

Rosetta Scoring (or Energy) Function

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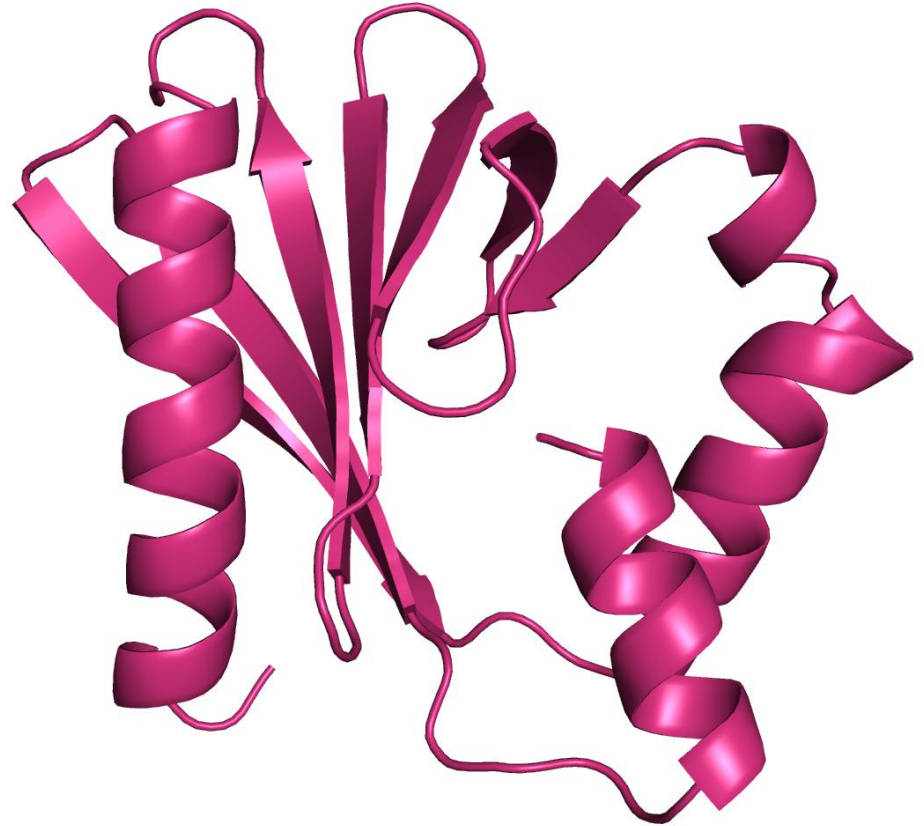
Rosetta Workshop

Nov 2018

Rosetta Scoring (or Energy) Function

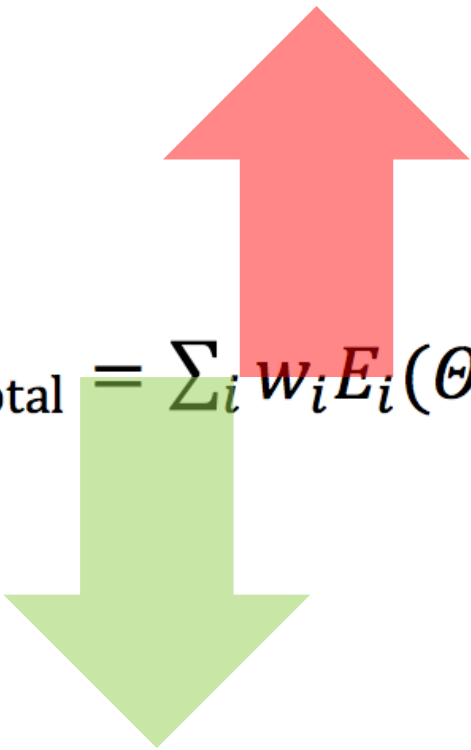
... or how models are evaluated in
Rosetta

Evaluating Models in Rosetta



An algorithm must assign a quantitative number to a model to decide whether it is “good” or “bad”

