

Research article

Open Access

SELECTpro: effective protein model selection using a structure-based energy function resistant to BLUNDERS

Arlo Randall^{1,2} and Pierre Baldi*^{1,2}

Address: ¹School of Information and Computer Sciences, University of California, Irvine, CA, 92697, USA and ²Institute for Genomics and Bioinformatics, University of California, Irvine, CA, 92697, USA

E-mail: Arlo Randall - arandall@ics.uci.edu; Pierre Baldi* - pfbaldi@ics.uci.edu;

*Corresponding author

Published: 03 December 2008

Received: 26 June 2008

BMC Structural Biology 2008, 8:52 doi: 10.1186/1472-6807-8-52

Accepted: 3 December 2008

This article is available from: <http://www.biomedcentral.com/1472-6807/8/52>

© 2008 Randall and Baldi; licensee BioMed Central Ltd.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<http://creativecommons.org/licenses/by/2.0>), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

Background: Protein tertiary structure prediction is a fundamental problem in computational biology and identifying the most native-like model from a set of predicted models is a key sub-problem. Consensus methods work well when the redundant models in the set are the most native-like, but fail when the most native-like model is unique. In contrast, structure-based methods score models independently and can be applied to model sets of any size and redundancy level. Additionally, structure-based methods have a variety of important applications including analogous fold recognition, refinement of sequence-structure alignments, and de novo prediction. The purpose of this work was to develop a structure-based model selection method based on predicted structural features that could be applied successfully to any set of models.

Results: Here we introduce SELECTpro, a novel structure-based model selection method derived from an energy function comprising physical, statistical, and predicted structural terms. Novel and unique energy terms include predicted secondary structure, predicted solvent accessibility, predicted contact map, β -strand pairing, and side-chain hydrogen bonding.

SELECTpro participated in the new model quality assessment (QA) category in CASP7, submitting predictions for all 95 targets and achieved top results. The average difference in GDT-TS between models ranked first by SELECTpro and the most native-like model was 5.07. This GDT-TS difference was less than 1% of the GDT-TS of the most native-like model for 18 targets, and less than 10% for 66 targets. SELECTpro also ranked the single most native-like first for 15 targets, in the top five for 39 targets, and in the top ten for 53 targets, more often than any other method. Because the ranking metric is skewed by model redundancy and ignores poor models with a better ranking than the most native-like model, the BLUNDER metric is introduced to overcome these limitations. SELECTpro is also evaluated on a recent benchmark set of 16 small proteins with large decoy sets of 12500 to 20000 models for each protein, where it outperforms the benchmarked method (I-TASSER).

Conclusion: SELECTpro is an effective model selection method that scores models independently and is appropriate for use on any model set. SELECTpro is available for download as a stand alone application at: <http://www.igb.uci.edu/~baldig/selectpro.html>. SELECTpro is also available as a public server at the same site.

Background

Selecting the most native-like model from a set of possible models is a crucial task in protein structure prediction. A variety of Model Quality Assessment Programs (MQAPs) have been developed that assign numeric scores to models in a set, and then use the scores to rank the models and ultimately select a single model. MQAP methods can be divided roughly into three categories based on the type of information they use: evolutionary methods use sequence or profile similarity between target sequence and template, consensus methods use similarity between models, and structure-based methods use model coordinates [1]. Each category of methods has inherent strengths and weaknesses.

Evolutionary methods can provide quality scores that have been shown to correlate with structural similarity to native [2]. However, for lower confidence alignments the scores do not correlate well with structural similarity. Furthermore, identification of the best template and specific alignment can be difficult. In addition, models built from multiple templates or template-free methods cannot be scored appropriately by evolutionary methods alone.

Consensus methods take advantage of the observation that similar models produced by different predictors tend to be more accurate than those that are structural outliers. In practice, consensus methods outperform the methods they draw from, and they rarely pick a very poor model. The disadvantage, however, is that when the best model is a structural outlier it will be overlooked for lack of popularity [1]. Also, consensus methods are not appropriate for selecting from small sets of structurally diverse models, especially in the extreme case of a two-model set.

While consensus methods depend on similarity between models, structure-based methods calculate scores on each model independently. For this reason, structure-based methods can be applied to model sets of any size and diversity, and will produce the same score for a model regardless of the other models in the set. Structure-based methods can also be used for template-free modeling [3-6] and model refinement procedures [7, 8]. One weakness of high resolution structure-based methods, including protein free energy approximation functions [9-12] and physics based approaches [13, 14], is their sensitivity to local structural irregularities such as steric clashes and chain breaks, which can significantly bias scores on otherwise accurate models. Even slight differences in model backbones can produce significantly different scores [15]. Lower resolution structure-based methods, such as statistical potentials [6, 16, 17],

are more robust to backbone variation, but are sensitive to extended low contact-order regions in the models.

Here we describe SELECTpro, a novel structure-based MQAP that combines high and low resolution energy terms into a model selection method that is effective on model sets of variable size, diversity, and target difficulty. Most of our assessment is calculated from the CASP7 model quality assessment category (QA) results published online [18]. The QA category provides a framework for the unbiased evaluation of MQAPs on ensembles of models produced by diverse automated prediction methods.

Results and discussion

We analyze the CASP7 quality assessment category predictions with a focus on the quality of the model ranked first by each predictor and the recovery of the most native-like model in the set. Only *SetAll* is used in the assessment of the quality of the model ranked first by each group (Table 1). The results are very similar when using *SetComplete* (data not shown) because QA groups rarely rank an incomplete model first.

The assessment of the recovery of the most native-like model, is performed on both *SetAll* and *SetComplete* (Table 2) because the few cases where an incomplete model is the most native-like have a significant effect on the average recovery metrics of all QA groups. Incomplete and irregular models are especially challenging for structure-based methods. A comparison of the average Pearson Correlation on *SetAll* and *SetComplete*, highlights these issues (Table 3). The frequency of recovering the most native-like model is calculated on *SetComplete* (Figure 1).

The utility of SELECTpro for selecting the best model from a small set is demonstrated by selecting from the five models submitted for each target by the top automated predictors. These small set selection results are calculated using *SetAll* (Figure 2). SELECTpro is also evaluated on a recent benchmark set of 16 small proteins with large decoy sets of 12500 to 20000 models for each protein and compared to I-TASSER (Figure 3).

To make fair comparisons to groups participating on only a subset of targets, common subset comparisons between SELECTpro and each of these groups are included in Tables 1 and 2. Only groups participating on at least half of the targets are included, and for groups with multiple submissions only the best one is shown. In the results tables any value that is better than SELECTpro is underlined.

For multiple domain targets, the sum of GDT-TS over all domains is used as the GDT-TS of the model. Since the

Table 1: Quality of Model Ranked First (M_{QA1}) Relative to Most Native-Like Model (M_{max})

Group	Targets ^a	Summary Results				Common Subset Results				p-value
		$\Delta GDT_{QA1} = 0$	$\Delta GDT_{QA1\%} < 1$	$\Delta GDT_{QA1\%} < 10$	$\overline{\Delta GDT}_{QA1}$	$\Delta GDT_{QA1} = 0$	$\Delta GDT_{QA1\%} < 1$	$\Delta GDT_{QA1\%} < 10$	$\overline{\Delta GDT}_{QA1}$	
699_1	95 (124)	12	18	66	5.07	-	-	-	-	-
713_1	95 (124)	7	11	63	5.44	12	18	66	5.07	2.5E-01
634_1	95 (124)	7	15	53	7.75	12	18	66	5.07	1.6E-03
704_1	95 (124)	5	8	49	7.76	12	18	66	5.07	3.5E-04
178_1	95 (124)	8	12	59	8.44	12	18	66	5.07	3.0E-03
633_1	95 (124)	6	9	52	10.12	12	18	66	5.07	1.8E-06
692_1	95 (124)	6	9	52	10.16	12	18	66	5.07	1.2E-06
657_1	95 (124)	1	5	40	12.71	12	18	66	5.07	1.8E-08
691_1	95 (124)	0	1	24	15.10	12	18	66	5.07	2.2E-13
091_1	94 (123)	11	18	61	7.93	12	18	65	5.10	2.1E-03
026_1	94 (123)	1	2	40	9.30	12	18	65	5.10	1.2E-07
338_5	93 (122)	2	3	37	15.10	12	18	65	5.05	1.3E-09
556_1	93 (121)	10	15	51	6.83	12	18	64	5.15	1.8E-02
734_1	92 (120)	4	4	36	16.16	12	18	64	5.10	5.6E-11
718_1	92 (119)	1	3	32	14.04	11	17	64	5.19	1.6E-10
717_1	87 (112)	3	7	36	10.15	10	15	59	5.31	4.3E-08
016_1	86 (111)	5	9	49	7.93	10	16	58	5.26	1.4E-03
038_1	85 (108)	3	7	60	5.75	11	16	58	5.34	1.2E-01
276_1	80 (104)	5	5	<u>39</u>	8.94	11	17	54	5.21	7.7E-07
013_1	78 (100)	4	6	41	9.86	10	15	56	4.87	2.0E-05
703_1	69 (86)	3	6	35	8.74	9	15	45	5.35	1.2E-04
191_1	61 (78)	2	5	32	9.35	7	10	39	6.04	2.3E-03
066_1	55 (72)	1	2	14	23.19	7	10	45	4.09	4.3E-10

^a The number of targets where the QA group made a valid prediction (N_T) with the number of domains of these targets (N_D) in parentheses.

* SELECTpro (699_1) results appear in bold face and all results that are better than SELECTpro are underlined. Statistically significant p-values ($p < .05$) are also in bold.

