Protein Structure Prediction Using Basin-Hopping

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Abstract

Associative memory Hamiltonian structure prediction potentials are not overly rugged, thereby suggesting their landscapes are like those of actual proteins. In the present contribution we show how basin-hopping global optimization can identify low-lying minima for the corresponding mildly frustrated energy landscapes. For small systems the basin-hopping algorithm succeeds in locating both lower minima and conformations closer to the experimental structure than does molecular dynamics with simulated annealing. For large systems the efficiency of basin-hopping decreases for our initial implementation, where the steps consist of random perturbations to the Cartesian coordinates. We implemented umbrella sampling using basin-hopping to further confirm when the global minima are reached. We have also improved the energy surface by employing bioinformatic techniques for reducing the roughness or variance of the energy surface. Finally, the basin-hopping calculations have guided improvements in the excluded volume of the Hamiltonian, producing better structures. These results suggest a novel and transferable optimization scheme for future energy function development.
1 Introduction

The complexity of the physical interactions that guides the folding of biomolecules presents a significant challenge for atomistic modeling. Many current protein models use a coarse grained approach to remove degrees of freedom, such as non-polar hydrogens, which increases the feasible time step in molecular dynamics simulations.\textsuperscript{1,2} For a more dramatic improvement of the computational efficiency solvent degrees of freedom can be reduced.\textsuperscript{3} In this case more severe approximations can prevent the model from reproducing experimental results. Another option is to reduce the number of degrees of freedom of the solute. The associative memory Hamiltonian (AMH),\textsuperscript{4–6} is a coarse-grained molecular mechanics potential inspired by physical models of the protein folding process, but flexibly incorporates bioinformatic data to predict protein structure. The AMH is first optimised using the minimal frustration principle in terms of the $T_f/T_g$ ratio, which estimates the separation in energy relative to the variance for the misfolded ensemble. Along with using the energy of the native structure to estimate $T_f$, a random energy model\textsuperscript{7} estimate of the glass transition temperature, $T_g$, is used based on a set of decoy structures. $T_g$ represents a characteristic temperature scale at which kinetic trapping in misfolded states dominates the dynamics. An improved potential is next obtained that uses better estimates of the $T_f/T_g$ ratio obtained by maximizing the normalized difference between the native state and a sampled set of misfolded decoys which are self-consistently obtained from the potential itself. The potential so obtained is transferable for the prediction of structures outside of the training set. The ratio $T_f/T_g$ has provided a powerful metric for the optimisation of both this bioinformatically informed energy function,\textsuperscript{8,9} as well as other types of energy functions incorporating only physical information.\textsuperscript{10–12}

While the optimisation\textsuperscript{13} of parameters using a training set of evolved natural proteins smooths the energy landscape from what it would be for a random hetero-polymer, the common problem of multiple competing minima persists even when using a reasonably accurate structure prediction potential, such as this one. Simulated annealing with molecular dynamics has previously been used to search the rugged landscapes of optimised structure prediction potentials.\textsuperscript{14} While free energy profiles indicate that better structures actually are present at low temperatures, the slow kinetics
of a glass-like transition during annealing has prevented these minima from being reached.\textsuperscript{15} To quantitatively investigate the origin of the sampling difficulties it is desirable to use different search strategies.

Here we implement the basin-hopping global optimisation algorithm,\textsuperscript{16–18} which has proved capable of overcoming large energetic barriers in a wide range of different systems. Basin-hopping is an algorithm where a structural perturbation is followed by energy minimisation. This procedure effectively transforms the potential energy surface, by removing high barriers, as shown in Fig. 1. Moves between local minima are accepted or rejected based upon a Monte Carlo criterion. Avoiding barriers by employing a numerical minimisation step not only facilitates movement between local minima, but also broadens their occupation probability distributions, which overlap over a wider temperature range, thereby increasing the probability of interconversion.\textsuperscript{19} Furthermore, it does not alter the nature of the local minima since the Hamiltonian itself is not changed, enabling the comparison between molecular dynamics and basin-hopping generated minima. This method has previously been applied to find global minima in atomic and molecular clusters,\textsuperscript{20, 21} biopolymers,\textsuperscript{22, 23} and solids.\textsuperscript{24} Since the algorithm only requires coordinates, energies, and gradients, it can be transferred between different molecular systems such as binary Lennard-Jones clusters, all-atom biomolecules, or coarse-grained proteins models as in this study.

