Reintroducing Electrostatics into Macromolecular Crystallographic Refinement: Application to Neutron Crystallography and DNA Hydration

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SUMMARY

Most current crystallographic structure refinements augment the diffraction data with a priori information consisting of bond, angle, dihedral, planarity restraints, and atomic repulsion based on the Pauli exclusion principle. Yet, electrostatics and van der Waals attraction are physical forces that provide additional a priori information. Here, we assess the inclusion of electrostatics for the force field used for all-atom (including hydrogen) joint neutron/X-ray refinement. Two DNA and a protein crystal structure were refined against joint neutron/X-ray diffraction data sets using force fields without electrostatics or with electrostatics. Hydrogen-bond orientation/geometry favors the inclusion of electrostatics. Refinement of Z-DNA with electrostatics leads to a hypothesis for the entropic stabilization of Z-DNA that may partly explain the thermodynamics of converting the B form of DNA to its Z form. Thus, inclusion of electrostatics assists joint neutron/X-ray refinements, especially for placing and orienting hydrogen atoms.

INTRODUCTION

The initial implementation of crystallographic refinement by simulated annealing (Brünger et al., 1987) used an early version of the CHARMM20 force field (Brooks et al., 1983) that included electrostatics. The benefits of including electrostatics with respect to hydrogen bonding in crystallographic refinement were noted in the refinement of influenza virus hemagglutinin (Weis et al., 1990), although incorrect hydrogen bonds were observed when electrostatics were used during the simulated annealing stages, especially for charged groups such as the head groups of arginine residues. As a compromise, electrostatics was only used during the minimization stages, which resulted in favorable electrostatic interactions while preventing formation of incorrect hydrogen bonds. A likely explanation for the formation of incorrect hydrogen bonds during simulated annealing is that continuum solvent electrostatics are not included in the CHARMM20 force field to dampen the strength of interactions between solvent exposed residues (e.g., Poisson-Boltzmann or Generalized Kirkwood) (Schnieders et al., 2007; Schnieders and Ponder, 2007). Furthermore, the CHARMM20 force field also lacks polarization (Ponder and Case, 2003) and it uses a spherical cutoff rather than Ewald lattice summation (Sagui and Darden, 1999; Karttunen et al., 2008). For simplicity, it has therefore become the practice in most refinement programs to exclude electrostatics during all refinement stages and assume the diffraction data is capable of supplying this excluded a priori information (Adams et al., 1997). More recently, there have been attempts to reintroduce electrostatics in the refinement of NMR structures (Linge et al., 2003) and crystal structures (Moulinier et al., 2003; Korostelev et al., 2004; Schnieders et al., 2009; Fenn et al., 2010). Furthermore, force fields, lattice summation, and continuum electrostatics have matured and computing power increased considerably over the last 20 years. It is therefore important to re-examine the question of the general use of electrostatics in routine crystallographic refinements, especially with respect to the increasing use of neutron diffraction and joint neutron/X-ray diffraction studies where hydrogen bonding is a motivation and electrostatics is expected to have a significant effect. We have therefore focused here on the effect of including electrostatics in the force field on joint neutron/X-ray diffraction crystal structure refinements.

The elastic interaction of X-rays with electrons gives rise to the coherent scattering phenomenon used in macromolecular crystallography experiments. The scattering power of a given atom type correlates with the number of electrons it possesses, and thus increases proportionally with atomic number. This puts light atoms at a disadvantage, especially in the case of hydrogen relative to the common atom types of organic biomolecules. Combined with the low signal to noise ratio of macromolecular
Electrostatic forces are particularly important for atoms that have several degrees of freedom after application of geometric chemical restraints, such as hydrogen atoms involved in hydroxyl groups and solvent molecules. In the case of water molecules modeled as D$_2$O, even when relatively high resolution X-ray and neutron data are available, interpretation of scattering density in terms of oriented or partially oriented water molecules can be difficult, laborious, and subjective (Chatake et al., 2003). Until recently, computation of the electrostatic energy typically involved a conditionally convergent lattice summation based on spherical cutoffs, which is problematic for systems with long range correlations such as crystals (York et al., 1999), large proteins in water (Schreiber and Steinhauser, 1992), nucleic acids (York et al., 1995), and hydrogen-bonding geometry (Morozov et al., 2004). A rigorous and efficient solution termed the particle mesh Ewald (PME) summation was devised, which employs the classical Ewald method (Ewald, 1921) of dividing the problem into rapidly converging real and reciprocal space sums and applies fast Fourier transforms (FFTs) to the latter portion of the summation method (Darden et al., 1993; Sagui and Darden, 1999). We have recently developed a refinement method for subatomic resolution crystal structures (Schnieders et al., 2009; Fenn et al., 2010) that includes a PME approach to rigorously calculate the electrostatic energy and forces, thereby avoiding the problems associated with spherical cutoffs. Further, this method replaces the simple geometric force field that is customarily used in macromolecular refinement (Engh and Huber, 1991) with the polarizable Atomic Multipole Optimized Energetics for Biomolecular Applications (AMOEBA) force field (Ren and Ponder, 2003, 2004; Ponder et al., 2010), which includes a water model that matches experimental observations in varied environments (Ren and Ponder, 2004; Schnieders et al., 2007; Schnieders and Ponder, 2007). We have previously used this approach to refine several subatomic resolution small molecule (Schnieders et al., 2009) and protein (Fenn et al., 2010) X-ray structures, which yielded improved statistical agreement with the diffraction data and more chemically informative models. In this work, we study the effect of including electrostatics using the OPLS-AA nonbonded parameters with a spherical cutoff (Jorgensen and Tirado-Rives, 1988) in conjunction with

