

## COMMUNICATION

## Rosetta Custom Score Functions Accurately Predict $\Delta\Delta G$ of Mutations at Protein-Protein Interfaces Using Machine Learning

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Sumant Shringari,<sup>‡a</sup> Sam Giannakoulias,<sup>‡a</sup> John J. Ferrie,<sup>\*a</sup> and E. James Petersson<sup>\*a</sup>

**Protein-protein interfaces play essential roles in a variety of biological processes and many therapeutic molecules are targeted at these interfaces. However, accurate predictions of the effects of interfacial mutations to identify “hotspots” have remained elusive despite the myriad of modeling and machine learning methods tested. Here, for the first time, we demonstrate that nonlinear reweighting of energy terms from Rosetta, through the use of machine learning, exhibits improved predictability of  $\Delta\Delta G$  values associated with interfacial mutations.**

Protein-protein interactions mediate many essential biological processes. Cellular signaling, spatial and temporal regulation, and metabolism are deeply rooted in the formation of higher order protein quaternary structures.<sup>1</sup> Complex formation is governed by the complementary structural and chemical features displayed by residues at the protein-protein interface, and mutations of these residues are highly correlated with dysfunction and disease.<sup>2</sup> Moreover, the development of new biomaterials and catalysis strategies are largely dependent on the binding affinity of the involved protein partners.<sup>3, 4</sup> Therefore, a computational model capable of rapidly and accurately predicting the energy differences ( $\Delta\Delta G$ s) associated with mutations would aid in the identification of protein-protein hotspots, providing insight in disease and design.<sup>5, 6</sup>

To date, several approaches have been developed towards the accurate prediction of protein  $\Delta\Delta G$  values. These include the use of statistical and contact potentials<sup>7-10</sup>, design of novel sampling schemes,<sup>11, 12</sup> generation of weighted energy or score functions,<sup>13-16</sup> and employment of supervised machine learning techniques.<sup>17-21</sup> Additionally, within the Rosetta Modelling Suite, new sampling schemes, designed to mimic protein

motions observed in solution, have afforded increased predictive accuracy.<sup>11, 22</sup> Though these methods have shown some notable success, there is still a need for a single, generalizable, and facile approach capable of accurately predicting  $\Delta\Delta G$ ’s of mutations at protein/protein interfaces.

To this end, we envisioned that reweighting of energy terms from Rosetta through machine learning will provide a platform with improved  $\Delta\Delta G$  prediction accuracy. The full-atom score function in Rosetta has been repeatedly improved through the introduction of new energy terms and optimization of term weighting. Although Rosetta-based simulations can generate accurate structural models, correlations between the canonical score functions and experimental data remain relatively poor<sup>23</sup>. This suggests that while the underlying set of terms may produce models with small RMSDs relative to experimental structures, energetically, they require differential weightings for specific applications like  $\Delta\Delta G$  prediction. Therefore, we designed the first reported Custom Score Function (CSF, named SRS2020), which is a score function devised purely through the reweighting of Rosetta energy terms for optimal prediction of an experimentally measurable variable of interest. This method allows for the traditional Rosetta score function to be used for structural refinement, while SRS2020 can be used to more accurately predict  $\Delta\Delta G$ s. This notion is not entirely novel as protein and small molecule design strategies in Rosetta have used supplementary criteria in the form of classifiers and filters to perform selection based on criteria not encompassed within scoring<sup>24</sup>. However, the approach presented herein is simpler in that it requires no additional terms to be constructed.

To test the utility of this approach, we focused on simulating the SKEMPI 2.0 database (<https://life.bsc.es/pid/skempiz/>) which is the largest curated database of protein-protein interfacial mutants. This database includes 348 different protein-protein complexes, and  $\Delta\Delta G$  values for 6193 unique interfacial mutations of this protein set.<sup>25, 26</sup> After removing complexes where the mutation location could not be accurately assigned due to ambiguities between the number of protein subunits and the generalized version of our computational protocol

<sup>a</sup> Department of Chemistry, University of Pennsylvania, 231 South 34th Street, Philadelphia, PA 19104, USA.

<sup>‡</sup> These authors contributed equally.

