Energy Landscape and Pathways for Transitions Between Watson-Crick and Hoogsteen Base Pairing in DNA

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Abstract

The recent discovery that Hoogsteen (HG) base pairs are widespread in DNA across diverse sequences, and positional contexts, could have important implications for understanding DNA replication and DNA-protein recognition. While evidence is emerging that the Hoogsteen conformation could be a thermodynamically accessible conformation of the DNA duplex, and provide a means to expand its functionality, relatively little is known about the molecular mechanism underlying the Watson-Crick to Hoogsteen transition. In this Perspective, we describe pathways and kinetics for this transition at an atomic level of detail, using the energy landscape perspective. We show that competition between the duplex conformations results in a double funnel landscape, which explains some recent experimental observations. The interconversion pathways feature a number of intermediates, with a variable number of Watson-Crick and Hoogsteen base pairs. The relatively slow kinetics, with possible deviations from two-state behavior, suggest that this conformational switch is likely to be a challenging target for both simulation and experiment.

DNA is one of the fundamental building blocks of life. Inside cells, DNA serves as the storehouse of genetic information, and transfers that information from one generation to the next, thereby providing a molecular basis for heredity and inheritance. DNA is not a static molecule, and its diverse range of functions are often accomplished via highly orchestrated motions of the helix, or large scale deviations from the canonical helical structure. The deformability of the DNA duplex is a key factor in determining nucleosome positioning, packaging of DNA within the cell nucleus, and sequence-specific recognition by proteins and ligands. There is a growing database highlighting the rich polymorphism exhibited by DNA, and strong dependence of the equilibrium structure of the double helix on sequence and environment. Repetitive sequences, in particular, have a high propensity to exhibit different duplex geometries, and assemble into higher order structures, such as triplexes and quadruplexes. Under specific environmental conditions, poly(G-C) sequences form Z-DNA, a left-handed duplex consisting of canonical Watson-Crick (WC)
Sequences rich in A-T repeats are known to exist in different polymorphic forms. For example, many crystal structures of A-T oligomers exhibit a B-type conformation, while poly(A-T) fibers can undergo a transition to C-type or D-type conformations, at low hydration and temperature conditions, respectively. Many of the alternate helical forms, and higher order structures, are often stabilized by base pairs that exhibit a different mode of hydrogen bonding from the original WC scheme. One such hydrogen bonding pattern, which was first reported by Hoogsteen in the 1960s from X-ray crystallography, results from the flipping of the purine base around the glycosidic torsion, changing an anti to syn orientation. The Hoogsteen (HG) base pairing mode is now known to be quite common for the A-T base pair. In fact, previous computational studies suggest that the WC pairing may not represent the optimal mode of interaction for A-T, and the HG orientation can sometimes be energetically more favorable. HG pairing is less common for the G-C base pair, as it requires protonation of the cytosine. The switch from the canonical WC to the HG orientation substantially modifies the chemical environment around the base pair in question, which can have major implications for DNA-protein recognition, damage repair, and replication. A prominent example includes the replication of DNA by Human DNA polymerase. In two separate studies, Satya Prakash and coworkers provide structural and biochemical evidence suggesting a replication mechanism dominated by HG base pairing. The authors argue that this mechanism helps to minimize interference during replication, circumventing lesions that may appear on the WC edge.

The realization that HG base pairing may provide an energetically viable route for key biological processes has fueled great interest in understanding the relative abundance of HG base pairs in diverse sequence and positional contexts, as well as the molecular mechanism underlying the WC to HG switch. A systematic survey of the DNA crystal structures deposited in the PDB database reveals that the repertoire of HG bonds is more extensive than previously thought, and the formation of HG base pairs can induce significant DNA
Figure 1. Representative structures for the Watson-Crick (WC) and Hoogsteen (HG) duplexes considered in this study. The WC base pairs are colored red, and the HG base pairs are colored blue. This coloring scheme is used in all the following figures. Also shown are the WC and HG orientations for the adenine-thymine base pair.
bending. In another study, Al Hashimi and coworkers employed NMR relaxation dispersion to show that the kinetics and HG base pair formation exhibit a rather complex dependence on sequence and position. They further predict that structurally stressed regions within DNA, such as kinks and turns, are likely to be hotspots for HG base pair formation.

Although HG pairing is more widespread within damaged or distorted DNA, as well as DNA-protein and DNA-antibiotic complexes during the last few years several reports of undistorted DNA and bare DNA duplexes containing HG base pairs have emerged. Subirana and coworkers were the first to report the crystal structure of a bare poly(A-T) duplex, in which all the base pairs exhibit HG base pairing. Interestingly, in solution, the poly(A-T) sequence exclusively formed the canonical B-DNA conformation. It was suggested that crystal packing forces may play a role in stabilizing the HG duplex. Although the overall topology of the HG duplex resembles the canonical B-form, it exhibits a narrower minor groove due to the shortening of the C1′-C1′ distances across the base pairs. The experimental results were corroborated in subsequent work by Orozco and coworkers. Using molecular dynamics simulations they suggested that both the HG and B forms are thermodynamically accessible states for the poly(A-T) duplex, and are remarkably similar in terms of structure and dynamics. In a followup study, Orozco and coworkers predicted that chimeras of HG duplexes embedded in B-DNA and HG-WC junctions are also thermodynamically stable under physiological conditions, and retain a strong memory of the shape, and the recognition properties of the original helices.

