

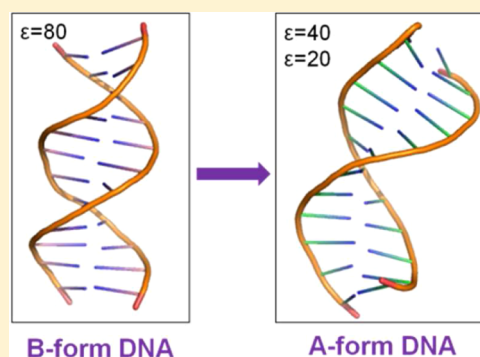
Conformational Preferences of DNA in Reduced Dielectric Environments

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S Supporting Information

ABSTRACT: The effect of reduced dielectric environments on the conformational sampling of DNA was examined through molecular dynamics simulations. Different dielectric environments were used to model one aspect of cellular environments. Implicit solvent based on the Generalized Born methodology was used to reflect different dielectric environments in the simulations. The simulation results show a tendency of DNA structures to favor noncanonical A-like conformations rather than canonical A- and B-forms as a result of the reduced dielectric environments. The results suggest that the reduced dielectric response in cellular environments may be sufficient to enhance the sampling of A-like DNA structures compared to dilute solvent conditions.



DNA is an essential biomolecule due to its role in numerous biological processes. Its structure, flexible and sensitive to environmental conditions, is one of the most crucial physical determinants of its biological roles. The major conformations of DNA, B-form and A-form, are well-known from in vitro studies; the B-form is the major conformation in solution¹ while the A-form is seen for GC-rich sequences under low-humidity environments and, sometimes, when bound to other biomolecules.² The structural preferences and transitions of DNA between these two forms have been studied extensively for many years not only to gain insight into the structure–function relationship of DNA, but also because DNA is a fascinating biophysical model system that exquisitely balances electrostatic and solvent interactions.^{3–27} Experimental studies on DNA structure have shown that B- to A-form transitions occur under conditions of “low water activity”, for example, when ethanol is introduced as a cosolvent^{4,9,12,14} and also when salt is added to solution which bridges the phosphate groups in the major groove of A-DNA.^{6,11,15} These findings have also been reproduced by computational approaches, especially by molecular dynamics (MD) simulation techniques for the exploration of these conformational preferences in atomistic detail.^{16–27} These MD studies showed that DNA maintained its B-DNA form in aqueous solvent, while A-DNA was stabilized in solutions containing explicit salt ions or cosolvents such as ethanol as expected from the experiments.^{16,18,21–25} The effect of salt on DNA structure is easily rationalized by increased electrostatic screening that allows subsequent phosphate groups along the backbone to come closer to each other so that A-DNA can be formed. It is less clear, however, whether cosolvents affect DNA structure through specific molecular

interactions or via a more general physical effect such as an altered dielectric response of the environment.

Reduced dielectric environments are also interesting in the context of understanding DNA structure in the cellular context. Cellular environments are highly crowded systems with high concentrations of proteins, nucleic acids, and numerous cosolvents. It has been reported that 20–30% of the cellular volume is occupied by macromolecules.²⁸ Previous studies have shown that the effect of such crowding on protein structure and dynamics can be significant.²⁹ Crowded environments alter the balance between enthalpic and entropic contributions to the conformational free energy either directly through macromolecular interactions or indirectly by modifying solvent properties.^{29–42} Essentially, the effect of crowding is threefold: (1) volume exclusion by surrounding macromolecules resulting in an entropic penalty for assuming more expanded states;^{30,39} (2) reduced dielectric response of the environment due to the displacement of 20–30% of the water with less polar macromolecules^{43–48} and due to slowed dynamics of the water itself;^{33,49} (3) specific interactions between different macromolecules. Therefore, studying the effect of reduced dielectric environments on DNA structure addresses one major consequence of cellular crowding. Such an approach neglects the effect of specific interactions between DNA and protein crowder molecules as well as the volume-exclusion effect, but it allows for more fundamental insight about how DNA structure is altered by different environments.

