The 12-6 Lennard-Jones (LJ) model was used for van der Waals interactions. The values of non-bonded interaction parameters for

![Figure 1. Structural variations for the C:Ag:C base pair: (a) transoid and (b) cisoid conformations. N3* defines the N3 atom in complementary base. DNA double helix: (c) with pure C:C mismatch and (d) with C:Ag:C formed during the classical MD. A snapshot of the double helix at t = 100 ns is shown. (e) The fitted Morse potential curve. The red point corresponds to the equilibrium value r$_{Ag-N3}$ in the MD simulation at a temperature of 300 K. (f) The histogram (population in percentage) of the distance between silver ion and cytosine nitrogen r$_{Ag-N3}$ at temperature 300 K. Histogram was constructed using 75 bins between 2.05 and 2.35.](image-url)
the LJ potential for Ag$^+$ ($\sigma = 2.731 \ \text{Å}$ and $\varepsilon = 0.16316922 \ \text{kJ/mol}$) and the partial charge of a silver ion ($q_{\text{Ag}^+} = 0.38272$) were taken from [57], which previously were validated for silver-mediated base pairing in double-stranded DNA in [29,51].

To simulate the bond dissociation of the silver ion from the N3 atom of cytosine, we chose the Morse potential. The Morse potential explicitly includes bond-breaking effects such as the existence of unbound states:

$$V(r) = D_e \left(1 - e^{-\alpha(r-r_e)}\right)^2,$$

where $r$ is the distance between N3 and Ag atoms, $r_e$ is the equilibrium bond distance, $D_e$ is the depth of the well, and $\alpha$ controls the “width” of the potential. The equilibrium bond distance $r_e = 2.14 \ \text{Å}$ is known from quantum mechanical calculations and has been successfully used in MD simulations of both transoid and cisoid pairs C:Ag:C [29,51]. The parameters of the Morse potential $\alpha$ and $D_e$ were obtained as a result of an optimization procedure described below.

The optimization procedure of the Morse potential parameters consisted of two steps. At first, we determined the value of the parameter $\alpha$. The idea is to keep the geometry of the C:Ag:C pair determined from quantum mechanical calculations [51], in particular, by fitting the equilibrium value of the angle N3-Ag$^+\cdot$N3$^*$ to one derived by the DFT method. To carry out this procedure, we set the parameter $D_e = 200 \ \text{kJ/mol}$, which exceeds the energy of the hydrogen bond. Such a large value of $D_e$ is required to prevent the C:Ag:C pair from spontaneous dissociation upon thermal oscillations. The initial value of parameter $\alpha$ was chosen so that the “width” of the Morse potential was similar to the “width” of the harmonic potential from [51].

Authors of ref. [51] introduced harmonic bonded potential as well as angular and dihedral ones in order to keep the angle N3-Ag$^+\cdot$N3$^*$ close to quantum chemical values. We used Morse potential rather than these ones and, therefore, had to optimize the “width” of the potential well.

We selected a range of values of parameter $\alpha$. For each value $\alpha$, we conducted an MD simulation at a temperature of 300 K with a length of 10 ns and determined the equilibrium value of the valent angle N3-Ag$^+\cdot$N3$^*$. It turned out that $\alpha = 2.0 \ \text{Å}^{-1}$ gives the closest to quantum chemical values [51].

The next step was the parameter $D_e$ optimization. The parameter $D_e$ is connected with dissociation energy. We conducted a series of MD simulations heating the system as described above. We reduced the parameter $D_e$ for each simulation. We calculated the melting curves at each value of the parameter $D_e$. The optimized value is $D_e = 150 \ \text{kJ/mol}$. Further decrease was accompanied by the melting of the double helix with the silver ion at the same temperatures as without a silver ion. Figure 1e shows the fitted curve of the potential Morse with optimized parameters.

We conducted two types of MD simulations of a DNA double helix: equilibrium dynamics at 300 K and simulation of melting. Both simulations began with a relaxation stage followed by a production run. The relaxation stage consisted of energy minimization using the conjugate gradient method and MD simulations under conditions of constant volume (NVT ensemble) and constant pressure (NPT ensemble) at a constant pressure of 1.0 bar and temperature of 300 K for 10 ns. After relaxation, the dynamics of the system were calculated for 100 ns at a temperature of 300 K. The simulation of melting started from the configuration of the system taken from the equilibrium dynamics at 300 K. The system was heated every 20 ns with a temperature step of 10 K. MD simulations were run at temperatures ranging between 300 K and 460 K.

Based on the obtained MD trajectories, the hydrogen bonds (H-bonds) were calculated. In detecting H-bonds, the well-known geometric criterion was used: H-bond between base pairs exists if the distance between two electronegative atoms is less than 3.5 Å and the H-bond angle is less than 30° [58].
3. Results

3.1. Dynamics and Stability of the DNA Double Helix with Either C:C or C:Ag:C Mismatch

Since we offer a new potential, we examined the stability of the dodecamer double helix in the presence of a silver ion. Therefore, after the equilibration process was completed, an MD simulation was carried out under constant temperature (300 K) and pressure (NPT) for 100 ns, of which the last 50 ns were used for analysis. The original potential was used to describe the interaction of the silver ion with cytosine atoms. These changes do not affect any other interaction parameters and should not affect the structure of the double helix outside the C:C pair mismatch. The double helix in the presence of the silver ion at 300 K was stable and had an average number of hydrogen bonds even greater than in the simulation with a C:C mismatch in the absence of a silver ion. The presence of silver ions in a system with a C:C mismatch leads to an increase in the stability of the double helix. Figure 1c,d shows typical snapshots at the end of the simulation in the absence and presence of the silver ion.

