DOMAC: an accurate, hybrid protein domain prediction server

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ABSTRACT

Protein domain prediction is important for protein structure prediction, structure determination, function annotation, mutagenesis analysis and protein engineering. Here we describe an accurate protein domain prediction server (DOMAC) combining both template-based and *ab initio* methods. The preliminary version of the server was ranked among the top domain prediction servers in the seventh edition of Critical Assessment of Techniques for Protein Structure Prediction (CASP7), 2006. DOMAC server and datasets are available at: http:// www.bioinfotool.org/domac.html.

INTRODUCTION

Protein domains are structural, functional and evolutionary units of proteins. The prediction of domains from sequence information can improve tertiary structure prediction (1), enhance protein function annotation (2), aid structure determination (3) and guide protein engineering (4) and mutagenesis (5).

A number of different methods have been developed to identify domains starting from primary sequences. These methods can be roughly classified into four categories: template-based methods (6–10), *ab initio* (template-free) methods (11–22), the hybrid approach combining template-based and *ab initio* methods (23), and meta-domain prediction methods (24).

Here we describe an accurate, hybrid domain prediction server (DOMAC) that integrates homology modeling, domain parsing and *ab initio* methods together. The preliminary implementation of the server [under the name: FOLDpro (25)] participated in the domain evaluation in the seventh edition of Critical Assessment of Techniques for Protein Structure Prediction (CASP7) (26,27). It was ranked among the top domain prediction servers in CASP7.

IMPLEMENTATION

Our hybrid approach uses the template-based method to predict domains for proteins having homologous template

structures in Protein Data Bank (PDB) (28), and the *ab initio* method based on neural networks (29) to predict domains for *de novo* proteins. It predicts protein domains in two steps.

First, it uses the PSI-BLAST (30) to search the target sequence against NCBI Non-Redundant sequence database to construct a profile. The profile is used to search a template structure library built from the proteins in PDB to identify templates, similarly as PDB-BLAST approach (31).

Second, if some significant templates are identified (*e*-value ≤ 0.001), it generates a structure model for the target using Modeller (32) based on the template structures. Multiple significant templates are combined to improve model quality if available. Then it uses an accurate domain parsing tool PDP (33) to parse the model into domains. If the parsed domains do not cover the whole target sequence, DOMAC will assign uncovered regions to adjacent domains.

If no significant homologous template is found, DOMAC will invoke the *ab initio* domain predictor DOMpro (29) to predict domains. DOMpro uses neural networks in conjunction with sequence profile, predicted secondary structure, and relative solvent accessibility to predict domain boundary. The secondary structure and relative solvent accessibility are predicted by SSpro (34) and ACCpro (35) in the SCRATCH suite (36). DOMpro tries to identify domain boundary positions based on the composition bias of sequence and structural features in domain linker regions.

The preliminary implementation of DOMAC participated in CASP7 and was ranked first among 13 domain prediction servers. Since then, we have significantly speeded up the template identification process without sacrificing accuracy and added a module to update the template library weekly to incorporate the newly released proteins in PDB.

RESULTS

Here we firstly describe the performance of the preliminary implementation of DOMAC in CASP7 (under server name: FOLDpro). We compare it with 12 other server

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Table 1. The performance of 13 domain prediction servers in CASP7

Method	Target Num	Domain Num Acc. (%)	CASP7 Score	
FOLDpro (DOMAC)	95	93.7	0.963	
Baker-RosettaDom (23)	94	86.2	0.940	
Ma-OPUS-DOM	94	87.2	0.933	
ROBETTA-GINZU (23)	94	84.0	0.932	
DomSSEA (7)	94	78.7	0.910	
HHpred3 (38)	95	75.8	0.910	
Meta-DP (24)	95	74.7	0.907	
HHpred1 (38)	93	75.3	0.902	
DomFOLD	95	75.8	0.898	
DPS(13)	93	75.3	0.889	
Chop (22)	83	56.6	0.827	
Distill (39)	95	70.5	0.819	
NN_PUT-Lab	92	58.7	0.795	

The second column (target num) lists the number of targets for which a predictor made predictions.

predictors in CASP7 using two evaluation metrics: CASP evaluation metric (37) and domain number accuracy.

CASP metric (NDO: normalized domain overlap score) is to compute the overlapping score of domains without explicitly checking domain number and domain boundary (37). It computes the numbers of correctly and wrongly overlapped residues between true domains and predicted domains, respectively. It summarizes the numbers of the overlapping residues into a single score to evaluate domain prediction. The best score for a target is 1 and the worst score is 0. The domain number accuracy is defined as the percentage of targets with correct domain number predictions.