Important Note

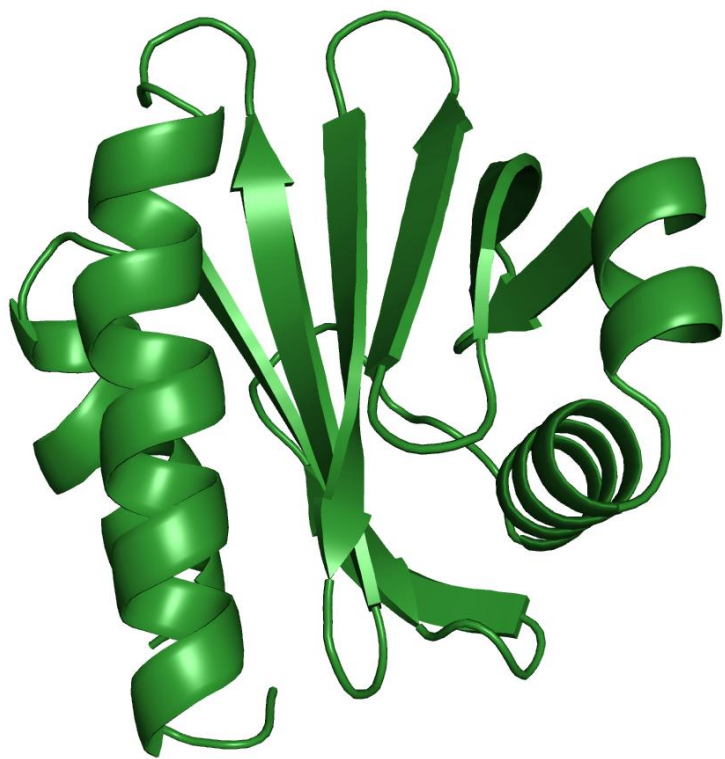

$$\Delta E_{\text{total}} = \sum_i w_i E_i(\theta_i, \text{aa}_i)$$

Energy is currently given in:
“kcal/mol”

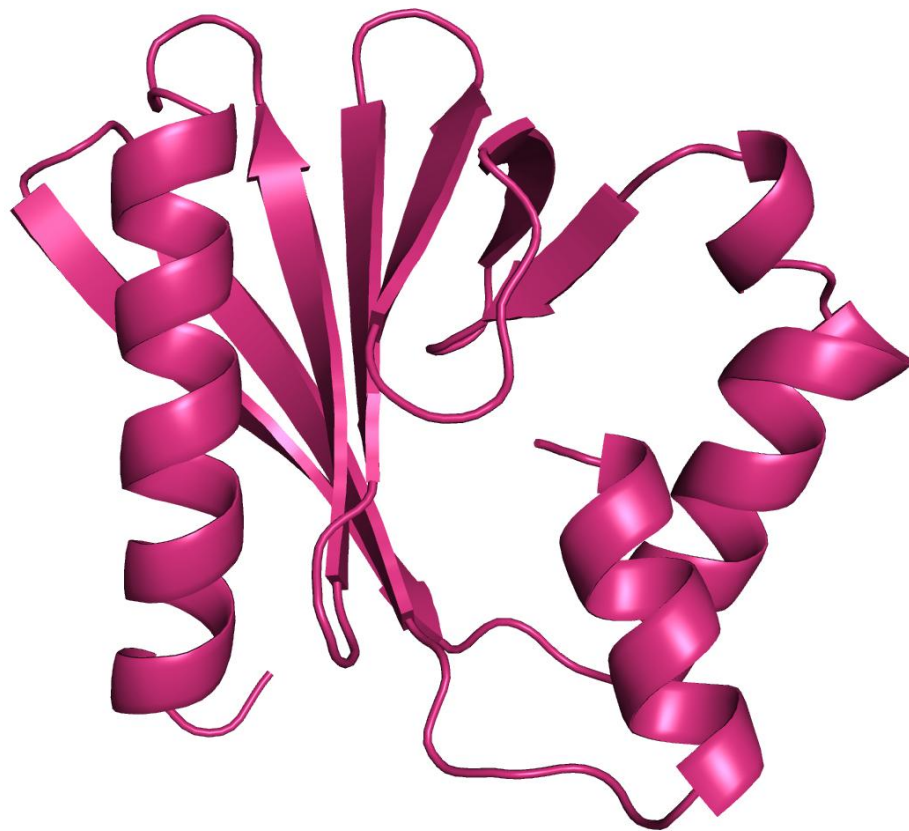
Previously in: “REU”