2 Theory and Computational Details

The AMH energy function used in the present work has previously been optimised over a set of non-homologous \( \alpha \) helical proteins, and consists of a backbone term, \( E_{\text{back}} \), and an interaction term, \( E_{\text{int}} \), which has an additive form.\textsuperscript{25, 26} This model is sometimes termed the AMC model (associative memory contact) to distinguish it from one that uses nonadditive water mediated interactions termed the AMW model.\textsuperscript{14, 27} Since this model has been described in detail before,\textsuperscript{15, 28} we will only summarize its form here. We employ a version of the coarse-grained model where the twenty letter amino acid code has been reduced to four, and the number of atoms per residue is limited to three (\( C_\alpha, C_\beta, \) and O), except for glycine residues. The units of energy and temperature
were both defined during the parameter optimisation. The interaction energy $\epsilon$ was defined in terms of the native state energy excluding backbone contributions, $E_{\text{amc}}^{N}$, via

$$
\epsilon = \frac{|E_{\text{amc}}^{N}|}{4N},
$$

where $N$ is the number of residues of the protein being considered. Temperatures are quoted in terms of the reduced temperature $T_{\text{amc}} = k_{B}T/\epsilon$. While $E_{\text{back}}$ creates self-avoiding peptide-like stereochemistry, $E_{\text{int}}$ introduces the majority of the attractive interactions that produce folding. The interactions described by $E_{\text{int}}$ depend on the sequence separation $|i - j|$. The interaction between residues less than 12 amino acids apart were defined by Eqs. (2).

$$
E_{\text{local}} = -\frac{\epsilon}{a} \sum_{\mu=1}^{N_{\text{mem}}} \gamma(P_{i}, P_{j}, P_{\mu}^{i}, P_{\mu}^{j}, x(|i - j|)) \exp \left[ -\frac{(r_{ij} - r_{ij}^{\mu})^2}{2\sigma_{ij}^2} \right],
$$

The index $\mu$ runs over all $N_{\text{mem}}$ memory proteins to which the protein has previously been aligned using a sequence-structure threading algorithm\textsuperscript{29} (i.e. each $i$-$j$ pair in the protein has an $i'$-$j'$ pair associated with it in every memory protein; if, due to gaps in the alignment, there is no $i'$-$j'$ pair associated with $i$-$j$ for a particular memory then this memory protein simply gives no contribution to the interaction between residues $i$ and $j$). The interaction between $C_{\alpha}$ and $C_{\beta}$ atoms is a sum of Gaussian wells centred at the separations $r_{ij}^{\mu}$ of the corresponding memory atoms. The widths of the Gaussians are given by $\sigma_{ij} = |i - j|^{0.15}$ Å. The scaling factor $a$ is used to satisfy Eq. (1).

The weights given to each well are controlled by $\gamma(P_{i}, P_{j}, P_{\mu}^{i}, P_{\mu}^{j}, x(|i - j|))$, which depends on the identities $P_{i}$ and $P_{j}$ of the residues to which $i$ and $j$ are aligned, as well as the identities $P_{i}$ and $P_{j}$ of $i$ and $j$ themselves. The self-consistent optimisation calculates the $\gamma$ parameter which originates the cooperative folding in the model. A three well contact potential [Eq. (3)] is used for residues separated by more than 12 residues as described by

$$
E_{\text{contact}} = -\frac{\epsilon}{a} \sum_{i<j-12} \sum_{k=1}^{3} \gamma(P_{i}, P_{j}, k)c_{k}(N)U(r_{\text{min}}(k), r_{\text{max}}(k), r_{ij}).
$$
Here, the sequence indices \( i \) and \( j \) sum over all pairs of \( C_\beta \) atoms separated by more than 12 residues. The sum over \( k \) is over the three wells which are approximately square wells between \( r_{\text{min}}(k) \) and \( r_{\text{max}}(k) \) defined by,

\[
U(r_{\text{min}}(k), r_{\text{max}}(k), r_{ij}) = \frac{1}{4} \left\{ [1 + \tanh \left( 7\frac{r_{ij} - r_{\text{min}}(k)}{\AA} \right)] + [1 + \tanh \left( 7\frac{r_{\text{max}}(k) - r_{ij}}{\AA} \right)] \right\}.
\]

(4)

The parameters \((r_{\text{min}}(k), r_{\text{max}}(k))\), are \((4.5\,\AA, 8.0\,\AA)\), \((8.0\,\AA, 10.0\,\AA)\), and \((10.0\,\AA, 15.0\,\AA)\) for \( k = 1, 2 \) and \( 3 \) respectively. In order to approximately account for the variation of the probability distribution of pair distances with number of residues in the protein \( (N) \) a factor \( c_k(N) \) has been included in \( E_{\text{long}} \). It is given by \( c_1 = 1.0, c_2 = 1.0/(0.0065N + 0.87) \) and \( c_3 = 1.0/(0.042N + 0.13) \).

