Modeling large regions in proteins: Applications to loops, termini, and folding

Aashish N. Adhikari,1,2 Jian Peng,3 Michael Wilde,5 Jinbo Xu,3 Karl F. Freed,1,2,4* and Tobin R. Sosnick4,5,6*

1Department of Chemistry, The University of Chicago, Chicago, Illinois 60637
2The James Franck Institute, The University of Chicago, Chicago, Illinois 60637
3Toyota Technological Institute at Chicago, Chicago, Illinois 60637
4Computation Institute, The University of Chicago and Argonne National Laboratory, Chicago, Illinois 60637
5Department of Biochemistry and Molecular Biology, The University of Chicago, Chicago, Illinois 60637
6Institute for Biophysical Dynamics, The University of Chicago, Chicago, Illinois 60637

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Abstract: Template-based methods for predicting protein structure provide models for a significant portion of the protein but often contain insertions or chain ends (InsEnds) of indeterminate conformation. The local structure prediction “problem” entails modeling the InsEnds onto the rest of the protein. A well-known limit involves predicting loops of \( \leq 12 \) residues in crystal structures. However, InsEnds may contain as many as \( \sim 50 \) amino acids, and the template-based model of the protein itself may be imperfect. To address these challenges, we present a free modeling method for predicting the local structure of loops and large InsEnds in both crystal structures and template-based models. The approach uses single amino acid torsional angle “pivot” moves of the protein backbone with a \( C_b \) level representation. Nevertheless, our accuracy for loops is comparable to existing methods. We also apply a more stringent test, the blind structure prediction and refinement categories of the CASP9 tournament, where we improve the quality of several homology based models by modeling InsEnds as long as 45 amino acids, sizes generally inaccessible to existing loop prediction methods. Our approach ranks as one of the best in the CASP9 refinement category that involves improving template-based models so that they can function as molecular replacement models to solve the phase problem for crystallographic structure determination.

Keywords: long loops; insertions; loop modeling; local protein structure prediction; molecular replacement

Introduction
Homology-based methods use known structures as templates and have proven extremely successful in modeling larger proteins in a computationally efficient fashion. The success of these methods, however, depends on the quality of the alignments between the target sequence and those of the templates.1 Frequently, the
sequence alignments contain gaps that correspond to regions in the sequence where no reliable structural information can be extracted from the templates. These gaps may be insertions or additions at the termini (Fig. 1). Inevitably, the model assembled from the templates lacks these local regions. In order to model the entire structure, alternative methods are required. The problem of reconstructing local regions in a protein is neither new nor exclusive to homology modeling. Experimentally determined structures from crystallography often contain regions that are difficult to characterize, because they are flexible or mobile. Consequently, crystal structures can contain loops that have weak or missing electron density. This issue is particularly significant because protein function is often mediated by loops; for example, loops often act as molecular recognition or binding sites and play a crucial role in executing the protein’s function.2–4 The specificity of protein interactions as mediated by active sites and binding pockets is also a consequence of local protein structure. These issues highlight the need for reliable methods to reconstruct local regions in protein structures.

Three important problems arise in developing methods for predicting local spatial structure. First, the local regions must be modeled subject to the constraints imposed by the rest of the protein structure. For example, the loop termini must end at the correct anchor positions. Some approaches to this long-standing “loop closure problem” seek analytical solutions to bond angles that properly position the ends.5–7 Although exact solutions have been found for short polypeptide segments, no general analytical solution is possible for segments containing more than a few amino acids in proteins. Other robotics-inspired algorithms for loop closure8,9 likewise experience decreasing accuracy as the size of the loops increases. Additionally, analytical approaches to the closure problem very often yield solutions that place backbone dihedral angles in disallowed regions of Ramachandran (Rama) space and thus generate sterically forbidden conformations.

Second, irrespective of how the loop closure is performed, a procedure is required for sampling various conformations of the local region. Existing approaches for predicting local regions in protein structures can be broadly categorized into two classes: database and de novo (free modeling) methods.10,11 Database methods search for loop fragments that best match the anchor geometries,10,12 but these approaches usually are confined to short insertions because of poor database coverage for larger fragments. Although these methods tend to be fast, the speed comes at the cost of the greater flexibility in exploring the conformational space of the loops permitted by free modeling methods. The applicability of these methods is further challenged in the modeling of InsEnds in template-based models because the regions are likely to correspond to parts of the sequence that are inaccessible to the homology methods.

In contrast, de novo methods sample sterically feasible loop conformations that are scored with physics-based or statistical potentials. For example, MODELLER places loop atoms uniformly between the anchor positions and optimizes the atom positions...
using conjugate gradient and molecular dynamics with simulated annealing, scoring the loops using a combination of the CHARMM22 force field and statistical preferences of the dihedral angles and atom contacts. Other free modeling methods such as RAPPER and PLOP build loop fragments by sampling from a dihedral angle library for each residue, beginning from one or both anchors and eventually attempting to close the loop while avoiding steric clashes.

The third challenge is associated with the scoring of various conformations. Because the number of residues whose conformation varies between the different structural candidates is small, accurate energy functions can be crucial to guide the conformational search and score the final structures. Both statistical potentials and physics-based force fields have been used as scoring functions in loop-building. Some methods use statistical potentials only for filtering, while the final ranking uses all atom force fields. Other methods focus on all atom energy functions designed specifically for loop modeling. However, energy functions that are good at guiding the conformational search during the loop-building stage might be inadequate for the final ranking of the decoys, especially in methods where the loop building is performed incrementally and separately from closure.

