Predicting Transcription Factor Specificity with All-atom Models

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Accessibility
Predicting transcription factor specificity with all-atom models

Sahand Jamal Rahi¹, Peter Virnau²*, Leonid A. Mirny¹,³ and Mehran Kardar¹

¹Department of Physics, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA, ²Staudinger Weg 7, Institut für Physik, 55099 Mainz, Germany and ³Harvard-MIT Division of Health Sciences and Technology, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA

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ABSTRACT

The binding of a transcription factor (TF) to a DNA operator site can initiate or repress the expression of a gene. Computational prediction of sites recognized by a TF has traditionally relied upon knowledge of several cognate sites, rather than an ab initio approach. Here, we examine the possibility of using structure-based energy calculations that require no knowledge of bound sites but rather start with the structure of a protein–DNA complex. We study the PurR Escherichia coli TF, and explore to which extent atomistic models of protein–DNA complexes can be used to distinguish between cognate and noncognate DNA sites. Particular emphasis is placed on systematic evaluation of this approach by comparing its performance with bioinformatic methods, by testing it against random decoys and sites of homologous TFs. We also examine a set of experimental mutations in both DNA and the protein. Using our explicit estimates of energy, we show that the specificity for PurR is dominated by direct protein–DNA interactions, and weakly influenced by bending of DNA.

INTRODUCTION

Binding of cognate sites of DNA is central to many essential biological processes. Most of DNA-binding proteins have the ability to recognize and tightly bind cognate DNA sequences (sites). To find sites bound by a particular DNA-binding protein, one needs to calculate the free energy of binding for the protein and possible DNA sites and then select sites that provide sufficiently low-binding energy. A widely used approach to find sites for a DNA-binding protein is to assume a form of the energy function, infer its parameters and to then calculate the energy for all sites in a genome. To infer parameters one needs to have a set of known sites bound by the protein. Given these known sites, the parameters are inferred using either a widely used Berg–von Hippel approximation (1), or by other recently proposed methods (2–4). This constitutes a physical basis for many widely used bioinformatics techniques that rely on a particular form of the energy function known as a position-specific weight matrix (PWM).

All these methods require a priori knowledge of the sites (or at least longer sequences containing these sites) bound by the protein. These data are available for only a small number of DNA-binding proteins. For many DNA-binding proteins, however, their sequence of amino acids is well known. Sufficiently high-evolutionary conservation of DNA-binding domains, and the availability of crystal structures for many of them, makes it possible to construct 3D models for a broad range of DNA-binding proteins. Can such protein structures be used to predict sites recognized by a DNA-binding protein? The basic procedure for structure-based methods is to compute the binding energy of the protein–DNA complex. The structure of the complex for an arbitrary DNA sequence can be modeled by replacing (‘mutating’) the DNA sequence in the protein structure containing its cognate site, followed by energy minimization and/or molecular dynamics (MD) to allow the protein–DNA complex to adjust to the new DNA sequence. After several minimization steps, the interaction energy can be calculated using either standard molecular mechanics force fields like AMBER (5) or CHARMM (6) with an implicit solvent, or a knowledge-based force field optimized for the particular complex (7).

Several recent studies have significantly elaborated upon the above procedure. Lafontaine and Lavery (8), for example, pioneered a very efficient process termed ADAPT in which they replace the DNA in the structure by a ‘multicopy’ or ‘average’ piece of DNA. From this, only the energy of the unbound DNA must be subtracted. The unbound protein
energy is the same for all DNA sequences and hence irrelevant for comparisons. This approach is so efficient that all possible sequences \(4^N\) for \(N\) bases for short DNA operator sites can be evaluated. Their results successfully identify the experimental consensus sequence for a variety of DNA-binding proteins \((9,10)\), and the ordering of binding free energies for DNA point mutations in several complexes \((9)\). In this context, it was also noted that the actual binding energy computed via minimizations is incorrect and cannot be compared to experiments quantitatively.

