

The Rosetta All-Atom Energy Function for Macromolecular Modeling and Design

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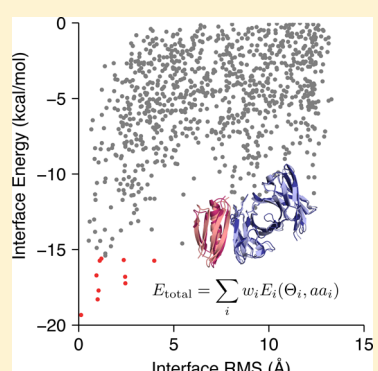
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S Supporting Information

ABSTRACT: Over the past decade, the Rosetta biomolecular modeling suite has informed diverse biological questions and engineering challenges ranging from interpretation of low-resolution structural data to design of nanomaterials, protein therapeutics, and vaccines. Central to Rosetta's success is the energy function: a model parametrized from small-molecule and X-ray crystal structure data used to approximate the energy associated with each biomolecule conformation. This paper describes the mathematical models and physical concepts that underlie the latest Rosetta energy function, called the Rosetta Energy Function 2015 (REF15). Applying these concepts, we explain how to use Rosetta energies to identify and analyze the features of biomolecular models. Finally, we discuss the latest advances in the energy function that extend its capabilities from soluble proteins to also include membrane proteins, peptides containing noncanonical amino acids, small molecules, carbohydrates, nucleic acids, and other macromolecules.



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■ INTRODUCTION

Proteins adopt diverse three-dimensional conformations to carry out the complex mechanisms of life. Their structures are constrained by the underlying amino acid sequence¹ and stabilized by a fine balance between enthalpic and entropic contributions to non-covalent interactions.² Energy functions that seek to approximate the energies of these interactions are fundamental to computational modeling of biomolecular structures. The goal of this paper is to describe the energy calculations used by the Rosetta macromolecular modeling program:³ we explain the underlying physical concepts, mathematical models, latest advances, and application to biomolecular simulations.

Energy functions are based on Anfinsen's hypothesis that native-like protein conformations represent unique, low-energy, thermodynamically stable conformations.⁴ These folded states reside in minima on the energy landscape, and they have a net favorable change in Gibbs free energy, which is the sum of contributions from both enthalpy (ΔH) and entropy ($T\Delta S$), relative to the unfolded state. To follow these heuristics, macromolecular modeling programs require a mathematical function that can discriminate between the unfolded, folded, and native-like conformations. Typically, these functions are linear combinations of terms that compute energies as a function of various degrees of freedom.

The earliest macromolecular energy functions combined a Lennard-Jones potential for van der Waals interactions^{5–7} with harmonic torsional potentials⁸ that were parametrized using force constants from vibrational spectra of small molecules.^{9–11} These formulations were first applied to investigations of the structures of hemolysin,¹² trypsin inhibitor,¹³ and hemoglobin¹⁴ and have now diversified into a large family of commonly used energy functions such as AMBER,¹⁵ DREIDING,¹⁶ OPLS,¹⁷ and CHARMM.^{18,19} Many of these energy functions also rely on new terms and parametrizations. For example, faster computers have enabled the derivation of parameters from ab initio quantum calculations.²⁰ The maturation of X-ray crystallography and NMR protein structure determination methods has enabled the development of statistical potentials derived from per-residue, inter-residue, secondary-structure, and whole-structure features.^{21–28} Additionally, there are alternate models of electrostatics and solvation, such as the generalized Born approximation of the Poisson–Boltzmann equation²⁹ and polarizable electrostatic terms that accommodate varying charge distributions.³⁰

The first version of the Rosetta energy function was developed for proteins by Simons et al.³¹ Initially, it used statistical potentials describing individual residue environments and frequent residue-pair interactions derived from the Protein Data Bank (PDB).³² Later the authors added terms for packing of van der Waals spheres and hydrogen-bonding, secondary-structure, and van der Waals interactions to improve the performance of ab initio structure prediction.³³ These terms were for low-resolution modeling, meaning that the scores were dependent on only the coordinates of the backbone atoms and that interactions between the side chains were treated implicitly.

To enable higher-resolution modeling, in the early 2000s Kuhlman and Baker³⁴ implemented an all-atom energy function that emphasized atomic packing, hydrogen bonding, solvation, and protein torsion angles commonly found in folded proteins. This energy function first included a Lennard-Jones term,³⁵ a

pairwise-additive implicit solvation model,³⁶ a statistically derived electrostatics term, and a term for backbone-dependent rotamer preferences.³⁷ Shortly thereafter, several terms were added, including an orientation-dependent hydrogen-bonding term,³⁸ in agreement with electronic structure calculations.³⁹ This combination of traditional molecular mechanics energies and statistical torsion potentials enabled Rosetta to reach several milestones in structure prediction and design, including accurate ab initio structure prediction,⁴⁰ hot-spot prediction,^{41,42} protein–protein docking,⁴³ small-molecule docking,⁴⁴ and specificity redesign⁴⁵ as well as the first de novo designed protein backbone not found in nature⁴⁶ and the first computationally designed new protein–protein interface.⁴⁷

The Rosetta energy function has changed dramatically since it was last described in complete detail by Rohl et al.⁴⁸ in 2004. It has undergone significant advances ranging from improved models of hydrogen bonding⁴⁹ and solvation⁵⁰ to updated evaluation of backbone⁵¹ and rotamer conformations.⁵² Along the way, these developments have enabled Rosetta to address new biomolecular modeling problems, including the refinement of low-resolution X-ray structures and use of sparse data^{53,54} and the design of vaccines,⁵⁵ biominerization peptides,⁵⁶ self-assembling materials,⁵⁷ and enzymes that perform new functions.^{58,59} Instead of arbitrary units, the energy function is now also fitted to estimate energies in kilocalories per mole. The details of the energy function advances are distributed across code comments, methods development papers, application papers, and individual experts, making it challenging for Rosetta developers and users in both academia and industry to learn the underlying concepts. Moreover, members of the Rosetta community are actively working to generalize the all-atom energy function for use in different contexts^{60,61} and for all macromolecules, including RNA,⁶² DNA,^{63,64} small-molecule ligands,^{65,66} noncanonical amino acids and backbones,^{67–69} and carbohydrates,⁷⁰ further encouraging us to reexamine the underpinnings of the energy function. Thus, there is a need for an up-to-date description of the current energy function.

In this paper, we describe the new default energy function, called the Rosetta Energy Function 2015 (REF15). Our discussion aims to expose the physical and mathematical details of the energy function required for rigorous understanding. In addition, we explain how to apply the computed energies to analyze structural models produced by Rosetta simulations. We hope this paper will provide critically needed documentation of the energy methods as well as an educational resource to help students and scientists interpret the results of these simulations.

■ COMPUTING THE TOTAL ROSETTA ENERGY

The Rosetta energy function approximates the energy of a biomolecule conformation. This quantity, called ΔE_{total} , is computed as a linear combination of energy terms E_i that are calculated as functions of geometric degrees of freedom (Θ) and chemical identities (aa) and scaled by a weight on each term (w_i), as shown in eq 1:

$$\Delta E_{\text{total}} = \sum_i w_i E_i(\Theta_i, \text{aa}_i) \quad (1)$$

Here we explain the Rosetta energy function term by term. First, we describe the energies of interactions between nonbonded atom pairs, which are important for atomic packing, electrostatics, and solvation. Second, we explain the

Table 1. Summary of Terms in REF15 for Proteins

term	description	weight	units	ref(s)
fa_atr	attractive energy between two atoms on different residues separated by a distance d	1.0	kcal/mol	5, 6
fa_rep	repulsive energy between two atoms on different residues separated by a distance d	0.55	kcal/mol	5, 6
fa_intra_rep	repulsive energy between two atoms on the same residue separated by a distance d	0.005	kcal/mol	5, 6
fa_sol	Gaussian exclusion implicit solvation energy between protein atoms in different residues	1.0	kcal/mol	36
lk_ball_wtd	orientation-dependent solvation of polar atoms assuming ideal water geometry	1.0	kcal/mol	50, 71
fa_intra_sol	Gaussian exclusion implicit solvation energy between protein atoms in the same residue	1.0	kcal/mol	36
fa_elec	energy of interaction between two nonbonded charged atoms separated by a distance d	1.0	kcal/mol	50
hbond_lr_bb	energy of short-range hydrogen bonds	1.0	kcal/mol	38, 49
hbond_sr_bb	energy of long-range hydrogen bonds	1.0	kcal/mol	38, 49
hbond_bb_sc	energy of backbone–side-chain hydrogen bonds	1.0	kcal/mol	38, 49
hbond_sc	energy of side-chain–side-chain hydrogen bonds	1.0	kcal/mol	38, 49
dslf_fa13	energy of disulfide bridges	1.25	kcal/mol	49
rama_prep	probability of backbone ϕ, ψ angles given the amino acid type	(0.45 kcal/mol)/kT	kT	50, 51
p_aa_pp	probability of amino acid identity given backbone ϕ, ψ angles	(0.4 kcal/mol)/kT	kT	51
fa_dun	probability that a chosen rotamer is native-like given backbone ϕ, ψ angles	(0.7 kcal/mol)/kT	kT	52
omega	backbone-dependent penalty for cis ω dihedrals that deviate from 0° and trans ω dihedrals that deviate from 180°	(0.6 kcal/mol)/AU	AU ^a	72
pro_close	penalty for an open proline ring and proline ω bonding energy	(1.25 kcal/mol)/AU	AU	51
yhh_planarity	sinusoidal penalty for nonplanar tyrosine χ_3 dihedral angle	(0.625 kcal/mol)/AU	AU	49
ref	reference energies for amino acid types	(1.0 kcal/mol)/AU	AU	1, 51

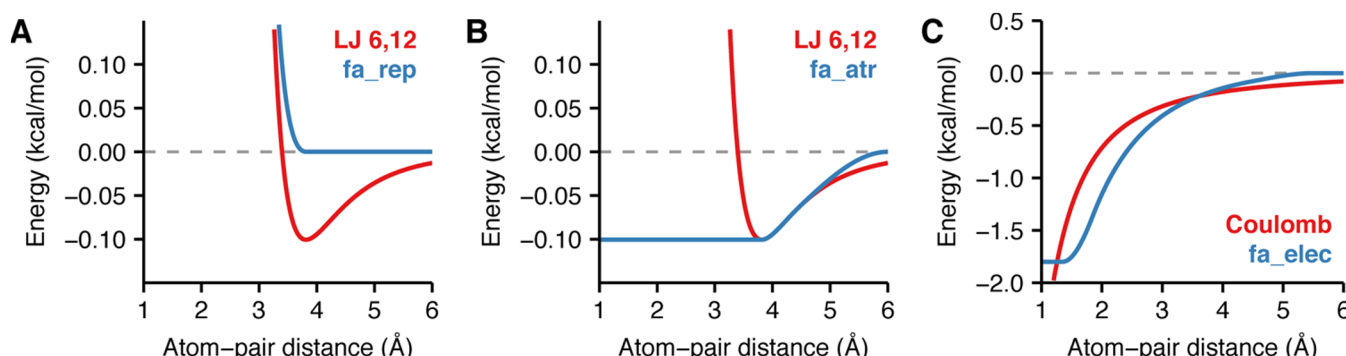
^aAU = arbitrary units.

