Development of a new physics-based internal coordinate mechanics force field and its application to protein loop modeling

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INTRODUCTION

All-atom force fields represent a very important tool used for theoretical studies of biomolecular systems. They are essential for many areas of computational chemistry including the prediction of protein structure and function, the study of protein–protein interactions, and the prediction of structure and binding affinities of protein–ligand complexes. Although the large size of conformational space and the complexity of the energy landscapes make protein structure prediction using all-atom force-fields prohibitive for all but smallest proteins and peptides, all-atom force fields are emerging as a major tool for the refinement of protein models generated using comparative modeling methods. Although modern comparative modeling methods are able to produce models resembling closely the native conformation when protein structures with reasonable percentage of sequence identity are available from the protein data bank (PDB),1 the accuracy of those models varies significantly between the regular secondary structure portions of a protein and the more flexible regions, such as long loops. Accurate loop modeling still remains a challenge for comparative modeling methods because loops differ in both sequence and structure even within the same protein family. On the other hand, loop regions are involved in a number of biochemical processes, such as protein recognition, ligand binding, and enzymatic activity, which makes accurate prediction of loop conformations a very important and challenging problem.

Role of all-atom force fields as a major tool in theoretical studies of biomolecules as well as importance of accurate prediction of loop conformations in proteins provided motivation for this work which was aimed at the development of a new, highly accurate, internal-coordinate all-atom force field and its evaluation in loop modeling. A significant effort has been made in the last two decades to develop accurate loop prediction methods, and a number of algorithms have been proposed. The prediction methods can be roughly divided into three groups: comparative, ab initio and a combination of both. Comparative methods rely on availability of...
a suitable template loop structure from the PDB. Ab initio methods use rigorous conformational sampling, physics-based all-atom force fields, and accurate solvation models. Although comparative loop modeling methods can be accurate when a specific class of loops is considered,\(^2,3\) the ab initio approach has been significantly more successful when applied to a variety of loops of different length.\(^4-6\) Table I presents the loop modeling results reported in the literature by various groups and obtained with ab initio or with combination modeling methods. It should be emphasized that the results shown in Table I are intended to give a general idea about state of the art in loop modeling. Direct comparison of the methods used to obtain these results is difficult because different loop sets were used by the majority of authors and the effect of crystal packing was taken into account in some of the studies. Data from Table I show that conformations of short loops (<7–8 residues) can be predicted with high accuracy.\(^6,8\) Longer (11–13 residue) loops may require consideration of the crystal contacts\(^4\) (PLOP and PLOP II), although the sophisticated hierarchical loop prediction method (HLP\(^5\)) demonstrated certain success for longer loops even without the help of crystal contact data. Such data is unlikely to be available in practically relevant applications.

In method development, loop prediction is usually understood as reconstruction of a loop conformation in its native crystal environment. A realistic refinement of protein loops in comparative models, when conformation of the rest of the protein may still contain structural defects, would require prediction of, at least, side chain conformations of the residues surrounding a given loop. This task is still too difficult for most of the existing methods, although some progress was reported recently by Sellers et al.\(^5\) who examined how loop refinement accuracy is affected by errors in surrounding side chains. Although the majority of prediction methods focus on individual loops, Danielson and Lill\(^12\) proposed a method for simultaneously predicting interacting loop regions.

Success of an ab initio loop prediction method depends on two main factors: the conformational search algorithm and the accuracy of the energy (scoring) function. Many sampling algorithms of different complexity have been proposed. Their extensive review can be found elsewhere.\(^7,11\) Sampling methods can be grouped into knowledge-based methods,\(^13-15\) ab initio strategies,\(^7,8,16-24\) and combined approaches.\(^14,25\) Conformational search algorithms include molecular dynamics simulated annealing,\(^7,22,26\) Monte Carlo simulated annealing,\(^26\) genetic algorithm,\(^27,28\) exhaustive enumeration or heuristic sampling of a discrete set of \((\phi,\psi)\) angles [or dihedral angle-based build-up methods],\(^6,17,21,23,24,29\) random tweak\(^8,30-32\) or analytical methods.\(^24,31-33\)

Energy or scoring functions that have been used for loop modeling are very diverse and include statistical\(^9,12,34,35\) and physics-based\(^4,5,7\) potentials or their combination.\(^7,10,11\) Physics-based potentials usually consist of an all-atom force field, such as OPLS,\(^4,5,11\) CHARMM,\(^7,36\) or AMBER,\(^9\) and a variety of treatments of electrostatics and solvation.\(^6,8,9,20-22,36-39\) Because of the high computational cost of loop modeling, continuum solvation models, such as the solvent accessible surface area (SA) model and the generalized born (GB) model have been the methods of choice in most of the studies. It was also shown\(^4,20\) that the accuracy of loop predictions can be increased by optimizing solvation parameters specifically for protein loops. Parameterization is carried out using the assumption that the optimal parameter set should stabilize the native loop conformation against a set of loop decoys. Thus, Das and Meirovitch\(^20\) obtained improved parameters of their GB/SA model using a “training” group of nine loops. By comparison, Zhu et al.\(^4\) optimized the parameter of an additional hydrophobic term used with a GB model.

Table I

<table>
<thead>
<tr>
<th>Loop length</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
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<tr>
<td>No crystal contacts</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MODELLER(^a)</td>
<td>0.0</td>
<td>1.0</td>
<td>1.8</td>
<td>2.0</td>
<td>2.5</td>
<td>3.5</td>
<td>3.5</td>
<td>5.5</td>
<td>6.2</td>
<td>6.6</td>
</tr>
<tr>
<td>LOOP(^i)</td>
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<td>0.85</td>
<td>0.92</td>
<td>1.23</td>
<td>1.45</td>
<td>2.68</td>
<td>2.21</td>
<td>3.52</td>
<td>3.42</td>
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<tr>
<td>RAPPER(^c)</td>
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<td>0.90</td>
<td>0.95</td>
<td>1.37</td>
<td>2.28</td>
<td>2.41</td>
<td>3.48</td>
<td>4.94</td>
<td>4.99</td>
<td>na</td>
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<tr>
<td>Rosetta(^d)</td>
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<td>na</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td>1.45</td>
<td>na</td>
<td>na</td>
<td>3.62</td>
<td>na</td>
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<tr>
<td>Loop builder(^h)</td>
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<td>na</td>
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<td>na</td>
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<td>na</td>
<td>1.31</td>
<td>1.88</td>
<td>1.93</td>
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<tr>
<td>HLP(^b)</td>
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<td>0.7</td>
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<td>na</td>
<td>0.6</td>
<td>na</td>
<td>1.2</td>
<td>na</td>
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<tr>
<td>Crystal contacts</td>
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<td></td>
<td></td>
<td></td>
<td></td>
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<td>PLOPI(^g)</td>
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<td>0.52</td>
<td>0.61</td>
<td>0.84</td>
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<td>1.22</td>
<td>1.63</td>
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</tr>
<tr>
<td>PLOP II(^h)</td>
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<td>na</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td>1.00</td>
<td>1.15</td>
<td>1.25</td>
<td>na</td>
</tr>
</tbody>
</table>

\(\text{aData taken from Figure 9 of Fiser et al.}^7\)
\(\text{bData taken from Table I of Xiang et al.}^8\)
\(\text{cData taken from Table III of de Bakker et al.}^9\)
\(\text{dData taken from Tables IV and V of Rohr et al.}^{10}\)
\(\text{eData taken from Table V of Soto et al.}^{11}\)
\(\text{fData taken from Table I of Sellers et al.}^5\)
\(\text{gData taken from Table IV of Jacobson et al.}^6\)
\(\text{hData taken from Table II of Zhu et al.}^4\)
Development of a New ICM Force-Field

Use of all-atom force fields often leads to high accuracy of loop modeling; however, the best predictions were achieved at the expense of significant computational time because of the large size of conformational space (especially for longer loops), which has to be explored, and the complexity of the energy landscapes. One way to reduce the conformational space is to use a rigid covalent geometry approximation, that is, internal coordinate (torsion angle) representation. The advantage of this representation is not only in the smaller (~10-fold) dimensionality of the sampling space and faster energy evaluation at each step but also in more efficient local gradient minimization, which has much larger radii of convergence than Cartesian space local minimization.40

The internal coordinate representation was originally introduced in the empirical conformational energy program for peptides (ECEPP) algorithm41 used for conformational energy computations of peptides and proteins. In this representation, only torsional angles of each residue are allowed to vary while all other internal coordinates, that is, bond lengths and valence angles are fixed at their standard values. The torsional angle representation combined with the ECEPP/342,43 force field has been applied successfully to a variety of problems43–45 from ab initio folding of small proteins (46-residue protein A and the 36-residue villin headpiece) to discrimination of native from non-native protein conformations. A new (ECEPP05) force field46 based on the ECEPP algorithm and designed specifically for torsional angle representation was reported recently. It was developed using high level ab initio calculations combined with global energy optimization for experimental crystal structures of organic molecules. The ECEPP05 force field combined with a surface area solvation model optimized using protein decoys was also successful47 in discriminating native-like from non-native conformations for a large set of proteins. Good results were achieved when rigid covalent geometry was applied to flexible docking of organic compounds.48,49

Internal coordinate force fields consider torsion angles as the only degrees of freedom while keeping all bond lengths and bond angles fixed at standard values. The analysis of a nonredundant set of ultrahigh-resolution protein structures carried out recently50 confirmed the earlier observation51,52 that the backbone covalent geometry should not be considered as ideal and context-independent because it varies systematically as a function of the φ and ψ backbone dihedral angles. It was demonstrated that variations occur for all bond angles adjacent to the central residue and throughout all the most populated areas of the Ramachandran plot. The largest (from 107.5 to 114.0 for a nonproline and nonglycine residue) variations within the most populated regions of the Ramachandran map occur for ∠NC-C angle suggesting that allowing flexibility of this angle should improve the force field’s ability to correctly describe the energetic balance between different conformations.