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Table 1. Helicoidal Parameters for the GC-Rich Dodecamer Compared to Experimental Results^a

	X-ray ^d	canonical ^b		MD simulations ^c		
		A-DNA	B-DNA	$\epsilon = 20$	$\epsilon = 40$	$\epsilon = 80$
propeller (deg)	−12.49 (2.0)	−9.00 (7.1)	−11.72 (5.8)	2.16 (15.5)	−2.85 (12.0)	−7.26 (14.1)
slide (Å)	−1.71 (0.4)	−1.67 (0.4)	0.17 (0.7)	−1.48 (0.8)	−1.23 (0.9)	−0.52 (1.4)
twist (deg)	29.59 (3.5)	30.24 (4.0)	34.81 (5.7)	27.50 (10.0)	29.70 (7.3)	36.27 (17.7)
X-displacement (Å)	−5.01 (1.1)	−4.54 (1.3)	−0.13 (1.1)	−2.68 (3.6)	−2.80 (2.9)	−1.19 (2.5)
helical rise (Å)	2.66 (0.6)	2.77 (0.5)	3.25 (0.2)	3.42 (1.0)	3.36 (0.8)	3.36 (0.8)
inclination (deg)	20.71 (11.4)	16.00 (10.7)	4.01 (6.7)	0.45 (20.7)	3.73 (15.1)	5.43 (16.4)

^aAll values are averaged over nonterminal base pairs with standard deviations given in parentheses. ^bAverages over the A-form structures 3V9D, 3QK4, 2B1B, 1ZEX, 1ZEY, 1ZF1, 1ZF6, 1ZF8, 1ZF9, 1ZFA and the B-form structures 2M2C, 4AGZ, 4AH0, 4AH1, 3U05, 3U08, 1VTJ, 3U2N, 3OIE, 3BSE. ^cAverages over snapshots at 300 K from replica exchange MD simulations. ^dPDB ID code: 399D.

Table 2. Helicoidal Parameters for the Drew-Dickerson Dodecamer Compared to Experimental Results^a

	X-ray ^d	canonical ^b		MD simulations ^c		
		A-DNA	B-DNA	$\epsilon = 20$	$\epsilon = 40$	$\epsilon = 80$
propeller (deg)	−13.34 (5.9)	−9.00 (7.1)	−11.72 (5.8)	−6.93 (12.5)	−10.67 (12.7)	−11.32 (13.7)
slide (Å)	0.07 (0.5)	−1.67 (0.4)	0.17 (0.7)	−0.86 (0.8)	−0.50 (0.7)	−0.32 (0.7)
twist (deg)	34.22 (5.7)	30.24 (4.0)	34.81 (5.7)	30.62 (6.4)	32.36 (5.9)	33.12 (6.7)
X-displacement (Å)	−0.23 (0.5)	−4.54 (1.3)	−0.13 (1.1)	−2.33 (2.8)	−1.65 (2.3)	−1.23 (2.4)
helical rise (Å)	3.29 (0.1)	2.77 (0.5)	3.25 (0.2)	3.24 (0.7)	3.25 (0.6)	3.18 (0.7)
inclination (deg)	4.02 (7.2)	16.00 (10.7)	4.01 (6.7)	7.26 (19.1)	8.00 (16.7)	6.30 (19.8)

^aAll values are averaged over nonterminal base pairs with standard deviations given in parentheses. ^bAverages over the A-form structures 3V9D, 3QK4, 2B1B, 1ZEX, 1ZEY, 1ZF1, 1ZF6, 1ZF8, 1ZF9, 1ZFA and the B-form structures 2M2C, 4AGZ, 4AH0, 4AH1, 3U05, 3U08, 1VTJ, 3U2N, 3OIE, 3BSE. ^cAverages over snapshots at 300 K from replica exchange MD simulations. ^dPDB ID code: 1BNA.

In this study, we rely on MD simulations using an implicit continuum dielectric model based on the Generalized Born (GB) methodology^{50–52} to directly observe the effect of reduced dielectric environments on the conformational sampling of DNA. The GB methodology approximates the solvation free energies obtained from Poisson(Boltzmann) theory in a numerically convenient and computationally efficient manner that is suitable for the application in MD simulations. This methodology has been applied previously to study peptides in reduced dielectric environments.^{40,53} We are focusing here on reduced dielectric constants of 20 and 40 that are compared with $\epsilon = 80$. The reduced values cover the effective dielectric response of ethanol–water mixtures as well as that of cellular environments. As mentioned above, 20–30% of water in the cell is replaced by macromolecules with an internal dielectric constant of 2–20^{43–47} and cosolvents.⁴⁸ Furthermore, water itself was found to have a reduced dielectric constant in crowded environments.^{33,49} As a result, the effective total dielectric constant of cellular environments is assumed to be in the range of 20–60 depending on the degree of crowding.^{40,53,54} We considered only scalar, static dielectric constants here as the dielectric response of aqueous solvent is isotropic and largely independent of frequency for the range of dynamics considered here (ps– μ s). Implicit solvent simulations of DNA using GB models with $\epsilon = 80$ were previously shown to closely approximate the conformational sampling of DNA seen in explicit solvent.^{55–58} Therefore, we have confidence in applying the implicit solvent methodology to nucleic acids.