Table 2: Recovery of Top GDT-TS Model (M_{\max})

SetAll							SetComplete						
Summary Results			Common Subset Results				Summary Results s			Common Subset Results			
Group	Targets ^a	<u>rank</u>	<u>$\Delta GDT_{BLUNDER}$</u>	<u>rank</u>	<u>$\Delta GDT_{BLUNDER}$</u>	p-value	Group	Targets	<u>rank</u>	<u>$\Delta GDT_{BLUNDER}$</u>	<u>rank</u>	<u>$\Delta GDT_{BLUNDER}$</u>	p-value
699_1^b	95 (124)	29.8	11.8	-	-	-	699_1	95 (124)	17.8	10.4	-	-	-
704_1	95 (124)	46.5	17.8	29.8	11.8	2.7E-06	633_1	95 (124)	20.7	11.8	17.8	10.4	4.7E-02
178_1	95 (124)	42.3	19.6	29.8	11.8	2.9E-04	634_1	95 (124)	29.5	12.7	17.8	10.4	5.7E-02
657_1	95 (124)	78.5	37.0	29.8	11.8	3.9E-20	704_1	95 (124)	24.1	13.1	17.8	10.4	1.1E-02
634_1	94 (121)	52.0	16.5	29.2	11.7	1.3E-02	178_1	95 (124)	24.1	13.7	17.8	10.4	6.5E-03
091_1	94 (123)	27.2	17.4	29.3	11.9	2.2E-05	657_1	95 (124)	53.5	32.0	17.8	10.4	8.6E-18
633_1	94 (121)	<u>39.0</u>	20.6	29.2	11.7	1.3E-08	713_1	94 (122)	18.3	10.9	17.9	10.4	2.0E-01
026_1	94 (123)	55.9	22.7	29.4	11.6	3.2E-10	692_1	94 (122)	20.6	11.6	17.7	10.3	6.7E-02
556_1	93 (121)	33.8	<u>11.7</u>	29.0	11.7	*	091_1	94 (123)	<u>16.8</u>	12.3	17.4	10.4	2.4E-02
692_1	93 (119)	38.7	<u>20.6</u>	29.2	11.6	1.1E-08	026_1	94 (123)	<u>37.3</u>	18.3	17.6	10.2	1.5E-07
691_1	93 (120)	98.1	28.6	28.6	11.7	9.6E-19	691_1	94 (123)	54.4	22.2	17.4	10.4	2.4E-14
338_2	93 (122)	60.4	30.2	30.2	11.9	2.7E-15	556_1	93 (121)	21.2	10.3	17.2	10.2	4.9E-01
713_1	92 (116)	<u>26.4</u>	12.8	29.6	11.8	3.2E-01	338_2	93 (122)	28.2	16.8	18.0	10.4	1.5E-08
734_1	89 (116)	<u>55.2</u>	31.5	29.3	11.2	1.6E-15	734_1	88 (115)	28.9	18.1	17.3	9.6	7.0E-09
718_1	83 (105)	81.6	31.9	30.5	12.0	1.6E-14	718_1	83 (105)	46.4	26.9	17.6	10.4	4.5E-13
717_1	78 (98)	46.8	22.8	30.9	12.0	3.4E-09	717_1	78 (98)	28.4	16.4	17.6	10.3	2.1E-05
013_1	78 (100)	60.1	27.5	30.2	12.0	1.5E-09	013_1	78 (100)	32.4	17.6	18.5	10.3	3.7E-06
276_1	78 (102)	52.9	28.9	29.6	11.6	3.3E-12	276_1	78 (102)	29.0	18.7	17.5	10.2	8.5E-10
038_1	70 (87)	<u>25.9</u>	11.9	27.6	11.8	4.6E-01	038_1	74 (95)	19.8	10.7	17.4	10.4	3.6E-01
703_1	69 (86)	<u>37.2</u>	20.6	31.5	11.9	5.1E-07	703_1	69 (86)	20.6	14.5	17.6	10.5	4.7E-04
191_1	61 (78)	45.5	21.9	26.2	12.6	1.0E-06	191_1	61 (78)	27.6	15.2	16.7	11.1	2.8E-03
066_1	55 (72)	91.1	54.6	30.5	10.5	5.1E-24	066_1	55 (72)	48.0	46.5	18.5	9.1	1.8E-18
016_1	53 (72)	<u>30.9</u>	20.0	31.2	12.5	2.0E-05	016_1	53 (70)	18.2	18.5	17.8	11.0	1.4E-05

^a The number of targets where the QA predictor scored M_{\max} (N_T) with the number of domains of these targets (N_D) in parentheses.

^b In the CASP7 submission SELECTpro did not have a score for M_{\max} of target T0356 due to a processing error. We added in the score for this analysis in order to make complete common subset comparisons.

* SELECTpro (699_1) results appear in bold face and all results that are better than SELECTpro are underlined. Statistically significant p-values ($p < .05$) are also in bold.

Table 3: Correlation of Selected Groups

Group	Targets	SetAll \overline{PC}	SetComplete \overline{PC}	ΔPC
634_I (Pcons) ^a	95	0.811	0.847	0.036
713_I (Circle-QA) ^b	95	0.765	0.823	0.058
633_I (ProQ) ^b	95	0.716	0.781	0.064
699_I (SELECTpro) ^b	95	0.676	0.763	0.087
556_I (LEE) ^c	93	0.814	0.792	-0.023

^a Consensus method.

^b Structure based method.

^c QA scores calculated as GDT-TS similarity to human predictor of LEE.

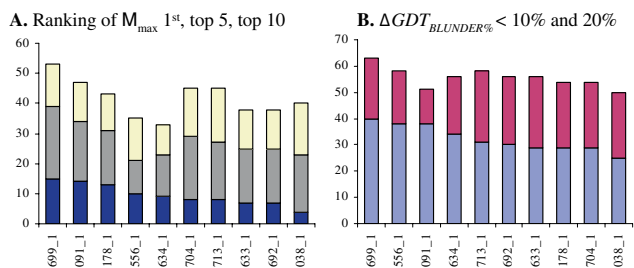


Figure 1
Recovery of M_{max} using SetComplete. (A) number of targets where M_{max} is ranked first (top of dark blue bar), in the top five (top of gray bar), and in the top ten (top of white bar). (B) number of targets where ΔGDT_{BLUNDER%} is less than 10% (top of light blue bar) and less than 20% (top of purple bar). Only the first ten groups are shown in both graphs.

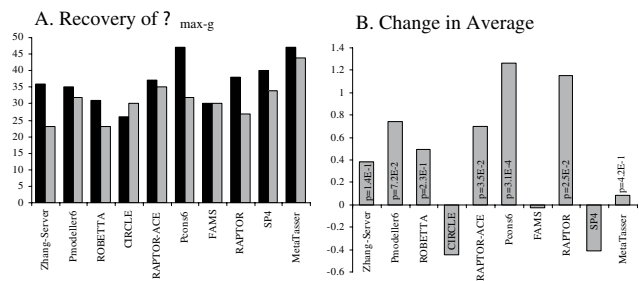


Figure 2
Reranking models from top servers. Each server predictor submitted five models per target, with the highest confidence model ranked first. (A) the number of targets where each server's highest GDT-TS model is ranked first is shown with gray bars, and black bars when the models are reranked with SELECTpro. (B) shows the change in average GDT-TS for each group when SELECTpro is used to select model 1. P-values of paired t-tests are shown above the horizontal axis when SELECTpro demonstrates improved model selection and statistically significant improvements ($p < .05$) are in bold.

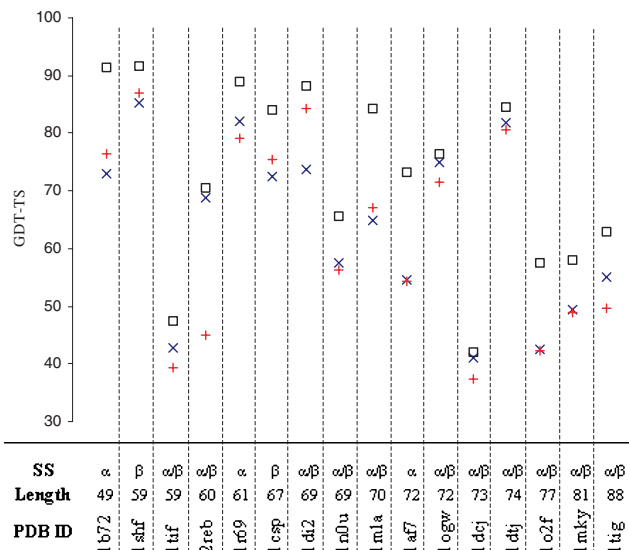


Figure 3
Large Decoy Set Model Selection. Large decoy set model selection with SELECTpro on I-TASSER benchmark set. This set of 16 small proteins was used as one of the benchmark sets for evaluating the I-TASSER method [19]. The complete decoy sets can be downloaded from [20]. Each protein has from 12500 to 20000 decoy models. For each protein different symbols are used to indicate the GDT-TS of M_{max} (□), SELECTpro's M_{QA1} (x), and I-TASSER's M_{QA1} (+).

QA predictions correspond to the entire structures, it is impossible to fairly assess the domains independently.

To assess the significance of the summary statistics compared in Table 1, Table 2, and Figure 2, we performed paired t-tests between SELECTpro each other group on common subsets of targets (or targets and models when appropriate). All p-values from the tests appear in the tables and figure, but only statistically significant p-values ($p < .05$) are shown in bold.

The following notations are used throughout the results section:

- M_{\max} : The model with the highest GDT-TS among all server models.
- M_{QA1} : The model with the highest QA score.
- N_T : The number of targets a group made valid predictions on.
- N_D : The number of domains a group made valid predictions on.

The recovery of M_{\max} by a QA predictor can only be evaluated if M_{\max} was scored by the predictor. In most cases QA predictors did not provide scores for all available server models, and frequently there is no score for M_{\max} . For example, predictor 016_1 (AMBER/PB) made submissions on 86 targets, but M_{\max} is only scored for 53 of these targets – so only these targets ($N_T = 53$) can be evaluated for this predictor.