Internal coordinate based modeling is at the core of the ICM program,40,53 an integrated molecular modeling and bioinformatics program that until now has largely relied on the ECEPP/3 internal coordinate force field. The capabilities of ICM include, among others, Monte Carlo molecular mechanics for ligand docking and virtual ligand screening53,54 (VLS) and homology modeling.55,56

The goal of this work was to develop, in the framework of the ICM package, an accurate and computationally efficient internal coordinate force field, ICMFF. ICMFF builds upon the approach first developed in ECEPP/05. ECEPP/05 was parameterized using a combination of experimental data (including small molecule crystal structure) for nonbonded parameter fitting and quantum mechanics (QM) calculations for derivation of other parameters. To improve accuracy of the force field, several new features were introduced: duplicate sets of van der Waals parameters for heavy atom-hydrogen interactions; flexibility of the ∠NC-C angle (and ∠NC-CO and ϕ angles in proline) for better representation of the backbone geometry; and special functional form of φ/ψ torsional potential for better description of the φ/ψ energy surface.

The new force field was tested in loop modeling simulations. Because of their remarkable structural diversity, protein loops provide a sample of the polypeptide conformational space and energy hyper-surface. Therefore, the close correspondence of the global minima of the force field energy and the experimental native loop conformations for a large set of proteins can be an indication of force field’s accuracy and relevance for practical applications in protein modeling. Because proper treatment of solvation is a critical element to obtain accurate conformational energies, the SA solvation model was optimized to stabilize native conformations relative to alternative low-energy structures. Finally, loop simulations also represent a stringent test for conformational sampling allowing us to evaluate efficiency of the biased probability Monte Carlo53 (BPMC) conformational search method implemented in ICM.

METHODOLOGY

Form of the potential

ICMFF (as well as ECEPP/3 which we use for comparison in loop simulations) is an internal coordinate force field, that is, its intramolecular energy is a function of torsional degrees of freedom (with certain exceptions). The two force fields also use the same standard residue geometry.19 Details of the ECEPP/3 force field are given in Ref. 42.
The total energy of a molecule in ICMFF, $E_{\text{intra}}$, consists of nonbonded (van der Waals plus electrostatics), $E^{\text{nbe}}$, torsional, $E^{\text{tor}}$, and angle bending terms:

$$E = E^{\text{nbe}} + E^{\text{tor}} + E_{\text{bb}}$$  \hspace{1cm} (1)

**Nonbonded potential**

The van der Waals and electrostatic parts are calculated as a sum of the Buckingham potential (instead of the Lennard-Jones potential used in the ECEPP/3 force field$^3$) and the Coulomb contribution:

$$E^{\text{nbe}} = \frac{1}{k_{14}} \sum_{i<j>0} \left[ -A_{ij}r_{ij}^{-6} + B_{ij} \exp\left(-C_{ij}r_{ij}\right) \right] + \sum_{ij>0} \frac{332q_iq_j}{rij^6}$$  \hspace{1cm} (2)

where $r_{ij}$ is the distance between atoms $i$ and $j$ separated by at least three bonds; $A_{ij}$, $B_{ij}$, and $C_{ij}$ are van der Waals parameters; $q_i$ and $q_j$ are point charges (in e.u.) localized on atoms [an additional point charge with zero van der Waals parameters assigned is used to model the lone-pair electrons of sp$^3$ nitrogen in histidine (see Ref. 57 for details)]. The summation runs over all pairs of atoms $i < j$. $k_{14}$ and $k_{14}^{el}$ are scale factors for 1–4 van der Waals and electrostatic interactions, respectively. The dielectric constant $\varepsilon = 2$ was used. In loop simulations, distance-dependent dielectric constant $\varepsilon = 2r_{ij}$ was used to account for solvent screening of electrostatic interactions.

The following combination rules for the van der Waals parameters $A_{ij}$, $B_{ij}$, and $C_{ij}$ were applied:

$$A_{ij} = \sqrt{A_iA_j}, \quad B_{ij} = \sqrt{B_iB_j}, \quad \text{and} \quad C_{ij} = (C_i + C_j)/2. \hspace{1cm} (3)$$

The van der Waals and electrostatic interactions described by Eq. (2) are included for 1–4 or higher order atom pairs. The 1–4 interactions are treated in a special way by introducing $k_{14}$ and $k_{14}^{el}$ scaling factors. $k_{14}^{el} = 1$, $k_{14} = 2$ was chosen based on our preliminary studies of the terminally blocked alanine where nonscaled 1–4 repulsion resulted in excessively high-energy barrier at $0 \sim 180^\circ$.

Unlike the ECEPP/3 force field,$^3$ ICMFF has no explicit hydrogen-bonding term in the potential function. This interaction is represented by a combination of electrostatic and van der Waals interactions with a separate set of parameters for heavy atom-hydrogen pairs.

Several force fields (e.g., ECEPP-05) rely solely on electrostatic and van der Waals terms to reproduce hydrogen-bonding interactions. However, use of the dielectric constant greater than one (as in ICMFF) leads to a considerably smaller electrostatic contribution to the interaction energy and may result in inability to reproduce short equilibrium contact distances characteristic to hydrogen bonds. In principle, because the partitioning of the total energy into different contributions is arbitrary, to some extent, appropriate fitting of van der Waals parameters should be able to lead to the results comparable to those for $\varepsilon = 1$ by redistributing interaction energy from the electrostatic to the van der Waals term. However, combination rules [Eq. (3)], used to compute van der Waals parameters for interactions between different types of atoms, restrict the ability of the nonbonded potentials to describe adequately unusually strong nonbonded interactions, such as hydrogen bonds. It was shown$^{38}$ that reasonable accuracy can be achieved in simulations of small molecule crystals even without use of electrostatic term if combination rules are abandoned. However, this approach can be applied only if the number of atom types is small because of the limited amount of experimental data available and the strong correlation between $A$, $B$, and $C$ parameters.

To solve this problem, we used an alternative approach. A second set of $A$, $B$, and $C$ parameters was introduced for all atoms except aliphatic hydrogens and carbons (see Supporting Information Table S1). Although the first set is used for interactions between either hydrogen atoms or heavy atoms, the second set describes interactions between heavy atoms and polar or aromatic hydrogens. The combination rules [Eq. (3)] are still used for computing van der Waals parameters for a pair of atoms of different type. Although this approach still requires larger number of parameters (maximum of $2n$) compared with $n$ parameters in the traditional approach, it is much smaller than the $n^2$ parameters necessary to avoid completely the use of combination rules. The atom types included in the new force field are listed in Supporting Information Table S1.

**Torsional potential**

The torsional energy term for all dihedral angles $\theta$, except $\phi$ and $\psi$ angles in Ac-Ala-NMe and $\phi$ angle in Ac-Pro-NMe, is computed as follows:

$$E^{\text{tor}} = k_{\theta}^1[1 + \cos(\theta)] + k_{\theta}^2[1 - \cos(2\theta)] + k_{\theta}^3[1 + \cos(3\theta)]$$  \hspace{1cm} (4)

where $\theta$ is a torsional angle varying from $0^\circ$ to $180^\circ$, and $k_{\theta}^i$ are the torsional parameters.

Torsional potential for Ac-Ala-NMe consists of two terms: (a) a “hump”-shaped function designed to correct the profile of the potential within certain regions of $\phi/\psi$ map, and (b) a “stripe”-shaped function that corrects the energy for a range of $\psi$ values as being near-flat beyond that range.

The “hump” potential was constructed as a product of cosine waves truncated to zero beyond the region of interest:
where $k'$ is the amplitude parameter, $n'_0$ and $n''_0$ are the frequencies of the cosine waves which determine the dimensions of the “bell” on $\phi/\psi$ plane, $\phi'_0$ and $\psi'_0$ are the coordinates of the bell's peak, and $i = 1$ or 2 is the index of the “bell.” It should be mentioned that a similar approach based on 2D $\phi/\psi$ corrections to the torsional energy was applied to development of the CHARMM force field.\(^{59}\) Molecular dynamics simulations carried out for several proteins in their crystallographic environment\(^ {59,60}\) and studies of helical peptides\(^ {61}\) demonstrated that introduction of $\phi/\psi$ dihedral crossterms or a $\phi/\psi$ grid-based energy correction term leads to significant improvement of force-field accuracy compared with the same force field but with a traditional 1D torsional potentials (Fourier series). It was also shown\(^ {62}\) that some distortions of the MM $\phi/\psi$ energy map (compared with QM map) resulting from the use of fixed bond lengths and valence angles can be removed by crossterm corrections.

The functional form of the “stripe” potential is as follows:

$$E_{\text{stripe}}(\psi) = k_{\text{stripe}} \left[ 3/5 + \cos \left( \psi - \psi_0^{\text{stripe}} \right) \right] + 2/5 \cos \left( 2 \left( \psi - \psi_0^{\text{stripe}} \right) \right)$$

where $k_{\text{stripe}}$ is the amplitude parameter and $\psi_0^{\text{stripe}}$ is the position of the stripe’s center.

The angle bending term, $E_{\text{bb}}$, introduced to account for conformation-dependent changes in $\angle NC_\alpha C$ angle (and $\angle CNC_\alpha$ angle of proline) is computed as follows:

$$E_{\text{bb}} = \frac{k_{\text{bb}}}{2} (\theta_{ijk} - \theta_{ijk}^0)^2,$$

where $k_{\text{bb}}$ is the angle-bending force constant (in kcal/deg\(^2\)) and $\theta_{ijk}$ is a reference $\angle NC_\alpha C$ (and $\angle CNC_\alpha$ for proline) angle in degrees.