As test systems, we considered the Drew-Dickerson dodecamer d(CGCGAATTCGCG)₂⁵⁹ known to be exceptionally stable in B-form and the GC-rich dodecamer d(CGCCGCGGGCG)₂⁶⁰ which tends toward A-form conformations under conditions of high salt, low humidity, or in the presence of cosolvents. Simulations of both systems were started from the standard B-form experimental structure of the

Drew-Dickerson dodecamer and we hypothesized that a reduced dielectric response may favor A-like structures akin to the effect of cosolvents shifting the A/B balance toward A. This is indeed what we observe as described in more detail below, although the A-like conformations we observe here appear to be distinct from salt-induced A-DNA. In the following, results are described and discussed.

DNA conformations are distinguished by the orientation of their bases, captured by helical parameters, and by backbone torsional angles. Both were analyzed from our simulations and will be described in the following. The most distinctive helical parameters to characterize A- and B-type conformations are the base inclination relative to the helical axis, the twist between subsequent base pairs, and the x-displacement of base pairs from the helical axis. Secondary parameters of common interest are the relative displacement between subsequent base pairs along the base pair axis (slide) and along the helical axis (rise) as well as the propeller twist between bases forming a base pair. The average values from the simulation are given in Tables 1 and 2. Distributions are shown as Supporting Information Figures S1 and S2. The simulation results are compared with averaged helical parameters for ten A-form and ten B-form DNA duplexes and for the X-ray structures of the exact sequences that were studied here. In all analyses, only the eight inner base pairs were taken into consideration, because of structural distortions at the duplex ends due to occasional base fraying.

X-displacement is one of the clearest indicators of A- vs B-DNA conformations. It is expected to have near zero values for B-DNA and significantly negative values for A-DNA, where bases are displaced from the helical axis. We find that, for both the GC-rich and the Drew-Dickerson dodecamers, reduced dielectric environments resulted in a clear shift toward more negative x-displacement values that are about halfway between canonical B- and A-DNA. The shift was slightly greater for the

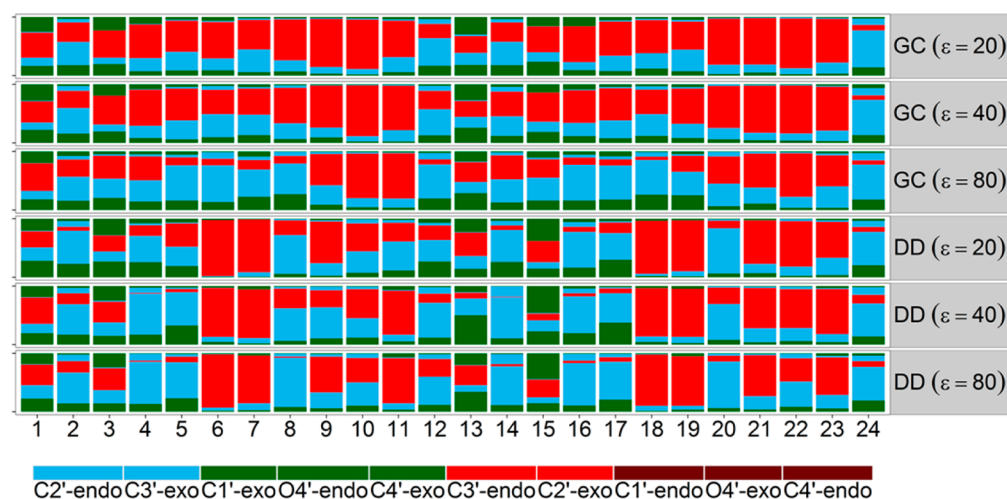


Figure 1. Sugar pucker conformations of each base from simulations of the GC-rich dodecamer (GC), the Drew-Dickerson dodecamer (DD) in different dielectric environments with $\epsilon = 20$, 40, and 80.