Figure 1. Van der Waals and electrostatics energies. Shown are comparisons between the pairwise energies of nonbonded atoms computed by Rosetta and the forms computed by traditional molecular mechanics force fields. Here the interactions between the backbone nitrogen and carbon are used as examples. (A) Lennard-Jones van der Waals energy with well-depths $\epsilon_{\text{Nbb}} = 0.162$ kcal/mol and $\epsilon_{\text{Cbb}} = 0.063$ kcal/mol and atomic radii $r_{\text{Nbb}} = 1.763$ Å and $r_{\text{Cbb}} = 2.011$ Å (red) and Rosetta fa_rep (blue). (B) Lennard-Jones van der Waals energy (red) and Rosetta fa_atr (blue). As the atom-pair distance approaches 6.0 Å, the fa_atr term smoothly approaches zero and deviates slightly from the original Lennard-Jones potential. (C) Coulomb electrostatic energy with a dielectric constant $\epsilon = 10$ and partial charges $q_{\text{Nbb}} = -0.604e$ and $q_{\text{Cbb}} = 0.090e$ (red) compared with Rosetta fa_elec (blue). The fa_elec model is shifted to reach zero at the cutoff distance of 6.0 Å.

empirical potentials used to model hydrogen and disulfide bonds. Next, we explain the statistical potentials used to describe backbone and side-chain torsional preferences in proteins. Then we explain a set of terms that accommodate features that are not explicitly captured yet are important for native structural feature recapitulation. Finally, we discuss how the energy terms are combined into a single function used to approximate the energy of biomolecules. For reference, items in the fixed-width font are names of energy terms in the Rosetta code. The energy terms are summarized in Table 1.

Terms for Atom-Pair Interactions. *Van der Waals Interactions.* Van der Waals interactions are short-range attractive and repulsive forces that vary with atom-pair distance. Whereas attractive forces result from the cross-correlated motions of electrons in neighboring nonbonded atoms, repulsive forces occur because electrons cannot occupy the same orbitals by the Pauli exclusion principle. To model van der Waals interactions, Rosetta uses the Lennard-Jones (LJ) 6–

12 potential,^{5,6} which calculates the interaction energy of atoms i and j in different residues given the sum of their atomic radii, σ_{ij} (σ_{ij} in Rosetta has the same definition as r_{ij}^{\min} in CHARMM), the atom-pair distance, d_{ij} , and the geometric mean of their well depths, ϵ_{ij} (eq 2):

$$E_{\text{vdw}}(i, j) = \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{d_{ij}} \right)^{12} - 2 \left(\frac{\sigma_{ij}}{d_{ij}} \right)^6 \right] \quad (2)$$

The atomic radii and well depths are derived from small-molecule liquid-phase data optimized in the context of the energy model.⁵⁰

Rosetta splits the LJ potential at the function's minimum ($d_{ij} = \sigma_{ij}$) into two components that can be weighted separately: attractive (fa_atr) and repulsive (fa_rep). By decomposing the function this way, we can alter component weights without changing the minimum-energy distance or introducing any derivative discontinuities. Many conformational sampling

protocols in Rosetta take advantage of this splitting by slowly increasing the weight of the repulsive component to traverse rugged energy landscapes and to prevent structures from unfolding during sampling.⁷³

The repulsive van der Waals energy, E_{fa_rep} , varies as a function of atom-pair distance. At short distances, atomic overlap results in strong forces that lead to large changes in the energy. The steep $1/d_{ij}^{12}$ term can cause poor performance in minimization routines and overall structure prediction and design calculations.^{74,75} To alleviate this problem, we weaken the repulsive component by replacing the $1/d_{ij}^{12}$ term with a softer linear term for $d_{ij} \leq 0.6\sigma_{ij}$. The term is computed using the atom-type-specific parameters m_{ij} and b_{ij} , which are fit to ensure derivative continuity at $d_{ij} = 0.6\sigma_{ij}$. After the linear component, the function transitions smoothly to the 6–12 form until $d_{ij} = \sigma_{ij}$, where it reaches zero and remains zero (eq 3 and Figure 1A):

$$E_{fa_rep}(i, j) = \sum_{i,j} w_{ij}^{\text{conn}} \begin{cases} m_{ij}d_{ij} + b_{ij} & d_{ij} \leq 0.6\sigma_{ij} \\ \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{d_{ij}} \right)^{12} - 2 \left(\frac{\sigma_{ij}}{d_{ij}} \right)^6 + 1 \right] & 0.6\sigma_{ij} < d_{ij} \leq \sigma_{ij} \\ 0 & \sigma_{ij} < d_{ij} \end{cases} \quad (3)$$

Rosetta also includes an intraresidue version of the repulsive component, $E_{fa_intra_rep}$, with the same functional form as the E_{fa_rep} term (eq 3). We include this term because the knowledge-based rotamer energy (E_{fa_dun} ; see below) underestimates intraresidue collisions.

The attractive van der Waals energy, E_{fa_atr} , has a value of $-\epsilon_{ij}$ for $d_{ij} \leq \sigma_{ij}$ and then transitions to the 6–12 potential as the distance increases (eq 4 and Figure 1B):

$$E_{fa_atr} = \sum_{i,j} w_{ij}^{\text{conn}} \begin{cases} -\epsilon_{ij} & d_{ij} \leq \sigma_{ij} \\ \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{d_{ij}} \right)^{12} - 2 \left(\frac{\sigma_{ij}}{d_{ij}} \right)^6 \right] & \sigma_{ij} < d_{ij} \leq 4.5 \text{ \AA} \\ f_{\text{poly}}(d_{ij}) & 4.5 \text{ \AA} < d_{ij} \leq 6.0 \text{ \AA} \\ 0 & 6.0 \text{ \AA} < d_{ij} \end{cases} \quad (4)$$

For speed, we truncate the LJ term beyond $d_{ij} = 6.0 \text{ \AA}$, where the van der Waals forces are small. To avoid derivative discontinuities, we use a cubic polynomial function, $f_{\text{poly}}(d_{ij})$, for $d_{ij} > 4.5 \text{ \AA}$ to transition the standard Lennard-Jones functional form smoothly to zero. These smooth derivatives are necessary to ensure that bumps do not accumulate in the distributions of structural features at inflection points in the energy landscape during conformational sampling with gradient-based minimization (Sheffler, unpublished, 2006).

Both terms are multiplied by a connectivity weight, w_{ij}^{conn} , to exclude the large repulsive energetic contributions that would otherwise be calculated for atoms separated by fewer than four chemical bonds (eq 5).

$$w_{ij}^{\text{conn}} = \begin{cases} 0 & n_{ij}^{\text{bonds}} \leq 3 \\ 0.2 & n_{ij}^{\text{bonds}} = 4 \\ 1 & n_{ij}^{\text{bonds}} \geq 5 \end{cases} \quad (5)$$

To ensure the connectivity rules include both atoms in a dipole, if an atom is part of a strong dipole, then we count bonds (n_{ij}^{bonds}) to include both atoms in the dipole. Such weights are common to molecular force fields that assume that covalent bonds are not formed or broken during a simulation. Rosetta uses four chemical bonds as the “crossover” separation when w_{ij}^{conn} transitions from 0 to 1 (rather than three chemical bonds as used by traditional force fields) to limit the effects of double counting due to knowledge-based torsional potentials.

The comparison between eq 2 and the modified LJ potential (eqs 3 and 4) is shown in Figure 1A,B.

Electrostatics. Nonbonded electrostatic interactions arise from forces between fully and partially charged atoms. To evaluate these interactions, Rosetta uses Coulomb’s law with partial charges originally taken from CHARMM and adjusted via a group optimization scheme (Table S3 in the Supporting Information).⁵⁰ Coulomb’s law is a pairwise term commonly expressed in terms of d_{ij} , the dielectric constant, ϵ , the partial atomic charges for each atom, q_i and q_j , and Coulomb’s constant, $C_0 (=322 \text{ \AA kcal/mol } e^{-2}$, where e is the elementary charge) (eq 6):

$$E_{\text{Coulomb}}(i, j) = \frac{C_0 q_i q_j}{\epsilon} \frac{1}{d_{ij}} \quad (6)$$

To approximate electrostatic interactions in biomolecules, we modify the potential to account for the difference in dielectric constant between the protein core and the solvent-exposed surface.⁷⁶ Specifically, we substitute the constant ϵ in eq 6 with a sigmoidal function $\epsilon(d_{ij})$ that increases from $\epsilon_{\text{core}} = 6$ to $\epsilon_{\text{solvent}} = 80$ when d_{ij} is between 0 and 4 Å (eqs 7 and 8):

$$\epsilon(d_{ij}) = g\left(\frac{d_{ij}}{4}\right)\epsilon_{\text{core}} + \left[1 - g\left(\frac{d_{ij}}{4}\right)\right]\epsilon_{\text{solvent}} \quad (7)$$

$$g(x) = \left(1 + x + \frac{x^2}{2}\right) \exp(-x) \quad (8)$$

As with the van der Waals term, we make several heuristic approximations to adapt this calculation for simulations of biomolecules. To avoid strong repulsive forces at short distances, we replace the steep gradient with the constant $E_{\text{elec}}(d_{\min})$ for $d_{ij} < 1.45 \text{ \AA}$. Next, since the distance-dependent dielectric assumption results in dampened long-range electrostatics, for speed we truncate the potential at $d_{\max} = 5.5 \text{ \AA}$ and modify the Coulomb curve by subtracting $1/d_{\max}^2$ to shift the potential to zero at $d_{ij} = d_{\max}$ (eq 9).

$$E_{\text{elec}}(i, j, d_{ij}) = \frac{C_0 q_i q_j}{\epsilon(d_{ij})} \begin{cases} \frac{1}{d_{ij}^2} - \frac{1}{d_{\max}^2} & d_{ij} \leq d_{\max} \\ 0 & d_{\max} < d_{ij} \end{cases} \quad (9)$$

We use the cubic polynomials $f_{\text{poly,low}}(d_{ij})$ and $f_{\text{poly,high}}(d_{ij})$ to smooth between the traditional form and our adjustments while avoiding derivative discontinuities. The energy is also multiplied by the connectivity weight, w_{ij}^{conn} (eq 5). The final

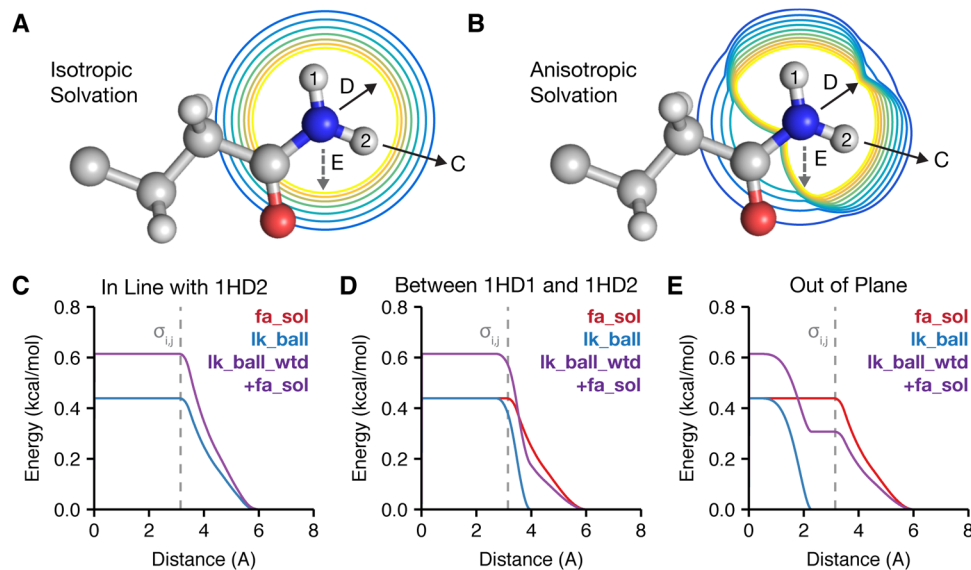


Figure 2. Two-component Lazaridis–Karplus solvation model. Rosetta uses two energy terms to evaluate the desolvation of protein side chains: an isotropic term (*fa*_sol) and an anisotropic term (*lk*_ball_wtd). (A) and (B) demonstrate the difference between isotropic and anisotropic solvation of the NH₂ group by CH₃ on the asparagine side chain. The contours vary from low energy (blue) to high energy (yellow). The arrows represent the approach vectors for the pair potentials shown in (C–E), where we compare the *fa*_sol, *lk*_ball, and *lk*_ball_wtd + *fa*_sol energies for the solvation of the NH₂ group on asparagine for three different approach angles: (C) in line with the 1HD2 atom, (D) along the bisector of the angle between 1HD1 and 1HD2, and (E) vertically down from above the plane of the hydrogens (out of plane).

