Subject Section

SAAMBE-SEQ: A Sequence-based Method for Predicting Mutation Effect on Protein-protein Binding Affinity

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Abstract

Motivation: Vast majority of human genetic disorders are associated with mutations that affect protein-protein interactions by altering wild type binding affinity. Therefore, it is extremely important to assess the effect of mutations on protein-protein binding free energy to assist the development of therapeutic solutions. Currently the most popular approaches use structural information to deliver the predictions, which precludes them to be applicable on genome-scale investigations. Indeed, with the progress of genomic sequencing, researchers are frequently dealing with assessing effect of mutations for which there is no structure available.

Results: Here we report a Gradient Boosting Decision Tree (GBDT) machine learning algorithm, the SAAMBE-SEQ, which is completely sequence-based and does not require structural information at all. SAAMBE-SEQ utilizes 80 features representing evolutionary information, sequence-based features and change of physical properties upon mutation at the mutation site. The approach is shown to achieve Pearson correlation coefficient (PCC) of 0.83 in 5-fold cross validation in a benchmarking test against experimentally determined binding free energy change (ΔΔG). Further a blind test (no-STRUC) is compiled collecting experimental ΔΔG upon mutation for protein complexes for which structure is not available and used to benchmark SAAMBE-SEQ resulting in PCC in the range of 0.37 to 0.46. The accuracy of SAAMBE-SEQ method is found to be either better or comparable to most advanced structure-based methods. SAAMBE-SEQ is very fast, available as webserver and stand-alone code, and indeed utilizes only sequence information, and thus it is applicable for genome-scale investigations to study the effect of mutations on protein-protein interactions.

Availability: SAAMBE-SEQ is available at http://compbio.clemson.edu/saambe_webserver/indexSEQ.php#started.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Mutations introduce diversity in genome that can be either advantageous or cause diseases. Their effect on molecular level is manifested as alterations of wild type properties of the corresponding macromolecules such as proteins, DNAs and RNAs (Kucukkal, et al., 2015; Petukh, et al., 2015). Of particular interest is the effect of mutations on protein-protein interactions, since protein-protein interactions are essential for a wide
range of cellular processes such as signal transductions, cell metabolism, regulation of gene expression, transport, and muscle contractions (Bustin, 2015; Jones and Thornton, 1996; Keskin, et al., 2008). Therefore, understanding the effect of mutations on protein-protein interactions at molecular level is crucial for protein engineering (Orii and Ganapathiraju, 2012), developing novel therapeutics (Pett, et al., 2016; Wells and McClendon, 2007) and revealing molecular mechanism of diseases (Kuzmanov and Emili, 2013; Nibbe, et al., 2011). This prompted numerous investigations, both experimental (Fragoza, et al., 2019) and computational (Das, et al., 2012), explore the impact of mutations on protein-protein interactions.

Computational methods for predicting the effect of mutations on protein-protein binding energy are alternative to experimental techniques, since they are less time consuming and do not require biochemical work to prepare the samples. Because of that, various computational methods (described below) were developed, however, most of them require structural information. This is a severe limitation for genome-scale approaches, since it is estimated that only about 6.5% of known human interactome has structural information (Mosca, et al., 2013).

Among various computational methods, some are based on physical energy-based features or knowledge-based features, some use machine learning algorithms others linear combination of energy terms. For example, FoldX(Guerois, et al., 2002; Schymkowitz, et al., 2005), a machine learning method uses physical energies such as van der Waals, electrostatic energy, hydrogen bond and solvation energy. Additionally, this method considers conformational changes of side chains using rotamer approach. In addition to physical energy, knowledge-based energy terms were also used to determine $\Delta \Delta$G. For example, SAAMBE(Petukh, et al., 2016; Petukh, et al., 2015) uses combination of MM/PBSA and knowledge-based energy terms. The specialty of SAAMBE is that it uses amino acid specific dielectric constants to mimic the conformational flexibility caused by mutation. BindProfX(Xiong, et al., 2017) is another method, which combines conservation profile with the FoldX to improve the prediction of $\Delta \Delta$G. In 2018, a statistical energy based $\Delta \Delta$G predictor based on a coarse-grained model, the BeAtMuSiC(Dehouck, et al., 2013), was developed. All these methods based on either physical energy or knowledge based potential or combination of both were reported to achieve Pearson correlation coefficient (PCC) ranging from 0.38 to 0.68 as benchmarked on SKEMPI v1.1 database(Moaal and Fernández-Recio, 2012).