Quality of Model Ranked First (M_{QA1}) Relative to Most Native-Like Model (M_{\max})

In this section on the assessment of the model ranked first, and the corresponding Table 1, we use the following three metrics:

- $\Delta GDT_{QA1} = GDT-TS(M_{\max}) - GDT-TS(M_{QA1})$: The GDT-TS difference between M_{\max} and M_{QA1} measures how much is lost by selecting M_{QA1} rather than M_{\max} for a single target.
- $\overline{\Delta GDT_{QA1}} = \Sigma \Delta GDT_{QA1} / N_D$: The average ΔGDT_{QA1} is a simple way of assessing the quality of M_{QA1} over all targets.
- $\Delta GDT_{QA1\%} = \Delta GDT_{QA1} / GDT-TS(M_{\max})$: The GDT-TS difference percentage allows for comparison across targets with different numbers of domains and difficulty levels.

The columns of Table 1 are: (1) group number; (2) number of targets the group made predictions on; (3) number of targets such that $\Delta GDT_{QA1} = 0$; (4) number of targets such that $\Delta GDT_{QA1\%} < 1\%$; (5) number of targets such that $\Delta GDT_{QA1\%} < 10\%$; and (6) $\overline{\Delta GDT_{QA1}}$. The common subset results section has an additional column for the p-value of the paired t-test using ΔGDT_{QA1} . The rows are sorted first by the number of targets and then by $\overline{\Delta GDT_{QA1}}$. Of the groups participating on all 95 targets, SELECTpro has the lowest average ΔGDT_{QA1} , with a value of 5.07, followed closely by group 713_1 (Circle-QA), with

a value of 5.44. Predictor 038_1 (GeneSilico) has an average ΔGDT_{QA1} of 5.75, with predictions on 85 targets. In common subset comparisons with these two groups SELECTpro is not significantly better, with p-values of .25 and .12 respectively. In common subset comparisons with all remaining groups SELECTpro is significantly better.

Another way to assess the quality of M_{QA1} over many targets is to count the number of targets such that M_{QA1} is the best model, or nearly the best, in the set. A method that performs very well on most targets, but very poorly on a few, would still be recognized by this criteria. SELECTpro recovers the best model for 12 targets, selects a model with $\Delta GDT_{QA1\%} < 1\%$ for 18 targets, and selects a model with $\Delta GDT_{QA1\%} < 10\%$ for 66 targets. Group 091_1 (Ma-OPUS) also performs well, with 11, 18, and 61 targets in the respective categories. Only the 60 targets with $\Delta GDT_{QA1\%} < 10\%$ of predictor 038_1 (GeneSilico) on its 85 target subset are better than SELECTpro in common subset comparison (58 for SELECTpro).

The BLUNDER Measure Recovery of M_{\max}

How well does a QA predictor recover M_{\max} ? The traditional metric to assess M_{\max} recovery is the *rank* of M_{\max} , and the average *rank* over many targets (\overline{rank}). While *rank* captures some important information, it ignores the redundancy of models and the quality of models ranked better than M_{\max} . Consider the following hypothetical situation: group A ranks M_{\max} 10th and all nine models ranked above it are redundant with ΔGDT of ~ 2.0 , group B ranks M_{\max} 5th and the four models ranked above it are diverse with a ΔGDT between 10.0 and 20.0. Which group has done a better job of recovering M_{\max} ? In this example, the *rank* metric favors group B, although group A ranks only a single redundant model above M_{\max} . In addition, the models ranked better than M_{\max} by group A have only slightly lower GDT-TS than M_{\max} , while the models ranked better than M_{\max} by group B are significantly worse than M_{\max} . To address these weaknesses of the *rank* metric, we introduce the BLUNDER metric, which focuses on the worst model ranked better than M_{\max} (the most embarrassing blunder). This measure is not affected by model redundancy and measures the quality of models ranked above M_{\max} . The BLUNDER metric is defined using the following notation, and used in the assessment of the recovery of M_{\max} and the corresponding Table 2 and Figure 1:

- $M_{BLUNDER}$: The model with the minimum GDT-TS among models ranked better than M_{\max} .
- $\Delta GDT_{BLUNDER} = GDT-TS(M_{\max}) - GDT-TS(M_{BLUNDER})$: The GDT-TS difference between M_{\max} and $M_{BLUNDER}$ measures the size of the worst blunder.

- $\overline{\Delta GDT_{BLUNDER}} = \Sigma \Delta GDT_{BLUNDER} / N_D$: The average $\Delta GDT_{BLUNDER}$ measures how well a method robustly recovers M_{max} over many targets.
- $\Delta GDT_{BLUNDER\%} = \Delta GDT_{BLUNDER} / GDT-TS(M_{max})$: The $\Delta GDT_{BLUNDER}$ percentage allows for comparison across targets with different numbers of domains and difficulty levels.

Figure 1 contains graphs of the frequency of recovering M_{max} using the *rank* (A) and $\Delta GDT_{BLUNDER\%}$ (B) measures on *SetComplete*. SELECTpro ranks M_{max} first for 15 targets, in the top five for 39 targets, and in the top ten for 53 targets. SELECTpro's $\Delta GDT_{BLUNDER\%}$ values are less than 10% of $GDT-TS(M_{max})$ for 40 targets and less than 20% for 63 targets. These results are best among all QA participants. The average M_{max} recovery results are summarized in Table 2. The results columns are (1) average *rank* (\overline{rank}) and (2) average $\Delta GDT_{BLUNDER}$ ($\overline{\Delta GDT_{BLUNDER}}$) on *SetAll* and *SetComplete*. The common subset results section also includes a column for the p-value of a paired t-test using $\Delta GDT_{BLUNDER}$ (*p-value*). Rows are sorted separately for each dataset by N_T first and then $\overline{\Delta GDT_{BLUNDER}}$. On *SetComplete* SELECTpro has a $\overline{\Delta GDT_{BLUNDER}}$ of 10.4. In common subset comparisons one group has a lower \overline{rank} : group 091_1 (Ma-OPUS) with \overline{rank} of 16.8 on 94 targets compared to 17.4 for SELECTpro. On *SetAll* SELECTpro did not submit a score for M_{max} of target T0356 (HHpred2_TS1) due to a processing error. In order to make complete common subset comparisons when possible we added in the SELECTpro score for HHpred2_TS1. SELECTpro ranks it 86th and $\Delta GDT_{BLUNDER} = 50.0$. Both results are significantly worse than the SELECTpro averages.

Pearson Correlation for Individual Proteins

The assessor evaluation of the quality assessment category [18] focused on the Pearson Correlation between the QA scores and GDT-TS. Here we use the Pearson Correlation only to highlight some of the difficulties for structure-based methods in dealing with incomplete models, as well as basic non-protein like structural features. Approximately half of the models in *SetAll* are incomplete, with backbone coordinates missing for one or more residues.

Incomplete models present a challenge to SELECTpro and other structure-based methods because the scores for each model are only comparable when calculated on coordinates for the same set of residues. Another issue is that some complete models have severe chain-breaks, severe steric clashes, or significant portions modeled only as extended chains. These local problems can overwhelm the energy of what may otherwise be a good model. Consensus methods

do not suffer from these local structure problems. Given this rationale, one would expect structure-based methods to see the most improvement in terms of average Pearson Correlation on *SetComplete* relative to *SetAll*. Table 3 shows the average Pearson Correlation of five selected groups. Predictors 713_1 (Circle-QA), 633_1 (ProQ), and SELECTpro are structure-based MQAPs, while 634_1 (Pcons) is a consensus method and 556_1 (LEE) scored structures based on the GDT-TS similarity to their human Model 1 CASP7 prediction [18]. As expected, the structure-based MQAPs improve more than the structural similarity-based methods. The even greater increase in Pearson Correlation for SELECTpro can be accounted for by the failure to generate appropriate complete models for some of the incomplete models resulting in QA scores calculated on extended chains.

Reranking Top Server Group Models

Predictors in CASP may submit up to five models, but CASP evaluation focuses on the model designated as Model 1. Clearly, the selection of Model 1 is critical in the CASP setting and for protein structure prediction in general. Figure 2 contains the results when SELECTpro is used to rerank the five models submitted by each of the top ten servers from CASP7, compared to each server's results. In the following assessment M_{max-g} is the model with the highest GDT-TS of the five models submitted by a server. Figure 2 (A) shows that SELECTpro recovers M_{max-g} more frequently than 8 of the top 10 server groups; in addition, when SELECTpro is used to select Model 1 the average GDT-TS increases for 7 of 10 sever groups; however, the increase is only statistically significant for 3 groups. SELECTpro improves using both criteria for the top 3 server groups (Zhang-Server, Pmodeller6, and ROBETTA). These results highlight the utility of SELECTpro for the task of model selection. The comparisons made here are fair because structure-based methods can be applied in the server setting to any number of models.

Large Decoy Set Model Selection

Here we analyze SELECTpro's model selection capability on the large decoy sets for 16 small proteins from a recent I-TASSER benchmark set [19]. The I-TASSER prediction method generates 12500 to 20000 different backbone conformations. The complete decoy sets can be downloaded from [20]. The consensus method SPICKER [21] is used to cluster the models and a centroid model is built from the first cluster. A second round of simulation resolves the steric clashes in the centroid model and results in the final predicted model. The centroid model and final model are not part of the decoy set. In order to make a fair model selection comparison the decoy model closest to the centroid is used as I-TASSER's M_{QA1} .

On the benchmark set SELECTpro has an average GDT-TS of 63.7, while I-TASSER has an average GDT-TS of 62.1. SELECTpro's average ΔGDT_{QA1} is 9.2 and I-TASSER's ΔGDT_{QA1} is 10.7. Figure 3 displays the GDT-TS results for the individual proteins in the benchmark set. Different symbols are used to indicate the GDT-TS of M_{max} (\square), the GDT-TS of SELECTpro's M_{QA1} (\times), and the GDT-TS of I-TASSER's M_{QA1} (+) for each protein. A paired t-test of the hypothesis that SELECTpro and I-TASSER's mean performance are equal produces a p-value of .19, which is not statistically significant, but does give some evidence that SELECTpro can select a very good model from a large set of decoys at least well as an established method that utilizes consensus methods.

