## Structural bioinformatics

# ProteinFishing: a protein complex generator within the ModelX toolsuite 

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Received on March 18, 2020; revised on May 11, 2020; editorial decision on May 14, 2020; accepted on May 18, 2020


#### Abstract

Summary: Accurate 3D modelling of protein-protein interactions (PPI) is essential to compensate for the absence of experimentally determined complex structures. Here, we present a new set of commands within the ModelX toolsuite capable of generating atomic-level protein complexes suitable for interface design. Among these commands, the new tool ProteinFishing proposes known and/or putative alternative 3D PPI for a given protein complex. The algorithm exploits backbone compatibility of protein fragments to generate mutually exclusive protein interfaces that are quickly evaluated with a knowledge-based statistical force field. Using interleukin-10-R2 co-crystalized with interferon-lambda-3, and a database of X-ray structures containing interleukin-10, this algorithm was able to generate interleukin-10-R2/interleukin-10 structural models in agreement with experimental data. Availability and implementation: ProteinFishing is a portable command-line tool included in the ModelX toolsuite, written in C++, that makes use of an SOL (tested for MySOL and MariaDB) relational database delivered with a template SOL dump called FishXDB. FishXDB contains the empty tables of ModelX fragments and the data used by the embedded statistical force field. ProteinFishing is compiled for Linux-64bit, MacOS-64bit and Windows-32bit operating systems. This software is a proprietary license and is distributed as an executable with its correspondent database dumps. It can be downloaded publicly at http://modelx.crg.es/. Licenses are freely available for academic users after registration on the website and are available under commercial license for for-profit organizations or companies. Contact: javier.delgado@crg.eu or luis.serrano@crg.eu Supplementary information: Supplementary data are available at Bioinformatics online.


## 1 Introduction

The ModelX toolsuite (Delgado Blanco et al., 2019) has been developed, among other purposes, for modelling biomolecular interactions. ModelX uses fragment libraries generated by in silico digestion of Protein Data Bank (PDB) structures (Berman et al., 2000) and stored in SQL databases. This strategy has proven successful when applied to the design of DNA-protein and RNA-protein interfaces (Blanco et al., 2018; Delgado Blanco et al., 2019). In the protein-protein interactions (PPI) prediction field, few examples of tools performing fast large-scale docking exist. MEGADOCK 4.0 (Ohue et al., 2014) is one, but it requires sophisticated heterogeneous supercomputing environments equipped with hardware accelerators such as GPUs. Another example is InterPred (Mirabello et al., 2017), which uses homology modelling of binding partners
and whole protein superimposition to gather interaction templates. Here, we present ProteinFishing, a tool based on the ModelX philosophy that enables the fast generation of 3D interaction models from observed protein-protein interfaces while fulfilling the requirements for local backbone compatibility.

## 2 New ModeIX tools

In addition to ProteinFishing, the latest ModelX release contains two more commands: GeneratePeptides, which is needed to populate the FishXDB database, and FishingLure, an automatized version of ProteinFishing. The three mentioned commands can be used with any type of PDB file containing standard amino acids and/or nucleotides, including X-ray, nuclear magnetic resonance (NMR), homology models or any other PDB model created by users.


Fig. 1. Algorithm description. (A) The IFN-lambda-R1/IFN-lambda-3/IL-10-R2 complex (PDB: 5T5W) containing the 'Hook' (IFN-lambda-3: red), the 'Fisher' (IL-10-R2: light blue) and IFN-lambda-R1 (grey); (B) The IFN-lambda-R1/IL-10/IL-10-R2 virtual complex superimposed with the 'Hook' window (red); (C) The IL-10/IL-10-R2 or 'Fish/ Fisher' complex (IL-10: dark blue; IL-10-R2: light blue); (D) A comparison between the reported binding levels (first row) and the $\Delta \Delta G$ of interaction as calculated by FoldX (rows 2-12). A unique colour scale has been used to make energies and percentages comparable. The binding loss (\%) numerical scale corresponds to $100 \%$-binding levels' for experimentally measured point mutations (Yoon et al., 2006) and the $\Delta \Delta G(\mathrm{kcal} / \mathrm{mol})$ numerical scale corresponds to FoldX interaction energy

### 2.1 GeneratePeptides command

The GeneratePeptides command allows ModelX users to customize their fragment library. It takes PDB structures as input and digests them into protein fragments of user-defined length in an overlapping sliding-window fashion. These fragments are stored in FishXDB and are therefore available for the ProteinFishing algorithm.