In recent years, several machine learning-based methods have been developed with structure-based features to predict $\Delta \Delta$G upon mutations. The first developed machine learning based predictor is mCSM(Pires, et al., 2013), which uses atomic distance pattern surrounding the mutation site to represent the neighboring environment and achieved a high correlation of 0.80 on 2317 single mutations from SKEMPI v1.1 database. Recently published iSEE(Geng, et al., 2019) method is based on 31 features involving position specific scoring matrix, structure interface profile and energy-based features and utilizes a random forest model to predict $\Delta \Delta$G caused by a given mutation. iSEE achieved a high correlation of 0.8 on single mutations in dimeric complexes from SKEMPI v1.1. MutaBind (Li, et al., 2016) is another predictor, which obtained a correlation of 0.68 on the single point mutations in SKEMPI 1.1. Recently, MutaBind2 (Zhang, et al., 2020) was developed and reported to achieve PCC of 0.82 against experimental $\Delta \Delta$G from SKEMPI v2.0(Jankauskaitė, et al., 2019). It is important to mention that MutaBind2 can predict $\Delta \Delta$G caused by multiple mutations as well. BindProfX(Xiong, et al., 2017) combines its interface profile with the FoldX score to improve the prediction of $\Delta \Delta$G using random forest model and achieved PCC of 0.74 on 1131 single mutations from SKEMPI v1.1. However, BindProfX can only predicts $\Delta \Delta$G for mutations located at the interface of the protein complexes. Recently, an improved version of mCSM method, called mCSM-PPI2(Rodrigues, et al., 2019), was reported. In mCSM-PPI2 method, the graph-based signature framework of mCSM is combined with additional inter-residue complex network, evolutionary information and energetic terms. Another recent innovative algorithm is TopNetTree(Wang, et al., 2020), which integrates topological features and a deep learning algorithm, represented by a topology-based network tree. The method achieved a PCC of 0.82 on single mutations from SKEMPI v2.0 database. Our recently published method, SAAMBE-3D(Pahari, et al., 2020) is a structure-based machine learning algorithm, which utilizes several knowledge-based features representing the physical environment surrounding mutation site. SAAMBE-3D is the fastest method available so far for predicting $\Delta \Delta$G caused by single mutation and comes as stand-alone code as well. Moreover, in addition to predicting $\Delta \Delta$G, the method predicts whether the mutation is disruptive or non-disruptive, which enables identification of disease-causing mutations.

The important thing to note here is that all the above-mentioned methods require a 3D structure of the protein complex as input to predict $\Delta \Delta$G upon mutations. However, as mentioned above, only 6.5% of known human interactome has structural information(Mosca, et al., 2013). Therefore, the applicability of these structure-based methods is limited. A partial solution that can extend their applicability is to predict the structures of protein complexes from sequence by using homology modeling. However, generating high-quality 3D structures is not always possible which makes the predictions much less accurate. Therefore, it is crucial to develop a method, which can predict $\Delta \Delta$G caused by mutations using only sequence information.

Currently, there is only one sequence-based method, the ProAffiMuSeq(Jemimah, et al., 2019), which takes as input the sequence of two interacting chains. However, ProAffiMuSeq is intended to only predict $\Delta \Delta$G caused by mutations located at the interfaces of the protein-protein complexes, and thus still requires structural information. Our attempt to use it for predicting $\Delta \Delta$G for non-interfacial mutations resulted in negative PCC as benchmarked against experimental data (see results section below). The ProAffiMuSeq is a machine learning based method, achieves PCC of 0.75 in benchmarking test (90% training and 10% testing sets) taken from 1173 interfacial mutations in protein-protein complexes from PROXIMATE database (Jemimah, et al., 2017).

