Protein stability: A single recorded mutation aids in predicting the effects of other mutations in the same amino acid site

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ABSTRACT

Motivation: Accurate prediction of protein stability is important for understanding the molecular underpinnings of diseases and for the design of new proteins. We introduce a novel approach for the prediction of changes in protein stability that arise from a single-site amino acid substitution; the approach uses available data on mutations occurring in the same position and in other positions. Our algorithm, named **Pro-Maya** (**Prot**ein **M**utant st**A**bilit**Y A**nalyzer), combines a collaborative filtering baseline-model, Random Forests regression and a diverse set of features. Pro-Maya predicts the stability free energy difference of mutant vs. wild type, denoted as $\Delta\Delta G$.

Results: We evaluated our algorithm extensively using cross-validation on two previously utilized datasets of single amino acid mutations and a (third) validation set. The results indicate that using known $\Delta\Delta G$ values of mutations at the query position improves the accuracy of $\Delta\Delta G$ predictions for other mutations in that position. The accuracy of our predictions in such cases significantly surpasses that of similar methods, achieving, e.g., a Pearson correlation coefficient of 0.79, and a root mean square error of 0.96 on the validation set. Because Pro-Maya uses a diverse set of features, including predictions using two other methods, it also performs slightly better than other methods in the absence of additional experimental data on the query positions.

Availability: Pro-Maya is freely available via web server at http://bental.tau.ac.il/ProMaya.

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1 INTRODUCTION

Understanding the mechanisms by which mutations affect protein stability is important for characterizing disease mechanisms and for protein design (Bromberg and Rost, 2009). Hence, the energetics of mutants has been studied extensively through experimental and theoretical approaches.

The methods for predicting the change in a protein's stability $(\Delta\Delta G)$ that results from a single amino acid mutation can be roughly classified according to the types of effective potentials they rely on: physical effective potentials (PEP), statistical effective potentials (SEP) and empirical effective potentials (EEP).

Notably, none of these potentials explicitly takes into consideration relevant known mutations at the query position. PEP-based methods use atomic-level representations to capture the underlying physical phenomena affecting protein stability, e.g. van der Waals interactions and dihedral (torsion) angle (Prevost, et al., 1991; Seeliger and de Groot, 2010). These techniques are computationally demanding and are not applicable to large datasets (Kollman, et al., 2000). SEP-based methods are based on the inverse Boltzmann law, which states that probability densities and energies are closely related quantities. Hence, these methods use datasets of proteins of known structures to calculate conditional probabilities that certain residues or atoms will appear in different contexts. Most SEP-based methods use pairwise potentials (Bahar and Jernigan, 1997; Samudrala and Moult, 1998; Sippl, 1995), though some studies have employed higher-order potentials; for example, (Vaisman, et al., 1998) used a four-body potential. SEP-based methods are computationally efficient, more robust than PEPbased methods to low-resolution protein structure prediction and are suitable to include known and unknown physical effects (Lazaridis and Karplus, 2000). Methods in the third category (EEP-based) use experimental energy data to calibrate the weights of the energy function terms. The types of energy terms used can vary and might be SEP-, PEP-, physicochemically- or evolutionbased (Bloom and Glassman, 2009; Gilis and Rooman, 1997; Masso and Vaisman, 2010; Shen, et al., 2008). For example, PoP-MuSiC-2.0 utilizes a neural network algorithm with SEP features that couple between the identity of the amino acid, secondary structure, accessibility and the spatial distance between amino acids (Dehouck, et al., 2009). Conversely, FoldX's (Guerois, et al., 2002) energy function consists of PEP energy terms calibrated using a grid search method on experimental data. The recently developed Prethermut tool (Tian, et al., 2010) incorporates the energy terms of FoldX and MODELLER (Sali and Blundell, 1993) into a Random Forests machine regression, and has reached impressive results. The use of a machine learning algorithm enables non-energy-like terms to be incorporated into the scoring function (Capriotti, et al., 2005; Cheng, et al., 2006; Montanucci, et al., 2008). For example, both I-Mutant2.0 (Capriotti, et al., 2005) and MUpro (Cheng, et al., 2006) encode the identities of the wild type and mutant amino acids in addition to the quantity (in I-Mutant2.0) or frequency (in MUpro) of the residue type found inside a sphere centered at the mutated residue. Both methods also offer sequence-based predictions in cases where the protein structure is not available. For instance, Capriotti et al. added a description of the amino-acid frequency within a symmetrical sequence window centered at the mutated residue and reached a prediction accuracy that was only slightly lower than that achieved using a structurally-based approach (Capriotti, et al., 2005).