Until recently, investigations of local protein structure have largely centered on predicting loops in defined crystal structures. However, InsEnd predictions are made in the context of template-based models where the structures for the remainder of the protein may be imperfect, being constructed from one or more crystal structures and relying on a sequence alignment. As a result of this imperfection, the structure prediction algorithm must be lenient, thereby fundamentally distinguishing this problem from traditional loop modeling.

Although the treatment of both loops and InsEnds involves local protein modeling, they can present different sets of challenges. Whereas loops in crystal structures are defined as regions connecting different secondary structure elements, InsEnds are defined as regions devoid of information extracted from sequence alignments. Hence, InsEnd may include regions with complete secondary structure elements. In addition, the length of loops is governed by the structural context, and, consequently, the loops usually contain a limited number of residues. InsEnds, on the other hand, can be of arbitrary lengths. Furthermore, the boundaries of loops are generally well defined whereas the boundaries of InsEnds are determined by the gaps in the alignments. When multiple templates are combined to generate one model, the gap regions may appear with different boundaries in different templates, thereby rendering the InsEnds boundaries ambiguous.

Our method is designed to address these issues. We demonstrate the robustness of our methods by successfully predicting the structures of long loop regions in crystal structures as well as providing blind structural predictions of InsEnds in the top homology models from our submissions to CASP9. We present a fragment-free method for local structure prediction.

**Approach**

Our approach assumes that the principles governing the folding of proteins are equally applicable for modeling InsEnds. We have shown that single backbone pivot moves provide an effective way to sample conformations, provided the moves are contingent upon the identities and conformations of the nearest neighbors (NNs). These moves have been used successfully in the fragment-free de novo prediction of the structures of single-domain proteins.

Our local structure prediction method generates random local conformations using the same single pivot move set as for our global structure prediction scheme (Fig. 2). The interaction energy is calculated both within the local regions and between the local region and the rest of the protein. The total energy is used to guide the conformational search, an approach that differs from many methods in
which the loop fragment is constructed one residue at a time while simultaneously trying to satisfy the loop closure constraint at the end. In contrast to some existing methods that separate loop building and closure into two subsequent stages, our approach integrates the two into a single Monte Carlo simulated annealing (MCSA) scheme, thus retaining the tertiary context of the entire protein during the simulation while attempting to rapidly find the best local conformation. This tertiary context can be critical for identifying crucial loop–protein interactions, thus greatly reducing the search space. The algorithm is designed to handle multiple loops in the same MCSA trajectory. Hence, when two loops are close enough to interact, they are modeled simultaneously.

The conformational search proceeds by using an MCSA scheme (described in detail in the Methods section), guided by a combination of the pair wise additive, orientationally dependent statistical potential DOPE-PW, along with a harmonic ligation energy term to close the loop (Figure 3). The relative weight of the ligation energy increases during the MCSA to enforce loop closure. Explicit side chains are absent during the sampling stage of the simulation since the DOPE-PW statistical potential and backbone torsional move set implicitly incorporates sufficient information concerning the side groups.20 Final conformations are scored using a combination of structural clustering and accessible surface area of the hydrophobic residues to select the best solutions. The standard deviation in the positions of a given loop residue in a cluster (i.e., the tightness of the cluster) provides a metric for assessing the local quality of the predictions for the loop.

**Results and Discussion**

Three different modeling scenarios are considered. First, we address the traditional loop modeling problem in crystal structures where the structure surrounding the loop is known. We next address InsEnds modeling as applied in the CASP9 blind prediction competition, where the InsEnds are as large as 45 aa regions in template-based models generated by Xu’s RAPTOR-X algorithm.22 The third scenario is for the CASP9 refinement category in which the InsEnds algorithm is applied to the best structure from the server predictions and where the starting model and boundaries are specified by the organizers.

**Loops in crystal structures**

In order to demonstrate the applicability to larger loops in crystal structures, 26 loops of lengths 8–12 have been randomly selected from standard loop benchmarking studies.15 Loop boundaries for each target are taken as previously specified, and the loops are modeled using our method. Figure 4 illustrates the process of selecting the top five predictions, and Table I presents the best and the remaining four top predictions. After the predictions are clustered according to the RMSD between the loop structures, the largest five clusters are ranked using a linear combination of the Z-scores for the cluster tightness (RMSD between structures in the cluster), size, and average DOPE-PW energy, defined as $Z_s$, $Z_a$, and $Z_E$, respectively,

$$\text{Cluster Score} = Z_s - Z_d - Z_t,$$

$$Z_i = \frac{X_i - <X_i>}{\sigma_i},$$

where the Z-score for the property of structure $X$, is $Z_i = (X_i - <X_i>)/\sigma_i$ and $<X_i>$ and $\sigma_i$ are the mean...
and standard deviation, respectively. After ranking the clusters, one representative from each cluster is selected by using a combination of the DOPE-PW energy and the SASA to obtain the top five predictions. Although DOPE-PW is very successful in guiding the protein backbone into a proper conformation based on the orientation of the $C_{\alpha}-C_{\beta}$ vectors, it is unable to resolve the details of solvation at an atomistic level because it is parameterized only at the $C_{\beta}$ level. Hence, explicit SASA calculations are necessary to properly account for the solvation energy.