The work of Nikolova et al. provided a significant advance in understanding of the WC to HG transition. Employing NMR relaxation dispersion spectroscopy, they demonstrated that WC base pairs in a DNA duplex could reshuffle due to thermal motion, and transiently adopt the HG conformation. These ‘excited states’ have an equilibrium population of around 0.1%, and are short-lived, with a lifetime of approximately 5 ms. The authors speculate that the WC to HG equilibrium is intrinsically encoded within the DNA double helix, and could be exploited by cellular machinery to expand the functionality of DNA beyond what is accessible.
with WC base pairing. The conjugate peak refinement method (CPR)\textsuperscript{54} was used to provide atomistic insight into the WC to HG transition pathway. Based on a two-dimensional free energy surface, Nikolova \textit{et al.} predict that the anti to syn transition of the purine base involves base-flipping along the major groove, through small opening angles. In recent work, Pak and coworkers proposed that the WC to HG transition can occur via multiple routes, with contributions from both major and minor groove flipping.\textsuperscript{42} The switch from WC to HG conformations could also be triggered by chemical modifications. For example, Isaksson \textit{et al.}\textsuperscript{55} showed that substituting the O4' oxygen atom of the sugar ring to a CH\textsubscript{2} moiety results in a dynamic equilibrium between the WC duplex conformation, and a partial HG duplex exhibiting embedded Hoogsteen base pairs at the sites of chemical modification.

The WC to HG transition is characterized by high potential and free energy barriers, which makes it relatively difficult to obtain microscopic insight using unbiased simulations.\textsuperscript{23,39,42} Although previous computational studies\textsuperscript{39,42} suggest transition pathways, the use of predefined reaction coordinates restricts the configuration space. In previous contributions\textsuperscript{39,42,55} the WC to HG transition has been investigated in the context of a single target base pair, embedded in a B-DNA duplex. To the best of our knowledge, the present contribution may be the first to provide mechanistic and kinetic insight into the complete switch from the WC to the HG helix conformations including all the nucleic acid degrees of freedom.

In this Perspective, we discuss how the computational energy landscape framework,\textsuperscript{56} based on geometry optimization with post-processing using tools of statistical mechanics and unimolecular rate theory,\textsuperscript{57,58} can provide unprecedented mechanistic and kinetic insight into the complete switch in helix morphology from WC to HG. As a prototypical example, we consider a DNA duplex with the sequence (ATTAAT)\textsubscript{2}, whose crystal structure was reported in recent work by Campos and coworkers.\textsuperscript{59} This sequence adopts the HG conformation in the crystal lattice, and does not require additional ligands to maintain stability. Interestingly, no trace of the WC duplex was found under the experimental conditions.\textsuperscript{59} Here, we explain
this key result in detail in terms of the underlying energy landscape.

**The computational energy landscape framework:** In chemical physics, the potential energy landscape perspective serves as a unifying theme in the study of structure, thermodynamics and kinetics. For biomolecules, the landscape can be quite complex. The presence of competing structures often leads to features in the heat capacity profile, and to characterize interconversion pathways usually requires tools of rare event dynamics. For many practical applications, it is often sufficient to consider a coarse-grained description of the energy landscape, in terms of stationary points (i.e. potential energy minima and transition states). This coarse-graining eliminates the need for explicit dynamics: stationary points on the underlying landscape are located using geometry optimization techniques, and thermodynamic properties and kinetics are predicted using established tools (and approximations) of statistical mechanics and unimolecular rate theory. The resulting simulation framework is largely independent of the magnitude of the corresponding barriers and time scales, and unaffected by the presence of kinetic bottlenecks.

The potential energy landscape framework has been developed as both a conceptual and computational tool. Usually the first step in this analysis is basin-hopping global optimization, which has proved effective in structure prediction for a diverse range of systems, spanning atomic and molecular clusters, glass formers, and biomolecules. For quantitative prediction of thermodynamic properties we have described basin-sampling and nested-sampling as alternatives to parallel tempering. The discrete path sampling (DPS) technique is used to systematically construct databases of stationary points, corresponding to kinetic transition networks. This approach has been applied to problems where the presence of high energy barriers and broken ergodicity makes the exploration of the landscape, and hence the prediction of pathways and kinetics, challenging. Our efforts largely complement the recent advances in the area of rare event sampling made by other groups, which include innovative schemes based on explicit dynamics. Reviewing these methods is beyond the scope of this Perspective, and readers are referred elsewhere for such
details. Here, we focus on how various observable properties of interest can be extracted from stationary point databases constructed from DPS simulations.

