Simulations of Nucleic Acids and Their Complexes[†]

EMMANUEL GIUDICE AND RICHARD LAVERY*

Laboratoire de Biochimie Théorique, CNRS UPR 9080, Institut de Biologie Physico-Chimique, 13, rue Pierre et Marie Curie, Paris 75005, France

Received July 16, 2001

ABSTRACT

Recent years have seen considerable progress in simulations of nucleic acids. Improvements in force fields, simulation techniques and protocols, and increasing computer power have all contributed to making nanosecond-scale simulations of both DNA and RNA commonplace. The results are already helping to explain how nucleic acids respond to their environment and to their base sequence and to reveal the factors underlying recognition processes by probing biologically important nucleic acid—protein interactions and medically important nucleic acid—drug complexation. This Account summarizes methodological progress and applications of molecular dynamics to nucleic acids over the past few years and tries to identify remaining challenges.

Introduction

This Account attempts to summarize the methodological developments and practical applications of molecular dynamics simulations applied to nucleic acids. We have considered studies involving both DNA and RNA, on isolated molecules and on complexes with drugs or proteins. In each case, we have tried to underline recent breakthroughs and continuing difficulties. Each section is illustrated with references drawn from the past four years of publication which, without being exhaustive, should cover a sufficiently broad spectrum to demonstrate how this field of molecular simulation is progressing. For other points of view and further references, the reader is referred to some recent review articles.^{1–3}

Force Fields and Methodology

Thanks to considerable progress in force fields over the past few years, stable multi-nanosecond simulations of nucleic acids are now routine. AMBER parameters Parm94 (or the slightly revised Parm98),⁴ CHARMM27,^{5,6} and BMS⁷ force fields all yield reasonable results for the A and B conformations of DNA (Figure 1). It is also worth remarking that stable B-DNA simulations^{8,9} have also been achieved with the latest GROMOS implementation.¹⁰ A number of tests have already been performed on both the

AMBER and CHARMM force fields in solution^{11,12} and in crystalline environments¹³ on both DNA and RNA. More stringent tests of energy barriers for conformational transitions remain to be carried out.¹⁴ A key element in all nucleic acid simulations is electrostatics. Brutal truncation methods lead to problems, but both particle mesh Ewald¹⁵ and atom-based force-shift approaches¹⁶ can yield stable structures. While force field parameters are continuously being improved thanks to accurate experimental data and quantum calculations on small systems, good results on macromolecules still require some less rigorous retouching in many cases. Force field refinements such as off-atom charges and the inclusion of polarization energy should help to improve this situation.¹⁷

In addition to the improvement of force fields, one of the main computational challenges remains the simulation of large systems over long times. One approach to this goal is the replacement of explicit solvent with hybrid explicit/implicit^{18,19} or entirely implicit models.^{20,21} Poisson-Boltzmann (PB) approaches have proved to be capable of reliably predicting both solvation energies and solvent-dependent conformational changes, but their computational complexity hinders their use in MD simulations. More recent efforts have focused on Generalized Born (GB) models which are much faster and can be formulated to include both salt and cavity contributions.^{20,22} Such methods can be parametrized to yield reasonable solvation energies^{22,23} and pK_a shifts,^{22,24} and they have already been used for MD simulations on nucleic acids,²⁵ although our own experience suggests that some base sequences can lead to the appearance of deformed conformations (Figure 2). Further efforts in parametrization, and perhaps in formulation, are probably necessary. Despite these caveats, both PB and GB approaches, combined with structural snapshots coming from explicit solvent MD simulations, have been used to very good effect in estimating free energies in a variety of systems.^{26,27} These so-called MM/PBSA or MM/GBSA approaches open an exciting route to calculating free energy changes for processes that are not accessible to conventional free energy algorithms.

Treating very large nucleic acids or dramatically extending simulation times will require replacing conventional Cartesian coordinate atom-level dynamics with simplified representations. A first step is the introduction of collective variables,²⁸ which allow certain degrees of freedom to be frozen, decreasing the dimension of the conformational space to be searched and allowing longer time steps to be used. Such methods have already been developed and applied to DNA with time steps around 10 fs, with good results in terms of structure¹⁸ and sequence-dependent curvature.³⁰ Other attempts have been made using a Lagrangian and quaternion formula-

Emmanuel Giudice, born in Corbeil-Essonnes, France, in 1976, is a doctoral student at the Laboratory de Biochimie Théorique (LBT) in Paris. His research concerns dynamic simulations of DNA deformation.

Richard Lavery, born in Aldershot, UK, in 1952, is Director of the Laboratory de Biochimie Théorique (LBT) in Paris. After training in quantum chemistry at the University of Manchester, he has spent the past 24 years in France working in the field of macromolecular modeling and simulation.

[†] This Account is dedicated to Peter Kollman, whose recent death is a great loss to our field. His contributions to the development of macro-molecular simulations and, notably, applications to nucleic acids have been instrumental in many areas of development.

 $[\]ast$ To whom correspondence should be addressed. E-mail: rlavery@ ibpc.fr.

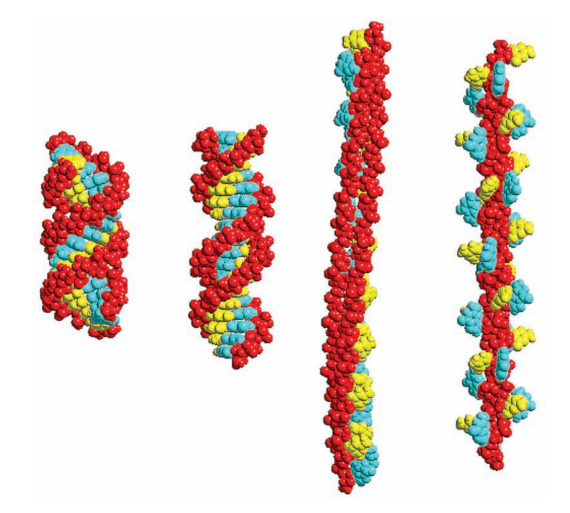


FIGURE 1. Canonical and mechanically distorted forms of helical DNA (from left to right: A-DNA, B-DNA, overstretched S-DNA,³² overtwisted P-DNA³³).

