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The stability of Seeman JX DNA topoisomers of paranemic crossover (PX) molecules as a function of crossover number

Prabal K. Maiti, Tod A. Pascal, Nagarajan Vaidehi and William A. Goddard III\*

Materials and Process Simulation Center (MSC), MC 139-74, California Institute of Technology, Pasadena, CA 91125, USA

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ABSTRACT

We use molecular dynamics simulations in explicit water and salt (Na+) to determine the effect of varying the number of crossover points on the structure and stability of the PX65 paranemic crossover DNA molecule and its JXM topoisomers (M denotes the number of missing crossover points), recently synthesized by the Seeman group at New York University. We find that PX65, with six crossover points, is the most stable, and that the stability decreases monotonically with the number of cross over points PX65 > JX1 > JX2 > JX3 > JX4, with 6, 5, 4, 3 and 2 crossover points, respectively. Thus, for PX65/JX1, the strain energy is 3 kcal/mol/bp, while it is 13 kcal/mol/bp for JX2, JX3 and JX4. Another measure of the stability is the change in the struc ture from the minimum energy structure to the equilibrium structure at 300 K, denoted as root mean-square deviation in coordinates (CRMSD). We findthat CRMSD is3.5A˚ for PX65,increasesto6A˚ for JX1 and increases to 10 A˚ for JX2/JX3/JX4. As the number of crossover points decreases, the distance between the two double helical domains of the PX/JX molecules increases from 20 A˚ for PX65 to 23 A˚ for JX4. This indicates that JX2, JX3 and JX4 are less likely to form, at least in with Na+. However, in all the cases, the two double helical domains have average helicoidal parameters similar to a typical B-DNA of similar length and base sequence.

INTRODUCTION

DNA double helix structures are emerging as a useful scaffold for creating nanostructures and as components for nanomecha nical devices (1,2). Self-assembly of various branched DNA motifs is emerging as an important route for constructing 2D and 3D periodic or aperiodic arrays (3–6), with potential appli cation in DNA-based computation (7–9). Recently, Yan et al. (10) reported the construction of DNA-based nanogrids that

provide an excellent scaffold for the production of highly con ductive, uniform width silver nanowires. A variety of unusual DNA motifs has been synthesized for constructing nanomecha nical devices (11–13). To facilitate the construction of robust DNA nanomechanical devices, the Seeman laboratory at New York University recently invented the new paranemic crossover (PX) class of DNA motifs (14,15) and their JXM topoisomers.

The PX DNA is a four-stranded molecule, in which two parallel double helices are joined by reciprocal exchange of strands at every point where the strands come together (15,16) (see Figure 2). The JXM structure is related to PX by contain ing M adjacent sites where backbones of the two parallel double helices juxtapose without crossing over. Seeman and co-workers (15) have demonstrated that interconversion between PX and JX2 states leads to robust DNA mechanical devices. However, no detailed structural characterizations have been made for these JXM molecules. Earlier, we demon strated (Maiti,P.K., Pascal,T.A., Vaidehi,N., Heo,J. and Goddard,W.A.I., submitted) the use of molecular dynamics (MD) simulations to characterize the thermodynamic stability of PX motifs, where we found the PX65 DNA motif, to be particularly stable. This paper focuses on the JXM topoisomers related to PX65. To assess the relative stability of PX65 and its various JX isomers, we used MD to extract thermodynamic and structural parameters of these molecules as a function of the number of crossovers.

The PX65 molecule has six crossover points at positions 5, 11, 16, 22, 27 and 33, leading to the structure shown in Figure 1. Removing the middle crossover point (position 22) leads to the JX1 molecule (Figure 2). Similarly omitting two, three and four contiguous crossover points leads to the JX2, JX3 and JX4 motifs (also in Figure 2). The details of the MD simulation methods and of building the structures of the PX/JX motifs are given in Methods. The results from MD simulation on these DNA motifs are presented in Results and Discussion. The conclusions are in Summary and Conclusions.

METHODS

Building atomic-level PX nanostructures

The base pair sequences used for building the PX65/JXM molecules are shown in Figure 1. Each PX/JX structure

\*To whom correspondence should be addressed. Tel: +1 626 395 2731; Email: wag@wag.caltech.edu Present address:

Prabal K. Maiti, Department of Physics, Indian Institute of Science, Bangalore 560012, India Nucleic Acids Research, Vol. 32 No. 20 ª Oxford University Press 2004; all rights reserved

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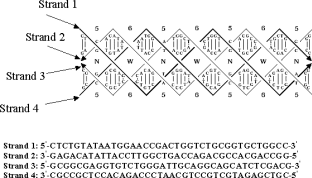


Figure 1. The base pair sequences used in the generations of PX65, JX1, JX2, JX3 and JX4 crossover molecules.

