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Coarse-Grained Model Of Entropic Alloster y

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Many signaling functions in molecular biology require proteins to bind to substrates such as DNA in response to environmental signals such as the simultaneous binding to a small molecule. Examples are repressor proteins which may transmit information via a conformational change in response to the ligand binding. An alternative entropic mechanism of “allostery” suggests that the inducer ligand changes the intramolecular vibrational entropy, not just the mean static structure. We present a quantitative, coarse-grained model of entropic alloster y, which suggests design rules for internal cohesive potentials in proteins employing this effect. It also addresses the issue of how the signal information to bind or unbind is transmitted through the protein. The model may be applicable to a wide range of repressors and also to signaling in trans-membrane proteins.

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Introduction.—It is becoming increasingly clear that dynamics, as well as static structure, are important in molecular biology. For example, simulations of dynamical transitions in proteins [1] suggest that collective global modes are correlated with protein function. This Letter focuses on repressor proteins which bind to DNA to “turn off” genes when the cell does not require their expression. The binding is “allosteric”: it is activated depending on the presence of inducer ligands, small molecules which themselves bind to the protein at a site distant from the active site. The “holorepressor” (“aporepressor”) is the protein with (without) a bound ligand. In allosteric repressor proteins, the ligand binding site is distant from that of the DNA. For this reason ligand binding has often been assumed to cause a conformational change within the repressor protein, decreasing its affinity for DNA in one state compared to the other state. However, there is evidence that dynamically induced entropic changes may contribute to alloster y [2–4].

A classic example of a repressor system is the E-coli lac repressor [5–9]. In this case the aporepressor binds to DNA, suppressing the genes for the metabolism of lactose. A second example is the E-coli trp repressor [10–12] that, on binding, prevents the expression of the gene for tryptophan synthesis. But in contrast to the lac, trp-type holorepressor proteins bind to DNA and the aporepressors do not. There are many such repressor systems but the lac and the trp will act as representative cases for this Letter. Our challenge is to explore whether the Brownian fluctuations in protein structure may carry information between the two binding sites, thereby producing cooperative lac-type or trp-type behavior. This mechanism of cooperativity is one of the key questions in understanding protein function [13].

As generally true for the functional roles of protein dynamics, the lower-frequency softer modes will dominate [14]. Although higher modes are more numerous, they are spatially localized due to elastic disorder [15]. Ligand binding at sites where high frequency modes have significant amplitude will therefore generally have only local effects; long distant allosteric signaling will be exponentially suppressed beyond the localization length of the mode. Focusing on the slower, global modes additionally motivates a spatially coarse-grained model.

A coarse-grained model.—In a coarse-grained representation (motivated by the common dimer motif) we model a repressor protein dimer as two rigid plates of length $l$ and width $w$, representing the two protein monomers (see Fig. 1). We parameterize the relative motion of the plates by three relative translation displacements $(x, y, z)$ and three rotation angles $(\theta_x, \theta_y, \theta_z)$. The stabiliz-
ing contact interaction between the two dimers is characterized by five quadratic (3D) potential wells. Figure 1 shows five of the effective springs that arise, $\lambda^2$ which are perpendicular to the plates. We find that a minimal model requires just four other springs in the plane of the plates (summations of diagonal springs between the plates), which we label as $\lambda^2_{00}, \lambda^2_{01}, \lambda^2_{10},\lambda^2_{11}$, representing the resolved $y$ and $z$ relative displacements. These local interactions represent a minimal set of “sticky patches” which could arise from hydrophobic, side chain, or electrostatic forces. The corresponding spring constants $\lambda^2_i$ could be calculated in principle from the details of these interactions. We allow only local modifications of the contact interactions on binding. The springs local to the ligand binding site will be affected by the binding of an inducer or corepressor. (Lac actually binds two inducer ligands, which we simplify here with a single bound state.) Similarly springs local to the DNA binding site will be affected by binding to DNA. The other springs act as anchoring potentials.

We consider vibrations of the plates in three different planes: (i) in the $x,y$ plane (translational vibrations along the $x$ axis and rotations about the $z$ axis); (ii) in the $x,z$ plane (translation along the $x$ axis and rotation about the $y$ axis); and (iii) in the $y,z$ plane (translation along the $z$ axis and rotation about the $x$ axis).

The Trion potential [16] replaces the full molecular dynamics potential with a single pair wise quadratic potential of universal strength and this is found to be sufficient to describe the low frequency modes involving coherent motion of large groups of atoms. In our model we similarly look only at these low frequency modes and describe them by harmonic potentials between the protein domains, but allow the potentials between the protein monomers to acquire locally specific values.