GC-rich sequence and was equally large at $\epsilon = 40$ and $\epsilon = 20$. On the other hand, the Drew-Dickerson dodecamer values changed more gradually as the dielectric constant was reduced. A similar observation was made for the twist angle that changed from values of $33\text{--}36^\circ$, typical for B-DNA, to $28\text{--}31^\circ$, typical for A-DNA, as the dielectric constant was changed from $\epsilon = 80$ to $\epsilon = 20$. Again, the change was already complete at $\epsilon = 40$ for the GC-rich sequence but occurred more gradually for the Drew-Dickerson dodecamer. The inclination of base pairs relative to the helical axis in A-type conformations is typically quite pronounced compared to B-type conformations, which are mostly perpendicular to the helical axis. Interestingly, we find that the average for the GC-rich dodecamer shifted to more B-like values as the dielectric constant was reduced. This appears to be in part due to increased sampling of negative inclination angles at $\epsilon = 20$ (see Supporting Information Figure S1). On the other hand, the Drew-Dickerson dodecamer showed a slightly increased inclination angle upon reduction of the dielectric response of the environment. In contrast to what x-displacement and twist values suggest, the changes in base inclination angles are not fully consistent with a classical transition from B-DNA to A-DNA. However, as commented by Dickerson and Ng,⁶¹ the inclination angle can be problematic in short helical segments for distinguishing A- from B-form due to the difficulties in separating inclination from local helix bending.

The slide of a base pair along its long axis relative to its neighboring base pair is another measure that can discriminate between A- and B-form DNA structures. While a significant slide is not observed for base pairs in typical B-DNA, A-DNA base pairs have a more pronounced propensity to slide along their long axes. We observed a significant increase in negative slide values for the GC-rich dodecamer at reduced dielectric constants and to a lesser extent for the Drew-Dickerson dodecamers. Another discriminative helical property is helical rise, which is smaller for typical A-DNA, around $2.7\text{--}2.8$ Å, compared to B-DNA, where it is typically around $3.3\text{--}3.4$ Å. Interestingly, we found the helical rise to remain unchanged upon reduction of the dielectric constants. Finally, the propeller twist of bases in a base pair with respect to each other is known to be highly sequence dependent with more negative values for A/T base pairs than for C/G base pairs, but it is also reduced in A-DNA vs B-DNA. As expected, the propeller twist was less

pronounced for the GC-rich dodecamers than for the AT-base pair containing Drew-Dickerson dodecamer. There was a trend, however, toward less negative propeller twist values upon a decrease of the dielectric constant. This is again indicative of a transition toward A-like structural features. Overall, the analysis of the helical parameters suggests a tendency toward A-type conformations upon a reduction in the dielectric constant. The tendency toward A-DNA was more pronounced for the GC-rich dodecamer and it manifested itself at higher dielectric constants compared to the Drew-Dickerson dodecamer. The helical parameters in the X-ray structure of the B-form Drew-Dickerson dodecamer agree closely with the simulation results at $\epsilon = 80$ while the helical parameters of the A-form X-ray structure of the GC-rich dodecamer are reproduced best by the simulation results at $\epsilon = 20$ and $\epsilon = 40$ rather than $\epsilon = 80$.

Among backbone torsion angles, the ribose sugar pseudorotation angle is the most significant indicator of A- vs B-DNA conformations. While A-DNA structures commonly have C3'-endo or C2'-exo sugar conformations, sugars in B-DNA structures are more often found in C3'-exo or C2'-endo conformations. The preferred sugar conformations for each base in the GC-rich and the Drew-Dickerson dodecamers in different dielectric environments are depicted in Figure 1. As expected from DNA simulations with the CHARMM force field,⁶² there was extensive sampling of both C2'-endo and C3'-endo conformations at $\epsilon = 80$ but with an overall preference for C2'-endo conformers. With a decreasing dielectric constant of the environment, sugar conformations in the GC-rich dodecamer were switched largely to C3'-endo or C2'-exo conformations, further indicating a B to A transition for this sequence. In contrast, the ribose sugars in the Drew-Dickerson dodecamer were largely unaffected by a change in the dielectric environment, suggesting that the backbone retained B-like features.

