What Drives Proteins into the Major or Minor Grooves of DNA?

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The energetic profiles of a significant number of protein–DNA systems at 20 °C reveal that, despite comparable Gibbs free energies, association with the major groove is primarily an enthalpy-driven process, whereas binding to the minor groove is characterized by an unfavorable enthalpy that is compensated by favorable entropic contributions. These distinct energetic signatures for major versus minor groove binding are irrespective of the magnitude of DNA bending and/or the extent of binding-induced protein refolding. The primary determinants of their different energetic profiles appear to be the distinct hydration properties of the major and minor grooves; namely, that the water in the A+T-rich minor groove is in a highly ordered state and its removal results in a substantial positive contribution to the binding entropy. Since the entropic forces driving protein binding into the minor groove are a consequence of displacing water ordered by the regular arrangement of polar contacts, they cannot be regarded as hydrophobic.

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Structural and energetic characterizations of protein–nucleic acid complexes are important for a better understanding of the molecular interactions that govern transcriptional regulation. Of particular importance are the energetic profiles of DNA-binding domains (DBDs) interacting with their target recognition sites. DBDs are known to interact specifically with either the major or minor grooves of DNA, with binding-induced structural effects ranging from negligible perturbation of the B-DNA conformation to substantial distortions, such as bending and kinking. One can then ask if there are qualitative differences in the forces driving protein binding to the different grooves of DNA. Comparing the association constants of these two types of DBDs does not furnish a satisfactory answer, since both categories contain examples of stronger and weaker binding interactions. An answer to this question therefore requires a detailed analysis of the forces involved in the formation of the specific protein–DNA complexes. This assumes measurement of the association constant and determination of the Gibbs energy and its enthalpic and entropic components over a broad range of conditions, particularly temperature and ionic strength. In this review, we analyze the thermodynamic characteristics of protein binding to DNA published over the last several years. This overall consideration has revealed qualitative differences in the energetic signatures of binding to the minor and the major grooves of DNA, indicating intrinsic differences between the grooves.

Methodology

Studying the binding reaction at different temperatures is necessary, since the state of the reaction components depends on temperature, and this is particularly true for DNA-binding proteins, which in their unbounded state are usually not highly stable but partly unfolded even at ambient temperatures.1–7 Association with DNA results in their refolding (Figure 1(a)) and this has to be taken into account.
accounts in order to correlate the binding characteristics with structural information obtained for the complexes with fully folded proteins (Figure 1(b)). Correction for refolding is particularly important when comparing the binding of various proteins, which in their free unbound state differ somewhat in stability, i.e. the extent of unfolding under the conditions of the binding experiments. Correction for refolding can be made in practice using differential scanning calorimetric (DSC) data on the partial heat capacities of the reacting species over the temperature range in which unfolding of the free protein occurs.\textsuperscript{7,8} Figure 1(b) illustrates that correction for refolding linearizes the ITC measured binding enthalpy dependence on temperature and can change the magnitude of the enthalpy and its dependence on temperature considerably, i.e. the heat capacity effect of binding. The binding Gibbs free energies do not require correction for refolding, since the free energy of temperature-induced conformational changes of proteins at nearly ambient temperatures are usually small and may be neglected. Correspondingly, the binding entropy (derived from the difference between the Gibbs free energy and the corrected binding enthalpy) is corrected for protein refolding. Therefore, calorimetry has principal importance in studying the energetics of protein–DNA interaction.