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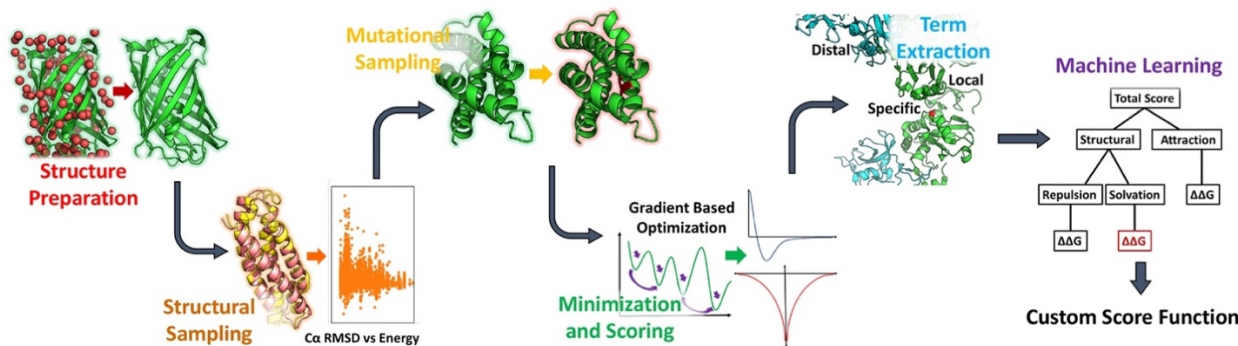


Fig. 1 Schematic of computational workflow for developing a custom score function to predict  $\Delta\Delta G$  values of mutations at protein/protein interfaces.

employed within PyRosetta.<sup>27</sup> First, wild-type complexes are cleaned (removal of solvent, ligands, or ions), renumbered, and subjected to initial minimization. Subsequently, mutations are introduced, and both the wild-type and mutant complexes are subjected to another round of structural optimization followed by the computation of Rosetta energies which in turn are fed into a variety of machine learning protocols. Sampling was varied both in the initial, structural stage where only wild-type complexes were considered, as well as in the mutational stage where both the wild-type and mutants were sampled under a uniform scheme. In the structural sampling stage, we assessed the impact of relaxing the input structure compared to using the structure directly from the SKEMPI 2.0 server. Additionally, we tested the impact of local and global sampling during the mutational sampling stage, where either only the mutant residue was packed, or repacking was performed on the entire complex. Lastly, we computed energies for each sampling combination using both REF2015,<sup>28</sup> the most recently published score function, and BETA\_NOV16,<sup>29</sup> the newest score function available in Rosetta.

We first focused on assessing the performance of the traditional Rosetta score function in predicting mutational  $\Delta\Delta G$ s from the SKEMPI 2.0 database. As expected, differences in sampling impacted the correlation of Rosetta total energy

scores with  $\Delta\Delta G$ . Initial minimization of wild-type complexes, prior to mutational sampling, was found to improve correlation between total Rosetta Energy Units (REUs), and experimental  $\Delta\Delta G$  values for both the REF2015 and BETA\_NOV16 score functions. This was unsurprising as it is widely recognized that structures determined from crystallographic data require initial relaxation prior to sampling within Rosetta to produce more correlative simulations.<sup>30</sup> Although we observe an improvement in the predictive capacity following minimization during initial structural sampling, which is likely due to the approximate five REU reduction in the average residue score, we observe only a minimal change in conformation (ESI, Fig. S1).

Across all sampling and scoring schemes, we see a maximum average RMSD of 0.45 Å compared to input structures from SKEMPI 2.0. Additionally, score values derived from local packing only at the mutation site prior to minimization showed a higher predictive capacity than global repacking following mutation. A Wilcoxon t-test was performed to identify Rosetta energy terms that differed between these simulations. The difference in predictability between these models is likely due to the drastic differences in Lennard-Jones, Dunbrack, and solvation terms produced by these simulations (see ESI). Lastly, it is notable that the most recent score function, BETA\_NOV16, afforded a higher correlation with experimental data than the