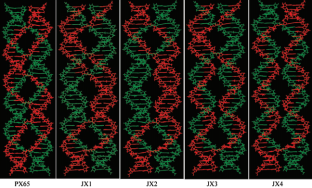
PX65 JX1 JX2 JX3 JX4

Figure 2. Generation of PX and JX DNA by reciprocal exchange. This illustrates the consequences of performing a crossover at various positions. PX65 has six crossover points. JX1, JX2, JX3 and JX4 have 5, 4, 3 and 2 crossover points, respectively.

has two double helical domains and has crossover points between strands of the same polarity. In the notation PXMN (say PX65), the first integer, M = 6, indicates the number of base pairs in the major groove; the second integer, N = 5, indicates the number in the minor groove. Thus, as shown in Figure 2, PX65 has two green strands and two red strands that intertwine each other with six crossover points. The five cases considered here (PX65, JX1, JX2, JX3 and JX4) all have 5 nucleotides in the minor groove and 6 nucleotides in the major groove.

The construction of these five PX/JX DNA motifs used the nucleic acid builder program Namot2 (17) (version 2.2.). The procedure for constructing these structures is as follows:

(i) Building the DNA double helices. First we create two regular B-DNA molecules, with the sequence given in Figure 1. Each of the double helices has 11 bp per turn. Table 1 shows the twist angles used for building the various PX/JX structures. We assigned the same twist angle for all the base pairs in the helical half turn. The

helical rise value of 3.4 s was used to build all the cross over structures.

(ii) Building the crossover points. When a double helix is built in Namot2, the molecules are oriented so that the 50 and 30 ends of the double helices are parallel to the y-axis. To create realistic crossover structures, it is necessary to rotate the individual helices so that the de sired crossover points are closest to each other (rotation angles shown in Table 1). To find this point, we wrote a program that starts with the first crossover point and rotates the first helix in 1increment to find the rotation leading to the shortest distance between these crossover points. Then the first helix is fixed at this prescribed value, while the second helix is rotated to achieve the shortest distance between the crossover points. The sec ond helix is rotated 180more than the first helix so that the helices are arranged as shown in Figure 2. The cross overs were then created using the ‘nick’ and ‘link’ commands in Namot2. These structures are saved in the PDB file format.

Table 1. Helical twist and rotation angles (in degrees) used in building various PX/JX starting structures

PX Twist angle Base pairs Rotation angles(about z-axis) structure (degrees) per turn Helix 1 Helix 2

PX65 30 11 60 240 JX1 30 11 60 240 JX2 30 11 60 240 JX3 30 11 60 240 JX4 30 11 60 240

Simulation details for the PX–JX structures

All MD simulations reported in this paper used the AMBER7 software package (18) with the all-atom AMBER95 force field (FF) (19). AMBER95 FF has been validated for MD simula tions of B-DNA in explicit water with salt, starting from the crystal structure (20–24). These validation studies found that the CRMS deviation from the crystal structure for a dodecamer structure is typically <4 s.

The electrostatics interactions were calculated with the Particle Mesh Ewald (PME) method (25,26) using a cubic B-spline interpolation of order 4 and a 104 tolerance set for the direct space sum cut-off. A real space cut-off of 9 s was used both for the electrostatics and van der Waals interactions with a non-bond list update frequency of 10.

Using the LEAP module in AMBER, the PX/JX nanostruc tures were immersed in a water box using the TIP3P model for water. The box dimensions were chosen in order to ensure a 10 s solvation shell around the DNA structure. In addition, some quantity of water was replaced by Na+ counter ions to neutralize the negative charge on the phosphate groups of the backbone of the PX/JX structures. This procedure resulted in solvated structures, containing 37 000 atoms. The solvated structures were then subjected to 1000 steps of steepest des cent minimization of the potential energy, followed by 2000 steps of conjugate gradient minimization. During this mini mization, the PX/JX DNA nanostructures were fixed in their starting conformations using harmonic constraints with a force constant of 500 kcal/mol/s2. This allowed the water molecules to reorganize to eliminate bad contacts with the PX structures.

The minimized structures were then subjected to 40 ps of MD, using a 2 fs time step for integration. During the MD, the system was gradually heated from 0 to 300 K using weak 20 kcal/mol/s2 harmonic constraints on the solute to its starting structure. This allows for slow relaxation of the built PX structures. In addition, SHAKE constraints (27) using a geometrical tolerance of 5 · 104 s were imposed on all covalent bonds involving hydrogen atoms. This is needed to dynamically prevent changes in the NH and OH bonds from disrupting associated hydrogen bonds. Subsequently, MD was performed under constant pressure–constant temperature con ditions (NPT), with temperature regulation achieved using the Berendsen weak coupling method (28) (0.5 ps time constant for heat bath coupling and 0.2 ps pressure relaxation time). This was followed by another 5000 steps of conjugate gradient minimization while decreasing the force constant of the harmonic restraints from 20 kcal/mol/s2 to zero in steps of 5 kcal/mol s2.