To assess the performance of prediction methods and to calibrate weights in EEP-based methods, several datasets of experimental energy values have been compiled. The main source is the ProTherm dataset (Kumar, et al., 2006). Capriotti et al. compiled a dataset of 1615 single-site mutations that has been used for crossvalidation procedures in several studies (Capriotti, et al., 2005; Cheng, et al., 2006; Masso and Vaisman, 2010). However, as previously indicated by Cheng et al., this dataset is highly redundant and may lead to unreliable predictions. Recently, two large nonredundant datasets have been compiled by Potapov et al. (2009; Potapov-DB) and by Dehouck et al. (2009; PoPMuSiC-DB), containing 2155 and 2648 mutations, respectively. The datasets comprise of $\Delta\Delta G$ measurements from thermal and denaturant denaturation experiments. To avoid redundancy, each dataset considers only one $\Delta\Delta G$ value per mutant. In cases where numerous values have been obtained for a single mutant, Potapov et al. set the mutant's $\Delta\Delta G$ as the mean of the measures, whereas Dehouck et al. determine this value using a weighted average, giving higher weights to measurements taken in physiological conditions (pH close to 7, temperature close to 25 °C and without additives). Thus, although the two datasets share 1405 common mutations, the $\Delta\Delta G$ values assigned to some of these differ.

Preliminary examination of the PoPMuSiC-DB indicated that $\Delta\Delta G$ values of mutations occurring at the same protein position tend to cluster (data not shown), i.e., $\Delta\Delta G$ values of mutations in a given position are closer to each other, on average, than to $\Delta\Delta G$ values in other positions. This suggests making explicit use of known $\Delta\Delta G$ values to predict the effects of new mutations. To this end, we developed an approach based on adaptation of the baseline-model of the BellKor collaborative filtering algorithm (CF) (Koren, 2008). To improve its accuracy we combined the baseline-model algorithm with a content-based model. The content-based model takes into account features of the mutation and its surrounding comprising of various sequence, structure, SEP- and EEP-based features. We benchmarked our algorithm extensively by carrying out cross-validation on the PoPMuSiC-DB and Potapov-DB datasets and by running it on an additional validation set. Statistical analysis of the results indicates that Pro-Maya surpasses all the compared methods both when additional $\Delta\Delta G$ values for the query position are available and when they are not.

2 METHODS

Our algorithm treats differently mutations at positions for which a $\Delta\Delta G$ value for a different mutant is known (denoted MRPM, Multi-Replacement Position Mutation) and at positions with no additional known recorded mutations at the query position (SRPM, Single-Replacement Position Mutation). Given a query mutation of SRPM we follow the traditional machine learning scheme. Specifically, the query mutations is fed to a pre-calculated

Random Forests regression model (Breiman, 2001) to predict the query's $\Delta\Delta G$, denoted as $\Delta\Delta G^{RF}$ (described in 2.1). For MRPMs, as detailed in figure 1, the predicted $\Delta\Delta G^{RF}$ is utilized as an input to an additional prediction step using the integrated baseline and content-based model, denoted as the CFCB algorithm. The $\Delta\Delta G^{RF}$ for the MRPMs is calculated using a Random Forests model retrained on a dataset comprising of the training dataset and the user reported $\Delta\Delta G$ records of mutations at the query position. The input to the CFBC algorithm also includes a matrix representation of the known $\Delta\Delta G$ (described in 2.2) and a set of the features. Note, that the $\Delta\Delta G^{RF}$ in our algorithm is utilized both for the prediction of SRPM mutations and as an input to the CFCB algorithm. The Pro-Maya algorithm predicts the ΔG change of the mutant vs. the wild type protein (i.e. Mutant-WT). Thus, indicating both the magnitude of the stability change and its sign, i.e., whether the mutant is more or less stable than the wt.

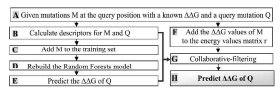


Figure 1. Prediction scheme for a query mutation with known $\Delta\Delta G$ values for additional mutations at the same position. (A) The input for this prediction scheme includes query (Q) and known (M) mutations at the query position. The $\Delta\Delta G$ value for M is known. (B-E) Calculate the predicted $\Delta\Delta G$ of Q using the Random Forests algorithm. (F) Add the $\Delta\Delta G$ values of M to the appropriate elements in the energy matrix r, according to the MU identity and position of M. (G) Given the training set (matrix r), and the features (including the $\Delta\Delta G$ predicted by Random Forests) start the stochastic gradient descent and calculate the $\Delta\Delta G$ of Q (H).