The individual wells are also weighted by \( \gamma \) parameters which depend on the identities of the amino acids involved, using the 4-letter code defined above. In contrast to the interactions between residues closer in sequence, this part of the potential does not depend on the database structures that define local-in-sequence interactions.

To pinpoint the effects of frustration or favorable non-native contacts always present in any coarse gained protein model, we simulated a perfectly smooth energy function often called a Gō model.\(^{30}\) Gō models are an essential tool for understanding protein folding kinetics.\(^{31,32}\) While having the same backbone terms,\(^{33}\) in this single structure based Hamiltonian [Eq. (5)], all of the interactions, \( E_{\text{int}} \) are defined by Gaussians whose minima are located at the pair distribution found in the experimental structure:

\[
E_{\text{Gō}}^{\text{AM}} = -\frac{\epsilon}{d_{\text{Gō}}} \sum_{i \leq j < 3} \gamma_{\text{Gō}}(x(|i - j|)) \exp \left[ -\frac{(r_{ij} - r_{ij}^N)^2}{2\sigma_{ij}^2} \right].
\]

(5)

The global minima of such an energy function should be the input structure.

Many have used additional constraining potentials to characterise unsampled regions of coordinate space while using molecular dynamics.\(^{15,34}\) To characterize the landscape sampled with basin-hopping, we also used a structure constraining potential to identify ensembles with fixed but varying fractions of native structure. Using such a potential allows us to access interesting
configurations that are unlikely to be thermally sampled. The constraining potential also called umbrella potentials are centered on different values of an order parameter to sample along the collective coordinates. One of the collective coordinates is $Q$, an order parameter that measures the sequence-dependent structural similarity of two conformations by computing the normalized summation of C-alpha pairwise contact differences, as defined in Eq. (6).^{15}

$$Q = \frac{2}{(N-1)(N-2)} \sum_{i<j-1} \exp \left[ -\frac{(r_{ij} - r_{ij}^N)^2}{\sigma^2_{ij}} \right].$$  \hspace{1cm} (6)

The resulting order parameter ranges from zero, where there is no similarity between structures, to one, which represents an exact overlap. The form of the potential is $E(Q) = 2500 \epsilon (Q - Q_i)^4$, where $Q_i$ may be varied in order to sample different regions of the chosen order parameter. As in equilibrium sampling, simulations were initiated at the native state and the $Q_i$ parameter was reduced throughout the sampling.

We have also studied the potential energy landscape when multiple surfaces are superimposed upon each other by the use of multiple homologous target proteins. This manipulation of the energy landscape has been shown to further reduce local energetic frustration that arises from random mutations in the sequence away from the consensus optimal sequence for a given structure. By reducing the number of non-native traps, this averaging often improves the quality of structure prediction results.\textsuperscript{35–38} As seen in Eq. (7), the form and the parameters of the energy function are maintained from Eqs. (2) and (3), but the normalized summation is taken over a set of homologous sequences:

$$E_{AM} = \frac{1}{N_{\text{seq}}} \sum_{k=1}^{\text{seg}} \sum_{i<j}^N E_{\text{int}}(P^k_i, P^k_j).$$  \hspace{1cm} (7)

Since proteins are not random heteropolymers, the differences in the energy function for homologous proteins are randomly distributed, therefore the mean over multiple energy functions should have less energetic variation than the original function. Indeed, performing this summation is a way of incorporating to the optimisation of the $T_f/T_g$ criterion into any energy function. The target sequences of the homologues can be identified using PSI-Blast with default parameters.\textsuperscript{39,40}
Some classes of proteins have a large number of sequence homologues, therefore performing a multiple sequence alignment can be impractical. Removing redundant sequences from within the set of identified homologues also removes biases that can be introduced where there are few homologues available. This is done by preventing sequences in the collected sequences from having greater than 90% sequence identity. The remaining sequences are aligned in a multiple sequence alignment.\(^{41}\) Gaps within the sequence alignment can be addressed within the AMH energy function in a variety of ways. In the present work, gaps in the target sequence were removed, while gaps within homologues were completed with residues from the target protein. While this procedure may introduce small biases toward the target sequence, it is preferable to ignoring the interactions altogether.