Conclusion

A MQAP that can select the most native-like model from a set of possibilities has a variety of applications in protein structure prediction. The new quality assessment category introduced in CASP7 allows for the unbiased assessment of MQAPs on the models produced by automated predictors. This category allows researchers to focus on the model scoring aspect of protein structure prediction.

The results presented in this work demonstrate that SELECTpro, a structure-based model selection method, consistently selects one of the best models from the large diverse sets of models produced by automated predictors, across all levels of target difficulty. On these large diverse sets of models, SELECTpro also recovers the single most native-like model well compared to other methods. On the small sets of five models submitted for each target by the top automated predictors, in most cases SELECTpro selects better models than the predictors themselves.

Since SELECTpro and other structure-based methods score models independently, they can be incorporated into the model selection pipelines of individual protein structure prediction servers. For this reason, it may help predictors if the CASP organizers distinguished methods that score models independently from those that do not.

Consensus and structure-based methods can be combined to achieve improved results. For example, the meta-server method Pmodeller [22] combines consensus (Pcons [23]) and structure-based methods (ProQ [24]) to predict protein structures more accurately than either method in isolation. The assessment of the QA category by CASP assessors recognized the consensus method Pcons (group 634_1) for the high Pearson Correlation between their scores and model GDT-TS on most targets [18]. In their own assessment the authors of Pcons recognized that while consensus methods perform well in

most cases, "when most of the models are incorrect and the few correct models are outliers a consensus based approach cannot be expected to make an optimal choice." [1] For instance, they identified three particular targets in CASP7 where their consensus method failed: T0283, T0350, and T0351 [1]. The Pcons average ΔGDT_{QA1} on these three targets is 30.8. The same research group's structure-based method ProQ (group 633_1) has an average ΔGDT_{QA1} of 17.2. In contrast, on these three targets SELECTpro has an average ΔGDT_{QA1} of only 7.1. This example highlights the potential of combining SELECTpro with existing model selection methods.

SELECTpro has been made publicly available as a server, where users may submit from 2 to 100 models for evaluation. In addition to the global confidence scores, the scores of individual energy terms are also returned to the user by email for each model submitted. SELECTpro is one of several protein structure tools in the SCRATCH suite of predictors [25], and is available through: <http://www.igb.uci.edu/~baldig/selectpro.html>.

Methods

Datasets

All of the comparative analysis in this work is performed on the server models and quality assessment predictions submitted in the CASP7 [26] experiment. The CASP QA experiment is particularly relevant for the evaluation of model selection methods for several reasons: (1) the QA predictors were blind to the true structures at the time of prediction making it impossible for methods to be tuned to improve results; (2) the set of proteins is diverse: the 95 targets range in size from 68 to 530 amino acids, come from a variety of organisms, and span the full range of prediction difficulty; (3) each target has more than 200 predicted models that contain the types of errors that occur in automated structure prediction; (4) the protein set is not selected by any of the participating QA groups; (5) the models are scored by a variety of methods and the results are publicly available. We perform analysis on the set of all models (*SetAll*) and a subset of models (*SetComplete*) that are complete and free of gross structural irregularities, as described below. All of the ABIPro models and some of the 3Dpro models were optimized using the exact energy function of SELECTpro. These models are removed because of the obvious bias towards these models. In recent CASP experiments the GDT-TS [27] has been used as the primary automatic structural similarity measure. The published GDT-TS values from the CASP7 website are the only structural similarity measure used in this work.

SetAll

The *SetAll* dataset consists of the server models with a GDT-TS value published on the CASP7 website, a total of 23,423

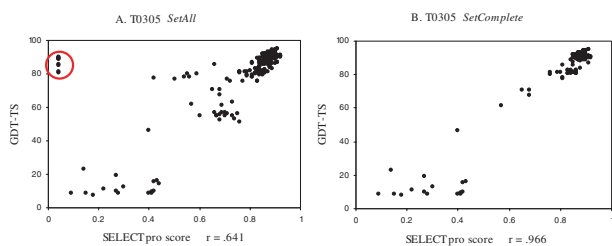


Figure 4
SetAll versus SetComplete. Plots of SELECTpro scores versus GDT-TS scores for T0305 models from *SetAll* (A) and *SetComplete* (B). The Pearson correlation is .641 for *SetAll* and .996 for *SetComplete*. This large difference is mainly due to the extended chain models (circled in plot A) scored by SELECTpro due to an error in our use of Modeller to generate complete models from incomplete ones.

models. To calculate a score on a protein model SELECTpro requires the backbone coordinates (N, C_α, C) for all model residues as input. A total of 8,812 models in *SetAll* have only a C_α trace or have no coordinates for one or more residues. Modeller8v1 [28-30] was used to generate complete models from the incomplete ones, and then the complete models were scored by SELECTpro. In most cases the complete models were built appropriately from the incomplete models; however, in some cases the final model was a fully extended chain due to an error in our application of Modeller. We failed to identify this problem until after the completion of the CASP7 competition. The SELECTpro scores versus GDT-TS scores for all models of target T0305 are displayed in plot A of Figure 4. The circled outliers with very low confidence scores and high GDT-TS scores are models that were incomplete and the complete models generated by Modeller were fully extended chains. The Pearson correlation on the set of all models for T0305 is .641. The SELECTpro scores versus GDT-TS scores for complete models only are displayed in plot B of Figure 4, and the Pearson correlation is .966.

SetComplete

The scores produced by SELECTpro are comparable on complete models of the same sequence. There is no standard for the handling of incomplete models and we assume that participating groups took a variety of approaches. Using only complete models ensures that the MQAP scores are calculated from the same coordinates. Thus, the models retained in *SetComplete* are screened first for completeness. Models missing backbone coordinates for one or more residues are removed. This leaves 14,611 models.

Structure-based MQAPs are susceptible to local structural irregularities in models, and will tend to score such models poorly. This is why methods developed to select near-native models from sets of decoys remove such

models from consideration [31]. We apply additional filters (described below) for C_α-C_α clashes, C_α-C_α chain breaks, and expanded termini to remove an additional 1,217 models leaving 13,494 more plausible models in *SetComplete*.

The C_α-C_α clash model filter enforces a squared difference penalty for C_α-C_α distances less than 3.6 Å. The distance between the C_α atoms of residue *i* and *j* is denoted by $r_{i,C\alpha,j,C\alpha}$ and *N* is the protein length. The constant 13.52 in the threshold below corresponds to two severe clashes where $r_{i,C\alpha,j,C\alpha} = 1.0$ Å. Models with a sum of squared differences greater than 13.52 per 100 residues are filtered out.

$$\sum_{i>j} \max\{0, 3.6 - r_{i,C\alpha,j,C\alpha}\}^2 > 13.52(N/100)$$

The C_α-C_α chain break model filter enforces a squared difference penalty for $r_{i,C\alpha,i+1,C\alpha}$ distances greater than 4.0 Å. The constant 16.0 in the threshold below corresponds to a single chain break where $r_{i,C\alpha,i+1,C\alpha} = 8.0$ Å. Models with a sum of squared differences greater than 16.0 per 100 residues are filtered out.

$$\sum_i \max\{0, r_{i,C\alpha,i+1,C\alpha} - 4.0\}^2 > 16.0(N/100)$$

The expanded termini filter removes models where a large portion of the structure is modeled as expanded chain with no non-local interactions. The screening procedure is: scan from the N-terminus until three consecutive residues have a contact number of at least 10, and repeat from the C-terminus. The contact number of a residue is defined here as the number of other C_β atoms within 10 Å of the residue's C_β [3]. If the sum of low contact number termini residues is at least 20% of *N*, the model is filtered out.

Model Representations

Reduced representation

In the reduced representation the heavy backbone atoms, carbonyl oxygen, amide hydrogen (N, C_α, C, O, H), and C_β are represented explicitly. For glycine residues a pseudo C_β is calculated. The side-chain atoms are represented by a single united point (centroid) [32, 33]. The centroid is calculated as the mean of the position of the heavy side-chain atoms. For glycine and alanine the centroid (CT) is set to the C_β atom. Only the heavy backbone atoms (N, C_α, C) are used as input to SELECTpro and the positions of additional atoms and centroids are calculated from these.

All heavy-atom representation

In the all heavy-atom representation the centroid is removed and the heavy side chain atoms are represented explicitly. The side-chains are initially placed onto the backbone of the reduced representation in their most likely conformation according to the SCWRL backbone-dependent rotamer library [34]. The side-chain placements are then optimized using the SELECTpro all-atom energy terms (described below) in conjunction with the rotamer library.

Energy Functions Overview

$E_{REDUCED}$ is the combined energy calculated from the reduced representation. $E_{REDUCED}$ is a linear combination of predicted ($E_{PRED-SS}$, $E_{PRED-SA}$, $E_{PRED-CM}$), physical ($E_{VDW-REP}$), and statistical (E_{CT-REP} , $E_{STAT-ENV}$, $E_{STAT-PW-CI}$, $E_{STAT-PW-CD}$, E_{ROG}) terms:

$$E_{REDUCED} = w_1 E_{PRED-SS} + w_2 E_{PRED-SA} + w_3 E_{PRED-CM} + w_4 E_{BETA} + w_5 E_{VDW-REP} + w_6 E_{CT-REP} + w_7 E_{STAT-ENV} + w_8 E_{STAT-PW-CI} + w_9 E_{STAT-PW-CD} + w_{10} E_{ROG}$$

$E_{ALL-ATOM}$ consists of the energy terms that depend on the all heavy-atom representation. $E_{ALL-ATOM}$ is a linear combination of the following physical terms:

$$E_{ALL-ATOM} = w_{11} E_{SC-HB} + w_{12} E_{LEN-JONES} + w_{13} E_{SOLVATION} + w_{14} E_{ELECTRO}$$

E_{FINAL} is the sum of $E_{REDUCED}$ and $E_{ALL-ATOM}$, and is used for the final scoring of models by SELECTpro. The individual energy terms are outlined briefly below and the detailed description of the novel terms follow in the remainder of this section. Underlined terms are adapted from previously described energy terms their details are included in the Appendix.

