Sequence-based prediction of protein crystallization, purification and production propensity

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ABSTRACT

Motivation: X-ray crystallography-based protein structure determination, which accounts for majority of solved structures, is characterized by relatively low success rates. One solution is to build tools which support selection of targets that are more likely to crystallize. Several in silico methods that predict propensity of diffraction-quality crystallization from protein chains were developed. We show that the quality of their predictions drops when applied to more recent crystallization trails, which calls for new solutions.

Results: The proposed PCCpred (predictor of protein Production, Purification, Crystallization and Crystallization) predict propensity for production of diffraction-quality crystals, production of crystals, purification and production of the protein material. PCCpred utilizes comprehensive set of inputs based on energy and hydrophobicity indices, composition of certain amino acid types, predicted disorder, secondary structure and solvent accessibility, and content of certain buried and exposed residues. Our method significantly outperforms alignment-based predictions and several modern crystallization propensity predictors. Receiver operating characteristic (ROC) curves show that PCCpred is particularly useful for users who desire high true positive (TP) rates, i.e. low rate of mispredictions for solvable chains. Our model reveals several intuitive factors that influence the success of individual steps and the entire crystallization process, including the content of Cys, buried His and Ser, hydrophobic/hydrophilic segments and the number of predicted disordered segments.

Availability: http://biomine.ece.ualberta.ca/PCCpred/.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION

Structural genomics (SG) is an international initiative that aims at solving 3D shapes of important biological macro-molecules, primarily focusing on proteins, and which significantly contributes to the overall protein structure determination efforts (Chandonia and Brenner, 2006). This initiative shifts the focus from one-by-one determination of individual protein structures to protein family-directed structure analyses where a group of proteins is targeted and structure(s) of representative members are determined (Terwilliger et al., 1998). The process of choosing the representative proteins, which is known as target selection, encompasses restricting the candidate proteins to those that are tractable and of unknown structure and prioritizing them according to the expected interest and accessibility (Brenner, 2000). For instance, in case of the protein structure initiative (PSI), target selection concentrates on representatives from large, structurally uncharacterized protein domain families, and from structurally uncharacterized subfamilies in very large and diverse families with incomplete structural coverage (Dessailly et al., 2009). The most recent PSI:Biology phase (PSI-Nature Structural Biology Knowledgebase, 2011), includes four major SG centers that will continue high-throughput structure determination defined through community nomination process. Importantly, these approaches allow for flexibility in the selection of the targets and would benefit from tools that support target selection.

Protein structures are solved using X-ray crystallography, NMR spectroscopy, electron microscopy and (recently) by application of computational approaches, such as homology modeling. The most popular method, which accounts for about 87% of the structures deposited into the Protein Data Bank (PDB), is the X-ray crystallography (Berman et al., 2000). One of the main challenges the SG faces is that only about 2-10% of protein targets pursued yield high-resolution protein structures (Service, 2005). A more recent analysis shows that about 4.6% of targets produce the diffraction-quality crystals (Kurgan and Mizianty, 2009). Moreover, estimates show that >60% of the cost of structure determination is consumed by the failed attempts (Slibinski et al., 2007a), while crystallization is characterized by a significant rate of attrition and is among the most complex and the least understood problems in the structural biology (Hui and Edwards, 2003). This provides motivation for further research in this area. Several strategies have been proposed to improve the success rates including obtaining one representative structure per protein family and working with multiple orthologs (Brenner, 2000; Chandonia and Brenner, 2005; Hui and Edwards, 2003). In spite of the advances made in the context of protein crystallization (McPherson, 2004), the production of high-quality crystals is one of the major bottlenecks in the structure determination pipelines (Biertumpef et al., 2005; Chayen, 2004; Puesy et al., 2005). This problem is usually tackled using an empirical trial and error approach, where a large number of experiments is brute-forced to find a suitable setup, and through understanding of the fundamental principles that govern crystallization (Chayen, 2004). The latter is used to design new and improved methodologies that produce the high-quality crystals.

One of the steps taken to alleviate difficulties in solving the structures via the X-ray crystallography was to create databases that record information concerning both successful and failed attempts to produce the structures (Rodrigues and Hubbard, 2003). The largest and most comprehensive database, TargetDB (Chen et al.,...
2004), which was launched July 2001, builds upon the work on the PRESEAGE database (Brenner et al., 1999). TargetDB consolidates data from 28 SG centers in USA, Canada, Germany, Israel, Japan, France and UK, including nine PSI centers. The PepDB (Protein Expression Purification and Crystallization DataBase) was established around 2004 as an extension to the TargetDB to collect more detailed status information and experimental details for each step in the protein structure production pipelines (Kouranov et al., 2006). This database stores a complete history of the status of the experimental steps in each production trial, the current status and stop conditions, which are collected from 15 SG centers in USA.