Here, we report a new development of SAAMBE, the SAAMBE-SEQ, which is a truly sequence-based machine learning algorithm to predict the binding affinity changes upon single mutation in protein-protein complexes. Unlike other existing methods, SAAMBE-SEQ does not either require a 3D complex structure as input or knowledge of interfacial residues. Therefore, this method can be applied to protein complexes without known structure. The prediction of $\Delta \Delta$G using SAAMBE-SEQ is found to be either more accurate or comparable to leading structure-based methods. The method is available as a webserver as well as stand-alone code. SAAMBE-SEQ utilizes 80 features representing evolutionary information using position specific scoring matrix, sequence-based features and change in some physical properties of mutation site. SAAMBE-SEQ is trained on 2398 single point mutations from 200 complexes taken from SKEMPI v2.0. The method
SAAMBE-SEQ

uses the Gradient Boosting Decision Tree (GBDT) machine learning algorithm and achieves a PCC of 0.83. Furthermore, SAAMBE-SEQ is also trained to discriminate disruptive from non-disruptive mutations and achieves accuracy of 0.81, precision of 0.65, sensitivity and specificity of 0.81 as benchmarked against Cornell University dataset(Fragoza, et al., 2019; Pahari, et al., 2020).

2 Methods

2.1 Dataset creation

The amino acid sequences and experimentally measured binding free energies in this work were taken from the recently updated version of SKEMPI, SKEMPI v2.0 database(Jankauskaite, et al., 2019), which compiles experimentally measured binding affinity values for wild-type as well as mutant protein-protein complexes. SKEMPI 2.0 contains binding affinity data for 7085 mutations from 389 protein complexes. Only cases of single point mutations from dimeric complexes were considered, resulting in 2446 mutations from 207 different protein-protein complexes. Then, the binding free energy (\( \Delta G \)) was calculated from the binding affinity:

\[
\Delta G = RT \ln(K_D)
\]

where \( R \) is the ideal gas constant, \( T \) is temperature in Kelvin and \( K_D \) is binding affinity of the given protein complex. The \( \Delta G \) is calculated for both wild type and mutant protein complexes. Then, the change in binding free energy upon mutation (\( \Delta \Delta G \)) is calculated by subtracting \( \Delta G \) for wild type from that of mutant

\[
\Delta \Delta G_{\text{Mutant}} - \text{Wild} - \text{type} = \Delta G_{\text{Mutant}} - \Delta G_{\text{Wild} - \text{type}}
\]

For some mutations, multiple measurements were performed, and all the measured binding affinity values were reported in SKEMPI v2.0 database. If the standard deviation of \( \Delta \Delta G \) for a particular mutation is less than 1.0 kcal mol\(^{-1}\), we considered those cases and used average value for developing and benchmarking our model. We removed all mutations with standard deviation greater than 1 kcal mol\(^{-1}\). Further, we removed the complexes for which any chain contains less than 20 amino acid residues. Therefore, the final compiled dataset consists of 2398 single point mutations from 200 different dimeric complexes.

2.2 Model development

Our methodology of predicting binding free energy changes due to mutation in protein complexes incorporates only sequence-based features. Our machine learning model is based on GBDT algorithm. We used Overall, we used 80 features which include average Position Specific Scoring Matrix (PSSM) for mutant and interaction chain, conservation score at mutation site, change in molar volume, hydrophobicity, flexibility, hydrogen bonds, polarity, mutation type, chemical nature, and size of the mutated amino acid. Label encoding method is used for incorporating mutation type, change in polarity, chemical properties, hydrogen bond donor/acceptor and size features. We describe the features in detail in the following section. We also analyzed the importance of each feature using XGBoost machine learning software. In order to avoid overfitting and make a robust model, we performed 100 times 5-fold cross validations. We created two models: one using 80% and another using 90% of the compiled dataset to train the model and remaining 20% or 10% is used for testing the performance of the model. For a more accurate estimation, we repeated the whole process 100 times and then averaged the PCC and Mean Square Error (MSE). Figure 1 represents a schematic illustration of the SAAMBE-SEQ method.