The databases of stationary points constitute a kinetic transition network, which can be analyzed to extract global thermodynamics and kinetics using statistical mechanics and unimolecular rate theory. The superposition approach enables us to define global equilibrium thermodynamic properties, such as the heat capacity and equilibrium populations, in terms of contributions from local minima. Here we often employ the harmonic approximation, using normal mode analysis to estimate the canonical partition function corresponding to the catchment basin of each minimum. This approach provides us with local free energies, or values for states defined by lumping, and anharmonic as well as quantum corrections can be included. The ensemble of transition paths in the transition network, which are described in terms of interconnected minimum-TS-minimum triples, reflect the connectivity of the landscape and encode the global dynamics. Each minimum-to-minimum rate constant is usually determined employing harmonic transition state theory. However, other formulations, such as Kramer’s theory, can be used. The kinetically relevant pathways on the landscape, which make the most significant contributions to the global dynamics, are extracted from the network using Dijkstra’s shortest path algorithm with appropriate edge weights. Phenomenological reactant to product rate constants are obtained from the kinetic transition network using the new graph transformation (NGT) method. Compared to other approaches NGT scales better with temperature as well as the size of the network, and is robust in terms of numerical precision.

In experiments, the relaxation time scales usually correspond to transitions between ensembles, rather than isolated minima. To compare with experimental kinetics, we employ a self-consistent regrouping scheme, which exploits the contrasting time scales between local equilibration within states that can interconvert via low barriers, and the slower conformational transition of interest. In the regrouping, minima separated by free energy barriers below a certain threshold are lumped into a single macrostate. This approach is similar in
spirit to various kinetic clustering schemes that have been recently developed.\textsuperscript{115–118}

Disconnectivity graphs\textsuperscript{119–121} play a key role in conceptualizing energy landscapes, hence providing connections between apparently unrelated systems. The disconnectivity graph construction segregates the landscape into disjoint sets of minima, known as superbasins,\textsuperscript{119,120} at a regular series of threshold energies. The minima within each superbasin are mutually accessible at a given threshold, whereas transitions between superbasins involve higher potential or free energy barriers. In contrast to representations relying on low-dimensional projections of the energy landscape onto structural order parameters, principal components, or reaction coordinates,\textsuperscript{115–117,122–124} disconnectivity graphs capture the full complexity of the landscape, and preserve kinetic information. This construction has proved to be very insightful. For example, efficient self-organization is associated with a ‘palm tree’ structure in the disconnectivity graph,\textsuperscript{121} while double funnel landscapes,\textsuperscript{121,125} which support competing morphologies, are associated with broken ergodicity\textsuperscript{70,97,126–131} and rare event dynamics.\textsuperscript{62,78,132}

**Conformational ensembles of the WC and HG duplexes:** To explore the thermodynamic stability of the WC and HG DNA duplex conformations, molecular dynamics simulations in explicit water were first carried out at 300 K for 200 ns. The duplexes were modeled using a properly symmetrized version\textsuperscript{133} of the AMBER99bsc0 force field,\textsuperscript{134} employing the $\chi_{\text{OL4}}$ corrections for DNA, as suggested by Šponer and coworkers.\textsuperscript{135} Further details regarding the preparation of the initial structures and the simulation protocol are included in the supporting information.

The key results from the simulations are summarized in Figure 2. The evolution of the root-mean-square-deviation (RMSD) with respect to the initial conformation (supporting information, Figure S1) indicate that the WC and HG duplexes are thermodynamically stable, with no major structural changes occurring on the time scale of the simulation. In the WC duplex, the terminal adenines undergo a $\text{anti} \rightarrow \text{syn}$ transition about the glycosidic torsion angle at around 20 ns, after fraying out of the helix. Our observation is consistent
with previous experiments\textsuperscript{136,137} and simulations,\textsuperscript{138–140} which indicate that base pair fraying is particularly common at the helix terminals. After the $\chi$ torsional transition, the adenines flip back into the helix, establishing a HG base pair with the corresponding thymines. The HG base pairs at the duplex terminals remain stable throughout the rest of the trajectory, and do not revert back to the canonical WC orientation. It is likely that our specific choice of force field, in particular the $\chi$ parametrization, affects the $anti \leftrightarrow syn$ equilibrium, as observed in previous work.\textsuperscript{138,141} In contrast to the WC duplex, no fraying at the helix terminals was observed for the HG duplex, and all adenines exclusively sampled the $syn$ region.