Endres et al. \((11)\) allowed protein side chains to explore rotamer conformations in their study of Zif268. Interestingly, the agreement with experiments becomes worse when rotamers are considered, which points to a potential bias of the approach towards sequences similar to the one on which the underlying experimental structure is based. Morozov et al. \((12)\) using energy measurements as well, keep their structures rigid or allow them to relax and compare the two approaches. However, instead of considering their binding energies to be approximately equal to free energies as we do, they fit their energies to a few experimentally known free energies. They assign different weights to the energies involved, e.g. the Lennard-Jones or the electrostatic energy, and optimize the weights so that the sum matches the free energy. They proceed to study several transcription factors (TFs) and even find consensus sequence logos for two TFs whose structures they construct by homology modeling. In recent work, Donald et al. \((7)\) focus on direct protein–DNA interactions. They study and compare a number of potentials and propose some that outperform the standard Amber potential. All these efforts represent pioneering work in the emerging field of structure-based predictions of TF specificity.

Here, we explore whether widely available MD force fields can be used to calculate the binding free energy from all-atom models of the protein–DNA complex. In contrast to some of the previous studies, we: (i) assess the power and limitations of the method in dealing with the roughly \(10^6\) decoy sites of bacterial genomes (by computing binding energies for representative mutations and assembling an energy-based weight matrix (EBWM), which is then used for the task); (ii) explore whether energy-minimization methods utilizing MD fields can predict protein–DNA binding when DNA sites, or the protein, are mutated.

For our study, we focus on the purine repressor, PurR, from Escherichia coli, a well-characterized TF with more than 20 known sites in the genome. The purine repressor is a member of the sizable LacI family, which is often regarded as a model system for transcription regulation. The abundance of both experimental \((13)\) and bioinformatics \((14)\) data makes this an ideal target for testing structure-based prediction techniques, and to study their assets and drawbacks.

We demonstrate that generic MD tools predict favorable binding energies for known cognate sites. To quantify the power and limitations of this approach, we investigated the following: (i) can we recognize the cognate sites from a large set of decoys, and estimate the number of false positives? (ii) How does the performance in the above test compare with that of a motif obtained from the set of cognate sites by bioinformatic methods? By calculating binding energies we can also answer the following questions which are not addressable by bioinformatic means: (i) what is the relative importance to recognition of direct binding energies to indirect factors such as DNA bending? (ii) can the computed results for \(\Delta G_{\text{binding}}\) of mutations in DNA, and more importantly in the protein, be compared to experiment? [Bioinformatics data can also be converted to compute \(\Delta G_{\text{binding}}\) for DNA, but not protein mutations as in ref. (1–4)].

To test the ability of the force field to discriminate between cognate sites and random decoys, we developed a procedure to speed up calculations and the screening of many sites. We find that a single cognate site can be discriminated from about 7000 random decoy sites. While such performance is impressive, it is insufficient to detect sites from the whole bacterial genome. In the comparisons of our results with experimental binding free energies for DNA and amino acid point mutations, we obtain the correct order of binding free energies of the mutants.

### MATERIALS AND METHODS

The change in free energy due to protein–DNA interactions can be decomposed as

\[
G_{\text{binding}} = G_{\text{protein–DNA complex}} - G_{\text{free DNA}} - G_{\text{free protein}}
\]

Clearly, \(G_{\text{binding}}\) depends on both the particular DNA sequence and the protein. In order to simplify the problem from a computational point of view, it is often assumed that the differences in \(G_{\text{binding}}\) for two different DNA sequences are dominated by differences in enthalpy. Entropic contributions are usually ignored since the entropy losses upon binding for both the fragment of DNA and the protein are likely not to depend significantly on the DNA sequence; hence

\[
G_{\text{binding}} \approx E_{\text{binding}} = E_{\text{protein–DNA complex}} - E_{\text{free (straight) DNA}} - E_{\text{free (unbound) protein}}
\]

Furthermore, if DNA sequences bound by the same protein are compared and \(\Delta E(DNA1, DNA2)_{\text{binding}} = E(DNA1, Protein) - E(DNA2, Protein)\) is of interest, the term \(E_{\text{free (unbound) protein}}\) cancels out.