tion coupled with an effective force field.³¹ These methods naturally pose the problem of correctly treating solvent and counterion effects. They have, however, been very useful in modeling extreme deformations, such as those induced during single-molecule experiments on DNA (Figure 1).^{32,33} More extreme simplifications involving the use of elastic rod or bead models allow simulations to be extended to genome-sized fragments but also pose major questions concerning the degrees of freedom to be maintained and the parametrization of the models.^{34,35}

Beyond time constraints, the difficulty of sampling conformational space can also lead to difficulties, notably in searching for complex folded conformations or in attempting to dock macromolecules together. In such cases, the use of multicopy algorithms can be very helpful,^{28,29} and this has recently been demonstrated with the locally enhanced sampling approach now available within AMBER.³⁶ Tools for controlling a dynamic trajectory are also of considerable importance, especially for studying phenomena that would normally occur too slowly. Digital filtering, which can target elements of a structure on the basis of their characteristic frequencies of their dynamics, is a recent and interesting addition to such methods.³⁷

DNA Structure and Flexibility

MD simulations have been used to look at the factors underlying DNA structure and flexibility and also the sequence dependence of these properties (Figure 3). At the local level, the free energy of both base pairing³⁸ and base stacking³⁹ has been calculated using isolated bases or dinucleotide monophosphate fragments, while other studies have probed these interactions within the double helix via induced base pair opening (Figure 4).⁴⁰ The latter results show that both bases can open into either groove of the duplex and that once a base is fully opened, water can fill the gap left within the duplex. Similar behavior is seen in the case of abasic nucleotides.⁴¹ Although normal bases open spontaneously on the millisecond time scale, unforced opening into the major groove has been seen on a subnanosecond time scale for a modified AT pair. where thymine was replaced with a more weakly binding analogue, difluorotoluene.42

Base sequence effects on stacking have been demonstrated with short MD runs, which confirm that ApA steps seem to have well-defined geometries, while CpA steps can show bimodal behavior.⁴³ Other studies have generalized the malleability of CpA to other pyrimidine-purine steps.^{44,45} Similarly, in line with NMR and CD data, runs

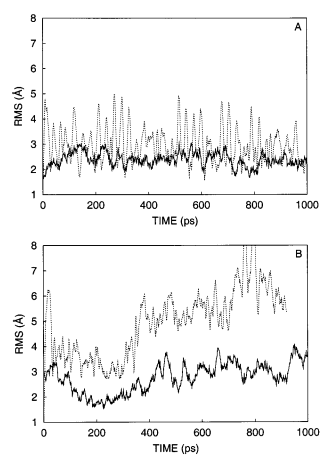


FIGURE 2. RMS deviations from a canonical B-DNA structure during dynamic simulations of 12-base-pair oligomers using explicit (solid lines) or implicit²⁵ (dashed lines) solvent: (a) an alternating GA sequence, (b) an alternating AT sequence (A. Lebrun, LBT Paris).

of GC base pairs have been shown to alter the stacking pattern to resemble that found in A-DNA, even if the backbone geometry remains close to that of B-DNA.⁴⁶⁻⁴⁸

Studies of backbone fluctuations have shown that phosphate movements associated with the B_{II} state ($\epsilon \zeta$ tg⁻ \rightarrow g⁻t) occur for most nucleotides, although they are facilitated by YpR steps.^{49,50} These transitions are linked to base unstacking and water rearrangement and are also favored by lower water activity.⁵¹ B_{II} states can last from a few picoseconds to several nanoseconds. However, even when such backbone conformations are relatively long-lived, they only weakly influence the helical conformation of DNA.⁵²

On a more global scale, a number of simulations have looked at the nature of the A \rightarrow B transition. This transition, which is directly coupled to changes in sugar puckering,⁵³ has been shown to occur spontaneously in water, although the reverse transition does not occur in an ethanol/water mixture, presumably because of a significant energy barrier.⁵⁴ Poisson–Boltzmann or Generalized Born solvation energy calculations using MD snapshots confirm the experimental observation that GCrich sequences are more A-philic⁴⁶ and show that the transition can be linked to a shift in the balance of interphosphate repulsion, desolvation, and ion–DNA interaction terms.⁵⁵ Studies of sequence effects on global DNA structure and flexibility have mainly dealt with A-tract-induced curvature, where the consensus view is now that A-tracts are straight and that curvature is explained by a junctiontype model with roll toward the major groove concentrated at the 5'-end of the A-tract.^{44,56–58} In line with experiment, simulations of A-tracts show minor groove narrowing in the 5' \rightarrow 3' direction, while TpA junctions widen the minor groove and disfavor curvature. ^{56,59}

Other AT sequences which have attracted the attention of several groups are the TATA box sites which are targeted, and strongly deformed, by the TATA box binding protein (TBP). In line with experiment, MD simulations have confirmed that AT sequences showing stronger and more anisotropic DNA bending toward the major groove are correlated with stronger TBP binding,⁶⁰ while PMF calculations also show a correlation with the deformation free energy necessary to reach the so-called TA conformation induced by this protein.^{61,62} Beyond these cases, only limited studies of sequence effects have been made so far, although one study of simple homopolymeric and dinucleotide repeating sequences has used a harmonic analysis of MD simulations to show that only relatively small, sequence-dependent variations occur in global rodlike flexibility (twisting, bending, stretching).63

DNA—Ligand Complexes

Studying ligand binding with MD simulations is apparently proving to be a reliable way of understanding the static and dynamic aspects of these complex conformations in solution, avoiding difficulties which can occur in a crystalline environment, such as modified backbone conformations (e.g., B_I/B_{II} ratios) or hindered curvature.^{64–66} Free energy calculations also allow reasonable calculations of binding strengths and can expose unexpected cancellations in the factors contributing to binding.⁶⁷

Simulations remain an integral part of structure determination by NMR spectroscopy. One can note a considerable increase in the quality of such calculations, which now frequently include explicit solvent, reliable treatment of long-range electrostatics, and time-averaged constraints. Ligand–DNA complexes can still hold a number of surprises, especially as concerns dynamics, as evidenced by recent studies of a bis-intercalator which suggest that the intercalated moieties can flip over rapidly (1800 s⁻¹ at 36 °C) without unstacking.⁶⁸

