Protein annotation and modelling servers at University College London


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ABSTRACT
The UCL Bioinformatics Group web portal offers several high quality protein structure prediction and function annotation algorithms including PSIPRED, pGenTHREADER, pDomTHREADER, MEMSAT, MetSite, DISOPRED2, DomPred and FFPred for the prediction of secondary structure, protein fold, protein structural domain, transmembrane helix topology, metal binding sites, regions of protein disorder, protein domain boundaries and protein function, respectively. We also now offer a fully automated 3D modelling pipeline: BioSerf, which performed well in CASP8 and uses a fragment-assembly approach which placed it in the top five servers in the de novo modelling category. The servers are available via the group web site at http://bioinf.cs.ucl.ac.uk/.

INTRODUCTION
As the rate of deposition of new protein sequences outstrips the rate at which new protein structures are deposited in the public databases, there will be a continued need for accurate protein structure prediction. It is also arguable that the need for novel function annotation tools is even more pressing. The UCL Bioinformatics Group web portal aggregates a range of methods, developed and maintained at UCL, which predict key structural features of proteins from either their primary structure (sequence) or tertiary structure. We have recently revamped all of our web tools and have taken the opportunity to port all of our services to a new unified server architecture. The services are now presented with a user-friendly common look and feel and we have greatly increased our capacity to serve requests to the community.

ALGORITHMS
This section gives an overview of the algorithms and services available via the UCL Bioinformatics Group web server (http://bioinf.cs.ucl.ac.uk/web_servers/).

PSIPRED is a secondary structure prediction method (1) which uses a two-stage neural network to predict secondary structure using PSI-BLAST output (2). PSIPRED V3.0 currently offers the highest available three-state accuracy (Q3) for secondary structure prediction of 81.4% (±0.6%). This year, in addition to improvements to the underlying algorithm, we have also made some minor updates to the PSIPRED web output, where we now provide coloured alignment schematic diagrams alongside the original secondary structure ‘cartoons’.

MEMSAT3 and MEMSAT-SVM
Transmembrane topology prediction is provided by our MEMSAT methods. Our server now supports two algorithms: MEMSAT3 (3) and MEMSAT-SVM (4). Predictions made with MEMSAT3 and MEMSAT-SVM begin with the generation of a PSI-BLAST profile (PSSM). Both algorithms employ dynamic programming to enumerate all possible transmembrane topologies, but differ in the method used to score the different topologies. MEMSAT3 uses a neural network for scoring, and MEMSAT-SVM makes use of a support vector machine (SVM) along with additional target features such as signal peptides and re-entrant helices. Benchmarked with full cross-validation on a data set composed solely of sequences with crystal structures available to validate their topologies, MEMSAT3 and MEMSAT-SVM achieved 76 and 89% accuracy, respectively. Additionally, MEMSAT-SVM was able to predict both signal peptides (93% accuracy) and re-entrant helices (44% accuracy). When run via the portal, both MEMSAT3 and MEMSAT-SVM are run simultaneously and the results

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produced allow the user to compare both predictions. Results may be returned by email and the web presentation displays both the schematic and cartoon diagrams of the predictions (Figure 1).

**DISOPRED2** allows the user to predict disordered regions of proteins (5). Disordered regions are known to play important roles in protein–protein interactions, cell signalling, enzyme kinetics, signalling and a range of other processes. Disordered regions are characterized as those regions which do not have static structure and are believed to move continually through different configurations when the protein is in its functional configuration. **DISOPRED2** generates a **PSSM** matrix and then analyses this with an appropriately trained **SVM**. **DISOPRED** accuracy has previously been shown to be ~93.1%. Our updated results presentation on the web now features a simple schematic which clearly displays the location of the disordered residues.