Investigation of the effect of ionic strength is necessary in studying the interaction of such highly charged molecules as DNA and DNA-binding proteins in order to assess the contribution of electrostatic forces in this process.\textsuperscript{9–11} Since specific protein–DNA complexes are characterized by dissociation constants below micromolar, investigation of their interaction requires use of very dilute solutions and can be studied only by various optical methods sensitive for binding (e.g. fluorescence anisotropy or FRET titration). Analysis of the optically obtained binding isotherms (Figure 2(a)) permits determination of the binding constant ($K_a$) and its dependence on the ionic strength (Figure 2(b)). The dependences of the logarithm of the association constants, $\log(K_a)$, on the logarithm of the

\begin{equation}
\Delta H^{\text{observed}}(T) = \Delta H^{\text{corrected}}(T) - \Delta H^{\text{protein refolding}}(T)
\end{equation}

Figure 1. (a) The heat capacity functions of a typical DBD, the HMG box from LEF-1, its target DNA duplex and their complex, determined by differential scanning calorimetry (DSC). This shows that the protein starts to unfold from very low temperatures but on association with DNA it refolds and forms a stable complex that dissociates and unfolds cooperatively at about 62 °C. The heat of protein refolding at any given temperature can be determined by integrating the differences between the summed heat capacity of the free protein and DNA (blue dot-dash line) and the observed heat capacity of the complex (black continuous line). (b) The observed enthalpy of association of the HMG box from LEF-1 with its target DNA measured by isothermal titration calorimetry (ITC) and the function corrected for heats of protein refolding upon binding; the corrected function corresponds to the enthalpy of association of the fully folded DBD with the DNA.\textsuperscript{4,5}

Figure 2. (a) Binding isotherms of the HMG box from Lef-1 (Lef86) and its truncated form lacking the eight-residue C-terminal extension (Lef79), measured by fluorescence anisotropy titration.\textsuperscript{4} These provide the value of the association constant, $K_a$ and thus the Gibbs free energy of association, $\Delta G^{\text{obs}}(T) = -RT \ln(K_a)$. Using the ITC-measured enthalpy of association, after correction for refolding, one can then determine the binding entropy factor, $T\Delta S^{\text{obs}}(T) - \Delta H^{\text{corrected}}(T) = \Delta G^{\text{corrected}}(T)$. (b) The dependences of the logarithm of the association constants, $\log(K_a)$, on the logarithm of the salt concentration is described effectively by the linear equation: $\log(K_a) = \log(K_{\text{salt}}) - N \log[\text{Salt}]$. The first term results from the non-electrostatic interactions between protein and DNA, and the second salt-dependent term represents the electrostatic component of protein–DNA interaction. Extrapolating this function to $\log[\text{Salt}]=0$, where the second term becomes zero, one can determine the non-electrostatic component of the Gibbs energy of association.

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The first term results from the non-electrostatic interactions between protein and DNA, and the second term reflects the entropy of mixing the released counterions with the ions in bulk solution. This salt-dependent second term, which does not depend on temperature, determines the electrostatic component of protein–DNA interactions. Extrapolating the log\(K^\theta\) function to log [Salt] = 0, where the second term vanishes, one can determine the non-electrostatic component of the Gibbs energy of association, \(\Delta G_{\text{nel}} = -2.3 \ RT \ \log \ (K_{\text{nel}})\) and then the electrostatic component of the Gibbs energy of association as \(\Delta G_{el} = \Delta G - \Delta G_{\text{nel}}\). The latter is equivalent to \(-T \Delta S_{el}\), since the enthalpy of electrostatic interactions is zero. The non-electrostatic association entropy factor is then obtained from the relation:

\[
T \Delta S_{\text{nel}} = T \Delta S^0 - T \Delta S_{el}
\]

**Thermodynamic Parameters of Protein–DNA Interactions**

The energetics of association of various DNA-binding proteins with their target DNAs have been studied calorimetrically by many groups over the years. For the present analysis, we have selected data obtained under similar conditions (20 °C, near neutral pH, 100 mM NaCl) and analyzed by the same approach; namely, correcting for refolding effects and resolving the electrostatic and non-electrostatic components of the binding characteristics, as outlined in Figures 1 and 2. All these data are illustrated in Figure 3 and summarized in Table 1 in Supplementary Data.