Future possibilities are illustrated by an ab initio dynamics study of *cis*-platin binding to a dinucleotide monophosphate in aqueous solution. Even if such simulations currently span only a few picoseconds, there is no doubt that such approaches will give detailed insight into the chemistry of DNA interactions.⁶⁹

DNA—Protein Complexes

Both the rapid accumulation of X-ray and NMR data and increasing computer power have contributed to an increasing number of simulations of protein–DNA complexes. Depending on whether CHARMM or AMBER is

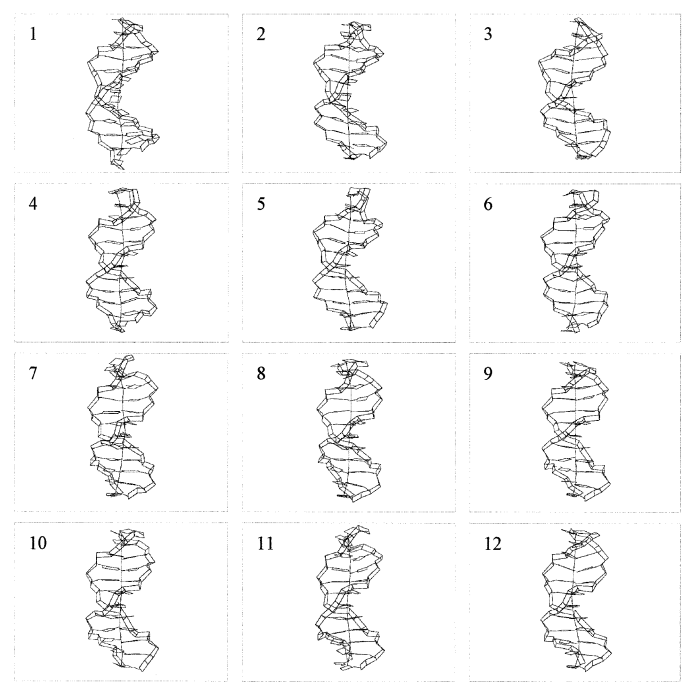


FIGURE 3. Conformational fluctuations of a B-DNA oligomer with an alternating GA sequence. The snapshots (100 ps intervals) from a simulation at 300 K using explicit solvent and counterions show axis and backbone fluctuations (E. Giudice, LBT Paris).

employed, the systems modeled generally solvate the complex within a water sphere and either use shift/switch technology for long-range interactions or use periodic boundary conditions and particle-mesh Ewald electrostatics. Recent studies belonging to the first category have dealt with complexes of EcoRI,^{70,71} the minor groove binding protein SRY,^{72,73} the 434cI repressor,⁷⁴ the glucocorticoid receptor,⁷⁵ and the non-sequence-specific protein HMG-D.⁷⁶ Examples of the second category include Hin-recombinase,⁷⁷ three zinc fingers belonging to TFIIIA,⁷⁸ and the Trp repressor.⁷⁹ Systems solvated with water spheres typically involve of the order of 20 000 atoms, while this number can rise to 40 000 when periodic

boundary conditions are used on large systems.⁷⁹ Simulations have reached the nanosecond range only in recent years, and we can assume, as for smaller systems, that longer simulations will reveal new categories of dynamic movement. With this caveat, present simulations nevertheless result in stable protein–DNA interfaces, reproduce most, or all, of the specific interactions observed experimentally, and, often, emphasize the role played by watermediated hydrogen bonding between the two partners.^{70,74,79} In the case of the zinc fingers of TFIIIA, a particularly impressive agreement was achieved with NMR data on the placement of solvating waters at and around the protein–DNA interface.⁷⁸

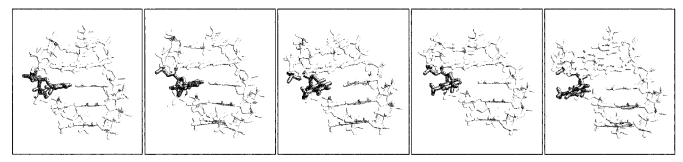


FIGURE 4. Induced base opening within B-DNA. Images show the conformational changes associated with moving thymine (bold) into the major groove of an oligomer with an alternating GA sequence.⁴⁰

Computer simulations are also powerful tools for analyzing the components of recognition processes. This can involve looking at structural adaptation upon binding^{70,72–74} or the consequences of point mutations.^{71,73,75} Another interesting approach used a simulation of the largely hydrophobic TBP–DNA complex to interpret hydroxyl radical cleavage data in terms of both sugar accessibility and fluctuations of the protein side chains contacting the DNA backbone.⁸⁰ A challenge which still lies largely in the future is the development of reliable protein–DNA docking procedures.⁷⁶

DNA Triple and Quadruple Helices

Three- and four-stranded DNA conformations continue to attract a great deal of attention because of their potential applications in artificial gene regulation and in cancer therapy. Dynamic simulations in this area have clearly helped in understanding the structure and stability of such structures.^{81–84} Recent refinements include studies of the stabilization of runs of GCC⁺ triplets by the replacement of some cationic cytosines with neutral amino tautomers^{85,86} and also studies of ion binding to quadruplexes, which can occur along the axis of the base tetrads, in the grooves, or in the loops of intramolecular complexes.^{83,87,88} Simulations are also being used actively in the design of modified nucleotides, such as 8-amino guanine, which reinforces Hoogsteen GC pairs,^{89,90} or of modified backbones which replace anionic phosphates with neutral linkers, as in PNA,⁹¹ or even with cationic groups, such as S-methylthiourea.⁹² Ligands which can specifically stabilize triplexes or quadruplexes, such as the family of disubstituted amidoanthraquinones, 93,94 are also being investigated.