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We then carried out 100 ps of unconstrained NPT MD to equilibrate the system at 300 K. We have found for other systems that the above equilibration protocol produces very stable MD trajectories for simulating large DNA nano structures (Maiti,P.K., Pascal,T.A., Vaidehi,N., Heo,J. and Goddard,W.A.I., submitted). Finally, for analysis of structures and properties, we carried out 2 ns of NVT MD using a heat bath coupling time constant of 1 ps.

Methods used for calculating various properties of the PX nanostructures

Flexibility of the PX nanostructures from MD. To obtain the structure of each PX/JX nanostructures equilibrated in salt and water, we averaged the coordinates of each MD snapshot from 1 to 2 ns at 1 ps time intervals. This averaging was performed only for the last 1 ns to ensure that the structure had converged. This average structure represents the ‘solution structure’ of the PX/JX nanostructures.

Then, to obtain a measure of the flexibility of these struc tures, we calculated the root-mean-square deviation in coor dinates, CRMSD, from the average solution structure for all atoms at each time step. This was performed at every 1 ps time interval in the final 1 ns of MD trajectory. This CRMSD is a measure of the overall flexibility of the PX/JX structures in solution. We also calculated the CRMSD for each base pair from the minimized starting structure using the time average over the last 200 ps for each base pair. This CRMSD from the minimized starting structure shows the flexibility of various regions of the PX/JX structure in solution.

Strain energy or the thermodynamic stability of the crossover motifs. To obtain a measure of the strain energy, we first partition the potential energy into a sum over atoms. This is performed by assigning half the energy for every two-body interaction to each of the two atoms, all the energy for each three-body interaction and each four-body inversion term to the central atom, and half the energy for every four-body dihedral (torsion) interaction to each of the two central atoms. Then, we collect these atomic energies together for each base of the DNA. Since the reference energy in the FF based simulations is not well defined, we use the reference energy for each base pair of the double helix formed by remov ing the crossover points between the double helices as a refer ence state. Then, for each base of the crossover nanostructures, we define the strain energy as the change from the reference structure. Then, summing over all bases, we obtain the total strain energy in each PX/JX structure, which we consider to determine the relative stability between the various crossover structures. Thus, the strain energy is defined as,

DDHstrainð Þ crossover structure

= DHð Þ crossover structure 2DHð Þ double helix 1 where

(i) DDHstrain is the sum of the strains in a given crossover nanostructure (PX/JX molecules),

(ii) DH(crossover structure) is the potential energy of the crossover structure (which is the sum of the strains), (iii) DH(double helix) is the potential energy of the corre sponding double helix without the crossover points (sum of the strains).

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This strain energy represents the energy cost for making a crossover structure and does not include the dependence of the strain energy on the length of the PX/JX structures or the sequence. The average strain energy is calculated by averaging over 200 snapshots uniformly distributed over the last 200–400 ps of the 2 ns MD run. The strain energy per base pair is obtained by dividing the total strain energy from Equation 1 by the number of base pairs in each of the crossover structures. The experimental measure for stability is the melting temperature. This is straightforward to measure but difficult to calculate, since the final state of melting is not so well defined.

The vibrational density of states (DoS) of PX/JX DNA. We also calculated the vibrational DoS of PX/JX nanostructures from the MD as follows (29). First, we calculated the velocity auto-correlation function C(t), defined as the mass weighted sum of the atom velocity autocorrelation functions

the solution phase, there are no reliable experimental struc tures that can be compared with the simulations, which generally lead to RMSD differences of 3.6–4.2 s from the crystal (22,24).

We carried out MD simulations for 2 ns in explicit salt and water for each of the five PX/JX nanostructures (PX65, JX1, JX2, JX3 and JX4) at 300 K. In each case, we defined the average MD structure by averaging the coordinates for various snapshots for the last 1 ns at an interval of 1 ps. This structure represents the time-averaged solution structure of the PX nanostructures (that one would compare to an NMR structure). These averaged structures for various PX structures are shown in Figure 3a and b. For JX2/JX3/JX4 (decreasing number of crossover points), we see that the two double helical domains move further apart, a feature highly undesirable for the application of these DNA motifs to construct periodic arrays. They move further apart and one helical domain gets twisted with respect to the second helical

C tð Þ =XN j=1

X3 k=1

mjckjð Þt 2

domain, leading to large writhing in the structure (see the side view in Figure 3b).

To obtain some idea about the stability as well as the

where ckjð Þt is the velocity autocorrelation of atom j in the k direction

R t

flexibility of these structures, Figure 4a shows the time evolu tion of the CRMSD of instantaneous PX snapshots from the initial minimized canonical structure.

c kjð Þt = lim

tvkj t0 ð Þ + t vkj t0 ð Þdt0

(i) For PX65, the CRMSD increases up to 400–500 ps and

t¥

R ~~t~~

tdt0

then stabilizes between 3 and 4 s over the rest of the three

= lim t¥

1

2t

Z t t

vkj t0 ð Þ + t vkj t0 ð Þdt0

ns trajectory.