Inspection of the data reveals no dramatic differences in the Gibbs free energies of protein binding to the major and minor grooves of DNA (Figure 3(a)). Therefore, the systematic qualitative difference in binding enthalpies for the two grooves is particularly striking (Figure 3(b)). Specifically, the enthalpies of binding to the minor groove are always positive, while the enthalpies of binding to the major groove are invariably negative. Positive binding enthalpies have been reported by other authors for minor groove DBDs, including: Sso7d,20 a thermophilic TATA box-binding protein,15,22 and Sac7d.24,25 Conversely, negative binding enthalpies have been reported for other major groove DBDs including: the λcI repressor,26 c-Myb,23 trp repressor,18 MM17, a single-chain dimer of transcription factor MASH-1,20 and the double homeobox Oct.21

Since the Gibbs free energies of binding at 20 °C do not vary greatly, the enthalpy differences are essentially balanced by entropic factors. Indeed, the entropic contribution (i.e. \(T \Delta S\)) of minor groove binding is significantly larger than that of major groove binding (Figure 3(c)). Positive entropies have been reported for binding of small molecules to the minor groove of DNA, in contrast with drug intercalators, for which binding is typically enthalpy-driven.31

The initial observation of a different sign in the enthalpies first arose when comparing the unfavorable enthalpy of the minor groove DBD from Sox-5 with the favorable enthalpies measured for major groove DBDs, a finding that was interpreted at the time as reflecting the work required to bend DNA and has been corroborated by other groups. Recent studies on additional protein–DNA systems now suggest that this interpretation is not always warranted. In fact, attempts to establish a correlation for the magnitudes of DBD-induced bending reveal that the enthalpies do not simply scale with the bending angles, as seen from Figure 3(d). For example, Lef-86 bends DNA by 117° yet exhibits a binding enthalpy of only +10 kJ/mol.4 The AT-hook and Hoechst interactions are even more striking, in that neither bends DNA, although both bind deeply in the minor groove with association enthalpies of +15 kJ/mol and +30 kJ/mol, respectively. Studies of several Sac7d mutants that differ in the number of intercalating residues in the DNA minor groove demonstrate clearly that, while such binding events induce a range of DNA bending angles (52–68 deg), their association enthalpies are similar (~+35 kJ/mol) at 20 °C.25 The finding that binding enthalpies do not obviously scale with bending angles is evident when comparing the association of sequence-specific HMG box DBDs with sub-optimal (as opposed to cognate) DNA sequences: an overall reduction in bending angle is accompanied by an increased positive binding enthalpy.4

These comparisons suggest that the work required to bend DNA cannot represent the main source of the positive enthalpy associated with protein/ligand binding to the minor groove. The only other possible source of such a large positive enthalpy is the removal of water and/or specifically bound ions from the protein/DNA interface. Specifically bound ions have been observed crystallographically in the A+T-rich minor groove of DNA in flash-frozen samples at very low temperatures. More appropriate to the present solution conditions, NMR measurements conducted at ambient temperatures clearly demonstrate that the occupancy of DNA by sodium ions is not high, comprising about 0.5 ion per AT-tract in 0.2 M NaCl at 4 °C, and decreasing to 0.2 ion per AT-tract as the temperature is increased to 27 °C.35 Thus, removal of these ions might be associated with a positive enthalpy. However, even if we assume that the enthalpy of removing a single ion is similar to that of releasing one water molecule (i.e. approximately 6 kJ/mol), the Na⁺ occupancy of the AT-tract is so low that ion displacement into the bulk solution cannot explain the large positive enthalpy of protein binding to the minor groove. It appears, therefore, that unlike what occurs in the major groove, the enthalpy of dehydrating the
minor groove far exceeds the favorable enthalpic contributions from newly formed binding interactions. This conclusion is supported strongly by the observed protein–DNA binding entropies, particularly their non-electrostatic component.

Components of the Binding Entropy

The binding entropy consists of electrostatic and non-electrostatic components, the first of which arises from the release of counterions from the DNA phosphate groups and their mixing with ions into bulk solution.\cite{9,10} Accordingly, this electrostatic component enhances the affinity but not the specificity of binding, which is determined by the non-electrostatic component. This non-electrostatic component of the binding entropy (corrected for protein refolding) includes a negative contribution resulting from the decrease in translational/rotational degrees of freedom,\cite{38} and the changes in hydration of the reaction components. The non-electrostatic entropy resulting from the dehydration of contacting groups is positive, but a negative contribution might arise if binding results in incorporation of water at the newly formed interface.