**Atomic partial charges**

Separate sets of small model molecules were used for deriving (a) van der Waals parameters from crystal data and (b) torsional parameters for the side chains of all 20 naturally occurring amino acids (both charged and neutral forms were considered for ionizable side chains). Their atomic charges, $q_i$ [Eq. (2)], were fitted to reproduce the molecular electrostatic potential, calculated with the Hartree-Fock wave function and the 6-31G* basis set using the GAMESS program.\(^ {43}\) The fitting was carried out using the Restrained Electrostatic Potential (resp) method\(^ {44}\) implemented in the AMBER 6.0 program.\(^ {44}\) Molecules from the first set were considered rigid and, therefore, only one, experimental, conformation was used for charge fitting. The electrostatic potentials of molecules from the second set were computed for all minimum-energy conformations of a given molecule characterized by different values of the dihedral angles that terminate in heavy atoms (C, N, O, and S) or polar hydrogens. The resp method was also used to obtain a single set of charges using several conformations of a given molecule (multiple-conformation fitting).

The charges for all 20 naturally occurring amino acid residues were taken from the ECEPP-05 force field.\(^ {46}\) The ECEPP-05 charges were derived following the same approach as described above for the side-chain model molecules.

**Solvation model**

In the loop simulations, the solvation free energy, $\Delta G_{\text{solv}}$ of each protein structure is estimated by using a solvent-accessible surface area model,

$$\Delta G_{\text{solv}} = \Sigma \sigma_i A_i,$$

where $A_i$ represents the solvent-accessible SAs of various atom types calculated as described in\(^ {40}\) and $\sigma_i$ is the solvation parameter for each type.

**Experimental data used for deriving van der Waals parameters**

X-ray and neutron diffraction data were used for deriving van der Waals parameters and for assessing the transferability of the resulting force field to related compounds. As our goal was to derive parameters for the atom types encountered in the naturally occurring amino acids, we focused on the following classes of compounds: aliphatic and aromatic hydrocarbons, alcohols, amines, azabenzones and imidazoles, arnides, carboxylic acids, sulfides and thiols, and unblocked amino acids. Amino acids crystallize as zwitterions and, therefore, can be used for parameterization of potentials describing interactions between charged groups of ionizable residues (Asp, Glu, Lys, and Arg).

A search of the Cambridge structural database\(^ {63}\) (CSD) carried out for structures with maximum R-factor of 7.0% and containing no ions (except zwitterions) or water molecules, yielded a large number of structures. Ideally, the discrepancy factor, $R$, should be less than 5% for the molecules used for parameter refinement; however, some structures with larger $R$ factors were used for
Table II
Results of the Crystal Calculations Carried Out Using the ICMFF Nonbonded Parameters

| Compound                  | Number of structures in training set | Number of structures in test set | \( \Delta \text{cell} (\%)^a \) | \( \Delta \Omega (\)^b | \( |E_{\text{unit}} - \Delta H_{\text{subl}}| \text{(kcal/mol)} \) |
|---------------------------|--------------------------------------|----------------------------------|-------------------------------|------------------------|-----------------------------------------|
| Hydrocarbons              | 10                                   | 19                               | 1.89                          | 2.6                    | 0.9                                      |
| Alcohols                  | 7                                    | 10                               | 1.23                          | 3.8                    | 0.5                                      |
| Aza compounds             | 10                                   | 11                               | 2.0                           | 3.5                    | 2.5                                      |
| Carboxylic acids          | 6                                    | 7                                | 2.4                           | 3.9                    | 3.8                                      |
| Amides                    | 7                                    | 7                                | 2.65                          | 3.6                    | 1.04                                     |
| Sulfur                    | 7                                    | 13                               | 2.44                          | 5.5                    | 0.2                                      |
| Unblocked amino acids     | 8                                    | 5                                | 1.89                          | 2.66                   | 10.5                                     |

\( ^a \) Average deviation of the unit cell parameters obtained after the energy minimization with ICMFF from their experimental values.

\( ^b \) Molecular rotations from the experimental positions cause by the energy minimizations.

evaluation of the parameters especially for crystalline amino acids. The following criteria were applied to choose crystal structures for our calculations:

1. The coordinates of all the atoms including hydrogen have to be provided;
2. the observed structures should have no disorder;
3. if several structures are available for a given molecule, the one with the lowest \( R \)-factor was used (except when they correspond to different polymorphs);
4. if several structures obtained at different temperatures are available, the one corresponding to the lowest temperature was used to minimize the errors due to the neglect of temperature effects;
5. only structures with one molecule in the asymmetric unit were selected.

A total of 127 crystal structures and heats of sublimation for 38 crystals were considered in this work. The numbers of crystal structures used for parameter optimization and testing for each class of molecules are shown in Table II. The complete list of structures is given in Supporting Information Tables S2–S8.

The experimental heats of sublimation for the compounds with available experimental crystal structures were retrieved from the NIST Chemistry WebBook. Whenever more than one experimental heat of sublimation was reported, more recent or higher values were selected, because experimental inaccuracies are more likely to produce lower than–true values.

**QM and MM \( \phi/\psi \) maps for terminally blocked alanine, glycine, and proline**

Three model systems, namely, terminally blocked alanine, glycine, and proline were used for deriving the \( \phi \) and \( \psi \) backbone torsional parameters. We considered Ac-Ala-NMe as a model for deriving backbone torsional parameters for all the amino acids except glycine and proline. \( \phi \) and \( \psi \) torsional parameters for glycine and proline were computed using Ac-Gly-NMe and Ac-Pro-NMe as model molecules. In contrast with the ECEPP/3 rigid geometry, the \( \phi \) angle of proline was allowed to vary and the corresponding torsional parameters were obtained by fitting \( \phi/\psi \) maps for both trans and cis conformations of \( \omega \) \( \omega \) pertains to the Ac-Pro peptide group) with “down” puckering of the pyrrolidine ring.

All quantum mechanical calculations were carried out using GAMESS software. The QM \( \phi/\psi \) maps were calculated in two steps. First, all conformations generated in two-dimensional \( \phi/\psi \) space on a \( 10^\circ \) grid were geometry-optimized at the Hartree-Fock level with the 6-31G** basis set and with the \( \phi \) and \( \psi \) angles constrained. Next, single-point energy calculations were carried out for each of the optimized geometries using the more accurate MP2 method with the 6-31G** basis set and the polarizable continuum model (PCM) implemented in GAMESS. The PCM model was used to take into account the solvation free energy for consistency with our nonbonded-energy calculations carried out with the effective dielectric constant \( \varepsilon = 2 \). Heptane (\( \varepsilon = 2 \)) was used as a solvent.

The resulting \( \phi/\psi \) energy maps were compared with the corresponding maps obtained with the ICM force field. The MM energy maps were computed using standard ICM geometries and minimizing the energy of each conformation with the main-chain torsion angles constrained at the designated values; all other (i.e., \( \omega \), side-chain, and end-group) torsions were allowed to vary.

For terminally blocked alanine, we considered the following \( \phi/\psi \) regions:

\[
\begin{align*}
-180^\circ < \phi < -40^\circ; \quad -180^\circ < \psi < 180^\circ \) \quad & \text{and} \\
40^\circ < \phi < 80^\circ; \quad -150^\circ < \psi < 60^\circ \)
\end{align*}
\]

The energies of the other regions are higher and cannot be reproduced using rigid geometry. For terminally blocked glycine, which does not exhibit large high-energy regions, the entire \( \phi/\psi \) map was generated. The \( -160^\circ < \phi < -30^\circ; \quad -180^\circ < \psi < 180^\circ \) region was considered for trans (“down”) and cis (“down”) conformations of Ac-Pro-NMe.

**Optimization of van der Waals parameters using experimental crystal data**

Van der Waals parameters of ICMFF were obtained using experimental data (crystal structures and sublimation enthalpies) for crystals of small molecules containing the same atom types as the 20 naturally occurring amino acids.
The CRYSTALG program\textsuperscript{66} was used for all crystal calculations. In CRYSTALG, the lattice energy, $E_{\text{latt}}$, of a crystal structure is considered as a function of unit cell parameters, positions and orientations (Euler angles) of the molecules in the unit cell, and torsional angles of the molecule. In this work, all molecules are considered rigid, and the lattice energy is calculated as a sum of atom–atom interactions

$$E_{\text{latt}} = \frac{1}{2} \sum_{i} \sum_{j} E_{ij}^{\text{inter}} = \frac{1}{2} \sum_{i} \sum_{j} \left( E_{ij}^{\text{bond}} + E_{ij}^{\text{electr}} \right)$$

where the summation is carried out over atoms in different molecules within a certain cutoff.

The intermolecular electrostatic energy was calculated with the Coulomb term of Eq. (2) and the Ewald summation\textsuperscript{53} without including a dipole moment correction term.\textsuperscript{54} The details for calculating lattice energies are given elsewhere.\textsuperscript{66}

The molecular geometries were taken as those in the experimental (X-ray or neutron diffraction) structures, except for X-H bond lengths which were adjusted to the average experimental (neutron diffraction) values\textsuperscript{67} (1.083 Å for C–H; 1.009 Å for N–H; 0.983 Å for O–H; 1.338 Å for S–H) because of the uncertainties in the X-ray determination of hydrogen positions.

To evaluate structural changes caused by a given set of nonbonded parameters, lattice energy of each experimental crystal structure was locally minimized. The experimental space group symmetry was used to generate coordinates of all the atoms in the unit cell. However, no symmetry constraints were used during the minimization; that is, the positions and orientations of all the molecules in the unit cell were allowed to vary for each molecule independently.

An optimized set of van der Waals parameters was obtained by minimizing the following target function:

$$F = \frac{1}{N} \sum_{i=1}^{N} F_{\text{cell}}^{\text{pred}} + \frac{1}{N_{\text{subl}}} \sum_{j=1}^{N_{\text{subl}}} E_{\text{ene}}^{\text{pred}}$$

Two terms in Eq. (10) reflect deviations between (1) the computed and experimental crystal structures and (2) the lattice energies and sublimation enthalpies, respectively, caused by relaxation under the action of current force field parameters. $N$ is the number of crystal structures used in parameter optimization, and $N_{\text{subl}}$ is the number of crystal structures with known sublimation enthalpies.