**Figure 1.** The diagram produced by the MEMSAT-SVM algorithm available via the PSIPRED server. (a) A schematic diagram of the transmembrane regions is presented. Directly below this a trace of the Kyte–Doolittle hydropathy plot (18) and below that four traces of the outputs for the four **SVMs** used to make the transmembrane region assignment are shown. (b) A cartoon of the transmembrane helix topology summarizing the linear coordinates for the helices and indicating where the protein’s extra- and intercellular regions are.
DomPred

DomPred is a protein domain boundary prediction service which presents the results of two different domain boundary prediction methods, DPS (Domains Predicted from Sequence) and DomSSEA (6). DPS is a protein domain homology based method which uses PSI-BLAST to find sequential discrete regions of domain alignments along a query sequence and is capable of resolving domain boundaries even when no close structural relatives are known. DomSSEA is a domain recognition method which attempts to estimate the location of domain boundaries using distant homologous domains. This method requires that a given domain already be present in the domain databases. Our new server makes only minor cosmetic changes to the presentation of these results.

pGenTHREADER and pDomTHREADER

The latest implementations of the GenTHREADER algorithm combine profile–profile alignments with secondary structure- based gap penalties, pair-wise and solvation-based potentials. A linear regression SVM is used to predict protein fold. Our new method, pDomTHREADER (7), is capable of discriminating protein domain superfamilies within a query sequence. The performance of pDomTHREADER exceeds that of most related methods. We have greatly improved the web-based presentation of the results. Results now feature external links to other structural resources including PDBSum (8), SCOP (9) and CATH (10). The sequence alignments are now presented in a fully interactive form using JalView (11); and we also now include links which allow users to explore functional and structural data derived from both the CATH and Gene3D databases (12). The integration with other resources gives the user a richer experience, enabling researchers to move easily from predicted fold through to putative function. In the future we hope to integrate a much wider range of structural and functional annotations into the results.

BioSerf

BioSerf is our new fully automated de novo and homology modelling pipeline available via the World Wide Web or as an XML-driven web service (manuscript in preparation). BioSerf uses a selection of algorithms including PSI-BLAST, PSIPRED and pGenTHREADER to attempt to intelligently select appropriate template structures for homology modelling. Where a suitable template or templates can be found, BioSerf uses MODELLER (13) to build a model. Where no suitable template structure or insufficient coverage can be found, FRAGFOLD (14) is used to create an ab initio template (Figure 2). As this service uses MODELLER, a valid MODELLER licence key is required to submit sequences to the service. Users receive a PDB file as the primary result along with the intermediate results used to generate the model. These intermediate results and the model itself can be inspected via a web page.

Figure 2. The process flow when analysing a sequence using the BioSerf pipeline is shown. An incoming sequence is initially analysed by both PSI-BLAST, PSIPRED and pGenTHREADER. The PSI-BLAST run allows the pipeline to identify whether the query sequences closely matches the sequence of a known structure in the PDB. PSIPRED and pGenTHREADER attempt to identify any remote structural homologues of the sequence should the PSI-BLAST run fail to find any related sequences. These data are then combined and potential template structures for modelling are selected given suitable cut-offs. Where there is at least one suitable template, the pipeline then uses MODELLER to generate a homology model. Where there are no suitable template structures, FRAGFOLD is used to build a possible model structure. A typical run of the pipeline that finds a suitable template structure takes around 10 min. In those cases where no template can be found and FRAGFOLD is used, runs can take up to 3 h.

MetSite

MetSite (15) is a fully automatic method for predicting clusters of metal-binding residues. It is highly sensitive and works well even on protein models of moderate resolution. Sequence profile information derived from PSI-BLAST is analysed using several neural network classifiers, one classifier for each metal atom type. These classifiers are capable of distinguishing metal binding sites from non-metal sites at a mean accuracy of ~94.5%. Users receive a PDB file as the main output with the residue relative-temperature feature used to annotate the prediction. Via the web site, users can explore the annotation in the Jmol applet provided.

FFPred

The FFPred server (16) offers a method for predicting protein function using a machine learning approach. FFPred attempts to assign reliable Gene Ontology (GO) classes (17) to queries using a series of SVM classifiers. Query sequences are annotated with a large set of possible features including such data as amino acid content, transmembrane regions and disordered regions. Every GO class is represented by five SVM classifiers and the sequence is scored for each class. Final GO class assignment is determined by majority rule wherever at least three out of the five classifiers agree. The method achieves a specificity of >90% at sensitivities >30%. Results are
Use case scenario

In this section, we describe a typical use case scenario for our methods which uses a range of our new servers. Typically we envisage that researchers with a single or a small number of uncharacterized or partially characterized proteins will be interested in making one or more predictions of structural features using our methods. We would advise users wishing to perform high-throughput studies to download the software which we make available via our web site (http://bioinf.cs.ucl.ac.uk/software_downloads/).