![Figure 1: Schematic illustration of SAAMBE-SEQ method](image)

2.3 Features

2.3.1 Features based on the position specific scoring matrix (PSSM)

The corresponding protein sequence is utilized as input to search and align homologous sequences from Uniref50 (Suzek, et al., 2014) database (https://www.uniprot.org/downloads) using the PSI-BLAST program (Camacho, et al., 2009) with 3 iterations and a cutoff E-value of 0.001. Then the PSSM is constructed through a multiple sequence alignment of the highest scoring hits. As a result, we obtain a \( L \times 20 \) PSSM for each protein sequence, where \( L \) is the length of each protein sequence. Each row of the PSSM matrix represents the log likelihood score for amino acid substitutions at the corresponding positions in the input sequence:

\[
P_{\text{PSSM}} = \begin{pmatrix}
p_{1,1} & p_{1,2} & \cdots & p_{1,20} \\
p_{2,1} & p_{2,2} & \cdots & p_{2,20} \\
\vdots & \vdots & \ddots & \vdots \\
p_{L,1} & p_{L,2} & \cdots & p_{L,20}
\end{pmatrix}
\]

where \( p_{i,j} \) represents the score of the amino acid residue in the \( i \)th position of the protein sequence being changed to amino acid type \( j \) during the evolution process.

2.3.2 Evolutionary features of mutated and interaction chains

Protein sequences of different size have different length of PSSM. To make the PSSM descriptor become a size-uniform matrix, one approach is to represent a protein sample \( P \) by

\[
P_{\text{PSSM}} = (P_1 P_2 \cdots P_{20})^T
\]
where
\[ P_j = \frac{1}{\sum_{i=1}^{20} P_{ij}} (j = 1, 2, \ldots, 20) \]  

and \( P_j \) is the composition of the amino acid type \( j \) in the PSSM and represents the average score of the amino acid residues in the protein \( P \) being mutated to amino acid type \( j \) during the evolution process. All values in PSSM of each protein sequence are normalized to be between 0 and 1 by sigmoid function:
\[ f(x) = \frac{1}{1 + e^{-x}} \]  

where \( x \) is the original value of PSSM.

Using this method, we obtained 20 uniform average conservation scores for mutated and interaction chains, respectively.

### 2.3.3 Conservation score at mutation site

We select the rows belonging to the given mutation site from PSSM to obtain 20 conservation scores features for the mutation site.

### 2.3.4 Sequence neighbors feature

We labeled ten residues including five on the left and 5 on the right side from the mutation site. There can be 20 possibility of each label representing 20 different amino acids. These are the ten residues which can have a significant influence on mutation site.

### 2.3.5 Features related to mutation site

We used 9 features related to mutation site: net volume, net hydrophobicity, mutation type, net flexibility, chemical property, size, polarity, hydrogen bond and label hydrophobicity. For detail description of each feature, refer to our previous paper (Pahari, et al., 2020) and supplementary material.

### 2.4 Feature importance analysis

We analyzed the importance of each selected feature for the prediction performance of SAAMBE-SEQ method. To evaluate the feature importance, we used XGBoost algorithm from python package. Figure 2 represents the importance level of each feature. Figure 2 reveals that average PSSM of mutant chain (MC avePSSM) and conservation score at mutation site (CS Mutation Site) are the two most important features in our model. The third highest contributing feature is average PSSM for interaction chain (IC avePSSM). These three features capture the evolutionary conservation of a given amino acid at the mutation site as well as of surrounding of mutation site and it’s change upon mutation. PSSM has already been established for providing crucial information in hot-spot prediction (Moreira, et al., 2017) and binding site prediction (Walia, et al., 2014). The next important feature is sequence neighbor, where we took into account 10 amino acids near mutation site according to primary sequence. Sequence neighbor feature captures the influence of neighboring amino acid residues on the mutation site. The next three important features are mutation type and change in molar volume and hydrophobicity of amino acid residues upon the mutation. We applied feature selection protocol on the training set with 5-fold cross-validation when tested on 20% of dataset. Table S7 displays the performances of the models using additive feature groups in each iteration. The final model achieves a PCC of 0.833, higher than the 0.820 using all features. For comparison, we also trained our model by removing each feature candidate from our final model to test the robustness of SAAMBE-SEQ.