Finally, we made several \textit{ad hoc} changes to the backbone potential, \(E_{\text{back}}\). Eliminating some compromises necessary for rapid molecular dynamics simulations allowed the AMH potential to be adapted to basin-hopping. Another goal was to prevent the over-collapse of the proteins by altering the excluded volume energy term, which should reduce the number of states available during minimisation. The terms shown in Eq. (8) are used to reproduce the peptide-like conformations in the original molecular dynamics energy function:

\[
E_{\text{back}} = E_{\text{ev}} + E_{\text{harm}} + E_{\text{chain}} + E_{\text{chi}} + E_{\text{Rama}}.
\]  

\(E_{\text{ev}}\) maintains a sequence specific excluded volume constraint between the \(C_\alpha-C_\alpha\), \(C_\beta-C_\beta\), O-O, and \(C_\alpha-C_\beta\) atoms that are separated by less than \(r_{\text{ev}}\). Previously, \(^{26}\) we have seen that modifying \(E_{\text{back}}\) can produce a less frustrated energy surface when using thermal equilibrium sampling, but slow dynamics was often found to result since the local barrier heights became too large. The ability of basin-hopping to overcome such large, but local barriers allows us therefore to consider a potential whose dynamics would otherwise be too slow for molecular dynamics. In the final part
of the paper, we altered the excluded volume term, as shown in Eq. (9) to prevent over-collapse:

\[
E_{ev} = \epsilon \lambda_{C_{EV}} \sum_{x,y} \sum_{i<j} \theta(r_{C}^{C}(j-i) - r_{C_{EV}}^{C})(r_{C}^{C}(j-i) - r_{C_{EV}}^{C})^2 + \epsilon \lambda_{O_{EV}} \sum_{i<j} \theta(r_{O}^{O}(O_{EV} - r_{O_{EV}}^{O}) (r_{O}^{O}(O_{EV} - r_{O_{EV}}^{O}))^2,
\]

(9)

by changing the default molecular dynamics parameters, \(\lambda_{C_{EV}} = 20\), \(\lambda_{O_{EV}} = 20\), \(r_{C}^{C}(j-i < 5) = 3.85\ \text{Å}\), \(r_{C}^{C}(j-i \geq 5) = 4.5\ \text{Å}\), and \(r_{O}^{O} = 3.5\ \text{Å}\), to \(\lambda_{C_{EV}} = 250\), \(\lambda_{O_{EV}} = 250\), \(r_{C}^{C}(j-i < 5) = 3.85\ \text{Å}\), \(r_{C}^{C}(j-i \geq 5) = 3.85\ \text{Å}\), and \(r_{O}^{O} = 3.85\ \text{Å}\). The force constant are over an order of magnitude larger than those used in the molecular dynamics, and the radii of the \(\text{C}_\alpha\), \(\text{C}_\beta\), and \(\text{O}\) atoms are also 10% larger than previously used values. This increase in excluded volume slows the onset of chain collapse, but improves steric interactions. The other change to the backbone potential involves terms which maintain chain connectivity. In molecular dynamics annealing, covalent bonds are preserved using the SHAKE algorithm,\(^{42}\) which permits an increase of the molecular dynamics time step. For basin-hopping in all parts of this paper, we removed the SHAKE method and replaced it with a harmonic potential, \(E_{\text{harm}}\), between the \(\text{C}_\alpha-\text{C}_\alpha\), \(\text{C}_\alpha-\text{C}_\beta\), and \(\text{C}_\alpha-\text{O}\) atoms. This replacement permits the location of local minima without requiring an internal coordinate transformation, and avoids discontinuous gradients. When minimised, the additional harmonic terms typically contribute only only .015 \(k_B T\) per bond. The remaining terms of the original backbone potential are maintained. Depending on the sidechain, the neighbouring residues in sequence sterically limit the variety of positions the backbone atoms can occupy, as evidenced in a Ramachandran plot.\(^{43}\) This distribution of coordinates is reinforced by a potential, \(E_{\text{Rama}}\), with artificially low barriers to encourage rapid local movements. The planarity of the peptide bond is ensured by a harmonic potential, \(E_{\text{chain}}\). The chirality of the \(\text{C}_\alpha\) centres is maintained using the scalar triple product of neighbouring unit vectors of carbon and nitrogen bonds, \(E_{\text{chi}}\).