For a given crystal structure $i$, the structural part of the target function was computed as follows:

$$F_{\text{cell}}^{\text{pred}} = w_{a,b,c} \left[ \left( \frac{\Delta a}{a} \right)^2 + \left( \frac{\Delta b}{b} \right)^2 + \left( \frac{\Delta c}{c} \right)^2 \right] + w_{a,b,c} \cdot (\Delta \alpha)^2$$

$$+ (\Delta \beta)^2 + (\Delta \gamma)^2 + w_x (\Delta x)^2 + w_b (\Delta \theta)^2$$

where $\Delta a$, $\Delta b$, $\Delta c$, $\Delta \alpha$, $\Delta \beta$, and $\Delta \gamma$ are changes in the unit cell parameters (in Å and deg.), $\Delta x$ is rigid-body translational displacement, and $\Delta \Omega$ is rigid-body rotational displacement (in deg.). $w_s$ are empirical weights that were taken from Ref. 58 and are introduced to ensure that all observed quantities had approximately the same relative deviations ($w_{a,b,c} = 100$; $w_{a,b,c} = 0.5$; $w_x = 10$; $w_b = 0.5$). The energy discrepancy

$$E_{\text{ene}}^{\text{pred}} = \frac{1}{N_{\text{subl}}} \sum_{j=1}^{N_{\text{subl}}} |E_j - \Delta H_{\text{subl}}|$$

is calculated for crystal structures with known experimental enthalpies of sublimation.

Besides minimizing the target function, an optimal parameter set was required to satisfy two other conditions. Potential energy curves for all the parameters and their combinations [computed using the combination rules from Eq. (3)] must have a minimum between 0 and 5 Å and van der Waals radius for each element must be within ±0.2 Å from the tabulated value (we use van der Waals radii\textsuperscript{58} of 1.7 for C; 1.2 for H; 1.55 for N; 1.52 for O; 1.8 for S). Because van der Waals radii of polar and aromatic hydrogens may depend on their chemical environment, they were allowed to assume any value.

We used an iterative parameter optimization procedure. At each iteration, one van der Waals parameter [A, B, or C from Eq. (2)] is selected at random from the set of parameters allowed to vary and a short search in the vicinity of its current value is carried out. This search consisted of a small number of steps (usually 6) each of which included: (1) perturbation of the selected parameter by adding a random number within ±10% of the current parameter value, (2) computation of van der Waals radii and positions of the minima for all combinations of the selected parameter with other parameters, and (3) local energy minimizations of all training crystal structures with the resulting set of van der Waals parameters and calculation of the target function [Eq. (9)]. The parameter value which gives the lowest value of the target function is accepted, and the updated set of van der Waals parameters is used as a starting point for the next iteration. The optimization procedure was terminated when changes in the target function did not exceed an empirical threshold of 0.5. Several optimization runs were carried out starting from the different initial sets of parameters generated by random perturbation of the ECEPP-05 van der Waals parameter set. Each optimization run consisted of ~100 iterations.

Once an optimized set of parameters was obtained, its accuracy and transferability was evaluated by lattice energy minimizations carried out for a test set of crystal structures for a given class of compounds.

The following three measures were used to assess performance of the optimized parameters:
An average percent deviation of the unit cell parameters from their experimental values ($\Delta_{\text{cell}}$) calculated using the formula

$$\Delta_{\text{cell}} = 100\% \cdot \frac{\sum_{i=1}^{N} |x_{i} - x_{i,\text{exp}}|}{x_{\text{exp}}} (1/N)$$  (13)

where $x_{i,\text{exp}}$ and $x_{i,\text{m,exp}}$ are the unit cell parameters of the experimental and the minimized experimental structures, respectively; $N$ is the number of unit cell parameters;

2. Rotational angle, $\Omega$, characterizing similarities of molecular orientations in the experimental and minimized experimental structures and computed as described in Ref. 57;

3. Deviation of lattice energy from the experimental sublimation enthalpy.

An accurate parameter set should provide structural deviations below 5%. Taking into account that there are many uncertainties in comparing lattice energies and corresponding sublimation enthalpies and that an average experimental error is about 2 kcal/mol or more, a deviation of a few kcal/mol between the lattice energy computed with a given parameter set and the corresponding sublimation enthalpy was considered to be acceptable.

**Derivation of parameters of the side-chain torsion potentials**

Our derivation of the torsional potential-energy terms relied on fitting the molecular-mechanical (MM) energy profiles for rotation around a specific bond against the corresponding QM profiles. The corresponding torsional potential energy terms were obtained by fitting a cosine series [Eq. (4)] to the difference between the QM and MM profiles (the latter consisting of nonbonded and electrostatic terms).

A set of model molecules containing the same types of torsional angles as those present in the naturally occurring amino acids and in the protein backbone was used. The four atoms (defining each type of torsional angle) with their covalently bound neighbors replaced by hydrogen atoms defined the molecules selected for the calculations. Thus, the torsional terms were parameterized to reproduce the properties of the simplest molecules possible and then applied to larger and more complex ones.

The QM and MM profiles of the model molecules were computed adiabatically, that is, by constraining the appropriate torsions for each of the torsional angles on a 10° grid and minimizing the energy with respect to all the other degrees of freedom. All the ab initio calculations were carried out at the MP2 level of theory with a 6-31G** basis set implemented in the GAMESS program. The corresponding MM torsional profiles were computed using the ICM program. The molecular geometries (bond lengths and bond angles) were optimized by QM calculations, and the lowest energy QM conformations were used for calculating the MM torsional profiles. Some functional groups such as methyl, phenyl, and amino groups can have higher symmetry than the geometries obtained from QM calculations on fixed rotamers of these groups; hence, the corresponding bond lengths and bond angles of these groups were averaged to conform to the highest symmetry possible for a particular group.

**Refinement of the backbone torsional parameters using QM $\phi/\psi$ maps**

Two different approaches were employed to obtain the $\phi/\psi$ backbone torsional parameters for the blocked Ala and Gly.

For parameterization of the backbone torsional potential for blocked glycine, the MM parameters were derived by minimizing the following target function:

$$F(k_0; A; B; C) = \sum_{i=1}^{N} w_i (\Delta E_i^{\text{MM}} - \Delta E_i^{\text{QM}})^2$$  (14)

with respect to the $k_0$ coefficients of the Fourier expansion [Eq. (4)]. The summation runs over all $N$ points of the $\phi/\psi$ map taken into consideration. $\Delta E_i^{\text{MM}}$ and $\Delta E_i^{\text{QM}}$ are the relative MM and QM energies, respectively, for a given point $i$; $w_i$ are empirical weights. The weights were computed according to the formula

$$w_i = \exp(-c \cdot ||(\Delta E_i^{\text{QM}} - \Delta E_i^{\text{MM}}) - c_i (\Delta E_i^{\text{QM}} - \Delta E_i^{\text{MM}})||),$$  (15)

where $c$ and $c_i$ are empirical parameters introduced to provide additional de-emphasis of high-energy regions. The value of $c$ was chosen so as to give higher weights to those of the fitting points located at or near the energy minima.

Because this fitting method did not produce acceptable results in our preliminary studies of Ace-Ala-NMe, an alternative empirical approach described in detail in the Results and Discussion Section was used. It was designed to reproduce main features of the QM $\phi/\psi$ map (such as shape and relative stability of the low-energy regions), while focusing on the low-energy regions of the QM energy surface that are also the most populated areas of the Ramachandran map obtained from the analysis of the experimental protein structures.

**Parameterization of the angle-bending term**

The force constant and reference angle of the angle-bending term [Eq. (7)] were obtained by minimizing RMSD between the QM and MM values of the $\angle NC\alpha C$ angle ($\theta$) for a set of conformations, that is,
\[
\text{RMSD(} \angle \text{N}^\alpha \text{C}) = \sqrt{\frac{\sum_{i=1}^{N}(\theta_i^{\text{QM}} - \theta_i^{\text{MM}})^2}{N}},
\]

where \(\theta_i^{\text{QM}}\) are the QM values of \(\angle \text{N}^\alpha \text{C}\) taken from the conformations of the Ac-Ala-NMe, Ac-Gly-NMe, and Ac-Pro-NMe generated to compute QM \(\phi/\psi\) energy maps. These conformations represent all of the most important regions of the Ramachandran map and, therefore, are suitable for parameterization of the term associated with the \(\angle \text{N}^\alpha \text{C}\) bending which was shown to be strongly influenced by changes in \(\phi/\psi\) backbone torsional angles. \(N\) is a number of structures (see QM and MM \(\phi/\psi\) maps for terminally-blocked alanine, glycine, and proline section). \(\theta_i^{\text{MM}}\) angles were calculated by minimizing MM energy of a given blocked amino acid (Ala, Gly, or Pro) as keeping the \(\phi\) and \(\psi\) angles fixed at the same values as those in the corresponding QM conformations.

As was reported by Karplus, \(\text{Q}^2\) QM results show larger deviations of bond angles ranging farther both positively and negatively than in the experimental protein structures; however, they follow very similar trends. In the case of an internal-coordinate force field, use of a “softer” \(\angle \text{N}^\alpha \text{C}\) angle bending potential parameterized using QM data may have an advantage because it may compensate partially for the rigidity of all other bond angles.

Parameter optimization was carried out via a systematic search on the \(k_0/\theta_0\) grid. Grid points were obtained by scanning the 200–500 kcal/deg\(^2\) range for \(k_0\) and 104–115° range for \(\theta_0\) with step of 20 kcal/deg\(^2\) and 1° for the force constant and the reference angle, respectively. The same range of \(k_0\) values and \(\theta_0\) from 110 to 120° was considered for parameterization of \(\angle \text{C}^\alpha \text{N}^\alpha \angle\) bending term in proline. The final \(\angle \text{N}^\alpha \text{C}\) angle bending parameters, \(k_0\) (in kcal/deg\(^2\)) and \(\theta_0\), are as follows: 405.0 and 108° for all amino acids except Gly and Pro; 440.0 and 110° for Gly; 330.0 and 110° for Pro. \(k_0\) is 250.0 kcal/deg\(^2\) and \(\theta_0\) is 116° for \(\angle \text{C}-\text{N-Co} \angle\) term in Pro.