The energies of the molecules were measured after minimizing the energy of their structures using the AMBER software package, its force field and an implicit water model. The reference structure in this study is 1qpz \((15)\), a wild-type PurR structure bound to DNA. The sequence of the DNA is also the consensus sequence obtained in the bioinformatics study of ref. (14) and we shall thus refer to it as the consensus sequence. The structure, depicted in Figure 1, was reduced to its 60 amino acid headpiece, and the DNA was trimmed to the 16-bp consensus sequence.
The first amino acid is missing and was not inserted artificially. The reference for straight DNA was taken and trimmed from the first model of the noncognate Lac–DNA binding complex 1osl (16). DNA sequences were exchanged with the 3DNA computer application (17). The experimental DNA backbone remained in place, only the base pairs were replaced. The free DNA molecule obtained in this manner deviates from a ‘perfect’ B-DNA molecule by about 1 Å RMS. While the experimentally derived straight DNA molecule was preferred over one molecule by about 1 Å RMS. While the experimentally obtained in this manner deviates from a ‘perfect’ B-DNA only the base pairs were replaced. The free DNA molecule by about 1 Å RMS. While the experimentally derived straight DNA molecule was preferred over one with average coordinates, this choice had no significant impact on our results. The SD of the energy difference between the canonical B-DNA structures and ours is merely 1.2 kcal/mol for the 50 random DNA sequences discussed subsequently does not change at all.

Protein mutants were generated with the Mutator 1.0 plugin built into VMD (18). The software uses psfgen to build a new side chain from pre-defined parameters for the CHARMM force field; this structure is not relaxed further by VMD. But the mutated side chain assumes a low-energy conformation during energy minimization because—unlike the original residues—mutated residues were not constrained to remain close to the coordinates of the original structure (see subsequently). For the study of DNA point mutations, the respective structures 1qp0, 1qp4, 1bdh, 1q6b, 1q7p and 1qqa from Ref. (15) were used in addition to 1qmz. We applied psfgen to combine and prepare structures for minimization. It should be noted that the conformational energy of the free unbound protein structure [Equations (2) and (6)] was not considered in most cases because we were only interested in differences between complexes. For example, $\Delta \Delta E_{\text{protein deform}}$ [Equation (6)] is simply the energy difference between the two bound protein structures. In our investigation of amino acid mutations, we approximated $E_{\text{unbound protein}}$ [Equation (2)] with $E_{\text{bound protein}}$, i.e. the self-energy of the protein in the bound complex. Again, this approximation is reasonable as we were only interested in differences of the binding energy between mutant complexes.

For all computations, we used the Amber 9 program with the parm99 force field (5), and the second implicit water model from Ref. (19). No cut-off was applied. Hydrogen atoms and the nucleic bases, as well as substituted residues in our amino acid mutation study, were allowed to rearrange freely to eliminate steric clashes. The movement of the protein and the DNA backbone was restricted by springs with a spring constant of 1.0 kcal/(molÅ²). To each configuration, 2500 steepest-descent and 2500 conjugate-gradient minimization steps were applied before energies were calculated to ensure convergence. A typical minimization run for a protein–DNA complex took about 4 h on a 3 Ghz Pentium 4 desktop computer.

While the relaxation of the structures is an essential element of our method, we cannot allow energy minimization to proceed unhindered. This is because: (i) we do not fully trust the potentials and (ii) the finite temperature fluctuations (not included) may prevent the structure moving into certain energy wells. As mentioned in the Introduction Section, previous work has indicated that the more the complex is allowed to move away from the known experimental structure (11,12), the less reliable are the energy-based methods in predicting binding specificity. The springs introduced in the previous paragraph limit the drift of the structure, but their strength is an additional parameter of the problem. In practice, for the spring constant we employ, the RMSD of the protein backbone changes by about 0.4 Å from the native structure. Fortunately, we find that the relevant aspects of the binding, namely the relative preferences to different sequences, are independent of the choice of the spring constant as long as the structures’ integrity is preserved. This conclusion was reached after performing studies with spring constants of 1.0, 2.5, 5.0 and 7.5 kcal/(molÅ²).