(ii) On the other hand, for JX1, the CRMSD fluctuates between 3 and 4 s up to 1 ns, but then increases to 5 s

vkjð Þt is the velocity of the atom j in the k-th direction at time t. The atomic spectrum density skjð Þ u is simply the Fourier transform of ckjð Þt and is given by

over the next 1 ns.

(iii) For JX2/JX3, the CRMSD increases with time, going up to 8 s in 2 ns, while for JX4 the CRMSD increases to

skjð Þ u = lim t¥

1

2t

Z t t

vkjð Þt ei2putdt

10 s. These results suggest that the JX2, JX3 and JX4

2

structures will not retain their helical DNA structures and

perhaps fall apart (but proving this would require much

= lim t¥

1

2t

Z t t

Z t t

vkjð Þt vkj t + t0 ð Þdt0ei2putdt

longer simulation times).

= lim t¥

Z t t

ckjð Þt ei2putdt 3

Experimentally it has not been possible to form any of these structures including PX65 in the presence of Na+ (N. Seeman, personal communication). They have been formed only in the

From which we determined the vibrational DoS (power spec trum) as

presence of Mg2+. On the other hand, in our simulation we see that in the presence of Na+, PX65 is a very stable molecule, and we suggest continued experimental studies in Na+ to

Sð Þ u = 2kTXN j=1

X3 k=1

mjskjð Þ u 4

clarify the apparent disagreement between experiment and theory. We also plan to study these nanostructures in the presence of Mg2+ (Maiti,P.K., Pascal,T.A., Vaidehi,N. and

where mj is the mass of atom j.

RESULTS AND DISCUSSION

Differences in flexibility of the PX/JX motifs

Previous MD simulations have been reported on the crystal structure of B-DNA using explicit salt and water, validating the AMBER FF (20) and the PME method for calculating the non-bond forces (20–22,24). Simulations have also been per formed in solution. The simulations on crystalline B-DNA lead to an overall calculated CRMSD for all atoms of 1.0– 1.5 s (20–22,24). This validates the accuracy of the FF. For

Goddard,W.A.I., submitted). In addition to the comparison to the initial minimized structure, we also calculated the CRMSD of these nanostructures with respect to the time aver aged solution structures as a function of time (see Figure 4b). This gives a better measure of the fluctuations in the structures. This CRMSD was calculated for the whole 2 ns MD runs. We see that the average solution structure for JX2, JX3 and JX4 deviates significantly from their average structure. To provide a quantitative measure of how the separation of the two helical domains varies as a function of the number of crossover points, Figure 5 shows the distance between the center of mass of the two helices during the dynamics. For PX65 the separation is

20 s, which is expected for such a structure; the diameter of normal B-DNA. As the number of crossover points decreases,

**(a) (b)**

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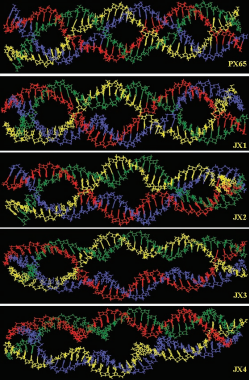
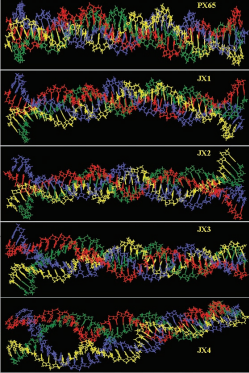


Figure 3.(a) Averaged dynamics structure for various PX molecules. Water molecules and counter ions are not shown for clarity. Note that with decreasing number of crossover points, the two double helical domains move further apart from each other. (b) Averaged dynamics structure for various PX molecules (side view). For clarity, water molecules and counter ions are not shown. With decreasing number of crossover points, there is significant bending of the two helical axis in the opposite direction leading to large writhing in the structure (see Table 4).

the separation increases reaching a maximum of 24 s for JX3. Such increase in separation of the two helices as a function of the crossover separation has been found experimentally for DX molecules as well (30). Surprisingly, for JX4, we see a decrease in the separation to 22.5 s. Very large difference in the bending angle of the two helical axes (2for PX65 and 40for JX4) and resulting writhing in the structure (see Table 4) might be the reason behind this behavior.

Helicoidal parameters and groove dimensions for the PX/JX motifs

Table 2 provides details of the conformational helicoidal parameters of the PX/JX structures averaged over the last 400 ps of the 2 ns long dynamics. For comparison, we also give the values for the two double helices in their non-crossover form. Most helicoidal parameters for the two helices in the PX/JX structures have average values very similar to the corresponding B-DNA form. For example,

the average helical twist for all the PX/JX structures fluctuates between 30 and 32, the range expected for normal B-DNA. This indicates that even though the two helical domains of the crossover structures move further apart as the number of cross over points decreases, the individual double helix regions preserve their B-DNA form. However, the presence of the crossovers points influences the helical conformation signifi cantly. Figure 6 shows the values of rise, tilt, roll and twist averaged over the MD as a function of the sequence along the backbone. At or near the crossover points, we see very large variations in these parameters from the values expected for a B-DNA (denoted by horizontal solid lines in Figure 6). Also for various JX motifs, we see similar trends: the values for rise, tilt, roll and twist for the individual helices remain close to the values expected for normal B-DNA. Table 3 compares the major and minor groove width of the PX/JX structures with the normal B-DNA of same length and sequence. We see that for PX65 and JX1 minor groove width narrows by 10% (from 6.95 to 6.3 s) while major groove depth narrows by 47%