For basin-hopping simulations, whose algorithm is outlined in Fig. 2 the most important sampling parameters are the temperature used in the accept/reject steps for local minima \(T_{bh}\), and the maximum step size for perturbations of the Cartesian coordinates \(d\). A higher temperature allows transitions to an increased energy minima to be accepted, and also creates a larger the number of iterations typically required to minimise the greater perturbed configurations. Too high a tem-
perature leads to insufficient exploration of low-energy regions. The temperature ($T_{bh}$) for these simulations was $10 T_{amc}$. Lower temperatures resulted in slower escape rates from low energy traps, while higher temperatures prevented adequate exploration of low energy regions. The step size needs to be large enough to move the configuration into the basin of attraction of one local minimum to a neighbouring one, but not be so large that the new minimum is unrelated to the previous state. Every Cartesian coordinate was displaced up to a maximum step size ($d$) of 0.75 A, the optimum value determined from preliminary tests. Each run consisted of 2500 basin-hopping steps saving structures every 5 basin-hopping steps. The convergence condition ($\delta E_{\text{min}}$) on the root-mean-square (RMS) gradient for each minimisation was set to $10^{-3} \epsilon$, and the 5 lowest-lying minima from each run were subsequently converged more tightly ($\delta E_{\text{final}}$) to an RMS gradient of $10^{-5} \epsilon$. It is important to note that basin-hopping does not provide equilibrium thermodynamic sampling. In structure prediction there, however, is no rigorous need for the search to obey detailed balance since the global energy minimum is the primary interest. Basin-hopping provides a means for the optimal global search of the energy landscape, however other methods must be used when calculating free energy and entropy.

In previous structure prediction studies with the AMH, low energy structures were identified using off-lattice Langevin dynamics with simulated annealing, employing a linear annealing schedule of 10000 steps from a temperature of 2.0 to 0.0, starting from a random configuration. The number and length of simulations needed in both strategies were determined by the number of uncorrelated structures encountered. The current basin-hopping method with the AMC energy function encounters roughly one deep trap per run. In order to sample 100 independent structures in molecular dynamics 20 separate runs were needed, because simulated annealing samples about five independent states before the glass transition temperature is encountered, as measured by the rapid decay of structural correlations. We compared several $\alpha$ helical proteins, both from within and outside the training set of the AMH energy function.
3 Results and Discussion

We performed initial calculations with a Gō potential for the 434 repressor (protein data bank (PDB) ID 1r69). In Fig. 3 we show this model accurately represents the native basin. Steps where the energy increases are allowed by the sampling method and are not examples of frustration. Studies on the Gō model provide a useful benchmark for comparing the computer time required for the different global optimisation strategies. Using the sampling parameters used in this report, we compared the time for initial collapse between the molecular dynamics and basin-hopping runs. The initial collapse required about 7 minutes for the annealing runs and 31 minutes for basin-hopping on a desktop computer. However, these values do not reflect the actual performance of the two approaches in locating global minima, which will depend upon the move sets, step size, temperature, and convergence criteria.

While using the AMC structure prediction Hamiltonian, we found that basin-hopping was often able to locate lower energy structures and also identified minima that have greater structural overlap with the native state than annealing. These results are produced for structure predictions for proteins both inside and outside the training set, as demonstrated in Table 1. The first three proteins (PDB ID 1r69, 3icb, 256b) in Table 1 are in the training set of the Hamiltonian25, while the other three are not, and can therefore be considered as predictions from the algorithm. The minima located with basin-hopping show an increase in structural overlap with the native state [Eq. (6)] when compared to the Langevin dynamics approach. $Q$ scores of 0.4 for single domain proteins generally correspond to a low resolution root mean square deviation (RMSD) of around 5 Å or better. $Q$ scores of 0.5 and higher have still more accurate tertiary packing and are of comparable quality to the experimentally derived models. The high quality structures obtained suggest the form of the backbone terms is appropriate, since the physically correct stereochemistry is reproduced. Lower energy structures are sampled by basin-hopping for the non-training set proteins, but the structural overlap improvement found in these deeper minima was smaller. Larger proteins pose a greater challenge for basin-hopping with this Hamiltonian due to the random steps in Cartesian coordinates. Dihedral coordinate move steps would probably be
more efficient, and will be considered in future work.