To evaluate force field effects on the conformational dynamics of the DNA duplexes, we carried out additional molecular dynamics simulations with the parmbsc1 force field, a relatively new parametrization introduced by Orozco and coworkers.\textsuperscript{142} We find that the terminal adenines in the WC duplex undergo multiple transitions about the glycosidic torsion angle (supporting information, Figure S2). Between 10 and 40 ns the terminal adenines undergo $anti \rightarrow syn$ transitions, and switch to the Hoogsteen orientation. The bases revert back to the Watson-Crick geometry after around 100 ns. In contrast to the results for the $\chi$OL4 parametrization, the HG duplex undergoes significant distortion along the trajectory (supporting information, Figure S3). Strand slippage occurs at around 40 ns, and the terminal adenines flip to the WC orientation after 160 ns. In their original report describing the parmbsc1 force field,\textsuperscript{142} Orozco and coworkers suggest that such geometric distortions arise due to the metastable nature of the HG conformation. They also found that the equilibrium configuration for the HG duplex displayed significant deviation with respect to the experimental structure. As the insight provided in our study critically hinges upon a proper description of the experimental structure,\textsuperscript{59} our focus in this Perspective is on the $\chi$OL4 parametrization.

The distributions depicted in Figure 2 indicate that HG pairing requires a constriction of the C1$'$–C1$'$ distances across the corresponding base pairs. On average, the C1$'$–C1$'$ distances
Figure 2. Summary of the key results from the molecular dynamics simulations. Top panel: evolution of the $\chi$ torsions for the adenine bases corresponding to the HG duplex (blue lines), and the WC duplex (red lines). Middle panel: distribution of the $C1'-C1'$ distances across the base pairs for the HG duplex (blue), and the WC duplex (red). Bottom panel: the estimated B factors for the HG duplex (blue), and the WC duplex (red).
in the HG duplex are shorter by approximately 2 Å, compared to the WC duplex. In the
distribution corresponding to the WC duplex ensemble, a minor peak appears due to the
HG base pairing at the helix terminals.

The relatively high B factors indicate that the WC duplex is intrinsically more flexible
than the HG duplex. The entropic stabilization of the WC conformation under physiological
conditions, compared to the HG form, has been noted in earlier studies, and it is thought
to play a key role in modulating DNA-protein recognition.

Configurations from the molecular dynamics trajectories were saved every 10 ps, and
subsequently locally minimized after removing the solvent molecules and the counterions.
This quenching was performed to generate initial samples of minima from which to construct
the WC and HG duplex ensembles. The lowest potential energy minimum corresponding to
the WC ensemble includes HG base pairs at the helix terminals. Several low-lying minima
were also identified in which WC base pairs are found throughout the helix. The lowest
energy structure from this ensemble is destabilized by approximately 4 kcal/mol, relative to
the putative global minimum for the WC duplex. For the HG ensemble, the lowest energy
minimum exhibits a near perfect alignment with respect to the X-ray structure, as expected
from the limited conformational variability of the duplex on the MD time scale.

The local minima identified from MD were systematically used to construct stationary
point databases (kinetic transition networks), employing the discrete path sampling (DPS)
technique. The network was expanded using standard database refinement schemes, until
the phenomenological rate constant for the WC $\leftrightarrow$ HG transformation converged to within
an order of magnitude, with respect to the addition of new stationary points. The technical
details of the DPS procedure and the refinement process are included in the supporting
information.

In the following sections, we discuss the organization of the free energy landscape and
its emergent features, along with the interconversion mechanism and kinetics underlying the
WC $\rightarrow$ HG conformational switch.
Free Energy Landscape and Equilibrium Thermodynamics: The harmonic superposition approach,\(^94-97\) in conjunction with self-consistent regrouping,\(^83,114\) was used to construct the free energy landscape at 300 K; the corresponding disconnectivity graph is shown in Figure 3. To obtain this graph, we recursively lumped together minima separated by free energy barriers less than 4 kcal/mol. This regrouping threshold seems appropriate, as free energy graphs exhibiting qualitatively similar topological features were produced for a range of values around this selection, consistent with the ungrouped results (data not shown).

Each branch of the disconnectivity graph is colored according to the number of adenine nucleobases adopting the *syn* orientation in the corresponding free energy minimum. This particular order parameter enables us to separate the two prominent funnels on the landscape precisely, which involve duplex structures rich in HG and WC base pairs, respectively. The inability of geometrically defined order parameters to faithfully represent the complexity of the landscape is well documented, and here this failure is manifested by intermixing of different colored branches in some regions of the disconnectivity graph. In a previous study, Wales and Head-Gordon show that using the radius of gyration \(R_g\) as an order parameter for protein L can result in misclassification of folded and unfolded states, leading to artificially high folding rates.\(^{143}\) We stress that the use of an order parameter in our analysis is merely to aid visualization, and we calculate transition rates between states defined by exploit kinetic criteria based on recursive lumping.