**RESULTS**

**Comparison with bioinformatics scores**

In order to assess the quality of binding predictions based on the all-atom calculations, we compared them with predictions made using a bioinformatic technique. The PurR TF has been studied extensively and is therefore particularly well suited for this task. Mironov and co-workers (14) compiled a collection of 21 binding sites to which PurR is known to bind in *E. coli*. Assuming independence of the influence of different base pairs on specificity, they set up a PWM that we use to calculate bioinformatics scores for various DNA sequences. Given a sufficient number of known sites, PWM scores provide a good approximation of experimentally measured binding energies (20–22) and have sufficient specificity to detect binding sites in bacterial genomes (23).

We challenged our structure-based approach, which uses only one known site that is a part of the crystal structure, to detect cognate sites among random ones using the binding energies after minimization. These energies were also compared to the PWM scores. In particular,
we examined the consensus sequence, the 21 binding sequences, 50 random sequences and several binding sites of closely related TFs FruR, GalR/GalS and MalI. Sites of homologous TFs were chosen because they constitute particularly challenging sites that are similar to PurR cognate sites and share the same palindromic structure. As shown in Figure 2, bioinformatics scores and binding energies correlate well with a linear correlation coefficient of −0.6 for the random sequences and −0.8 for all sequences displayed. The bioinformatics consensus sequence has the second lowest energy and random non-cognate sequences are generally well separated from random cognate sequences. While the separation between cognate sites and the 50 random decoys is reassuring, it is important to find out whether the procedure is able to find cognate sites among 10^6 other sites (decoys) on the bacterial genome.

Assuming a Gaussian distribution of the binding energies for random sites, we can estimate the number of decoys that have binding energies comparable to the cognate sites. The distance between the average of the random sequences (red line) and the third worst cognate site (black line) is 3.63 σ. (We chose the third worst cognate site because the two next sequences are the PurA operator sites, see next paragraph.) This roughly amounts to one false positive hit in 7000 random sites. Note that 50 random sequences can only yield a rough estimate for this number. This number is quite encouraging although it should only be considered a rough estimate. For comparison, the corresponding PWM bioinformatics scores from Ref. (14) are separated by 4.55 σ which would amount to one false positive hit in 370,000 sequences. (The K12 E. coli genome (24) consists of 4.64 Mbps).

The two cognate sequences with the highest energies are the two PurA operators (unfilled black circles). Indeed, the suggestion that the PurA operon may be regulated by PurR is controversial (14,25,26). Although at the lower end of the spectrum, the bioinformatics scores for these sites are comparable with other cognate sites, while our computations give distinctly higher binding energies.

Testing the energy-based approach on operator sites regulated by other members of the LacI family is a particularly challenging task. Although the FruR and MalI binding sequences are energetically well separated from the PurR cognate sequences, GalR/GalS binding sequences are not. (The bioinformatics score appears to have less difficulty with these sites.)

Finally, we would like to point out that the absolute energy scale is incorrect, in line with the conclusions of Ref. (9). Excluding PurA, the range of binding energies for cognate sequences is 10 kcal/mol which is clearly too large. The underlying assumptions and approximations of the method are, however, quite considerable and quantitative agreement cannot really be expected.

**Direct and indirect contributions to the binding energy, sequence logos**

The binding free energies can be subdivided into two parts: direct interactions between the TF and DNA, and indirect contributions due to sequence-specific DNA bending. In recent work, Paillard and Lavery (9) noted that the level of each contribution varies significantly from complex to complex. Their method is based on a careful analysis of a subset of sequences with particularly low-binding energies, after having computed the energies of all possible 4^n sequences. Here, we propose a simple method which can distinguish between contributions of bending and protein–DNA interactions on the basis of a rather limited set of measurements.