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**(a)**

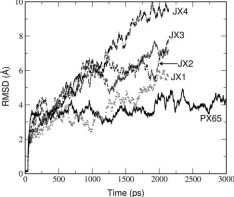
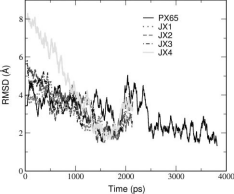
**(b) **

Figure 4. (a) Variation of the CRMSD of all atoms of various snapshots from the MD simulation run with respect to the starting minimized canonical structures. (b) RMSD with respect to average dynamics structure for different PX/JX molecules. The averaged structures were generated by averaging the coordinates for the last 1 ns of the 2 ns long MD runs. PX65/ JX1 with CRMSD 3–5 s is a stable molecule. Large CRMSD for JX2, JX3 and JX4 (8–10 s) suggests that these structures will not retain their helical DNA structures and perhaps fall apart.

(from 7.9 to 4.2 s) compared with its normal B-DNA counterpart. However, for JX2, JX3 and JX4, the minor groove width and depth approaches the canonical B-DNA value, while there is slight increase in the major groove width. These changes might affect the binding of various regulatory proteins to these crossover structures and might be used to physically separate the various generations of crossover structures.

End-to-end distances, strand shortening and bending of helical axis

The variation of end-to-end distance as a function of the number of crossover points is a measure of rigidity for the

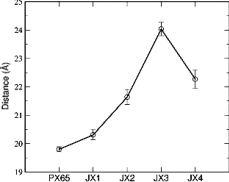


Figure 5. Distance between the center of mass of the two helices. With decreasing number of crossover points, the two helices move apart. The lowering of the distance for JX4 might be due to the large writhing during dynamics. The data has been averaged over the last 400 ps of the 2 ns long dynamics.

crossover molecules. Another closely related quantity is the variation of ‘strand shortening’ for various PX/JX structures over the dynamics. End-to-end distance and strand shortening are calculated as follows: CURVE algorithm outputs the vec torial direction of each local helical axis segment U and its reference point P. The path length between successive helical axis reference points is calculated as

path = X jP~i P~i1j

and the end-to-end distance of the DNA fragment is calculated as

Re = jP~1 P~Nj

where P~1 and P~N are the reference points for the two end helical axis corresponding to two terminal nucleotides. The difference between the sum of all the path lengths and the total end-to-end distance is a measure of the strand shortening. Table 5 reports the average end-to-end distance and strand shortening for both helices over last 200–400 ps of the 2 ns long dynamics.

We see that PX65 has the largest end-to-end extension (126 s for helix1 and 128 s for helix2) and smallest strand shortening (6.24 s for helix1 and 5.05 s for helix2). On the other hand, JX4 shows the smallest end-to-end extensions (116 s for helix1 and 119 s for helix2) and the large strand shortening (12.3 s for helix1 and 9.8 s for helix2). This is consistent with the writhing observed for JX4. The general increase in the end-to-end extension and decrease in strand shortening with increasing number of crossover points implies that increased number of crossover points induces enhanced rigidity in these molecules. This is also observed in the vibrational mode analysis presented in Normal mode analysis section. The larger strand shortening is accompanied by the significant bending in the DNA helical structure, as seen in Table 4.

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Table 2. Helicoidal parameters for the PX/JX molecules averaged over the last 400 ps of the 2 ns MD runs

Parameter B-DNA PX65 JX1 Helix1 Helix2 Helix1 Helix2 Helix1 Helix2

Shift (s) 0.00 (0.3) 0.01 (0.5) 0.02 (0.7) 0.00 (0.8) 0.03 (0.6) 0.01 (0.5) Slide (s) 0.15 (0.3) 0.15 (0.3) 0.05 (0.8) 0.07 (0.7) 0.10 (0.4) 0.12 (0.4) Rise (s) 3.39 (0.4) 3.39 (0.4) 3.53 (0.6) 3.57 (0.4) 3.47 (0.4) 3.48 (0.4) Tilt (degrees) 0.27 (3.4) 0.17 (2.1) 0.25 (5.0) 0.64 (4.3) 0.07 (3.4) 0.03 (4.3) Roll (degrees) 6.37 (5.9) 6.16 (8.1) 2.08 (9.8) 3.34 (11.9) 4.75 (9.2) 4.30 (12.6)

Twist (degrees) 29.71 (3.7) 30.46 (4.5) 32.10 (5.3) 31.7 (7.4) 32.49 (4.9) 32.18 (6.6)