### 2.5 no-STRUC dataset

From literature, we collected the experimental \( \Delta \Delta G \) values upon mutation in protein-protein complexes, for which structure is not available. We utilized Uniprot database to obtain the sequence from protein name and used these. We provided the Uniprot ID and the corresponding reference in Tables S1 and S2 in supporting information for homodimer and heterodimer complexes, respectively.

### 3 Results

We trained SAAMBE-SEQ on a large and diverse dataset containing experimental \( \Delta \Delta G \) for 2398 single point mutations from 200 protein-protein dimeric complexes. For predicting \( \Delta \Delta G \) upon a given mutation, we developed a regression model using 80 knowledge-based features, representing evolutionary information and physical environment surrounding the mutation site. In order to build a reliable and robust model, we performed 100 times 5-fold cross validation. Selection of the training and test sets were repeated 100 times randomly, and average PCC and MSE are taken into account. We trained our model against 80% as well as 90% of the 2398 mutations present in our compiled dataset and tested against the remaining 20% or 10% data. In 5-fold cross-validation, our model shows a correlation of 0.83 and MSE of 1.44 kcal/mol when tested on 20% of the database (Figure 3a). On the other hand, we obtained a PCC of 0.91 and MSE of 0.90 kcal/mol when tested on 10% of the database (Figure 3c). In Figure 3, we also plotted the distribution of both experimental as well as predicted \( \Delta \Delta G \)s for the corresponding test sets. In both cases, it can be seen that the distribution of predicted \( \Delta \Delta G \)s using SAAMBE-SEQ is remarkably similar to corresponding experimental \( \Delta \Delta G \)s. To avoid any bias and overfitting, we
chose the model, which is trained on 80% of the dataset for the rest of the paper.

Furthermore, a detailed comparison between predicted and experimental ΔΔGs upon mutations associated with different type of amino acid are plotted in Figure 4. All mutations from the 20% test set were evaluated. In Figure 4, x-axis and y-axis represent the amino acid residue type for wild type and mutant, respectively. The value (in kcal/mol) of variance of ΔΔG upon each type of mutation is shown in color code – the darker the color the large is the variance. We categorized the amino acid residue types in three categories depending on their physico-chemical characteristics: (a) size of the residue (small, medium and large); (b) polarity (polar and nonpolar) and (b) hydrophobicity (hydrophobic, neutral and hydrophilic).

One can see in Figure 4 that both experimental data as well as predicted data using SAAMBE-SEQ indicate that variance of binding energy changes associated with mutations from small residue type in wild type to small residue type in mutant is usually low, while mutations from small to large residue type result in large change in binding energy. Another interesting fact is that if the amino acid residue type is alanine in the wild type, then irrespective of any residue type in the mutant, the binding energy change is always small. One can also see that in both experimental as well as SAAMBE-SEQ predicted data, mutations involving hydrophobic to hydrophobic residue are usually associated with small binding energy change. Overall, comparing the patterns on the left and right panels in Figure 4, one can easily notice that they are very similar indicating that SAAMBE-SEQ predicted variances of ΔΔG is very similar to those of experimental data.
3.1 Performance comparison on blind datasets

For validation, we used three recently published datasets (for more details see Ref. (Geng, et al., 2019)): MDM2-p53, NM and s487 and one compiled by us, termed no-STRUC, which contains mutations from protein complexes whose 3D structures are not available.