The availability of the databases motivated the development of analytical and predictive models that either support or directly predict protein crystallization (Rupp and Wang, 2004). Initial work concentrated on finding sequence-derived factors that are useful to determine crystallization propensity. For instance, using the data from the TargetDB, Goh et al. (2004) found that conservation of the sequence across organisms, composition of charged residues, occurrence of hydrophobic patches in the sequence, number of binding partners and chain length influence the feasibility for the high-throughput structure determination. The isoelectric point calculated from the sequence was used to suggest optimal pH ranges for the crystallization screening (Kantardjieff and Rupp, 2004; Kantardjieff et al., 2004). The Berkeley’s SG center utilized several protein features including chain length and predicted transmembrane helices, coiled coils and low-complexity regions to eliminate the intractable targets (Chandona et al., 2006). A recent study shows that the crystallization propensity can be computed from the knowledge of predicted disordered residues, side-chain entropy of predicted exposed residues and the amount of Phe and predicted buried Gly in the input sequence (Price et al., 2009). These works demonstrate that the crystallization propensity can be successfully predicted from the protein chain. However, these studies usually concern data from a single SG center, and propose simple predictive models (that could be outperformed by more advanced models) that are rarely made available, e.g. via web servers, to the community.

To this end, several computational sequence-based crystallization propensity predictors which utilize the data that span multiple SG centers and more advanced predictive models were proposed in the recent years. They include SECRET (Smialowski et al., 2006), OB-Score (Overton and Barton, 2006), CRYSTALP (Chen et al., 2007), XtalPred (Slabinski et al., 2007a, b), ParCrys (Overton et al., 2008), CRYSTALP2 (Kurgan et al., 2009), MetaPPCP (Mizianty and Kurgan, 2009), Pcx (Price et al., 2009), SVMCrys (Kandaswamy et al., 2010) and MCSG-Z score (Babrigi and Joachimiak, 2010). Some of them, including the OB-score and XtalPred, were already used by the SG centers. Details concerning these methods can be found in a recent review (Kurgan and Mizianty, 2009). The above predictors have a few drawbacks, which motivate this work. They are built and tested using outdated data, which results in a relatively poor performance for recent data (we demonstrate that in Fig. 2 in Section 3), the annotation of the chains used in their training database (crystallizable versus crystallization resistant) is based on an incomplete/inaccurate protocol, and they address the prediction of the success of the entire crystallization process without pointing out which of the steps is responsible for the failure. Our objective is to alleviate these drawbacks by proposing a novel predictor that (i) is built using a recent and large dataset, (ii) uses improved annotation protocol, (iii) targets prediction of the success of the entire crystallization process and also predicts which of the steps results in the failed attempts and (iv) uses a compact and comprehensive range of sequence-derived inputs to generate accurate predictions. In collaboration with the TargetDB and PepDB curators, we formulate a more precise and comprehensive protocol to annotate proteins. Also, our method can be used to predict propensity of a given chain for (i) production of diffraction quality crystals, (ii) production of crystals, (iii) purification and (iv) production of the protein material; the existing methods target only one of these outcomes. The proposed method, called PPCpred (predictor of protein Production, Purification and Crystallization) provides individual predictions for each of the four steps, and it also provides an integrated output that predicts whether the chain would produce the diffraction quality crystals, and if not then it predicts which of the steps is most likely to cause the failure.

2 METHODS

2.1 Current annotation protocols

The existing sequence-based crystallization propensity predictors assign one of the two labels, crystallizable or non-crystallizable, to each protein chain. The only exception is the XtalPred which defines five classes that range between ‘easy to crystalize’ and ‘hard to crystalize’. However, the five classes are equivalent to the probabilities/scores generated by the other methods. The annotations used in SECRET are primarily based on the data from the PDB and they were improved by the subsequent works. Most of the subsequent predictors, including CRYSTALP, OB-Score, ParCrys, CRYSTALP2, MetaPPCP and SVMCrys, utilize data extracted from PepDB and TargetDB databases using approach described in Overton and Barton (2006) to annotate proteins as crystallizable and non-crystallizable and to derive their training and test datasets. As it was pointed out in Overton and Barton (2006), this annotation protocol has a few shortcomings. The authors of XtalPred use a different approach in terms of how the crystallizable samples were extracted. In XtalPred (Slabinski et al., 2007b) only trials deposited in PDB were marked as the crystallizable trials, whereas in Overton et al. (2008) those trials were excluded. The two remaining predictors use data from only one SG center; Pcx is based on data from the Northeast SG Consortium and MCSG-Z score from the Midwest Center for SG.

Our aim is to deliver more comprehensive and precise annotations of the crystallization propensity (and the other steps in the crystallization process). First, we consider only the proteins with the known outcome of the experiment, i.e. completed stop status, as opposed to the previous approaches in which the experiment was considered as finished if there was no update for the specific amount of time or it was annotated as work stopped. We note that there are instances where work on a given target was discontinued for unspecified reasons, not necessarily related to the crystallization protocol, e.g. the targets were re-prioritized and administratively abandoned. This is often done without evaluating/reporting the success or failure at the trial level, and thus it is difficult to associate an interpretation to an unqualified ‘work-stopped’ status. Although the authors of Overton and Barton (2006) were aware of this, a small number of trials with the completed stop status at the time when they developed their method did not allow them to utilize a more accurate annotation protocol. Secondly, we filter our dataset to further improve the quality of annotations for the selected samples; details are explained in Section 2.2. Thirdly, we divide the chains annotated as the non-crystallizable to indicate the reason for the crystallization failure.