### 2.2 ProteinFishing command

ProteinFishing uses protein complex structures as input, and requires the user to select one molecule to be part of the output complex ('Fisher', Fig. 1A, light blue) and another molecule to be used as the structural template for the retrieval of new docking partners ('Hook', Fig. 1A, red). The algorithm requires the user to define an amino acid window from the 'Hook' molecule to query the Fish $X D B$ protein fragment database with fragment windows interacting with the 'Fisher'. When the geometrical backbone compatibility and sequence similarity-according to user-configurable options-matches the peptide window with a FishXDB fragment, the full PDB model ('Fish', Fig. 1A) from which the fragment was obtained is placed over the 'Hook' fragment by local fitting (Fig. 1B). In this way, complexes containing both the 'Fisher' and the 'Fish' molecules are built (Fig. 1C). Finally, the generated complexes go through two energy filters: the first filter evaluates the presence of atomic clashes between the backbones of the two molecules, and the second filter uses a customizable threshold for free energy values calculated over the generated models. Free energies (representing backbone compatibility) are obtained using a statistical force field embedded in ModelX. The force field is based on a Boltzmann device (Sippl, 1990) with the Kono modification (Kono et al., 1999) of the Sippl method. The models that pass these filters are later returned as PDB files, together with a summary file showing the number of intermolecular contacts, backbone clashes and energy values.

### 2.3 FishingLure command

The FishingLure command represents a fully automated, multi-thread version of ProteinFishing in which the algorithm itself determines all
possible overlapping sliding windows around the 'Hook' residues contacting the 'Fisher'. The FishingLure command allows the use of ProteinFishing over multiple scanning windows computed in parallel.

## 3 Demonstration

To test the utility of our tool, we focused on the complexes of interleukin-10 (IL-10) with its two receptors (IL-10-R1 and IL-10R2). While structures are available for the IL-10/IL-10-R1 complex (PDB: 1J7V, 1Y6K), the structure of the IL-10/IL-10-R2 complex has not been elucidated. We chose the crystallographic structure of IL-10R2 complexed with interferon-lambda-3 and interferon-lambda-receptor-1 (PDB: 5T5W) as input. IL-10-R2 was used as the 'Fisher' molecule and interferon-lambda-3 was used as the 'Hook' (Fig. 1A, red). Defining the scanning window between residues 89-94 of the 'Hook', ProteinFishing yielded 11 models that were then energetically minimized. Next, using the BuildModel command of FoldX (Delgado et al., 2019), five point mutations experimentally reported to modify 'binding levels' between IL-10 and IL-10-R2 (Yoon et al., 2006) were modelled. For each mutation, we computed the FoldX free energy variations ( $\Delta \Delta G[\mathrm{kcal} / \mathrm{mol}]$ of interaction) between the 'Fisher' and the mutated 'Fish'. Finally, we compared the variations between the 'binding levels' of the five IL-10 mutants with IL-10-R2, as reported in literature, with those predicted by FoldX in each of the 11 models (Fig. 1D). The two best-fitting models, as ranked by the statistical force field of ModelX (Supplementary Table S1; 5T5W_1Y6K_8 and 5T5W_1J7V_7), were found to have the best agreement between FoldX energy values and the experimental results (Yoon et al., 2006) (Fig. 1D and Supplementary Table S3). Complete details of the entire process, including the specific parameters used, can be found in Supplementary Appendix: User Tutorial.

## 4 Conclusions

The tools presented here enable the fast structural modelling of PPI suitable for protein design. The ProteinFishing algorithm can be applied in two types of scenarios. The first scenario, described
above, allows the user to model a protein complex for which there is no structure available. Depending on the structures with which the user populates the Fish $X D$ B, the possible interactors 'fished' can be restricted to specific desired targets, or can be exploratory, using all structures from the PDB. In a second scenario, the tool could be used to model different possible conformations between two members of a complex for which a structure already exists. This second scenario could be useful for performing energetic filtering of different conformations, redesigning interfaces by mutagenesis or identifying putative small-molecule binding pockets in the interface between complex members, for example.

## Acknowledgements

The authors would like to thank Professor Jesús Delgado Calvo for the inspiring mathematical discussions that helped us improve the computing efficiency of the algorithm, Tony Ferrar for manuscript revision and language editing, the Centre for Genomic Regulation (CRG) Technology \& Business Development Office (TBDO) for support with licensing information, the CRG Tecnologías de Información y Comunicación (TIC) for assistance with web hosting and the Scientific Information Technologies (SIT) for distributed computing. We appreciate all the feedback from the members of the L.S. lab, especially from Samuel Miravet-Verde.

## Funding

This work was supported by funding from the Spanish Ministry of Science and Innovation (Plan Nacional BFU2015-63571-P). The authors also acknowledge the support of the Spanish Ministry of Science and Innovation to the EMBL
partnership, the Centro de Excelencia Severo Ochoa and the Centres de Recerca de Catalunya (CERCA) Programme/Generalitat de Catalunya. The project that gave rise to these results was supported by a fellowship from 'la Caixa' Foundation (ID 100010434; fellowship code LCF/BQ/DI19/11730061).

Conflict of Interest: none declared.

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