We analyzed the ϵ and ζ backbone torsion angles as an indicator of the relative sampling of BI and BII conformations. A value of $\epsilon - \zeta$ near -90° corresponds to the BI conformation whereas $+90^\circ$ characterizes BII conformations. The potential of mean force as a function of ϵ/ζ is given in Figure 2. Frequent BI/BII transitions were observed in both dodecamers for all dielectric constants as expected from the most recent version of the CHARMM nucleic acid force field⁶² used in this study. Closer inspection shows that the sampling of BII conformations

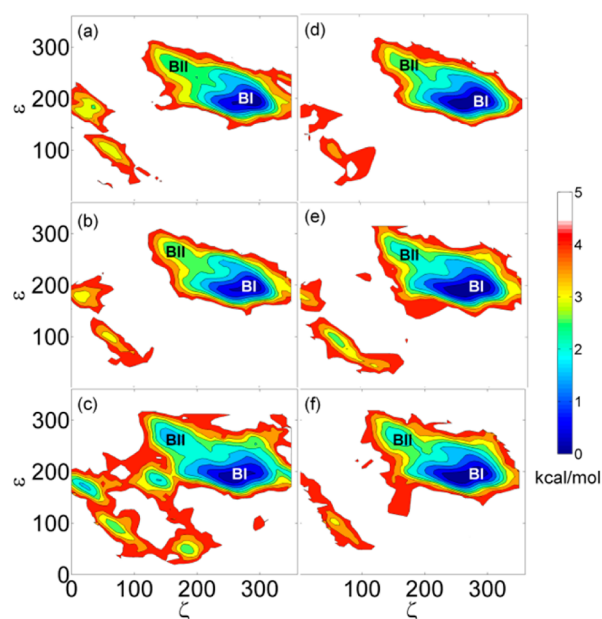


Figure 2. Potential of mean force (kcal/mol) from simulations as a function of ϵ and ζ torsion angles for the GC-rich at $\epsilon = 20$ (a), $\epsilon = 40$ (b), and $\epsilon = 80$ (c) and for the Drew-Dickerson dodecamer at $\epsilon = 20$ (d), $\epsilon = 40$ (e), and $\epsilon = 80$ (f).

diminished at $\epsilon = 20$ and $\epsilon = 40$ for the GC-rich dodecamer and at $\epsilon = 20$ for the Drew-Dickerson dodecamer. This suggests that excursions to the less-populated BI conformer may be further suppressed in reduced dielectric environments. We also found two minima at $\epsilon = 80$ corresponding to noncanonical conformations of the GC-rich dodecamers that appear to be missing at the lower dielectric constants. However, it is likely that this observation is due to limited sampling in our simulations and/or structural distortions due to the implicit solvent model used here (see Figure 2).

Next, we clustered snapshots from each of the simulations to obtain representative structures at different dielectric environments. The resulting structures for both studied dodecamers are shown in Figure 3. Using a clustering radius of 3 Å resulted in five major conformations for the GC-rich dodecamer and three major conformations for the Drew-Dickerson dodecamer. The average helical parameters of these clusters were calculated and compared with canonical A-, B-, and C-DNA forms (Supporting Information). According to this analysis, GC1 and GC2, GC5 and DD3, and DD1 and DD2 have A-like, B-like, and mixed A- and B-type features, respectively, further confirming the prevalence of B-form conformations (GC5 and DD3) at $\epsilon = 80$ and a shift toward A-like structures at lower dielectric constants. We also observed a minor population of a distorted conformation (GC4) at $\epsilon = 80$ that may be a simulation artifact. At the lowest dielectric, a new conformation appears (GC3) that has mixed characteristics of A- and B-type structures. As for the Drew-Dickerson dodecamer, the B-form DD3 conformation persists at $\epsilon = 40$, but disappears at $\epsilon = 20$. Instead, A-like structures (DD1 and DD2) appear with a small percentage at $\epsilon = 40$ while becoming dominant at $\epsilon = 20$. None of the structures observed here resemble other previously characterized conformations of DNA, such as C-DNA.