We hypothesize that success/failure in each of the crystallization process steps may be associated with different protein properties. Fourthly, previous methods used the training and test datasets with substantially reduced sequence similarity (<25%). This removes hard to predict samples, as sometimes even relatively minor changes in the protein sequence (e.g. point mutations and usage of C- and N-terminals tags to ease purification) may
change the final outcome of the crystallization. In contrast, we remove similar sequences only within each class (e.g. similar crystallizable chains), but we do not reduce the sequence similarity between the classes (e.g. between crystallizable and non-crystallizable chains). Finally, we use new data (from beginning of 2006 and 2009) to address recent changes in the crystallization protocols; the prior predictors were built on older data from before 2006.

2.2 Annotations and datasets

The protein chains were extracted from the PepcDB (Kouranov et al., 2006). We used PepcDB downloaded on November 17, 2010, which includes 261 572 targets. A target defines the objective of the crystallization attempt for either a single protein or a collection of proteins. Targets in PepcDB can have either one or multiple trials, each trial representing a set of procedures used to crystallize a target. There are 817 099 trials in our PepcDB dataset. Each trial has information about its current status and, in case if work was finished, the stop status, see Table 1. The stop statuses indicate the step at which the work on a given trial was stopped and the reason of the failure (which divides the work covered in the current status into substeps) or the fact that the trial produced a proper outcome. The majority of trials have the stop status field empty, which makes it impossible to deduce the final outcome of the trial, we cannot be sure whether the experiment was finished, abandoned or is still in progress. Therefore, we selected trials with the completed stop status field, with the exception for trials with the current status ‘in PDB’ or ‘crystal structure’, as they clearly indicate the successful crystallization attempts. Since each trial may concern more than one sequence, we considered each sequence from each trial as a separate trial.

For the set of the non-crystallizable proteins (NCDB), we considered 45/924 trials with the completed stop status field, which includes any of the following: ‘sequencing failed’, ‘cloning failed’, ‘expression failed’, ‘purification failed’, ‘crystallization failed’ or ‘poor diffraction’.

The set of the crystallizable proteins (CDB) was developed using 15 412 trials with the stop status equal to ‘structure successful’, ‘TargetDB duplicate target found’ or ‘PDB duplicate found’ and the trials with the current status ‘crystal structure’ or ‘in PDB’. We did not use trials corresponding to the NMR structures and we disregarded the trials with the ‘other’ or ‘duplicate target found’ stop status.

We filtered both sets to remove the trials with duplicate sequences, i.e. so far we collected all trials with the complete stop (or current) status irrespective of the sequence. Given two trials with the same sequence, we removed the trial with an earlier stop status (Table 1), e.g. given two trials with the same sequence marked with ‘sequencing failed’ and ‘purification failed’ stop statuses, we remove the trial with the ‘sequencing failed’ stop status. In case of two trials with the same stop status, we removed the older trial.

Next, we filtered all chains in the NCDB set against the CDB set and the chains in the PDB, i.e. we remove a given chain from the NCDB set in case if this sequence occurs in the CDB set or in the PDB.

In the next step, we filtered the chains in the NCDB set against all trials in the PepcDB based on their current status field. We removed each non-crystallizable trial for which there is a trial with the same sequence and the current status further along the crystallization process (Table 1). In this case, the current status indicates that the trial succeeded with the stop (stop status) which was used to enter it into the NCDB set.

Next, we removed all trials from before January 1, 2006 and after December 31, 2009. We removed the older samples to accommodate for the latest advances in the crystallization protocols. For example, our analysis of the PepcDB shows that before 2006, i.e. in the first PSI phase, a large number of failures corresponded to problems with cloning, whereas after 2005 the problems with cloning subsided. The samples from 2010 could not be used since some of them may not be yet completed or updated in the database.

We assigned the following classes to the remaining trials: (i) production of the protein material failed (MF) for all trials with stop status ‘sequencing failed’, ‘cloning failed’ and ‘expression failed’; (ii) purification failed (PF) for the ‘purification failed’ stop status; (iii) crystallization failed (CF) (for the ‘crystallization failed’ and ‘poor diffraction’); and (iv) CRYStallizable (CRYS) (for the stop statuses ‘structure successful’, ‘TargetDB duplicate target found’ and ‘PDB duplicate found’; or the current statuses ‘crystal structure’ and ‘in PDB’).

Finally, using BLASTCLUS we reduced the sequence identity among chains within the same class, i.e. for each class we kept only the sequences below 25% sequence identity threshold. This is consistent with the threshold used in the prior studies (Overton and Barton, 2006), but we did not reduce the sequence identity between trials from different classes.

We created three datasets to build predictors for each of the non-crystallizable classes (MF, PF and CF). Each of these datasets includes trials which failed to proceed through a given step (as the samples in the negative set), and trials which passed this step (as the positive set). In the dataset for the prediction of the production of the protein material (DB_MF), the negative set contains all trials labeled as MF and the positive set contains the remaining proteins; for the purification dataset (DB_PF), the negative set contains trials marked as PF and the positive set includes trials from the CF and CRYs classes; for the crystallization dataset (DB_CF), the negative set contains trials with the CF class, and the positive set includes trials from the CRYs class. For the DB_PF and DB_CF datasets, we did not include trials from the MF and the MF and PF classes, respectively, due to the fact that we do not know whether these trials would pass the purification or crystallization steps since they did not pass the previous steps, e.g. we do not know whether the MF trials would purify if they pass the production of the protein material step. We also created the DB_CRYs dataset with the class labels similar with the previous predictors, which indicate the success of the entire process, i.e. production of the diffraction-quality crystals. The DB_CRYs dataset includes two labels, non-crystallizable chains (MF, PF and CF) versus crystallizable chains (CRYs). The chains in

| Table 1. List of stop statuses and current statuses in PepcDB |
|---------------------------------------------------------------|
| Class dedicated from PepcDB annotation | Stop status | Current status |
| Production of protein material failed | Sequencing failed, cloning failed | Cloned |
| Purification failed | Purification failed | Soluble |
| Crystallization failed | Crystallization failed | Crystallized |
| Crystalizable Structure successful, TargetDB duplicate target found, PDB duplicate found | | Crystal structure |
Crystallization propensity prediction