3.1.1 Performance of SAAMBE-SEQ on MDM2-p53 dataset

MDM2-p53 dataset contains 33 mutations among which 7 were reported as mutation for which ΔΔG exceeded experimental detection limit. Therefore, we removed these 7 entries from our validation test set, resulting in 26 mutations from a single protein complex (PDB ID is 1YCR, however, in our benchmarking, we used only sequence information). These mutations were not used in our training and test dataset. We compared the correlation between experimental and predicted ΔΔG on these 26 mutations. SAAMBE-SEQ achieved a PCC of 0.35 and MSE of 0.45 kcal/mol. We compared the performance of SAAMBE-SEQ with the only existing sequence-based method, ProAffiMuSeq, which obtained a PCC of 0.16 and MSE of 0.99 kcal/mol. We also compared the prediction of SAAMBE-SEQ with other existing high performing structure-based methods such as iSee, mCSM, BindProfX, FoldX, mCSM-PP12, MutaBind2 and SAAMBE-3D. Figure 5 shows the performance of SAAMBE-SEQ and other methods on MDM2-p53 validation dataset. We can see in Figure 5 that SAAMBE-SEQ outperforms iSee, FoldX, mCSM and MutaBind2 and achieved similar performances (PCC=0.35) as of BindProfX (PCC = 0.36) and mCSM-PP12 (PCC = 0.35). Figure 5 indicates that SAAMBE-3D outperforms all the existing methods and achieved a PCC of 0.41. However, we need to keep in mind that SAAMBE-SEQ is a sequence-based method and the comparable performances of the method with the already established high performing structure-based methods makes SAAMBE-SEQ an outstanding ΔΔG predictor. Moreover, Figure 5b reflects that SAAMBE-SEQ has the second lowest MSE of 0.45 kcal/mol after mCSM-PP12. Furthermore we performed Fisher-T statistical significance test of correlation coefficient (Table S3) for each method and also evaluated whether the difference in other methods compared to SAAMBE-SEQ is statistically significant or not (Table S4) using Fisher-Z test.

3.1.2 NM dataset

The second validation dataset was taken from Benedix el al.’s NM dataset (Benedix, et al., 2009). We only selected single mutations that were not present in our training dataset, and we removed cases in which more than two chains were present in the PDB structure. In this way, we filtered out 27 single mutations from a single protein complex (PDB ID: 1IAR, however, in the benchmark we used only sequence information). Unfortunately, we could not compare the prediction performance of SAAMBE-SEQ with ProAffiMuSeq as these mutations from NM dataset are present in their training dataset. However, we compared the performance of SAAMBE-SEQ with existing structure-based methods, mCSM-PP12 (Rodrigues, et al., 2019), SAAMBE-3D and MutaBind2 (Zhang, et al., 2020) on these selected 27 mutations. Figure 5 presents the correlation between experimental and predicted ΔΔG for the 27 mutations for all the above-mentioned methods. We could not calculate ΔΔG using iSee, mCSM and FoldX because the corresponding web servers were unavailable. Also, we were unable to compare the prediction with BindProfX method as this method only can predict ΔΔG for interfacial mutations. Figure 5 indicates that SAAMBE-SEQ outperforms SAAMBE-3D and mCSM-PP12 in predicting ΔΔG upon mutations from NM dataset. SAAMBE-SEQ achieved PCC of 0.57 and MSE of 2.06 kcal/mol. It should be mentioned that MutaBind2 is the highest performer with PCC of 0.70 and MSE of 1.35 kcal/mol. The results of statistical significance are shown in Table S3 and Table S4.

3.1.3 S487 dataset

The third validation dataset is s487, compiled by Geng et al. (Geng, et al., 2019) The dataset contains 487 single mutations from 56 complexes and all mutations are located at protein-protein interfaces. Figure 5 represents a prediction comparison in the form of PCC and MSE obtained using different structure-based ΔΔG predictors along with SAAMBE-SEQ. The PCC and MSE values, achieved by iSee, BindProfX, FoldX, mCSM, MutaBind2 and SAAMBE-3D are taken from our previous paper (Pahari, et al., 2020). We couldn’t compare the prediction of mCSM-PP12 as some or all of these 487 mutations are present on their training dataset (SKEMPI v2.0). As shown in Figure 5, BindProfX and MutaBind2 both achieve the highest PCC of 0.41 followed by SAAMBE-3D (PCC=0.39) and SAAMBE-SEQ (PCC=0.34).