Finally, we estimated relative conformational free energies using the MMPB/SA approach²⁵ and compared the effect of

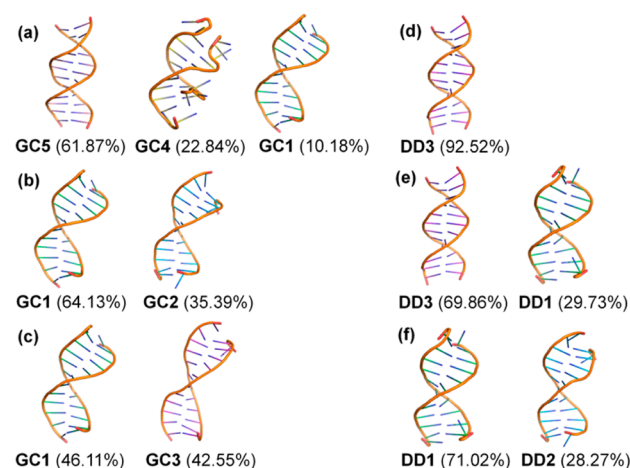


Figure 3. Representative conformations from clustering analysis in different dielectric environment for GC-rich dodecamer with $\epsilon = 80$ (a), 40 (b), and 20 (c) and for the Drew-Dickerson dodecamer with $\epsilon = 80$ (d), 40 (e), and 20 (f). Cluster populations are given in the parentheses.

reduced dielectric environments to previously described effects of increased salt concentrations on the A- vs B-DNA equilibrium.^{6,11,15,18,25} Calculated free energy differences between A- and B-form structures are given in Tables 3 and 4 with positive values indicating a preference for the A-form while negative values indicate a preference for the B-form.

The total free energy differences ($\langle E(\text{total}) \rangle$) at $\epsilon = 80$ between simulated B- and A-forms and between simulated B- and canonical A-structures were more positive for the GC-rich dodecamer (Table 3) than for the Drew-Dickerson dodecamer (Table 4). This indicates that the A-form is relatively more favorable in aqueous medium for the GC-rich dodecamer as we would expect. Furthermore, at $\epsilon = 80$, both dodecamers appear to be more favorable in canonical A-DNA than in the A-form structures observed in our simulations. This is also consistent with expectations since past *in vitro* studies have established the canonical A-DNA structure as the relevant A-form in noncellular environments instead of the somewhat different A-form reported here. We note that the energetic differences are relatively large (tens of kcal/mol instead of physically probably more realistic values of less than 10 kcal/mol). This is likely a result of the approximate nature of the MMPB/SA methodology and/or limitations in sampling. Overall, we estimate that the errors of the MMPB/SA estimates are on the order of tens of kcal/mol, consistent with previous studies.^{21,25,66} Nevertheless, the MMPB/SA estimates appear to give at least qualitatively the correct picture.

Addition of salt to the environment is known to favor A-DNA over B-DNA structures.^{6,11,15,18,25} Within the MMPB/SA framework, salt can be considered by solving the Poisson–Boltzmann equation with added salt.²⁵ While the overall MMPB/SA estimate is prone to uncertainties because, in part, the vacuum force field term is highly sensitive to small conformational changes, and because entropic effects are not fully taken into account, the relative change in the electrostatic solvation term as a result of added salt or a changed dielectric constant can be estimated with much higher accuracy.⁶⁷ We find that canonical A-DNA was favored with increasing salt as expected. However, the trend was reversed when the A-like structures from our simulations were considered instead of the canonical forms. Increasing salt concentration appeared to

Table 3. Conformational Free Energies from MMPB/SA Analysis for the GC-Rich Dodecamer^a

	A ^b	B ^c	canonical A ^d	(B – A)	(B – Can.A)
$\langle E(\text{gas}) \rangle^e$	2181.2 (4.1)	2331.1 (4.1)	2371.2 (4.2)	149.9	–40.1
$\langle E(\text{nonpolar}) \rangle^f$	26.9 (0.0)	26.9 (0.0)	26.3 (0.0)	0.0	0.6
$\langle E(\text{PB}), \epsilon = 80 \rangle^g$	–5837.5 (3.2)	–5965.5 (3.5)	–6069.4 (3.7)	–128.0	103.9
$\langle E(\text{total}), \epsilon = 80 \rangle$	–3629.4 (4.0)	–3607.5 (3.9)	–3672.0 (2.3)	21.9	64.5
$\langle \Delta E(\text{total}), \text{salt: 0.1 M} \rangle$	–40.6 (0.0)	–41.1 (0.0)	–42.9 (0.0)	–0.5	1.8
$\langle \Delta E(\text{total}), \text{salt: 1.0 M} \rangle$	–53.4 (0.1)	–54.5 (0.1)	–58.5 (0.1)	–1.1	4.0
$\langle \Delta E(\text{total}), \epsilon = 20 \rangle$	242.6 (0.2)	247.3 (0.2)	261.1 (0.3)	4.7	–13.8