The prediction is performed in two steps: (i) the input sequences are converted into a set of numerical features that describe certain, relevant characteristics of the protein chain; and (ii) the feature values are fed into four predictive models that output the predicted propensity for material production, purification, crystallization and diffraction quality crystallization, respectively; we use a support vector machine (SVM), which was previously shown to provide high-quality predictions in this area (Kandaswamy et al., 2010; Smialowski et al., 2006), to implement these four models and their outputs are aggregated together to provide a four-class prediction. The architecture of the PPCpred method is shown in Figure 1.

2.3 Architecture of the proposed predictor

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2.3.2 Features

We combine information based on the amino acids indices, predicted secondary structure predicted with PSIPRED 3.2 (Jones, 1999). The importance of the information derived directly from the protein chain, including the composition of certain amino acids, the isoelectric point, etc., for prediction of the crystallization success was demonstrated in numerous studies (Chandonia et al., 2006; Chen et al., 2007; Goh et al., 2004; Kandaswamy et al., 2010; Kantardjieff and Rupp, 2004; Kantardjieff et al., 2004; Kurgan et al., 2009; Overton and Barton, 2006; Overton et al., 2008; Slabinski et al., 2007b; Smialowski et al., 2006). The evaluation was performed per-protein at two levels: (i) the binary values that define whether a given trial/sequence is positive (passes a given step) or negative (fails to pass the step) are evaluated using the Matthews correlation coefficient (MCC), accuracy (ACC), Sensitivity and Specificity measures; and (ii) the real values that quantify the probability of prediction which correspond to the propensity of the chain to pass the test, i.e. to produce the protein material, to purify, and to produce crystals, are evaluated with the receiver operating characteristic (ROC) and area under the ROC curve (AUC) measures. Detailed explanation of the above-mentioned quality measures is given in the Supplementary Material. The aggregated predictions of four classes are assessed based on the overall accuracy (number of correctly predicted trials over all trials) and mean MCC (over the four MCC values computed for each class label).

We analyze statistical significance of the differences between PPCpred and the other methods. For each test set, we compared 100 paired results for MCC and ACC obtained using the bootstrapping with 25% of randomly selected trials. Since the measurements follow normal distribution, as tested using Shapiro–Wilk test at the 0.05 significance, we use paired t-test and we measure significance of the differences at the 0.01 level. We designed the proposed predictor, which includes filtration of the considered features, parameterization of the SVM-based classifiers, feature selection and selection of the thresholds for the aggregation of the four predictors (which are described below), using 5-fold cross-validation on the training datasets. In this test, we randomly divide the training dataset into five equal-sized subsets of protein chains. We use four of these subsets to form a training fold that is utilized to compute the model and the fifth subset constitutes the testing fold that is used to perform the evaluation. This is repeated five times, each time choosing a different subset to be the test fold. The tests on the independent test datasets was performed using the model trained on the corresponding training dataset using the parameters and feature sets that were established based on the cross-validation.

Fig. 1. The overall architecture of the proposed PPCpred method.

the DB_CRYS dataset were annotated with the four class labels, i.e. the non-crystallizable chains were annotated as MF, PF or CF, to create the DB_4CL dataset. The number of trials in each dataset is shown in Supplementary Table 1.

We randomly divided the MF, PF, CF and CRYS sets into two equal sized subsets, the training and the test sets. We used the training subsets to create the corresponding training DB_MF, DB_PF, DB_CF and DB_CRYS datasets, and the test subsets to create the independent test datasets. We designed our predictor based on the training datasets (using 5-fold cross-validation protocol) and then we performed evaluation and comparison with the existing methods on the independent test datasets. We note that the sequence identity between chains from the same class in the training and test sets is <25%.

A. The predicted propensity for the material production

B. The predicted propensity for the purification

C. The predicted propensity for the crystallization

D. The predicted propensity for diffraction quality crystallization

E. Aggregation of the predicted propensities

F. The overall architecture of the proposed PPCpred method.
solvent accessibility is <25%; otherwise a given residue is assumed to be
solvent exposed. In total, we generated 828 features, which include:

- AA {AA} Composition of the 20 standard amino acid (AA) types, i.e.
  the count divided by the sequence length, where AA stands for one
  of 20 AAs (20 features).
- AA {exp,bur},{AA} Composition of the exposed/buried AAs (count
  of the exposed/buried AA divided by the number of all exposed/buried
  residues in a given chain) (40 features).
- p The p-electrolyte point (1 feature).
- [AAIndex] The average value of a given amino acids index [AAIndex]
  over the whole sequence (65 features).
- [AAIndex] {min,max},{5,10,15,20} The minimal/maximal average
  value of the amino acid index [AAIndex] among all sliding windows
  of sizes 5, 10, 15 and 20. For chains shorter than a given window
  size, we use the window size equal the length of the sequence. These
  features are motivated by the work in (Iliev and Jachinmaci, 2010)
  (65 × 4 × 2 = 520 features).
- [AAIndex] {exp,bur} The summed value of the amino acid index
  [AAIndex] for exposed/buried residues, divided by the number of
  exposed/buried residues in the sequence. These features are motivated
  by the work in (Price et al., 2009) (65 × 2 = 130 features).
- DIS_AVG_VAL The average value of the predicted disorder
  probabilities (1 feature).
- DIS_SEG Number of the predicted disorder segments (1 feature).
- DIS_RES_{seg},{1,5,10,15,20} Number of the predicted disorder
  residues in the disorder segments which are at least 1, 5, 10, 15 and
  20 residues long, divided by the sequence length. For segments with
  at least one residue, this feature represents content of the predicted
  disorder. (5 features).
- DIS_avg The average length of the predicted disorder segments divided
  by the sequence length (1 feature).
- DIS_max The maximal length of the predicted disorder segment
divided by the sequence length (1 feature).
- DIS_{exp,bur} Number of the predicted exposed/buried residues
  divided by the number of exposed/buried residues (2 features).
- DIS_{exp,bur} AVG_VAL The summed value of the predicted disorder
  probability for the predicted exposed/buried residues divided by the
  number of predicted exposed/buried residues (2 features).
- SS {SS},RES_{seg},{1,5,10,15,20} Number of residues in the predicted
  coil/helix/strand segments, SS i ∈ {C, H, E}, which are at least 1, 5,
  10, 15 and 20 residues long, divided by the sequence length. For
  segments with at least one residue, these features represent content of
  the predicted coils, helices and strands (15 features).
- SS {SS},avg The average length of the predicted SS segments divided
  by the sequence length (3 features).
- SS {SS},max The maximal length of the predicted SS segments
  divided by the sequence length (3 features).
- SS {exp,bur},SS AVG_VAL The average predicted probability be in
  the secondary structure state SS i (3 features).
- SS {exp,bur},SS Number of the predicted exposed/buried residues
  in the secondary structure state SS i divided by the number of
  exposed/buried residues (6 features).
- RSA_AVG_VAL The average value of predicted relative solvent
  accessibility (1 feature).
- EXP {BUR},RES_{seg},{1,5,10,15,20} Number of the predicted
  exposed/buried residues in the exposed/buried segments which are
  at least 1, 5, 10, 15 and 20 residues long divided by the sequence
  length. For segments with at least 1 residue, these features represent
  content of the exposed/buried residues. We note that there were no predicted
  exposed segments with over 15 residues, and thus the corresponding
  two features were removed (8 features).

2.3.3 Filtration of the considered features We note that some of the
considered features may be correlated with each other and may not be useful
to differentiate between the considered class labels (i.e. the annotation of
the protein production, purification, crystallization and diffraction-quality
crystalization, respectively). We performed filtration to remove the highly
correlated and low-quality features. The training dataset was divided at
random into five training and test fold (as in the 5-fold cross-validation)
and we ranked the features according to their average, over the five training
folds, bieral correlation with the class labels. We selected the feature
with the highest average bieral correlation, and we added the next ranked
feature into the set of the selected features only if the Pearson correlation
coefficient of this feature with every feature in the selected feature set
was <0.7. This step removed the highly correlated features. Next, we
computed the average value of the average (over the 5-folds) absolute
bieral correlations (with the class labels) for the selected features, and
we removed the features with the correlations below the average. The latter
step removes the low-quality features. At the end, we selected 86, 100, 115
and 95 features for the DB_MF (material production), DB_PF (purification),
DB_CF (crystallization) and DB_CRYS (diffraction-quality crystallization)
training datasets, respectively.

2.3.4 Parametrization and feature selection For each dataset we built
three SVM models, implemented in the LibSVM package (Chang and Lin,
2001), that are based on the three available kernels including the radial basis
function (RBF), polynomial (POLY), which also includes the linear kernel,
and sigmoid (SIG). We computed total of 12 models (three different kernels
for four datasets: DB_MF, DB_PF, DB_CF and DB_CRYS). Each model
was built using the same procedure. First, the model for a given dataset was
parameterized using 10 features with the highest average bieral correlation
from the features selected in Section 2.3.3. Using 5-fold cross-validation
on the training set, we performed grid search to find parameters that maximize
the MCC. For each parameter, except the degree of the polynomial, we
considered consecutive powers of 2. In case when the selected parameter
values were at the border of the search grid, we extended the search to the
next consecutive power of 2. The selected parameters were used to perform
the wrapper-based feature selection (Hall and Smith, 1999) using the best
first search strategy. Supplementary Material provides detailed description
of the considered parameters and feature search procedure. After the feature
selection is completed, we parameterized the SVMs using the selected feature
set and the same grid search as above. Finally, we select the SVM that
provides the highest MCC (among the three kernel types) for each dataset,
see Supplementary Table 3, i.e. we use the POLY kernels for the prediction of
diffraction-quality crystalization, purification and crystalization, and RBF
kernel for the prediction of the material production.