High Energy = Bad
Low Energy = Good

Evaluating Models in Rosetta

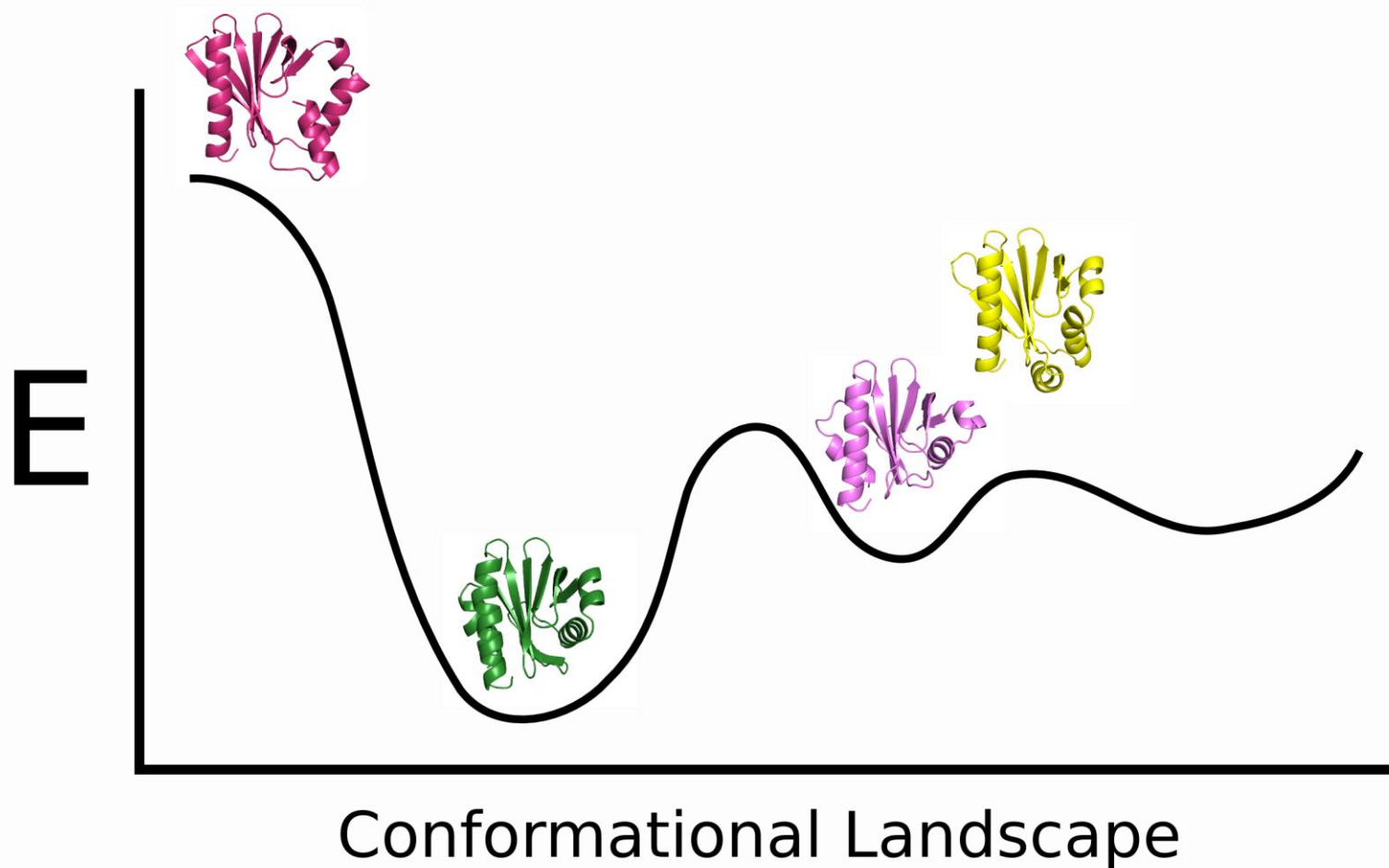


-400 kcals/mol



-350 kcals/mol

Score is Central to Monte Carlo Selection

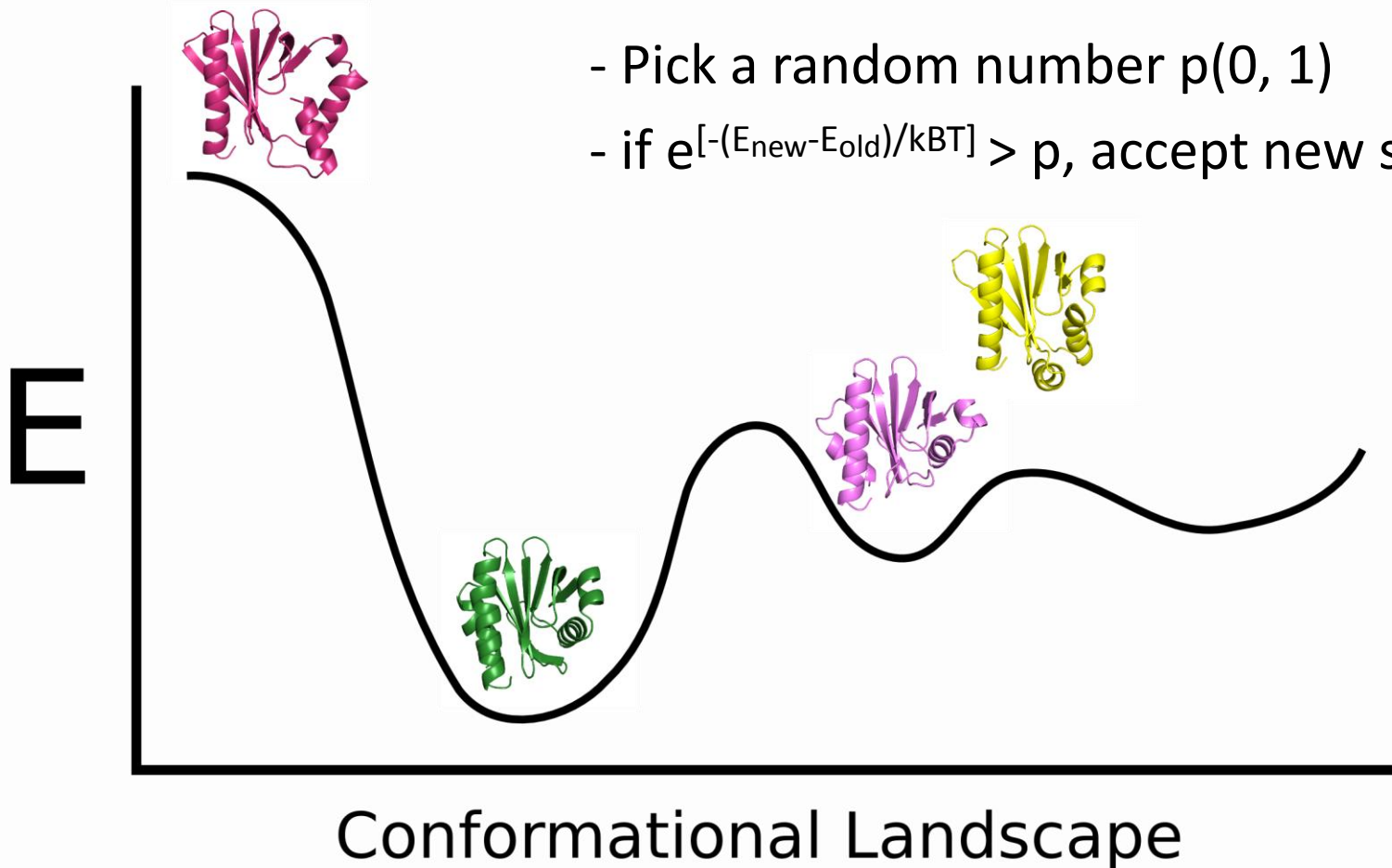


Metropolis Criterion

If $E_{\text{new}} < E_{\text{old}}$: Accept new structure

If $E_{\text{new}} > E_{\text{old}}$:

- Pick a random number $p(0, 1)$
- if $e^{[-(E_{\text{new}} - E_{\text{old}})/kBT]} > p$, accept new structure