2.1 Calculation of $\Delta\Delta G^{RF}$

The $\Delta\Delta G^{RF}$ is calculated using the Random Forests R implementation (Liaw and Wiener, 2002) of WEKA (Frank, et al., 2004). The number of trees to grow was set to 650 since the addition of more trees did not change the performance. The number of random features to be searched at each tree node was the square-root of the number of features, i.e. 6.

The Random Forests regression utilizes a total of 11 descriptors (F1-F11) with 30 dimensions, which can be roughly divided into sequence- and structure-based features as follows:

Sequence-based features The multiple sequence alignment (MSA) holds important information regarding the physicochemical preference of the position in the protein. From the MSA we calculated the Position Specific Scoring Matrix (indicating the frequency of the amino acids in each MSA column) and used a physicochemical scale matrix to calculate the weighted average and standard deviation of a physicochemical property. Given a mutation, we measured the degree to which its physicochemical properties deviated from the mean physicochemical preference at the query position. Each query mutation was evaluated according to the following physicochemical properties (F1-F3): hydrophobicity scale (Kessel and Ben-Tal, 2002), molecular weight and isoelectric point (Table S1, supplementary material). In addition, we added into the model the number of sequences in the alignment (F4).

Based on a related study (Wainreb, et al., 2010), we added an additional descriptor measuring the sequence identity of the query protein to the closest homolog bearing the mutant amino acid (denoted SIDCH) (F5). For example, mutation I48A in the *Hordeum vulgare* chymotrypsin (Uni-ProtKB/Swiss-Prot ID: ICI2_HORVU) (The_UniProt_Consortium., 2010) was shown by (Jackson, et al., 1993) to cause a major destabilization of the protein. Fifteen homologous proteins with sequence identities of 31%-to-47% to ICI2_HORVU feature the amino acid serine in the corresponding position. Here we set the SIDCH of I48A to 47%. We also included an array of 20 features (for 20 residue types) to encode the identity of the wild

type and mutant amino acids (F6). The features of the wild type and mutant amino acids were set to 1 or -1, respectively, and the rest of the features were set to 0.

Structure-based features

Average solvent accessibility: The side chain accessible surface area (calculated by NACCESS (Hubbard, et al., 1991)) was averaged over all the protein structures of the query protein (F7). In proteins for which an X-ray crystal structure existed, all structures determined through nuclear magnetic resonance (NMR) were disregarded.

Protein flexibility: To reflect the mobility of the protein's backbone at the mutated positions we used the B-factors of the crystal structure (F8).

PEP-based features: We made use of $\Delta\Delta G$ values predicted by the Prethermut tool (Tian, et al., 2010) (F9). Prethermut uses a Random Forests machine learning algorithm and combines the energy terms of FoldX and MODELLER (Sali and Blundell, 1993). The energy terms are translated into units of standard deviation from the average of the energy terms calculated over all possible mutations of the whole protein. To calculate the Prethermut prediction value, we conducted a Random Forests regression over the original energy terms (calculated using the Prethermut scripts). As suggested by Tian et al. the number of trees to grow was set to 650 and the number of random features to be searched at each tree node was the square-root of the number of features, i.e., 8.

SEP-based features: The amino acid-specific torsion angle potential were calculated according to (Parthiban, et al., 2006) (F10). In addition, we utilized the PoPMuSiC-2.0 predicted $\Delta\Delta G$ value, calculated using the energy terms in (Dehouck, et al., 2009) and the Gaussian regression (Rasmussen and Williams, 2006) implementation of Weka (Frank, et al., 2004) (F11). The Gaussian regression cross-validation results of PoPMuSiC-2.0 were comparable with the published results. The predicted PoPMuSiC-2.0 $\Delta\Delta G$ values for mutations that were absent from the Potapov-DB were calculated using the PoPMuSiC-2.0 web server.

2.2 Collaborative filtering and content-based algorithm (CFCB)

Collaborative filtering and content-based recommender systems are used by many websites to generate personalized recommendations. For example, when a customer purchases an item on a retail website, such algorithms try to predict which other items the user would enjoy, on the basis of his/her past behavior and similarity to the behavior of other users. CF algorithms use only user—item data to make predictions. Conversely, content-based algorithms rely on the features of users and items for prediction.

In recent years, the main driving force behind the development of CF algorithms has been Netflix's million-dollar prize for improving the performance of the site's recommendation system. Here, we chose to utilize a part of the CF solution of the winning group (named BellKor) (Koren, 2008). In order to improve the model's performance we extended it using a content-based-model to take into account biological information regarding the mutations.