The free energy global minimum corresponds to an ensemble of duplexes, in which the complementary bases are engaged in HG base pairing. Although this ensemble is largely homogeneous in terms of overall helicity, in some of the constituent structures one of the terminal bases is frayed out of the helix, adopting a *syn* orientation about the glycosidic torsion angle (see supporting information, Figure S4). The terminal adenines have a higher propensity to flip out, and we could identify only one structure within this ensemble in which a terminal thymine was displaced out of the helix.

Due to the coexistence of HG and WC base pairs within the DNA helix, a combinatorial
Figure 3. Free energy landscape obtained for a regrouping threshold of 4 kcal/mol at 300 K. The branches are colored according to the number of adenines in syn orientation (from red to blue, with red representing structures in which all the adenines are in the anti orientation, and blue representing structures in which all the adenines are in the syn orientation. Some representative snapshots from the different conformational ensembles are shown. The HG base pairs are colored in blue, and the WC base pairs in red, following the same coloring scheme as in Figure 1.
number of mixed duplex configurations could exist. In the stationary point databases, we identified around 90% of all the possible combinations, corresponding to stable duplex structures exhibiting base pairing between the complementary nucleotides. However, only a small subset of structures are associated with appreciable equilibrium occupation probabilities at 300 K (Figure 3, and supporting information, Figures S5–S9). Snapshots corresponding to some of these minima are shown superimposed on the disconnectivity graph.

Mixed duplexes occupying the low-lying regions of the blue funnel consist of either one or two WC base pairs (denoted 5HG+1WC, and 4HG+2WC, respectively), embedded within a HG base-paired helix. In particular, there is a significant population of HG duplexes exhibiting WC base pairing at the helix terminals. The mixed duplex ensembles within the blue funnel are separated from the HG duplex by low free energy barriers, and are likely to be late-stage intermediates during the WC $\rightarrow$ HG conformational switch. Ensembles of duplexes having an equal number of HG and WC base pairs (denoted as 3HG + 3WC) occupy the low-lying and intermediate regions of the landscape. Interestingly, structures in which the HG and WC base pairs are symmetrically distributed do not exhibit any particular energetic advantage over those in which the different types are more randomly distributed.

Duplexes exhibiting primarily WC base pairing occupy the bottom of the red funnel. The lowest free energy group within this funnel corresponds to an ensemble of duplexes in which the terminal base pairs exhibit a diverse range of conformations (Figure S10). The most prominent members of this group are structures in which either one or both the terminal base pairs adopt a HG orientation. In addition, we also identified structures in which one of the terminal bases (primarily adenines) is frayed out of the helix. Canonical WC duplex structures constitute the second-lowest free energy group within the red funnel. At a regrouping threshold of 6.8 kcal/mol, the two WC duplex ensembles described above merge into a single state, implying that the constituent structures, having different configurations for the terminal bases, would interconvert on a time scale of about 14.4 ns. The predicted time scale is consistent with our initial MD simulations, where the anti $\rightarrow$ syn torsional transitions
for the terminal adenines were observed at around the 20 ns mark in the trajectory. The presence of one or two embedded HG base pairs (1HG+5WC, and 2HG+4WC, respectively) in the interior of the helix destabilizes the WC duplex structure, and the corresponding minima are located in the intermediate region of the disconnectivity graph.

To compare the contrasting flexibilities of the HG and WC duplexes, we computed the log product of the normal-mode frequencies (\( \Lambda \)) for minima defining the corresponding free energy ensembles. Within the harmonic approximation, vibrational entropy is proportional to \(-\Lambda\), and therefore flexible structures would be associated with lower (more negative) \( \Lambda \) values. As shown in Figure 4, the \( \Lambda \) values for the WC duplex conformations are lower than most HG duplex structures, implying that the WC duplex is entropically favorable. Although the harmonic approximation does not account for well anharmonicity, which is implicitly included in MD simulations, the predicted trend in flexibilities is in line with the estimated B factors for the WC and HG duplexes.

The WC duplex is destabilized by around 10 kcal/mol compared to the HG duplex in terms of potential energy (supporting information, Figure S11). In contrast, the free energy difference between the two conformations at 300 K is approximately 3 kcal/mol. Hence en-
tropic contributions play a key role in modulating the stability gap (free energy difference) between the HG and WC conformations at physiological temperatures. As discussed above, the presence of soft normal mode vibrations makes the WC conformation favorable in terms of vibrational entropy. Additional stability for the WC form is due to landscape entropy (supporting information, Figure S12), which is determined by the density of high-lying local potential energy minima. The effect of the landscape contribution can be quantified by calculating the heat capacity excluding the vibrational degrees of freedom. The superposition approach includes both the landscape and vibrational contributions, but it is often not possible to characterize sufficient high-lying minima to provide a quantitative description of landscape entropy. In this case an alternative approach, such as the basin-sampling scheme, is required.