Parameter JX2 JX3 JX4 Helix1 Helix2 Helix1 Helix2 Helix1 Helix2

Shift (s) 0.02 (0.6) 0.01 (0.5) 0.01 (0.5) 0.03 (0.4) 0.05 (1.0) 0.01 (0.4) Slide (s) 0.12 (0.4) 0.13 (0.4) 0.09 (0.5) 0.08 (0.6) 0.10 (0.6) 0.09 (0.5) Rise (s) 3.48 (0.5) 3.47 (0.5) 3.48 (0.5) 3.49 (0.5) 3.5 (0.4) 3.40 (0.3) Tilt (degrees) 0.26 (3.5) 0.33 (3.3) 0.05 (3.3) 0.12 (3.2) 2.8 (6.2) 0.04 (3.1) Roll (degrees) 5.11 (8.4) 5.50 (10.3) 4.78 (8.9) 4.26 (12.5) 4.30 (12.1) 3.90 (9.7) Twist (degrees) 32.3 (5.4) 31.85 (5.7) 31.58 (5.5) 31.94 (6.0) 30.70 (7.8) 32.39 (6.9)

Here, the PX/JX molecules are analyzed in terms of its two double helices. For comparison, we have also tabulated the helical parameters for normal B-DNA of same length and sequence. The RMS deviation from the MD is shown in parenthesis. Note that the individual double helices of the crossover structure retain their B-DNA form quite well.

Table 3. Average major groove and minor groove width for all the PX/JX

molecules

Molecules Major groove width (s)

B-DNA

Minor groove width (s)

Major groove depth (s)

Minor groove depth (s)

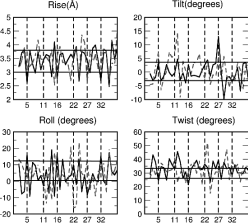


Figure 6. Average rise, tilt, roll and twist for PX motif. Solid line is for helix1 and broken line is for helix2. The vertical lines correspond to the crossover points. The horizontal solid lines give the upper bound and lower bound for the corresponding quantities expected for the helices in their B-DNA form (non-crossover form) during the dynamics. The data has been averaged over last 400 ps of the 3 ns long dynamics. In general, the two double helical domains in the crossover structure keep their B-DNA form quite well. However, at or near crossover points the helical parameters deviate significantly from the values expected in their B-DNA form.

This increased bending with decreased crossovers is further confirmed by the changes in the bending angle of each double helixineveryPX/JXstructure.Thebendingangleiscalculatedas the angle between the successive U~i vector and is defined as

q = cos1 U~i U~iþ1

Figure 7 shows the bending angle variation for i-th and (i + 5)-th base pair calculated for the two double helices for

Helix1 13.65 (0.4) 6.79 (0.2) 7.32 (0.4) 4.11 (0.7) Helix2 14.53 (0.4) 7.1 (0.2) 8.43 (0.3) 3.81 (0.1) PX65

Helix1 13.93 (0.3) 6.43 (0.2) 4.52 (0.3) 4.66 (0.2) Helix2 13.94 (0.3) 6.27 (0.3) 3.82 (0.3) 4.91 (0.1) JX1

Helix1 14.61 (0.2) 5.98 (0.2) 4.93 (0.3) 4.78 (0.1) Helix2 14.94 (0.3) 6.16 (0.2) 5.26 (0.3) 4.53 (0.1) JX2

Helix1 14.75 (0.3) 6.25 (0.2) 7.03 (0.4) 4.27 (0.1) Helix2 14.84 (0.5) 5.96 (0.3) 6.29 (0.3) 4.33 (0.2) JX3

Helix1 14.55 (0.2) 6.42 (0.1) 5.20 (0.3) 4.43 (0.1) Helix2 14.06 (0.2) 5.99 (0.2) 4.92 (0.3) 4.72 (0.1) JX4

Helix1 14.22 (0.3) 6.67 (0.3) 6.68 (0.3) 4.18 (0.1) Helix2 14.42 (0.3) 6.54 (0.2) 6.72 (0.6) 4.19 (0.2)

The data has been averaged over the last 400 ps of the 2.5 ns long MD runs. The width and depths are the sequence averaged values computed with program Curves (32). We see narrowing of minor groove width and major groove depth for PX65/JX1 compared to its normal B-DNA counterpart. However, as the number of crossover points decreases, for JX2/JX3/JX4, the minor groove width and depth approaches the canonical B-DNA value.

various PX/JX nanostructures averaged over the MD simulations. There is no appreciable bending visible for PX65. However,increasedbendingoccursatorneareachmissingcross over point. Thus, for JX1, the missing crossover at the 16th nucleotide position leads bending at or near this crossover point that is 50% larger than the other parts of the structure. This increased bending is also evident in Figures 3, showing snapshots from the MD simulations for each PX structure.