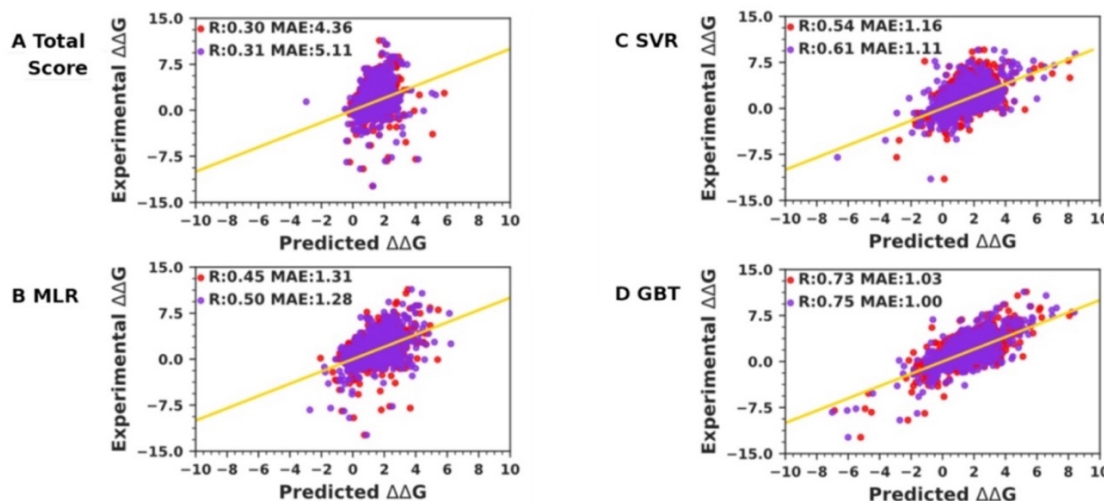


Fig. 2 Models for predicting interfacial  $\Delta\Delta G$ , REF2015 (red), BETA\_NOV16 (purple). Rosetta total score in Rosetta energy units, or REU (A), multiple linear regression (MLR) of Rosetta energy terms (B), polynomial support vector regression (SVR, C), and gradient boosted random forest (GBT) regression (D). MAE: Mean absolute error in kcal/mol.

benchmarked REF2015 score function. This may be due to increased accuracy in weightings or the additional terms added in the BETA\_NOV16 score function.

We sought to improve the predictive power of the models generated by both the REF2015 and BETA\_NOV16 score functions using machine learning. Given the results of the structure preparation comparisons, we elected to focus exclusively on improving the predictability of simulations where structures are initially relaxed and only locally sampled following mutation. Multiple linear regression (MLR) is a basic machine learning technique that uses several explanatory variables to predict a single output. Here, those variables are the Rosetta energy terms generated during simulation which will be reweighted to optimize the correlation to experimental  $\Delta\Delta G$ . As illustrated in Fig. 2, even this simple MLR approach (Fig. 2B) results in an improved correlation with experimental data compared to the traditional score functions (Fig. 2A). Using both five and ten-fold cross validation, we determined that using an MLR improved testing set correlation by a factor of at least 1.58 (ESI, Table S1, S2). We also observed, again, that application of the BETA\_NOV16 score function displayed a higher predictability than the REF2015 score function. Interestingly, the most important terms in the MLR scoring correlated well with the terms that differed between sampling schemes (ESI, Table S3). While this improvement was encouraging, the usability of this reweighted score function is still poor as the Pearson correlation coefficient was only 0.51 and the mean absolute error (MAE) was 1.28 kcal/mol.

In order to further improve our  $\Delta\Delta G$  predictions, additional inputs were considered, as well as the introduction of more complex machine learning algorithms. Inputs were extracted from simulated structures as the change in energy following sampling. Several different values were computed to capture global vs. specific and local vs. distal differences. Global terms correspond to the change of the total values of the decomposed Rosetta energy terms across the whole mutant and wild-type complexes. Specific values refer directly to the differences between the decomposed Rosetta energy terms of only the mutated and wild-type residues. To distinguish local and distal terms, an 8 Å contacting shell was created around the mutation site. This was used for the calculation of local and distal terms, which correspond to the change in total decomposed scores within or beyond this sphere. In addition to alternative inputs, we employed more sophisticated machine learning algorithms: Kernel Ridge Linear Regression (KRR, see S10, S11), support vector regressions (SVRs), and Gradient Boosted Random Forrest Regression (GBT). Our reasons for choosing these methods are outlined in ESI. Training and testing sets were specifically designed to ensure that no mutational redundancy existed between the sets. Curation of training and testing sets in this manner allows for the greatest predictive power of generated models<sup>31</sup>. Additionally, more rigorous investigation describing the robustness of our models as a function of training and testing sets is found in ESI Table S9.

SVRs were performed using various kernels, including Polynomial (degrees 2, 3, 4, and 5), radial base function, and Sigmoid. Using these algorithms, correlation to  $\Delta\Delta G$  from

**Table 1.** Machine Learning Models Using the SKEMPI Database

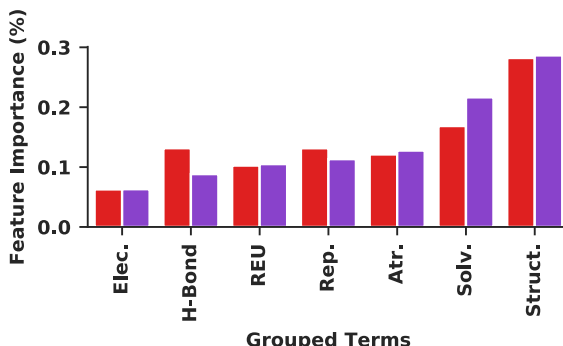
Method	R Value	MAE
FoldX	0.34	1.33
Pred1	0.45	1.14
BeAtMuSiC	0.46	1.09
Pred2	0.54	1.07
MLR	0.31	1.18
SVR	0.53	1.09
<b>SRS2020</b>	<b>0.65</b>	<b>0.92</b>