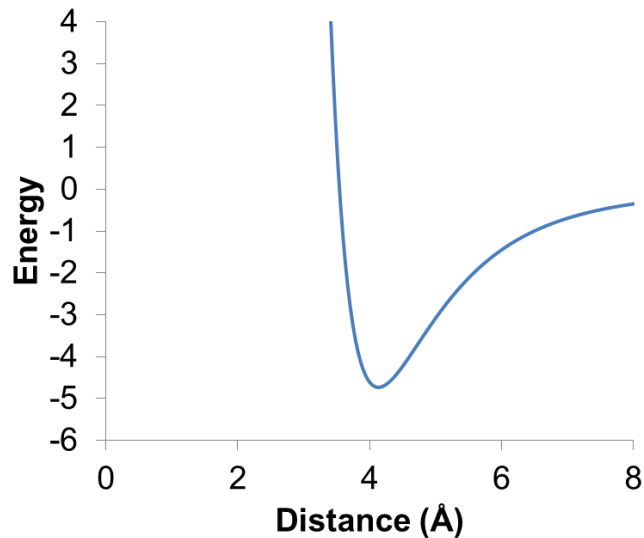
Rosetta Combines Physics-Based and Knowledge-Based Potentials to Build the Energy Function

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Lennard-Jones Potential

$$\sum_{i < j} \left[\frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^6} \right]$$

van der Waals Energy



Rosetta Combines Physics-Based and Knowledge-Based Potentials to Build the Energy Function