RNA and RNA Complexes

Simulations involving RNA are accumulating rapidly, although the size, charge, and structural variety of these molecules continue to pose significant challenges. Several studies have addressed the dynamics of the basic motifs found in folded RNA, including mispairs such as flexible water-bridged UC pairs,⁹⁵ or coaxial stacking, where the absence of backbone continuity causes kinking and untwisting but does not enhance flexibility.⁹⁶ Studies of stem loops show that nanosecond dynamics maintain stable conformations but cannot sample extensively enough

to identify conformational substates.⁹⁷ Similar problems occur with bulged bases, where energy barriers prevent sampling of stacked and expelled base conformations during single trajectories.⁹⁸ Despite some courageous attempts,⁹⁹ many problems confront the use of simulations for structural predictions on larger fragments. One positive step is the estimation of relative free energies using MM/PBSA or MM/GBSA approaches, which has already correctly predicted RNA tetraloop conformations and, in the case of PB calculations, hairpin/duplex equilibria.^{100,101}

Despite sampling problems, it is impressive to see the size of the systems which are now being treated, the most striking being a stable simulation of $tRNA^{Phe}$ with more than 8000 waters—although some rearrangement in the core region raises the question, and the difficulties, of adding Mg^{2+} ions to such simulations.¹⁰²

Most protein-RNA studies have targeted the U1A-RNA system where structural data exist for both the U1 stem loop and the UTR internal loop binding sites. Simulations show that stable complexes are maintained on the nanosecond time scale in low salt^{103,104} and can be dissociated in high salt.¹⁰⁵ Recognition processes depend on initial electrostatic interactions but also involve the deformation and stiffening of the macromolecular partners and specific hydrogen bonding with bases. MM/PBSA studies have again been used successfully to estimate binding free energies and the consequences of site mutations.^{106,107} RNA-drug complexes are also attracting much attention. An interesting example involves a general model for binding aminoglycoside antibiotics, where the ammonium groups of the drug substitute for Mg²⁺ binding sites.¹⁰⁸ The readers are referred to a recent review for other examples.101

Water and Ion Distributions

Longer and more stable simulations have sparked a deeper analysis of the role of water and ions in stabilizing both DNA and RNA conformations. First-shell water residence times can extend to the nanosecond range, and this is also true for monovalent ions.^{109–112} Waters are found around the backbones and in both grooves of the helix, although long-lived hydration patterns are mainly confined to the narrower groove of both DNA and RNA and appear more sequence dependent in the case of DNA.^{109,113} Ions occupy both major and minor groove sites and are strongly coupled to DNA bending¹¹⁰ via major groove interactions with CpG steps and possibly to minor groove width in AT-rich tracts, although this conclusion depends on the definition of ion binding.^{110,111,114} First-shell hydration and ion binding have also been studied with respect to the $B \rightarrow A$ transition, where the electrostatics dominate, although many factors contribute to a subtle sequence-dependent free energy balance.^{110,114,115}

Conclusions

Stable multi-nanosecond trajectories of both helical and folded nucleic acids showing good agreement with experimental data are now within reach. The effects of base sequence and of environmental changes are also becoming accessible. Such studies provide an atomic-scale view of thermally induced fluctuations, allow the characterization of conformational substates and the location of transition pathways, and, in optimal cases, allow a detailed structural and energetic analysis which goes beyond current experimental possibilities. Given the polyelectrolyte nature of nucleic acids, particularly interesting developments involve a better understanding of how solvent and counterions influence the behavior of these environmentally sensitive macromolecules. Outstanding challenges concern the simulation of comparatively slow and complex processes such as the formation of nucleic acidprotein complexes or the folding of RNA domains, the simulation of much larger molecular systems, and the treatment of chemical reactions. Solving these problems will require effort on many fronts, including the definition of better continuum solvent models, the formulation of mesoscopic models (which implies decisions on crucial degrees of freedom and the development of reliable effective force fields), and the creation of interfaces between classical and quantum dynamics.

References

- Cheatham, T. E.; Kollman, P. A. Molecular dynamics simulation of nucleic acids. Annu. Rev. Phys. Chem. 2000, 51, 435–471.
- (2) Beveridge, D. L.; McConnell, K. J. Nucleic acids: theory and computer simulation, Y2K. *Curr. Opin. Struct. Biol.* 2000, 10, 182– 196.
- (3) MacKerell, A. D., Jr.; Nilsson, L. Nucleic Acid Simulations. In *Computational Biochemistry and Biophysics*; Becker, O., MacKerell, A. D., Jr., Roux, B., Watanabe, M., Eds.; Marcel Dekker: New York, 2001; pp 441–463.
- (4) Cheatham, T. E., III; Cieplak, P.; Kollman, P. A. A modified version of the Cornell *et al.* Force field with improved sugar pucker phases and helical repeat. *J. Biomol. Struct. Dyn.* **1999**, *16*, 845–862.
- (5) Foloppe, N.; MacKerell, A. D., Jr. All-atom empirical force field for nucleic acids: I. Parameter optimization based on small molecule and condensed phase macromolecular target data. J. Comput. Chem. 2000, 21, 86–104.
- (6) MacKerell, A. D., Jr.; Banavali, N. K. All-atom empirical force field for nucleic acids: II Application to molecular simulations of DNA and RNA in solution. *J. Comput. Chem.* 2000, *21*, 105–120.
- (7) Langley, D. R. Molecular dynamic simulations of environment and sequence dependent DNA conformations: The developments of the BMS nucleic acids force field and comparison with experimental results. J. Biomol. Struct. Dyn. 1998, 16, 487–509.
- (8) Bonvin, A. M. J. J.; Sunnerhagen, M.; Otting, G.; van Gunsteren, W. F. Water molecules in DNA recognition II: A molecular dynamics view of the structure and hydration of the *trp* operator. *J. Mol. Biol.* **1998**, *282*, 859–873.