modified electrostatic potential is given by eq 10 and compared with the standard form in Figure 1C.

$$E_{\text{fa_elec}} = \sum_{i,j} w_{i,j}^{\text{conn}} \begin{cases} E_{\text{elec}}(i, j, d_{\min}) & d_{i,j} < 1.45 \text{ \AA} \\ f_{\text{poly}}^{\text{elec,low}}(d_{i,j}) & 1.45 \text{ \AA} \leq d_{i,j} < 1.85 \text{ \AA} \\ E_{\text{elec}}(i, j, d_{i,j}) & 1.85 \text{ \AA} \leq d_{i,j} < 4.5 \text{ \AA} \\ f_{\text{poly}}^{\text{elec,high}}(d_{i,j}) & 4.5 \text{ \AA} \leq d_{i,j} < 5.5 \text{ \AA} \\ 0 & 5.5 \text{ \AA} \leq d_{i,j} \end{cases} \quad (10)$$

Solvation. Native-like protein conformations minimize the exposure of hydrophobic side chains to the surrounding polar solvent. Unfortunately, explicitly modeling all of the interactions between solvent and protein atoms is computationally expensive. Instead, Rosetta represents the solvent as bulk water based upon the Lazaridis–Karplus (LK) implicit Gaussian exclusion model.³⁶ Rosetta’s solvation model has two components: an isotropic solvation energy, called *fa*_sol, which assumes that bulk water is uniformly distributed around the atoms (Figure 2A), and an anisotropic solvation energy, called *lk*_ball_wtd, which accounts for specific waters near polar atoms that form the solvation shell (Figure 2B).

The isotropic (LK) model³⁶ is based on the function *f*_{desolv} that describes the energy required to desolvate (remove contacting water) an atom *i* when it is approached by a neighboring atom *j*. In Rosetta, we exclude the LK Δ*G*^{ref} term because we implement our own reference energy (discussed later). The energy of the atom-pair interaction varies with the separation distance, *d*_{*i,j*}, the experimentally determined vapor-to-water transfer free energy, Δ*G*^{free}_{*i*}, the sum of the atomic radii, *σ*_{*i,j*}, the correlation length, *λ*_{*i*}, and the atomic volume of the desolvating atom, *V*_{*j*} (eq 11):

$$f_{\text{desolv}} = -V_j \frac{\Delta G_i^{\text{free}}}{2\pi^{3/2}\lambda_i\sigma_i^2} \exp\left[-\left(\frac{d_{i,j} - \sigma_{i,j}}{\lambda_i}\right)^2\right] \quad (11)$$

At short distances, *fa*_rep prevents atoms from overlapping; however, many protocols briefly downweight or disable the *fa*_rep term. To avoid scenarios where *f*_{desolv} encourages atom-pair overlap in the absence of *fa*_rep, we smoothly increase the value of the function to a constant at short distances where the van der Waals spheres overlap (*d*_{*i,j*} = *σ*_{*i,j*}). At large distances, the function asymptotically approaches zero; therefore, we truncate the function at 6.0 Å for speed. We also transition between the constants at short and long distances using the distance-dependent cubic polynomials *f*_{poly}^{solv,low} and *f*_{poly}^{solv,high} with constants *c*₀ = 0.3 Å and *c*₁ = 0.2 Å that define a window for smoothing. The overall desolvation function is given by eq 12:

$$g_{\text{desolv}} = \begin{cases} f_{\text{desolv}}(i, j, \sigma_{i,j}) & d_{i,j} \leq \sigma_{i,j} - c_0 \\ f_{\text{poly}}^{\text{solv,low}}(i, j, d_{i,j}) & \sigma_{i,j} - c_0 < d_{i,j} \leq \sigma_{i,j} + c_1 \\ f_{\text{desolv}}(i, j, d_{i,j}) & \sigma_{i,j} + c_1 < d_{i,j} \leq 4.5 \text{ \AA} \\ f_{\text{poly}}^{\text{solv,high}}(i, j, d_{i,j}) & 4.5 \text{ \AA} < d_{i,j} \leq 6.0 \text{ \AA} \\ 0 & 6.0 \text{ \AA} < d_{i,j} \end{cases} \quad (12)$$

The total isotropic solvation energy, *E*_{fa,sol}, is computed as a sum of the energies for atom *j* desolvating atom *i* and vice versa, scaled by the previously defined connectivity weight (eq 13):

$$E_{\text{fa,sol}} = \sum_{i,j} w_{i,j}^{\text{conn}} [g_{\text{desolv}}(i, j) + g_{\text{desolv}}(j, i)] \quad (13)$$

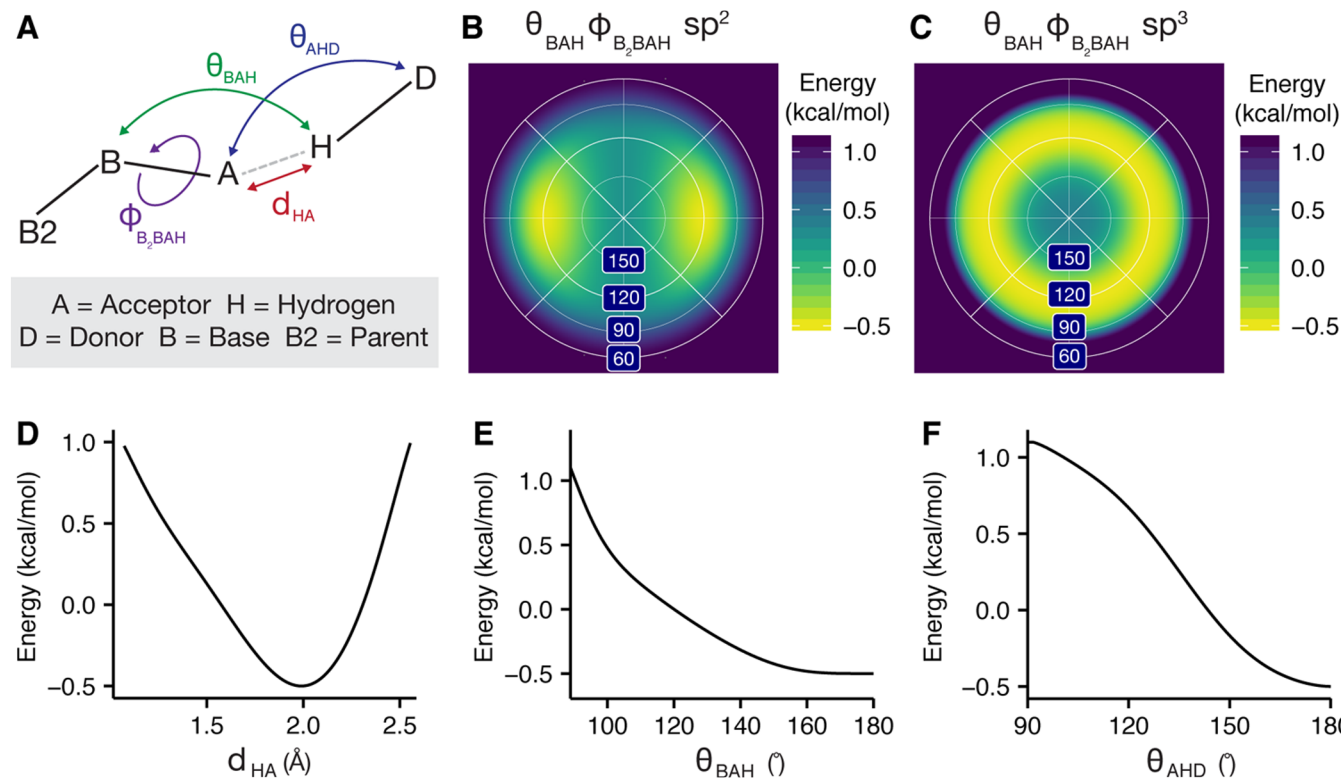


Figure 3. Orientation-dependent hydrogen bonding model. (A) Degrees of freedom evaluated by the hydrogen-bonding term: the acceptor–donor distance, d_{HA} ; the angle between the base, acceptor, and hydrogen, θ_{BAH} ; the angle between the acceptor, hydrogen, and donor, θ_{AHD} ; and the dihedral angle corresponding to rotation around the base–acceptor bond, ϕ_{B_2BAH} . (B) Lambert azimuthal projection of the $E_{hbond}^{B_2BAH}$ energy landscape for an sp^2 -hybridized acceptor.⁴⁹ (C) $E_{hbond}^{B_2BAH}$ energy landscape for an sp^3 -hybridized acceptor. (D–F) Example energies for hydrogen bonding of the histidine imidazole ring acceptor with a protein backbone amide: (D) energy E_{hbond}^{BAH} vs the acceptor–donor distance d_{HA} ; (E) energy E_{hbond}^{BAH} vs the base–acceptor–hydrogen angle θ_{BAH} ; (F) energy E_{hbond}^{AHD} vs the acceptor–hydrogen–donor angle θ_{AHD} .

Rosetta also includes an intraresidue version of the isotropic solvation energy, fa_intra_sol , with the same functional form as the fa_sol term (eq 13).

A recent innovation (2016) is the addition of the energy term lk_ball_wtd to model the orientation-dependent solvation of polar atoms. This anisotropic model increases the desolvation penalty for occluding polar atoms near sites where waters may form hydrogen-bonding interactions. For polar atoms, we subtract off part of the isotropic energy given by eq 13 and then add the anisotropic energy to account for the position of the desolvating atom relative to hypothesized water positions.