The interplay of enthalpy and entropy results in a heat capacity peak profile (Figure 4), which is typical for solid-solid type phase transitions observed in double funnel systems, such as the LJ$_{38}$ cluster. At low temperature, the blue funnel is thermodynamically favored, and relaxation down the energy landscape would preferentially lead to the HG duplex. Nonetheless, trapping in the competing red funnel, corresponding to the WC type duplexes, is likely to impede the overall kinetics. At higher temperatures, or under conditions in which entropy dominates, the landscape shifts in favor of the WC duplex. It is likely that the inherent limitations of the harmonic approximation, and incompleteness of the superposition sum, lead to an overestimation of the transition temperature for the HG$\leftrightarrow$WC switch. Both effects lead to a systematic underestimation of the entropy, which is greater for the WC ensemble, so a higher temperature is required to shift the equilibrium.

With the parmbsc1 parametrization, some reorganization of minima within the free energy basins is anticipated, but the key topological features should be preserved. To check this intuition we reoptimized some key minima in the red and blue funnels with the parmbsc1 force field. The probability distribution of relative energies (supporting information, Figure S13) reveals that, although the relative ordering of the basins is consistent with the
\(\chi_{OL4}\) parametrization, the energy gap between the HG and WC conformations is somewhat reduced. In addition, several distinct potential energy minima located with the \(\chi_{OL4}\) parametrization, collapse to a single minimum on the parmbsc1 surface (supporting information, Figure S14). This trend suggests that the parmbsc1 force field results in a smoother potential energy surface, which could have implications in terms of sampling efficiency. A similar trait was reported by Somani et al. in their benchmark study on alanine peptides, comparing the AMBER and CHARMM force fields.\(^{146}\) Furthermore, shifts in the relative potential energies of the minima, with respect to a fixed reference structure (in this case, the free energy global minimum identified with the \(\chi_{OL4}\) parametrization), provides clear-cut evidence of reorganization within the funnels. Details are provided in the supporting information. Interestingly, the magnitude of vibrational entropies for the different conformations, as determined by \(\Lambda\), is in quantitative agreement with the results obtained using the \(\chi_{OL4}\) parametrization (supporting information, Figure S15). As a final remark on force field effects, we note that a complete characterization of the key differences between the parmbsc1 and the \(\chi_{OL4}\) landscapes, would require systematic benchmarking, not only in terms of structure,\(^{147}\) but also thermodynamics and kinetics.

The topography of the free energy landscape, and the equilibrium thermodynamics, suggest that at low temperatures, or crystallization conditions that constrain the DNA helix, similar to those employed by Campos and coworkers,\(^{59}\) \((ATTAAT)\)\(_2\) preferentially adopts the HG conformation. The high free energy barriers separating the HG and WC duplex configurations will make it challenging to observe transitions between them, and probably preclude the observation of both conformations in a single experiment if the observation time scale is relatively short. The landscape analysis therefore also explains the results of Subirana and coworkers,\(^{47,51}\) who isolated different polymorphs for the AT repeat sequence, using X-ray crystallography and NMR.

The organization of the landscape provides key insight into the chain of events that occur during the WC\(\rightarrow\)HG conformational switch. The transition is likely to proceed in a
stepwise fashion, involving mixed duplex intermediates, with a varying number of HG and WC base pairs.

**Transition Pathways and Interconversion Kinetics:** The diversity of mixed duplex intermediates on the free energy landscape suggests that multiple routes between the HG and WC duplex conformations are feasible. For brevity, we consider only those discrete paths in the network that are associated with the highest statistical weights, which contribute maximally to the flux. The conformational switch takes place via two dominant mechanisms (Figure 5), differing primarily in the nature of the mixed duplex intermediates visited en route to the HG duplex.

Both mechanisms (paths P1 and P2) are essentially characterized by six elementary steps, in which a WC base pair switches to the HG orientation, resulting in a mixed duplex intermediate. Recursive enumeration analysis\(^{148}\) identifies P1 as the ‘fastest pathway’ in the network, while P2 appears around position one thousand. The pathways ranked in between, in terms of their contribution to the overall interconversion rate, correspond to minor variations of P1, with recrossing of some barriers. Hence we assign them to the same P1 mechanism and pathway ensemble. The highest energy transition states along the pathways correspond to structures in which the adenine that flips breaks away from its partner thymine, and moves out of the helix through small opening angles. Although the order of adenine flips is remarkably different in paths P1 and P2, in both cases base flipping is initiated at the TA step, which is known to stack quite poorly in the WC conformation.\(^{59,149}\) The transition from WC to the HG orientation for the central adenines results in significant kinetic bottlenecks, in contrast to the terminal adenines, which are more labile. The characteristic energy barriers for flipping of adenine bases within the center of the helix cover quite a broad range, from 8 to 18 kcal/mol. Our predictions are in excellent agreement with the relaxation dispersion experiments and CPR simulations of Nikolova et al.,\(^{39}\) which indicate that the transition state ensemble primarily features flipped-out states in which the purine base unstacks out of the helix via the major groove. A direct comparison with experiment, which provides insight
Figure 5. Transition pathways for the WC to HG conformational switch that make the largest contributions to the global dynamics. Some representative structures encountered at different stages of the transformation are also shown. Top panel: Pathway P1 where the order of base flipping is $A_4 \rightarrow A_2 \rightarrow A_1 \rightarrow A_3 \rightarrow A_5 \rightarrow A_6$. Bottom panel: Pathway P2 where the order of base flipping is $A_3 \rightarrow A_2 \rightarrow A_5 \rightarrow A_4 \rightarrow A_1 \rightarrow A_6$. The flipped-out adenine, and its partner thymine are colored green in the transition state structures. The WC base pairs are colored red, and HG base pairs colored blue.
at the single base pair level, is only feasible in this case because both pathways P1 and P2 involve step-wise base flipping, and no signatures of cooperative switching are found. In scenarios where the transformation pathways are dominated by multiple base flips occurring concurrently, a meaningful comparison with the experiment of Nikolova et al. would be more difficult.