- (9) Bonvin, A. M. J. J. Localisation and dynamics of sodium counterions around DNA in solution from molecular dynamics simulation. *Eur. Biophys. J.* 2000, *29*, 57–60.
- (10) Scott, W. R. P.; Hunenberger, P. H.; Tironi, I. G.; Mark, A. E.; Billeter, S. R.; Fennen, J.; Torda, A. E.; Huber, T.; Kruger, P.; van Gunsteren, W. F. The GROMOS biomolecular simulation program package. J. Phys. Chem. A **1999**, *103*, 3596–3607.
- (11) Feig, M.; Pettitt, B. M. Structural Equilibrium of DNA represented with different force fields. *Biophys. J.* **1998**, *75*, 134–149.
- (12) MacKerell, A. D., Jr. Observations on the A versus B equilibrium in Molecular dynamics simulations of duplex DNA and RNA. In *Molecular Modeling of Nucleic Acids*; Ceontis, N. B.; SantaLucia, J., Jr., Eds.; ACS Symposium Series 682; American Chemical Society: Washington, DC, 1998; pp 304–311.
- (13) Bevan, D. R.; Li, L. P.; Pedersen, L. G.; Darden, T. A. Molecular dynamics simulations of the d(CCAACGTTGG)₂ decamer: Influence of the crystal environment. *Biophys. J.* 2000, 78, 668–682.
- (14) Bosh, D.; Foloppe, N.; Pastor, N.; Pardo, L.; Campillo, M. Calibrating nucleic acids torsional energetics in force-field: insights from model compounds. *J. Mol. Struct.*: THEOCHEM 2001, 537, 283– 305.
- (15) Sagui, C.; Darden, T. A. Molecular dynamics simulations of biomolecules: Long-range electrostatic effects. *Annu. Rev. Biophys. Biomol. Struct.* **1999**, *28*, 155–179.
- (16) Norberg, J.; Nilsson, L. On the truncation of long-range electrostatic interactions in DNA. *Biophys. J.* 2000, 79, 1537–1553.
- (17) Wang, J.; Cieplak, P.; Kollman, P. A. How well does a restrained electrostatic potential (RESP) model perform in calculating conformational energies of organic and biological molecules? *J. Comput. Chem* **2000**, *21*, 1049–1074.
- (18) Mazur, A. K. Accurate DNA dynamics without accurate long-range electrostatics. J. Am. Chem. Soc. 1998, 120, 10928–10937.
- (19) Kimura, R. S.; Brower, R. C.; Zhang, C.; Sugimori M. Surface of active polarons: a semiexplicit solvation method for biomolecular dynamics. *J. Chem. Phys* **2000**, *112*, 7723–7734.
- (20) Bashford, D.; Case, D. Generalized Born models of macromolecular solvation effects. Annu. Rev. Phys. Chem. 2000, 51, 129– 152.
- (21) Orozco, M.; Luque, F. J. Theoretical methods for the description of the solvent effect in biomolecular systems. *Chem. Rev.* 2000, 100, 4187–4225.
- (22) Onufriev, A.; Bashford, D.; Case, D. A. Modification of the generalized Born model suitable for macromolecules. *J. Phys. Chem. B* 2000, 104, 3712–3720.
- (23) Jayaram, B.; Sprous, D.; Beveridge, D. L. Solvation free energy of biomacromolecules: Parameters for a modified generalized Born model consistent with the AMBER force field. *J. Phys. Chem. B* 1998, 102, 9571–9576.
- (24) Jayaram, B.; Liu, Y.; Beveridge, D. L. A modification of the generalized Born theory for improved estimates of solvation energies and pK shifts. J. Chem. Phys. **1998**, 109, 1465–1471.
- (25) Tsui, V.; Case, D. A. Molecular dynamics simulations of nucleic acids with a generalized Born solvation model. J. Am. Chem. Soc. 2000, 122, 2489–2498.
- (26) Srinivasan, J.; Cheatham, T. E., III; Cieplak, P.; Kollman, P. A.; Case, D. A. Continuum solvent studies of the stability of DNA, RNA, and phosphoramidate–DNA helices. *J. Am. Chem. Soc.* **1998**, *120*, 9401–9409.
- (27) Kollman, P. A.; Massova, I.; Reyes, C.; Kuhn, B.; Huo, S.; Chong, L.; Lee, M.; Lee, T.; Duan, Y.; Wang, W.; Donini, O.; Cieplak, P.; Srinivasan, J.; Case, D. A.; Cheatham, T. E., III. Calculating structures and free energies of complex molecules: Combining molecular mechanics and continuum models. *Acc. Chem. Res.* 2000, *33*, 889–897.
- (28) Lafontaine, I.; Lavery, R. Collective variable modeling of nucleic acids. Curr. Opin. Struct. Biol. 1998, 9, 170–176.
- (29) Elber, R.; Karplus M. Enhanced sampling in molecular dynamics: use of the time-dependent Hartree approximation for a simulation of carbon monoxide diffusion through myoglobin *J. Am. Chem. Soc.* **1990**, *112*, 9161–9175.
- (30) Mazur, A. K. Theoretical studies of the possible origin of intrinsic static bends in double helical DNA. J. Am. Chem. Soc. 2000, 122, 12778–12785.
- (31) Rudnicki, W. R.; Bakalarski, G.; Lesyng, B. A mezoscopic model of nucleic acids. Part 1. Lagrangian and quaternion molecular dynamics. *J. Biomol. Struct. Dyn.* 2000, *17*, 1097–1108.
- (32) Lebrun, A.; Lavery, R. Modeling the Mechanics of a DNA Oligomer. J. Biomol. Struct. Dyn. 1998, 16, 593–604.
- (33) Allemand, J. F.; Bensimon, D.; Lavery, R.; Croquette, V. Stretched and Overwound DNA forms a Pauling-like Structure with Exposed Bases. *Proc. Natl. Acad. Sci. U.S.A.* 1998, *95*, 14152–14157.
- (34) Schlick, T.; Beard, D. A.; Huang, J.; Strahs, D. A.; Qian, X. Computational challenges in simulating large DNA over long times.*Comput. Sci. Eng.* 2000, 38–51.