**RESULTS**

**B-DNA**

Early X-ray work by Dickerson on DNA described and refined a proposal of hydration in the minor groove of DNA referred to as the zigzag spine (Wing et al., 1980; Drew and Dickerson, 1981). The hydration pattern is defined as a water molecule bridging a base with the base following on the opposing strand in the 5' → 3' direction (e.g., see water W2 in Figure 1). The water
Molecules that form these bridges are joined by a second hydration shell, creating a zigzag pattern (Figure 1). \(^{1}\)H NMR NOESY and ROESY spectra confirm this configuration via the presence of strong, long-lived (>1 ns) NOE signals arising from adenine H2 protons in close proximity to water protons in the primary spine (Kubinec and Wemmer, 1992; Liepinsh et al., 1992; Fawthrop et al., 1993).

Our experiments have independently observed the zigzag spine or long-lived water molecules in a zigzag pattern, including molecular dynamics (Chuprina et al., 1991; Chen et al., 2008; Young et al., 1997a, 1997b) and atomic resolution crystallography studies (Chiu et al., 1999; Minasov et al., 1999; Valls et al., 2004; Woods et al., 2004), and the spine is a critical part of DNA stability (Lan and McLaughlin, 2000). Further, re-refinement of a crystal structure of a B-DNA 9-mer dGCGAATTCG (Soler-Lopez et al., 2000) using the Repel force field (using a real space spherical cutoff of 8.5 Å) produced all the features of the zigzag spine and only forms one hydrogen bond (shown as dashed lines in all figures) in the region shown, suggesting that the particular

![Figure 2. Neutron Diffraction-Based Hydration Structure in B-DNA](image)

(A) Minor groove of DNA resulting from the 2.5 Å refinement with neutron data (PDB ID 1WQZ, sequence dCCATTAATGG) and corresponding 1.6 Å X-ray data (PDB ID 1WQY) using the Repel force field (i.e., without the inclusion of electrostatics).

(B) As in (A), but with electrostatics computed using the OPLS-AA-X force field (using a real space spherical cutoff of 8.5 Å).

(C) As in (A) but using the AMOEBA force field. In all figures, the purple mesh represents X-ray, \(\alpha\), weighted \(2F_{o}-F_{c}\), electron density contoured at 2.0 \(\alpha\) and light blue mesh represents \(\alpha\), weighted \(2F_{o}-F_{c}\), nuclear density contoured at 1.5 \(\alpha\). Density is contoured around the water molecules only for the sake of clarity. Nucleotide bases in white are from the 5' → 3' strand, and those in black correspond to the partner 3' → 5' strand. Figures were generated with POVScript+ (Fenn et al., 2003) and rendered using POVRay.

The results near the zigzag spine are presented in Figures 2A–2C, and the overall statistics are presented in Table 1. Joint neutron/X-ray refinement with the Repel force field (Figure 2A) does not produce any of the features of the zigzag spine and only forms one hydrogen bond (shown as dashed lines in all figures) in the region shown, suggesting that the particular

### Table 1. Joint Neutron/X-Ray Refinement Statistics

<table>
<thead>
<tr>
<th>Structure</th>
<th>PDB ID (N/X)</th>
<th>(d_{\text{im}}(N/X, \text{Å}))</th>
<th>Force Field</th>
<th>Rmsd Bonds(^b)</th>
<th>Rmsd Angles(^b)</th>
<th>(R/R_{\text{free}}(N)) (^a)</th>
<th>(R/R_{\text{free}}(X)) (^a)</th>
<th>HB ratio(^d)</th>
<th>(O_{\nu}-O_{\nu}) (d, \text{Å}) (^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B-DNA 1WQZ/1WQY</td>
<td>2.5/1.6</td>
<td>Repel</td>
<td>0.003 (0.001)</td>
<td>0.60 (0.05)</td>
<td>32.47/35.32</td>
<td>31.90/37.21</td>
<td>0.07</td>
<td>2.98 ± 0.08</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OPLS-AA-X</td>
<td>0.004 (0.008)</td>
<td>0.63 (0.46)</td>
<td>33.15/37.39</td>
<td>32.04/37.00</td>
<td>0.5</td>
<td>2.86 ± 0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>AMOEBA</td>
<td>0.012 (0.007)</td>
<td>2.64 (2.33)</td>
<td>32.55/35.05</td>
<td>31.64/36.24</td>
<td>0.56</td>
<td>2.90 ± 0.20</td>
<td></td>
</tr>
<tr>
<td>Z-DNA 3QBA</td>
<td>1.4/1.5</td>
<td>Repel</td>
<td>0.006 (0.002)</td>
<td>0.76 (0.12)</td>
<td>31.45/32.21</td>
<td>21.89/23.14</td>
<td>0.18</td>
<td>2.94 ± 0.18</td>
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<tr>
<td></td>
<td></td>
<td>OPLS-AA-X</td>
<td>0.007 (0.008)</td>
<td>0.83 (0.38)</td>
<td>31.60/33.28</td>
<td>21.81/23.49</td>
<td>0.44</td>
<td>2.90 ± 0.18</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>AMOEBA</td>
<td>0.011 (0.008)</td>
<td>2.81 (2.51)</td>
<td>30.51/31.95</td>
<td>21.05/24.46</td>
<td>0.55</td>
<td>2.92 ± 0.16</td>
<td></td>
</tr>
<tr>
<td>Xylose isomerase</td>
<td>3KCL/3KBM</td>
<td>2.0/2.0</td>
<td>0.005 (0.001)</td>
<td>0.76 (0.13)</td>
<td>23.69/26.08</td>
<td>14.93/17.95</td>
<td>0.37</td>
<td>2.90 ± 0.14</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OPLS-AA-X</td>
<td>0.006 (0.008)</td>
<td>0.79 (0.40)</td>
<td>23.87/25.57</td>
<td>14.99/17.81</td>
<td>0.68</td>
<td>2.87 ± 0.16</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)N refers to model statistics with respect to the neutron diffraction data, X refers to model statistics with respect to the X-ray diffraction data.