In our CF scenario, there is a list of possible mutation outcomes (MU) (i.e. all possible amino acids), a list of mutation positions (defined by the protein and the residue number) and the experimental $\Delta\Delta G$ values for some of the mutations at these positions. The data can be stored in a sparse matrix r of size $n \times m$, where n denotes the number of MUs and m denotes the number of positions. Each cell r_{ui} of the matrix r indicates the $\Delta\Delta G$ of a mutation to amino acid u at position i (see, for example figure S1A, supplementary material).

For clarity, special indexing letters u and i are reserved for distinguishing MUs and positions, respectively.

2.2.1 The prediction models The BellKor CF algorithm (Koren, 2008) tries to model the relations between the known data points in matrix r. The model's parameters are learned during the training procedure. The optimal model is later utilized to predict $\Delta\Delta G$ values of unknown mutations in positions with known $\Delta\Delta G$ values for other mutation.

The BellKor model integrates three types of approaches to CF: a baseline model, a neighborhood model and the latent factor model. Our CFCB algorithm integrates the BellKor baseline estimator model with a content-based model. We also implemented the neighborhood and latent factor models but according to our analysis their incorporation into the model does not improve the prediction accuracy significantly, although it might in certain cases (supplementary material). A schematic representation of all models can be seen in figure S1 in the supplementary material.

2.2.1.1 The baseline estimator model Different MUs and positions have different $\Delta\Delta G$ tendencies. For example, the $\Delta\Delta G$ of a mutation at a buried position in a protein is usually larger than that of the same mutation at an exposed position. Similarly, we would expect that in most cases the consequences of mutation to proline would be more severe than a mutation to alanine. Hence, each position and MU is ascribed unique baseline estimators, denoted b_i and b_u , respectively. Thus, for every r_{ui} we define a baseline estimator $b_{ui}=\mu+b_i+b_u$, with μ denoting the overall average of all $\Delta\Delta G$ in r. The variables b_i and b_u are learned during the training stage of the algorithm (described in section 2.2.2).

2.2.1.2 The content-based model The baseline model does not use any explicit description of the mutation. In order to describe the biological aspects of the mutation we use a linear regression solution (with no intercept) (equation 1) with a subset of the features (described in section 2.2): solvent accessibility, torsional statistical force field, Prethermut MODELLER-based features, the SIFT predicted compatibility of the mutated amino acid to the query position (Ng and Henikoff, 2003), and $\Delta\Delta G$ predictions by PoPMuSiC-2.0, Prethermut. In addition we also use as a feature the $\Delta\Delta G^{RF}$.

In equation 1, X_{ui} is the set of d features ($X_{ui,1}$, $X_{ui,2}$... $X_{ui,d}$), describing the mutation whose $\Delta\Delta G$ indices in matrix r are u and i. F denotes a set of d descriptor coefficients. As is often done in linear regression, each descriptor is normalized across all positions and MUs so that its average is zero and the standard deviation is 1. F is learned during the training stage described in section 2.2.2 using the stochastic gradient descent.

Equation 1:
$$r_{ui}^{Con} = X_{ui}F$$

2.2.1.3 The integrated model The integrated model (equation 2) combines the baseline- and content-based models. y_{ui} denotes the predicted $\Delta\Delta G$

Equation 2:
$$y_{ui} = b_{ui} + \sum_{g=0}^{d} X_{ui,g} F_g$$

2.2.2 The CFCB training and prediction procedures As in any machine learning algorithm the aim of the training procedure is to obtain parameters that fit the model to the observed data best. Unconventionally, the CFCB model is retrained for every server query in order to identify the parameters of the newly added user-reported mutations, e.g. the baseline estimator of the newly added position. The model with the optimized set of parameters presumably describes best the relations between the known $\Delta\Delta$ Gs in matrix r and is used to predict the unknown MRPM queries.

The training procedure is performed using a stochastic gradient descent algorithm that attempts to minimize the associated regularized squared error function (equation 3) and determines the following parameters: b_{uv} b_{iv} and F. Thus, starting with random values for the parameters, it randomly loops over all the known $\Delta\Delta G$ values in r (which is composed of all known mutations across all proteins in the training dataset) and modify the parameters by moving in the opposite direction of the gradient (equations 4). The descent iterations continue until the difference between the e_{ui} of the current iteration and the previous iteration is smaller than ε . During the training we used the following meta parameters: (learning rate) γ =0.02, (regularization factor) λ =0.025 and ε =0.00001.