2.3.5 The 4-class prediction We aggregate outputs from the best four
SVM models developed in Section 2.3.4 to perform the 4-class prediction.
We investigated two approaches to aggregate predictions: (i) by selecting
the class with the maximal predicted probability (max-based), and (ii) by
choosing the class based on the order of the steps in the crystallization
protocol (order-based). In the latter case, we select the class by checking
the outputs of the SVMs in the order defined in Table 1, from MF to
PF, to CF and to CRYS. We predict the corresponding outcome if the
predicted probability is above a cut-off threshold, e.g. we predict that the
material production fails if the output from the corresponding SVM, which
quantifies the probability of the material production failure, is greater that
the threshold. If none of the probabilities are above the threshold then we
select the outcome with the highest probability. We tried the cut-off values
from between 0.01 and 1 with step 0.01 and selected the value = 0.43 that
maximizes the mean MCC over the four classes on the DB_APL training
dataset. The max-based aggregation obtains the mean MCC and accuracy
equal 0.293 and 49.0%, respectively, while the order-based method obtains...
The proposed PPCpred method, which utilizes the order-based crystallization propensity including OBScore (Overton and Barton, 1997), and we use the class label of the most similar chain as the prediction. In the case when no alignments are found, the test chain is predicted with the label of the most probable crystallizable chain. This is repeated for each of the five training and test dataset pairs, DB_MF, DB_PF, DB_CF, DB_Crys and DB_4CL.

3 RESULTS

The proposed PPCpred method, which utilizes the order-based aggregation of the propensities predicted by the four SVM models, was evaluated on the test datasets for the predictions of each of the four outcomes, i.e. prediction of material production, purification, crystallization and diffraction-quality crystallization, as well as for the 4-class prediction. We compare the results generated by the PPCpred with the existing methods for the prediction of the crystallization propensity, with the BLAST-based solution, and with the maximum-based aggregation scheme.

3.1 Comparison of the diffraction-quality crystallization propensity predictions

The PPCpred is compared with the recent predictors of the crystallization propensity including OBScore (Overton and Barton, 2006), XtalPred (Slabinoski et al., 2007b), ParCrys (Overton et al., 2008), CRYSTALP2 (Kurgan et al., 2009), MetaPPCP (Mizianty and Kurgan, 2009) and SVMCrys (Kandaswamy et al., 2010), with the BLAST-based method, and with our SVM predictor of the diffraction-quality crystallization (SVM_POLY), see the results on the DB_Crys dataset in Table 2. The PPCpred outperforms the existing solutions in both the binary prediction (based on the MCC and ACC scores) and the real-valued propensities (based on the AUC values). The best, existing predictor is XtalPred, which is likely due to the usage of the sequence alignment against the PDB and nr databases, followed by SVMCrys and MetaPPCP. The PPCpred improves over the SVM_POLY method, which demonstrates that aggregation of the results from the four SVMs is helpful. Also, the maximum-based aggregation is shown to be inferior to the order-based aggregation used in the PPCpred for the binary predictions, but the magnitude of this difference is relatively small. Table 2 shows that the improvements in MCC and ACC offered by PPCpred are statistically significant. The binary predictions from PPCpred are characterized by high specificity (high success rate among the native non-crystallizable proteins) at about 85%. This means that we relatively rarely mispredict these chains to be crystallizable, which would save resources to solve other chains.

The ROC curves of the considered predictors, except for the BLAST and SVMCrys that provide only the binary predictions, are shown in Supplementary Figure 1. PPCpred outperforms the other three classes considered in this work. Therefore, we consider a baseline score implemented using sequence alignment to comparatively evaluate the predictive quality of our method. Each test trial chain was aligned against the sequences in the corresponding training dataset using PSI-BLAST (Altschul et al., 1997) and we use the class label of the most similar chain as the prediction. In the case when no alignments are found, the test chain is predicted with the label of the most probable crystallizable chain. This is repeated for each of the five training and test dataset pairs, DB_MF, DB_PF, DB_CF, DB_Crys and DB_4CL.

# Table 2. Summary of results for the prediction of the propensity of the diffraction-quality crystallization success (based on the DB_Crys test dataset), the prediction of the propensity of the material production failure (DB_MF test set), the prediction of the propensity of the purification failure (DB_PF test set) and the prediction of the propensity of the crystallization failure (DB_CF test set)