Figure 1f shows a histogram of the $r_{\text{Ag-N3}}$ distance between the silver ion and cytosine nitrogen. One can see that the MD simulation equilibrium distance $r_{\text{Ag-N3}} = 2.18 \pm 0.003$ Å is scattered in close proximity around the equilibrium distance between the silver ion and cytosine nitrogen, in agreement with the literature data [51]. We also compared this equilibrium distance with the potential minimum (Figure 1e). The equilibrium value is close to the potential minimum. The displacement is due to the presence of other highly charged cytosine atoms in the immediate vicinity of the nitrogen–silver bond.

The increase in the stability of the C:C pair in the presence of the silver ion is well characterized by the $r_{\text{N3-N3}}$ distance between the N3 nitrogen atoms of cytosines in the silver-mediated pair. Figure 2a shows the $r_{\text{N3-N3}}$ distance. The average distance between N3 atoms of cytosines in the absence of silver ions at temperature 300 K the N3-N3 distance is $r_{\text{N3-N3}} = 3.97 \pm 0.27$ Å. In the presence of the silver ion, the N3-N3 distance is $r_{\text{N3-N3}} = 4.31 \pm 0.07$ Å, which is higher than in the C:C pair, but the C:Ag:C pair is more stable, as indicated by the standard deviation from the average. In this case, we observed large deformations of the C:C pair and the temporary formation of a hydrogen bond between the NH$_2$ group of one cytosine and the N3 atom of another cytosine. However, such a hydrogen bond turns out to be unstable.

We also calculated the local base pair parameters of the DNA double helix in the absence and presence of the silver ion. The local base pair parameters for the C:C pair are significantly different compared to adjacent pairs based on the absence of a silver ion. Local base pair parameters such as opening, shear and stretch turned out to be the most noticeable (see Figure 3a–c).

In the presence of the silver ion, the profile of shear and stretch parameters along the helix is almost flat (see Figure 3d,e). This indicates greater stability in the silver-mediated C:C pair. The non-typical value of opening is evidence of the cisoid conformation C:C pair and reflects the particular features of this non-canonical pair.

The structure of the silver-mediated pair of cytosines in the transoid and cisoid conformations can be different, as was demonstrated in [51] by the DFT method. In our case, the C:C pair is in the cisoid conformation, as in antiparallel helices. The initial cisoid conformation does not transform into a transoid one during simulation. The angles formed between the silver ion and cytosines in the coordination bond are shown in Table 1. A slight shift in the equilibrium angles is observed, indicating a displacement of the silver ion towards the cytosine oxygens.
Figure 2. The distance $r_{N3}$ between nitrogen atoms N3 of cytosines in the absence of a silver ion (thick gray line) and in the presence of the silver ion (thin black line) reflects the stability of the C:C and C:Ag:C pairs: (a) A 50 ns interval at the end of the analyzed time interval for the temperature 300 K. (b–f) A 20 ns time interval for the temperatures 320 K (b), 340 K (c), 360 K (d), 380 K (e), 400 K (f).

Figure 3. Local base pair parameters: shear, stretch and opening for case of absence (a–c) and presence (d–f) of the silver ion.
Table 1. Angles formed by the coordination bond between the silver ion and nitrogen N3 atom of cytosine. Standard atom notations for nucleotide bases were used.

<table>
<thead>
<tr>
<th>Angle</th>
<th>$\theta$, (deg)</th>
</tr>
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<tbody>
<tr>
<td>N3-Ag$^+$-N3$^+$</td>
<td>163 ± 8</td>
</tr>
<tr>
<td>Ag$^+$-N3-C2</td>
<td>103 ± 7</td>
</tr>
<tr>
<td>Ag$^+$-N3-C4</td>
<td>138 ± 7</td>
</tr>
</tbody>
</table>

3.2. Melting of the DNA Double Helix with Either C:C or C:Ag:C Mismatch

We performed an MD simulation of DNA heating in the range from 300 to 400 K with a temperature step of 10 K. With increasing temperature, the C:C mismatch distance fluctuates strongly, which is often accompanied by the opening of a C:C pair. Figure 2 shows the $r_{N3-N3}$ for some temperatures in the range from 300 to 400 K. At a temperature of 400 K, the C:C pair is already completely destroyed (Figure 2f). In the presence of the silver ion at a temperature of 400 K, the C:Ag:C pair is stable. The destruction of this pair coincides with the melting of the double helix at 410 K.