**Loop datasets**

Loops with lengths from 4 to 13 residues were considered in this work (Supporting Information Tables S9–S18). To facilitate comparison with the work of other authors, we chose the loop databases used previously by Jacobson \(\text{et al.}\) (the filtered sets for 4–12 residue loops) and Zhu \(\text{et al.}\) (12 and 13 residue loops). All the loops were taken from high resolution (2 Å or better) and diverse (<20–60% sequence identity) protein crystal structures. Complete lists of the selection criteria used for compiling these databases are given in Refs. 4 and 6–8. We excluded the loops containing cysteine residues involved in disulfide bonds with the rest of the protein because the additional covalent bond constraint radically reduces the conformational space of the loop making it nonrepresentative of the particular loop length. Because neither of the loop databases selected for this work contains structures with cis-prolines in the loop regions, we added two loops with cis-proline [from 1w0n (six-residue loop) and 2ixt (nine-residue loop); see Supporting Information Tables S11 and S14] to test accuracy of our proline torsional potential.

We did not consider crystal packing in the simulations presented in this work.

The standard protonation state at pH 7.0 was assigned to all titratable groups (histidine and tyrosine were considered as uncharged). Only the δ tautomer of histidine was used.

Accuracy of the loop modeling results was assessed using the average and median backbone root-mean-square deviation (RMSD) computed after superimposing the body (i.e., all the residues except those from the loop region) of the protein.

To evaluate efficiency of the sampling algorithm, energy of the lowest energy conformation was compared with that of the optimized native structure. Thus, the positive energy gap between the lowest energy and the optimized native conformations was considered as an indication of possible sampling errors, that is, an inability of the procedure to locate the global minimum. On the other hand, non-native predicted conformation with the energy lower than that of the optimized native conformation may indicate force field errors. We did not attempt using energy of the native conformation as a reference for evaluation of the force field. Taking into account the roughness of the energy surfaces of large all-atom systems characterized by significant energy variations corresponding to small changes in structural parameters, the lower energy of a predicted near-native conformation compared with the native one does not necessarily mean inadequate accuracy of the force field.

When comparing energies of the native and predicted structures, it is important to eliminate the noise originating from the minor differences in covalent geometry and sub-optimal van der Waals contacts that are due to the different force fields used in X-ray structure refinement. Therefore, the native structure was optimized by conversion to the standard ICM covalent geometry (which included rebuilding of all hydrogens), by carrying out a systematic search for torsional angles defining positions of polar hydrogens, and by local energy minimization as a function of loop degrees of freedom.

**Optimization of solvation parameters for loop simulations**

To derive optimal solvation parameters, we split the SA-based solvation term into four components according to the following classes of atoms: aliphatic carbons, aromatic carbons, charged (ionized) atoms (side-chain oxygens in glutamate and aspartate and nitrogens in arginine and lysine), and other polar atoms. Four weights for the
contributions of each component were introduced into the final free energy calculation.

Conformational ensembles for 58 loops of nine residues (Supporting Information Table S14) taken from the benchmark were used as a training set to optimize the solvation energy function. The ensembles were generated using the loop simulation protocol (described below) with the initial solvation parameter set.73 The ensembles contained 960–1656 conformations per loop, for a total of 80,855 conformations. The conformational ensembles are characterized by RMSDs distributed across the ~0.2 to 12.0 Å range.

The goal of the optimization was to find the parameters resulting in an energy function that gives best ranks to near-native loop conformations. Therefore, to quantitatively evaluate the performance of the energy function, we constructed a score that combines RMSDs and rankings in such a way that it rewarded placing low-RMSD conformations at low ranks:

\[
S = \sum_{j \in \text{Loops}} \left( \sum_{i \in \text{Confs}} e^{-\text{RMSD}_i^j}/i \right)
\]  

(17)

Initial values of solvation energy components, the remaining force-field energy terms and RMSDs were pre-calculated for all conformations in the training set, so that total free energies could be re-evaluated rapidly for any combination of the weights of the solvation components without resorting to calculations on actual 3D structures. The simplex method74 was used to minimize score \( S \) as a function of weights.

**Loop modeling protocol**

We used the BPMC conformational search procedure53 as implemented in ICM40 for the sampling and global optimization of the loop conformations. To facilitate sampling of the loop backbone conformational space, a two-stage conformational search protocol was devised. During the first stage, only the loop was explicitly present in molecular mechanics, whereas the rest of the protein was represented by a simple steric exclusion potential precalculated on a grid. The steric exclusion potential was calculated as standard van der Waals energy for an aliphatic carbon atom probe placed at each node of the grid. The calculated energy values were trimmed to the [0.0, 4.0] kcal/mol range, resulting in a potential with no attraction and with the maximum penalty of 4 kcal/mol per atom for entering sterically excluded regions. To make the protocol applicable in simulations with flexible side chains on the static part of the protein, this steric exclusion potential was generated without the sidechains. Furthermore, the loop was reduced to a simplified Glycine-Alanine-Proline (GAP) model by substituting all other amino acid residues by alanine. BPMC sampling for this model allowed us to generate rapidly an ensemble of low-energy backbone conformations of the loop that were free of gross clashes with the rest of the protein backbone. Temperature parameter for the Metropolis criterion in MC was set to 1000 K. Up to 300 steps of local gradient minimization were allowed after each random step.

During the second stage of our loop simulation protocol, a full-atom model of the protein was rebuilt. Static parts of the protein were then explicitly present, and the original sequence of amino-acid residues in the loop was restored based on GAP backbone conformation. Short gradient energy minimizations for \( \chi \) angles of the loop side-chains were performed to resolve clashes (where possible without backbone movement). The conformations in the resulting ensemble were used as starting points for a new series of BPMC runs: the first simulation was started from the lowest energy conformation in the ensemble. Whenever no further progress was detected in the current trajectory, a different conformation was chosen from the ensemble to start a new trajectory. Lack of progress was determined using a visit count mechanism.54

A simple empiric rule was used to determine the total length of the BPMC simulation: the simulation was terminated after 8000 \( \times 2^L \) energy evaluations, where \( L \) is the loop length. We evaluated convergence to the global minimum by performing five independent runs of the full protocol in parallel.

Covalent attachment of the loop to the N- and C-terminal parts of the polypeptide chain was maintained by adding virtual “shadow” C and C atoms at the junction of the static and flexible segments of the polypeptide chain (Fig. 1). The “shadow” and real atoms were tethered to each other with harmonic constraints \( E_{\text{constr}} = k_{\text{constr}} (r_{\text{shadow}} - r_{\text{real}})^2 \). The force constant, \( k_{\text{constr}} \), was

![Figure 1](image_url)  

**Figure 1**  
Junction of the flexible and static segments of the polypeptide chain in loop simulations. The arrows show virtual bonds that are parts of the internal coordinate trees of the two segments. Virtual C and C atoms at the termini of the two segments are constrained to their physical counterparts. These constraints, in conjunction with rigid covalent geometry within the two segments, maintain (near) ideal geometry of the physical C—C bond.
**RESULTS AND DISCUSSION**

**Optimization and evaluation of van der Waals parameters**

Optimized values of the van der Waals parameters for the atom types present in 20 naturally occurring amino acids are given in Supporting Information Table S1. The average deviations of unit cell parameters, molecular rotations, and lattice energies from the corresponding experimental values computed for each class of compounds are shown in Table II, and the complete list of deviations for all the molecules considered in this work can be found in Supporting Information Tables S2–S8.

Results reported in Table II show that energy minimizations carried out with the optimized van der Waals parameters result only in minor changes of the unit cell as compared with the experimental crystal structures for all types of compounds. Thus, for the majority of molecules, the original space group symmetry of the experimental structure was preserved after local-energy minimization. Average deviations of unit cell parameters did not exceed 3% (Table II and Supporting Information Table S2–S8). Changes in molecular orientations were also small (<4°) for sulfur-containing compounds (5.5°, Supporting Information Table S7). Computed lattice energies are in agreement with the experimental sublimation enthalpies within the expected differences for hydrocarbons, alcohols, amines and imidazoles, amides, and sulfur-containing compounds, whereas lattice energies are somewhat lower than $\Delta H_{\text{subl}}$ for carboxylic acids (Supporting Information Table S4). Only one heat of sublimation (that of glycine) is available for crystalline amino acids, and it is somewhat underestimated by the new parameterization (see discussion below).

In general, the agreement between the structures energy-minimized using the new force field and the experimental data is very good. The average deviation of the unit cell parameters from their experimental values for hydrocarbons was found to be less than 1.9%. Excellent agreement between the energy-minimized and experimental structures (with the average deviation of 1.2%) was also obtained for alcohols (Supporting Information Table S3). The larger than average deviations observed for some crystals [e.g., for propane and pentane (Supporting Information Table S2)] with the deviations of the unit cell parameters of $-5.7\%$ and $-6.1\%$, respectively, and for molecular rotation in methanol crystal (CSD ID methol02, Supporting Information Table S3)] can be explained by physical effects such as thermal motion which has stronger effect on crystal of very small molecules and on those with relatively weak intermolecular interactions.

All the experimental structures of azabenzenes and amines, except purine (CSD ID PURINE) and pyrazine (CSD ID PIRAZI01), were reproduced very well (Supporting Information Table S5). On average, the deviations of the unit cell parameters from the experimental values did not exceed 2%. The deviations were larger for PURINE (maximum of 7.6%) and PIRAZI01 (maximum of 6.5%). The nonbonded parameters derived in this work performed well for carboxylic acids and amides (Supporting Information Tables S4 and S6). Thus, the average deviation of the unit cell parameters did not exceed 2.5% and 2.7% for carboxylic acids and amides, respectively. The largest deviations of unit cell parameters were obtained for formic (6.6%) and acetic (7.8%) acids and formamide (FORMAM02, 8.7%). A comparison with the results reported for carboxylic acids by other groups shows that all the force fields (AMBER, Dreiding, OPLS, ECEPP/05) that use partial charges located on atomic sites to describe electrostatic interactions give similar results, that is, energy minimization always led to significant changes in the unit cell parameters. On the other hand, it was shown that the potentials, which make use of the distributed multipole analysis (DMA) to describe electrostatic interactions, performed significantly better in maintaining the experimentally correct structure. This suggests that an accurate description of the anisotropy of the electrostatic interactions is very important for modeling the crystal structure of carboxylic acids and amides.