\(^b\)Root mean square deviations from ideal bond lengths and bond angles are given for all atoms and for water molecules alone (in parentheses).

\(^c\)\(R = \sum|F_{\text{calc}}| |F_{\text{obs}}| / \sum|F_{\text{obs}}|\) 10% of reflections in the B-DNA case and 5% of reflections in the Z-DNA and xylose isomerase data sets were set aside for computation of \(R_{\text{free}}\).

\(^d\)Criteria used for hydrogen bonds: hydrogen to water oxygen distance at most 2.5 Å, and \(O_{\nu}-H_{\nu}\)-A angle of 180 ± 40°. Includes bonds to symmetry mates, and only water-water hydrogen bonds are considered.

\(^e\)Expected \(O_{\nu}-O_{\nu}\) distance based on condensed phase small angle neutron/X-ray diffraction value is 2.80 ± 0.04 Å (Soper, 2007).
neutron and X-ray diffraction data do not provide enough information to uniquely determine the hydrogen-bonding pattern. In contrast, joint neutron/X-ray refinement with the OPLS-AA-X force field (Figure 2B) improves the hydrogen bonding, as one of the primary spine hydrogen bonds is reproduced on water molecules W1 and W2. Joint neutron/X-ray refinement using the AMOEBA force field further improves the agreement with the zigzag spine model: water W2 (Figure 2C), for example, rotates to function as a hydrogen-bond donor to the O2 and N3 atoms of the thymine and adenosine, respectively, as evidenced by NMR experiments (Kubinec and Wemmer, 1992; Liepinsh et al., 1992; Fawthrop et al., 1993). Likewise, water molecules W3 and W4 form secondary spine interactions that yield part of the classical zigzag pattern. Water W1 fails to orient correctly. Furthermore, both the neutron and X-ray based $R_{\text{free}}$ values are decreased by 2.3 and 0.7%, respectively compared to refinement with the OPLS-AA-X force field, albeit at the expense of larger deviations from ideal bonds and angle values for the AMOEBA refinements. These differences may be related to differences in the relative weighting of the various energy terms rather than differences in the hydrogen-bonding network, as we did not perform $R_{\text{free}}$-based optimizations of the weights ($\omega_a$ and $\omega_b$, Equation 1). The $2F_o-F_c$ nuclear density maps are of rather poor quality (light blue mesh in Figures 2A–2C); this poor quality is likely due to the low resolution (2.5 Å) and low completeness (59%) of the neutron diffraction data. Thus, the positions of the oxygen atoms of the water molecules are largely determined by the X-ray data, whereas the orientation of the water molecules, and therefore the arrangements of the hydrogen bonds, is largely determined by the force field used in the joint neutron/X-ray refinement.

Z-DNA
The patterns of hydration in Z-DNA have been characterized by a number of crystallographic studies, but have received relatively little attention by computer simulation and nuclear magnetic resonance studies. As a result, the hydrogen-bonding characteristics of this form of DNA have not been experimentally verified. We therefore applied joint neutron/X-ray refinement to high-resolution neutron and X-ray diffraction data of Z-DNA crystals (dCGCGCG) (Langan et al., 2006). Importantly, we were able to obtain the Z-DNA crystals without the use of duplex stabilizing polyamines which can potentially disrupt or stabilize hydration networks, or alternatively facilitate interconversion between DNA structural forms (Hou et al., 2001). We collected a 90% complete neutron data set to 1.4 Å (Table 2), yielding a data to parameter ratio of approximately 2.3 (including hydrogen and deuterium atoms and assuming four independent parameters per atom) and corresponding X-ray diffraction data collected on the same crystal (to 1.5 Å resolution). The overall results of the joint neutron/X-ray refinements are presented in Table 1. In the following, we have restricted our structural references to those of unmodified and identical nucleotide sequence to that studied here (dCGCGCG) for the sake of brevity, although similar conclusions have been reached with other Z-DNA sequences (see, for example, Gessner et al., 1994 and references therein).

A common observation in Z-DNA structure is the hydration of the phosphate group with the preceding (i.e., in the 5’ direction) guanosine N2 amino group (Wang et al., 1984; Gessner et al., 1994; Chatake et al., 2005). Such a configuration is associated with lower interaction energies with phosphate groups as compared with water self-interactions in molecular mechanics studies (Swamy and Clement, 1987), and with long-lived hydrogen bonds in molecular dynamics trajectories (Laaksonen et al., 1989; Eriksson and Laaksonen, 1992). An example of phosphate group solvation resulting from the joint neutron/X-ray refinements is shown in Figure 3. As with the B-DNA results, refinement with the Repel force field (Figure 3A) results in little to no hydrogen bonding of the water molecules with the surrounding environment. The refinement based on the OPLS-AA-X force field (Figure 3B) orients the water molecules to form a network from the guanosine N2 with the following phosphate, as predicted. The AMOEBA-based results agree with the OPLS-AA-X based refinement in this instance (Figure 3C). Also note that the nuclear density agrees with the electrostatics-based results at water molecule W1, although it is weakly defined around water molecule W2 (green mesh in Figure 3). Thus, the introduction of electrostatics with the OPLS-AA-X and AMOEBA force fields are primarily responsible for orienting the deuterium atoms in the latter case.