- (35) Gonzalez, O.; Maddocks, J. H. Extracting parameters for basepair level models of DNA from molecular dynamics simulations. *Theor. Chem. Acc.* 2001, 106, 76–82.
- (36) Simmerling, C.; Miller, J. L.; Kollman, P. A. Combined locally enhanced sampling and particle mesh Ewald as a strategy to locate the experimental structure of a nonhelical nucleic acid. *J. Am. Chem. Soc.* **1998**, *120*, 7149–7155.
- (37) Phillips, S. C.; Essex, J. W.; Edge, C. M. Digitally filtered molecular dynamics: The frequency specific control of molecular dynamics simulations. *J. Chem. Phys.* 2000, *112*, 2586–2597.
- (38) Stofer, E.; Chipot, C.; Lavery, R. Free energy calculation of Watson–Crick base pairing in aqueous solution. J. Am. Chem. Soc 1999, 121, 9503–9508.
- (39) Norberg, J.; Nilsson, L. Solvent influence on base stacking. Biophys. J. 1998, 74, 394–402.
- (40) Giudice, E.; Varnai, P.; Lavery, R. Energetic and conformational aspects of A:T base opening within the DNA double helix. *Chem. Phys. Chem.* 2001, *2*, 673–677.
- (41) Barsky, D.; Foloppe, N.; Ahmadia, S.; Wilson, D. M., III; MacKerell, A. D., Jr. New insights into the structure of abasic DNA from molecular dynamics simulations. *Nucleic Acids. Res.* 2000, 28, 2613–2626.
- (42) Cubero, E.; Sherer, E. C.; Luque, F. J.; Orozco, M.; Laughton, C. A. Observation of spontaneous base pair breathing event in the molecular dynamics simulation of a difluorotoluene-containing DNA oligonucleotide. J. Am. Chem. Soc. **1999**, *121*, 8653–8654.
- (43) Bandyopadhyay, D.; Bhattacharrya, D. Effect of neighboring bases on base-pair stacking orientation: A molecular dynamics study. *J. Biomol. Struct. Dyn.* 2000, *18*, 29–43.
- (44) Pastor, N.; MacKerell, A. D., Jr.; Weinstein, H. TIT for TAT: The properties of inosine and adenosine in TATA box DNA. J. Biomol. Struct. Dyn. 1999, 16, 787–810.
- (45) Derreumaux, S.; Fermandjian, S. Bending and adaptability to proteins of the cAMP DNA-responsive element: Molecular dynamics contrasted with NMR. *Biophys. J.* 2000, 79, 656–669.
- (46) Cheatham, T. E., III; Srinivasan, J.; Čase, D. A.; Kollman, P. A. Molecular dynamics and continuum solvent studies of the stability of polyG-polyC and polyA-polyT DNA duplexes in solution. *J. Biomol. Struct. Dyn.* **1998**, *16*, 265–280.
- (47) Trantirek, L.; Štefĺ, R.; Vorlickova, M.; Koca, J.; Sklenar, V.; Kypr, J. An A-type double-helix of DNA having B-type puckering of the deoxyribose rings. *J. Mol. Biol.* 2000, 297, 907–922.
- (48) Štefl, R.; Trantirek, L.; Vorlickova, M.; Koca, J.; Sklenar, V.; Kypr, J. A-like guanine-guanine stacking in the aqueous DNA dupluex of d(GGGGCCCC). J. Mol. Biol. 2001, 307, 513–524.
- (49) Winger, R. H.; Lledl, K. R.; Rudisser, S.; Pichler, A.; Hallbrucker, A.; Mayer, E. B-DNA's B₁ → B₁₁ conformer substate dynamics is coupled with water migration. *J. Phys. Chem. B* **1998**, *102*, 8935– 8940.
- (50) Pichler, A.; Rudisser, S.; Winger, R. H.; Liedl, K. R.; Hallbrucker, A.; Mayer, E. The role of water in B-DNAs B₁ to B₁₁ conformer substates interconversion: a combined study by calorimetry, FT-IR spectroscopy and computer simulation. *Chem. Phys.* 2000, 258, 391–404.
- (51) Winger, R. H.; Liedl, K. R.; Pichler, A.; Hallbrucker, A.; Mayer, E. B-DNA's B_{II} conformer substate population increases with decreasing water activity: I. A molecular dynamics study of d(CGC-GAATTCGCG)₂. J. Phys. Chem. B 2000, 104, 11349–11353.
- (52) Winger, R. H.; Liedl, K. R.; Pichler, A.; Hallbrucker, A.; Mayer, E. Helix morphology changes in B-DNA induced by spontaneous B_I → B_{II} substate interconversion. *J. Biomol. Struct. Dyn.* **1999**, 17, 223–235.
- (53) Soliva, R.; Luque, F. J.; Alhambra, C.; Orozco, M. Role of the sugar re-puckering in the transistion of A and B fomrs of DNA in solution. A molecular dynamics study. *J. Biomol. Struct. Dyn.* **1999**, *17*, 89–99.
- (54) Sprous, D.; Young, M. A.; Beveridge, D. L. Molecualr dynamics studies of the conformational preferences of a DNA double helix in water and ethanol/water mixture: Theoretical considerations of the A ↔ B transistion. J. Phys. Chem. B 1998, 102, 4658–4667.
- (55) Jayaram, B.; Sprous, D.; Young, M. A.; Beveridge, D. L. Free energy analysis of the conformational preferences of A and B forms of DNA in solution. *J. Am. Chem. Soc.* **1998**, *120*, 10629– 10633.
- (56) Sprous, D.; Young, M. A.; Beveridge, D. L. Molecular dynamics studies of axis bending in d(G₅-(GA₄T₄C)₂-C₅) and d(G₅-(GT₄A₄C)₂-C₅): Effects of sequence polarity on DNA curvature. *J. Mol. Biol.* **1999**, *285*, 1623–1632.
- (57) Sherer, E. C.; Harris, S. A.; Soliva, R.; Orozco, M.; Laughton C. A. Molecular dynamics studies of DNA A-tract structure and flexibility. J. Am. Chem. Soc. 1999, 121, 5981–5991.
- (58) Strahs, D.; Schlick, T. A-tract bending: Insights into experimental structures by computational models. J. Mol. Biol. 2000, 301, 643– 663.