The distribution of minima encountered from multiple simulations for both search methods is shown in Fig. 4 where a greater density of high quality structures is obtained by the basin-hopping algorithm. The potential energy surface still includes, therefore, significant residual frustration in the near-native basin in the form of low-lying minima separated by relatively high barriers. Without the parameter optimisation to reduce frustration, folding would exhibit more pronounced glassy characteristics. Most of the cooperative folding occurs during collapse until $Q$ values of around 0.4 are reached. While the structures from simulated annealing are accurate enough for functional determination, we see basin-hopping can better overcome barriers that are created after collapse. The density of the high quality structures is also important for post-simulation k-means clustering analysis. Another way of representing the data of a set of independent basin-hopping simulations is by selecting the lowest energy structures from each simulation of the 434 repressor (PDB ID 1r69) and HDEA (PDB ID 1bg8) proteins and ordering them with respect to their structural overlap. As shown in Fig. 5, the protein in the training set (434 repressor) produces better results than the non-training protein, as expected.

We have decomposed the different energy terms in the Hamiltonian in Table 2, to examine which interactions are most effectively minimised. The AMC potential has three different distance classes in terms of sequence separation, and these are defined as short ($|i - j| < 5$), medium ($5 \leq |i - j| \leq 12$), and long ($|i - j| > 12$). Most importantly, the long-range AMH interactions are successfully minimised in the basin-hopping runs, due to the ability of basin-hopping to overcome large energetic barriers. This term will govern the quality of structures sampled using an approximately smooth energy landscape. The other terms that define secondary structure formation are not as well minimised. This result is due to the disruption of helices by the random Cartesian perturbation move steps. These move steps benefit favorable steric packing and therefore do well at minimising the excluded volume energy term of the Hamiltonian. A combined minimisation approach might be more efficient, where larger dihedral steps could be made early during minimisation to sample a wider number of structures, followed by random Cartesian steps.
to optimise the steric interactions.

While we sampled high quality structures, we would like to confirm that we have completely sampled the global minima of the energy surface. To access these unsampled states we used umbrella potentials. When constraining a set of simulations to different values of $Q$, we have obtained energy minima for cytochrome c, roughly 15 $\epsilon$ deeper than those from unconstrained minimisations starting with a randomized structure, as shown in Fig. 6. For the 434 repressor the minima obtained from randomized states and those found with the $Q$ constraints applied differ by only a few $k_B T$. This shows basin-hopping does indeed act as a global optimisation method, by accurately identifying the global energy minimum from multiple independent unconstrained simulation. This behavior is predictable from the choices that governed the design of the Hamiltonian. Low energy barriers between structures are desirable during a molecular dynamics simulation because they accelerate the dynamics. However, for basin-hopping these low barriers encourage tertiary contact formation before secondary structure units condense for sequences greater than 110 amino acids.

**Superposition of Multiple Energy Landscapes**

Constructing a Hamiltonian by calculating the arithmetic average of the potential over a set of homologous sequences increased the quality of predictions in both equilibrium and annealing simulations. We have found this approach also improved the performance in basin-hopping simulations. For two different proteins, 100 independent basin-hopping runs were performed with both the standard and sequence-averaged Hamiltonians. By the superposition of multiple energy landscapes we saw a reduction in the number of competing low energy traps around $Q$ values of 0.3 for both the 434 repressor and uteroglobin (PDB ID 1UTG), as shown in Fig. 7. Improvement of structure prediction Hamiltonians can be statistically described by the average energy gap between the native basin and a set of unfolded structures, and by the roughness of the energy surface, which corresponds to the variance of the energy. The sequence based energy function summations limited the energetic variance of the sampled landscapes, thereby reducing the glass transition temperature. This improvement, even at the low temperatures sampled in basin-hopping, is predicted
from theory, but difficult to observe in conventional equilibrium simulations due to the emergent
glassy dynamics, which slows the kinetics. The energy gap improvement was smaller than the
reduction of the energetic variation of the Hamiltonian. In terms of the goal of maximizing the
ratio of $T_f/T_g$, this increase came primarily from to reducing the glass transition temperature $T_g$.
In the low energy region we saw fewer competing states, and an increased correlation between $E$
and $Q$ for the sequence-averaged Hamiltonian compared to the original Hamiltonian. For the 434
repressor the lowest energy structure had the highest $Q$ value encountered.