To compute the anisotropic energy, we first calculate the set of ideal water sites around atom i , $\mathcal{W}_i = \{\nu_{i1}, \nu_{i2}, \dots\}$. This set contains one to three water sites, depending on the atom type of atom i . Each site is 2.65 Å from atom i and has an optimal hydrogen-bonding geometry, and we consider the potential overlap of a desolvating atom j with each water. The overlap is considered negligible until the van der Waals sphere of the desolvating atom j (with radius σ_j) touches the van der Waals sphere of the water (with radius σ_w) at site k , and then the term smoothly increases over a zone of partial overlap of approximately 0.5 Å. Thus, for each water site k with coordinates $\nu_{i,k}$ we compute an occlusion measure d_k^2 to capture the gap between the hypothetical water and the desolvating atom j , using the offset $\Omega = 3.7$ Å² to provide the ramp-up buffer (eq 14):

$$d_k^2 = \| \mathbf{r}_j - \mathbf{r}_{i,k} \|^2 - (\sigma_w + \sigma_j)^2 + \Omega \quad (14)$$

Next, we find the soft minimum of d_k^2 over all water sites in \mathcal{W}_i by computing the logarithmic average:

$$d_{min}^2(i, j) = -\ln \left[\sum_{k \in \mathcal{W}_i} \exp(-d_k^2) \right] \quad (15)$$

Then d_{min}^2 and Ω are used to compute a damping function, f_{lkfrac} , that varies from 0 when the desolvating atom is at least a van der Waals distance from any preferred water site to 1 when the desolvating atom overlaps a water site by more than ~0.5 Å (eq 16):

$$f_{lkfrac}(i, j) = \begin{cases} 1 & d_{min}^2(i, j) < 0 \\ \left[1 - \left(\frac{d_{min}^2(i, j)}{\Omega} \right) \right]^2 & 0 \leq d_{min}^2(i, j) < \Omega \\ 0 & \Omega \leq d_{min}^2(i, j) \end{cases} \quad (16)$$

We calculate the anisotropic energy for desolvating a polar atom, E_{lk_ball} , by scaling the desolvation function g_{desolv} by the damping function f_{lkfrac} and an atom-type-specific weight, w_{aniso} , which is typically ~0.7 (eq 17):

$$E_{lk_ball}(i, j) = w_{aniso,i} g_{desolv}(i, j) f_{lkfrac}(i, j) \quad (17)$$

The amount of isotropic solvation energy subtracted is g_{desolv} multiplied by w_{iso} , where w_{iso} is an atom-type-specific weight, typically ~0.3 (eq 18):

$$E_{lk_ball_iso}(i, j) = -w_{iso}g_{desolv}(i, j) \quad (18)$$

The total weight on the isotropic contribution through both the `fa_sol` and `lk_ball_wtd` terms is thus ~ 0.7 . The isotropic and anisotropic components are then summed to yield a new desolvation function, h_{desolv} (eq 19):

$$h_{desolv}(i, j) = E_{lk_ball_iso}(i, j) + E_{lk_ball}(i, j) \quad (19)$$

Like `fa_sol`, the energies for desolvation of atom i by atom j and desolvation of atom j by atom i are summed to yield the overall `lk_ball_wtd` energy, but counting only the desolvation of polar, hydrogen-bonding heavy atoms (O, N), defined as the set \mathcal{P} (eq 20):

$$E_{lk_ball_wtd} = \sum_{i \in \mathcal{P}} w_{i,j}^{conn} h_{desolv}(i, j) + \sum_{j \in \mathcal{P}} w_{i,j}^{conn} h_{desolv}(j, i) \quad (20)$$

Figure 2 shows comparisons of `fa_sol`, `lk_ball` (eq 17), and the sum of `fa_sol` and `lk_ball_wtd` for the example of an asparagine NH₂ desolvated from three different approach angles. As the approach angle varies, the sum of `lk_ball_wtd` and `fa_sol` creates a larger desolvation penalty when water sites are occluded and a smaller penalty otherwise, relative to `fa_sol` alone.

Hydrogen Bonding. Hydrogen bonds are partially covalent interactions that form when a nucleophilic heavy atom donates electron density to a polar hydrogen.⁷⁷ At short ranges (< 2.5 Å), they exhibit geometries that maximize orbital overlap.⁷⁸ The interactions between hydrogen-bonding groups are also partially described by electrostatics. While this hybrid covalent–electrostatic character is complex, it is crucial for capturing the structural specificity that underlies protein folding, function, and interactions.

Rosetta calculates the energies of hydrogen bonds using `fa_elec` and a hydrogen-bonding model that evaluates energies on the basis of orientation preferences of hydrogen bonds found in high-resolution crystal structures.^{38,49} To derive this model, we curated intraprotein polar contacts from ~ 8000 high-resolution crystal structures (the Top8000 data set⁷⁹) and identified features using adaptive density estimation. We then empirically fit the functional form of the energy such that the Rosetta-generated polar contacts mimic the distributions from Top8000. The resulting hydrogen-bonding energy is evaluated for all pairs of donor hydrogens, H, and acceptors, A, as a function of four degrees of freedom (Figure 3A): (1) the distance between the donor and acceptor, d_{HA} ; (2) the angle θ_{AHD} formed by the acceptor, the donor H, and the donor heavy atom, D; (3) the angle θ_{BAH} formed by the acceptor's parent atom ("base"), B, the acceptor, and the donor; and (4) the torsion angle ϕ_{B_2BAH} formed by the donor H, the acceptor, and two subsequent parent atoms B and B₂. B, the parent atom of A, is the first atom on the shortest path to the root atom (e.g., C_α). The B₂ atom of A is the parent atom of B (e.g., the sp² plane is defined by B₂, B, and A). For convenience, the hydrogen-bonding energy is subdivided into four separate terms: long-range backbone hydrogen bonds (`hbond_lr_bb`), short-range backbone hydrogen bonds (`hbond_sr_bb`), hydrogen bonds between backbone and side-chain atoms (`hbond_bb_sc`), and hydrogen bonds between side-chain atoms (`hbond_sc`).

To avoid overcounting, side chain to backbone hydrogen bonds are excluded if the backbone group is already involved in

a hydrogen bond to prevent formation of too many i to $i\text{-}4$ or i to $i\text{-}3$ hydrogen bonds for helical serines and threonines. For speed, the component terms have simple analytic functional forms (Figure 3B–F and eqs S1–S7 in the Supporting Information). The term is also multiplied by two atom-type-specific weights, w_H and w_A , that account for the varying strength of hydrogen bonds. The overall model is given by eq 21, in which the $E_{hbond}^{B,BAH}$ term depends on the orbital hybridization of the acceptor, ρ , and the function $f(x)$ (eq 22) is used for smoothing to avoid derivative discontinuities and ensure that edge-case hydrogen bonds are considered:

$$\begin{aligned} E_{hbond} = & \sum_{H,A} w_H w_A f(E_{hbond}^{HA}(d_{HA}) + E_{hbond}^{AHD}(\theta_{AHD}) \\ & + E_{hbond}^{BAH}(\theta_{BAH}) + E_{hbond}^{B,BAH}(\rho, \phi_{B_2BAH}, \theta_{BAH})) \end{aligned} \quad (21)$$

$$f(x) = \begin{cases} x & x < -0.1 \\ -0.025 + \frac{x}{2} - 2.5x^2 & -0.1 \leq x < 0.1 \\ 0 & 0.1 \leq x \end{cases} \quad (22)$$

Disulfide Bonding. Disulfide bonds are covalent interactions that link sulfur atoms in cysteine residues. In Rosetta we typically rely on a tree-based kinematic system^{3,80} to keep bond lengths and angles fixed so that we may sample the conformation space by changing only torsions. For this reason, we do not generally need terms that evaluate bond-length and bond-angle energetics. However, with disulfide bonds and proline (discussed below), the extra bonds cannot be represented with a tree (since a tree graph is acyclic) and thus must be treated explicitly. Therefore, disulfide bonds are a special case of inter-residue covalent contacts that requires a representation with more degrees of freedom. To evaluate disulfide-bonding interactions, Rosetta identifies pairs of cysteines that have covalent bonds linking the S_γ atoms and computes the energies of these interactions using an orientation-dependent model called `dslf_fa13`.⁴⁹ The model was derived by curating intraprotein disulfide bonds from Top8000 and identifying features using kernel density estimates. For speed, the feature distributions are modeled using skewed Gaussian functions and a mixture of one, two, and three von Mises functions (eqs S8–S11).

The overall disulfide energy is computed as a function of six degrees of freedom (Figure 4) that map to four component energies. First, the component due to the sulfur–sulfur distance, d_{SS} , is evaluated as $E_{dsf}^{SS}(d_{SS})$. Second, the components due to the angles formed by C_{β1} and C_{β2} with the S–S bond, $\theta_{C_{\beta1}SS}$ and $\theta_{C_{\beta2}SS}$, respectively, are evaluated as $E_{dsf}^{CSS}(\theta_{C_{\beta1}SS})$. Third, the components due to the dihedral angles formed by C_{α1}C_{β1} and C_{α2}C_{β2} with the S–S bond, $\phi_{C_{\alpha1}C_{\beta1}SS}$ and $\phi_{C_{\alpha2}C_{\beta2}SS}$, respectively, are evaluated as $E_{dsf}^{C_{\alpha}C_{\beta}SS}(\phi_{C_{\alpha1}C_{\beta1}SS})$. Finally, the dihedral angle formed by C_{β1}, C_{β2}, and the S–S bond, $\phi_{C_{\beta1}SSC_{\beta2}}$, is evaluated as $E_{dsf}^{C_{\beta}SSC_{\beta}}(\phi_{C_{\beta1}SSC_{\beta2}})$. The complete disulfide bonding energy evaluated for all S–S pairs is given by eq 23:

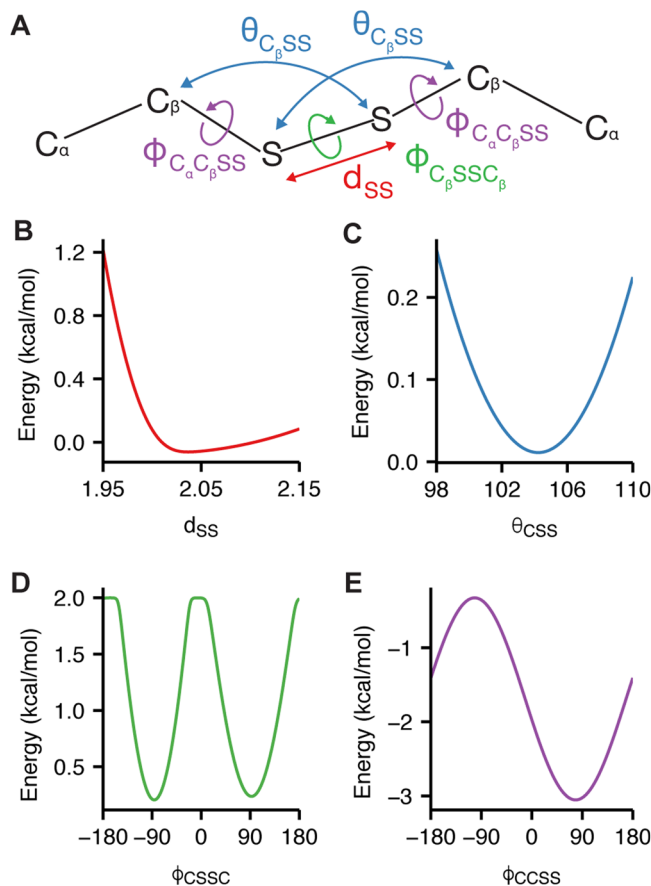


Figure 4. Orientation-dependent disulfide bonding model. (A) Degrees of freedom evaluated by the disulfide bonding energy: the sulfur–sulfur distance, d_{ss} ; the angle formed by C_β and the two sulfur atoms, $\theta_{C_\beta SS}$; the dihedral angle corresponding to rotation about the C_β –sulfur bond, $\phi_{C_\alpha C_\beta SS}$; and the dihedral angle corresponding to rotation about the S–S bond, $\phi_{C_\beta SSC_\beta}$. (B–E) Plots of the energy terms (B) $E_{\text{dslf}}^{\text{SS}}(d_{ss})$, (C) $E_{\text{dslf}}^{\text{CSS}}(\theta_{C_\beta SS})$, (D) $E_{\text{dslf}}^{C_\beta SSC_\beta}(\phi_{C_\beta SSC_\beta})$, and (E) $E_{\text{dslf}}^{C_\alpha C_\beta SS}(\phi_{C_\alpha C_\beta SS})$.

$$\begin{aligned} E_{\text{dslf_fa13}} = & \sum_{S_1, S_2} E_{\text{dslf}}^{\text{SS}}(d_{SS}) + E_{\text{dslf}}^{\text{CSS}}(\theta_{C_\beta SS}) + E_{\text{dslf}}^{\text{CSS}}(\theta_{C_\beta SS}) \\ & + E_{\text{dslf}}^{C_\alpha C_\beta SS}(\phi_{C_\alpha C_\beta SS}) + E_{\text{dslf}}^{C_\alpha C_\beta SS}(\phi_{C_\alpha C_\beta SS}) \\ & + E_{\text{dslf}}^{C_\beta SSC_\beta}(\phi_{C_\beta SSC_\beta}) \end{aligned} \quad (23)$$

Terms for Protein Backbone and Side-Chain Torsions.