To understand the source of the sequence specificity of PurR, we partitioned its binding energy as follows:

\[ E_{\text{binding}} = E_{\text{interaction}} + E_{\text{DNA deform}} + E_{\text{protein deform}} \]

where

\[ E_{\text{interaction}} = E_{\text{protein–DNA complex}} - E_{\text{bound protein}} - E_{\text{bound DNA}} \]

and

\[ E_{\text{DNA deform}} = E_{\text{bound DNA}} - E_{\text{straight DNA}} \]

\[ E_{\text{protein deform}} = E_{\text{bound protein}} - E_{\text{unbound protein}} \]
We next computed direct and indirect contributions for both cognate and random sequences and compare the differences: on average, \( E_{\text{interaction}} \) was lower by 34 kcal/mol for the cognate sites compared to random ones and \( E_{\text{DNA deform}} \) lower by 7 kcal/mol. Assuming that the force field reproduces the correct ratios of direct and indirect contributions, specificity towards PurR is predominantly determined by protein–DNA interactions. It is interesting to note that \( E_{\text{protein deform}} \) was slightly higher for cognate sites (2 kcal/mol) indicating that the interactions were strong enough to bend the protein towards a slightly unfavorable position.

To study the contribution of individual base pairs to specificity, and to significantly speed up computations, we used energy minimization to calculate a position-specific energy matrix, analogous to PWM (14). As specificity towards PurR is dominated by direct, pairwise interactions, we computed the change in \( E_{\text{interaction}} \) due to each possible single mutation of the consensus sequence and set up an EBWM (Table 1). (While it is in principle possible to construct a statistical weight matrix based on the top 21 sites identified by energy minimization, this would effectively reproduce the experimental PWM of Ref. (14). The interaction energy for an arbitrary DNA sequence can now be computed by adding the appropriate base pair energies. This requires only a limited number of computations at the cost of being less accurate.

Although computationally efficient, both EBWM and PWM methods are based on the assumption that the contributions of individual base pairs are independent from each other, neglecting many-body effects, such as due to solvation. EBWM calculated using \( E_{\text{interaction}} \) also ignores sequence-dependent contributions of DNA deformation.

Table 1. Position-specific energy matrices based on direct interaction energies and interaction energies plus bending corrections

<table>
<thead>
<tr>
<th>( \Delta E_{\text{interaction}} )</th>
<th>( \Delta E_{\text{interaction}} + \Delta E_{\text{DNA deform}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>C</td>
</tr>
<tr>
<td>-----</td>
<td>-----</td>
</tr>
<tr>
<td>0.0</td>
<td>-0.5</td>
</tr>
<tr>
<td>1.3</td>
<td>0.0</td>
</tr>
<tr>
<td>4.7</td>
<td>14.3</td>
</tr>
<tr>
<td>-1.0</td>
<td>0.0</td>
</tr>
<tr>
<td>0.0</td>
<td>2.1</td>
</tr>
<tr>
<td>0.0</td>
<td>3.5</td>
</tr>
<tr>
<td>0.0</td>
<td>2.0</td>
</tr>
<tr>
<td>6.2</td>
<td>0.0</td>
</tr>
<tr>
<td>5.3</td>
<td>0.1</td>
</tr>
<tr>
<td>2.2</td>
<td>0.6</td>
</tr>
<tr>
<td>3.9</td>
<td>3.4</td>
</tr>
<tr>
<td>3.4</td>
<td>3.8</td>
</tr>
<tr>
<td>0.9</td>
<td>2.3</td>
</tr>
<tr>
<td>8.4</td>
<td>0.0</td>
</tr>
<tr>
<td>-0.6</td>
<td>0.6</td>
</tr>
<tr>
<td>-0.7</td>
<td>-0.6</td>
</tr>
</tbody>
</table>

The energies are normalized to the consensus sequence, which has, accordingly zero binding energy. This is why a ‘\( \Delta \)’ appears in front of the energies. All contributions from each base pair (including bending) were considered to be independent of the other base pairs. Energies are given in units of kilo calories per mole. Only the first decimal place is shown.