**Characterisation of Polymer Collapse**

When we annealed the Hamiltonian using molecular dynamics we observed some over-collapse of
the polypeptide chain, producing a smaller radius of gyration than the experimental structure.
In basin-hopping runs we also found structures exhibiting a larger number of contacts than the
experimental structure, as show in Fig. 8 where a contact is defined as a $C_\alpha-C_\alpha$ distance of
less than 8 Å. While the low-energy structures may be native-like, these structures were more
compact than those observed experimentally. To investigate this behavior, we examined the
backbone and interaction terms of the Hamiltonian separately using the Gō Hamiltonian in Eq. (5).
Somewhat surprisingly, the Gō model also produces over-collapse, as shown in Fig. 9. Hence the
interaction parameters of the structure prediction Hamiltonian were not responsible for all of
the over-collapse. These minimal model-dependent frustrations were only eliminated in the final
stages of minimisation. The most effective technique for reducing over-collapse was to increase the
force constant and the atomic radius in the excluded volume terms [Eq. (9)]. The barrier crossing
capabilities of basin-hopping steps produce more over-collapse than do the annealing minimisations
without these parameter changes. The glass-like transition seen in simulated annealing prevents
further collapse in molecular dynamics, as the rearrangement rates slow down exponentially with
temperature. The improved parameter set of Fig. 10 shows more native-like collapse, but the
lowest energy structures had $Q$ values of 0.36 and the best $Q$ value was 0.45, which are worse than
basin-hopping simulations with the original parameters.
4 Conclusion

In this report we have demonstrated that minima with lower energy and higher quality structures can often be located for the AMH potential using basin-hopping global optimisation compared to annealing. Encouragingly, the long range in sequence energy contributions are better minimised than with simulated annealing. Umbrella sampling using basin-hopping can also show when the global minima are reached for a selected order parameter. Previous techniques for reducing the energetic variance of the energy surface in simulated annealing are also applicable to basin-hopping. Using basin-hopping also permits improvements in certain backbone terms of the Hamiltonian. These changes would make the kinetics too slow in molecular dynamics annealing runs, but larger barriers can be easily crossed using basin-hopping.

These results highlight future optimisation strategies where the deep non-native traps found by basin-hopping could be used as decoys for further parameter refinement, rather than the higher-lying minima obtained by quenching with simulated annealing. This re-optimisation of the potential results in getting a better estimate for $T_f/T_g$ now possible because of the efficiency of the basin-hopping algorithm at identifying low energy decoys. Another future direction would be evaluating the equilibrium properties of low-lying structures identified by basin-hopping also provides a means to calculate free energy barriers, which would be difficult to characterise via conventional simulations.

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**Figure Captions**

1. In the basin-hopping approach the original potential energy surface (solid) is transformed into a set of plateaus (dashed). The local minima are not changed, but the transition state regions are removed.

2. The basin-hopping algorithm is defined by a few parameters that allow its transfer to different systems.

3. Variation of the energy of the current minimum as a function of $Q$ for minima encountered in the Markov chain during a basin-hopping run using a Gō model. Steps that increase the energy are sometimes allowed by the Monte Carlo criterion, which employed a temperature of $10 \ k_b/\epsilon$.

4. Energy as a function of $Q$ for local minima of 434 repressor encountered during 100 independent basin-hopping optimisations (top) and 20 annealing simulations (bottom).

5. The lowest energy structures of the training set protein, 434 repressor (top) and the blind prediction proteins, HDEA (bottom) identified from 100 independent basin-hopping simulations. Each minimum has values for energy illustrated by the dots and structural overlap to the native state $Q$ represented by lines. These minima are ordered with respect to their structural overlap $Q$ with the native state (Index). The data shows correlations between the energy and $Q$, while the number of high quality structures is superior for the training protein.