Equation 3:

$$\min_{\substack{b_u, b_i, F \\ i \in Positions}} \sum_{\substack{u \in MU. \\ i \in Positions}} (r_{ui} - r_{ui}^{\hat{}})^2 - \lambda (b_u^2 + b_i^2 + \sum_{g=0}^d F_g^2)$$

Equations 4:

- $e_{ui} \leftarrow r_{ui} r_{ui}$
- $b_u \leftarrow b_u + \gamma \cdot (e_{ui} \lambda \cdot b_u)$
- $b_i \leftarrow b_i + \gamma \cdot (e_{ui} \lambda \cdot b_i)$
- $F \leftarrow F + \gamma \cdot (e_{ui} \cdot X_{ui} \lambda \cdot F)$

2.3 The datasets and performance measurements

To train and assess our algorithm we utilized two publicly available datasets: the PoPMuSiC-DB with 2648 mutations in 137 proteins, and the Potapov-DB with 2155 mutations in 79 proteins. Both datasets include $\Delta\Delta G$ values of non-redundant single site mutations (apart from a single mutation in Potapov-DB that was disregarded). Several Protein Data Bank (PDB) structures (NMR and $C\alpha$ only structures) were replaced by others (Table S2, supplementary material). Both datasets have been previously used as benchmarks: Potapov-DB for Prethermut (Tian, et al., 2010) and PoPMuSiC-DB for PoPMuSiC-2.0 (Dehouck, et al., 2009).

To fairly compare our method with Prethermut and PoPMuSiC–2.0, we followed their cross-validation protocols and used a 5- and 10-fold cross-validation on the PoPMuSiC-DB and Potapov-DB sets, respectively. The randomly selected folds were maintained throughout the prediction scheme, i.e. the calculation of the Prethermut, PoPMuSiC-2.0, $\Delta\Delta G^{RF}$ and CFCB prediction values. To calculate the average and SD for the performance measures, we used a bootstrap procedure with 1000 iterations. For each iteration we randomly selected 60% of the cross-validation $\Delta\Delta G$ predictions.

To further evaluate and compare our performance to that of other prediction methods we also utilized the validation set compiled by Dehouck et al. (2009). This validation set includes 350 mutations from 67 different proteins that were not included in any of the training databases of current methods (specified in Table S3, supplementary material). Here, the predicted $\Delta\Delta G$ values of Prethermut and PoPMuSiC-2.0, used as features in Pro-Maya's prediction scheme, were calculated using a 5-fold cross-validation on PoPMuSiC-DB after removing the validation set.

To assess how the number of mutations with known $\Delta\Delta G$ values in the query position affect the prediction accuracy we compared the performance of two leave-one-out (LOO) cross-validation variations named LOO-all and LOO-neglected. In each iteration of both procedures one query mutation was kept as a test and the rest of the mutations were used for training. However, during the LOO-neglect, randomly selected mutation occurring at the query position was neglected from the training set.

To empirically estimate how well Pro-Maya can be generalized to unseen mutations it is important that the training and testing sets are as dissimilar as possible. Therefore, we performed an additional LOO variation, we name LOO-unseen. During each iteration of the LOO-unseen a single mutation was kept for testing and the rest of the mutations in the query position were used for training. Next, all the rest of the mutations that occur at proteins with a low sequence identity to the query protein (sequence identity < 30%) were added to the training set.

At each iteration of LOO-all, LOO-neglected and LOO-unseen the $\Delta\Delta G$ prediction models of Prethermut and PoPMuSiC-2.0 had to be retrained with the modified training set. Since for the Potapov-DB we do not have the PoPMuSiC-2.0 statistical force field components (needed for the retraining) all the LOO procedures were conducted solely on the PoPMuSiC-DB for which we have the required PoPMuSiC-2.0 statistical force field components.

To evaluate performance we used two standard measures: the Pearson correlation coefficient (PCC) and root mean square error (RMSE) between the measured and predicted $\Delta\Delta G$ values (Equations S7 and S8, Supplementary material).

2.5 Data collection

Both the sequences and the PDB file names required were extracted from the corresponding SWISS-PROT entries (Jain, et al., 2009). The MSAs and the PDB files were downloaded from the ConSurf-DB (Goldenberg, et al., 2009) and PDB (Berman, et al., 2000) databases, respectively.

3 RESULTS

3.1 Cross-validation results

According to the PCC and RMSE, Pro-Maya exhibits better performance than FoldX, Prethermut and PoPMuSiC-2.0 for both the Potapov-DB and the PoPMuSiC-DB sets (Table 1; Figures S2, S3, supplementary material). Pro-Maya reached a PCC of 0.77 for both sets (column RF ∩ CF) and RMSE values of 1.09 and 0.94 for the Potapov-DB and PoPMuSiC-DB sets, respectively. These results are also superior to those obtained by CC/PBSA (Benedix, et al., 2009), EGAD (Pokala and Handel, 2005), FoldX (Guerois, et al., 2002), Hunter (Tian, et al., 2009), I-Mutant2.0 (Capriotti, et al., 2005), Rosetta (Rohl, et al., 2004) and the combined method used by Potapov et al. (Potapov, et al., 2009) on the Potapov-DB (Table S4, supplementary material).