The DNA bending energy cannot be easily decomposed into contributions of individual base pairs; the energetic contribution of a single base pair to DNA bending not only depends on its neighbors but also on the whole sequence. This makes the exact treatment of the problem computationally challenging. Lavery and co-workers (8–10) address this issue by constructing an ‘average structure’ into which they can substitute all possible sequences and create a sequence logo based on the sequences with the lowest energy. Endres et al. (11,27) employ an efficient scheme to screen sequences and only compute those which look promising. Thus they afford to compute bending energies for each individual sequence.

We sought to improve the EBWM approximation by making a ‘zeroth order’ estimate of the bending energy. The bending energy can also be subdivided into two parts: interactions between base pairs and the backbone, which are approximately independent of the other base pairs, and interactions between nucleic bases, which are not. Nevertheless, we tested if the results can be improved by including an additive bending term to \( E_{\text{interaction}} \). Such a treatment is tantamount to considering interactions of single base pairs in the ‘mean-field’ environment of the consensus sequence. Similar to the case of \( E_{\text{interaction}} \), we computed the change in \( E_{\text{DNA deform}} \) due to each possible single base pair substitution in the consensus sequence and set up a second EBWM (Table 1). Thus, for every position along the DNA sequence, the change in bending energy due to a point mutation is measured. Then, the total bending energy for an arbitrary sequence is approximated as the sum of the changes in bending energy at each position. This approximation only captures the interactions of the base pairs and the backbone and some mean-field portion of the interaction between base pairs, but leaves out base stacking energies which are explicitly not pairwise additive. More precisely, to find \( E_{\text{DNA deform}} \), as defined in Equation (5), \( E_{\text{bound DNA}} \) and \( E_{\text{straight DNA}} \) are needed. The former is computed by excising the DNA from the energy-minimized protein–DNA complex structure and measuring its energy without the surrounding protein. \( E_{\text{straight DNA}} \) is, of course, the energy of the DNA in its free form. Subtracting the two energies yields the energy of deformation, \( E_{\text{DNA deform}} \).

In an effort to gauge the usefulness of the two matrices, we computed the energy difference between the worst cognate sequence and all the sequences encountered in scanning the E. coli genome excluding the PurA operator sites. We repeated this to find the energy difference between the worst cognate sequence and the 50 nonspecific random sequences discussed above. If only protein–DNA interactions are taken into account, the separation between the lowest and the average of all sequences is 2.74 \( \sigma \) (3.2 \( \sigma \) based on the 50 nonspecific sequences). If the additive bending correction is included the distance is also 2.74 \( \sigma \) (3.38 \( \sigma \) based on the 50 nonspecific sequences). The 50 nonspecific random sequences are too few to allow reliable conclusions; clearly, the genome scan is more significant. These results indicate that a simplified EBWM approach which only considers additive interactions is not sufficient to the binding energy. Can we improve upon this by taking the bending energy into account?
to provide accurate discrimination of sites. Similarly, we expect that the experimentally obtained PWM based on $\Delta G_{\text{binding}}$ of single base pair mutants of the consensus sequences (14) to suffer from a similar lack of discrimination power.

To visualize the contribution of individual base pairs to the PurR motif, and hence to the specificity of recognition, we converted the information contained in the EBWM into a sequence logo (Figure 3). This is done by using Boltzmann weights to represent the frequency of occurrence of each base pair at each position. Room temperature was used in the Boltzmann factors, that is, $k_B T = 0.59 \text{ kcal/mol}$. Comparison with the bioinformatics logo from Ref. (14) (Figure 3a) indicates that the structure-based method is able to reproduce the specificity of most of the positions in the PurR motif. In particular, base pairs at positions 3 (G), 5 (A), 6 (A), 11 (T), 12 (T) and 14 (C) are identified correctly, but the method cannot distinguish between the consensus CpG versus GpC in positions 8 and 9. Base pairs 8 and 9 play an important role in the binding of DNA because PurR bends DNA by intercalating a lysine side chain between these two base pairs (28). Either CpG is selected in nature for reasons that cannot be explained by binding energy considerations or the force field cannot capture a subtle difference in bending between CpG and GpC in the center of the binding sequence. Nevertheless, it is surprising that although the computed energy differences are too big compared to the experimental energy differences, the sequence logo is recreated rather accurately.