Parameter Weights

The parameter weights were determined by repeatedly varying individual weights and maximizing the sum of the GDT-TS of the lowest E_{FINAL} models on a training set built from CASP6 protein domains. For each CASP6 protein domain a set of 500 decoy models was generated using fragment assembly with the RMSD to native as the dominant term in the objective function [3].

$E_{REDUCED}$

$E_{PRED-SS}$: predicted secondary structure

$E_{PRED-ACC}$: predicted solvent accessibility

$E_{PRED-CM}$: predicted contact map

E_{BETA} : sheet formation

E_{BB-REP} : backbone repulsion

E_{CT-REP} : centroid repulsion

$E_{STAT-ENV}$: residue environment potential [3]

$E_{STAT-PW-CI}$: context independent pair-wise potential [3, 16]

$E_{STAT-PW-CD}$: context dependent pair-wise potential [6]

E_{ROG} : compactness

$E_{ALL-ATOM}$

E_{SC-HB} : side-chain hydrogen bonding

$E_{LEN-JONES}$: van der Waals forces [10]

$E_{SOLVATION}$: solvation effects [35]

$E_{ELECTRO}$: electrostatic interactions

Throughout this work the convention of all capital letters referring to global energy and all lower case referring to local energy is used. For instance, $E_{PRED-CM}$ refers to the global contact map energy and $E_{pred-cm}(i,j)$ refers to the contact map energy between residues i and j .

Parameter notation used in energy equations

Model variables

$r_{i,x,j,y}$: distance between atom x of residue i and atom y of residue j

$r_{x,y}$: distance between atom x and atom y

$v_{i,x,j,y}$: vector from atom x of residue i to atom y of residue j

$u_{i,x,j,y}$: unit vector calculated from $v_{i,x,j,y}$

N_i : number of residues in contact with residue i , with contact defined as $r_{i,C\beta,j,C\beta} < 10 \text{ \AA}$

ϕ_i : Phi angle of residue i

ψ_i : Psi angle of residue i

Protein specific input parameters

aa_i : amino acid type of residue i

ss_i : predicted secondary structure of residue i (H,E,C)

acc_i : predicted solvent accessibility of residue i ('e': exposed, '-', buried)

$cmap_{i,j}$: predicted contact/non-contact between residues i and j , with contact defined as $r_{i,C\alpha,j,C\alpha} < 12 \text{ \AA}$

Protein independent parameters

I_{value} : ideal parameter value for a given calculation

σ_{value} : standard deviation value for a given calculation

vdw_x : van der Waals radius of atom x

vdw_{x+y} : $vdw_x + vdw_y$

$\Theta_{stat-env}$: pre-calculated statistics for use in $E_{STAT-ENV}$

$\Theta_{stat-pw-oi}$: pre-calculated statistics for use in $E_{STAT-PW-CI}$

$\Theta_{stat-pw-od}$: pre-calculated statistics for use in $E_{STAT-PW-CD}$

$D_{min,pw-od}$: minimum interaction distance for centroid pairs used in $E_{STAT-PW-CD}$

$D_{max,pw-od}$: maximum interaction distance for centroid pairs used in $E_{STAT-PW-CD}$

D_{min-CT} : minimum distances between centroids of amino acid pairs observed in pdb_select25 [36].

Reduced Representation Energy Term Details

The details of how the novel reduced representation energy terms are calculated are presented in this section. The predicted structural terms $E_{PRED-SS}$, $E_{PRED-ACC}$, and $E_{PRED-CM}$ and the β -strand pairing term, E_{BETA} , are novel and unique to SELECTpro. Additional reduced representation terms are adapted from previously published work and their details are included in the Appendix.

Predicted structural features overview

The predicted structural feature predictions used in $E_{PRED-SS}$, $E_{PRED-ACC}$, and $E_{PRED-CM}$ come from the SCRATCH suite of predictors [25]. Each predictor is trained in a supervised fashion using curated non-redundant datasets extracted from the PDB [37]. The secondary structure (SSpro [38]) and solvent accessibility (ACCpro [39]) predictors use ensembles of 1D-RNN (one dimensional-recursive neural network) architectures [40]. The contact map predictor (CMAPpro [41]) uses ensembles of 2D-RNN architectures [40].

 $E_{PRED-SS}$: predicted secondary structure

The predicted secondary structure term $E_{PRED-SS}$ penalizes deviation of the torsion angles from the torsion angle parameters for helices and strands predicted by SSpro. There is no penalty for predicted coils. The parameter values for helix residues are: $I_{H\phi} = -65.3$, $\sigma_{H\phi} = 11.9$, $I_{H\psi} = -39.4$, $\sigma_{H\psi} = 11.3$. The parameter values for strand residues are: $I_{E\phi} = -135.0$, $\sigma_{E\phi} = 15.0$, $I_{E\psi} = 135.0$, $\sigma_{E\psi} = 15.0$. Only torsion angles that are more than two

standard deviations from the ideal are penalized, with the penalty defined as follows:

$$E_{PRED-SS} = \sum_{ss_i=H} E_{pred-helix}(i) + \sum_{ss_j=E} E_{pred-strand}(j)$$

$$E_{pred-helix}(i) = \sqrt{\Theta_{H\phi}(i) \left(|phi_i - I_{H\phi}| - 2\sigma_{H\phi} \right)^2 + \Theta_{H\psi}(i) \left(|psi_i - I_{H\psi}| - 2\sigma_{H\psi} \right)^2}$$

$$\Theta_{H\phi}(i) = \begin{cases} 1, & \text{if } |phi_i - I_{H\phi}| > 2\sigma_{H\phi} \\ 0, & \text{otherwise} \end{cases}$$

$$\Theta_{H\psi}(i) = \begin{cases} 1, & \text{if } |psi_i - I_{H\psi}| > 2\sigma_{H\psi} \\ 0, & \text{otherwise} \end{cases}$$

The definition of $E_{pred-strand}(j)$ is equivalent to $E_{pred-helix}(i)$, but with $I_{E\phi}$, $\sigma_{E\phi}$, $I_{E\psi}$ and $\sigma_{E\psi}$ in place of the corresponding helical values.

 $E_{PRED-ACC}$: predicted solvent accessibility

The solvent accessibility predictor ACCpro predicts the percent of solvent accessibility in 5% increments for each residue. Using 25% exposure as a binary threshold the accuracy of the predictor is $\sim 77\%$ [39]. The binary exposure ('e')/burial ('-') prediction is used as the predicted solvent accessibility for $E_{PRED-ACC}$. In the reduced representation the solvent accessibility of residue i is estimated by its contact number (N_i), where $N_i > 16$ is considered buried [3]. If the predicted status of a residue is not realized in the model, the penalty is calculated as:

$$E_{PRED-ACC} = \sum_i E_{pred-acc}(i)$$

$$E_{pred-acc}(i) = \begin{cases} (17 - N_i)^2, & \text{if } acc_i = '-' \text{ and } N_i \leq 16 \\ (N_i - 16)^2, & \text{if } acc_i = 'e' \text{ and } N_i > 16 \\ 0, & \text{otherwise} \end{cases}$$

 $E_{PRED-CM}$: predicted contact map

The contact map predictor CMAPpro predicts the probability of contact or non-contact between C_{α} atoms, with a contact threshold of 12 Å. The strategy utilized to infer predicted contacts from the probability matrix [41] results in maps that are sparse when compared to those of real proteins; thus, unrealized contacts are penalized while non-contacts are not. The constant 1.0 is added to the penalty to ensure that all unrealized contacts make a significant contribution to $E_{PRED-CM}$.

$$E_{PRED-CM} = \sum_{j>i} \Theta(i, j) E_{pred-cm}(i, j)$$

$$\Theta(i, j) = \begin{cases} 1, & \text{if } r_{i,C_{\alpha},j,C_{\alpha}} > I_{cm-thresh} \\ 0, & \text{otherwise} \end{cases}$$

$$E_{pred-cm}(i, j) = cmap_{i,j} \left(1.0 + \frac{r_{i,C_{\alpha},j,C_{\alpha}}^2}{I_{cm-thresh}^2} \right)$$

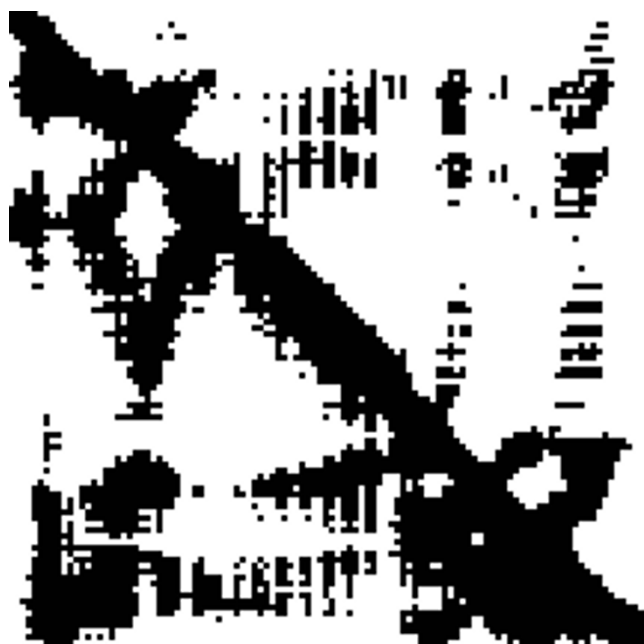


Figure 5
Contact map comparison. True contact map of target T0304 in lower left, predicted contacts upper right. Contact is defined as $C\alpha$ atoms within 12 Å. For predicted contacts with a sequence separation of at least six, 651 of 915 (71%) are correct.

The predicted contact map can help identify the highest GDT-TS models in the set, even when they are not highly similar to native. A good example of this is CASP7 target T0304 is a 122 residue α/β protein where the highest GDT-TS model in the set is Zhang-Server_TS1 (GDT-TS = 45.55). Most secondary structure predictors (including SSpro) failed to predict the first two strands making this target especially difficult. No QA method ranked the highest GDT-TS model first; however, SELECTpro ranked it second and the model ranked first by SELECTpro (T0304.Zhang-Server_TS4) has the second highest GDT-TS. These models have the lowest $E_{PRED-CM}$ of any models in the set, but the native structure has an even lower $E_{PRED-CM}$. Figure 5 compares the native and predicted contact maps for target T0304.