Table 1 reports the performance of 13 servers on 95 targets in CASP 7. The CASP score is the average domain overlap score across all predicted targets. The domain number accuracy is computed by comparing the domain number predictions with the official domain definitions released by CASP7. In terms of the two evaluation metrics, the preliminary implementation of DOMAC (FOLDpro) yielded the best performance.

We also evaluate DOMAC on the three categories of CASP7 targets: highly homologous, homologous and analogous/*ab initio*. The domain number prediction accuracy of DOMAC is 96%, 94% and 88% in the three categories, respectively.

However, because the majority (68 out of 95) of CASP7 targets is single-domain proteins, the domain prediction accuracy is very likely *over-estimated*.

Thus, we evaluate DOMAC on a larger, balanced, highquality dataset manually curated by Holland *et al.* (2). The publicly released version of the Holland's benchmark2 dataset has 156 proteins consisting of 54 single-domain proteins, 69 two-domain proteins, 25 three-domain proteins, 4 four-domain proteins, 3 five-domain proteins and 1 six-domain protein. We evaluate both templatebased and *ab initio* methods on the whole dataset, respectively. Table 2 reports the specificity and sensitivity of each method in each category in terms of domain numbers. The overall domain number prediction accuracy of the template-based and *ab initio* methods is 75% and 46%, respectively.

Table 2.	The s	pecificit	y and	sens	sitivity	of doma	in nu	mber	pree	diction	on
the Holl	and's	dataset	using	the	templa	ate-based	and	ab in	itio	metho	ds

Method	Acc. (%)	1-dom	2-dom	3-dom	4-dom	5-dom	6-dom
Template	Sens. Spec.	96.1 74.2	66.7 88.0	56.0 70.0	75.0 42.9	66.7 33.3	_
Ab initio	Sens. Spec.	88.5 46.5	31.3 48.8	12.0 30.0	_	_	_

Moreover, we assess the accuracy of the domain boundary prediction, which is important for generating hypotheses for crystallizing individual protein domains. Following the same convention (7,22), a predicted boundary within 20 residues away from a true domain boundary is considered correct.

The domain boundary specificity and sensitivity is 50% and 76.5% for the template-based method, and 27% and 14% for the *ab initio* method. Thus, the accuracy are sufficient for guiding the crystallization experiment, whereas the *ab initio* method is not always reliable enough for the general, practical use.

USE OF WEB SERVICE

The use of DOMAC are intuitive through a simple input form. Since the reliability assessment of domain predictions is still an open issue, the user is advised to use the accuracy on the Holland's dataset to decide how to use these predictions. The input form requires only three inputs: email address, target name, and protein sequence. DOMAC usually can make predictions within 15 min and send the results back to users through email.

Domain prediction results include the user-defined target name, the protein sequence, the predicted domain number, the start and end positions of each domain and the method (template-based or *ab initio*). For template-based prediction, it also reports the PDB codes of the templates. Figure 1 shows an output example for the CASP7 target T0324.

CONCLUSION AND FUTURE WORK

We have developed a hybrid domain prediction web service integrating template-based and *ab initio* methods. The template-based method is accurate enough for guiding protein structure prediction, structure determination, function annotation, mutagenesis analysis and protein engineering. However, the *ab initio* method still needs to be improved for practical use. Since protein domain architecture is largely shaped by gene recombination events, such as gene fusion, fission, domain swapping and exon exchange, leveraging the evolutionary gene recombination signals embedded in the multiple sequence alignment of a protein family and exon boundaries (or splicing sites) in its gene structure, may help improve *ab initio* domain prediction significantly.

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Target: T0324

Sequence: MTYQALMFDIDGTLTNSQPAYTTVMREVLATYGKPFS PAQAQKTFPMAAEQAMTELGIAASEFDHFQAQYEDVMASHYDQI ELYPGITSLFEQLPSELRLGIVTSQRRNELESGMRSYPFMMRMAV TISADDTPKRKPDPLPLLTALEKVNVAPQNALFIGDSVSDEQTAQA ANVDFGLAVWGMDPNADHQKVAHRFQKPLDILELFK

domain num: 2 domain 1: 1-16, 82-208 domain 2: 17-81

The domain prediction is made by the template-based method using the following protein templates in PDB: 2HSZA, 1JUDA, 1ZRNA, 2GFHA, 2AH5A, 2HDOA, 1008A, 1Z4NA, 1Z4OB.

Figure 1. Domain prediction result of CASP7 target T0324. The protein is predicted to have two domains. Domain 1 has two non-continuous segments, spanning from residues 1 to 16 and residues 82 to 208, respectively. Domain 2 spans from residues 17 to 81. The templates used to make the domain prediction are identified by PDB code + chain id. The chain in a single-chain protein is always assigned chain id 'A' instead of '-'.

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