Unfortunately, we could not compare the ΔΔG prediction on the s487 validation dataset using SAAMBE-SEQ with the only existing sequence-based method, the ProAffiMeSeq, as some of these mutations are already included in their training dataset. Therefore, we considered their compiled validation dataset, s473, which is a combination of above mentioned three datasets. They removed all the mutations which are not present at the protein-protein interfaces as ProAffiMeSeq is trained to predict ΔΔG only for interfacial mutations. Thus, on the s473 dataset, SAAMBE-SEQ achieved PCC of 0.35 and MSE of 1.73 kcal/mol where ProAffiMeSeq obtained PCC of 0.20. Furthermore the statistical significance was evaluated and results are shown in Table S3, S4.

3.1.4 No-STRUC dataset

The last validation test was done on no-STRUC dataset compiled by us (see method section and Tables S1 and S2). This dataset is comprised of experimentally measured changes of the binding free energy of protein-protein complexes for which there is no available experimentally determined 3D structure. Because of that, all above mentioned structure-based methods can not be tested. The only methods that can handle such dataset are SAAMBE-SEQ and ProAffiMeSeq.

We divided the no-STRUC dataset into two categories: homodimer and heterodimer. ProAffiMeSeq was trained on some of the entries in homodimer dataset since some of the no-STRUC cases are taken from PROXiMATE database (Jemimah, et al., 2017). Nevertheless, the
benchmarking was carried out and the results are shown in Table 1. Among the 30 mutations in homodimer dataset, ProAffiMuSeq could not predict for 5 non-interfacial mutations. Therefore, we discarded those 5 mutations while comparing the performance of SAAMBE-SEQ with ProAffiMuSeq on homodimer dataset and reported in Table 1. SAAMBE-SEQ achieved a correlation of 0.35 and MSE of 1.42 kcal/mol when considered all the 30 mutations from homodimer dataset.

<table>
<thead>
<tr>
<th>dataset</th>
<th>PCC</th>
<th>MSE (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homodimers (Table S1)</td>
<td>0.37</td>
<td>1.34</td>
</tr>
<tr>
<td>Heterodimers (Table S2)</td>
<td>0.47</td>
<td>0.73</td>
</tr>
</tbody>
</table>

One can see from Table 1 that SAAMBE-SEQ drastically outperforms ProAffiMuSeq, despite that some of the cases in the homodimer dataset were used for training of ProAffiMuSeq model. In Table S5, we evaluated the statistical significance of the correlation obtained using SAAMBE-SEQ and ProAffiMuSeq for both homodimers and heterodimers. The p-value indicates that for homodimers, the correlation obtained using SAAMBE-SEQ is indeed statistically significant while this is not the case for ProAffiMuSeq. Further we evaluated whether there is significant statistical difference in PCC of SAAMBE-SEQ and ProAffiMuSeq using Fisher-Z test. We obtained p-value of 3.63E-2 for homodimers and 1.20E-2 for heterodimers (Table S6).

3.2 Performance of SAAMBE-SEQ on identifying disruptive and non-disruptive mutations both for homodimer as well as heterodimer

We explored the success of the prediction of SAAMBE-SEQ in classifying disruptive and non-disruptive mutations using only sequence information in case of both homodimer as well as heterodimer complexes. As mentioned in our previous paper (Pahari, et al., 2020), Cornell University dataset contains 2500 single mutations from 300 homodimer protein complexes and 245 single mutations from 50 heterodimeric complexes. Yeast two-hybrid (Y2H) experiments were performed at Cornell University (Fragoza, et al., 2019) and the mutations were scored either disruptive or non-disruptive. The dataset was purged to remove identical sequences and cases where any of the two chains has less than 20 amino acid residues. We combined both homodimer and heterodimer complexes together and ended up with 342 mutations from 90 protein complexes. These 342 mutation entries were split into 80% training and 20% test sets. We used the same features for this classification as described in the methods section for our SAAMBE-SEQ model. We performed ROC analysis and found that our method is 84% successful in classifying disruptive and non-disruptive mutations for the 342 mutations for both homodimer and heterodimer complexes. We plotted ROC in Figure 6 and further prediction performance is measured by area under the curve, accuracy, precision and sensitivity. SAAMBE-SEQ achieved an accuracy of 0.81, precision of 0.65, sensitivity and specificity of 0.81 in classifying disruptive and non-disruptive mutation.