As discussed in the Methods section, the SASA scores are determined from a combined ranking of the hydrophilic and hydrophobic ASAs. Similarly, the structures are ranked by using the DOPE-PW energy function as well. The structure with the lowest total DOPE-PW + SASA rank in a given cluster is taken as the predicted structure from that cluster. Models are discarded when the distance between the free end and the anchor point fails to return to within 1.5 Å of the initial distance. If the largest cluster contains less than 5% of the structures, the scoring for the top five candidates uses only the sum of DOPE-PW and SASA scores. As shown in Supporting Information Figure 1, the inclusion of SASA to the DOPE-PW energy improves the selection of the top structure in most cases compared with simply using DOPE-PW energy to select the top structure.

A residue-specific deviation quantifies the local confidence score of the prediction for each residue individually in each of the top five predictions. The local confidence scores are illustrated by color and thickness in Figure 4. The thicker (redder) portions in the predicted local region correspond to residues displaying the largest deviation within the cluster.

The simulations for loops of length 12 and 8–11 residues generate conformations with global loop RMSDs of 2.76 and 1.93 Å, respectively, where the RMSD is calculated for the loop residues after aligning the structures without the loop regions (Table II). These results can be compared with Table II of

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**Figure 4.** Selection of the top 5 loop predictions for 1xnb. After clustering, the largest 5 clusters are ranked based on Z-scores with respect to cluster tightness, size, and average DOPE-PW energy. Once ranked, a selection is made from each of the 5 clusters using DOPE-PW + SASA. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
Lee et al., which presents the minimum backbone RMSDs found using different existing loop sampling protocols. Simulations for 12 and 8 residue loops in crystal structures with the cyclic coordinate descent (CCD) protocol generates minimum RMSDs of 3.05 and 1.59 Å, the CJSD method obtains 2.34 and 1.01 Å, the self-organizing algorithm using an alternating scheme of pairwise distance adjustments (SOS) yields 2.29 and 1.19 Å and the FALC scheme finds 2.34 and 1.01 Å.

The average RMSDs of our top ranked predicted loops are 3.98 and 0.78 Å for loops with 12 and 8–11 amino acids, respectively (Table I). These results are comparable to those from two different methods, RAPPER and FALC, ranked by DFIRE as listed in Table IV in Lee et al., where the average RMSDs of the top ranked 12 residue loop decoys for RAPPER and FALC are 4.32 and 3.84 Å, respectively. Rossi et al. compare four commercial loop modeling packages—Prime (Schrodinger, LLC), MODELLER (Accelrys Software), ICM (Molsoft, LLC), and Sybyl (Tripos)—which obtain RMSD values ranging from 3 to 5 Å for loops with 10–12 amino acids. Our performance is comparable to these methods.

We also compare our results to a recent atomic level modeling study which has sub-angstrom level accuracy. Although our Cα level modeling certainly limits us in terms of obtaining sub-angstrom models, we still are able to obtain models for some of the same benchmark proteins that are better than or comparable to the high-resolution Kinematic Closure (KIC) protocol (Supporting Information Table 1 of Ref. 26). For instance, our top predicted model for 1hfc with 3.69 Å RMSD outperforms KIC’s 8.2 Å prediction for the same loop. Similarly, our top predictions for the other targets 4i1b, 1msc, 1cyo, and 1pmy from our benchmark set in Table I yield RMSDs of 2.03, 5.87, 2.47, and 2.97 Å that are better or comparable to the high resolution KIC method’s top predictions of 3.8, 3.2, 5.2, and 2.6 Å.