Nonbonded parameters derived for sulfur-containing compounds led to the average unit cell and rotational deviations of 2.4% and 5.5° (Supporting Information Table S7), respectively, and to the excellent agreement with the experimental sublimation enthalpy (that of S8). In general, agreement with the experimental crystal struc-
tures is somewhat worse than for other compounds. Abraha and Williams,81 who considered a set of crystal structures of elemental sulfur (S₄), presented evidence that the bonded sulfur atom in these structures is not spherical and that an aspherical van der Waals model was necessary to achieve acceptable agreement with the experimental data. Implementation of such an aspherical model may also increase accuracy of the potential for other sulfur-containing molecules albeit at a higher computational cost. Because the number of sulfur atoms in an average protein structure is usually low, accuracy of the new parameterization was considered acceptable for use in protein modeling.

A group of crystalline amino acids (Supporting Information Table S8) was used for deriving parameters for nitrogen and hydrogen in the –NH₃+ functional group. Both the experimental unit cell parameters and molecular orientations were reproduced very well (Table II) with no structures with deviations of unit cell parameters exceeding 5%. The sublimation enthalpy of γ-glycine was used as a reference for lattice energy. The sublimation enthalpy was reproduced with accuracy lower than that of other types of compounds (Table II) and a comparison with the results reported for glycine in other studies82 suggests that a more sophisticated electrostatic model may be required to get better agreement between the lattice energy and sublimation enthalpy of γ-glycine. Because unblocked amino acids considered in this work contain a variety of functional groups, results obtained for this group of compounds also serve as an indication of the overall high accuracy of the nonbonded part of the new force field.

Optimization of torsional parameters

Side chain torsional potentials

Supporting Information Tables S19–S23 contain the list of small molecules used for parameterization of the torsional energy terms for amino acid side chains. Ab initio and ICMFF energies of different conformations are in excellent agreement, that is, the average difference between them is less than 0.1 kcal/mol. There are only a few deviations greater than 0.3 kcal/mol with the majority of them taking place for sulfur-containing compounds (Supporting Information Table S20). The two largest deviations between the QM and MM energies were obtained for the cis conformation of H₂CSSCH₃ (0.98 kcal/mol) and for the cis conformation of H₃CH₂CH₂SCH₃ (0.89 kcal/mol, rotation ca. CH₂—CH₂ bond). ICMFF overestimates the energy of these strained conformations because of the rigid valence geometry employed and the larger size of sulfur atom. It should be mentioned that the current version of the ICM program utilizes a set of harmonic distance constraints rather then bonded parameters to describe the disulfide bridges.

As indicated by the results in Supporting Information Tables S19–S23, accuracy of the model is high enough to reproduce well the fine details of the QM results.

Torsional potential of χ₁ of amino acids is modeled by the third term of Fourier expansion [Eq. (5)] with the parameters reported in Supporting Information Table S24. An exception was made for the threonine side chain. We observed significant discrepancy between the χ₁, MM energy profile and the statistics of χ₁ angle distribution in X-ray protein structures.83 In particular, while trans- and meta-conformers are almost equally populated in X-ray structures, the MM energy difference was >1 kcal/mol when only the third term of the Fourier expansion was used to compute χ₁ torsional energy (φ and ψ angles were set to −160° and 140°, respectively). Therefore, the first term of the Fourier expansion (with an empirically adjusted parameter) was introduced to compensate for the difference (see Supporting Information Table S24). Similar analysis carried out for all other amino acids showed that the use of a single third term of Fourier expansion [Eq. (5)] leads to reasonable agreement with the distribution for χ₁ rotamers in PDB. The resulting parameters for the torsional potential for all types of torsional angles are given in Supporting Information Table S24.

Parameterization of the backbone torsional potential

Ac-Ala-NMe. Accuracy of the force-field energy function with respect to backbone φ/ψ angles is of extraordinary importance in protein modeling because these angles determine secondary structure and because the relatively small angular deviations result in large movements as they propagate along the polypeptide chain. Therefore, we paid special attention to the parameters and choice of the functional form for φ/ψ torsion potentials. To gain initial understanding of the behavior of ICMFF force-field energy function, we calculated φ/ψ energy map [Fig. 2(a)] for the nonbonded terms (van der Waals and electrostatics) on the model molecule, Ac-Ala-NMe, and compared it with the QM energy map [Fig. 2(b)]. We also visualized the difference between the MM and QM energies as a heatmap on the φ/ψ plane, emphasizing areas populated in protein structures [Fig. 2(c)].

Two regions on the φ/ψ energy map appeared to be significantly lower for the MM map as compared with the QM one: (1) the region below φ'<1 and α<1, centered at approximately (φ = −120°, ψ = −60°); and (2) the region corresponding to β, centered at approximately (φ = −150°, ψ = 70°). Another deviating region was the horizontal stripe region following ψ ~ 0°, where the MM energy is significantly higher. This difference is apparently due to some clashing of N° and N°+1 backbone nitrogens. A comparison to the Ramachandran plot [Fig. 2(d)] suggests that these three discrepancies are true artifacts of the MM energy indeed, the two areas where QM energy is significantly higher show low population
in the Ramachandran plot, and, conversely, the $\psi \sim 0^\circ$ region is well populated at least around $\phi \sim -90^\circ$.

To attenuate or eliminate these artifacts, we considered various functional forms of $\phi/\psi$ torsion potentials. Despite extensive efforts using various weighting schemes for fitting of traditional cosine potentials, including multiple harmonics (1–6) and sine/cosine combinations, we could not achieve satisfactory fit across the entire region of interest. A good fit for the characteristic minima points would invariably lead to “ripples” elsewhere, resulting in an unsatisfactory overall profile. We eventually converged on a special functional form that introduces “a hump” around specific $\phi/\psi$ value pair (see Form of the potential section). Use of the cosine-based function [Eqs. (5) and (6)] allowed us to evaluate the term quickly without additional trigonometric function calls because cosines of all torsions are calculated and stored anyway during geometry construction in ICM. We used two such “humps,” the first one of 4.0 kcal/mol at ($\phi = 120^\circ$, $\psi = -60^\circ$) and second one of 2.0 kcal/mol at ($\phi = -150^\circ$, $\psi = 70^\circ$), to compensate for the low-energy artifacts. The amplitudes of the two “humps” were chosen to correct the $E_{\phi' - C_\phi}$ and $E_{\beta - C_\beta}$ energy differences, respectively, to within 0.1 kcal/mol accuracy. $C_\phi$ rather than $C_7\text{eq}$ minimum was considered as a reference state because it is much more populated in protein structures and also because there are multiple indications that deep $C_7\text{eq}$ minimum might be an artifact of the QM calculations, associated with intramolecular hydrogen bonding. Indeed, a basis set superposition error (BSSE) results in the overestimation of the stabilization energy of hydrogen bonds by ab initio molecular orbital calculations. BSSE can be corrected using counterpoise calculations, but for intramolecular hydrogen bonds this approach is not directly applicable. Another smaller effect is associated with stronger vibrational force constants of the more rigid, ring-like conformations held tight by the internal hydrogen bonds. The estimated contributions from these two effects are as high as 1.5–4.4 kcal/mol for the BSSE and $\sim 0.3$ kcal/mol for the vibrational effects.

Figure 2
(a) Nonbonded plus electrostatics $\phi/\psi$ energy map for Ace-Ala-NMe; (b) QM $\phi/\psi$ energy map for Ace-Ala-NMe. The color code from purple to red corresponds to the 0–8 kcal/mol range; (c) heatmap of the deviations between the QM and MM (computed without torsion potential) energies for Ace-Ala-NMe. The size of the squares indicates the frequency (on a logarithmic scale) of occurrence of a particular $\phi/\psi$ value pair (within a $10^\circ \times 10^\circ$ bin); (d) Ramachandran plot for a set of 21 diverse ultrahigh resolution structures (resolution between 0.5 and 0.8Å, PDBs 1ejg, 1ucs, 2v61, 1us0, 2dsx, 1r6j, 2h97, 1x6e, 1gci, 1p7f, 1ius, 2ist, 1wbn, 2hsc, 1mrz, 1n55, 2o9a, 2jr6, 2pwa, 2o7a, and 2hs1), blue points. Background is colored by $\phi/\psi$ frequencies for a much broader protein set, calculated within each $10^\circ \times 10^\circ$ on the $\phi/\psi$ plane. (e) Contour map of the $\phi/\psi$ torsion potential. Color code from purple to red corresponds to the 0–4 kcal/mol range; (f) final energy map including the torsion potential. The color code from purple to red corresponds to the 0–8 kcal/mol range; (g) heatmap of the residual deviations between the QM and final MM energies for Ace-Ala-NMe. Size of the squares indicates the frequency (on a logarithmic scale) of occurrence of a particular $\phi/\psi$ value pair (within a $10^\circ \times 10^\circ$ bin). Contours in (a, b, e, and f) are drawn with 1 kcal/mol step.
A \( \psi \)-only two-cosine wave was used to compensate the high-energy artifact in the \( \psi \sim 0^\circ \) region. However, we found that full compensation of the MM/QM difference in \( E_{\text{MM}} - E_{\text{QM}} \) led to an unsatisfactory geometry and excessive stability of the alpha-helical conformation in our tests for a 20-residue polyalanine peptide. We, therefore, reduced the amplitude of this component of the potential until average \( \phi/\psi \) angle pairs in the energy minimized helix returned closer to typically observed values of \( \phi = 60^\circ \) and \( \psi = 45^\circ \). At the final amplitude of 1.4 kcal/mol, the minimum was at \((\phi = -66.2^\circ, \psi = -40.9^\circ)\). Importantly, when distance-dependent dielectric method rather than Colomb electrostatics is used, the average \( \phi/\psi \) angles in the minimum were at \((\phi = -65.6^\circ, \psi = -41.6^\circ)\). It should be noted that this component of the torsion potential function affects directly the energy difference between alpha-helical and beta-strand conformations in a polypeptide. We are currently testing the force field in peptide folding simulations to fine-tune it in conjunction with other terms including the solvation model.