E_{BETA} : strand pairing

The formation of hydrogen bonds between the residues of β -strand partners is a major determinant of the tertiary structure of β and α/β proteins. The β hydrogen bonding treatment described here favors realistic strand pairing and sheet formation. The treatment also efficiently accommodates bulges in strands because it does not force the register between two paired strands. E_{BETA} is the global strand pairing energy that penalizes the hydrogen bonding of β residues between strand pairs. $E_{beta-sp}$

$E_{beta-sp}(\beta_k \rightarrow \beta_w)$ is the strand pairing energy of strand β_k to strand β_w . $E_{beta-sp}$ is only commutative if the two strands have the same length. $E_{beta-hb}(i,j)$ is the hydrogen bonding penalty between residues i and j .

$E_{beta-sp}$ is calculated for all possible strand pairings, but only the two lowest energies from each strand are used in E_{BETA} . Other strand-strand interactions are ignored. In the equations below S is the set of all strands in the protein, β_{m1} is the strand with the minimum pairing energy from β_k , and β_{m2} is the strand with the next lowest pairing energy from β_k . If the strand count is less than six at least two of the strands must be edge strands. This is accounted for by only considering the single best strand partner for two strands.

$$E_{BETA} = \sum_{\beta_k \in S} E_{beta-sp}(\beta_k \rightarrow \beta_{m1}) + E_{beta-sp}(\beta_k \rightarrow \beta_{m2})$$

$$\beta_{m1} = \{\beta_x : \min_{\beta_x \in S / \{\beta_k\}} E_{beta-sp}(\beta_k \rightarrow \beta_x)\}$$

$$\beta_{m2} = \{\beta_\gamma : \min_{\beta_\gamma \in S / \{\beta_k, \beta_{m1}\}} E_{beta-sp}(\beta_k \rightarrow \beta_\gamma)\}$$

In the equations for $E_{beta-sp}$ below, S_k is the set of all residues in strand β_k . Each time $E_{beta-hb}$ is calculated the pair (i,j) is chosen with i from S_k and j from S_w , such that $E_{beta-hb}$ is minimized. Then residue i is removed from S_k , and residue j is removed from S_w . $E_{beta-hb}$ is calculated once for each residue in S_k . If S_k has more residues than S_w each unpaired residue is given maximum penalty of $E_{beta-hb}$.

$$E_{beta-sp}(\beta_k \rightarrow \beta_w) = \sum_{S_k \neq \emptyset} E_{beta-hb}(i, j)$$

$$(i, j) = \{(x, y) : \min_{x \in S_k, y \in S_w} E_{beta-hb}(x, y)\}$$

$$S_k = S_k / \{i\}, S_w = S_w / \{j\}$$

Between two anti-parallel strand partners, only every other pair of residues is hydrogen bonded. For the pairs that are not hydrogen bonded, a pseudo-bonding calculation is used. The hydrogen bonding energy and pseudo-bonding energy are both calculated and the minimum of the two is used in $E_{beta-hb}(i,j)$.

If residues i and j are paired in parallel strands, either i forms hydrogen bonds with $j-1$ and $j+1$, or j forms hydrogen bonds with $i-1$ and $i+1$. No hydrogen bonds are formed between the atoms of residues i and j . The hydrogen bonding energy is calculated for both possible conformations and only the minimum of the two is used in $E_{beta-hb}(i,j)$.

$$E_{beta-hb}(i, j) = \begin{cases} \min\{\Phi(i \rightarrow j) + \Phi(j \rightarrow i), \Phi_p(i \rightarrow j) + \Phi_p(j \rightarrow i)\}, & \text{if strands are anti-parallel} \\ \min\{\Phi(i \rightarrow j+1) + \Phi(j-1 \rightarrow i), \Phi(i-1 \rightarrow j) + \Phi(j \rightarrow i+1)\}, & \text{if strands are parallel} \end{cases}$$

$\Phi(a \rightarrow d)$ is the directional energy calculation for a single hydrogen bond where a is the index of the acceptor residue

and d is the index of the donor residue. Three geometrical measures are used to estimate the strength of hydrogen bonds: the distance between the acceptor and the hydrogen atoms ($r_{a,O,d,H}$), the angle at the acceptor atom ($u_{a,C,a,O} \cdot u_{a,O,d,H}$), and the angle between the acceptor and donor atom vectors ($u_{a,C,a,O} \cdot u_{d,N,d,H}$). The distance and acceptor atom angle parameters are motivated by the orientation-dependent hydrogen bonding potential described in [42]. The following parameters were set based on idealized hydrogen bonding between β residues, with standard deviation values set such that two standard deviations approximate the cut-off in true hydrogen bonds. The ideal distance from hydrogen atom to accepting oxygen is $I_{hb-dist} = 1.9 \text{ \AA}$, with standard deviation $\sigma_{hb-dist} = 0.5 \text{ \AA}$. The ideal angle at the acceptor atom is 0° , so the ideal ($u_{a,C,a,O} \cdot u_{a,O,d,H}$) is $I_{acc-dp} = 1.0$, with standard deviation $\sigma_{acc-dp} = 0.11$. The ideal angle between the acceptor and donor atom vectors is 180° , so the ideal ($u_{a,C,a,O} \cdot u_{d,N,d,H}$) is $I_{acc-don-dp} = -1.0$, with standard deviation $\sigma_{acc-dp} = 0.15$. The parameters for pseudo-bonded residues are as follows: the ideal distance for $r_{a,O,d,H}$ is $I_{ps-hb-dist} = 7.9 \text{ \AA}$, $I_{ps-acc-dp} = -1.0$, and $I_{ps-acc-don-dp} = -1.0$. The standard deviations from the corresponding hydrogen bonding parameters above are used in $\Phi_{ps}(a \rightarrow d)$.

$$\Phi(a \rightarrow d) = \begin{cases} 54.0, & \text{if } r_{a,O,d,H} > 7.0 \\ \Psi(r_{a,O,d,H}, I_{hb-dist}, \sigma_{hb-dist}) \\ + \Psi(u_{a,C,a,O} \cdot u_{d,N,d,H}, I_{acc-don-dp}, \sigma_{acc-don-dp}) \\ + \Psi(u_{a,C,a,O} \cdot u_{a,O,d,H}, I_{acc-dp}, \sigma_{acc-dp}), & \text{otherwise} \end{cases}$$

$$\Phi_{ps}(a \rightarrow d) = \begin{cases} 54.0, & \text{if } r_{a,O,d,H} > 10.0 \\ \Psi(r_{a,O,d,H}, I_{ps-hb-dist}, \sigma_{ps-hb-dist}) \\ + \Psi(u_{a,C,a,O} \cdot u_{d,N,d,H}, I_{acc-don-dp}, \sigma_{acc-don-dp}) \\ + \Psi(u_{a,C,a,O} \cdot u_{a,O,d,H}, I_{ps-acc-dp}, \sigma_{acc-dp}), & \text{otherwise} \end{cases}$$

The penalty for the observed value (x) increases up to 6 standard deviations from the ideal value (μ).

$$\Psi(x, \mu, \sigma) = \begin{cases} \frac{(x-\mu)^2}{2\sigma^2}, & \text{if } |x - \mu| < 6\sigma \\ \frac{(6\sigma)^2}{2\sigma^2} = 18.0, & \text{otherwise} \end{cases}$$

All-Atom Energy Term Details

The all-atom energy terms depend on atom-atom interactions when all heavy atoms are included in the model. In the all-atoms energy equations x and y refer to atoms in the model and the residue positions are not referenced. The van der Waals radii and well-depths (ϵ_x used in $E_{LEN-JONES}$) come from the CHARMM19 parameter set [43]. The side-chain hydrogen bonding term, E_{SC-HB} , is described in detail here because it is unique to SELECTpro. The details of $E_{LEN-JONES}$, $E_{SOLVATION}$, and $E_{ELECTRO}$ are provided in the Appendix.

E_{SC-HB} : side-chain hydrogen bonding

E_{SC-HB} penalizes unsatisfied hydrogen bond donor and acceptor atoms that are at least partially buried. There is no penalty for fully exposed donor or acceptor atoms. Exposure percent ($\Delta G_x^{slv} \%$) is calculated as $\Delta G_x^{slv} / \Delta G_x^{ref}$. The definitions of ΔG_x^{slv} and ΔG_x^{ref} are provided in the description of $E_{SOLVATION}$ in the Appendix. Atoms at least 75% exposed are considered fully exposed and atoms less than 25% exposed are considered fully buried. For 25% $< \Delta G_x^{slv} \% < 75\%$ the penalty weight is reduced linearly from 1.0 at 25% to 0 at 75%. The ideal distance from the acceptor atom to donor atom is $I_{hb-da-dist} = 2.9 \text{ \AA}$. In the equations below $donors$ is the set of all side-chain hydrogen donor atoms and $acceptors$ is the set of all side-chain hydrogen acceptor atoms.

$$E_{SC-HB} = \sum_{x \in acceptors} E_{hb-acc}(x) + \sum_{x \in donors} E_{hb-don}(x)$$

$$E_{hb-acc}(x) = \lambda(x) \min_{y \in donors} |r_{x,y} - I_{hb-da-dist}|^2$$

$$E_{hb-don}(x) = \lambda(x) \min_{y \in acceptors} |r_{x,y} - I_{hb-da-dist}|^2$$

$$\lambda(x) = \begin{cases} 1 & \text{if } \Delta G_x^{slv} \% < .25 \\ 0 & \text{if } \Delta G_x^{slv} \% > .75 \\ 2(.75 - \Delta G_x^{slv} \%) & \text{otherwise} \end{cases}$$

Appendix

In the interest of completeness and reproducibility we include the details of the energy terms that are adapted from previous work.