Rosetta evaluates backbone and side-chain conformations in torsion space to greatly reduce the search domain and increase the computational efficiency. Traditional molecular mechanics force fields describe torsional energies in terms of sines and cosines, which have at times performed poorly at reproducing the observed backbone dihedral distributions in unstructured regions.⁸¹ Instead, Rosetta uses several knowledge-based terms for torsion angles that are fast approximations of quantum effects and more accurately model the preferred conformations of protein backbones and side chains.

Ramachandran Maps. To evaluate the backbone ϕ and ψ angles, we define an energy term called `rama_prep` based on Ramachandran maps for each amino acid using torsions from 3985 protein chains with a resolution of ≤ 1.8 Å, R factor of ≤ 0.22 and sequence identity of $\leq 50\%$.⁸² Amino acids with low electron density (in the bottom 25th percentile of each residue type) were removed from the data set. The resulting $\sim 581\,000$ residues were used in adaptive kernel density estimates⁵² of Ramachandran maps with a grid step of 10° for both ϕ and ψ . Residues preceding proline are also treated separately because they exhibit distinct ϕ , ψ preferences due to steric interactions with the proline C_δ .⁸³ The energy, called $E_{\text{rama_prep}}$, is then computed by converting the probabilities to energies at the grid points via the inverted Boltzmann relation⁸⁴ (eq 24 and Figure 5):

$$E_{\text{rama_prep}} = \sum_i \begin{cases} -\ln[P_{\text{reg}}(\phi_i, \psi_i | \text{aa}_i)] & \text{C-terminus, } i+1 \neq \text{Pro} \\ -\ln[P_{\text{prep}}(\phi_i, \psi_i | \text{aa}_i)] & i+1 = \text{Pro} \end{cases} \quad (24)$$

The energies are then evaluated using bicubic interpolation. The [Supporting Information](#) includes a detailed discussion of why interpolation is performed on the backbone torsional

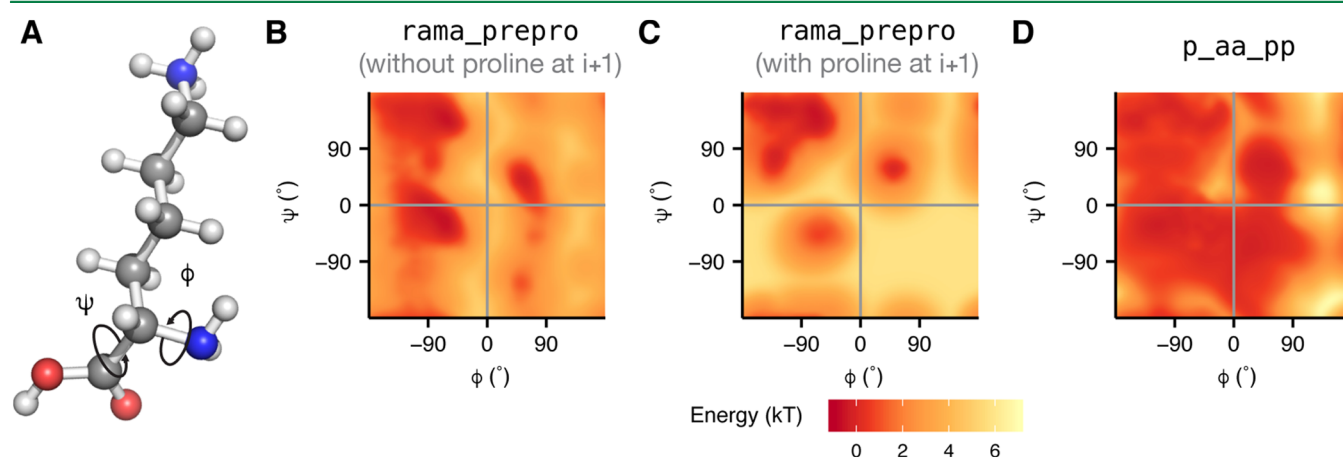


Figure 5. Backbone torsion energies. (A) The angle ϕ is defined by the backbone atoms C_{i-1} –N– C_α – C_i , and the angle ψ is defined by N– C_α – C_i – N_{i+1} . (B, C) Backbone-dependent torsion energies ($E_{\text{rama_prep}}$) for the lysine residue (B) without a proline at $i+1$ and (C) with a proline at $i+1$. (D) $E_{\text{p_aa_pp}}$ of lysine.

energies rather than the probabilities (Figure S3 and eqs S12 and S13).

Backbone Design Term. Rosetta also computes the likelihood of placing a specific amino acid side chain given an existing ϕ , ψ backbone conformation. This term, called $p_{\text{aa_pp}}$, represents the propensity to observe an amino acid relative to the other 19 canonical amino acids.⁸⁵ The knowledge-based propensity, $P(\text{aal}\phi, \psi)$ (eq 25), was derived using the adaptive kernel density estimates for $P(\phi, \psi|aa)$ and Bayes' rule, and the equation for $E_{p_{\text{aa_pp}}}$ is given in eq 26 (Figure 5D):

$$P(\text{aal}\phi, \psi) = \frac{P(\phi, \psi|aa)P(aa)}{\sum_{aa'} P(\phi, \psi|aa')} \quad (25)$$

$$E_{p_{\text{aa_pp}}} = \sum_r -\ln \left[\frac{P(aa_r|\phi, \psi_r)}{P(aa_r)} \right] \quad (26)$$

Side-Chain Conformations. Protein side chains mostly occupy discrete conformations (rotamers) separated by large energy barriers. To evaluate rotamer conformations, Rosetta derives probabilities from the 2010 backbone-dependent rotamer library (<http://dunbrack.fccc.edu/bbdep2010/>), which contains the frequencies, means, and standard deviations of individual χ angles for each χ angle k of each rotamer of each amino acid type.⁵² The probability has three components: (1) observing a specific rotamer given the backbone dihedral angles; (2) observing specific χ angles given the rotamer; and (3) observing the terminal χ angle distribution, which is either Gaussian-like or continuous when the terminal χ angle is sp^2 -hybridized (eq 27):

$$P(\chi|\phi, \psi, aa) = P(\text{rot}\phi, \psi, aa) \left[\prod_{k < T} P(\chi_k|\phi, \psi, \text{rot}, aa) \right] \\ P(\chi_T|\phi, \psi, \text{rot}, aa) \quad (27)$$

where T is the number of rotameric χ angles plus 1.

The 2010 rotamer library distinguishes between rotameric and nonrotameric torsions. A torsion is rotameric when the third of the four atoms defining the torsion is sp^3 -hybridized (i.e., preferring $\sim 60^\circ$, $\sim 180^\circ$, and $\sim -60^\circ$ with steep energy barriers between the wells). If the last χ torsion is rotameric, the probability $p(\chi_T|\phi, \psi, \text{rot}, aa)$ is fixed at 1. On the other hand, a torsion is nonrotameric if its third atom is sp^2 -hybridized: the library describes its probability distribution continuously instead. The category of semirotameric amino acids with both rotameric and nonrotameric dihedrals encompasses eight amino acids: Asp, Asn, Gln, Glu, His, Phe, Tyr, and Trp.⁸⁶

The probability of each rotamer, $P(\text{rot}\phi, \psi, aa)$, is derived from the same data set as the Ramachandran maps described above. The probabilities were identified using adaptive kernel density estimation, and the same data set is used to estimate the mean μ_χ and standard deviation σ_χ for each χ dihedral in the rotamer as functions of the backbone dihedrals, allowing us to compute a probability for the χ values using eq 28:

$$P(\chi_k|\phi, \psi, \text{rot}) = \exp \left[-\frac{1}{2} \left(\frac{\chi_k - \mu_{\chi_k}(\phi, \psi|rot, aa)}{\sigma_{\chi_k}(\phi, \psi|rot, aa)} \right)^2 \right] \quad (28)$$

This formulation is reminiscent of the Gaussian distribution, except that it is missing the normalization coefficient of $[2\pi\sigma_{\chi_k}(\phi, \psi|rot, aa)]^{-1/2}$. Taking the logarithm of this probability gives a term resembling Hooke's law with the spring constant given by $\sigma_{\chi_k}^{-2}(\phi, \psi|rot, aa)$.

The full form of $E_{fa_{\text{dun}}}$ is given by eq 29 as a sum over all residues r . The difference between the rotameric and semirotameric models is also shown in Figure 6.

$$E_{fa_{\text{dun}}} = \sum_r -\ln[P(\text{rot}_r|\phi_r, \psi_r, aa_r)] \\ + \sum_{k < T_r} \frac{1}{2} \left(\frac{\chi_{k,r} - \mu_{\chi_k}(\phi_r, \psi_r|rot_r, aa_r)}{\sigma_{\chi_k}(\phi_r, \psi_r|rot_r, aa_r)} \right)^2 \\ - \ln[P(\chi_{T_r,r}|\phi_r, \psi_r, rot_r, aa_r)] \quad (29)$$

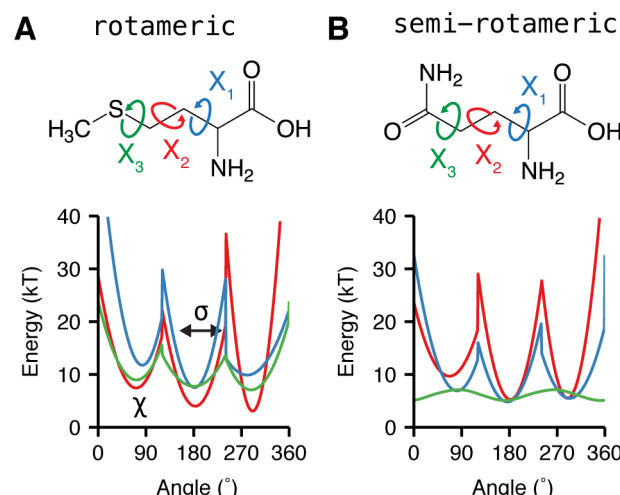


Figure 6. Energies for side-chain rotamer conformations. The Dunbrack rotamer energy, $E_{fa_{\text{dun}}}$, is dependent on both the ϕ and ψ backbone torsions and the χ side-chain torsions. Here we demonstrate the variation of $E_{fa_{\text{dun}}}$ when the backbone is fixed in an α -helical conformation with $\phi = -57^\circ$ and $\psi = -47^\circ$ and the χ values can vary. χ_1 is shown in blue, χ_2 in red, and χ_3 in green. (A) χ -dependent Dunbrack energy of methionine with an sp^3 -hybridized terminus. (B) χ -dependent energy of glutamine with an sp^2 -hybridized χ_3 terminus. χ_1 , χ_2 , and χ_3 of methionine and χ_1 and χ_2 of glutamine express rotameric behavior, while χ_3 of the latter expresses broad nonrotameric behavior.