The contour map of the full \( \phi/\psi \) torsion potential [Fig. 2(e)] illustrates the corrections introduced into the MM energy: oval and round red/yellow areas are the two “humps,” whereas the blue/purple band is the \( \psi \)-only component. A contour map of the final total MM energy as well as the MM-QM difference heatmap are shown on Figure 2(f,g), and the main minima of Ac-Ala-NMe are listed in Table III. Good correspondence of the location and shape of the minima can be observed, in particular, in highly populated areas.

**Ac-Pro-NMe.** In contrast with other internal coordinate force fields, we also allowed flexibility of the \( \phi \) angle in proline. Although this torsion is commonly kept fixed in torsion space modeling because it is constrained by the proline’s pyrrolidine ring, deviations of up to 15–20\(^\circ\) from the median value of \(-75^\circ\) are frequently observed in X-ray protein structures (Fig. 4). In the present version of the ICM force field we did not introduce any treatment of the internal flexibility of the proline ring, keeping internal variables of the side chain rigid. Although this approach does result in somewhat distorted ring geometry when \( \phi \) deviates far from the mean value, only minor distortions occur for \( \phi \) angle values within the range of interest in protein simulations. Indeed, we compared the conformation of the ring atoms together with two adjacent backbone C atoms (of the proline itself and the preceding residue) for an idealized Ac-Pro-NMe conformation with \( \phi \) angle fixed at \(-68.8^\circ \pm 20^\circ\) before and after QM geometry optimization (HF6/31G**) and found that RMSD did not exceed 0.12 \( \text{Å} \). We judged this an acceptable trade-off as compared with the optimized backbone torsional parameters. All the main features of the entire map are reproduced accurately. Notable differences include the shape and higher energy of the \(-30^\circ < \phi < 30^\circ\) region of the MM map, which is a result of the rigid internal geometry used. \( \phi \) and \( \psi \) torsional parameters for glycine are given in Supporting Information Table S24.
much larger inaccuracies associated with completely rigid pyrrolidine ring. We did not introduce any explicit torsional potential for the proline $\phi/\psi$ pair because the overall shape of the low-energy region was reproduced well by the non-bonded terms of the force field (Fig. 5). While the locations of the minima deviated somewhat from those observed in QM energy map [Fig. 5(d,e)], we judged that they may be influenced by the artifacts of QM calculations (see Ac-Ala-NMe section). Indeed, the reported experimental energy difference$^{86}$ between the cis- and trans-conformations of N-methylacetamide is 2.5 kcal/mol, which is much closer to our MM value of 2.7 kcal/mol than to the QM one of 3.89 kcal/mol (Table IV). Furthermore, the MM energy minima seem to correspond better with the Ramachandran plot for proline residue [Fig. 5(c)].

**Optimization of solvation parameters for loop modeling**

The SA solvation model was optimized using a training set of conformations generated for 58 loops of 9 residues as described in the Methodology Section. The goal of the optimization was to obtain the parameter set that would result in the most favorable ranking of the near-native solutions among a large number of decoy structures generated by the BPMC sampling. The rationale for this optimization is that the original parameters of the SA solvation model were derived by fitting to the vacuum/water transfer energies for small molecule analogues to amino-acid side-chains, which may not adequately capture some of the effects of the burial of a particular chemical moiety in the bulk protein interior.

The optimized solvation parameters are given in Supporting Information Table S25. A comparison with the initial parameter set (column 3, Supporting Information Table S25) shows that the optimization led to changes in all the parameters, although the most significant changes took place for aromatic carbon and oxygen and nitrogen atoms from ionized groups. Interestingly, the solvation parameter of aromatic carbon changed its sign, so the atom became hydrophobic rather than weakly hydrophilic as was the case in the original solvation model. The new sign of this parameter is in agreement with an empirical observation that aromatic side chains have a

**Table IV**

Backbone Torsional Angles and Intramolecular Energies for Local Minima of Ac-Pro-NMe Optimized Using QM (MP2/6-31G**+PCM/HF/6-31G**) and MM (ICMFF) Methods

<table>
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<th>Conformer</th>
<th>$\Delta E_{QM}$ (kcal/mol)</th>
<th>$\phi$ (°)</th>
<th>$\psi$ (°)</th>
<th>$\omega$ (°)</th>
<th>$\Delta E_{MM}$ (kcal/mol)</th>
<th>$\phi$ (°)</th>
<th>$\psi$ (°)</th>
<th>$\omega$ (°)</th>
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<td>96.4</td>
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<td>-77.4</td>
<td>151.9</td>
<td>-1.0</td>
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</tbody>
</table>
strong tendency to be buried rather than exposed on the protein surface.

In principle, the optimized parameters are not guaranteed to be optimal, even for the training set of loops, because low-energy conformations, not found during the generation of the training set, may exist. Therefore, the quality of the parameters was first assessed for the same set of nine residue loops by carrying out BPMC loop simulations. Average and median RMSDs for the lowest energy loop conformations produced by the new BPMC runs were in complete agreement with the corresponding values obtained at the end of the parameter optimization procedure, that is, a global conformational search with the optimized solvation parameters did not locate any new low-energy non-native conformations. This result can be attributed to the quality of the training set used in optimization and, therefore, to the efficiency of the search method that is able to explore thoroughly the conformational space of each loop.

Significant improvement over the results obtained with the initial solvation parameters was observed in both average and median RMSDs. Thus, average RMSD decreased from 1.53 Å for the initial solvation model to 0.98 Å for the optimized set, whereas median RMSD dropped by 0.31 Å from 0.75 Å RMSD of the initial model.

To make sure that the optimized solvation parameters are transferable to loops of different lengths and indeed lead to better accuracy of loop predictions, calculations were also carried out for 7 and 11 residue loops using the original SA parameters and the optimized values. In agreement with the results obtained for the training set, the optimized solvation model led to better accuracy of the predictions for both loop lengths. The improvement was particularly significant for 11 residue loops where the average/median RMSD decreased from 2.26/1.64 Å for the initial set of parameters to 1.45/1.00 Å for the optimized solvation model. For seven residue loops, the final average/median RMSD was 0.66/0.33 compared with 0.72/0.35 for the initial parameter set. Because longer loops are more exposed to the solvent, it could be expected that they should be more sensitive to the accuracy of the solvent model.

A comparison of the results obtained with the initial and optimized solvation models demonstrates that parameter optimization using a set of loop conformations and aimed at stabilizing native structure against a large number of decoys represents an efficient method to increase accuracy of a given force field and produces parameters transferable to loops of different lengths.

### Loop modeling results

This section contains discussion of the results obtained for protein loops of 4–13 residues and can be roughly divided into three parts. First, we compare the general performance of two different internal coordinate force fields implemented in the ICM package, that is, ECEPP/3, and the new force field, ICMFF, presented in this work. Performance of ICMFF is discussed in detail in the second part and compared with that of other methods reported in the literature in the last part.

#### Comparison of ECEPP/3 and ICMFF loop simulation results

The average and median RMSD’s computed for 4–11 residue loops with ECEPP/3 and for 4–13 residue loops with ICMFF are listed in Table V. We did not carry out simulations with ECEPP/3 for the longest and the most time-consuming 12 and 13 residue loops. ICMFF performs better for the entire (4–11 residue) range of loop lengths with the average RMSD of more than 20% lower than for the ECEPP/3 force field.

RMSDs obtained with the two force fields grow almost linearly with loop length. Two small peaks in RMSD take place for 9 and 11 residue loops and are more pronounced for ECEPP/3 than for ICMFF. A similar trend, at least for the loops with less than 11 residues, can be observed in the results of Jacobson et al. Because the 4–11 residue loop sets used in this work are the same as those considered by Jacobson et al., it is reasonable to suggest that the unexpectedly higher RMSD for 9 and 11 residue loops may be caused by some other feature of the set. For example, insufficient filtering could lead to a higher percentage of structures with lower experimental accuracy of the loop region. Unusual ionization states of the loop residues, or relatively close proximity of ligands or ions and so forth could also influence the outcome of the prediction.

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<table>
<thead>
<tr>
<th>Loop length</th>
<th>4</th>
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<th>6</th>
<th>7</th>
<th>8</th>
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<th>13</th>
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<td>0.74</td>
<td>1.53</td>
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PROTEINS
Accuracy of the ICMFF-based loop predictions

A detailed examination of the ICMFF results showed that the predictions carried out for short loops (4–7 residues) are in general very accurate with only a few outliers (lowest energy structures with RMSD > 2.0 Å). Thus, median RMSD is in the range of 0.2–0.4 Å which is comparable with the uncertainty of the experimental data. There are no outliers for four residue loops, that is, RMSD never exceeds 0.6 Å. Despite the increase in the average RMSD for longer loops (8–10 residues), the accuracy of the predictions in terms of the median RMSD is still quite high (<0.5 Å). The loops containing 11 and 13 residues appear to be the most challenging for ab initio prediction as indicated by higher values of both average and median RMSD (Table V). Examples of successful predictions obtained for 12-residue loop in 1oth and for 13-residue loop in 1p1m are given in Figure 6(a,b).