Figure 4 illustrates that binding to the minor groove proceeds with large positive non-electrostatic entropies, while binding to the major groove is characterized in most cases by small negative non-electrostatic entropies. Since the entropies associated with the decrease of translational/rotational freedom are similar for both categories, the only possible explanation for the large differences in the, corrected for refolding, non-electrostatic entropies of binding to the two grooves of DNA is the nature of their hydration.
There is indeed evidence that water is incorporated into the interfaces of certain homeodomain/DNA and IRF/DNA complexes. However, the value of the negative non-electrostatic entropy and enthalpy in most of these cases is rather small for assuming that water incorporation is associated with complex formation: the bound water found in these complexes is likely to be bound also in the free state of the protein and DNA. The only cases among those considered for which water ordering seems to occur upon complex formation are the bZIP/AP-1 and IRF1 complexes, both of which proceed with a large negative enthalpy and entropy (Figure 3(b) and (c)). In the case of the minor groove, judging from the predominance of AT sequences among its DBD binding sites, and bearing in mind the absence of the 2-amino group in adenosine (relative to guanosine), one might argue that this groove is more apolar than the major groove. Furthermore, it is known that apolar groups promote water ordering. It is therefore tempting to assume that the large positive non-electrostatic entropy of protein binding to the minor groove results from the dehydration of apolar groups on forming hydrophobic contacts between protein and DNA. This hypothesis, however, does not explain the large positive enthalpy of binding to the minor groove, since the enthalpy of formation of hydrophobic contacts is known to be close to zero at ~20 °C. Analysis of the heat capacity effects of association shows that it is not the formation of hydrophobic contacts that gives rise to the large positive non-electrostatic entropy.

**Heat Capacity Effects of Protein–DNA Association**

The heat capacity effect of protein–DNA association is invariably negative, as illustrated for the binding of c-Myc, c-Myb and Oct1 to the major groove, as well as for the binding of TBP, Sso7d and Sac7d to the minor groove. However, the net heat capacity effect due to binding-induced surface dehydration can be evaluated properly only after correction for protein refolding: this correction can result in changes in the magnitude of ΔC_p (see Figure 1(b)) and even in its sign. For comparative analysis of the role of binding-induced dehydration, the observed heat capacity effects have been normalized per Å² of interfacial surface area, since this varies substantially among the different protein–DNA complexes. Figure 5 presents surface-normalized ΔC_p values for the association of folded DBDs lacking long, charged extensions (thereby excluding NHP6A, HMG-B1, HMG-D100, and Lef86 from the analysis), since such extensions contribute a large positive ΔC_p component arising from multiple electrostatic interactions with DNA phosphate-groups. The DNA-binding helices of the GCN4 bZIP dimer are also highly charged, likewise resulting in a substantial positive contribution to ΔC_p, and consequently have been removed from the analysis depicted in Figure 5.

At first glance, an overall negative heat capacity change for both minor and major groove DBDs seems to support the assumption that in both cases protein binding results in extensive dehydration of apolar rather than polar groups, since it is well...
known that the dehydration of apolar groups results in negative heat capacity effects, in contrast to polar groups, for which the heat capacity effects of dehydration are positive. Of particular interest, however, are the individual contributions from protein and DNA to the overall measured heat capacity effect of complex formation. The heat capacity changes associated with the dehydration of proteins are well described by changes in the water-accessible surfaces areas ($\Delta ASA$) of polar and apolar groups. However, this approach cannot be used to estimate the heat capacity effects from the DNA, since the heat capacity changes associated with the dehydration of DNA groups have not been rigorously characterized to date. The only way therefore to derive the heat capacity effects of DNA dehydration is to calculate the contribution from protein dehydration and subtract it from the observed total $\Delta C_p$ (see Table 2 in Supplementary Data).