Alternative methods for predicting  $\Delta\Delta G$  using the SKEMPI database. R value is the Pearson correlation coefficient and MAE is mean absolute error in kcal/mol.

similarly to many other literature models.<sup>14</sup> (Fig. 2C and Table 1) SVR analysis also demonstrated that the BETA\_NOV16 score function performed better than the REF2015 score function (Fig. 2C). For GBT, we found that after an exhaustive grid search of tuneable parameters, this technique was the most predictive of all models tested as it produced the highest correlation as well as lowest MAE. Interestingly, GBT models were invariant to which Rosetta score function was used for simulation as BETA\_NOV16 and REF2015 score functions tested identically.

To further identify any potential differences between these two models, feature importance analysis was performed. The two models were extremely similar with the only notable differences coming from a slight enrichment in the importance of solvation and hydrogen bonding terms in the BETA and REF GBT models respectively. In both models, terms corresponding to phi-psi or rotameric preferences (Fig. 3, Struct. category) were found to be most important. These terms were followed by solvation (Solv.), van der Waals (Atr. and Rep.), the single value of Rosetta total energy (REU), electrostatic (Elec.), and hydrogen bonding terms (H-Bond). Considering that the database is primarily comprised of mutations to alanine, the typical reduction in size and increase in hydrophobicity associated with these changes likely explains the importance of solvation and nonpolar, attractive interactions over hydrogen bonding or electrostatics. Further analysis of how SRS2020 predicts specific subsets of SKEMPI2.0 data is found in ESI.

After identifying that the GBT-based CSF derived from BETA\_NOV16 optimized structures was our best model, we have named it SRS2020 (information on using this code can be found on our github [https://github.com/Sam-Giannakoulias/RML\\_ddG/](https://github.com/Sam-Giannakoulias/RML_ddG/) or automated prediction can be performed with our Jupyter web app deposited there. Table 1 compares our results to other machine learning models utilizing a subset of the SKEMPI database to predict  $\Delta\Delta G$ . All values in this table represent five-fold cross validation of the exact same subset of single alternative methods and are thus directly comparable. In addition to these baseline comparisons, a more extensive set of benchmarking, including comparable performance against standard hold-out sets such as S487 can be found in Tables S10-16<sup>31-39</sup>. SRS2020 represents not only the first reported CSF but is the most accurate predictor of  $\Delta\Delta G$  in both correlation and MAE, highlighting the potential utility of CSF-based approaches. SRS2020 was trained and regressed from the largest protein interaction data set available in the literature and has proven robust to alterations in sampling and scoring, demonstrating the strength of CSF approaches for specific applications.



**Fig. 3** Important features in GBT models derived from REF2015 (red) and BETA\_Nov16 (purple). Feature importance (%) determined as described in ESI.

Furthermore, while our algorithm may be slower than GPU accelerated approaches, the freedom from sampling optimization removes the need to find the perfect simulation and results in an incredibly rapid approach. Prediction of the  $\Delta\Delta G$  values upon mutation for interfacial residues can be computed in <60 s on a single intel core i5 8<sup>th</sup> generation CPU.

We intend to extend this methodology to encompass more complex sampling methods, such as the ensemble-based Backrub approach.<sup>33</sup> Although this CSF contains no additional energy terms or metrics, one can also easily introduce a variety of bioinformatics terms to further strengthen these models. Additionally, even more complex machine learning methods such as Extreme Gradient Boosted Random Forrest Regressions (XGBoost) or neural networks (NNs)<sup>32</sup> may be employed to further improve  $\Delta\Delta G$  prediction. Finally, we hope to extend the SRS2020 model beyond the prediction of interfacial  $\Delta\Delta G$  and use it to design protein-protein interfaces as well as peptides or peptidomimetics targeting such interfaces.

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## Conflicts of interest

“There are no conflicts to declare”.

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## Correction: Rosetta custom score functions accurately predict $\Delta\Delta G$ of mutations at protein–protein interfaces using machine learning

Sumant R. Shringari, † Sam Giannakoulias, † John J. Ferrie \* and E. James Petersson \*

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