Reduced Representation Energy Term Details

E_{BB-REP} : backbone repulsion

This term penalizes steric clashes between non-bonded atoms explicitly represented in the reduced representation. The penalty for overlapping atoms is the overlap distance squared as defined here:

$$E_{BB-REP} = \sum_{j>i} E_{bb-rep}(i, j)$$

$$E_{bb-rep}(i, j) = \sum_x \sum_y \Theta(i, x, j, y) (vdu_{x+y} - r_{i,x,j,y})^2$$

$$\Theta(i, x, j, y) = \begin{cases} 1, & \text{if } r_{i,x,j,y} < vdu_{x+y} \\ 0, & \text{otherwise} \end{cases}$$

E_{CT-REP} : centroid repulsion

A centroid-centroid repulsive term is used to reduce the overcrowding of side-chains in the reduced representation. The minimum distance between two centroids in

the calculation is the minimum observed for each pair of residue types - $D_{CT-min}(aa_i, aa_j)$ - in pdb_select25. The penalty for centroid-centroid overlaps is defined as the overlap distance squared:

$$E_{CT-REP} = \sum_{j>i} \Theta(i, j) [D_{CT-min}(aa_i, aa_j) - r_{i,CT,j,CT}]^2$$

$$\Theta(i, j) = \begin{cases} 1, & \text{if } r_{i,CT,j,CT} < D_{CT-min}(aa_i, aa_j) \\ 0, & \text{otherwise} \end{cases}$$

$E_{STAT-ENV}$: residue environment potential

The motivation for this term is to model the hydrophobic effect. The level of burial for each residue in the model is estimated by the number of other C_β atoms within 10 Å (the contact number N_i) [3]. The values in the table $\Omega_{stat-env}$ reflect the likelihood of observing a particular N_i for each residue type. For model residues near both termini the contact number is artificially increased to account for the missing neighbors along the chain.

$$E_{STAT-ENV} = \sum_i \Omega_{stat-env}(aa_i, N_i^*)$$

$$N_i^* = \begin{cases} N_i + 4 - i, & \text{if } i < 4 \\ N_i + 4 - |i - N|, & \text{if } |i - N| < 4 \\ N_i, & \text{otherwise} \end{cases}$$

$E_{STAT-PW-CI}$: context independent pair-wise interactions

This context independent pair-wise potential comes from Equation 6 of [3]. The potential considers the likelihood of observing the pair of centroids in a given distance bin relative to the background, with distance bins of < 5, 5-7, 7-10, 10-12, and > 12 Å. The advantage of a context independent pair-wise potential is that it is less vulnerable to over-fitting by a conformational search because of its generality.

$$E_{STAT-PW-CI} = \sum_{j>i} E_{stat-pw-ci}(i, j)$$

$$E_{stat-pw-ci}(i, j) = \Omega_{stat-pw-ci}[aa_i, aa_j, r_{bin}(i, j)]$$

$$r_{bin}(i, j) = \begin{cases} 0 - 5 & \text{if } r_{i,CT,j,CT} \leq 5.0 \\ 5 - 7 & \text{if } 5.0 < r_{i,CT,j,CT} \leq 7.0 \\ 7 - 10 & \text{if } 7.0 < r_{i,CT,j,CT} \leq 10.0 \\ 10 - 12 & \text{if } 10.0 < r_{i,CT,j,CT} \leq 12.0 \\ 12 + & \text{if } r_{i,CT,j,CT} > 12.0 \end{cases}$$

$E_{STAT-PW-CD}$: context dependent pair-wise potential

This context specific pair-wise potential is from [6]. This pair-wise potential depends on the local structure and relative orientation of both amino acids in the

interaction. The statistics are calculated independently for each combination of local structures and relative orientations. At each position the local structure is considered either compact or open and the relative orientation is determined by the dot product of the C_α to C_β unit vectors of each residue and divided into three classes: parallel, anti-parallel, and intermediate.

$$E_{STAT-PW-CD} = \sum_{j>i} E_{stat-pw-cd}(i, j)$$

$$E_{stat-pw-cd}(i, j) = \Theta(i, j) \Omega_{stat-pw-cd}[aa_i, aa_j, \lambda(i), \lambda(j), \Phi(i, j)]$$

$$\Theta(i, j) = \begin{cases} 1, & \text{if } r_{i,CT,j,CT} > D_{min,pw-cd}[aa_i, aa_j, \lambda(i), \lambda(j), \Phi(i, j)] \text{ and} \\ & r_{i,CT,j,CT} < D_{max,pw-cd}[aa_i, aa_j, \lambda(i), \lambda(j), \Phi(i, j)] \\ 0, & \text{otherwise} \end{cases}$$

$$\lambda(i) = \begin{cases} \text{compact,} & \text{if } r_{i-1,C_\alpha,i+1,C_\alpha} < 6.0 \\ \text{open,} & \text{otherwise} \end{cases}$$

$$\Phi(i, j) = \begin{cases} \text{parallel,} & \text{if } \mathbf{u}_{i,C_\alpha,i,C_\beta} \cdot \mathbf{u}_{j,C_\alpha,j,C_\beta} > .5 \\ \text{antiparallel,} & \text{if } \mathbf{u}_{i,C_\alpha,i,C_\beta} \cdot \mathbf{u}_{j,C_\alpha,j,C_\beta} < -.5 \\ \text{intermediate,} & \text{otherwise} \end{cases}$$

E_{ROG} : compactness

The radius of gyration is a simple measure of the global compactness of a domain. E_{ROG} penalizes models that are less compact than expected according to [44]. If the radius of gyration of the model (λ) is less than the expected value ($2.2N^{.38}$), there is no penalty. If it is greater, then the penalty is the squared difference between observed and expected. In the equation below $r_{i,mean}$ is the distance between the C_α of residue i and the mean of all C_α s in the model.

$$E_{ROG} = \Theta(\lambda - 2.2N^{.38})^2$$

$$\lambda = \sqrt{\frac{\sum r_{i,mean}^2}{N}}$$

$$\Theta = \begin{cases} 1, & \text{if } \lambda > 2.2N^{.38} \\ 0, & \text{otherwise} \end{cases}$$

All-Atom Energy Term Details

$E_{LEN-JONES}$: van der Waals forces

A fundamental characteristic of native globular protein structures is their efficient steric packing of atoms in the protein core. A Lennard-Jones 12-6 potential with damped repulsion ($E_{LEN-JONES}$) is used to measure the quality of steric packing. $E_{LEN-JONES}$ is the sum of local energy calculations $E_{len-jones}(x,y)$ performed on all pairs of non-bonded atoms. Since the repulsive portion of the standard Lennard-Jones 12-6 potential will overwhelm the entire energy function with a single significant atom-atom clash - repulsion is handled by a linear ramp from 0 to 10 as shown in the equation below [10]. Since $E_{len-jones}$

= 0 when $(vdw_{x,y}/r_{x,y}) = \sqrt[6]{2}$ independent of atom types, the switch to a linear ramp occurs when $(vdw_{x,y}/r_{x,y}) > \sqrt[6]{2}$.

$$E_{LEN-JONES} = \sum_{y>x} E_{len-jones}(x,y)$$

$$E_{len-jones}(x,y) = \begin{cases} 10.0 \left(1 - \frac{\sqrt[6]{2}}{vdw_{x,y}/r_{x,y}} \right), & \text{if } (vdw_{x,y}/r_{x,y}) > \sqrt[6]{2} \\ \sqrt{\epsilon_x \epsilon_y} \left[\left(\frac{vdw_{x,y}}{r_{x,y}} \right)^{12} - 2 \left(\frac{vdw_{x,y}}{r_{x,y}} \right)^6 \right], & \text{otherwise} \end{cases}$$

***E_{SOLVATION}*: solvation effects**

Solvation energy is calculated using the implicit solvation model described in [35] with the following adjustment: for overlapping atoms, the sum of their van der Waals radii is used in the calculation in place of the observed atom-atom distance in the model. This restricts the amount a single atom can contribute to the burial of another atom. Without this adjustment overlapping atoms will bias the calculation to indicate an atom is more buried than it would be otherwise. In the solvation model ΔG_x^{slv} is the observed solvation free energy of atom x in the model, calculated as the free energy of the fully exposed atom (ΔG_x^{ref}) minus the reduction in solvation caused by the surrounding atoms. ΔG_x^{free} was determined empirically by setting it equal to ΔG_x^{ref} and increasing its magnitude until ΔG_x^{slv} of deeply buried atoms became zero. λ_x is the correlation length of atom x . V_y is the volume neighboring atom y . The values of these parameters come from [3535], with the exception of ΔG_x^{ref} [45]. The equation for ΔG_x^{slv} below is the combination of Equations 5,6, and 7 of [35], with the atom overlap adjustment.

$$E_{SOLVATION} = \sum_x \Delta G_x^{slv}$$

$$\Delta G_x^{slv} = \Delta G_x^{ref} - \frac{\Delta G_x^{free}}{2\lambda_x \pi \sqrt{\pi}} \sum_{x \neq y} e^{-\left[\left(\frac{r_{x,y}^* - vdw_x}{\lambda_x} \right)^2 \right]} \frac{1}{r_{x,y}^{*2}} V_y$$

$$r_{x,y}^* = \begin{cases} vdw_{x+y}, & \text{if } r_{x,y} < vdw_{x+y} \\ r_{x,y}, & \text{otherwise} \end{cases}$$

***E_{ELECTRO}*: electrostatics**

Electrostatic interactions between charged atoms are treated by simple repulsion and attraction according to inverse distance squared. The use of distance squared rather than linear distance encourages the formation of salt bridges in the models. There is a correction for atom-atom distance below the minimum realistic value. The

ideal distance between oppositely charged atoms is $I_{hb-dist} = 2.75 \text{ \AA}$. In the equations below pos is the set of all positively charged atoms and neg is the set of all negatively charged atoms.

$$E_{ELECTRO} = \sum_{y>x \in pos \cup neg} \Theta(x)\Theta(y)/r_{x,y}^{*2}$$

$$\Theta(x) = \begin{cases} 1, & \text{if } x \in pos \\ -1, & \text{if } x \in neg \end{cases}$$

$$r_{x,y}^* = \begin{cases} I_{e-dist}, & \text{if } r_{x,y} < I_{e-dist} \\ r_{x,y}, & \text{otherwise} \end{cases}$$

Availability and requirements

- **Project home page:** <http://www.igb.uci.edu/~baldig/selectpro.html>
- **Operating system:** linux for stand alone version, server is platform independent
- **Programming language:** C++ and Perl
- **Software requirements:** Perl
- **Disk space requirements:** 1.6 Gb for full version, 13 Mb without feature predictors

Authors' contributions

AR and PB designed the novel energy terms. AR implemented the methods and carried out the experiments. AR and PB authored the manuscript. Both authors approved the manuscript.