![Figure 6: Prediction performance of SAAMBE-SEQ in identifying disruptive and non-disruptive mutations](image)

3.3 Webserver

We implemented SAAMBE-SEQ as a user-friendly web server, freely available at http://compbio.clemson.edu/saambe_webserver/indexSEQ.php#started. The server front end is built using JavaScript and backend using PHP. It is hosted on a Linux server running in Apache. SAAMBE-SEQ can be used in two different ways: i) predict the effect of mutation specified by the user in the given boxes. User needs to provide FASTA sequence of the protein-protein complex, which can be provided by uploading the sequence in the FASTA format or by inputting the sequence in appropriate box. User must provide two sequences corresponding to two protein chains. User need to make sure that they are uploading or putting the sequence in the appropriate place corresponding to mutated chain and interaction chain. Then, user require to provide mutation details in three different boxes: in position box, corresponding residue number according to FASTA sequence file should be provided. It is important to remember that the first residue number starts with 1 not 0. In the ‘Original Amino Acid’ box, user must specify one-letter code for the wild type residue as a string and similarly for ‘Mutated Amino Acid’, mutant residue in one-letter code must be mentioned. In this way, user can submit a single job. ii) If user wants to submit multiple jobs at the same time, in addition to uploading or inputting sequences of mutated and interaction chain in FASTA format, user need to upload a file called ‘List_Mutation.txt’. The file must contain a list of mutations information in a text file for batch processing. A sample ‘List_Mutation.txt’ file is provided in the submission page in order to assist the user for submission of jobs. iii) User can also directly download the SAAMBE-SEQ code by clicking the download option available via the top navigation bar. A readme file will also be downloaded which will guide the user how to use the code.

4 conclusion

Machine learning methods are the alternative to the first principle-based approaches such as quantum mechanics (QM) modeling, molecular dynamics (MD) and Monte Carlo (MC) simulations(Klepeis, et al., 2009;
Gen et al. (2019), molecular mechanics PB/GB surface area (MM/PB/GBSA), multiscale, and mesoscale methods. In terms of modeling the effects of amino acid substitutions on protein stability, binding and dynamics, one should mention methods as free energy perturbation (FEP), thermodynamics integration (TI) and molecular mechanics Poisson-Boltzmann/Generalized Born surface area (MM/PB/GBSA) (Getov et al., 2016; Li et al., 2014; Petukh et al., 2015). However, machine learning methods are more accurate in their predictions and require less computational time, making them primary choice for large-scale investigations. Indeed, the abovementioned first principle-based methods frequently require days of computation for a single case and since they require 3D structure, any small structural imperfection could result in very wrong predictions.

It should be mentioned that one of the most indicative measure of methods performance is the MSE. The best SAAMBE-SEQ MSE is 0.90 kcal/mol when tested on 10% of the training set. Other methods mentioned in the paper reported MSe ranging from 0.94 kcal/mol up to 2.89 kcal/mol. Thus, one should be careful in interpreting prediction results, since they come with an inherited error. In the simplest way the predictions should be considered on the background of reported MSE. However, different MSE were reported depending on the datasets used in the benchmarking. Therefore, the safest protocol should apply the largest reported MSE to investigations on new set of cases (for which there is no experimental data). Alternatively, one may want to utilize as many as possible predictors and seek a consensus.

Here we reported a method, the SAAMBE-SEQ method, which predicts the change of the binding free energy caused by single mutations utilizing sequence information only. Combined its computational efficiency, accuracy, and availability as a stand-alone code, the SAAMBE-SEQ is the only available method to be applied on genome-scale investigations. Indeed, genomic sequencing produces much more data than the efforts of structure determination, and this trend is not going to change. Therefore, there is a desperate need for machine learning methods that can make predictions using only genomic sequencing data, a need that SAAMBE-SEQ addresses for protein-protein interactions. Furthermore, it is demonstrated that SAAMBE-SEQ is capable of distinguishing disruptive from non-disruptive mutations. Since disruptive mutations are usually disease-causing, SAAMBE-SEQ can be used for early diagnosis by detecting the disruptive mutations.

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References