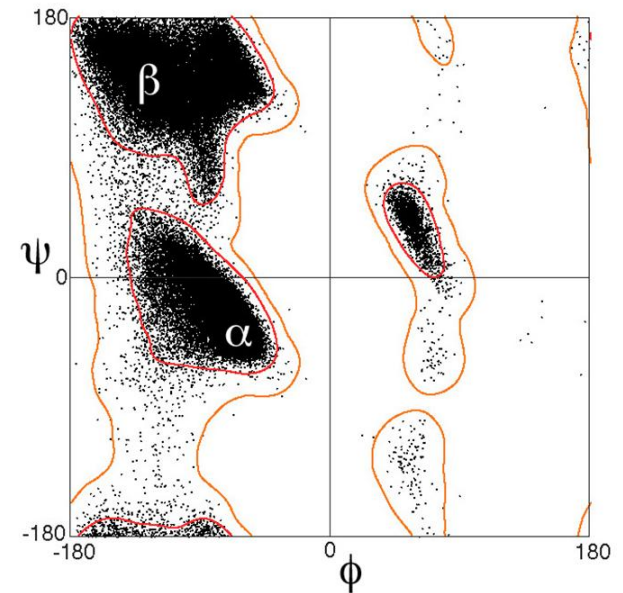
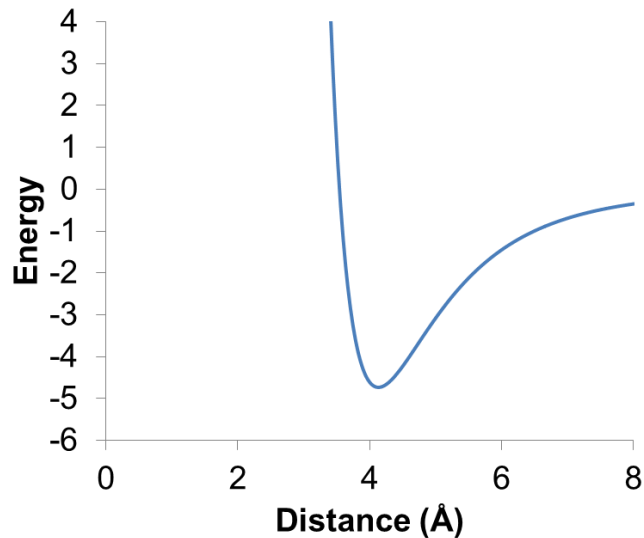
Lennard-Jones Potential

$$\sum_{i < j} \left[\frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^6} \right]$$

Ramachandran Plot

Statistical mining of
Protein Databank (PDB)

van der Waals Energy



The Score Function is a Weighted Linear Combination of Individual Score Terms

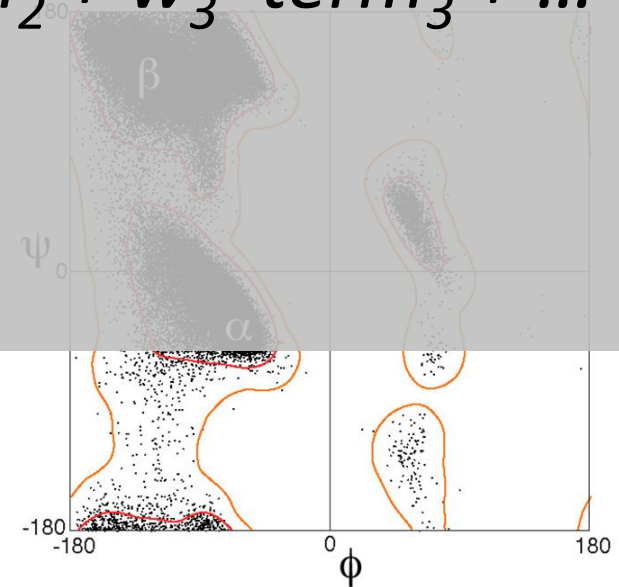
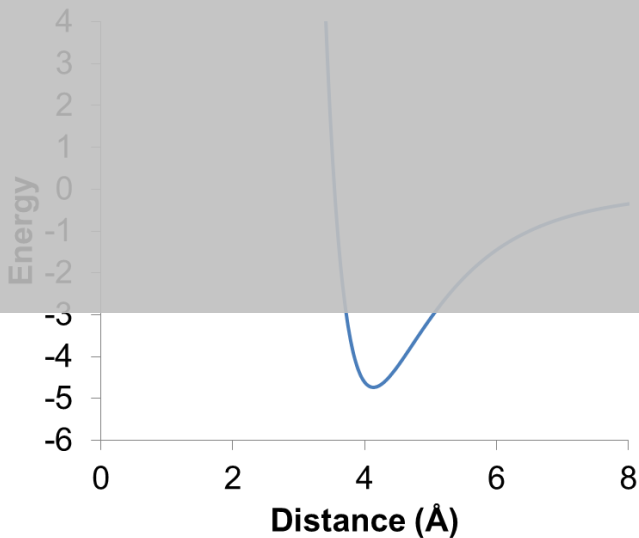
Lennard-Jones Potential

$$\sum_{i < j} \left[\frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^6} \right]$$