### Table I. Prediction of Loops of 8–12 Residues in Crystal Structures

<table>
<thead>
<tr>
<th>Target</th>
<th>Loop length</th>
<th>Local loop RMSD (Align Loop, RMSD loop)</th>
<th>Global loop RMSD (Align rest, RMSD loop)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Best Pred1 Pred2 Pred3 Pred4 Pred5</td>
<td>Best Pred1 Pred2 Pred3 Pred4 Pred5</td>
</tr>
<tr>
<td>1rce</td>
<td>12</td>
<td>1.81 2.67 3.39 3.27 3.38 2.79</td>
<td>2.61 3.79 5.29 4.63 4.64 4.53</td>
</tr>
<tr>
<td>1thw</td>
<td>12</td>
<td>1.84 4.61 2.4</td>
<td>4.53 5.94 5.87</td>
</tr>
<tr>
<td>2cpl</td>
<td>12</td>
<td>1.09 1.09 4.14 4.02 4.72 4.37</td>
<td>2.43 2.43 6.34 6.94 6.84 8.37</td>
</tr>
<tr>
<td>1cvo</td>
<td>12</td>
<td>0.75 1.97 3.46 1.67 3.94 5.11</td>
<td>1.23 2.47 4.95 4.09 5.15 7.03</td>
</tr>
<tr>
<td>1hfc</td>
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<td>1.91 2.08 2.42 3.38 3.06 3.39</td>
<td>3.69 3.69 4.58 5.6 4.29 7.01</td>
</tr>
<tr>
<td>1onc</td>
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<td>2.19 3.24 2.19 2.95 2.94 2.99</td>
<td>2.91 3.83 5.26 5.36 5.66 5.22</td>
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<tr>
<td>1pmy</td>
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<td>1.24 2.97 3.13 2.98 4.07 7.02</td>
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<tr>
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<tr>
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<td>3.1 4.22 7.87 6.95</td>
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<tr>
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<tr>
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<td>3.64 4.19 5.32 5 4.66 4.33</td>
</tr>
<tr>
<td>1eco</td>
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<td>0.85 2.65 3.01 1.99 3.45 3.09</td>
</tr>
<tr>
<td>1msc</td>
<td>12</td>
<td>1.96 3.64 5.08 3.57 3.96 5.17</td>
<td>2.57 5.5 7.3 11.9 12.6 7.9</td>
</tr>
<tr>
<td>1acf</td>
<td>12</td>
<td>1.8 2.05 3.59 3.52 3.88 2.22</td>
<td>2.34 2.86 4.21 4.09 4.41 3.19</td>
</tr>
<tr>
<td>1cid</td>
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<td>1noa</td>
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<td>1.68 4.14 4.35 4.7 3.43 4.35</td>
<td>2.59 7.22 6.7 8.3 6.51 7.7</td>
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<tr>
<td>1plc</td>
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<td>3.42 3.42 4.89 4.08 8.35 3.57</td>
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<td>1.39 2.41 3.42 4.45 3.67 4.6</td>
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<td>4i1b</td>
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<td>2.03 2.03 9.71 10.3 10.83</td>
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<tr>
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<td>2.89 2.89 4.79 6.19 6.37 6.39</td>
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<tr>
<td>1aaj</td>
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<td>5p21</td>
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<td>5p22</td>
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<td>1.93 2.79 3.42 2.5 3.77 3.92</td>
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</tr>
<tr>
<td>1chs</td>
<td>8</td>
<td>0.74 3.65 0.74</td>
<td>1.74 5.41 1.74</td>
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<tr>
<td>1xnb</td>
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<td>0.28 1.49 1.21 0.53 1.16 1.13</td>
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</tr>
<tr>
<td>1paa</td>
<td>8</td>
<td>0.44 0.54 0.97 1.01 3.94 2.71</td>
<td>0.76 1.23 2.1 1.59 5.56 4.84</td>
</tr>
</tbody>
</table>

* Cases where the top cluster contain less than 5% of the total structures. In those cases, the top 5 predictions are selected using DOPEPW+SASA instead; units are in Å. The bold font indicates the best out of the top five predictions.

### Table II. Statistics for Loops in Crystal Structures

<table>
<thead>
<tr>
<th>Length</th>
<th>Local loop RMSD (Å)</th>
<th>Global loop RMSD (Å)</th>
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</thead>
<tbody>
<tr>
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<td>Average Stdev</td>
<td>Best Pred1 Stdev</td>
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<tr>
<td></td>
<td>Best Pred1 Stdev</td>
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<td>12</td>
<td>1.59 0.62</td>
<td>2.80 0.85</td>
</tr>
<tr>
<td>8–11</td>
<td>1.31 0.54</td>
<td>1.97 1.09</td>
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</tbody>
</table>
respectively, for the same 10–12 amino acid loops. The results demonstrate that a Cβ level representation of the protein chain without a costly analytical closure constraint is sufficient to achieve accuracy comparable with existing methods for relatively long loops in the context of crystal structures.

**Ends in crystal structures**

Another challenge involves modeling the termini of protein structures, a challenge that has attracted only limited study.27,28 Unlike loops, end regions require no loop closure. To demonstrate that our method is also applicable to end regions, we have refolded the termini for six proteins (Table III). In each of the case, 20 residues in the C terminal end of the native proteins are first randomized while the rest of the structure is kept fixed. Starting from these pseudo-random structures, the end residues are sampled and clustered using the loop modeling protocol. Because no loop closure is required, the termini are folded using only the DOPE-PW energy function.

In three out of six cases (1af7, 1o2f, 1r69), the best and the predicted structures have a global RMSD of under 3 Å (Table III), with the best local RMSD under 5 Å. Although direct comparisons are unavailable for the same proteins, the results are comparable to another method28 for refolding of terminal secondary structures where the number of unknown residues that require modeling is determined by the gaps in the sequence alignment and where reliable information about secondary structure type or boundaries is often unavailable.

**CASP9 blind InsEnds predictions**

Methods designed for predicting the structure of internal loops may be inappropriate for termini of proteins because the energy functions and sampling generally used for loop modeling assume both ends are fixed. Furthermore, InsEnds can encompass whole secondary structure elements. The existing loop modeling methods have been benchmarked for loops in crystal structures where the remaining structure and loop boundaries are known. The situation for homology modeling, however, is more complex, being highly dependent on the quality of the sequence alignments, template identification, and boundary determination. Consequently, the starting point for InsEnds modeling is imperfect and inexact.

The biannual CASP experiments present a unique platform for testing new and benchmarking developed methods through blind predictions. Our

<table>
<thead>
<tr>
<th>CASP9 refinement target</th>
<th>GDT starting</th>
<th>RMSD starting</th>
<th>GDT MidwayFolding</th>
<th>RMSD MidwayFolding</th>
<th>Rank of MidwayFolding</th>
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<tr>
<td>TR569</td>
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<td>76.58</td>
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<tr>
<td>TR568</td>
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<td>6.963</td>
<td>56.7</td>
<td>5.108</td>
<td>6/127</td>
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<tr>
<td>TR517</td>
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<td>4.646</td>
<td>72.17</td>
<td>4.638</td>
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<tr>
<td>TR622</td>
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<td>72.78</td>
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The numbers reported are the GDT and RMSD values from the CASP9 website. The values in bold indicate targets with an improvement in the GDT score from the starting model.