A second notable motif in Z-DNA hydration is the two-water bridge that forms between N4 amino groups on opposite strands of neighboring cytosine residues (Eriksson and Laaksonen, 1992; Gessner et al., 1994; Chatake et al., 2005). This interaction is shown in Figure 4. In this case, the neutron data are relatively well defined (green mesh in Figure 4). However, only when electrostatics are used for the joint neutron/X-ray refinement (both OPLS-AA-X, Figure 4B, and AMOEBA, Figure 4C) do we observe the hydrogen bonds for the N4-water-water-N4 bridge (see water molecules W1 and W2). Interestingly, the OPLS-AA-X and AMOEBA-based refinements result in a different orientation of water molecule W2, the latter forming an additional hydrogen bond that is part of an extensive hydrogen-bonding network that bridge two symmetry mates (tan DNA molecules in Figure 4). We speculate that this cross-symmetry hydrogen-bonding network stabilizes crystal contacts in a manner similar to the role of ionic Z-DNA crystal forms (Minasov et al., 1999). This more extensive hydrogen-bonding network for the AMOEBA based refinement compared to the OPLS-AA-X based refinement is also reflected in the higher ratio of hydrogen bonds (Table 1). The difference was further analyzed by omitting water molecules W1 and W2 from the model and calculating annealed omit-maps.

### Table 2. Neutron Data Collection Statistics

<table>
<thead>
<tr>
<th>Neutron Data</th>
<th>TOF neutron Laue</th>
</tr>
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<tbody>
<tr>
<td>Diffraction protocol</td>
<td></td>
</tr>
<tr>
<td>$\lambda$ (Å)</td>
<td>0.6–0.7</td>
</tr>
<tr>
<td>Temperature (K)</td>
<td>293</td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>25.6–1.4 (1.45–1.4)</td>
</tr>
<tr>
<td>Unique reflections</td>
<td>4680 (407)</td>
</tr>
<tr>
<td>Completeness</td>
<td>90.1 (78.8)</td>
</tr>
<tr>
<td>Redundancy</td>
<td>5.5 (3.2)</td>
</tr>
<tr>
<td>$\langle d/\sigma \rangle$</td>
<td>5.6 (3.8)</td>
</tr>
<tr>
<td>Rmerge (%)</td>
<td>23 (49)</td>
</tr>
</tbody>
</table>

See also Table S1.

See also Table S1.
The difference nuclear density around both water molecules is in better agreement with the AMOEBA-based refinement (Figure S1B, related to Figure 4C) versus the OPLS-AA-X based refinement (Figure S1A). Thus, it would have been possible to deduce the hydrogen-bonding pattern from inspection of such omit difference neutron density maps alone, but it is gratifying that the AMOEBA-based refinement achieves this pattern without manual inspection.

Similar to the zigzag spine of hydration described for B-DNA, a spine of hydration has likewise been noted for Z-DNA. In the case of a CG dinucleotide repeat (as described here), the hydration spine takes the form of an interstrand water bridge between neighboring O2 oxygen atoms on cytosine residues (Wang et al., 1984; Egli et al., 1991; Bancroft et al., 1994; Gessner et al., 1994; Chatake et al., 2005). The consistency of this water bridge led to proposals regarding stability of CG over AT dinucleotide repeats based on free energy measurements and calculations (Ho and Mooers, 1997 and references therein), the latter of which does not indicate a hydration spine (Wang et al., 1984; Ho and Mooers, 1997). The results of the joint neutron/X-ray refinements in the region of the hydration spine are shown in Figure 5. The apparent feature in all three refinements, refined either with or without electrostatics, indicate that the hydration spine is not a prevalent feature in Z-DNA, as only one of five waters in the spine (the water molecule at the top of Figures 5B and 5C) forms the expected hydrogen-bonding pattern. Surprised by this result, we refined an independently determined Z-DNA structure with X-ray data to atomic resolution (0.95 Å, PDB ID 1ICK) (Dauter and Adamiak, 2001) using the AMOEBA force field. The resultant model from this re-refinement also does not support the hydration spine (Figure S2).

To further analyze this unexpected result, we compared the primary hydration spine bonding geometry in the minor groove of B- versus Z-DNA (Figure 6). In B-DNA (Figures 6A and 6B), the two thymine bases that act as hydrogen-bonding acceptors are offset from one another, which allows the bridging water molecule to interact with the lone pair electrons on both bases at a distance that is optimal for a hydrogen bond (approximately

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**Figure 3. Guanosine N2-Phosphate Hydration in Z-DNA**

(A) Hydration pattern resulting from the joint refinement with the 1.4 Å neutron diffraction data (PDB ID X, sequence dCGCGCG) and with the corresponding X-ray diffraction data to 1.5 Å (PDB ID Y) using the Repel force field (i.e., without the inclusion of electrostatics). Covalent bonds from neighboring nucleotides to the respective phosphate groups (magenta) are omitted for clarity.

(B) As in (A), but with electrostatics computed using the OPLS-AA-X force field (using a real space electrostatic cutoff of 8.5 Å).

(C) Results from refinement using the AMOEBA force field (using a PME-based electrostatic implementation). In all figures, the cyan mesh represents X-ray $\sigma_A$ weighted $F_o-F_c$ electron density contoured at 2 $\sigma$ and green mesh represents $\sigma_A$ weighted $F_o-F_c$ nuclear density contoured at 1.5 $\sigma$ prior to the inclusion of these particular water molecules in the model phase calculations. Density is contoured around the water molecules only for the sake of clarity. See also Figure S1.