Acknowledgements

Work supported by NIH grant LM-07443-01, NSF grants EIA-0321390 and IIS-0513376, and a Microsoft Faculty Research Award to PFB.

References

1. Wallner B and Elofsson A: **Prediction of global and local model quality in CASP7 using Pcons and ProQ.** *Proteins* 2007, **69** (Suppl 8):184-193.
2. Cozzetto D and Tramontano A: **Relationship between multiple sequence alignments and quality of protein comparative models.** *Proteins* 2005, **58**:151-157.
3. Simons KT, Kooperberg C, Huang E and Baker D: **Assembly of protein tertiary structures from fragments with similar local sequences using simulated annealing and Bayesian scoring functions.** *J Mol Biol* 1997, **268**:209-225.
4. Kihara D, Lu H, Kolinski A and Skolnick J: **TOUCHSTONE: An ab initio protein structure prediction method that uses threading-based tertiary restraints.** *Proc Natl Acad Sci USA* 2001, **98**:10125-10130.
5. Boniecki M, Rotkiewicz P, Skolnick J and Kolinski A: **Protein fragment reconstruction using various modeling techniques.** *J Comput Aided Mol Des* 2003, **17**:725-738.
6. Kolinski A: **Protein modeling and structure prediction with a reduced representation.** *Acta Biochim Pol* 2004, **51**:349-371.
7. Sanchez R and Sali A: **Comparative protein structure modeling. Introduction and practical examples with modeller.** *Methods Mol Biol* 2000, **143**:97-129.

8. Qian B, Ortiz A and Baker D: **Improvement of comparative model accuracy by free-energy optimization along principal components of natural structural variation.** *Proc Natl Acad Sci USA* 2004, **101**:15346–15351.
9. Lazaridis T and Karplus M: **Discrimination of the native from misfolded protein models with an energy function including implicit solvation.** *J Mol Biol* 1999, **288**:477–487.
10. Kuhlman B and Baker D: **Native protein sequences are close to optimal for their structures.** *Proc Natl Acad Sci USA* 2000, **97**:10383–10388.
11. Vorobjev Y and Hermans J: **Free energies of protein decoys provide insight into determinants of protein stability.** *Protein Sci* 2001, **10**:2498–2506.
12. Felts A, Gallicchio E, Wallqvist A and Levy R: **Distinguishing native conformations of proteins from decoys with an effective free energy estimator based on the OPLS all-atom force field and the Surface Generalized Born solvent model.** *Proteins* 2002, **48**:404–422.
13. Dominy B and Brooks C: **Identifying native-like protein structures using physics-based potentials.** *J Comput Chem* 2002, **23**:147–160.
14. Oldziej S, Czaplowski C, Liwo A, Chinchio M, Nianis M, Vila JA, Khalili M, Arnautova YA, Jagielska A, Makowski M, Schafroth HD, Kazmierkiewicz R, Ripoll DR, Pillardy J, Saunders JA, Kang YK, Gibson KD and Scheraga HA: **Physics-based protein-structure prediction using a hierarchical protocol based on the UNRES force field: Assessment in two blind tests.** *Proc Natl Acad Sci USA* 2005, **102**:7547–7552.
15. Shortle D, Simons KT and Baker D: **Clustering of low-energy conformations near the native structures of small proteins.** *Proc Natl Acad Sci USA* 1998, **95**:11158–11162.
16. Simons KT, Ruczinski I, Kooperberg C, Fox BA, Bystroff C and Baker D: **Improved recognition of native-like protein structures using a combination of sequence-dependent and sequence-independent features of proteins.** *Proteins* 1999, **34**:82–95.
17. Vendruscolo M, Najmanovich R and Domany E: **Can a pairwise contact potential stabilize native protein folds against decoys obtained by threading?** *Proteins* 2000, **38**:134–148.
18. Cozzetto D, Kryshchafovich A, Ceriani M and Tramontano A: **Assessment of predictions in the model quality assessment category.** *Proteins* 2007, **69**:175–183.
19. Wu S, Skolnick J and Zhang Y: **Ab initio modeling of small proteins by iterative TASSER simulations.** *BMC Biol* 2007, **5**:17.
20. Zhang 2007 Decoy Sets. <http://zhang.bioinformatics.ku.edu/~TASSER/decoys/>.
21. Zhang Y and Skolnick J: **SPICKER: A clustering approach to identify near-native protein folds.** *J Comput Chem* 2004, **25**:865–871.
22. Wallner B, Fang H and Elofsson A: **Automatic consensus-based fold recognition using Pcons, ProQ, and Pmodeller.** *Proteins* 2003, **53**(Suppl 6):534–541.
23. Lundstrom J, Rychlewski L, Bujnicki J and Elofsson A: **Pcons: a neural-network-based consensus predictor that improves fold recognition.** *Protein Sci* 2001, **10**:2354–2362.
24. Wallner B and Elofsson A: **Can correct protein models be identified?** *Protein Sci* 2003, **12**:1073–1086.
25. SCRATCH Cheng J, Randall AZ, Sweredoski M and Baldi P: **SCRATCH: a protein structure and structural feature prediction server.** *Nucleic Acids Res* 2005, **33** Web Server: W72–W76.
26. Moulton J, Fidelis K, Kryshchafovich A, Rost B, Hubbard T and Tramontano A: **Critical assessment of methods of protein structure prediction-Round VII.** *Proteins* 2007, **69**(Suppl 8):3–9.
27. Zemla A, Veclovas C, Moulton J and Fidelis K: **Processing and analysis of CASP3 protein structure predictions.** *Proteins* 1999, **37**(Suppl 3):22–29.
28. Sali A and Blundell TL: **Comparative protein modeling by satisfaction of spatial restraints.** *J Mol Biol* 1993, **234**:779–815.
29. Martin-Renom MA, Stuart A, Fiser A, Sanchez R, Melo F and Sali A: **Comparative protein structure modeling of genes and genomes.** *Annu Rev Biophys Biomol Struct* 2000, **29**:291–325.
30. Fiser A, Do RK and Sali A: **Modeling of loops in protein structures.** *Protein Sci* 2000, **9**:1753–1773.
31. Tsai J, Bonneau R, Morozov AV, Kuhlman B, Rohl CA and Baker D: **An Improved Protein Decoy Set for Testing Energy Functions for Protein Structure Prediction.** *Proteins* 2003, **53**:76–87.
32. Baker D, Bystroff C, Fletterick RJ and Agard DA: **PRISM: topologically constrained phased refinement for macromolecular crystallography.** *Acta Crystallogr D Biol Crystallogr* 1993, **49**:429–39.
33. Sun S: **Reduced representation approach to protein tertiary structure prediction: statistical potential and simulated annealing.** *J Theor Biol* 1995, **172**:13–32.
34. Canutescu AA, Shelenkov AA and Dunbrack RL: **A graph-theory algorithm for rapid protein side-chain prediction.** *Protein Sci* 2003, **12**:2001–2014.
35. Lazaridis T and Karplus M: **Effective Energy Function for Proteins in Solution.** *Proteins* 1999, **35**:133–152.
36. Hobohm U and Sander C: **Enlarged representative set of protein structures.** *Protein Sci* 1994, **3**:522–524.
37. Berman HM, Westbrook J, Feng Z, Gilliland G, Bhat TN, Weissig H, Shindyalov IN and Bourne PE: **The Protein Data Bank.** *Nucleic Acids Res* 2000, **28**:235–242.
38. Pollastri G, Przybylski D, Rost B and Baldi P: **Improving the prediction of protein secondary structure in three and eight classes using recurrent neural networks and profiles.** *Proteins* 2002, **47**:228–235.
39. Pollastri G, Baldi P, Fariselli P and Casadio R: **Prediction of coordination number and relative solvent accessibility in proteins.** *Proteins* 2002, **47**:142–153.
40. Baldi PF and Pollastri G: **The principled design of large-scale recursive neural network architectures—DAG-RNNs and the protein structure prediction problem.** *J Mach Learn Res* 2003, **4**:575–602.
41. Pollastri G and Baldi P: **Prediction of contact maps by GIOHMMs and recurrent neural networks using lateral propagation from all four cardinal corners.** *Bioinformatics* 2002, **18**:S62–S70.
42. Kortemme T, Morozov AV and Baker D: **An orientation-dependent hydrogen bonding potential improves prediction of specificity and structure for proteins and protein-protein complexes.** *J Mol Biol* 2003, **326**:1239–1259.
43. Neria E, Fischer S and Karplus M: **Simulation of activation free energies in molecular systems.** *J Chem Phys* 1996, **105**:1902–1921.
44. Skolnick J, Kolinski A and Ortiz AR: **MONSSTER: A method for folding globular proteins with a small number of distance restraints.** *J Mol Biol* 1997, **265**:217–241.
45. Privalov PL and Makhatadze GI: **Contribution of hydration to protein folding thermodynamics II. The entropy and Gibbs energy of hydration.** *J Mol Biol* 1993, **232**:660–679.

Publish with **BioMed Central** and every scientist can read your work free of charge

"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."

Sir Paul Nurse, Cancer Research UK

Your research papers will be:

- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- yours — you keep the copyright

Submit your manuscript here:
http://www.biomedcentral.com/info/publishing_adv.asp