The maximum number of hydrogen bonds in the complementary base pairs in the sequence under consideration is 25. However, during the course of MD simulation, a smaller number of hydrogen pairs is observed as a result of base pairs fraying, a known process in MD simulations. When a DNA double helix with a C:C pair without a silver ion is heated, melting begins from the terminal base pairs of the double helix and from the C:C pair. Without a stable connection between these cytosines, the double helix forms a bubble, and three melting hot spots form in this case: two terminal pairs and the C:C mismatch. With the silver ion between two cytosines, we observe a more stable double helix, and melting occurs only from the terminal base pairs. The melting curves of the DNA double helix in the presence and absence of a silver ion are presented in Figure 4. It can be seen that in the presence of the silver ion, the melting curve is shifted towards higher temperatures, just as observed in the experiment [28].

![Figure 4](image-url)  
*Figure 4. The presence of the silver ion shifts the melting curve towards higher temperatures. The temperature dependence of the number of hydrogen bonds $i_{hb}$ without (circles) and with (triangles) silver ion is presented.*

Table 2 shows the melting temperatures of the DNA double helix in the absence and presence of the silver ion. As usual for MD simulations, we obtain the higher absolute values of melting temperatures in both cases (due to the superheating, see Discussion); however, we are interested in the relative effect—the melting temperature shift due to the
silver ion presence. This shift, in our case, is 12 K, which is close to the experimental value of 7.5 K. When the DNA melts, the metal-mediated pair breaks down, and the silver ion is released into the solution. The metal-mediated pair C:Ag:C is thermally more stable than the hydrogen bonds of complementary bases, but in this case, it has one silver-mediated base pair, which is also subject to thermal destruction, as observed in the experiment [28].

Table 2. The melting temperatures for double helix in the absence and the presence of the silver ion in comparison with experimental data. The bottom row shows the shift in melting temperature due to the presence of silver ions.

<table>
<thead>
<tr>
<th>Silver Ion</th>
<th>Tm Experiment $^\dagger$, K</th>
<th>Tm MD, K</th>
</tr>
</thead>
<tbody>
<tr>
<td>absence</td>
<td>299.5</td>
<td>392</td>
</tr>
<tr>
<td>presence</td>
<td>307</td>
<td>404</td>
</tr>
<tr>
<td>temperature shift $\Delta$Tm</td>
<td>7.5</td>
<td>12</td>
</tr>
</tbody>
</table>

$^\dagger$ Experimental results were taken from [28] for the same sequence as studied in this work.

4. Discussion

We took this sequence as a robust object for potential validation since its experimentally found melting point is well known in both the absence and the presence of silver ions [28]. We chose the standard antiparallel double helix since, for such objects, the MD approach is well-validated.

By introducing bond potentials to describe the coordination bond between the silver ion and cytosine, we were able to directly model the dynamics of coordinated bond breaking, which was previously impossible while treating these bonds as covalent. The main feature of our potential in comparison with early works is that the structure of the C:Ag:C pair is not fixed near the equilibrium position but can fluctuate significantly near it up to the destruction of the pair with increasing temperature. Our calculations show that the structure of the pair corresponds to the early work [51].

The discrepancies in the absolute values of melting temperatures between experiments [28] and MD simulations are common and attributed to the methodology used to simulate the nanoscale samples [59]. In MD simulations, the melting point is determined by “slow” heating; thus, the rate of temperature change compared to a real experiment is very high and can exceed the experimental value by several orders of magnitude. Since the transition does not occur instantly but requires a small (on an experimental scale) time, a shift is observed. This effect can be corrected by using Monte Carlo simulations instead of molecular dynamics, but this requires more expensive calculations. In addition, the melting temperature discrepancy is affected by the size of the simulation cell and the thermostats used. In molecular dynamics, we are dealing with a small cell, nanometers in size. In this case, the temperature change occurs instantly throughout the entire volume. In the experiment, the measured cell is macroscopic, and temperature changes inside it at the molecular level occur non-uniformly. Thus, we believe that the discrepancy between the melting temperatures in the real experiment and our results is not a disadvantage. We obtained a qualitative agreement of the melting temperature shift in the presence of a silver ion compared to the case when a silver ion is absent. In this case, we observe a satisfactory shift in the melting temperature, comparable with the experimental results.

5. Conclusions

We proposed and verified the Morse potential for the silver-mediated base pair bond that can describe the destruction of the coordination bond between cytosine and a silver ion. The verified potential parameters allowed us to reproduce the process of melting of the DNA double helix with a silver-mediated pair and the destruction of the coordination bond between the silver ion and the cytosine base with increasing temperature. The results of the simulations are in good agreement with experimental data on the melting temperature shift of the DNA double helix with a C:C mismatch in the absence and the presence of a silver ion. Since the B-form double helix is a basic building block not only in the genetic code of
living organisms but also in DNA nanotechnology, metal-mediated double helices may be used for self-assembly dynamic DNA nanomachines. Our results open new possibilities for the development of new metal-mediated 3D DNA materials with increased resistance to thermal, chemical, and possibly enzymatic degradation due to the additional stability provided by silver-mediated base pairs.

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**Data Availability Statement:** Data are contained within the article.

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**References**