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**Figure 4. Cytosine N4-Water and Phosphate Hydration in Z-DNA**

(A) to (C) follow the same order and density rendering as in Figure 3. Symmetry related molecules are colored in tan. Covalent bonds from neighboring nucleotides to the respective phosphate groups (magenta) are omitted for clarity.
1.9 Å, Figure 6B). In Z-DNA, however, the differing geometry leads to more closely stacked cytosine bases (compare Figure 6C with Figure 6A). For the bridging water molecule to interact with both bases in this case, it must straddle the axis formed by the O2 atoms of the two cytosine bases in order to form an electrostatic interaction with the lone pair electrons, which primarily lie in the plane of the base. This also requires the water molecule to lie at a distance that is reasonable for the electrostatic interaction to take place, which is difficult to achieve given the stacked cytosine geometry. This results in an overall poor hydrogen-bonding geometry, evidenced by a calculated ~2 kcal/mol weaker hydrogen-bond interaction in Z-DNA versus B-DNA (based on comparing a primary spine water from the B-DNA result for PDB ID 1ENN as shown in Figure 1 and the Z-DNA result shown in Figure 5C, computed using the AMOEBA force field).

Xylose Isomerase

We applied joint neutron/X-ray refinement with electrostatics to a recently collected neutron and matching X-ray diffraction data set of xylose isomerase (Kovalevsky et al., 2010). The position of hydrogen atoms is of relevance given the importance of water/hydrogen in the mechanism of this enzyme. In brief, the enzyme catalyzes ring opening of sugar monomers, followed by metal-dependent isomerization, via a hydride shift mechanism, between the aldo and keto (with the carbonyl at C2) forms of the sugar (Suekane et al., 1978; Farber et al., 1989; Collyer and Blow, 1990; Collyer et al., 1990; Lee et al., 1990; Allen et al., 1994a; Hu et al., 1997). In general, xylose isomerases tend to favor the aldo form of pentoses (e.g., 88%–12% for xylose and xylulose at equilibrium, respectively), although hexoses (e.g., glucose) also serve as substrates, albeit at a two to three order of magnitude depressed efficiency (van Bastelaere et al., 1991). This latter point is of industrial importance, as column-immobilized whole cells expressing xylose isomerase is the principal method of generating high fructose corn syrup for use as a sugar substitute/food sweetener (for a review of the industrial aspects of xylose isomerase, see Bhosale et al., 1996). Also of crystallographic relevance is the hydride transfer step, which involves metal stabilization of a general acid/base water/hydroxide ion and metal-induced polarization of the substrate to lower the energy barrier for hydride transfer (Schray and...
The active site of xylose isomerase with bound perdeuterated, cyclic α-D-glucose is shown in Figure 7. Along with the $2F_o-F_c$ nuclear density map resulting from joint neutron/X-ray refinement using the Repel and OPLS-AA-X force fields. Unfortunately, limitations in the AMOEBA implementation in TINKER currently preclude the application to large structures such as xylose isomerase. The hydrogen bonding in the OPLS-AA-X based refinement is primarily dictated by the strong electric field at the two metal centers ($\text{Co}^{2+}$ in the case presented), which orients the sugar OD3 and OD4 deuterium atoms away from the ion (M1 in Figure 7). This orientation is in agreement with the result obtained using the Repel force field (Figure 7A). The majority of the water molecule orientations are unchanged upon refinement with and without electrostatics, resulting in a low number of hydrogen bonds at the active site. Inclusion of electrostatics tends to primarily affect regions where the nuclear density is not clearly defined (Figure S3), leading to an overall improvement in the water molecule hydrogen-bonding geometry (Table 1).

DISCUSSION

Macromolecular diffraction techniques are utilized in an increasing number of biological applications over an increasingly larger resolution range. Many of the questions posed and structural models that are generated stand to benefit from a physics-based force field, as the results here indicate. Although the presented examples are primarily restricted to solvent related hydrogen-bonding patterns/geometries due to the inclusion of electrostatics, it is important to point out the crucial role of electrostatics in alpha helix and beta sheet stabilization, and protein stabilization overall: studies have suggested approximately 82% of a protein hydrogen atoms are involved in hydrogen bonding (Baker and Hubbard, 1984; Sticke et al., 1992), with an energetic benefit of up to 2 kcal/mol for each hydrogen bond formed (Fersht and Serrano, 1992). Therefore, physics-based force fields are likely to significantly affect heavy atom positioning and assist in keeping secondary and tertiary structures intact during refinement. This is a significant issue for low-resolution crystal structures, for which additional restraints are sometimes used to maintain secondary structure and proper hydrogen bonding (Brunger, 2005).

Conversely, hydrogen orientation/bonding is often the motive for determining subatomic X-ray crystal structures, for which an improved force field is also of utility (Fenn et al., 2010).

The refinements with force fields that include electrostatics produce energetically more favorable hydrogen bonds and more canonical O-O water-water bond distances (Table 1). In the DNA examples, the AMOEBA-based refinements result in a further improvement, as determined by inspection of omit nuclear density maps (Figure 4; Figure S1) and hydrogen-bonding statistics (Table 1). Further, the models refined with electrostatics agree with theory and independent experiments regarding solvent structure in the minor groove of B-DNA (Figure 2) and offer new insight regarding the stability and structure of the hydration spine in Z-DNA (Figures 5 and 6; see discussion below). Conversely, structures refined with the commonly used repulsive force field without electrostatics do not converge to the same hydrogen-bonding network, especially for cases where the neutron diffraction data do not uniquely determine the hydrogen positions. Even at high resolution and with complete neutron diffraction data electrostatics is a useful complement and can assist careful modeling, as observed for xylose isomerase (Figure 7; Figure S3).