To gain a more comprehensive understanding, we also examined the results on the MRPMs and SRPMs subsets of each of the two datasets. The results for the MRPM sets exhibit how well Pro-Maya utilizes the $\Delta\Delta G$ data of known mutation(s) in a specific position to predict $\Delta\Delta G$ values of other mutations at the same site. As can be seen in Table 1, although all methods perform better on the MRPMs, our CFCB algorithm utilizes the training data best and reaches correlation values of 0.83 for the Potapov-DB set and 0.82 for the PoPMuSiC-DB set.

The results for the SRPM subset indicate the performance for mutations at positions that are absent from the training set. For this mutation subset, our prediction scheme does not involve the CFCB algorithm and relies solely on the Random Forests regression and on the quality of the features. Here, our prediction scheme performs slightly better than Prethermut and PoPMuSiC-2.0 on both

Table 1. Cross-validation results

	Mutation	Dataset	Performance measure	Pro-Maya			Prethermut	PoPMuSiC	E-LIV
	number			ΔΔG ^{RF}	CFCB	ΔΔG ^{RF} ∪ CFCB	Prememu	-2.0	FoldX
The whole dataset	2155	Potapov	PCC	0.74±0.01		0.77±0.01	0.72±0.01	0.62±0.01	0.55±0.02
		-DB	RMSE (kcal/mol)	1.13		1.09	1.20	1.35	1.64
	2648	PoPMuSiC	PCC	0.74±0.01		0.77±0.01	0.71±0.01	0.62±0.01	0.52±0.02
		-DB	RMSE (kcal/mol)	0.99		0.94	1.05	1.15	1.71
SRPM	752	Potapov	PCC	0.59±0.04			0.57±0.03	0.48±0.04	0.5±0.03
		-DB	RMSE (kcal/mol)	1.28			1.30	1.39	1.57
	913	PoPMuSiC	PCC	0.64±0.02			0.61±0.02	0.55±0.02	0.44±0.03
		-DB	RMSE (kcal/mol)	1.11			1.14	1.21	1.74
MRPM	1403	Potapov	PCC	0.80±0.01	0.83±0.01		0.77±0.01	0.69±0.01	0.58±0.02
		-DB	RMSE (kcal/mol)	1.07	0.98		1.14	1.32	1.67
	1735	PoPMuSiC	PCC	0.79±0.01	0.82±0.01		0.75±0.01	0.66±0.01	0.55±0.02
		-DB	RMSE (keal/mol)	0.92	0.85		0.99	1.12	1.69

The Pearson correlation coefficient (PCC) and root mean square error (RMSE) of current methods and Pro-Maya's CFCB and Random Forests ($\Delta\Delta G^{RF}$) prediction schemes on the PoPMuSiC-DB and Potapov-DB datasets and its subsets. The two subsets are mutations at positions absent from the training set (SRPM), and mutations at positions found in the training set (MRPM). The $\Delta\Delta G^{RF}$ \cap CFCB column reports the total performance for the $\Delta\Delta G^{RF}$ and CFCB results on the SRPM and MRPM subsets, respectively. The average and SD of the performance measures were obtained by a bootstrap procedure run for 1000 iterations performed on the cross-validation predictions. As can be seen, Pro-Maya outperforms the other methods. Moreover, the results for the MRPM set indicate that the incorporation of experimental data regarding mutations at the query position improved the prediction accuracy.

datasets. However all methods show major decline in the performance. Note that although the ranges of Prethermut's and our results coincide according to the average and the standard deviation, for all subsets created during the bootstrapping process our PCC showed an average (minor) improvement of 0.02±0.1 over the PCC of Prethermut, the best of the other methods.

Interestingly, each method achieved a lower RMSE for the PoP-MuSiC-DB set than for the Potapov-DB set. This trend is also seen in the cross-validation results of the 1405 mutations shared by the two datasets (data not shown). Possible explanations are suggested in the Discussion below.