In the kinetically relevant pathways, we do not find any clear-cut evidence of minor groove flipping, which was suggested by Pak and coworkers as a plausible mechanism of WC to HG transition for a single AT base pair within a DNA helix.\textsuperscript{42} In their analysis the transition pathways were based on a two-dimensional free energy surface. In such representations, projection errors are common, and the true complexity of the landscape can be masked. Energy barriers in orthogonal degrees of freedom may be averaged over, and as a result, the location of dividing surfaces (regions separating two basins) can be problematical. Hence low-dimensional projected surfaces may provide a misleading description of the kinetics; these issues have been discussed in a number of recent contributions.\textsuperscript{115,124} Our results for the free energy landscape, using recursive regrouping based on the calculated barriers, do not employ order parameters or projection, and all degrees of freedom are retained in the thermodynamic and kinetic analysis. Differences of interpretation may therefore originate from the alternative perspectives employed to extract the observables.

Pak and coworkers note that the free energy barrier for minor groove flipping is somewhat higher, compared to the major groove, during the initial stages.\textsuperscript{42} This situation is exacerbated when multiple flipping events are required to completely switch the topology of the duplex from the WC to the HG form. In our kinetic transition network, we do identify transition states in which adenine bases protrude out of the helix along the minor groove. However, the energy barriers connecting these transition states to the adjoining minima are substantial. Hence the corresponding pathways are energetically unfavorable, and are associated with relatively low statistical weights in the transition network. These results suggest that pathways proceeding via minor groove base flipping make little or no contribution to
the global dynamics.

![Arrhenius plot](image)

**Figure 6.** The variation of the rate constants corresponding to the conformational switch from WC to HG duplex ($k_{WC\rightarrow HG}$), and the reverse transformation ($k_{HG\rightarrow WC}$) with reciprocal temperature, are shown as solid lines with symbols. The fits to the Arrhenius equation, $k_{AB}(T) = A \exp\left(\frac{-E_a}{kT}\right)$ are depicted as dashed lines. From the fits, we obtain an activation energy, $E_a = 9.48 \pm 0.2$ kcal/mol, and a prefactor, $A = 2.19 \times 10^5$ s$^{-1}$ for the WC→HG transition. The reverse transformation has $E_a = 23.90 \pm 0.3$ kcal/mol, and $A = 8.69 \times 10^{13}$ s$^{-1}$.

The slow transition between the HG and WC duplex conformations was quantified by estimating the phenomenological rate constants. The WC→HG switch is associated with a rate constant ($k_{WC\rightarrow HG}$) of $2.78 \times 10^{-2}$ s$^{-1}$ at 300 K. The reverse transformation, HG→WC is orders of magnitude slower, with a rate constant ($k_{HG\rightarrow WC}$) of $2.70 \times 10^{-4}$ s$^{-1}$. The variation of $k_{WC\rightarrow HG}$, and $k_{HG\rightarrow WC}$ with temperature, depicted in the form of Arrhenius plots (Figure 6), provides further insight into the underlying kinetics. At 300 K, the Arrhenius prefactor associated with the WC→HG transition (Figure 6) is several orders of magnitude smaller than the time scale of typical molecular motions. Using the prefactor in standard thermodynamic expressions we obtain a negative value for the entropy of activation ($\sim -0.036$ kcal mol$^{-1}$ K$^{-1}$), which strongly suggests that the switch to the HG form is triggered via conformational restriction.
As temperature increases, $k_{HG\rightarrow WC}$ becomes higher than $k_{WC\rightarrow HG}$, indicating that entry into the WC funnel is an entropically driven process. The forward and backward rates are nearly equal at 365 K, which is close to the transition temperature predicted by the heat capacity profile. Interestingly, the temperature dependence of $k_{WC\rightarrow HG}$ becomes non-Arrhenius in the low temperature regime, which is reminiscent of protein and RNA folding.\textsuperscript{150–152} This deviation suggests that the conformational switch cannot be associated with a well defined transition state. At low temperatures, intermediates and kinetic traps that populate the free energy landscape are associated with distinct relaxation time scales. The transition from the WC to the HG conformation is likely to involve off-pathway excursions to trapped states, or periods in on-pathway intermediates, and a multi-state kinetic model seems more appropriate. At higher temperatures, the different basins merge, and an apparent two-state behavior is recovered. In contrast to $k_{WC\rightarrow HG}$, a small curvature is exhibited by $k_{HG\rightarrow WC}$ even at high temperatures, with the estimated rates being slightly lower than those predicted by the Arrhenius relationship. We speculate that this deviation could be due to the entropic stabilization of competing free energy minima close to the WC funnel, and the increased statistical weights of some slower routes in the pathway ensemble.