Liquid water hydrogen bonds break at a rate of approximately 1 ps (Luzar and Chandler, 1996; Steinel et al., 2004), whereas hydrogen bonds formed at the interior of proteins can be stable for tens of milliseconds (Otting et al., 1991). Therefore, an important aspect we have not discussed or addressed here regards the dynamic nature of hydrogen-bond networks, and the possibility that hydrogen atoms may not be well defined in terms of their position during the course of a crystallographic experiment. These networks can be modeled as discrete conformers (as is typically done in macromolecular crystallography), although the dynamic nature of some hydrogen bonds may not be adequately modeled in this fashion. Therefore, future methods utilizing ensemble-based approaches coupled with electrostatics may be of utility in this regard (Rice et al., 1998).

Our results show differences in hydrogen-bonding/orientation results between the OPLS-AA-X and AMOEBA force fields. It is possible that the differences result from the fixed charge OPLS-AA-X versus the polarizable atomic multipole AMOEBA force fields or the use of spherical cutoffs versus the particle mesh Ewald method of computing electrostatics. Both of these
factors play a role in the energetics and stability of a given water molecule orientation, particularly in periodic systems. The improved agreement versus the neutron diffraction data with the AMOEBA models (as assessed by annealed omit maps, Figure S1) and higher prevalence of hydrogen bonding (Table 1) suggests that the more physical representations available with this description is desirable over using the OPLS-AA-X force field with spherical cutoffs, although the latter force field offers benefits over not using electrostatics at all. However, it is important to recall that lattice summations computed with a spherical cutoff are expected to contain systematic errors due to lack of convergence of the electrostatic energy (Sagui and Darden, 1999; Karttunen et al., 2008 and references therein).

The ability of DNA to be stabilized in different conformations is widely recognized to be crucial to its biological function. The equilibrium between the right-handed B and left-handed Z forms is complex and poorly understood, but the thermodynamics is thought to be influenced by several factors including hydration, ionic strength, and topological stress (Fuertes et al., 2006). Early on, it was established that the formation of Z-DNA, in particular with the sequence described here, is most likely driven by entropy (Pohl and Jovin, 1972). However, molecular dynamics studies suggest Z-DNA is more rigid than B-DNA (Irikura et al., 1985), implying that solvent entropy must play a significant role in the thermodynamics of the equilibrium from B- to Z-DNA. The finding that the minor groove hydration spine is predominantly disordered, as observed here with all three methods of computing hydrogen-bonding geometry that include both X-ray and neutron diffraction data, is not expected (Wang et al., 1984; Laaksonen et al., 1989; Egli et al., 1991; Bancroft et al., 1994; Gessner et al., 1994; Chatake et al., 2005), but is reasonable given the differences in hydrogen-bonding geometry (Figure 6). Further, if the primary spine were ordered, it is reasonable to also expect a secondary spine bridging the primary water molecules as observed with B-DNA, but this is not the case. We therefore propose the primary water spine in Z-DNA is disordered, and most likely prefers to form transient hydrogen bonds with the cytosine O2 atoms (Figure 5; Figure S2). If this is the case, it is interesting to speculate as to the possibility that the increased disorder in the primary spine represents an increase in entropy upon transiting from B- to Z-DNA, thereby partly explaining the observed free energy difference (Pohl and Jovin, 1972). Recent simulations on B-DNA suggest a gain in entropic free energy of 0.9 kcal/mol for each water molecule moving from the minor groove to bulk solution, suggesting the water molecules that occupy the spine have a significant effect, providing some support to this theory (Jana et al., 2006). This conclusion would be difficult to attain without the combination of crystallographic data and rigorous electrostatics.

It has been noted for some time that medium resolution neutron density “is not sufficiently resolved to orient the water hydrogens precisely” (Finer-Moore et al., 1992), a shortcoming that can be partly compensated by use of more physics-based force fields during refinement, as described here. Even for high-resolution neutron structures, the joint neutron/X-ray refinement using the OPLS-AA-X or AMOEBA force fields provides a means of automating water molecule building, one of the more laborious and subjective tasks in structure refinements against neutron diffraction data and joint neutron/X-ray diffraction data. The improved force fields that are used in this work and elsewhere (Schneider et al., 2009; Fenn et al., 2010) suggest a more widespread “resurrection” of refinements that include electrostatics as had been initially used in crystallographic refinement (Weis et al., 1990). We expect that such refinements will improve hydrogen-bonding geometry and yield more chemically reasonable models for studies of catalysis, drug design, ligand bonding and molecular stability, systems for which electrostatics plays a major role. Indeed, the need for rigorous electrostatics methods (Warshel and Papazyan, 1998) and advanced force fields (Morozov et al., 2004) in structural biology has been stressed in the past, which we believe the results presented here partly fulfill. However, we note that the inclusion of implicit solvent to model disordered solvent and counter-ion atmospheres in crystals remains an area that needs further development in the context of crystallographic refinement, as the current exclusion of such phenomena may lead to spurious hydrogen-bonding patterns involving charged groups (Weis et al., 1990; Moulinier et al., 2003; Korostelev et al., 2004). This will be particularly important during the early stages of refinement, when solvent networks and side-chain orientations are incomplete and/or incorrect. Therefore, the proposed methods are best utilized at the final stages of refinement to avoid artifacts resulting from incomplete electrostatics.