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Table IV. *Prediction of 20 End Residues in Crystal Structures*

<table>
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<th>Target</th>
<th>Type</th>
<th>Ends length</th>
<th>Local ends RMSD (Align Ends, RMSD Ends)</th>
<th>Global ends RMSD (Align rest, RMSD Ends)</th>
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<td>5.50</td>
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</tbody>
</table>
participation in CASP9 as MidwayFolding (groups TS435 and TS477) focused on testing our local structure prediction method and on improving poorly predicted local regions in template-based models. Our analysis begins with models generated by the program RAPTOR-X, which utilizes homology to identify template structures appropriate to a target sequence through sophisticated sequence/structure alignments. The templates are processed by MODELER to generate our starting model. We also use the sequence alignments of RAPTOR-X to identify the InsEnds regions in the models. Five entries may be submitted to CASP9 for each target, and Figure 5 displays the best of the five blind models submitted to CASP9 for each target.

The CASP9 targets serve to illustrate several strengths of our method. Several of the insertion regions contain secondary structure elements in the targets. The target T0464 from CASP8 presents a case where the insertion region is a helix, which our method predicts correctly, improving the model’s RMSD from 9.6 to 4.5 Å, as exhibited in Figure 5(A). Another target, T0623 has a 25-residue gap in a region that is, in fact, a hairpin that is correctly predicted by our method as well (8.2-Å RMSD improved to 6.3 Å).

The largest InsEnds contains 45 residues (T0585), and the RAPTOR-X+MODELER programs describe them as a large loop. Our method correctly identifies that the missing region corresponds to three helices that pack into the protein core, thereby improving the model substantially from 15.1 to 9.1 Å overall RMSD as depicted in Figure 5(H). The target TR606 presents an example where the local modeling is performed for both termini simultaneously to form a pair of beta strands, thereby improving the overall RMSD from 4.9 to 3.8 Å for the target as a result of modeling the ends [Fig. 5(G)].

Other CASP targets contain InsEnds that are loops connecting different secondary structures. For instance, the targets T0520, T0594, and T0612 yield initial models with loops containing as many as 17 residues (identified from the gap boundaries in the sequence alignments). Use of our InsEnds protocol for these three loop regions improves the overall RMSDs from 3.2 → 2.6 Å, 2.2 →1.7 Å and 7.3 → 6.6 Å for T0520, 594, and 612, respectively [Fig. 5(C,D)]. The demonstration that we successfully model various types of InsEnds with the same protocol without any prior knowledge of whether they are loops or contain secondary structure elements highlights the robustness of the method.

Blind prediction of refinement targets in CASP9

The judges for the refinement category in the CASP experiment select the best of all submitted (template-based) models from all participating groups. The local regions that deviate most from the native structure are identified to the predictors as the refinement targets. From our perspective, the
Refinement category is distinct because the starting model is guaranteed to be the best of all CASP server models rather than one of RAPTOR-X’s model and because the boundaries for InsEnds are specified based on where the server model differs from the native structure (as identified by the organizers) rather than from RAPTOR-X’s sequence alignment. On average for the 12 refinement targets, the 24 different refinement methods in CASP8 yield no net improvement over the starting models.29 Table IV lists the RMSD as well as the Global Distance Test (GDT) changes from the starting models along with the ranking of our method with respect to all the other refinement methods. Our method proceeds by first initializing the InsEnds regions to a completely random conformation, so that no structural information about the InsEnds is retained from the starting model.

Unlike the RMSD which relies on a single alignment, the GDT scores reflect the structural similarities at different distance cutoffs and, therefore, are generally better at assessing improvements in local regions.30 We have attempted 11 targets for refinement in CASP9 (Fig. 6) and improve the GDT scores for 7 of them. Among all groups participating in CASP9 refinement, four of our 11 predictions (targets TR517, TR568, TR569, and TR517) fall in the top 10% of all submissions, and eight of the 11 reside in the top 25% of all submissions, thereby outperforming several of the more costly all atom refinement methods. The improvements are achieved for targets with a wide range of starting GDTs (>50). The GDT/RMSD for TR569 improves from 73.1/3.01 Å to 76.58/2.24 Å, and our method ranks fourth out of the 121 total submissions for this target. The starting values for TR568 are lower at 53.35/6.39 Å.

Figure 6. InsEnds predictions for CASP9 refinement targets. Difference between CA/CA distance across the sequence of the initial (starting) /native and final (refined using InsEnds method)/native after superposition using sequence-dependent LGA protocol. Official data from CASP9 official website (http://predictioncenter.org/casp9/). For each target, the arrows indicate the regions where the InsEnds modeling has been performed. The blue to green color change designates regions where the InsEnds modeling improves upon the given target based on LGA superposition to native structures.
and we improve them to 56.7/5.1 Å with an overall ranking of sixth out of the 127 submissions for the target. Our method performs much worse than the rest of the methods for one target, TR592, presumably because the starting structure is already extremely good (91.2/1.2 Å) so that our Cβ level representation is inadequate, and, consequently, an all atom side chain representation is required to improve the model further. Moreover, we have not refined the side chains in any of the cases, something that probably would have improved the results even more.