^aStandard errors are given in the parentheses. ^b100 snapshots selected from GC1 and GC2 clusters closest to cluster centers and with minimal distortions at terminal base pairs. ^c100 snapshots selected from GC5 cluster as in ^b. ^dSnapshots from 1 ns restrained implicit solvent simulation of canonical A-DNA structure taken at 5 ps intervals starting from 0.5 ns. ^eCHARMM force field energy in vacuum. ^fSolvent-accessible surface area term: $\gamma^* \text{SASA} + \beta$ with $\gamma = 0.00542 \text{ kcal/mol/\AA}^2$ and $\beta = 0.92 \text{ kcal/mol}$. ^gSolution of Poisson–Boltzmann equation using PBEQ module⁶⁵ in CHARMM program with a grid spacing of 0.25 Å.

Table 4. Conformational Free Energies from MMPB/SA Analysis for the Drew-Dickerson Dodecamer^a

	A ^b	B ^c	canonical A ^d	(B – A)	(B – Can.A)
$\langle E(\text{gas}) \rangle^e$	2742.6 (5.8)	2812.8 (7.3)	2944.2 (5.3)	70.2	–131.4
$\langle E(\text{nonpolar}) \rangle^f$	26.9 (0.0)	26.7 (0.0)	26.2 (0.0)	–0.2	0.5
$\langle E(\text{PB}), \epsilon = 80 \rangle^g$	–5818.1 (3.9)	–5931.7 (6.1)	–6090.8 (4.6)	–113.6	159.1
$\langle E(\text{total}), \epsilon = 80 \rangle$	–3048.6 (5.5)	–3092.3 (3.5)	–3120.4 (2.0)	–43.7	28.1
$\langle \Delta E(\text{total}), \text{salt: 0.1 M} \rangle$	–40.5 (0.0)	–41.1 (0.1)	–43.1 (0.0)	–0.6	2.0
$\langle \Delta E(\text{total}), \text{salt: 1.0 M} \rangle$	–53.1 (0.1)	–54.2 (0.1)	–59.1 (0.1)	–1.1	4.9
$\langle \Delta E(\text{total}), \epsilon = 20 \rangle$	239.2 (0.2)	245.0 (0.3)	260.4 (0.3)	5.8	–15.4

^aStandard errors are given in the parentheses. ^b100 snapshots selected from DD1 and DD1 clusters as in Table 3. ^c100 snapshots selected from DD3 cluster as in Table 3. ^dSnapshots from 1 ns restrained implicit solvent simulation of canonical A-DNA structure taken at 5 ps intervals starting from 0.5 ns. ^eCHARMM force field energy in vacuum. ^fSolvent-accessible surface area term: $\gamma^* \text{SASA} + \beta$ with $\gamma = 0.00542 \text{ kcal/mol/\AA}^2$ and $\beta = 0.92 \text{ kcal/mol}$. ^gSolution of Poisson–Boltzmann equation using PBEQ module⁶⁵ in CHARMM program with a grid spacing of 0.25 Å.

stabilize the B-form over our A-form structures. On the other hand, a reduced dielectric resulted in a stabilization of our A-like structures while canonical A-DNA structures were destabilized for both dodecamers when compared to $\epsilon = 80$. This suggests that the effects of increased salt and reduced dielectric environment are different and that, although both changes in the environment appear to lead to A-like structures, the resulting conformational ensembles appear to have distinct features.

The main goal of this study has been to examine the conformational sampling of DNA in reduced dielectric environments to both understand the origin of cosolvent induced A-DNA stabilization as well as explore the possible effects of cellular environments. At $\epsilon = 80$ we observe stable sampling of mostly B-DNA structures with the implicit solvent methodology used here in agreement with experimental expectations. This suggests that the implicit solvent methodology can reasonably describe the conformational sampling of DNA in aqueous solutions. Additional simulations were then carried out at lower dielectric constants. This approach emphasizes mean field solvation properties while neglecting specific molecular interactions with the environment. 30.68 (v/v) Ethanol/water solutions have dielectric constants of 30.⁶⁸ At the same time, while $\epsilon = 20$ may be at the low end of what is found in crowded environments, an effective dielectric near $\epsilon = 40$ is assumed to be a typical value for crowded biological cells.^{40,53,54}