**EXPERIMENTAL PROCEDURES**

**B-DNA and Xylose Isomerase Models**

The neutron diffraction data for the B-DNA (dGCCAATTCG) was obtained from PDB ID 1WQZ (at 2.5 Å resolution) and corresponding X-ray diffraction data from PDB ID 1WOY (at 1.8 Å resolution) (Arai et al., 2005). The neutron diffraction data for xylose isomerase was obtained from PDB ID 3KCL (at 2.0 Å resolution) and the corresponding X-ray diffraction data from PDB ID 3KMB (at 2.0 Å resolution) (Kovalevsky et al., 2010). In both cases, the deposited neutron models were used as starting models for the model refinement, detailed below. Water molecules lacking deuterium atoms were fully deuterated prior to refinement.

**Z-DNA**

Synthesis, crystallization and preliminary X-ray and neutron data collection for Z-DNA have been previously described (Langan et al., 2006). For this study additional neutron diffraction data were collected using the Protein Crystallography Station (PCS) at the Los Alamos Neutron Science Center (Langan et al., 2006) in order to improve completeness and resolution (resolution: 1.4 Å; completeness 90%). The data collection statistics are provided in Table 2. The X-ray data statistics are as given in Langan et al. (2006). The final, deposited atomic model (PDB ID 3OBA) included discrete disorder at three non-phosphate backbone positions, and the statistics for this model with respect to both the X-ray and neutron data is given in Table S1.

**Crystallographic Refinements**

Refinement of each model was performed using an updated version of nCNS (Langan et al., 2008; Adams et al., 2009) using CNS v1.3 as the base (Schroder et al., 2010) (the nCNS patch for CNS v1.3 is available upon request from M.M. and P.L.). In addition, an interface between this new version of nCNS and TINKER was implemented as previously described (Schnieders et al., 2009). In the case of B-DNA and xylose isomerase, hydrogen/deuterium assignments and positions were left unaltered from the deposited coordinates. All refinements included hydrogen atoms (including aliphatic groups). Refinement was carried out using a maximum likelihood amplitude based target function that combined the X-ray diffraction maximum likelihood term, neutron diffraction maximum likelihood term, and force field energy:

$$E_{total} = w_{o}E_{x-ray} + w_{n}E_{neutron} + E_{forcefield}$$  

**Table 1**

<table>
<thead>
<tr>
<th>Protein</th>
<th>Resolution (Å)</th>
<th>Completeness (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B-DNA</td>
<td>1.8</td>
<td>90</td>
</tr>
<tr>
<td>Z-DNA</td>
<td>2.0</td>
<td>90</td>
</tr>
</tbody>
</table>
Refinements were carried out with one of three force fields: the first force field used the all hydrogen parameters for proteins and nucleic acids as implemented in CNS v1.3 that incorporates non-hydrogen target and energy values for bond, angle, and dihedral potentials as described by Engh and Huber (1991) and Parkinson et al. (1996) along with a purely repulsive nonbonded potential function without electrostatics (Adams et al., 1997) (referred to as "Repel," the topology and parameter files "protein-allhydro5-4.4," "dna-ma-allatom-hij-opls," and "water-allhydro5-4.4" were used with no electrostatics and the "repel" option). The second force field used the same non-hydrogen-bond lengths, angles and dihedral terms as the "Repel" force field, but with the OPLS-AA nonbonded potential (Jorgensen and Tirado-Rives, 1988) which includes van der Waals and electrostatic potentials, computed using a real space spherical cutoff of 8.5 Å with a switching function that begins taking effect at 6.5 Å, and the CHARMM22 version of the TIP3P water model (but with stiff, not fixed bond lengths and angles with energy constants of 1000 kcal/mol Å² and 500 kcal/mol, respectively) as implemented in CNS v1.3, revision 3 (referred to as the "OPLS- AA-X" force field, the topology and parameter files "protein-allhydro5-4.4," "dna-ma-allatom-hij-opls," and "water-allhydro5-4.4," were used in conjunction with the "electrostatics.settings" file in CNS version 1.3, revision 3) (Linge et al., 2003; Nozovici et al., 2010). The third force field is the all-hydrogen polarizable AMOEBA force field (Ren and Ponder, 2003, 2004) as implemented in TINKER (which includes van der Waals and PME electrostatic terms (Sagui et al., 2004), referred to as "AMOEBA"). For the PME summation, we used a real space cutoff of 7.0 Å, an Ewald coefficient of 0.545, grid spacing of 1.2 Å⁻¹, and fifth order B-splines. The weights (wR and wP) were set to 1.0 in all cases, i.e., we did not perform Rfree-based optimization of these weights. Three cycles of refinement were carried out for each, which consisted of 250 rounds of positional and 100 rounds of B-factor based minimization. In the xylose isomerase test case, the system size was too large for TINKER when expanded to space group P1, and therefore only the Repel and OPLS-AA-X force fields could be tested for this case.

Contour levels of X-Ray and Nuclear Density Maps

The standard deviation (σ) of a Fourier map is often used as the unit for the contour levels in the map. For example, contours might be drawn at 1.5 σ, at 3 σ, and at other multiples of σ. The sigma value is calculated by the following equation:

\[ \sigma = \frac{1}{N} \sum (\rho - \langle \rho \rangle)^2 \]

where ρ is the electron density for XPC and the nuclear density for NPC. N is the number of grid points in a unit cell. For neutron density (2ρF+2ρF maps, the map is normalized to unit sum of the electron density in holes in the map. Therefore, the deviations with respect to the mean are larger for nuclear density maps, and as a result, σ becomes larger. This means that the same value of σ represents a stronger signal in NPC maps that in electron density maps. Therefore, all figures used lower contour levels when displaying nuclear density versus electron density.