The energy from $-\ln[P(\text{rot}_r|\phi_r, \psi_r, aa_r)]$ is computed using bicubic-spline interpolation; $P(\chi_{T_r,r}|\phi_r, \psi_r, rot_r, aa_r)$ is computed using tricubic-spline interpolation. To save memory, $\mu_{\chi_k}(\phi_r, \psi_r|rot_r, aa_r)$, and $\sigma_{\chi_k}(\phi_r, \psi_r|rot_r, aa_r)$ are computed using bilinear interpolation, though this has the effect of producing derivative discontinuities at the (ϕ, ψ) grid boundaries. These discontinuities, however, do not appear to produce noticeable artifacts.⁵¹

Terms for Special-Case Torsions. Peptide bond dihedral angles, ω , remain mostly fixed in a cis or trans conformation and depend on the backbone ϕ and ψ angles. Since the electron pair on the backbone nitrogen donates electron density to the electrophilic carbonyl carbon, the peptide bond has partial double-bond character. To model this barrier to rotation, Rosetta implements a backbone-dependent harmonic penalty

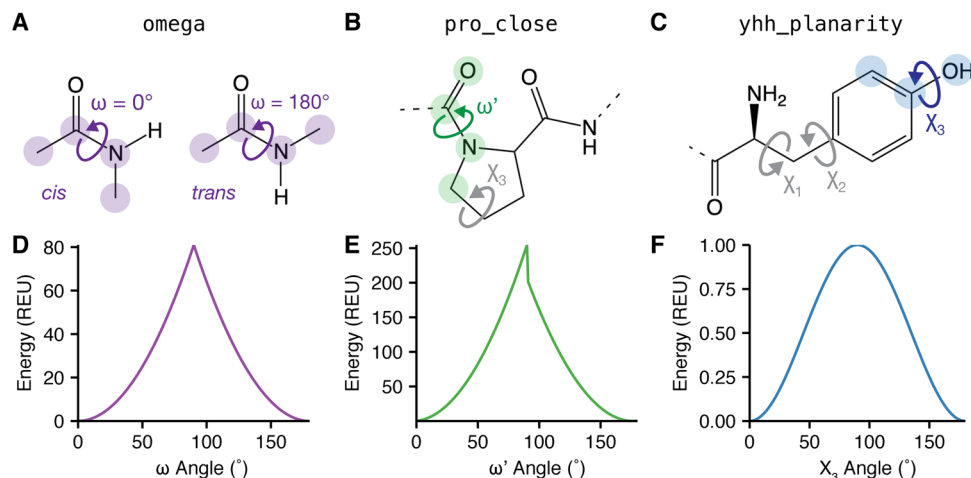


Figure 7. Special case torsion energies. (A–C) Rosetta implements three additional energy terms to model torsional degrees of freedom with acute preferences: (A) omega torsion, corresponding to rotation about C–N; (B) proline secondary omega torsion, corresponding to rotation about C–N related to the C_{δ} in the ring; (C) tyrosine terminal χ torsion. (D) Omega energy. (E) Proline closure energy. (F) Tyrosine planarity energy.

centered near 0° for cis and 180° for trans (Figure 7A). This energy, called omega, is evaluated on all peptide bonds in the biomolecule (eq 30):

$$E_{\text{omega}} = \sum_r \ln\left(\frac{1}{6\sqrt{2\pi}}\right) - \ln\left(\frac{1}{\sigma_{\omega}(\phi_r, \psi_r | \text{aa}_r)\sqrt{2\pi}}\right) + \frac{[\omega_r - \mu_{\omega}(\phi_r, \psi_r | \text{aa}_r)]^2}{2\sigma_{\omega}^2(\phi_r, \psi_r | \text{aa}_r)} \quad (30)$$

The means μ_{ω} and standard derivations σ_{ω} are backbone (ϕ, ψ)-dependent, as given by kernel regressions of ω on ϕ and ψ .⁷²

Most Rosetta protocols search over only simple torsions within chains and rigid-body degrees of freedom between chains. However, the side chain of proline requires special treatment because its ring cannot be represented by a kinematic tree.⁸⁷ Therefore, Rosetta implements a proline closure term, called pro_close (Figure 7B). There are two components to this energy, as shown in eq 31. First, there is a torsional potential that operates on the dihedral formed by $O_{r-1}-C_{r-1}-N_r-C_{\delta,r}$ (called ω'_r) given the observed mean $\mu_{\omega'_r}$ and standard deviation $\sigma_{\omega'_r}$, where r is the residue index. This term keeps the C_{δ} atom in the peptide plane. Second, to ensure the correct geometry for the two hydrogens bound to C_{δ} , we build a virtual nitrogen atom, N_v , off C_{δ} whose coordinate is controlled by χ_3 (Figure 7B). The pro_close term seeks to align the virtual atom N_v directly on top of the real backbone nitrogen. The $N-C_{\delta}-C_{\gamma}$ bond angle and the $N-C_{\delta}$ bond length are restrained to their ideal values.

$$E_{\text{pro_close}} = \sum_{r \in \text{Pro}} \begin{cases} \frac{(\omega'_r - \mu_{\omega'_r})^2}{\sigma_{\omega'_r}^2} & r \text{ is not the N-terminus} \\ + \frac{\|N_r - N_{v,r}\|^2}{\sigma_{N_r, N_{v,r}}^2} & \\ \frac{\|N_r - N_{v,r}\|^2}{\sigma_{N_r, N_{v,r}}^2} & r \text{ is the N-terminus} \end{cases} \quad (31)$$

Tyrosine also requires special treatment for its χ_3 angle because the hydroxyl hydrogen prefers to be in the plane of the aromatic ring.⁸⁸ To enforce this preference, Rosetta implements a sinusoidal penalty to model the barrier to a χ_3 angle that deviates from planarity. This tyrosine hydroxyl penalty is called yhh_planarity (eq 32 and Figure 7C):

$$E_{\text{yhh_planarity}} = \sum_i \frac{1}{2} [\cos(\pi - 2\chi_{3,i}) + 1] \quad (32)$$

Terms for Modeling Nonideal Bond Lengths and Angles. *Cartesian Bonding Energy.* Recently, modeling Cartesian degrees of freedom during gradient-based minimization has been shown to improve Rosetta's ability to refine low-resolution structures determined by X-ray crystallography and cryogenic electron microscopy⁵³ as well as its ability to discriminate near-native conformations in the absence of experimental data.⁸⁹ These data suggest that capturing nonideal bond lengths and angles can be important for accurate modeling of minimum-energy protein conformations. To accommodate, Rosetta now allows these "nonideal" angles and lengths to be included as additional degrees of freedom in refinement and includes a Cartesian minimization mode in which the atom coordinates are explicit degrees of freedom in optimization.

To evaluate the energetics of nonideal bond lengths, angles, and planar groups, an energy term called cart_bonded represents the deviation of these degrees of freedom from ideal using harmonic potentials (eqs 32–34):

$$E_{\text{cart_length}} = \frac{1}{2} \sum_{i=1}^n k_{\text{length},i} (d_i - d_{i,0})^2 \quad (33)$$

$$E_{\text{cart_angle}} = \frac{1}{2} \sum_{i=1}^m k_{\text{angle},i} (\theta_i - \theta_{i,0})^2 \quad (34)$$

$$E_{\text{cart_torsion}} = \frac{1}{2} \sum_{i=1}^l k_{\text{torsion},i} \left[f_{\text{wrap}} \left(\phi_i - \phi_{i,0}, \frac{2\pi}{\rho_i} \right) \right]^2 \quad (35)$$

In these equations, d_i is a bonded-atom-pair distance with $d_{i,0}$ as its ideal distance, θ_i is a bond angle with $\theta_{i,0}$ as its ideal angle, and ϕ_i is a bond torsion or improper torsion with $\phi_{i,0}$ as its

ideal value and ρ_i as its periodicity. The ideal bond lengths and angles^{90,91} were selected on the basis of their ability to rebuild side chains observed in crystal structures (Kevin Karplus and James J. Havranek, unpublished); they were subsequently modified empirically.⁵¹ The spring constants for the angle and length terms are from CHARMM32.¹⁹ Finally, all planar groups and the C _{β} “pseudotorsion” are constrained using empirically derived values and spring constants:

The function $f_{\text{wrap}}(x, y)$ wraps x to the range [0, y). To avoid double counting in the case of $E_{\text{cart_torsion}}$, the spring constant $k_{\text{torsion},i}$ is zero when the torsion ϕ_i is being scored by either the rama or fa_dun terms.

Terms for Protein Design. Design Reference Energy. The terms above are sufficient for comparing different protein conformations with a fixed sequence. However, protein design simulations compare the relative stability of different amino acid sequences given a desired structure to identify models that exhibit a large free energy gap between the folded and unfolded states. Explicit calculations of unfolded-state free energies are computationally expensive and error-prone. Rosetta therefore approximates the relative energies of the unfolded-state ensembles using an unfolded-state reference energy called ref .

Rosetta calculates the reference energy as a sum of individual constant unfolded-state reference energies, ΔG_i^{ref} , for each amino acid, aa _{i} (eq 36).¹

$$E_{\text{ref}} = \sum_i \Delta G_i^{\text{ref}} (\text{aa}_i) \quad (36)$$

The ΔG_i^{ref} values are empirically optimized by searching for values that maximize native sequence recovery (discussed below) during design simulations on a large set of high-resolution crystal structures.^{50,51} During design, this energy term helps normalize the observed frequencies of the different amino acids. When design is turned off, the term contributes a constant offset for a fixed sequence.

Bringing the Energy Terms Together. The Rosetta energy function combines all of the terms using a weighted linear sum to approximate free energies (Table 1). Historically, we have adjusted the weights and parameters to balance the energetic contributions from the various terms. This balance is important because the van der Waals, solvation, and electrostatics energies partially capture torsional preferences and overlap can cause errors as a result of double counting of atomic or residue-specific contributions.⁹² More recently, we have fixed the physics-based terms with weights of 1.0 and perturbed the other weights and atomic-level parameters using a Nelder–Mead scheme⁹³ to optimize the agreement of Rosetta calculations with small-molecule thermodynamic data and high-resolution structural features.⁵⁰ The energy function parameters have evolved over the years by optimization of the performance of multiple scientific benchmarks (Table 2).^{50,51,94} These benchmarks were chosen to test the recovery of native-like structural features, ranging from individual hydrogen-bond geometries to thermodynamic properties and interface conformations. In addition, and more recently, Song et al.,⁹⁵ Conway and DiMaio,⁹⁶ and O’Meara et al.⁴⁹ have fit intraterm parameters to recover features of the experimentally determined folded conformations. An in-depth review of energy function benchmarking can be found in Leaver-Fay et al.⁵¹ Table S3 lists the Rosetta database files containing the current full set of physical parameters for each score term.