Our results indicate an overall shift toward A-like conformations that was moderate at $\epsilon = 40$ and became more pronounced at $\epsilon = 20$ for both studied dodecamers. The tendency to form A-like conformations was greater for the GC-rich dodecamer, but even the classical B-DNA Drew-Dickerson dodecamer exhibited A-like features in low-dielectric environ-

ments. The transition to canonical A-form DNA was nearly complete for the GC-rich dodecamer, but the Drew-Dickerson dodecamers retained many B-like features even at $\epsilon = 20$. However, even for the GC-rich dodecamer at $\epsilon = 20$, the resulting structures differ somewhat from canonical A-form. MMPB/SA analytical results further show that increased salt appears to destabilize the A-like structures seen in our simulations, while it stabilizes canonical A-DNA. This suggests the important conclusion that salt effects and reduced dielectric environments, for example, by reducing “water activity” through cosolvents or crowding, affect DNA structure in similar but different ways. The results are not fully consistent with fiber diffractions studies where DNA remains in canonical B-DNA form with its two hydration layers.^{69,70} However, the fiber environment is likely not as packed as crowded environments, and hence, the effective dielectric environment may not actually be reduced dramatically. Furthermore, DNA–DNA interactions and the presence of salt in fibers may further explain differences between the fiber experiments and our findings. Specific interactions of DNA with proteins in crowded cellular environments as well as volume exclusion effects were also not considered here and may significantly modulate the results reported here. Nevertheless, any tendency toward A-DNA features under crowded cellular conditions could have significant energetic consequences for protein–DNA interactions since most DNA bound to proteins is in B-form.

It is clear that further studies are needed to fully understand DNA structure in crowded environments and in the presence of cosolvents. One specific question to be addressed is whether protein–DNA interactions could counteract the apparent tendency toward A-form DNA and modulate these findings. Furthermore, the studies performed here involved short oligomers vs the very long polymeric DNA found *in vivo*.

Further investigations are also required to see whether longer DNA molecules would exhibit the same tendencies. It is our hope that the results reported will also stimulate new experimental studies of DNA structure under crowded conditions—none of which are available to our knowledge.

■ COMPUTATIONAL DETAILS

Replica exchange molecular dynamics simulations⁷¹ of d(CGCGAATTCGCG)₂ and d(CGCCCGCGGGCG)₂ dodecamers were performed to enhance the sampling by accelerating barrier crossings at elevated temperatures. The Generalized Born with molecular volume (GBMV) implicit solvation model^{50–52} was used with dielectric constants 20, 40, and 80. The initial d(CGCGAATTCGCG)₂ structure was obtained from X-ray analysis (PDB ID code: 1BNA)⁵⁹ and the initial d(CGCCCGCGGGCG)₂ structure was obtained by mutating the base sequence in the X-ray structure of 1BNA. Structures were minimized and equilibrated before the replica exchange simulations. Replica exchange simulations were performed using eight replicas between temperatures 300 and 400 K. Although the replicas visit higher temperatures than 300 K, only the sampling at 300 K is considered and reported here. Exchange probabilities of the simulations were around 25–30%. Langevin dynamics⁷² was performed using a friction coefficient of 50 ps^{−1} to control the temperature of the system. Simulations were carried out for 50 ns for each replica with a total simulation time of 400 ns and replica exchange was attempted every 10 ps. The first 6 ns of each replica was excluded during the analysis of the simulations. Nonbonded interactions were cut off at 18 Å with a switching function becoming effective at 16 Å and a cutoff at 20 Å was used for the nonbonded list.

All simulations were performed using the CHARMM program package (v c37a2)⁶⁵ with CHARMM36 force field^{62,73} for nucleic acids. Replica exchange simulations were carried out using the Multiscale Modeling Tools for Structural Biology (MMTSB)⁷⁴ in combination with CHARMM. Cluster analysis of the conformations was carried out with the kclust program in MMTSB. Helicoidal and backbone parameters of dodecamers were analyzed by using the 3DNA program package.⁷⁵ VMD⁷⁶ and PyMOL⁷⁷ were used for the visualization of structures.

■ ASSOCIATED CONTENT

● Supporting Information

Distribution diagrams for helicoidal parameters. Supplemental figures S1–S2 and tables S1–S2. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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