| Test dataset (propensity target) | Method     | MCC     | ACC     | SPEC   | SENS   | AUC   |
|---------------------------------|------------|---------|---------|--------|--------|-------|
| DB_Crys                         | ParCrys    | 0.108   | 47.5    | 31.8   | 78.6   | 0.561 |
| (propensity of crystallization success) | OBSCore | 0.124   | 47.8    | 31.4   | 80.3   | 0.572 |
|                                 | BLAST-based| 0.188   | 65.6    | 79.5   | 58.0   | N/A   |
|                                 | CRYSTALP2  | 0.198   | 55.3    | 45.7   | 74.4   | 0.648 |
|                                 | MetaPPCP   | 0.195   | 59.9    | 59.0   | 61.7   | 0.620 |
|                                 | SVMCrys    | 0.213   | 56.3    | 46.7   | 75.2   | N/A   |
|                                 | XtalPred   | 0.278   | 63.9    | 62.3   | 67.0   | 0.683 |
|                                 | SVM_POLY   | 0.398   | 74.6    | 87.9   | 57.9   | N/A   |
|                                 | max-based  | 0.467   | 76.1    | 81.6   | 65.3   | 0.793 |
|                                 | PPCpred    | 0.471   | 76.8    | 84.8   | 61.2   | 0.789 |
| DB_MF (propensity of the material failure) | BLAST-based | 0.014   | 55.4    | 35.3   | 60.0   | N/A   |
|                                 | max-based  | 0.339   | 71.6    | 45.4   | 85.5   | 0.621 |
|                                 | PPCpred    | 0.462   | 75.0    | 69.2   | 78.0   | 0.755 |
| DB_PF (propensity of the purification failure) | SVM_RBF | 0.423   | 74.6    | 56.1   | 64.5   | 0.791 |
|                                 | PPCpred    | 0.462   | 75.0    | 69.2   | 78.0   | 0.755 |
|                                 | max-based  | 0.246   | 70.8    | 34.4   | 86.9   | 0.609 |
|                                 | SVM_POLY   | 0.290   | 73.2    | 30.8   | 91.8   | 0.741 |
|                                 | PPCpred    | 0.328   | 72.0    | 50.1   | 81.6   | 0.697 |
| DB_CF (propensity of the crystallization failure) | BLAST-based | 0.060   | 60.9    | 34.1   | 69.4   | N/A   |
|                                 | SVM_POLY   | 0.346   | 77.0    | 40.1   | 90.0   | 0.814 |
|                                 | PPCpred    | 0.457   | 76.6    | 70.8   | 87.3   | 0.811 |
|                                 | max-based  | 0.464   | 76.9    | 70.5   | 79.2   | 0.813 |

The proposed PPCpred is compared against results on the DB_Score, XtalPred, ParCrys, CRYSTALP2, MetaPPCP and SVMCrys on the DB_Crys dataset, and against the maximum-based aggregation method (max-based), the best performing SVM classifier (SVM_POLY or SVM_RBF), and the BLAST-based predictor on the four datasets. The methods are sorted in the ascending order based on their MCC scores, and the highest values for each quality index and dataset are shown in bold. The BLAST and SVMCrys provide only binary prediction and thus we could not compute their AUC. Results of tests of significance of the differences in MCC and ACC between PPCpred and the other methods are given in the ‘sig’ columns. The tests compare values over 100 bootstrapping repetitions. The ‘+’ and ‘-’ means that PPCpred is statistically significantly better/worse with P < 0.01, and ‘=’ means that results are not significantly different.
The results for each individual target outcome of the PPCpred, trials (crystallization propensity predictors with respect with the date of the test improvement with small magnitude that is statistically significant. with the maximum-based aggregator for the prediction of the crystallization, and it provides comparable predictive quality of the material production, purification and diffraction-quality significantly outperforms the other methods for the binary prediction are summarized in Table 2. Using the MCC measure, PPCpred maximum-based method for combining the four SVMs predictors (SVM_POLY and SVM_RBF), and the BLAST-based predictors, our four SVM-based predictors of the material production, purification, crystallization and diffraction-quality shows that the success/failure in the considered steps of the crystallization process depends on a combination of multiple factors. The overall accuracy of PPCpred equals 55.6%, which is higher by 5 and 21% than the accuracy of the other two solutions. The improvements are statistically significant. Only the PPCpred, the alignment based predictor, and the maximum-based aggregation method can be compared—the other methods predict only one of the outcomes. The overall accuracy of PPCpred and ACC between PPCpred and the other methods are shown in the 'sig' columns. The methods compare values over 100 bootstrapping repetitions. The ‘+’ and ‘-’ mean that PPCpred is statistically significantly better/worse with \( P < 0.01 \), and ‘=’ means that results are not significantly different.

PPCpred also provides well-balanced values of the sensitivity and specificity. Our method provides reasonably high values of MCC, between 0.32 and 0.47, which indicate that it provides useful outputs. The evaluation for the 4-class predictions on the DB_4CL test dataset is shown in Table 3. The output of the predictor indicates whether a given chain will provide high-quality crystal, will fail to crystallize, or whether the purification or material production will fail. The methods are evaluated using the overall accuracy (fraction of the correctly predicted chains) and mean MCC (over the four MCC values computed for each class/outcome). Only the PPCpred, the alignment based predictor, and the maximum-based aggregation method can be compared—the other methods predict only one of the outcomes. The overall accuracy of PPCpred equals 55.6%, which is higher by 5 and 21% than the accuracy of the other two solutions. The improvements are statistically significant. We believe that this level of predictive quality should be acceptable for the potential users given the current crystallization success rates, which are at about 4.6% (Kurgan and Mizianty, 2009).