6. Energy as a function of $Q$ for the 434 repressor and cytochrome c proteins obtained in basin-hopping calculations with the structure prediction Hamiltonian. These runs employed an additional umbrella potential that constrains the simulation to different values of $Q$. The results for the 434 repressor are similar to the unconstrained basin-hopping results, but the structures for cytochrome c are $15\epsilon$ lower in energy than those found in unconstrained basin-hopping runs.
7. Energies of local minima obtained using basin-hopping with the original and a sequence-averaged Hamiltonian for two training proteins. Importantly for both the top graph (434 repressor) and the bottom graph (uteroglobin) fewer non-native states are seen with the sequence averaged (red) Hamiltonian when compared to standard Hamiltonian (black).

8. Results of 100 independent basin-hopping runs for the 434 repressor using the set of backbone parameters that was optimised for molecular dynamics. Structures were saved every 20 basin-hopping steps. The ratio of contacts to native state contacts shows that most of the structures are more compact than the native state.

9. A Gō potential simulation for the 434 repressor shows a modest amount of over-collapse during a basin-hopping simulation, which is resolved as the structure approaches a $Q$ value of 1.0.

10. Results of 100 independent basin-hopping runs for the 434 repressor using the set of backbone parameters that was optimised for molecular dynamics. Structures were saved every 20 basin-hopping steps. An altered set of backbone parameters produces structures that have similar collapse behavior when compared to the native state.
Figures

Figure 1:
Basin Hopping Algorithm

*Monte Carlo Step* (n steps)

random Cartesian move step with maximum distance (d) and temperature (T_{bh})

*Minimisation*

L-BFGS quasi-Newtonian method for optimization

convergence condition ($\delta E_{min}$) is RMS gradient of $10^{-3} \epsilon/r$

*Minimisation with tight convergence* (after n steps)

convergence condition ($\delta E_{final}$) is RMS gradient of $10^{-5} \epsilon/r$

**Figure 2:**
$E(Q)/\epsilon$
Figure 4:
Figure 5:
Figure 6:
Figure 8:
Figure 9:
Figure 10:

Basin-Hopping Steps

Total Contacts/Native Contacts

0 10 20 30 40 50

0 0.6 1.0 1.4

0.6 0.8 1.0 1.2 1.4

Figure 10:
### Tables

**Table 1:** Minima located by molecular dynamics/annealing (MD) and basin-hopping (BH); the first three proteins are in the training set of the Hamiltonian, while the results for the second three proteins are predictions.

| PDB ID | length | Lowest $E$ | $Q$ | Highest $Q$ | Lowest $E$ | $Q$ | Highest $Q$ | $E$ |
|--------|--------|------------|-----|-------------|------------|-----|-------------|-----|
| 1r69   | 63     | -428.92    | 0.39| 0.53        | -307.96    | 0.39| 0.52        | -408.482|
| 3icb   | 75     | -536.98    | 0.47| 0.52        | -390.54    | 0.40| 0.49        | -518.92|
| 256b   | 106    | -735.02    | 0.42| 0.65        | -707.51    | 0.37| 0.40        | -716.51|
| 1uzc   | 69     | -457.55    | 0.36| 0.42        | -383.08    | 0.37| 0.45        | -433.41|
| 1bg8   | 76     | -469.49    | 0.25| 0.34        | -465.19    | 0.36| 0.39        | -461.50|
| 1bqv   | 110    | -737.91    | 0.21| 0.27        | -441.92    | 0.23| 0.27        | -481.22|
Table 2: Contribution of different energy terms in local minima obtained using molecular dynamics/annealing (MD) and basin-hopping (BH).

| PDB | Method | length | Ex Vol | Rama   | Short Range | Medium Range | Long Range |
|-----|--------|--------|--------|--------|-------------|--------------|------------|
| 1r69 | MD     | 63     | 9.77   | −101.64| −128.90     | −84.87       | −123.28    |
| 1r69 | BH     | 63     | 2.65   | −91.06 | −125.04     | −84.80       | −137.57    |
| 3icb | MD     | 75     | 11.74  | −127.70| −177.21     | −90.11       | −153.69    |
| 3icb | BH     | 75     | 4.40   | −115.76| −178.47     | −83.37       | −173.38    |
| 1uzc | MD     | 69     | 10.10  | −118.66| −134.00     | −90.75       | −124.24    |
| 1uzc | BH     | 69     | 2.22   | −106.20| −137.95     | −92.40       | −123.77    |
| 1bg8 | MD     | 76     | 11.68  | −136.39| −173.45     | −94.40       | −76.94     |
| 1bg8 | BH     | 76     | 2.72   | −112.13| −151.95     | −94.23       | −113.09    |