As an example of detailed calculations we take the $2 \times 2$ system of motions in the $y,z$ plane. We write $z_{01} = z_{00} - \frac{\theta_{z}}{2}$ and $z_{0-1} = z_{00} + \frac{\theta_{z}}{2}$ (where $\theta_{z}$ is the angle of rotation about the $x$ axis) to obtain the Hamiltonian in terms of the mutual translational vibration coordinate $z_{00}$ and mutual rotational vibration coordinate $\theta_{z}$.

$$H = \frac{1}{2} p M^{-1} p + \frac{1}{2} z K z$$

(1)

where the interaction matrix for the $y,z$ plane is given by

$$K = \begin{pmatrix}
-(\lambda_{01}^2 + \lambda_{01}^2 + \lambda_{01}^2) & \frac{1}{2}(\lambda_{01}^2 - \lambda_{01}^2)
\frac{1}{2}(\lambda_{01}^2 - \lambda_{01}^2) & -(\lambda_{01}^2 + \lambda_{01}^2)
\end{pmatrix}$$

the inertial matrix by

$$M = \begin{pmatrix}
m & 0 \\
0 & I
\end{pmatrix}, \quad z = \begin{pmatrix}
z_{00} \\
\theta_{z}
\end{pmatrix}$$

where $m$ is the reduced mass and $I$ is the reduced moment of inertia of the dimer pair. This leads to the partition function in the relevant classical limit.

$$Z = \int \cdots \int e^{-\frac{H}{kT}} dx_0, d\theta_i = \frac{2\pi kT}{(|M|^{-1}|K|)^{1/2}}.$$ 

(2)

Finally from (2) the entropy of the protein dimer for a single plane is

$$S = N k (2\pi T \sqrt{m} / l + 1 - 1/2 \ln[1 + \lambda \lambda_{-1}])$$

(3)

where $\lambda_{-1}$ has been abbreviated to $\lambda_{1}$ for convenience. We are interested in the difference between the change in entropy on binding at the DNA binding site of the two cases in which the protein is, and is not also bound to the inducer. We call this $\Delta S = S_{apo} - S_{holo}$. A result with $\Delta S \neq 0$ would signify cooperative behavior, i.e., the binding to DNA is affected by the binding to the inducer. We write $\Delta S$ in terms of dimensionless spring constants $\lambda_{1}$ and $\lambda_{-1}$ and bound to unbound ratios $\lambda_{1} = \frac{\lambda_{1}}{\lambda_{0}}$ and $\lambda_{-1} = \frac{\lambda_{-1}}{\lambda_{0}}$. This gives us

$$\Delta S = \frac{1}{2} N k \ln \left(\frac{4\lambda_{1} + 1}{4\lambda_{1} - 1}(4\lambda_{-1} + 1)\right)$$

(4)

The other modes can be modeled in the same way to give additional contributions to $\Delta S$. The two-plane model generates a $3 \times 3$ form of $M$ and $K$ for coupled rotations about $y$ and $z$, and translations along $x$ plus one simple $y$ translation.

We take $N = 2$ for the lac since it is a tetramer of two dimers and similarly for the trp since it represses as a dimer of dimers so also has two dimers.

$\Delta S > 0$ gives the trp case whereby the affinity for the holorepressor binding to DNA is greater than the apo repressor. $\Delta S < 0$ however gives the lac case since the apo lac repressor is the one with the higher affinity for DNA. Applying these inequalities to Eq. (4) gives the following rule determining which case arises.

$$\lambda_{1} \lambda_{-1} + 1 \begin{cases}
> 0 & \Rightarrow \lambda_{1} > \lambda_{-1} \text{ (trp)} \\
< 0 & \Rightarrow \lambda_{1} < \lambda_{-1} \text{ (lac)}
\end{cases}$$

(5)

The trp case occurs when both spring constants increase ($\lambda_{1} > 1, \lambda_{-1} > 1$) or decrease ($\lambda_{1} < 1, \lambda_{-1} < 1$) on ligand binding. The lac case occurs when one spring constant increases and the other decreases ($\lambda_{1} > 1$ and $\lambda_{-1} < 1$ or $\lambda_{1} < 1$ and $\lambda_{-1} > 1$). Figure 2 plots the function $\Delta S(\lambda_{1}, \lambda_{-1})$ (Eq. (4)). Biologically relevant values for the original spring constants were chosen using protein B-factor data [related to the root mean square (rms) positions $\sqrt{\langle u^2 \rangle}$ of the atoms $B = 8\pi^2\langle u^2 \rangle$ [17]] and steered simulations (see later) giving the case for a potential which is stronger at the inducer binding site ($\lambda_{1} > \lambda_{-1}$). It can be seen that the negative $\Delta S$ lac type effect is maximised when the ligand binding at the inducer binding site decreases the spring constant $\lambda_{1}$ but
\( \lambda_{-1} \) increases on DNA binding. The trp-type effect (positive \( \Delta \Delta S \)) is maximised when both spring constants decrease as much as possible on binding. This reduction will however be limited by the physical requirement for overall stability of the complex. Requiring that the rms displacement of the monomers be less than the average separation of the atoms leads to the estimation that \( \lambda_i > 0.1 \text{ kT} \lambda_i^{-2} \).