The resultant partitioned $\Delta C_p$ values presented in Figure 5 for several complexes of known structure reveal a number of unprecedented findings. The heat capacity change due to dehydration of the DBDs is always negative but its overall magnitude is significantly larger for minor groove DBDs, reflecting the fact that the interacting surface of minor groove DBDs is more hydrophobic than that of major groove DBDs. The contributions of DNA are not similar to those of the proteins, a finding that appears counterintuitive considering the complementarity of their contacting surfaces: the heat capacity effect of dehydrating the major groove is much larger than that of the protein, whereas the opposite effect is observed for minor groove binding. Thus, despite the fact that the minor groove is more apolar than the major groove, the surface normalized heat capacity effect of its dehydration is substantially less than that of the major groove (see Table 2 in Supplementary Data). It appears, therefore, that the water displaced from the minor groove has a substantially lower heat capacity than that of bulk water. This would be the case if such water is in an ice-like state, since the heat capacity of ice is about half that of liquid water.

**Hydration of the DNA Grooves**

The presumption that water ordering in the A+T-rich minor groove differs significantly from water hydrating the major groove has been a subject of discussion for quite some time. The presence of a spine of well-ordered water molecules in the minor groove of A+T-rich DNA sequences has been derived from high-resolution X-ray crystallography and neutron diffraction studies. The minor groove is unusually narrow in AT stretches and a primary shell of water molecules runs across the groove, bridging acceptor sites on adjacent AT/TA base-pairs (Figure 6). A secondary shell of water molecules runs along the groove, donating hydrogen bonds to the primary shell of oxygen atoms that assume the tetrahedral coordination characteristic of ice. There is further evidence of an outer spine of third and fourth-shell water molecules in a pattern of fused hexagons. Exchange of the bound water with bulk solvent has been studied by NMR, and its residence time is longer than that of water residing in the major groove. Studies of the interaction of bisbenzimide with DNA reveal that the dielectric constant in the minor groove is less than that estimated for the major groove, consistent with stronger hydrogen bonding in the minor groove.

In the major groove, the distribution of hydrogen bond donors and acceptors exhibits greater irregularity than in the minor groove, and no regular water superstructure has been identified.

Based on the assumption that the water hydrating the minor groove of A+T-rich DNA is more ordered than the water hydrating the major groove, its

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**Figure 6.** Display of primary (blue) and secondary (yellow) layers of the spine of water in the minor groove of the X-link dodecamer CGCAATTCCGC generated from the coordinates of NDB accession number bd0008. A similar structure has been obtained for the identical DNA sequence.
Conclusions

Inspection of the thermodynamic data summarized herein on structurally characterized protein–DNA and drug–DNA complexes reveals a clear qualitative difference in the energetic signatures of binding to the major and minor grooves that appears to be a consequence of their distinct hydration properties. It is important to note that protein binding to the minor groove, as well as the binding of small molecules to this groove,61 occurs principally at A+T-rich DNA sequences, the latter corresponding to those very regions in which water ordering is the most prevalent. The overall consequence is that minor groove binding is normally driven by the very large entropy of releasing the ordered water, despite an unfavorable enthalpy. However, this does not represent a hydrophobic force, since water ordering in the A+T-rich minor groove is determined not by the apolar groups of the DNA but by the regular arrangement of its polar groups that stabilize the ice-like organization of the water in this groove.36,53–55 Further investigation of the state of hydration in the DNA grooves is thus of primary importance for a more complete understanding of the mechanisms of protein binding to DNA.

Acknowledgements

This research was supported by National Institutes of Health Grants GM48036-12 (P.L.P.) and CA47995 (K.J.B.), and National Science Foundation Grant MCB0519381 (P.L.P.).

Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2006.09.059

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_Edited by P. Wright_

(Received 30 June 2006; received in revised form 21 September 2006; accepted 22 September 2006) Available online 27 September 2006