The Boltzmann weights of each base pair at any position along the DNA represent a probability distribution for 4 bp at that position. Two different probability distributions $p(i)$ and $q(i)$ can be compared using the relative entropy measure [Kullback–Leibler divergence (29)], $\sum_i p(i) \log p(i)/q(i)$. Here, $q(i)$ is the probability distribution of the base pairs $i = \text{AT, CG, GC or TA}$ at some position in the DNA derived from the bioinformatics weblogo and $p(i)$ is the probability distribution for the base pairs at the same position derived from the Boltzmann weight, which is computed with our method.

The more dissimilar two probability distributions are, the larger is their relative entropy. Excluding the two end base pairs and the two middle base pairs with the problematic CpG ambiguity, the distance between the probability distribution based on $E_{\text{interaction}}$ and the bioinformatics probability distribution is, on average, 0.8. Excluding the same base pairs, the distance between the probability distribution based on $E_{\text{interaction}} + E_{\text{DNA deform}}$ and the one based on bioinformatics is 0.3, which reflects a clear improvement.

In summary, we have shown that: (i) the contribution of indirect readout due to DNA bending is significant; (ii) this contribution cannot be easily accounted for by a site-specific approximation; (iii) the EBWM provides a fast way of estimating the binding energy but suffers from a significant loss of statistical power; and (iv) a structure-based energy calculation is able to capture most of the PurR motif, but fails to identify the central base pairs correctly. This suggests a hybrid strategy of first using EBWMs to scan for potential binding sites, and then following up by a more computationally intensive energy minimization for these candidates.

**Investigation of DNA and protein mutants**

In this section, we compare the binding energies derived from the structure-based approach with the experimental free energies of binding for a number of DNA and amino acid point mutations (15,30). In particular, we investigated the following sequences bound to the protein wild-type and K55A mutant:

- $\text{seq}_1 = \text{aagcaaa}cg(t)tctcg}$ (consensus),
- $\text{seq}_2 = \text{aagcaaa}cg(g)tctcg}$,
- $\text{seq}_3 = \text{aagcaaa}cg(c)tctcg}$,
- $\text{seq}_4 = \text{aagcaaa}cg(a)tctcg}$.

In addition, we studied protein mutants L54M, L54S, L54T and L54V bound to the consensus sequence. The results are summarized in Table 2.
Table 2. Calculated changes in binding energies of DNA and amino acid point mutations compared with experiments (15,30)

<table>
<thead>
<tr>
<th>Mutant</th>
<th>$\Delta E_{\text{binding}}$</th>
<th>$\Delta G_{\text{binding}}$ (experiment)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>L54M</td>
<td>5.79</td>
<td>0.38</td>
</tr>
<tr>
<td>L54S</td>
<td>16</td>
<td>larger, not measured</td>
</tr>
<tr>
<td>L54T</td>
<td>10.05</td>
<td>..</td>
</tr>
<tr>
<td>L54V</td>
<td>6.15</td>
<td>..</td>
</tr>
<tr>
<td>K55A</td>
<td>12.55</td>
<td>3.48</td>
</tr>
</tbody>
</table>

When only the DNA is mutated, the binding order is correct (top panel). When both DNA and the protein are mutated (middle panel), two DNA mutants are lower in binding energy and one higher than the original sequence. This is correctly identified by our method, but the binding preference to seq1 and seq2 is reversed. When only the protein is mutated, the binding preferences of the DNA to the mutants are correctly captured (bottom panel). Energies are given in kilo calories per mole and measured relative to the respective consensus protein–DNA complex.