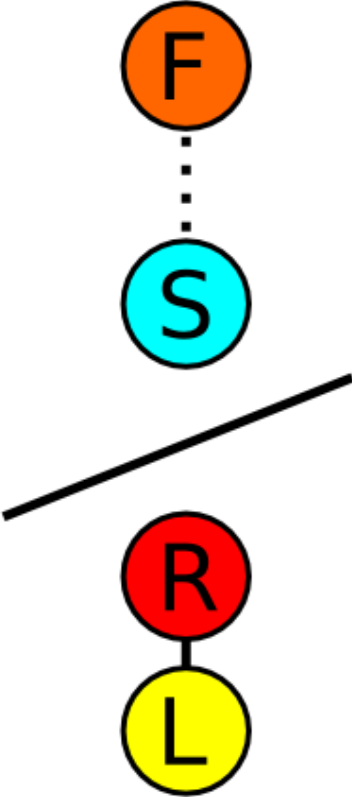
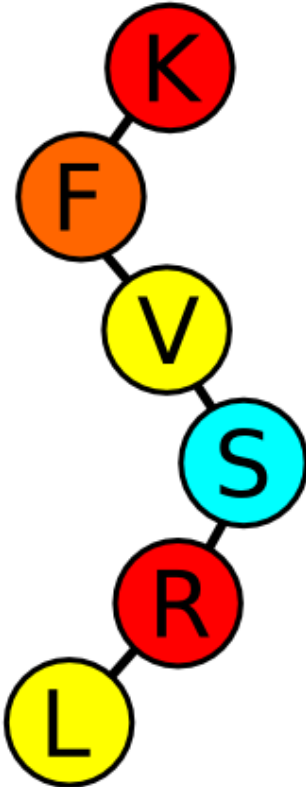
Ramachandran Plot

Statistical mining of
Protein Databank (PDB)

$$Energy = w_1 * term_1 + w_2 * term_2 + w_3 * term_3 + \dots$$



Rosetta is a Residue-Centric Scoring Function

One Body	Two Body	Whole Body
		

Rosetta is a Residue-Centric Scoring Function

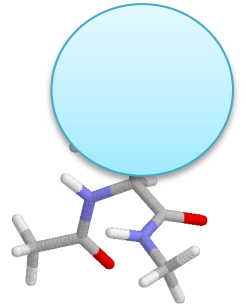
One Body	Two Body	Whole Body
Backbone - p_aa_pp - rama_prepro	Lennard-Jones - fa_atr - fa_rep	Radius of Gyration - rg
Side Chain - fa_dun - yhh_planarity	Solvation - fa_sol Hydrogen Bond - hbond_lr_bb - hbond_sr_bb - hbond_bb_sc - hbond_sc	Contact Order - co Structure Alignment - hs_pair - ss_pair - sheet
Reference - ref		

* NOTE: Not all score terms are listed here

Rosetta has 2 score function modes

Low resolution (or centroid) mode:

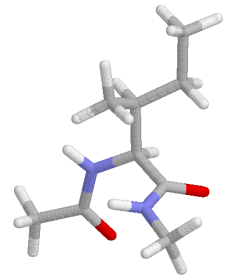
Reduced atom representation (centroid)
Simple energy function
Aggressively search conformational space



[database/chemical/residue_type_sets/centroid](#)

High resolution (or full atom) mode:

Full atom (FA)
More sophisticated energy function
“Local” search of conformational (and sequence) space



[database/chemical/residue_type_sets/fa_standard](#)

Breakdown of Full Atom Score Terms – REF2015

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_prepro	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.

What you'll actually see in your
output...

Output Score Table

Found in output score table (score.sc) and at the end of every output pdb (S_0001.pdb)

```
# All scores below are weighted scores, not raw scores.
#BEGIN_POSE_ENERGIES_TABLE dock_01_girk_1212_jon_ML297_0500.pdb
label fa_atr fa_rep fa_sol fa_intra_rep fa_elec total
weights 0.8 0.4 0.6 0.004 0.42 NA
pose -5400.73 638.748 2428.12 13.5145 -302.036 -4267.34
GLN:NtermProteinFull_1 -1.02843 0.12113 0.83426 0.00595 -0.05882 0.54065
ARG_2 -5.46203 0.44148 4.02307 0.01303 -1.13134 -3.04715
PHE_3 -5.25196 0.29047 1.51945 0.02354 -0.38757 -1.99927
VAL_4 -4.7212 0.38219 2.30519 0.01101 -0.33608 -4.69107
ASP_5 -6.18239 0.68119 4.3013 0.02667 -1.21805 -4.77894
LYS_6 -3.70129 0.51352 2.90682 0.00832 -0.90685 -3.25917
ASN_7 -2.97985 0.42136 1.9979 0.00362 -0.17245 -2.08876
GLY_8 -2.75353 0.28458 1.44949 1e-05 -0.20107 -4.00172
ARG:CtermProteinFull_1294 -2.34298 0.24806 1.73197 0.01216 -0.06632 -0.34395
```

Output Score Table

Score Terms:

All scores below are weighted scores, not raw scores.