Combining the effect of strand shortening with the bending, we infer that all JX structures show large writhing in solution

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Table 4. Average global bend, global roll and tilt angle calculated by Madbend (31)

angles for each base pair step computed by the Curves program (32). Bends in the helical axis defined by a negative roll angle

Molecules Global bend (degrees)

B-DNA

Global tilt (degrees)

Global roll (degrees)

indicate bending toward the minor groove, while bends defined by a positive roll angle correspond to bending toward the major groove (31). Table 4 gives the values of global bend, tilt and roll for the two helices for all the PX/JX molecules

Helix1 14.14 (8.3) 4.54 (9.1) 1.78 (12.8) Helix2 24.4 (14.2) 14.14 (14.9) 14.1 (13.1) PX65

Helix1 39.01 (13.1) 15.06 (13.4) 33.68 (12.3) Helix2 36.68 (12.3) 2.41 (12.2) 34.28 (12.9) JX1

Helix1 22.75 (11.59) 14.84 (11.2) 8.07 (15.5) Helix2 36.41 (10.87) 17.84(11.7) 29.08 (11.9) JX2

Helix1 15.53 (8.9) 5.75 (11.2) 3.27 (12.3) Helix2 35.06 (10.7) 17.86 (19.2) 20.43 (15.5) JX3

Helix1 27.62 (10.6) 24.22 (10.9) 8.0 (10.3) Helix2 59.09 (14.1) 34.11 (11.1) 45.83 (17.4) JX4

Helix1 71.74 (10.5) 69.28 (12.0) 8.0 (15.8) Helix2 32.58 (13.1) 12.42 (10.1) 27.98 (13.9)

The data has been averaged over last 400 ps of the 2 ns long MD runs. The SDs are shown in brackets. Bends in the helical axis defined by a negative roll angle indicate bending toward the minor groove, while bends defined by a positive roll angle correspond to bending toward the major groove (31). For PX65/JX1, similar bend angle in the same direction (toward minor groove) might be the reason for their greater stability. On the other hand, for JX2 and JX4, we see that the two helices have very different bend angle either in the same direction or in the opposite direction. This led to the very high strain in the all the JX structures as is evident from our strain energy calculation.

Table 5. End-to-end distance and strand shortening for PX/JX molecules

Molecules End-to-end distance (A˚ ) Strand shortening (A˚ ) Helix1 Helix2 Helix1 Helix2

B-DNA 119.31 (1.0) 117.54 (1.33) 6.89 (1.16) 8.79 (1.77) PX65 126.10 (1.36) 128.48 (1.64) 6.24 (0.97) 5.05 (0.96) JX1 123.75 (1.03) 124.25 (1.09) 7.51 (0.88) 7.23 (1.36) JX2 122.03 (0.92) 118.26 (0.88) 8.76 (1.34) 11.82 (2.15) JX3 124.42 (0.97) 123.73 (1.17) 6.29 (0.79) 8.47 (1.28) JX4 115.89 (1.35) 118.98 (0.86) 12.30 (0.89) 9.75 (1.25)

The data has been averaged over last 200–400 ps of the 2.1 ns long dynamics. The SDs are shown in brackets. B-DNA helix1 and helix2 corresponds to the case when the simulation has been performed with the B-DNA with 38 bp with the same sequence for the two double helical domain as shown in Figure 1. We see increase in end-to-end extension and decrease in strand shortening with increasing number of crossover points. These imply that higher number of crossover points induce enhanced rigidity to these molecules.

compared with the PX65 structure. This writhing is an impor tant structural feature to be taken into account in designing nanostructures. For example, the minimal writhing in PX65 makes it a better choice than any JX structures for construct ing 2D arrays using crossover nanostructures. Figure 3b com pares the side views (the average solution structure from the MD run) of the solution structure for PX65 with various JX structures. Clearly, JX2, JX3 and JX4 bend much more than PX65.

We also calculated the global helical bending for each of the two helices using the algorithm developed by Strahs and Schlick (31). This method computes the DNA curvature by summing the projected components of local base pair step tilt and roll angles after adjusting the helical twist. Our analysis for the global angles is based on the values of local tilt and roll

studied in this paper. For comparison, we also calculated the values for the helix1 and helix2 in their B-DNA form. The similar bend angle in the same direction (toward minor groove) observed for PX65, JX1 and B-DNA is consistent with their greater stability. On the other hand, for JX2 and JX4, we see that the two helices have very different bend angles sometimes in the same direction and sometimes in the opposite direction. This leads to the very high strain in the all JX structures found in our strain energy calculation.