Pro-Maya's performance was also evaluated on a validation set of mutations excluded from the PoPMuSiC-DB. This validation set has been previously used by Dehouck et al. to benchmark PoPMuSiC-2.0, Dmutant (Zhou and Zhou, 2002), Auto-MUTE (Masso and Vaisman, 2010), FoldX (Guerois, et al., 2002), CUPSAT (Parthiban, et al., 2006), Eris (Yin, et al., 2007) and I-Mutant-2.0 (Capriotti, et al., 2005). Both the PCC and RMSE values indicate that Pro-Maya performs better than these aforementioned methods (Table 2, Table S5; supplementary material) for the entire validation set and for its SRPM and MRPM subsets. As can be seen in Table 2, Pro-Maya's PCC on the entire validation set reaches a value of 0.79, constituting an improvement of 0.07 and of 0.1 over the PCCs obtained by Prethermut and by PoPMuSiC-2.0, respectively.

Table 2. Performance over the validation set

	Mutation number	Performance measure	Pro-Maya	Prethermut	PoPMuSiC- 2.0
The whole	350	PCC	0.79	0.72	0.69
dataset	330	RMSE (kcal/mol)	0.96	1.12	1.16
SRPM	196	PCC	0.69	0.65	0.65
SKI WI	190	RMSE (kcal/mol)	1.09	1.15	1.15
1 (DD) (4-4	PCC	0.89	0.79	0.75
MRPM	154	RMSE (kcal/mol)	0.77	1.09	1.18

The Pearson correlation coefficient (PCC) and root mean square error (RMSE) of Pro-Maya's(*), Prethermut's and PoPMuSiC-2.0's predictions schemes on the whole validations set, and the MRPM and SRPM subsets. As can be seen, Pro-Maya performs better on the entire validation set and subsets.

*Pro-Maya's final performance is the total performance for the Random Forests and collaborative filtering results on the SRPM and MRPM subsets, respectively (CFCB) $\Delta \Delta G^{RF}$.

To estimate how well Pro-Maya performs on query mutations at proteins that are not homologous to any of the proteins in the training set we compared the performance of the LOO-unseen with the performance of the LOO-all (table S4; supplementary material). Interestingly, although the performance of the $\Delta\Delta G^{RF}$ of the LOO-unseen declined both on the MRPM and SRPM subsets (PCC of 0.76±0.01 and 0.60±0.02, respectively) the CFCB algorithm was able to compensate and maintain a similar PCC in both LOO procedures, achieving a PCC of 0.83±0.01.

The results of the 5- and 10-fold and LOO-unseen cross-validation can be viewed online at the FAQ section of the Pro-Maya website. The FAQ section also contains a detailed description of Pro-Maya's training set e.g. number of proteins, number of mutated positions per proteins, functionality (SCOP classification (Andreeva, et al., 2008)) and physical properties of the proteins. An analysis of Pro-Maya's LOO-unseen versus the SCOP classifi-

cation (Table S6; supplementary material) of the proteins shows

that Pro-Maya performs similarly on the All α , All β , α + β and α/β SCOP classes with a PCC ranging from 0.59 to 0.64 for the SRPM and 0.8-0.83 for the MRPM. The PoPMuSiC-DB includes low number of mutations from the Coiled-coil, Multi-domain and Small proteins SCOP classes. Thus we cannot estimate Pro-Maya performance on these classes, although there is no reason to believe that the performance over them will differ significantly from the rest.

3.2 How do the number and type of mutations with known $\Delta\Delta G$ values in the query position affect the prediction accuracy?

Figure 2 shows that Pro-Maya's prediction accuracy increases significantly with the addition of a single or two known mutations at the query position, and that the accuracy does not improve further with the addition of more than two records.

Intuitively, we might expect that the prediction accuracy of the CFCB algorithm should be correlated with the level of similarity

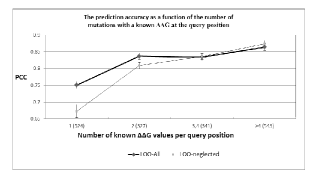


Figure 2. The PCC of Pro-Maya on the PoPMuSiC-DB vs. the number of known mutations at the query position using the LOO-all and LOO-neglect. The number of mutations in each group is shown in parentheses. For example, the second data point of the black curve indicates the performance of Pro-Maya on 327 query mutations with two known records for each in the training set. The second data point of the grey curve indicates the performance of Pro-Maya on the same subset of mutation after one of the two known mutations was removed from the training set. The first data point of the grey curve was calculated using the $\Delta\Delta G^{RF}$. The difference between the grey and black curves indicates the PCC improvement achieved by the addition of a single known mutation in the query position. The results suggest that the improvement in accuracy is facilitated by the incorporation of as few as 1-2 known $\Delta\Delta G$ values in the query position.