#BEGIN POSE ENERGIES TABLE dock_01_girk_1212_jon_ML297_0500.pdb

label	fa	atr	fa	rep	fa	sol	fa	intra	rep	fa	elec	total
-------	----	-----	----	-----	----	-----	----	-------	-----	----	------	-------

weights	0.8	0.4	0.6	0.004	0.42	NA						
---------	-----	-----	-----	-------	------	----	--	--	--	--	--	--

pose	-5400.73	638.748	2428.12	13.5145	-302.036	-4267.34						
------	----------	---------	---------	---------	----------	----------	--	--	--	--	--	--

GLN:NtermProteinFull_1	-1.02843	0.12113	0.83426	0.00595	-0.05882	0.54065						
------------------------	----------	---------	---------	---------	----------	---------	--	--	--	--	--	--

ARG_2	-5.46203	0.44148	4.02307	0.01303	-1.13134	-3.04715						
-------	----------	---------	---------	---------	----------	----------	--	--	--	--	--	--

PHE_3	-5.25196	0.29047	1.51945	0.02354	-0.38757	-1.99927						
-------	----------	---------	---------	---------	----------	----------	--	--	--	--	--	--

VAL_4	-4.7212	0.38219	2.30519	0.01101	-0.33608	-4.69107						
-------	---------	---------	---------	---------	----------	----------	--	--	--	--	--	--

ASP_5	-6.18239	0.68119	4.3013	0.02667	-1.21805	-4.77894						
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LYS_6	-3.70129	0.51352	2.90682	0.00832	-0.90685	-3.25917						
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ASN_7	-2.97985	0.42136	1.9979	0.00362	-0.17245	-2.08876						
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GLY_8	-2.75353	0.28458	1.44949	1e-05	-0.20107	-4.00172						
-------	----------	---------	---------	-------	----------	----------	--	--	--	--	--	--

ARG:CtermProteinFull_1294	-2.34298	0.24806	1.73197	0.01216	-0.06632	-0.34395						
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Output Score Table

Weights:

```
# All scores below are weighted scores, not raw scores.
#BEGIN POSE ENERGIES TABLE dock_01_girk_1212_jon_ML297_0500.pdb
label fa_atr fa_rep fa_sol fa_intra_rep fa_elec total
weights 0.8 0.4 0.6 0.004 0.42 NA
pose -5400.73 638.748 2428.12 13.5145 -302.036 -4267.34
GLN:NtermProteinFull_1 -1.02843 0.12113 0.83426 0.00595 -0.05882 0.54065
ARG_2 -5.46203 0.44148 4.02307 0.01303 -1.13134 -3.04715
PHE_3 -5.25196 0.29047 1.51945 0.02354 -0.38757 -1.99927
VAL_4 -4.7212 0.38219 2.30519 0.01101 -0.33608 -4.69107
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GLY_8 -2.75353 0.28458 1.44949 1e-05 -0.20107 -4.00172
ARG:CtermProteinFull_1294 -2.34298 0.24806 1.73197 0.01216 -0.06632 -0.34395
```

Output Score Table

Scores for individual terms:

```
# All scores below are weighted scores, not raw scores.
#BEGIN POSE ENERGIES TABLE dock_01_girk_1212_jon_ML297_0500.pdb
label fa atr fa rep fa sol fa intra rep fa elec total
weights 0.8 0.4 0.6 0.004 0.42 NA
pose -5400.73 638.748 2428.12 13.5145 -302.036 -4267.34
GLN:NtermProteinFull_1 -1.02843 0.12113 0.83426 0.00395 -0.05882 0.54065
ARG_2 -5.46203 0.44148 4.02307 0.01303 -1.10204 0.00000
PHE_3 -5.25196 0.29047 1.51945 0.02354 -0.38757 -1.99927
VAL_4 -4.7212 0.38219 2.30519 0.01101 -0.33608 -4.69107
ASP_5 -6.18239 0.68119 4.3013 0.02667 -1.21805 -4.77894
LYS_6 -3.70129 0.51352 2.90682 0.00832 -0.90685 -3.25917
ASN_7 -2.97985 0.42136 1.9979 0.00362 -0.17245 -2.08876
GLY_8 -2.75353 0.28458 1.44949 1e-05 -0.20107 -4.00172
ARG:CtermProteinFull_1294 -2.34298 0.24806 1.73197 0.01216 -0.06632 -0.34395
```

Total Score of Protein

Output Score Table

Scores by amino acid position:

```
# All scores below are weighted scores, not raw scores.
#BEGIN POSE ENERGIES TABLE dock_01_girk_1212_jon_ML297_0500.pdb
label fa atr fa rep fa sol fa intra rep fa elec total
weights 0.8 0.4 0.6 0.004 0.42 NA
pose -5400.73 638.748 2428.12 13.5145 -302.036 -4267.34
GLN:NtermProteinFull_1 -1.02843 0.12113 0.83426 0.00595 -0.05882 0.54065
ARG_2 -5.46203 0.44148 4.02307 0.01303 -1.13134 -3.04715
PHE_3 -5.25196 0.29047 1.51945 0.02354 -0.38757 -1.99927
VAL_4 -4.7212 0.38219 2.30519 0.01101 -0.33608 -4.69107
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LYS_6 -3.70129 0.51352 2.90682 0.00832 -0.90685 -3.25917
ASN_7 -2.97985 0.42136 1.9979 0.00362 -0.17245 -2.08876
GLY_8 -2.75353 0.28458 1.44949 1e-05 -0.20107 -4.00172
ARG:CtermProteinFull_1294 -2.34298 0.24806 1.73197 0.01216 -0.06632 -0.34395
```