Strain energy

As a reference for calculating the strain energy, we carried out MD simulations on the two separate double helices with the same sequence as the PX/JX structures (given in Figure 1). These explicit solvent simulations are used to extract the strain energy for just the DNA fragment and serves as our reference energy. Earlier (Maiti,P.K., Pascal,T.A., Vaidehi,N., Heo,J. and Goddard,W.A.I., submitted), we used the ANAL module of the AMBER7 (18) to obtain the energy for just the DNA. However, ANAL uses evaluate the energetic assuming a non periodic structure with real space cut-offs in computing non bond interactions, rather than the full periodic calculation using PME that was used in the dynamics. The result is a very inaccurate estimate of the electrostatics interactions. Now, we obtain the DNA energy from the explicit solvent runs using the MPSIM program (33), which uses the PME method for the non-bond coulomb interactions. The strain energy is calculated as described in Methods and plotted in Figure 8. We see that PX and JX1 have very little strain, indicating that they are very stable. On the other hand, the very high strains for JX2, JX3 and JX4 suggest that these molecules are very unstable. This indicates that increased number of crossover points increases stability. As the number of crossovers increases, these molecules become quite rigid, an essential feature for their use as building blocks for constructing such larger supramolecular aggregates as 2D or 3D arrays.

Normal mode analysis of the PX/JX structures

The density and distribution of vibrational states at low fre quencies provides valuable insight into the structural changes induced by the crossover points. Accordingly, we calculated the distribution of vibrational modes for each PX/JX structure using the analysis of velocity autocorrelation function, as described in Methods. The vibrational (power) spectra for all crossover structures are shown in Figure 9. For comparison, we also show the vibrational spectrum for one double helix corre sponding to the crossover structures.

The high frequency regime leads to quite similar vibrational spectra for all the PX/JX motifs. Since the AMBER FF requires use of the SHAKE algorithm to constrain the high frequency XH bond vibrations, we do not find any mode frequencies beyond 1400 cm1.

For frequencies below 100 cm1, we see that the DoS decreases with increased number of crossover points.

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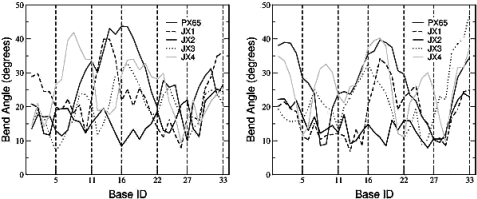


Figure 7. Bending angle between every i-th and (i + 5)-th nucleotide for (a) helix1 and (b) helix2 for each of the PX/JX structures. There is no appreciable bending visible for PX65. However, with decreasing number of crossover points, increased bending occurs at or near each missing crossover point.

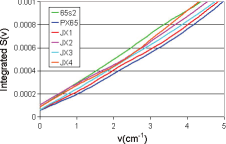
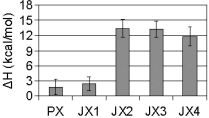


Figure 8. Strain energy for various PX/JX structures. The starin energy has been

calculated with respect to the two separate double helices. Very little strain

energy (3 kcal/mol/bp) for PX65/JX1 indicates that they are very stable. On the other hand the very high strains for JX2, JX3, and JX4 suggest that these molecules are very unstable.

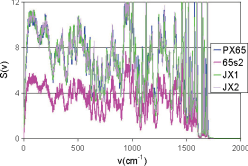


Figure 9. Power spectrum for various PXJX crossover molecules. For com parison, we have also shown the spectrum from 65S2, which is a B-DNA with same length and sequence as that of one of the double helix of PX65.

Figure 10. Intgrated DoS S(n) as a function of n (cm1) for various PX/JX crossover molecules. The population of low frequency modes gradually decreases as the number of crossover points increases making PX65 (with most number of crossover points) more rigid than the other JX structures.

Integrating the power spectrum leads to the integrated DoS shown in Figure 10 for all the PX/JX molecules. The popula tion of low frequency modes is a direct measurement of the rigidity of the DNA molecules since they dominate the overall global dynamics. Thus, the population of low frequency modes gradually decreases as the number of crossover points increases. This indicates the enhanced rigidity of the crossover molecules with increased number of crossover points. This observation is consistent with the experimental findings on another class of crossover molecules, namely the DX mole cules (34), which were found to be very rigid compared with linear DNA molecules. Among all the motifs studied here, PX65 turns out to be more rigid than other structures.

SUMMARY AND CONCLUSIONS

These fully atomistic studies on PX65 and its JXM (M = 1, 2, 3 and 4) topoisomers clearly demonstrate that increased number

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of crossover points provides increased stability to the cross over motifs. However, for the Na+ monovalent ions used in this study, the JX motifs are not able to maintain the parallel double helix crossover structures. Thus, the two helical domains of the crossover structure move increasingly far apart with decreasing number of crossover points. This leads to very large deviations [CRMSD (10 s)] of the JX2, JX3 and JX4 motifs from their initial canonical struc tures. Our strain energy calculations show that motifs with fewer crossover points have strain energies in the range of

12–14 kcal/mol/bp, compared with 3 kcal/mol/bp for PX65 with largest number of crossover points. Our results are consistent with experimental observations that stable JX motifs are found only when using divalent Mg2+ ions. It is plausible that Mg2+ ions might induce effective attraction between the two helical domains to maintain their crossover structures, and we plan to study the behavior of PX/JX motifs in the presence of divalent Mg ions.

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