Energy Function Units. Initially, Rosetta energies were expressed in a generic unit called the Rosetta energy unit

Table 2. Common Energy Function Benchmarking Methods

test	description	ref(s)
sequence recovery	percentage of the native sequence recovered after backbone redesign	1, 51
rotamer recovery	percentage of native rotamers recovered after full repacking	51
$\Delta\Delta G$ prediction	prediction of free energy changes upon mutation	97
loop modeling	prediction of loop conformations	98
high-resolution refinement	discrimination of native-like decoys upon refinement of ab initio protein models	99
docking	prediction of protein–protein, protein–peptide, or protein–ligand interfaces	44, 100–102
homology modeling	structure prediction incorporating homologous information from templates	103
thermodynamic properties	recapitulation of thermodynamic properties of protein side-chain analogues	17
recapitulation of crystal structure geometries	recapitulation of features (e.g., atom-pair distance distribution) from high-resolution protein crystal structures	50

(REU). This choice was made because some original Rosetta energy terms were not calibrated with experimental data, and the use of statistical potentials convoluted interpretation of the energy. Over time the physical meaning of Rosetta energies has been extensively debated within and outside the community, and several steps have been taken to clarify the interpretation. The most recent energy function (REF15) was parametrized on high-resolution protein structures and small-molecule thermodynamic parameters that were measured in kilocalories per mole.⁵⁰ The optimization data show a strong correlation between the experimental data and values predicted by Rosetta ($\Delta\Delta G$ upon mutation, $R = 0.994$; small-molecule ΔH_{vap} ; Figure S1). As a result, Rosetta energies are now a stronger approximation of energies in units of kilocalories per mole. Therefore, as is standard practice for molecular force fields such as OPLS, CHARMM, and AMBER, we now also express energies in kilocalories per mole.

■ ENERGIES IN ACTION: USING INDIVIDUAL ENERGY TERMS TO ANALYZE ROSETTA MODELS

Rosetta energy terms are mathematical models of the physics that governs protein structure, stability, and association. Therefore, the decomposed relative energies of a structure or ensemble of structures can expose important details about the biomolecular model. Now that we have presented the details of each energy term, we here demonstrate how these energies can be applied to detailed interpretations of structural models. In this section, we discuss two common structure calculations: (1) estimating the free energy change ($\Delta\Delta G$) of mutation⁹⁷ and (2) modeling the structure of a protein–protein interface.¹⁰¹

$\Delta\Delta G$ of Mutation. The first example demonstrates how Rosetta can be used to estimate and rationalize thermodynamic parameters. Here we present an example $\Delta\Delta G$ of mutation calculation for the T193V mutation in the RT-RH-derived peptide bound to HIV-1 protease (PDB entry 1kjj; Figure 8A).¹⁰⁴ The details of this calculation are provided in the Supporting Information.

Rosetta calculates the $\Delta\Delta G$ of the T193V mutation to be -4.95 kcal/mol, and the experimentally measured value is -1.11 kcal/mol.¹⁰⁴ Both the experiment and calculation reveal that T193V is stabilizing, yet these numbers alone do not reveal which specific interactions are responsible for the stabilization. To investigate, we used various analysis tools accessible in PyRosetta¹⁰⁵ to identify important energetic contributions to

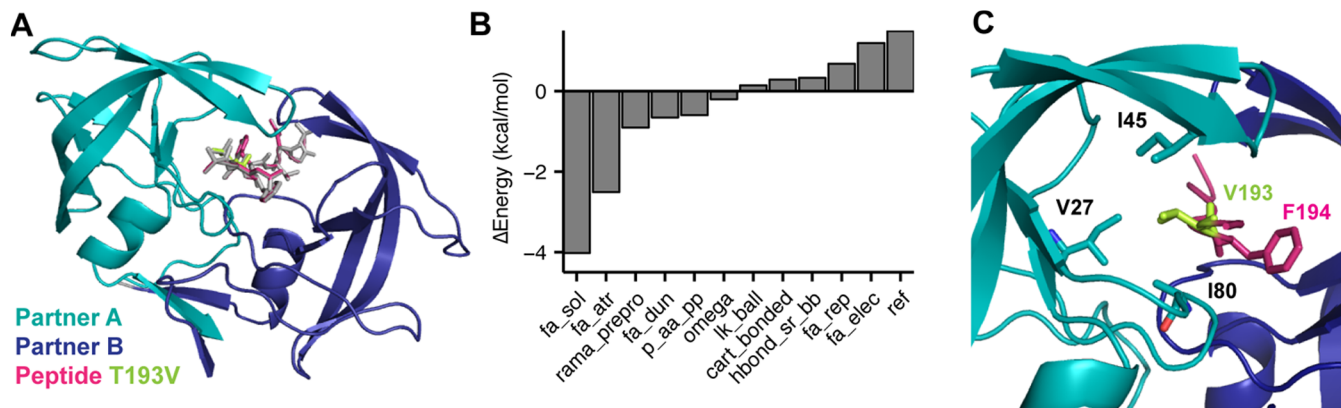


Figure 8. Structural model of the HIV-1 protease bound to the T193V mutant of RT-RH-derived peptide. (A) Structural model of the native HIV-1 peptidase (teal and dark blue) bound to the native peptide (gray) superimposed onto the T193V mutant peptide (magenta). (B) Contributions greater than ± 0.1 kcal/mol to the $\Delta\Delta G$ of mutation for T193V. The remaining contributions are $\text{dslf_fa13} = 0$ kcal/mol, $\text{hbond_lr_bb} = -0.09$ kcal/mol, $\text{hbond_bb_sc} = -0.05$, $\text{hbond_sc} = -0.0104$, $\text{fa_intra_rep} = 0.01$, $\text{fa_intra_sol} = -0.07$, and $\text{yhh_planarity} = 0$. (C) Hydrophobic patch of residues surrounding position 193 on the RT-RH-derived peptide.

the total $\Delta\Delta G$. First, we decomposed the $\Delta\Delta G$ into individual energy terms and observed the balance of terms, both favorable and unfavorable, that sum to the total (Figure 8B). To decompose the most favorable term, $\Delta\text{fa_sol}$, we used the `print_residue_pair_energies` function to identify residues that interact with the mutation site (in this case, residue 4) to produce a nonzero residue-pair solvation energy. With the resulting table, we found that a hydrophobic pocket around the mutation site formed by residues V27, I45, G46, and I80 on HIV peptidase and residue F194 on the peptide made a large (>0.05 kcal/mol) and favorable contribution to the change in solvation energy (Figure 8C).

We further investigated this result on the atomic level with the function `print_atom_pair_energy_table` by generating atom-pair energy tables (see the Supporting Information) for residues 5, 27, 45, 46, and 80 against both threonine and valine at residue 193 (an example for residue 80 is shown in Table 3). Here we find that the specific substitution

neutralizing antibody (PDB entry 1ztx; Figure 9A).¹⁰⁶ Calculation details can be found in the Supporting Information.

To evaluate the docked models, we examine the variation of the energies as a function of the root-mean-square deviation (RMSD) between the residues at the interface in each model and the known structure. For our calculation, interface residues are residues with a C_β atom less than 8.0 Å away from the C_β of a residue in the other docking partner. The plot of energies against RMSD values is called a *funnel plot* and is intended to mimic the funnel-like energy landscape of protein folding and binding.

As in the previous example, we decompose the energies to yield information about the nature of interactions at the interface. Here we observe significant changes in the following energy terms upon interface formation relative to the unbound state: fa_atr , fa_rep , fa_sol , lk_ball_wtd , fa_elec , hbond_lr_bb , hbond_bb_sc , and hbond_sc (Figure 9B). The change in the Lennard-Jones energy upon interface formation is due to the introduction of atom–atom contacts at the interface. As more atoms come into contact near the native conformation ($\text{RMSD} \rightarrow 0$), the favorable attractive energy (fa_atr) decreases whereas the unfavorable repulsive energy (fa_rep) increases. The change in the isotropic solvation energy ($\Delta\text{fa_sol}$) is positive (unfavorable), indicating that polar residues are buried upon interface formation. Balancing the desolvation penalty, the changes in polar solvation energy ($\Delta\text{lk_ball_wtd}$) and electrostatics ($\Delta\text{fa_elec}$) are negative because of the formation of polar contacts at the interface. Finally, the changes in the three hydrogen-bonding energies ($\Delta\text{hbond_lr_bb}$, $\Delta\text{hbond_bb_sc}$, and $\Delta\text{hbond_sc}$) reflect the formation of backbone–backbone, backbone–side-chain, and side-chain–side-chain hydrogen bonds at the interface.

■ DISCUSSION

The Rosetta energy function represents our collaboration's ongoing pursuit to model the rules in nature that govern biomolecular structure, stability, and association. This paper summarizes the latest version, which brings together fundamental physical theories, statistical-mechanical models, and observations of protein structures. This work represents almost 20 years of interdisciplinary collaboration in the Rosetta

Table 3. Changes in Atom-Pair Energies (in kcal/mol) between I80 and T193 versus V193

T193→V193 atoms	I80 atoms			
	CB	CG1	CG2	CD1
N	0.000	0.000	0.000	0.000
CA	0.000	0.000	0.000	0.004
C	0.000	0.000	0.000	0.008
O	0.000	0.000	0.000	-0.010
CB	0.000	0.054	0.000	-0.002
OG1 → CG1	0.008	-0.054	-0.316	-0.398
CG2 → CG2'	0.000	0.000	0.001	0.020

of the polar hydroxyl group on threonine with the nonpolar alkyl group on valine stabilizes the peptide in the hydrophobic protease pocket. This result is consistent with chemical intuition and demonstrates how breaking down the total energies can provide insight into characteristics of the mutated structures.

Protein–Protein Docking. The second example shows how the Rosetta energies of an ensemble of models can be used to discriminate between models and investigate the characteristics of a protein–protein interface. Below we investigate docked models of West Nile Virus envelope protein and a

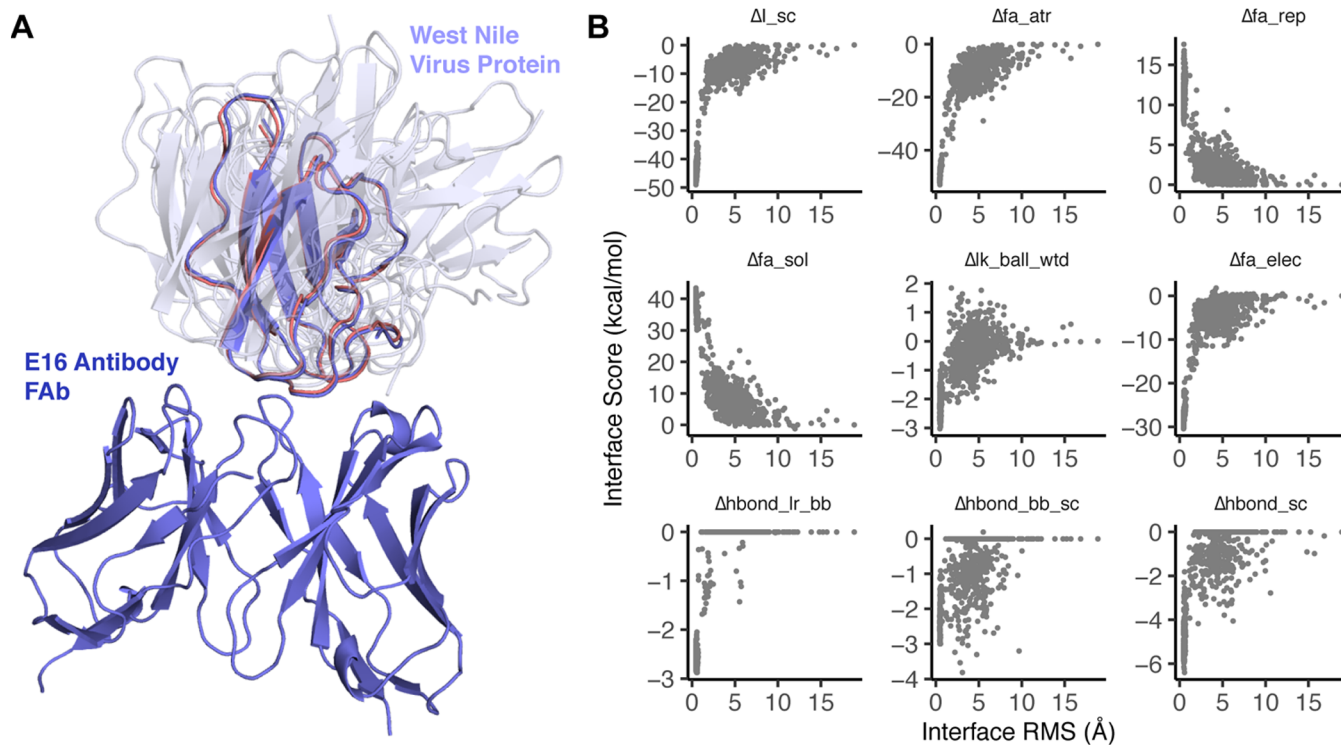


Figure 9. Using energies to discriminate docked models of West Nile Virus and the E16 neutralizing antibody. (A) Comparison of the native E16 antibody (purple) docked to the lowest-RMSD model of the West Nile Virus envelope protein and several other random models of varying energy to show sampling diversity (gray, semitransparent). (B) Change in the interface energy relative to the unbound state vs RMSD relative to the native structure. Models with low RMSD relative to the native interface have a low overall interface energy due to favorable van der Waals contacts, electrostatic interactions, and side-chain hydrogen bonds, as reflected by the Δf_{α_atr} , Δf_{α_elec} , and Δh_{bond_sc} energy terms.