Figure 6 displays all our predictions for the refinement targets in CASP9. The figure illustrates how well the model aligns to the native structure before refinement (initial) and after refinement (after) when superposed using the LGA program. The improvements introduced into the local region also help to align the remainder of the protein in several cases. For example, in TR614, even though the actual regions modeled are an insertion from 33 to 50 and the C terminal residues from 106 to 121, the local alignment of the N terminal residues improves over the starting model as indicated with blue in the LGA alignment for TR614 in Figure 6.

**Molecular replacement results for CASP9 refinement targets**

One of the CASP9 refinement metrics assesses how well the predicted models reproduce the experimental data. Recently, models generated by the structure prediction methods have been inserted into the molecular replacement likelihood algorithms for X-ray crystallographic refinement to solve the phase problem. The assessors for CASP9 refinement judge the quality of each submitted model in this regard by calculating the Z-score of the best orientation of the model in the unit cell of the crystal compared to placing it in a set of random orientations. Only models with Z-scores above 6 are considered good enough to solve the phase problem. Table 3 in Ref. 31 summarizes how often various groups improve the Z-score of the targets from likely unrefinable (<7) to likely refinable (>7). Our method performs as well or better than all the other groups in this test, with positive results in two of three cases attempted. As our approach uses a backbone + Cβ model with the side chains either missing or added simply using SCWRL4.0 with no further refinement, some of our submitted models were discarded in the analysis by assessors. Regardless, the fact that our method ranks at the top in the molecular replacement test proves its real value in X-ray crystal structure refinement.

In contrast to most other methods that expend considerable computing resources on including all-atom interactions, our method lacks explicit side chain atoms. This difference highlights the distinction between the refinement of crystal structures and template-based models. The all atom refinement of crystal structures benefits from having high-resolution information for the rest of the structure, whereas homology models are usually far from perfect. It is unclear whether the expensive modeling of all the atoms in an imperfect environment provides a computationally efficient strategy. In contrast, the first step of our approach is designed to obtain the proper backbone structure and orientation for the local region by using a coarse level of modeling that is less sensitive to the atomic level details for the rest of the homology model. Once the coarse level model is obtained for the local region, side chains may be added, and more detailed all-atom refinement can proceed.

**Global InsEnds RMSD versus local InsEnds RMSD**

RMSDs are calculated in three ways to help quantify the quality of the modeling of local InsEnds regions:

1. **Local InsEnds RMSD**: Align the loop and calculate the RMSD of only the InsEnds region.
2. **Global InsEnds RMSD**: Align all the residues besides the InsEnds, and then calculate the RMSD of the InsEnds region.
3. **Global structure RMSD**: Optimally align all the residues in the protein and calculate the RMSD of the full chain.

The local InsEnds RMSD is a measure of how well the InsEnds region itself is modeled, and the global InsEnds RMSD provides a measure of how well the modeled InsEnds is oriented with respect to the rest of the protein. The global InsEnds RMSD is the ideal measure of loop quality when predicting loops in crystal structures because the only difference between the native structure and the model can appear in the loop region. In contrast, InsEnds modeling of homology models begins from inexact structures; therefore, assessing the refinements requires accounting for the RMSD of the rest of the structure (besides the InsEnds) with respect to the native structure. If the starting homology model deviates significantly from the native structure, the alignment of the non-InsEnds region necessarily must skew the anchor regions, and therefore the global InsEnds RMSD would not provide as good a metric for reporting the accuracy of InsEnds modeling as does either the local InsEnds RMSD or the overall RMSD of the structure.

This utility of the different RMSDs is illustrated for six targets from CASP8 for which the initial RAPTOR models have variable RMSDs to the native structures. The 11–12 residue InsEnds regions in those models are chosen for (post-dictum) prediction using our method (Table V). Not surprisingly, the
global InsEnds RMSD is highly dependent on the quality of the initial model (i.e., the RMSD of all but the InsEnds region in the initial model). For target T0478D1, the RMSD of the non-InsEnds region in the starting model is 8.07 Å; the best local InsEnds RMSD decreases from 2.9 Å to 1.58 Å, whereas the best global InsEnds RMSD decreases from 12.2 Å to 8.4 Å. Target T0411D1 has a non-InsEnds RMSD of the starting model much closer to native structure at 2.74 Å, and our local InsEnds RMSD improves from 3.53 to 1.85 Å, similar to the local InsEnds RMSD improvement in T0478D1 (2.9 to 1.58 Å). However, the global InsEnds RMSD for this target improves from 10.2 to 2.78 Å, which is much more remarkable than the global InsEnds RMSD in T0478D1 (12.2 to 8.4 Å). The difference can be attributed to T0411D1’s starting model having the non-InsEnds region much closer to the native structure when compared with T0478D1. Figure 7 illustrates this behavior and indicates that the local InsEnds RMSD remains relatively unaffected, whereas the global InsEnds RMSD for the same targets is quite severely affected by the RMSD of the remaining region. The successes of the modeling also support our previous contention from protein structure predictions that the neighbor-dependent $\phi$, $\psi$ distributions capture local interactions reasonably well.\(^{20}\)

**Table V. Prediction of InsEnds in CASP8 structures (postdiction)**

<table>
<thead>
<tr>
<th>CASP8 target</th>
<th>InsEnds length</th>
<th>RMSD of InsEnds region</th>
<th>Initial model</th>
<th>Best</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>T0431D1</td>
<td>11</td>
<td>5.21</td>
<td>5.34</td>
<td>4.7</td>
<td>4.91</td>
</tr>
<tr>
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<td>2.03</td>
<td>2.04</td>
<td>1.91</td>
</tr>
<tr>
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<td>1.24</td>
<td>8.4</td>
<td>1.44</td>
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<tr>
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<td>1.13</td>
<td>7.3</td>
<td>1.25</td>
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<tr>
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<td>1.23</td>
<td>3.1</td>
<td>1.31</td>
</tr>
<tr>
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<td>1.54</td>
<td>0.91</td>
<td>1.54</td>
<td>0.83</td>
</tr>
</tbody>
</table>