between the physicochemical properties of the query and recorded mutations. To examine this hypothesis, for each of the mutations predicted by the CFCB algorithm in the PoPMuSiC-DB, we measured the shortest physicochemical distance (using the Miyata matrix (Miyata, et al., 1979)) from the query mutation amino acid to any of the recorded mutations. For example, given a query mutation to isoleucine at residue 29 in the apomyoglobin protein (PDB id: 1bvc chain A), we measured the shortest Miyata distance from isoleucine to any of the mutations, e.g. alanine, valine and methionine. Here, we set the shortest Miyata distance to 0.29, which is the Miyata distance between isoleucine and methionine. The correlation between the Miyata distances of all query mutations with the squared error ((predicted $\Delta\Delta G$ – observed $\Delta\Delta G$)²) reached only a low PCC of 0.14. This unexpected low correlation suggests that the performance of the CFCB algorithm is not affected by the

identity of the mutations with known $\Delta\Delta G$ values at the query position.

4 DISCUSSION

We tested Pro-Maya extensively using cross-validation on two datasets and an additional validation dataset, and found that it outperformed current methods for the prediction of mutation stability. Our results demonstrate that the availability of as few as one or two records in the query position improve the prediction accuracy of $\Delta\Delta G$ values of additional mutations in that position. This improvement is independent of the amino acid identity of these records and of the sequence identity of the query protein to the training set. Thus, a systematic alanine-scanning mutagenesis of all the amino acids in a protein could greatly increase Pro-Maya's prediction accuracy for any mutation in the protein.

The performance of our Random Forests prediction scheme on the SRPM subset is slightly better than that of the other methods we investigated. We attribute the improvement to the use of an inhomogeneous feature set comprising PEP-, SEP- and evolution-based features, including predictions by the Prethermut (Tian, et al., 2010) and PoPMuSiC-2.0 (Dehouck, et al., 2009) tools. Previous prediction methods, in contrast, have been based on features of a single type (e.g., only PEP).

Pro-Maya's RMSEs for mutations in the PoPMuSiC-DB set are consistently lower than those for the Potapov-DB set. This is presumably because of the different procedures used for compilation of each dataset. PoPMuSiC-DB's compilation procedure used a weighted average of the identical mutations occurring in different conditions to calculate the $\Delta\Delta G$ values that are most likely to occur at physiological conditions. Whereas, the Potapov-DB compilation procedure gives equal weight to the various conditions at which $\Delta\Delta G$ values are measured. Our prediction scheme does not take into account the conditions at which the $\Delta\Delta G$ was measured. Thus, it assumes that all measurements were taken under the same conditions. Therefore, the PoPMuSiC-DB mutation set, which is characterized by more homogenous experimental conditions, is presumably more suitable for our prediction scheme, as indicated by the low RMSE value. To achieve more accurate predictions, we trained the Pro-Maya web server using the PoPMuSiC-DB set. Thus, the server is best suited for predicting mutations at physiological conditions.

Pro-Maya's improved accuracy is facilitated by the use of a baseline estimator that utilizes known $\Delta\Delta G$ records to determine a position specific baseline $\Delta\Delta G$ (b_i) model. The underlying assumption of Pro-Maya is that the $\Delta\Delta G$ of a mutation is strongly dependent on properties that are inherent to the amino acid position in the protein (e.g. solvent accessibility, amino acid identity, interaction with the environment and secondary structure). Thus, on average all mutations at the same position are expected to have similar $\Delta\Delta G$ values. Therefore the position baseline $\Delta\Delta G$ which presumably reflects the inherent properties of the position can roughly model the query mutation. To fully model a mutation Pro-Maya also uses a content based- and a MU specific $\Delta\Delta G$ baseline based models. These models describe the mutation outcome attributes (e.g. physicochemical properties) and predict the $\Delta\Delta G$ shift from the position baseline. Nevertheless, it is expected that mutations with an irregular $\Delta\Delta G$ that differs much from the position $\Delta\Delta G$ baseline would be harder to predict.

By design, Pro-Maya is not very suitable as a classifier of whether a mutation would stabilize or destabilize the protein; a classifier should be trained to this end.

CF algorithms have been developed mainly for online electronic commerce applications and are particularly useful for exploiting large datasets very rapidly. To the best of our knowledge, their use in biology is quite scarce (Erhan, et al., 2006). The success of the CFCB algorithm in this study and the capability of the neighborhood- and latent-factor based models to identify biological properties (discussed in the supplementary material) suggest that the CF approach could be applied to additional problems in biology. Examples include the identification of deleterious mutations in single nucleotide polymorphism data, the detection of true protein-protein interactions in noisy yeast-two-hybrid and mass-spectrometry data, as well as the prediction of ligand and drug molecules that could bind target proteins. Our CFCB algorithm and its integration with the neighborhood- and latent factor-based models can be readily adapted to these problems.

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