More score term information

Constraints (actually “restraints”)

Supplements energy function with additional information

- Commonly from experimental information

Various types:

atom_pair_constraint, dihedral_constraint,
angle_constraint, coordinate_constraint,
residue_type_constraint ...

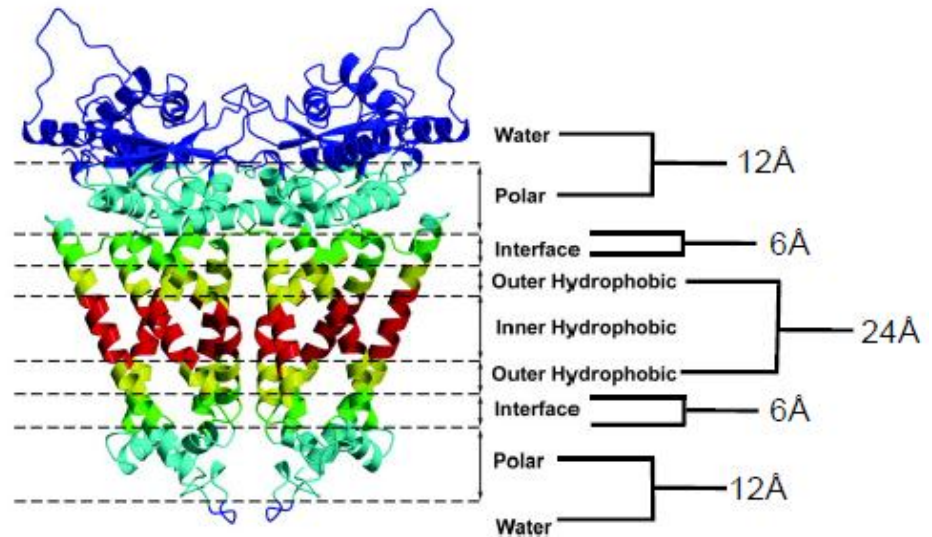
Interface with Rosetta using constraint files

Additional score terms

Membrane terms:

fa_mbsolv

fa_mbenv



Yarov-Yarovoy, Schonbrun, and Baker 2006

Over 100+ score terms

Most are turned off (weight is set to 0), not in REF2015

To turn on score term, set weight to non-zero value

Modifying Scorefunctions in Rosetta

Modifying using the command line

1 - score:weights <filename>

2 - score:set_weight <scoreterm₁> <wt₁> <scoreterm₂> <wt₂>

3 - score:patch <patchfile>

Patchfile example

```
fa_atr = 0.423  
fa_rep = 0.100
```

Modifying Scorefunctions in Rosetta

```
<ROSETTASCRIPTS>
  <SCOREFXNS>
    <ScoreFunction name="ligand_soft_rep" weights="ligand_soft_rep">
      <Reweight scoretype="fa_elec" weight="0.42"/>
    </ScoreFunction>
    <ScoreFunction name="hard_rep" weights="ligandprime">
    </ScoreFunction>
  </SCOREFXNS>
  <OUTPUT scorefxn="hard_rep" />
</ROSETTASCRIPTS>
```

XML script options

- Weights filename or path to file
- Reweight specific terms as needed (as in patch file)
- Must include top-level output tag to ensure proper scoring in output files

References:

REF2015 References:

Alford RF, et. al, **The Rosetta All-Atom Energy Function for Macromolecular Modeling and Design.** *Journal of Chemical Theory and Computation*, **2017**. 13 (6), 3031-3048

Park H, et. al **Simultaneous Optimization of Biomolecular Energy Functions on Features from Small Molecules and Macromolecules** *Journal of Chemical Theory and Computation*, **2016**. 12 (12), 6201-6212

Old Scorefxn (Talaris) References:

O'Meara MJ, et. al, **A Combined Covalent-Electrostatic Model of Hydrogen Bonding Improves Structure Prediction with Rosetta.** *Journal of Chemical Theory and Computation*, **2015**.

Leaver-Fay A, et. al **Scientific benchmarks for guiding macromolecular energy function improvement.** *Methods in enzymology*, **2013**. 523: p. 109.

Useful Links

https://www.rosettacommons.org/docs/latest/rosetta_basics/scoring/score-types

https://www.rosettacommons.org/docs/latest/rosetta_basics/scoring/scoring-explained