- (59) Bostock-Smith, C. E.; Laughton, C. A.; Searle, M. S. Solution structure and dynamics of the A–T tract DNA decamer duplex d(GGTAATTACC)₂: Implications for recognition by minor groove binding drugs. *Biochem. J.* **1999**, *342*, 125–132.
- (60) De Souza, O. N.; Onrstein, R. L Inherent DNA curvature and flexibility correlate with TATA box functionality. *Biopolymers* **1998**, 46, 403–415.
- (61) Pardo, L.; Pastor, N.; Weinstein, H. Selective binding of the TATA box-binding protein to the TATA box-containing promoter: Analysis of structural energetics factors. *Biophys. J.* 1998, 75, 2411– 2421.
- (62) Pardo, L.; Campillo, M.; Bosch, D.; Pastor, N.; Weinstein, H. Binding mechanisms of TATA box-binding proteins: DNA kinking is stabilized by specific hydrogen bonds. *Biophys. J.* 2000, 78, 1988–1996.
- (63) Lankas, F.; Sponer, J.; Hobza, P.; Langowski, J. Sequencedependent elastic properties of DNA. J. Mol. Biol. 2000, 299, 695– 709.
- (64) Wellenzohn, B.; Winger, R. H.; Hallbrucker, A.; Mayer, E.; Liedl, K. R. Simulation of EcoRI dodecamer netropsin complex confirms class I complexation mode. *J. Am. Chem. Soc.* 2000, *122*, 3927– 3931.
- (65) Garcia-Nieto, R.; Manzanares, I.; Cuevas, C.; Gago, F. Bending of DNA upon binding of ecteinascidin 743 and phthalascidin 650 studied by unrestrained molecular dynamics simulations. J. Am. Chem. Soc. 2000, 122, 7172–7182.
- (66) Wellenzohn, B.; Flader, W.; Winger, R. H.; Hallbrucker, A.; Mayer, E.; Liedl, K. R. Structural flexibility of the d(CCAGTACTGG)₂ B-DNA decamer and its complex with two polyamides. *J. Phys. Chem. B.* 2001, *105*, 3135–3142.
- (67) Singh, S. B.; Kollman, P. A. Calculating the absolute free energy of association of netropsin and DNA. J. Am. Chem. Soc. 1999, 121, 3267–3271.
- (68) Gallego, J.; Reid, B. R. Solution structure and dynamics of a complex between DNA and the antitumor bisnaphthalimide LU-79553: Intercalates ring flipping on the millisecond time scale. *Biochemistry* 1999, *38*, 15104–15115.
- (69) Carloni, P.; Sprik, M.; Andreoni, W. Key steps of the cis-platin-DNA interaction: Density functional theory-based molecular dynamics simulations. *J. Phys. Chem. B* 2000, 104, 823–835.
- (70) Sen, S.; Nilsson, L. Structure, interaction, dynamics and solvent effects on the DNA-*Eco*RI complex in aqueous solution from molecular dynamics simulation. *Biophys. J.* 1999, 77, 1782–1800.
- (71) Sen, S.; Nilsson, L. Free energy calculations and molecular dynamics simulations of wild-type and variants of the DNA-*Eco*RI complex. *Biophys. J.* **1999**, *77*, 1801–1810.
- (72) Tang, Y.; Nilsson L. Interaction of human SRY protein with DNA: A molecular dynamics study. *Proteins: Struct., Funct. Genet.* 1998, 31, 417–433.
- (73) Tang, Y.; Nilsson, L. Effect of G40R mutation on the binding of human SRY protein to DNA: A molecular dynamics view. *Proteins: Struct., Funct. Genet.* **1999**, *35*, 101–113.
- (74) Harris, L. F.; Sullivan, M. R.; Popken-Harris, P. D. Molecular dynamics simulation in solvent of the bacteriophage 434 cl repressor protein DNA binding domain amino acids (R1–69) in complex with its cognate operator (OR1) DNA sequence. J. Biomol. Struct. Dyn. 1999, 17, 1–17.
- (75) Eriksson, A. A. L.; Nilsson, L. Structural and dynamical effects of point mutations in the recognition helix of the glucocorticoid receptor DNA-binding domain. *Protein Eng.* **1998**, *11*, 589–600.
- (76) Balaeff, A.; Churchill, M. E. A.; Schulten, K. Structure prediction of a complex between the chromosomal protein HMG-D and DNA. Proteins Struct., Funct. Genet. **1998**, *30*, 113–135.
- (77) Komeiji, Y.; Uebayasi, M. Change in conformation by DNA– Peptide association: Molecular dynamics of the Hin-Recombinasehixl complex. *Biophys. J.* **1999**, 77, 123–138.
- (78) Tsui, V.; Radhakrishnan, I.; Wright, P. E.; Case, D. A. NMR and molecular dynamics studies of the hydratation of a zinc finger– DNA complex. J. Mol. Biol. 2000, 302, 1101–1117.
- (79) Suenaga, A.; Yatsu, C.; Komeiji, Y.; Uebayasi, M.; Meguro, T.; Yamato, I. Molecular dynamics simulation of *trp*-repressor/ operator complex: Analysis of hydrogen bond patterns of protein– DNA interaction. *J. Mol. Struct.* **2000**, *526*, 209–218.
- (80) Pastor, N.; Weinstein, H.; Jamison, E.; Brenowitz, M. A detailed interpretation of OH radical footprints in a TBP–DNA complex reveals the role of dynamics in the mechanism of sequencespecific binding. *J. Mol. Biol.* 2000, 304, 55–68.
- (81) Špackova, N.; Berger, I.; Šponer, J. Nanosecond molecular dynamics simulations of parallel and antiparallel guanine quadruplex DNA molecules. J. Am. Chem. Soc. 1999, 121, 5519–5534.
- (82) Štefl, R.; Špackova, N.; Berger, I.; Koca, J.; Šponer, J. Molecular dynamics of DNA quadruplex molecules containing inosine, 6-thioguanine and 6-thiopurine. *Biophys. J.* 2001, 80, 455–468.