Table 4. New Energy Terms for Biomolecules Other than Proteins

biomolecule	term	description	unit	ref
noncanonical amino acids	mm_lj_intra_rep	repulsive van der Waals energy between two atoms from the same residue	kcal/mol	67
	mm_lj_intra_atr	attractive van der Waals energy between two atoms from the same residue	kcal/mol	67
	mm_twist	molecular mechanics derived torsion term for all proper torsions	kcal/mol	67
	unfolded	energy of the unfolded state based on explicit unfolded state model	AU ^a	67
	split_unfolded_1b	one-body component of the two-component reference energy, lowest energy of a side chain in a dipeptide model system	AU	in the SI
	split_unfolded_2b	two-body component of the two-component reference energy, median two-body interaction energy based on atom-type composition	AU	in the SI
carbohydrates	sugar_bb	energy for glycosidic torsions	kcal/mol	70
DNA	gb_elec	generalized Born model of the electrostatics energy	kcal/mol	107
RNA	fa_stack	$\pi-\pi$ stacking energy for RNA bases	kT	113
	stack_elec	electrostatic energy for stacked RNA bases	kT	114
	fa_elec_rna_phos	electrostatic energy (fa_elec) between RNA phosphate atoms	kT	62
	rna_torsion	knowledge-based torsional potential for RNA	kT	62
	rna_sugar_close	penalty for opening an RNA sugar	kT	62

^aAU = arbitrary units.

community, which in turn builds on and incorporates decades of work outside the community.

After 20 years, we have improved physical theories, structural data, representations, experiments, and computational tools; nevertheless, the energy functions are far from perfect. Compared with the first torsional potentials, the energy functions are also now vastly more complex. There are countless ways to arrive at more accurate energy functions. Here we discuss grand challenges specific to development of the Rosetta energy function in the coming decade.

Modeling of Biomolecules Other than Proteins. The Rosetta energy function was originally developed to predict and

design protein structures. A clear artifact of this goal is the energy function's dependence on statistical potentials derived from protein X-ray crystal structures. Today the Rosetta community also pursues goals ranging from the design of synthetic macromolecules to the prediction of interactions and structures of other biomolecules such as glycoproteins and RNA. Accordingly, an active research thrust is to generalize the all-atom energy function for all biomolecules.

Many of the physically derived terms (e.g., van der Waals) have already been made compatible with noncanonical amino acids and nonprotein biomolecules (Table S5). Recently, Bhardwaj, Mulligan, Bahl, and co-workers⁶⁹ adapted the

Table 5. Energy Terms for Structure Prediction in Different Contexts

context	term	description	unit	ref(s)
membrane environment	fa_mpsolv	solvation energy dependent on the protein orientation relative to the membrane	kcal/mol	118, 121
	fa_mpenv	one-body membrane environment energy dependent on the protein orientation relative to the membrane	kcal/mol	118, 121
pH	e_pH	likelihood of side-chain protonation given a user-specified pH	kcal/mol	117

rama_prep, p_aa_pp, fa_dun, pro_close, omega, dslf_fa13, yhh_planarity, and ref terms to be compatible with mixed-chirality peptides. Several of Rosetta's statistical potentials have been validated against quantum-mechanical calculations to evaluate non-protein models (Table 4). Early work by Meiler and Baker⁴⁴ on Rosetta Ligand introduced new atom and residue types for non-protein residues. The first non-protein energy terms were added by Havranek et al.¹⁰⁷ and Chen et al.,¹⁰⁸ who modified the hydrogen-bonding potential to capture planar hydrogen bonds between protein side chains and nucleic acid bases. Renfrew and co-workers^{67,109} added molecular mechanics torsions and Lennard-Jones terms to model proteins with noncanonical amino acids, oligosaccharides, β -peptides, and oligopeptoids.⁶⁸ Labonte et al.⁷⁰ implemented Woods' CarboHydrate-Intrinsic (CHI) function,^{110,111} which evaluates glycan geometries given the axial-equatorial character of the bonds. Das and co-workers added a set of terms to model Watson-Crick base pairing, $\pi-\pi$ interactions in base stacking, and torsional potentials important for predicting and designing RNA structures.^{62,112-114} Bazzoli and Karanicolas¹¹⁵ recently developed a new polar solvation model that evaluates the penalty associated with displacing waters in the first solvation shell. In addition, Combs¹¹⁶ tested a small-molecule force field based on electron orbital models. Many of these terms are presented in detail in the Supporting Information.

Expanding Rosetta's chemical library brings new challenges. Currently there are separate energy functions for various types of biomolecules. Typically, these functions mix physically derived terms from the protein energy function with molecule-specific statistical potentials, custom weights, and possibly custom atomic parameters. If nature uses only one energy function, why do we need so many? Some discrepancies may result from features that we do not model explicitly, such as $\pi-\pi$, $n-\pi^*$, and cation- π interactions. Efforts to converge on a single energy function will therefore pose interesting questions about the set of universal physical determinants of biomolecular structure.

Capturing the Intra- and Extracellular Environments.

Rosetta traditionally models the solvent surrounding the protein using the Lazaridis-Karplus (LK) model, which assumes a solvent environment made of pure water. In contrast, biology operates under various conditions influenced by pH, redox potential, temperature, solvent viscosity, chaotropes, kosmotropes, and polarizability. Therefore, modeling more details of the intra- and extracellular environments would enable Rosetta to identify structures that are important in different biological contexts.

Rosetta currently includes two groups of energy terms to model alternate environments (Table 5). Kilambi and Gray¹¹⁷ implemented a method to account for pH by including a term called e_pH that calculates the likelihood of a protein side chain's protonation state given a user-specified pH; it requires the inclusion of both protonated and deprotonated side chains during side-chain rotamer packing. This model can predict pK_a

values with an RMS error under 1 unit,¹¹⁷ and it improves protein-protein docking, especially under acidic or basic conditions.⁶⁰ The accuracy of this model is limited by the distance-dependent Coulomb approximation and sensitivity to fine backbone rearrangements.

In addition, Rosetta implements Lazaridis' implicit membrane model (IMM) for modeling proteins in a lipid bilayer environment.^{36,118,119} The IMM terms provide a fast approximation of the nonpolar hydrocarbon core of the lipid bilayer and have been successfully applied to membrane protein folding,¹²⁰ docking, and early design tasks.⁶¹ This continuum model has a fixed thickness, omitting the detailed chemistry at the membrane interface and any dynamic bilayer rearrangements.

The Origin of Energy Models: Top-Down versus Bottom-Up Development. Traditionally, energy functions are developed using a bottom-up approach: experimental observables serve as building blocks to parametrize physics-based formulas. The advent of powerful optimization techniques and artificial intelligence has recently empowered the top-down category, where numerical methods are used to derive models and/or parameters. Top-down approaches have been used to solve problems in various fields, including structural biology and bioinformatics. Recently, top-down development was also applied to optimize the Lennard-Jones, LK, and Coulomb parameters in the Rosetta energy function (Tables S4-S6).^{50,93}

Top-down approaches have enormous potential to improve the accuracy of biomolecular modeling because more parameters can vary and the objective function can be minimized with more benchmarks. These approaches also introduce new challenges. With any computer-derived model there is a risk of overfitting, as validation via structure-prediction data sets reflect observable states, whereas simulations are intended to predict features of states that experiments cannot yet observe. Computer-derived parameters also introduce a unique kind of uncertainty. Consider the following scenario: the performance of scientific benchmarks improves as physical atomic parameters are perturbed away from the measured experimental values. As there is less physical basis for the parameters, are the predictions and interpretations still meaningful?

Top-down development will also provide power to develop more complicated energy functions. Currently, the Rosetta energy function advances by incrementally addressing weaknesses: with each new paper, we modify analytic formulas, add corrective terms, and adjust weights. As this paper demonstrates, the energy function is significantly more complicated than the initial theoretical forms. Given this complexity increase, an interesting approach to leverage the power of top-down development would be to simplify and subtract terms to evaluate their individual benefits.

A Highly Interdisciplinary Endeavor. The Rosetta energy function has advanced rapidly because of the Rosetta Community, a highly interdisciplinary collaboration among

scientists with diverse backgrounds located in over 50 laboratories around the world. The many facets of our team enable us to probe different aspects of the energy function. For example, expert computer scientists and applied mathematicians have implemented algorithms to speed up calculations. Dedicated software engineers maintain the code and maintain a platform for scientific benchmark testing. Physicists and chemists develop new energy terms that better model the physical rules found in nature. Structural biologists maintain a focus on created biological features and functions. We look forward to leveraging this powerful interdisciplinary scientific team as we head into the next decade of energy function advances.

CONCLUSION: A LIVING ENERGY FUNCTION

For the first time since 2004,⁴⁸ we have documented all of the mathematical and physical details of the Rosetta all-atom energy function, highlighting the latest upgrades to both the underlying science and the speed of calculations. In addition, we have illustrated how the energies can be used to analyze output models from Rosetta simulations. These advances have enabled Rosetta's achievements in biomolecular structure prediction and design over the past 15 years. Still, the energy function is far from complete and will continue to evolve long after this publication. Thus, we hope that this document will serve as an important resource for understanding the foundational physical and mathematical concepts in the energy function. Furthermore, we hope to encourage both current and future Rosetta developers and users to understand the strengths and shortcomings of the energy function as it applies to the scientific questions they are trying to answer.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: [10.1021/acs.jctc.7b00125](https://doi.org/10.1021/acs.jctc.7b00125).

Description of changes to the Rosetta energy function since 2000; data describing the calibration of Rosetta energies in kcal/mol; additional details of energy terms and details on smoothing of statistical terms; energy terms for D-amino acids, noncanonical amino acids, carbohydrates, and nucleic acids; and methods describing example energy calculations ([PDF](#))

Protocol capture within an interactive Python notebook demonstrating the usage of the `print_atom_pair_energy_table` function ([ZIP](#))

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V.K.M., J.W.L., R.B., and R.D. wrote, verified, and/or contributed figures on non-protein energy terms.

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Notes

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