Structural annotation

We selected a previously uncharacterized, putative membrane protein from UniProt, A8MYE5 (currently available via http://www.uniprot.org/uniprot/A8MYE5.fasta). This is a 552 residue protein derived from human gene MERTK. The annotation via the protein domain databases available at UniProt suggests this may be a putative protein tyrosine kinase. Initially we submitted the sequence to our FFPred server. FFPred confirms this tentative functional annotation, identifying a range of GO terms the most significant of which are: GO:0008152, GO:0004672, GO:0006468 for 'metabolic process', 'protein kinase activity' and 'protein amino acid phosphorylation', respectively. Structural annotation began by submitting the sequence for analysis with the MEMSAT-SVM service (Figure 3b). The MEMSAT-SVM prediction indicates the possible presence of two membrane spanning helices towards the C-terminus of the sequence, between positions 115 and 130, and 146 and 165. Presence of these helices could indicate that the putative description assigned by Uniprot is correct. However were this a receptor tyrosine kinase, we might expect there to be only a single helix such that a ligand binding domain might be presented to the extracellular space.
The next analysis performed was to submit the sequence to the DomPred server (Figure 3c). The output shows two strong peaks around residue 150 and 234, with the domain boundary prediction made at residue 234 as the first peak likely corresponds to the transmembrane region we had already predicted. As such, this annotation appears to be in reasonable agreement with the initial MEMSAT-SVM annotation.

Based on the MEMSAT-SVM and DomPred annotation, the sequence was then divided into three segments. The initial break point was around residue 234, as suggested by the DomPred analysis, gave two sub-sequences; an N-terminal sequence containing both the initial domain alongside the transmembrane region and a larger C-terminal region containing the second domain. The smaller N-terminal sequence was then divided again at residue 130 to cleave the putative initial domain from the predicted transmembrane region. Both the putative N-terminal domain and the larger C-terminal domain were then submitted to the BioSerf homology modelling server, (Figure 3d). This generated two good, compact homology models for each domain.

The smaller N-terminal domain appears to be a simple two-layer α–β sandwich but the larger domain appears to be more complex. The smaller domain has a large exposed beta sheet, this type of conformation is not common in extracellular domains and this lends some support to the MEMSAT-SVM annotation that predicts two transmembrane helices, which in turn places the N-terminal domain in the cytosolic region. The homology model of the larger C-terminal domain shows an initial region comprised of a well-resolved beta sheet with an attendant layer of alpha helices forming a two-layer α–β sandwich. The other portion of the model appears to form a further discrete region that contains a clear four helix bundle. These two discrete compact regions may indicate that the larger C-terminal domain is in fact comprised of two smaller domains. The homology model for the C-terminal domain has a similar topology to PDB entry 1apm. Searching the CATH database for 1apm reveals that this structure has been divided into two domains by the CATH classification. Finally, following the model all the way to its C-terminus, the tail end of the sequence loops back along the entire length of the domain as a pair of extended β strands. These primarily make contacts only with loop regions. This rather unnatural formation seems likely to be an artefact of the modelling process.

To further examine the terminal β strands of the larger domain, the whole C-terminal sequence was submitted to the DISOPRED2 server (Figure 3e). The DISOPRED2 prediction suggests that the final 35 residues of the C-terminal domain are likely to be disordered. This lends some credence to the proposition that the final two β strands of the protein are an artefact of the modelling process. This may instead be a highly mobile region which may contain features such as binding sites, signal sites or phosphorylation sites.

Our complete annotation suggests a model for this protein containing, in sequence, an initial domain, a transmembrane region, one large or two putative domains and a trailing disordered region (Figure 4). Our prediction of a tyrosine kinase potentially made up of three domains, comprising two compact α–β sandwiches and a compact all alpha domain, is in keeping with other tyrosine kinase structures already deposited at the PDB.

**DISCUSSION**

The UCL Bioinformatics Group server offers some of the best performing algorithms of their kind. They can be accessed via the web (http://bioinf.cs.ucl.ac.uk/web_servers/) or most of the programs themselves can be downloaded for use on a local machine (http://bioinf.cs.ucl.ac.uk/software_downloads/). These services will be useful to any bioinformatician or biologist looking to functionally and structurally characterize proteins.

**FUNDING**

Funding for open access charge: Biotechnology and Biological Sciences Research Council, UK.

Conflict of interest statement. None declared.

**REFERENCES**