**Figure 7.** Global versus local RMSD. The RMSD of non InsEnds region is plotted against the global InsEnds RMSD (red) and local InsEnds RMSD (blue) for six CASP8 targets. The global InsEnds RMSD is affected severely by the quality of the homology model. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

### Applications to protein folding simulations

Although loop modeling is often called the “mini-folding problem,” traditional approaches to loop modeling do not consider the folding mechanism when...
predicting loops. Our method on the other hand, views local modeling in a fashion that fits naturally into the larger problem of protein folding.

Experimental studies indicate that proteins fold through sequential stabilization of tertiary structure elements or foldons.34–37 Often, long-range contacts form early in the folding pathway and produce intermediate species where some entrained local regions are not yet folded. Hence, a computational scheme designed to predict structure by mimicking the natural stepwise fashion of folding pathways should encounter the problem of folding inside of loops.

Our InsEnds algorithm is well suited to address this problem because the undetermined local regions in the structure that arise during the folding pathway can correspond to distinct secondary structures, loops, or to combinations thereof. As a proof of principle, we test our method by predicting native structures of possible intermediates in the pathways for folding two proteins: ubiquitin and barnase.

The late-folding intermediate in ubiquitin lacks the $3_{10}$ helix and the $\beta_5$ strand, while the rest of the structure is well formed34,38 (Fig. 8(B)). Starting from a native-like structure for the intermediate, the InsEnds algorithm is used to fold the 18-residue insertion. The InsEnds refinement procedure successfully recovers the native structure to a global RMSD of 1.6 Å [Fig. 8(C)]. This illustrates an example where the local region is neither a loop nor a continuous secondary structure. Nevertheless, we still obtain the right topology, essentially completing the last step of the folding pathway to predict to the native structure.

Barnase is a 108-residue protein that is atypical for a small protein because it contains three distinct hydrophobic cores. The two hairpin loops depicted in Figure 8(D) are crucial to the structure because they are involved in formation of the protein’s cores, and, therefore, the correct prediction of the loops is essential for the prediction of the global structure. Experiments indicate that loop 2 is the last structure to form in the folding pathway.36 When the InsEnds method is used to fold both the 10 and 15 residue loops in barnase [Fig. 8(E,F)], our best predictions in both cases lie in the top clusters, and the best global RMSDs are 2.03 Å and 1.27 Å for loops 1 and 2, respectively.

The problem of folding inside of loops highlights two aspects of our method. The first is that our approach treats local and global structure prediction similarly by mimicking the natural protein folding mechanism. The second aspect is the demonstration that given the correct boundaries, our method is able to reconstruct the local structures irrespective of whether the local regions are well defined secondary structures or loops.

**Simultaneous folding of multiple InsEnds**

One crucial feature of our approach is the ability to simultaneously model multiple local regions. When the regions are interacting, simultaneous modeling...
can be essential because the context provided by one local region may be important in guiding the other into place. A good example is the CASP target TR606, where the InsEnds correspond to the two termini that form a hydrogen-bonded pair of β strands. The initial template model fails to identify the ends as strands, and, therefore, the ends are wrongly placed. Accurate modeling requires that they be folded simultaneously. Guided only by the orientationally dependent DOPE-PW energy function, we have modeled the free termini and correctly predicted the pair of strands in our top submission [Fig. 5(G)].

**Protein structure prediction pipeline**

One of our goals is to combine the respective strengths of free modeling with template-based modeling for an integrated structure prediction pipeline. This goal is realized through an automated server, created for CASP9 that integrates the InsEnds, RAPTOR-X and ItFix methods. Given a sequence, the pipeline begins by performing homology modeling using RAPTOR-X. If no templates are identified, the pipeline directs the sequence for free modeling using our existing ItFix algorithm for secondary and tertiary structure prediction. If RAPTOR is able to build a template-based model, the InsEnds are modeled to obtain a final structure. The pipeline has been used for the CASP9 structure predictions of the Midway-Folding group (CASP9 group numbers 435 and 477).

**Conclusions**

Loop modeling has been an on-going challenge in protein structure prediction. With the recent surge in template-based modeling, InsEnds modeling is a relatively new topic in need of novel approaches. Previous methods have focused on loops in the context of crystal structures and may not be generalizable to treat imprecise template-based models. InsEnds pose a more complicated situation where the poorly predicted local regions must be modeled without assumptions concerning the accuracy of the rest of the structure or the boundaries and secondary structure of the local regions being modeled. This work presents a novel free-modeling method for local protein structure prediction that is applicable for modeling large local regions in both exact and inexact environments, as demonstrated by results both for loops in crystal structures and for InsEnds in template-based models. We consider this result as a step towards the generalization of the local protein structure problem. The work also presents a framework in which free and template-based modeling are integrated, with the aim of closing the final gaps in protein structure prediction.