Regarding the efficiency of the search algorithm, the results in Table VI show that for short loops with up to eight residues, there was only one sampling error, that is, the situation where no conformation with the energy lower or equal to the energy of the optimized native loop was found (it occurred for one 7-residue loop). The percentage of sampling errors was still small (<6%) for 9–11 residue loops and increased considerably (>14%) for 12 residue loops. No sampling errors occurred for the longest 13 residue loops. A higher percentage of sampling errors for longer loops can be explained not only by the larger number of degrees of freedom but also by the fact that longer loops are usually located farther away from the body of the protein. This leads to the much larger size of the conformational space to be explored because loop conformations are less influenced by the interac-

### Table VI

Percentage of Error Attributable to Sampling

<table>
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<tr>
<th>Loop length</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
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<th>10</th>
<th>11</th>
<th>12</th>
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<tbody>
<tr>
<td>Percentage</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>2</td>
<td>3</td>
<td>6</td>
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<tr>
<td>of error (%)</td>
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</tbody>
</table>
tions with the rest of the protein. Second, on average, the solvent exposure of a loop grows with the length resulting in the higher sensitivity of the result to the accuracy of the solvation model used.

In general, the number of poorly predicted loops (i.e., those with RMSD greater than 2 Å) increases gradually with loop length. Thus, the percentage of incorrect predictions is 0% for 4 residue loops, 12% for 7 residue loops, ~20% for 11 and 12 residue loops, and reaches a maximum of 32% for 13 residue loops. Only one incorrect prediction has energy higher than that of the corresponding optimized native conformation and, therefore, can be explained by a sampling error. To study the origin of all other incorrect predictions, we carried out structural analysis of the corresponding experimental structures. In particular, we were looking for close (<4 Å) contacts between heavy atoms of the loop and any crystallographic neighbor of the protein molecule. Close intermolecular contacts were found in ~76% of cases with many of them involving hydrogen bonds. This result suggests that crystal packing may have significant influence on loop conformations for the cases with high predicted RMSD. Figure 6(c) showing the lowest energy predicted loop conformation for 2rn2 (7 residue loop) overlaid with the corresponding native structure illustrates the effect of crystal packing on the outcome of the loop simulations.

The remaining wrong predictions may be caused by a number of reasons. As mentioned above, highly solvent-exposed loops are more sensitive to the correct balance of intramolecular and solvation energy contributions. An example of this situation is shown in Figure 6(d) for the 11 residue loop in 2eng. This is a particularly interesting case because this loop is packed inside a cavity but with a layer of water molecules separating it from the rest of the protein. An explicit water model may be necessary for correct prediction of this loop.

Other possible factors that may affect the prediction outcome include minor defects of the experimental structures in the vicinity of the loop, as well as uniform protonation state assignment (corresponding to pH 7) regardless of the actual pH of the experiment and local structure environment.

Results obtained for the two loops containing cis-proline [lowest-energy conformations with RMSD of 0.39 Å and 0.27 Å for 1w0n and 2ixt, respectively (Supporting Information Table S11 and S14)] demonstrated that the ICMFF description of the internal energy of proline residue is accurate enough to reproduce the energetic balance between trans- and cis-isomers. Figure 7 shows the overlay of the native and the lowest energy predicted conformations of the 178–186 loop in 2ixt.

**Comparison with previously published methods**

In most cases, a comparison of our results with the previous studies is confounded by the differences in the composition of the test sets and the inclusion of crystallographic neighbors in loop modeling in other studies. However, results reported by Sellers et al. for 6, 8, 10, and 12 residue loops can be compared directly with those discussed in this work because they were obtained without considering the crystal environment. Moreover, our loop sets contain, among others, all the loops used by Sellers et al. We plotted the percentage of loops with RMSD below a given cutoff as a function of the cutoff computed based on the results of Sellers et al. (taken from Supporting Information Table S5 in Ref. 5) and those reported in this work for 6, 8, 10, and 12 residue loops [Fig. 8(a–d)]. For shorter loops (six and eight residues), performance of ICM is clearly superior to that of HLP in terms of average RMSD and very close in terms of median RMSD [average/median RMSD for 6 residue loops is 0.7/0.3 Å for HLP and 0.4/0.3 Å for ICM; RMSD for 8-residue loops is 1.2/0.6 Å for HLP and 0.5/0.5 Å for ICM (Table VII)]. For 10 residue loops, ICM performs somewhat worse than HLP mainly because of the larger number of outliers as indicated by a higher average (0.8 Å vs. 0.6 Å for ICM and HLP, respectively) and almost identical median RMSD (0.5 Å vs. 0.4 Å for ICM and HLP, respectively). Both ICM and HLP methods behave very similarly for 12 residue loops, with an average RMSD of 1.1 and 1.2 Å and a median RMSD of 0.7 and 0.6 Å, respectively. It should be mentioned that the size of the loop sets used for this comparison is relatively small (20 loops for each length) and therefore, the 0.1–0.2 Å difference in RMSD may not be statistically significant.

Results obtained here for 13 residue loops can be compared directly to those of Zhu et al. because the same set of loops was considered in both studies. The average RMSD of our predictions is ~30% higher than that of Zhu et al. (1.67 Å vs. 1.28 Å, Table VII); however, the
median RMSDs are virtually identical for both methods (0.74 and 0.72 for ICMFF and PLOP II, respectively). The analysis of our results for each loop showed that there are three outliers (1hnj, 1jp4, and 1ojq) with RMSD greater than 4 Å. In all three cases, the loop residues form multiple close contacts with crystallographic neighbors. Zhu et al.4 reconstructed the crystal packing (an asymmetric unit plus all atoms from the surrounding symmetric units within 30 Å) in their simulation, which likely helped their method to predict native-like loop conformations for all three proteins.

ICMFF is an internal coordinate force field and as such it relies on the use of standard bond lengths and bond angles (except $NC^\alpha C$). It is reasonable to ask whether the fixed standard residue geometry has any significant effect on the accuracy of the loop modeling. If such an effect exists, it can be expected to be more noticeable for the shortest loops (whereas for longer loops the errors may be obscured by the noise from non-bonded interactions or may compensate each other). The comparison (in terms of median RMSD) with the highly accurate results reported by Jacobson et al.6 for 4 and 5 residue loops and by Sellers et al.5 for 6 residue loops indicates that rigid geometry approximation, at least when it is used in combination with an accurate force field, is unlikely to have any significant effect on the accuracy of the loop modeling. However, this conclusion is based on the analysis of only three outliers, and further studies are needed to confirm this result.

Table VII
Comparison of the ICMFF Loop Prediction Results with Published Works

<table>
<thead>
<tr>
<th>Loop length</th>
<th>RMSD</th>
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aData taken from Table I of Sellers et al.5
bData taken from Table II of Zhu et al.4

Figure 8
Comparison of loop prediction results obtained with ICM (solid line) and reported5 for HLP (dashed line). The y-axis is the percentage of loops for which the backbone RMSD of the lowest energy conformation is at or below the RMSD on the x-axis.

Table VII
Comparison of the ICMFF Loop Prediction Results with Published Works

<table>
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<tr>
<th>Loop length</th>
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<td>na</td>
<td>na</td>
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<td>0.72</td>
</tr>
</tbody>
</table>

aData taken from Table I of Sellers et al.5
bData taken from Table II of Zhu et al.4
field and relaxation of $\angle NC^\alpha$C angle, can provide accuracy of loop predictions which is as good as that of high quality Cartesian potentials. Thus, the same median RMSD of 0.2 Å was obtained for 4 residue loops with ICMFF and reported by Jacobson et al. Both ICM and HLS$^5$ yielded median RMSD of 0.3 Å for 6 residue loops.

As pointed out by Chen et al., rigid covalent geometry in torsion space modeling affects not only the profile of the low-energy regions (which we correct by an appropriate torsion potential) but also the height of the barriers between them. Higher barriers are a significant issue in torsion angle molecular dynamics, and special torsion cross-term corrections have been proposed to recover the potential surface across both, low- and high-energy regions.$^{62}$ In Monte Carlo minimization simulations, where the main goal is to explore the low-energy minima, the barriers issue appears to be rather minor because the search procedure is able to “jump” from one low-energy region to another irrespective of the height of the barrier. Furthermore, introduction of the flexibility of $\angle NC^\alpha$C planar angle in ICMFF reduces the barriers for backbone flexing: indeed, the lowest saddle points on $\phi = 0^\circ$ ($\psi \sim 90^\circ$) and $\psi = 0^\circ$ ($\phi \sim -90^\circ$) rotation barriers are lowered by 6.0 kcal/mol and 0.8 kcal/mol, respectively, as compared with rigid $\angle NC^\alpha$C model. These lower barriers may further ease the sampling of various backbone conformations.

**Computational cost**

All loop simulations were carried out on a cluster of Intel Core2 processors at 2.13 GHz. The average CPU time per single run is 2.5 h for 8-residue loop; 12 h for 10-residue loop and 55 h for 12-residue loop. A sharp increase in computational time with the number of residues in the loop is due to the exponential dependence of the number of energy evaluations required for adequate sampling of the conformational space at the second stage of our protocol. We are currently developing an improved version of the loop modeling algorithm that will be characterized by a more efficient sampling at the first stage and, therefore, will enable us to decrease significantly the number of energy evaluations required for the second, more time consuming stage.

The results reported in this work were obtained by performing five independent runs for each loop. All five runs can be performed simultaneously on a cluster or even on multiple CPU cores of a single workstation.

**CONCLUSIONS**

Loop simulations represent a rigorous test of the force-field accuracy and the sampling method, and are important for many practical applications such as homology modeling and protein design. A comparison with other studies shows that the ICM loop modeling method provides high accuracy of predictions and is on par with the most accurate methods developed so far. Success in loop modeling simulations also provides the first demonstration of the quality of our novel torsion force field, ICMFF. Further tests including energy ranking of protein decoys or folding of small proteins and peptides are necessary for more comprehensive evaluation of the force field. We are planning to carry these tests out in the future.

**ACKNOWLEDGMENTS**

We would like to acknowledge the help of Prof. Harold Scheraga and Dr. Daniel Ripoll of Cornell University.

**REFERENCES**