Physically, “entropic allostery” allows the lac inducer binding to communicate via the large amplitudes of the internal modes of the protein to the “readhead” binding regions near the DNA which as a result move too much to be inserted into the DNA.

Estimating the spring constants.—We evaluate our model in the real example of the lac repressor as an illustration. First, we converted the protein data for the \( B \) factors \([7,8,18]\) into rms vibration values and estimated the spring constants \( \lambda_i \) and \( \Lambda_i \) (in this case averaged over the vibrations in the different planes) from the expression \( \lambda \sim 1/(\mu^2) \) giving \( \lambda_1 = 1.2, \lambda_{-1} = 0.1 \) (estimated from \([19]\)), \( \Lambda_1 = 0.07 \) and \( \Lambda_{-1} = 6.7 \) (so supporting our prediction that the lac case has \( \lambda_1 < 1 \) and \( \lambda_{-1} > 1 \)). We then calculated an estimate of \( \Delta \Delta S \) using Eq. (4). Including a factor of 3 due to the three planes of vibration, we obtain for our plate-dimer model a value of \( T\Delta \Delta S \sim -1.4 \text{ kT} \). Since the experimental values for the change in binding energy between holo and aporepressor binding to DNA are \( \Delta \Delta G \sim 6 \text{ kT} \) \([20–22]\) this indicates that the entropic contribution is likely to be significant since the crystal dynamics can only be a lower bound for the amplitudes of vibration.

To improve upon this rough estimate we calculated the interaction energy between the lac monomers in a fully atomistic computation using the software “cns” \([23]\) and steering the relative positions of the two monomers. We used the x-ray crystal structure coordinates PDB ID 1LBI \([7]\) for the aporepressor and 1TLF \([18]\) for the holorepressor. By relative translation along the three axes, and rotation about axes at the extremities of the protein dimer, and recalculating the total interaction energy at each increment, we were able to build up curves for the potential energy wells (see Fig. 3). By curve fitting the bottom of these wells to a quadratic (to fit with our harmonic approximation) we were able to extract the curvature and therefore the spring constants for each global mode of both the holo and aporepressors. We then used these to calculate \( S_{\text{inducer}} \) which is an estimate for \( -\Delta \Delta S \) in the case of large \( \lambda_{-1} \). We add together the motion in the \( x, y \) plane, \( x, z \) plane, and \( y, z \) plane to obtain in total an estimate for \( T\Delta \Delta S \sim -2.48 \text{ kT} \). Interestingly the softest mode, contributing most to the allostery, is the one which shifts the DNA read heads (which point in the plane perpendicular to the core) perpendicularly away from the DNA in the \( x, z \) plane.

To check how significant this entropy contribution is to the total \( \Delta \Delta G \sim 6 \text{ kT} \) we compared the probability of genes being repressed (bound by repressor) against lactose concentration with and without this entropic contribution (following Yildirim \([24]\)) (see Fig. 4). For 95% activation (operators not bound) 18 lactose molecules are required with the entropic contribution but 50 would be

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required if there was no entropic contribution. Note there are only of order 100 repressors in the cell [5]. This would imply that the entropic contribution to allostery is significant in controlling the lactose level at which the gene expression is turned on.

Conclusions.— We conclude that inducer binding affects the Brownian motion within the repressor protein and this entropic effect contributes to the allostetic mechanism in DNA binding proteins alongside static conformational changes. We can relate the communication of the signal across the protein to “design rules” for the potentials within it.

Several extensions of this approach suggest themselves. A discrete many-springs model naturally extends to the case of a continuous potential between the plates. Second, the case of multiple, sequential ligand binding will lead to additional structure. To make the model even more realistic we should also include bending modes of the protein monomers themselves. Significantly, addition of such bending modes must increase the predicted $\Delta \Delta S$ value if the ligand binding changes the bending rigidity. For lac any such increase might give values of $\Delta \Delta S$ that actually dominate the binding free energy.

We expect this model can be also applied to transmembrane proteins that transmit signals across membranes into cells and organelles [25, 26]. These systems are similar in that an inducer ligand (e.g., adrenaline) binds to the receptor trans-membrane protein which in turn allows it to bind or unbind proteins on the interior of the membrane (e.g. to bind to a G-protein in the control cycle for glycogen). Within the restricted environment of the membrane we expect entropic allostery to play an important role in the transmission of the signal through the receptor protein.

The calculated values of the contribution to the free energy change from the change in intramolecular vibrational entropy of the protein easily reach the order of a few kT per molecule, within the experimentally observed range for these systems.

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