**Methods**

All backbone heavy atoms are explicitly treated, whereas the side chains are represented by single C₆ atoms. The backbone bond lengths and angles are fixed at their ideal values, and only backbone torsional angles ϕ, ψ are sampled during the simulation. Loop closure is achieved by ligating the free ends of the loops to the beginning of the subsequent chain with a harmonic constraint whose strength increases as 1/Temperature during the MCSA procedure (Fig. 3).

**Ramachandran map (Pivot) move set and sampling**

The study uses our approach for sampling single-residue (ϕ,ψ) backbone torsional angles. A distribution of ϕ,ψ angles is generated from a high-resolution library of PDB structures for each amino acid (aa), conditional on the identity of the side chain amino acids. These NN-dependent torsional angle distributions are precalculated for all 20 aas, resulting in 8000 total Rama Maps that are divided into 5° × 5° bins. During each Monte Carlo step, a selected residue’s ϕ, ψ angles are changed. Besides the identity of the NNs, the Rama Maps can also be restricted according to secondary structure of the aa and its NNs. The data presented in the paper, however, are obtained without the imposition of this restriction, thereby enabling the exploration of all regions of torsional space allowed for a given amino acid based on its neighbor’s identity. The only exception to this is the CASP8 target T0464, where five of the 24 residues are restricted to helical angles as the PSIPRED program predicts them to be helical with high confidence.

**Energy functions**

The conformational search is guided through the simulation by an energy function that is a combination of the pairwise, orientation-dependent statistical potential DOPE-PW and a harmonic ligation term for the closure of the loop:

$$E = E_{DOPE-PW} + \frac{T_k}{T} [(D-D_0)^2 + (L-L_0)^2]$$

where $T$ is the simulation temperature, $D/D_0$ are the current/initial distances between the two anchor points, and $L/L_0$ are the distances between the free end and the anchor point at the site of the cut. The ligation term becomes stronger as the simulated annealing temperature decreases. The initial temperature of the simulations is set to 100, and $T_k$ is chosen such that the contributions from the DOPE-PW and ligation energies become comparable by the end of the simulation.

The interactions in DOPE-PW are parameterized based on the observed distance distributions in the PDB, contingent on neighbors, amino acid identities, secondary structures, and side chain orientations. DOPE-PW has been demonstrated to perform
well in guiding the conformational search during prediction of the structure of small proteins. The DOPE-PW term initially dominates the total energy and provides greater freedom for the conformational search, thereby aiding in properly orienting the loop with respect to the rest of the structure.

**Scoring**

Once the set of final conformations is generated from the MCSA simulations, the best candidate in this set of conformations is chosen by using a combination of quantities computed from clustering, DOPE-PW energies, and solvent accessibility.

**Clustering**

Clustering based on the Cα RMSD provides a very effective means of identifying dominant conformations. Hierarchical clustering proceeds with a distance cutoff of 5 Å, using the minimum distance method with the Cluster module in Biopython.40 Trials with distance cutoffs of 4 Å and 6 Å do not significantly alter the results. Clustering is used only when the largest cluster contains at least 5% of the total structures. The clusters are ranked as detailed in the Results section, while the best individual structures are selected according to the sum of the DOPE-PW energy and the solvent accessible surface area (SASA).

Loop regions reside mostly on the protein surface, and thus, solvent interactions can be crucial determinants of loop structures. Hence, most successful loop scoring schemes include an approximate measure for the extent of solvation as part of the scoring function.14,15,17 While the DOPE-PW energy function accurately describes the preferred orientations of the side chains of both hydrophilic and hydrophobic residues as being directed away and toward solvent, respectively, the interactions are still assumed to be pairwise additive between Cα–Cβ bond vectors and thus do not explicitly treat the solvent accessibility. Since explicit side chains are absent during the sampling stage, the program SCWRL 4.041 is used to add side chains to enable calculating the SASA using a rapid approximation with a water radius of 1.4 Å.42 The SASAs of each residue are assigned into hydrophobic and hydrophilic components, and the structure that minimizes the hydrophobic ASA and maximizes the hydrophilic components, and the structure that minimizes the hydrophilic ASA is presumed to have the best ASA score. For this purpose, the structures are ranked by using both the hydrophobic and hydrophilic ASAs, and the combined rank is taken as the net ASA score.

**MCSA simulation procedure**

The initial torsional angles of the InsEnds are randomly chosen so that no prior information is retained regarding its conformation, while the rest of the protein structure is kept fixed. A total of 700–1000 independent MCSA trajectories are run using the energy functions described above. Each step of the MCSA trajectory involves selection of a random amino acid in the InsEnds whose torsional angle is modified according to the pregenerated NN-dependent Rama map for that amino acid. This results in a new InsEnds conformation whose energy is evaluated, and the conformation is either accepted or rejected based on the Metropolis criteria at that temperature using the energy functions described above. The temperature is updated every 500 Monte Carlo steps, using a polynomial time cooling schedule.26 The simulation protocol has been implemented in a C library, called the Protein Library, and the input/output is handled by using the PDB tools from the Biopython package.

**Parallel scripting**

The InsEnds algorithm has been implemented for high throughput structure prediction using the parallel scripting language, Swift.43 Swift enables the algorithm to be expressed in a high-level logical manner independent of any specific computing resources. Swift automatically parallelizes the independent invocations of the lower-level protein structure manipulation programs, which are written in Python and C. Swift further provides the flexibility of running on multiple, different, parallel architectures by automating job scheduling and error handling, and it logs the provenance of all data objects produced.

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