DISCUSSION

It is clearly desirable to understand protein–DNA binding on a molecular level, and all-atom energy calculations based on minimizing experimental structures are a promising step towards this goal. In this work, we studied the feasibility of predicting the affinity of a TF to different sequences, by using off-the-shelf and widely used interaction potentials. Our main goal was to test whether computation of energies using such a potential allows discrimination of cognate sites from random decoys. Using the example of the PurR TF as a model system, and starting from the structure this protein bound to a specific DNA sequence, we tested our method in the following ways:

- estimated the number of random sites that have binding energies comparable to cognate sites, thereby assessing the potential of this method to detect sites in long genomic sequences;
- compared the performance of this structure-based method with the bioinformatic PWM technique that requires a priori knowledge of several cognate sites;
- examined PurR motifs obtained using structure-based calculations, and compared them with the motif inferred from the cognate sites; and
- calculated the change in the binding energy due to mutations in the protein and DNA and compared with experimentally measured $\Delta G_{\text{binding}}$.

We further investigated the contribution of the sequence-dependent DNA binding and tested whether computations can be accelerated using an EBWM approach. This systematic and diverse testing makes our study complimentary to other recent works (7–12,31).

Overall, the changes in energy of the minimized structures correlate well with corresponding bioinformatics scores and are accurate enough to discriminate between binding and random sequences. Unfortunately, they are not sufficiently discriminating to enable systematic scanning of entire genomes. The method can, however, distinguish between weak and strong binding sites and, to a lesser extent, between operator sites of related factors.

To highlight the contributions of individual base pairs, we compared a motif logo obtained using structure-based calculations with the logo for cognate sites. While most of the positions reflect the cognate motif correctly, the two central base pairs are predicted incorrectly with atomistic force fields—indicating no difference between G and C in these positions. This difficulty is likely due to a complicated binding mechanism through lysine intercalation used by PurR to bind the central base pairs. It is possible that sequence-dependent bendability of DNA makes CpG a preferred base pair in the center of a sharply bent PurR site. Understanding the molecular mechanism of recognition of the central base pairs requires further studies using MD. A method that can resolve this discrepancy is likely to provide a significant improvement to structure-based predictions for PurR and other TFs that bend DNA.

Our analysis provides a glimpse into the promise of structure- and interaction-based methods. Relatively crude computations are able to predict the correct order of binding energies. This is particularly useful for the study of amino acid mutants which cannot be investigated with standard bioinformatics methods. There are several reasons for the limited success of our approach. First, the force fields employed are likely not accurate enough to deliver precision of the binding energy at the level of a few kilo calories per mole, as required to discriminate cognate sites, especially when an implicit water model is used. [The ParmBSC0 force field, for example, could be used in the future because of its improved treatment of noncanonical backbone conformations compared to PARM99 (32)]. Second, our procedure crudely approximates the differences in the binding free energy by ignoring entropy contributions and by limiting the flexibility of

Although the overall energy scale is incorrect, we are able to reproduce (with one exception) the correct order of experimental binding free energies for all DNA and amino acid mutants. Qualitatively similar results were obtained in Ref. (9), which, however, only considered mutations of the DNA sequence. The sampling of amino acid mutants is particularly relevant because it allows us to predict whether simple modifications of TFs can lead to higher or lower binding affinity.
the protein–DNA complex through the use of fast energy minimization, thus not allowing for major rearrangements of the structure. Furthermore, the restraints on the protein and the DNA, which keep them from deviating too far from the native experimental structure, could be increasing the energy scales in the system, thus exaggerating the energy differences. The energies of the restraining springs are not included in the calculations, but the springs may keep the structure from relaxing to the equilibrium coordinates that the force field favors. The (qualified) success of this simple approach suggests that further optimization of the force field (7) and conformational sampling [e.g. similar to those of (11)] may lead to significant improvements. Resolution of these issues is necessary to gain a better quantitative understanding of protein–DNA binding.

Structure-based methods are more laborious than bioinformatics but less costly and elaborate than experiments. Although accuracy is still somewhat lacking in current implementations, the results are promising and still leave considerable room for improvements. Promising applications which already appear feasible include the study of sequence-dependent motion of proteins along DNA, and investigations of simple amino acid point mutations in conjunction with experiments.

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