- (83) Meyer, M.; Steinke, T.; Brandl, M.; Sühnel, J. Density functional study of guanine and uracil quartets and of guanine quartet/metal ion complexes. J. Comput. Chem. 2000, 109, 109–124.
- (84) Špackova, N.; Berger, I.; Šponer, J. Structural dynamics and cation interactions of DNA quadruplex molecules containing mixed guanine/cytosine quartets revealed by large-scale MD simulations. J. Am. Chem. Soc. 2001, 123, 3295–3307.
- (85) Soliva, R.; Laughton, C. A.; Luque, F. J.; Orozco, M. Molecular dynamics simulation in aqueous solution of triple helix containing d(G.C.C) trios. J. Am. Chem. Soc. **1998**, *120*, 11226–11233.
- (86) Soliva, R.; Luque, F. J.; Orozco, M. Can G-C Hoogsteen-wobble pairs contribute to the stability of d(G.C-C) triplexes? *Nucleic Acids Res.* 1999, *27*, 2248–2255.
- (87) Chowdhury, C.; Bansal, M. Effect of coordinated ions on structure and flexibility of parallel G-quadruplexes: a molecular dynamics study. J. Biomol. Struct. Dyn. 2000, 18, 11–28.
- (88) Chowdhury, S.; Bansal, M. A nanosecond molecular dynamics study of antiparallel d(G)₇ quadruplex structures: Effect of coordinated cations. *J. Biomol. Struct. Dyn.* 2001, 18, 647–669.
- (89) Garcia, R. G.; Ferrer, E.; Macias, M. J.; Eritja, R.; Orozco, M. Theoretical calculations, synthesis and base pairing properties of oligonucleotides containing 8-amino-2'-deoxyadenosine. *Nucleic Acids Res.* **1999**, *27*, 1991–1998.
- (90) Soliva, R.; Garcia, R. G.; Blas, J. R.; Eritja, R.; Asensio, J. L.; Gonzalez, C.; Luque, F. J.; Orozco, M. DNA-triplex stabilizing properties of 8-aminoguamine. *Nucleic Acids Res.* 2000, 28, 4531– 4539.
- (91) Shields, G. C.; Laughton, C. A.; Orozco, M. Molecular dynamics simulation of a PNA.DNA.PNA triple helix in aqueous solution. *J. Am. Chem. Soc.* **1998**, *120*, 5895–5904.
- (92) Luo, J.; Bruice, T. C. Structural properties of hybrid triplex of polycation deoxyribonucleic S-methylthiourea (DNmt) strands with a complementary DNA strand, probed by nanosecond molecular dynamics. J. Biomol. Struct. Dyn. 2000, 17, 629–643.
- (93) Keppler, M. D.; Read, M. A.; Perry, P. J.; Trent, J. O.; Jenkins, T. C.; Reszka, A. P.; Neidle, S.; Fox, K. R. Stabilization of DNA triple helices by a series of mono- and disubdtitued amidoanthraquinones. *Eur. J. Biochem.* **1999**, *263*, 817–825.
- (94) Read, M. A.; Neidle, S. Structural characterization of a guaninequadruplex ligand complex. *Biochemistry* 2000, 39, 13422–13432.
- (95) Schneider, C.; Brandl, M.; Suhnel, J. Molecular dynamics simulation reveals conformational switching of water-mediated uracilcytosine base-pairs in an RNA duplex. *J. Mol. Biol.* 2001, 305, 659–667.
- (96) Schneider, C.; Suhnel, J. A molecular dynamics simulation study of coaxial stacking in RNA. J. Biomol. Struct. Dyn. 2000, 18, 345– 352.
- (97) Lahiri, A.; Nilsson, L. Molecular dynamics of the anticodon domain of yeast tRNA(Phe): codon–anticodon interaction. *Biophys. J.* 2000, 79, 2276–2289.
- (98) Sarzynska, J.; Kulinski, T.; Nilsson, L. Conformational dynamics of a 5S rRNA hairpin domain containing loop D and a single nucleotide bulge. *Biophys. J.* 2000, 79, 1213–1227.

- (99) Guo, J.-X.; D, I.; Daizadeh, I.; Gmeiner, W. Structure of the Sm binding site from human U4 snRNA derived from a 3ns PME molecular dynamics simulation. *J. Biomol. Struct. Dyn.* 2000, 18, 335–344.
- (100) Srinivasan, J.; Miller, J.; Kollman, P. A.; Case, D. A. Continuum solvent studies of the stability of RNA hairpin loops and helices. *J. Biomol. Struct. Dyn.* **1998**, *16*, 671–682.
- (101) Zacharias, M. Simulation of the structure and dynamics of nonhelical RNA motifs. *Curr. Opin. Struct. Biol.* 2000, 10, 311– 317.
- (102) Auffinger, P.; Louise-May, S.; Westhof, E. Molecular dynamics simulations of solvated Yeast tRNA^{Asp}. *Biophys. J.* **1999**, *76*, 50– 64.
- (103) Reyes, C. M.; Kollman, P. A. Molecular dynamics studies of U1A– RNA complexes. *RNA* 1999, 5, 235–244.
- (104) Tang, Y., Nilsson, L. Molecular dynamics simulations of the complex between human U1A protein and hairpin II of U1 small nuclear RNA and of free RNA in solution. *Biophys. J.* 1999, 77, 1284–1305.
- (105) Hermann, T.; Westhof, E. Simulations of the dynamics at an RNAprotein interface. *Nat. Struct. Biol.* 1999, 6, 540–544.
- (106) Reyes, C. M.; Kollman, P. A. Investigating the binding specificity of U1A-RNA by computational mutagenesis. J. Mol. Biol. 2000, 295, 1–6.
- (107) Reyes, C. M.; Kollman, P. A. Structure and thermodynamics of RNA-protein binding: Using molecular dynamics and free energy analyses to calculate the free energies of binding and conformational change. J. Mol. Biol. 2000, 297, 1145–1158.
- (108) Hermann, T.; Westhof, E. Aminoglycoside binding to the hammerhead ribozyme: A general model for the interaction of cationic antibiotics with RNA. J. Mol. Biol. 1998, 276, 903–912.
- (109) Auffinger, P.; Westhof, E. Water and ion binding around RNA and DNA (C,G) oligomers. *J. Mol. Biol.* **2000**, *300*, 1113–1131.
- (110) McConnell, K. J.; Beveridge, D. L. DNA Structure: What's in charge? *J. Mol. Biol.* **2000**, *304*, 803–820.
- (111) Hamelberg, D.; McFail-Isom, L.; Williams, L. D.; Wilson, W. D. Flexible structure of DNA: Ion dependence of minor-groove structure and dynamics. *J. Am. Chem. Soc.* 2000, *122*, 10513– 10520.
- (112) Štefl, R.; Koca, J. Unrestrained molecular dynamics simulations of [d(AT)⁵]₂ duplex in aqueous solution: Hydration and binding of sodium ions in the minor groove. J. Am. Chem. Soc. 2000, 122, 5025–5033.
- (113) Auffinger, P.; Westhof, E. Water and ion binding around r(UpA)¹² and d(TpA)¹² oligomers. Comparison with RNA and DNA (CpG)¹² duplexes. J. Mol. Biol. 2001, 305, 1057–1072.
- (114) Feig, M.; Pettitt, B. M. Sodium and chlorine ions as part of the DNA solvation shell. *Biophys. J.* 1999, 77, 1769–1781.
- (115) Feig, M.; Pettitt, B. M. Modeling high-resolution hydration patterns in correlation with DNA sequence and conformation. *J. Mol. Biol.* **1999**, 286, 1075–1095.

AR010023Y