3.2 Comparison of the prediction of the propensity of material production, purification, crystallization and diffraction-quality crystallization

The results for each individual target outcome of the PPCpred, BLAST-based predictors, our four SVM-based predictors of the material production, purification, crystallization and diffraction-quality crystallization (SVM_POLY and SVM_RBF), and the maximum-based method for combining the four SVMs predictors are summarized in Table 2. Using the MCC measure, PPCpred significantly outperforms the other methods for the binary prediction of the material production, purification and diffraction-quality crystallization, and it provides comparable predictive quality with the maximum-based aggregator for the prediction of the crystallization, i.e. the maximum-based aggregator provides an improvement with small magnitude that is statistically significant.
Table 4. Summary of the features types selected for the prediction of the material production, purification and crystallization

| Features types       | Number of features selected for the prediction of material production | Purification | Crystallization | Diffraction-quality crystallization |
|----------------------|-----------------------------------------------------------------------|--------------|----------------|------------------------------------|
| Hydrophobicity index | 2                                                                     | 2            | 5              | 5                                  |
| Energy-based index   | 4                                                                     | 0            | 2              | 3                                  |
| Composition of AAs   | 1                                                                     | 3            | 1              | 1                                  |
| Isoelectric point    | 0                                                                     | 1            | 0              | 0                                  |
| Solvent accessibility| 3                                                                     | 4            | 1              | 3                                  |
| Disorder             | 1                                                                     | 0            | 1              | 1                                  |
| Secondary structure  | 0                                                                     | 0            | 0              | 1                                  |
| Considered AA types  | Arg, Cys, Asn, Cys, Ser, His, Cys, His, Ser                            |              |                |                                    |

also suggest the importance of Cys residues for the prediction of the material production and diffraction-quality crystallization, and buried Cys for the prediction of purification. This agrees with the observations in (Overton et al., 2008; Slabinski et al., 2007b), but these studies investigated only the propensity for the diffraction-quality crystallization and did not consider the influence of the solvent accessibility. Another factor related to the crystallization success is the content of the buried His. This agrees with results in Overton et al. (2008) and Kurgan et al. (2009), but these studies again considered the overall content of His, without the influence of the solvent accessibility.

Figure 3 shows scatter plots of three pairs of features that were selected for the prediction of the crystallization, diffraction-quality crystallization and purification, respectively. The two features used to predict crystallization, GOLD730101_min_10 and WERD780103_min_5 (Fig. 3A), and based on the minimal average values of the hydrophobicity (Goldsack and Chalifoux, 1973) and energy (specifically the energy of transfer in water of an isolated residue from a non-regular structure to the helical conformation) (Wertz and Scheraga, 1978) indices in the sliding windows of sizes 10 and 5, respectively. This means that the sequence segments with low hydrophobicity and transfer energy values are characteristic to chains that are difficult to crystallize. Importantly, combining these two features allows for improved separation between the successful and unsuccessful crystallization trials, i.e. trials for a given range of values of one index are further separated by the values of the other index. The diffraction-quality crystallization is impacted by the DIS_SEG and AA_bur_S features (Fig. 3B), which quantify the number of the predicted disorder segments and the content of the predicted buried Ser, respectively. The content of Ser was shown to be important for the prediction of crystallization propensity in Overton et al. (2008) and Kurgan et al. (2009), but these studies investigated the overall Ser content, while we show that the (predicted) buried Ser provides strong discriminatory power. Similarly, while the content of the predicted disordered residues was used in several related studies (Slabinski et al., 2007b; Price et al., 2009), our analysis reveals the strong influence of the number of disordered segments. The plot shows that chains with larger number of disordered segments and larger number of buried Ser are more difficult to crystallize. Finally, Figure 3C shows that chains with larger amount of buried Ser (AA_bur_S feature) and high hydrophobicity in a long-sliding window (GOLD730101_max_20 feature) are more challenging to purify.

Overall, the factors that we identified are intuitive, physically reasonable and they are well aligned with the existing ‘rules of thumb’. Our main contributions are in providing additional details (e.g. related to solvent accessibility of selected residues types) and the fact that our model provides a novel way of balancing these factors to obtain good predictive performance.
4 CONCLUSIONS

We developed a first-of-its-kind in silico method, PPCpred, which predicts the success/failure for four main steps in the protein crystallization protocols, including the material production, purification, crystallization and diffraction-quality crystallization. PPCpred significantly outperforms the alignment-based predictor as well as the several modern crystallization propensity predictors. Our method provides the overall accuracy at 56% and average MCC at 0.35, which given current low success rates in the experimental protocols should provide useful input for the SG centers and crystallographers/biologists who are interested in participation in the PSI:Biology phase. We also developed an improved protocol to annotate progress of protein chains along the crystallization process using the PepDB, and we shows/confirm several interesting markers (based on the features included in our predictors) that influence the success/failure of the above-mentioned steps.

The predictions generated by PPCpred could be used to guide crystallographers to select more feasible alternative targets or, in case when the target is already selected, to rank different constructs of the same target. The former application is evaluated in our work, while the evaluation of the latter one will be performed in a feature study when large enough amount of suitable data becomes available. Users of PPCpred could also find out which of the crystallization steps is the most likely obstacle in the crystallization process, and try to modify the target to increase chances to pass that step, e.g. the user may introduce tags at sequence termini to ease purification when the purification failure is predicted. The success of crystallization also depends on the crystallization protocols. Our method was designed using data from several SG centers, which allows us to generalize over multiple protocols. At the same time, our model takes into account only the intra-molecular factors that are encoded in the protein chain. Therefore, the PPCpred as well as the other crystallization propensity predictors may not provide reliable predictions when the inter-molecular factors such as the specific characteristics of the expression systems, proteins-protein and/or protein-precipitant interactions, buffer composition, precipitant diffusion method, gravity, etc., must be considered.

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