**Conclusions:** The potential energy landscape perspective has provided new insight into a wide range of diverse systems, including atomic and molecular clusters, soft and condensed matter, and biomolecules.\textsuperscript{56,66} In this Perspective, we have highlighted how coarse-graining the landscape in terms of stationary points (minima and transition states), provides a powerful alternative to conventional Monte Carlo and molecular dynamics simulations. Equilibrium thermodynamic and kinetic properties can be extracted from the underlying energy landscape using established tools and approximations of statistical mechanics and unimolecular rate theory. The inherently time-independent nature of the geometry optimization procedure makes it possible to probe biomolecular pathways that are not amenable to conventional simulations.\textsuperscript{153} Here, we have discussed one such application in detail, the conformational switch between the Watson-Crick and Hoogsteen forms of a DNA duplex.
The recent discovery that Hoogsteen base pairs are more widespread in DNA than previously thought, expands the repertoire of DNA functionality, with possible implications in DNA-protein recognition. Hence it is important to understand how DNA switches between different base pairing modes. Our results provide new insight at an atomic level of detail into the transition pathways between the WC and HG conformations of DNA, and the associated kinetics. We show that the competition between the two conformations results in a double funnel free energy landscape, which can be tuned as a function of temperature, or other environmental conditions. In particular, we can explain why complementary experiments usually identify only one of the polymorphs. The system exhibits a solid-solid-type heat capacity peak at low temperature, which is a characteristic property of multifunnelled landscapes. In the context of biology, such topographical features could naturally lead to multifunctionality, as we have recently suggested.

Our analysis reveals that multiple pathways exist between the HG and WC conformations, via mixed duplex intermediates. The transition state ensemble corresponds to base flipped-out states, in agreement with the experimental work of Nikolova et al. The interconversion kinetics exhibits non-Arrhenius behavior, suggesting that a two-state description is insufficient to describe the conformational switch. It will be interesting to see if any evidence of intermediate states and multistate kinetics emerge in future experiments. Sequence composition and oligonucleotide length could play a key role in modulating the transition pathways and kinetics. As shown by Orozco and coworkers, the entropic stabilization of the WC conformation increases with the chain length. On the other hand, energetic stabilization from base-stacking and hydrogen-bonding exhibits a strong linear dependence on length for the HG form. This interplay of competing interactions could provide a means to design a landscape that favors one form or the other, by tuning the environmental conditions. Furthermore, the energetics and kinetics of DNA base-flipping, which is a key step in the transformation between the HG and WC conformations, depends critically on the identity of the proximal bases. Al-Hashimi and coworkers previously suggested the appearance
of a late transition state in the conformational switch from WC to HG, based on the Ham-
mond postulate, and argued that the energetics exhibit a complex dependence on both the
sequence and position. Similarly, we speculate that oligonucleotide length and sequence
effects will play a role in modulating the barrier heights and interconversion rates for the
HG\rightleftharpoons WC conformations. These possibilities could be explored further in future studies.

In future work we envisage many new applications of the computational energy landscapes
methodology for biomolecules, particularly nucleic acids, whose polyelectrolyte nature, and
lower stability gap, makes conventional simulations challenging. Likely targets include
biophysical processes of fundamental importance, such as assembly mechanisms for higher
order RNAs, and conformational changes in riboswitches.

Supporting Information Available

Details regarding the preparation of initial structures, molecular dynamics protocol, and
discrete path sampling. The formalism describing the calculation of free energies, and global
dynamics. Figures showing the evolution of the RMSD for the WC and HG duplexes along
the MD trajectories, potential energy disconnectivity graph, benchmark calculations with
the parmbsc1 force field, plot of landscape entropy, representative structures constituting
the HG duplex ensemble, snapshots of mixed duplex intermediates exhibiting appreciable
occupation probabilities at 300 K, representative structures constituting the lowest energy
WC duplex ensemble. This material is available free of charge via the Internet at http://
pubs.acs.org/.

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