Model Evaluation

To construct Table 1, we wrote a computer program to systematically search for hydrogen-bond donors/acceptors with the assistance of the CCP4 coordinate library (Krissinel et al., 2004). Each water hydrogen atom (or deuterium in the case of neutron structures) was analyzed by searching for water oxygen atoms within 2.5 Å of the hydrogen, including symmetry mates. The O-H...O-A angle was then checked, and only angles within 180 ± 40° were considered for analysis. The program is available upon request.

SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures and one table and can be found with this article online at doi:10.1016/j.str.2011.01.015.

ACKNOWLEDGMENTS

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## Supplemental Information

### Reintroducing Electrostatics into Macromolecular Crystallographic Refinement: Application to Neutron Crystallography and DNA Hydration

Timothy D. Fenn, Michael J. Schnieders, Marat Mustyakimov, Chuanjie Wu, Paul Langan, Vijay S. Pande, and Axel T. Brunger

**Table S1, related to Table 2.** Final statistics for deposited Z-DNA model refined against both neutron and X-ray data.

<table>
<thead>
<tr>
<th>Data statistics: Neutron Data</th>
<th>Z-DNA</th>
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<tbody>
<tr>
<td>PDB ID</td>
<td>3QBB</td>
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<tr>
<td>Resolution range (Å)</td>
<td>100.0-1.4</td>
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<tr>
<td>No. of reflections (theoretical)</td>
<td>4680 (5186)</td>
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<tr>
<td>Completeness (highest resolution shell) (%)</td>
<td>90.2 (78.8)</td>
</tr>
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<td>$R$-factor ($R_{free}$) (%)</td>
<td>19.37 (23.59)</td>
</tr>
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</table>

<table>
<thead>
<tr>
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<th>Z-DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDB ID</td>
<td>3QBA</td>
</tr>
<tr>
<td>Resolution range (Å)</td>
<td>100.0-1.53</td>
</tr>
<tr>
<td>No. of reflections (theoretical)</td>
<td>3696 (4012)</td>
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<tr>
<td>Completeness (highest resolution shell) (%)</td>
<td>92.1 (65.3)</td>
</tr>
<tr>
<td>$R$-factor ($R_{free}$) (%)</td>
<td>30.07 (31.47)</td>
</tr>
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</table>

<table>
<thead>
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<th></th>
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</thead>
<tbody>
<tr>
<td>No. of solvent atoms</td>
<td>41</td>
</tr>
<tr>
<td>No. of disordered phosphate residues</td>
<td>3</td>
</tr>
<tr>
<td>Average B-factor</td>
<td>4.6</td>
</tr>
<tr>
<td>rmsd bonds</td>
<td>0.015</td>
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<tr>
<td>rmsd angles</td>
<td>4.33</td>
</tr>
<tr>
<td>HB ratio</td>
<td>0.52</td>
</tr>
<tr>
<td>Ow-Ow (d, Å)</td>
<td>2.90 ± 0.15</td>
</tr>
</tbody>
</table>
**Figure S1.**

**Figure S1, related to Figure 3.**
Simulated annealing neutron omit maps at the water molecule W1 and W2 positions around the cytosine N4 position of Z-DNA (main text Figure 3). In both panels, the simulated annealing neutron omit map was computed from a joint neutron/X-ray refinement using the OPLS-AA-X force field, and is contoured at $2 \sigma$ (green mesh in both panels). (A) Resultant model from electrostatics computed from a joint neutron/X-ray refinement using the OPLS-AA-X force field overlaid on the simulated annealing omit nuclear density map. A hydrogen atom from water molecule W1 sticks out of the plane of the image and is out of density, and both water molecule W2 hydrogen atoms are out of the neutron density. (B) Resultant model from joint neutron/X-ray refinement using the AMOEBA force field overlaid on the simulated annealing omit map. In this case, water molecule W1 is in nuclear density and one of the hydrogen atoms on water molecule W2 is in the nuclear density, while maintaining the necessary hydrogen bonding pattern with the cytosine N4 atoms.
Figure S2, related to Figure 5.
Z-DNA hydration spine in a 0.95 Å X-ray structure (PDB ID 1ICK) re-refined with hydrogen atoms included in the model using the AMOEBA force field. The $\sigma_A$ weighted 2Fo-Fc electron density map is shown contoured at 2 $\sigma$ in light blue. As with the Z-DNA structure reported in the main text, very few hydrogen bonds are formed with the cytosine O2 atoms. What hydrogen bonds that are formed are through a two water molecule bridge (at the top of the Figure), which is not predicted by the hydration spine model.
Figure S3.

**Figure S3, related to Figure 7.**

Hydrogen bonding at the periphery of xylose isomerase following joint re-refinement of the 2.0 Å X-ray (PDB ID 3KBM) and 2.0 Å neutron (PDB ID 3KCL) data. (A) Resultant model and nuclear density maps following refinement with the Repel force field. The water at the center of the figure only makes one hydrogen bond with the surrounding aspartate residues, and no water-water hydrogen bonds. The star on Asp336 denotes it originates from a symmetry mate. (B) As in (A), but with electrostatics computed using the OPLS-AA-X force field (using a real space spherical cutoff of 8.5 Å). In this case, the waters hydrogen bond and the center water rotates to form a favorable hydrogen bond with both aspartate residues. The $\sigma_A$ weighted 2Fo-Fc nuclear density map is shown contoured at 1.5 $\sigma$ in light blue.