Ensembler: Enabling high-throughput molecular simulations at the superfamily scale

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The rapidly expanding body of available genomic and protein structural data provides a rich resource for understanding protein dynamics with biomolecular simulation. However, simulations on an omics scale are not yet widely performed, partly because software infrastructure to enable this has not kept pace. For example, it should now be possible to study protein dynamics across entire (super)families, exploiting the entire variety of available structural biology data and conformational similarities across homologous proteins. Here, we present a new tool for enabling high-throughput simulation in the genomics era. Ensembler takes any set of sequences—from a single sequence to an entire superfamily of interest—and shepherds them through various stages of modeling and refinement to produce simulation-ready structures. This includes comparative modeling to all relevant PDB structures (which may span multiple conformational states of interest), reconstruction of missing loops, addition of missing atoms, culling of nearly identical structures, assignment of appropriate protonation states, solvation in explicit solvent, and refinement with molecular simulation to ensure stable simulation. The output of this pipeline is an ensemble of structures ready for subsequent parallel or distributed molecular simulations using clusters, supercomputers, or distributed computing projects like Folding@home. Ensembler automates much of the time-consuming process of preparing protein models suitable for simulation, while allowing scalability up to entire superfamilies. A particular advantage of this approach can be found in the construction of kinetic models of conformational dynamics such as Markov state models—which benefit from a diverse array of initial configurations that span the accessible conformational states to aid sampling. We demonstrate the power of this approach by constructing models for all catalytic domains in the human tyrosine kinase family, using all available kinase catalytic domain structures from any organism as structural templates.

Ensembler is free of charge, and is made available under the terms of the GNU General Public License (GPL) v2. It should run on all major operating systems, and has been tested on Linux and OS X. The latest release can be installed via the conda package manager, and the latest source can be downloaded from https://github.com/choderalab/ensembler.

Keywords: molecular dynamics simulation; comparative modeling

I. INTRODUCTION

II. DESIGN AND IMPLEMENTATION

Ensembler is written in Python, and can be used via a command-line tool (ensembler), or via the flexible Python API.

The Ensembler modeling pipeline entails a series of stages which are performed in a defined order. A visual overview of the pipeline is shown in Fig. 1, and a detailed description follows.

1. Target selection

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The output from this stage is simply a FASTA-formatted sequence file containing the selected target sequences and identifiers. The ensembler command-line tool provides methods for selecting targets from either UniProt (a freely accessible resource for protein sequence and functional data—uniprot.org) or TargetExplorer (a database framework for aggregating various types of biological data; work to be published). This allows the user to easily select a single protein, many proteins, or an entire superfamily. Alter-

natively, the targets file can be generated using any other
software, and stored at the appropriate filepath.

The method for selecting targets from UniProt is de-27 scribed here. A query string is required as input, using 28 the same syntax as the search function on the UniProt 29 website. For example, 'domain: "Protein kinase" AND 30 taxonomy:9606 AND reviewed:yes' would select all hu-31 man protein kinases which have been reviewed by a hu-32 man curator. Ensembler is designed to work with pro-33 tein domains, rather than full-length proteins, and the de-34 sired protein domain(s) can be selected using a regular expression. For example, the string 'Protein kinase(?!; 36 truncated) (?!; inactive)' would match domains an-37 notated as "Protein kinase", "Protein kinase; 1" or "Protein 38 kinase; 2", but would exclude the domains "Protein kinase; 39 truncated" and "Protein kinase; inactive". The program 40 then extracts sequences and identifiers from the UniProt data, and saves these to a FASTA-format text file.

2. Template selection

As for target selection, the ensembler tool provides methods for selecting templates from various resources— UniProt, the Protein Data Bank (PDB; pdb.org) or a TargetEx- plorer database. From the user perspective, selection from UniProt proceeds in a similar fashion as described above. The returned data for each UniProt entry includes an up-

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include the desired domain. Data from the SIFTS ser- 99 out poor quality models. vice (www.ebi.ac.uk/pdbe/docs/sifts) (CITE: Velankar Nu- 100 format coordinate files respectively.

Selection from the PDB simply requires specifying a list of 107 PDB IDs. These are matched to UniProt entries via the SIFTS template sequences and structures.

Unresolved template loops can optionally be remodeled with a kinematic closure algorithm, which is provided via the loopmodel tool of the Rosetta software suite (CITE: Rosetta and/or loopmodel). This tends to provide higherquality models following the subsequent modeling process.

3. Modeling

In this stage, models are generated for each targettemplate pair, using the Modeller automodel function (CITE: Modeller), which implements comparative structure modeling by satisfaction of spatial restraints (CITE: Sali Blundell J Mol Biol 1993; Fiser Sali Prot Sci 9 2000). Modeller requires the user to first provide a target-template sequence align-75 ment. This is implemented in Ensembler using the BioPy-76 thon pairwise2 module (CITE: BioPython)—which uses a dynamic programming algorithm—with the PAM 250 scoring matrix of Gonnet et al (CITE: Gaston Gonnet Science 1992).

Non-unique models are then filtered out using structural similarity-based clustering. The mdtraj (CITE: mdtraj) Python library is used to calculate RMSD with a fast quaternion characteristic polynomial (QCP) implementation, and the leader algorithm is then used to populate clusters. A 85 minimum distance cutoff (default: 0.6 Å) is used to retain 86 only a single model per cluster.

Refinement

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Models are then refined with a steepest descent energy minimization and a short molecular dynamics (MD) simulation with implicit solvent. This is implemented using the OpenMM molecular simulation toolkit (link and CITE: OpenMM), chosen for its flexible Python API, and high performance GPU-acclerated simulation code. The 94 Amber99SB-ILDN force field is used (CITE: amber99sbildn 95 refs) with a modified generalized Born solvent model (GBSA-138 96 OBC) (CITE: GBSA-OBC). The simulation is run for a default of

49 to-date list of PDB structures and their residue spans. PDB 97 100 ps. This refinement process helps to prepare models for files are downloaded from the PDB for structures which 98 subsequent production simulation, and also helps to filter

Ensembler also provides optional routines for solvating cleic Acids Res 2013), which provides residue-level map- 101 models with explicit solvent and performing a second MD repings between PDB and UniProt entries, is then used to fil- 102 finement. The solvation step solvates each model for a given ter out PDB chains with < 70% resolved residues within the 103 target with the same number of waters, as this is (currently) domain span. Template sequences and structures are then 104 a requirement for building MSMs from multiple independent extracted and written as FASTA-format sequence and PDB- 105 MD trajectories. The target number of waters is selected by 106 first solvating each model with a specified padding distance (default: 10 Å), then taking a percentile value from the distribution (default: 68th percentile). Models are resolvated service, and the same procedure is then followed to extract with the target number of waters by first solvating with zero padding, then incrementally increasing the box size and rem solvating until the target is exceeded, then finally deleting 112 sufficient waters to match the target value. The explicit solvent MD simulation is also implemented using OpenMM, with the Amber99SB-ILDN force field and TIP3P water.

Packaging

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Finally, Ensembler provides a packaging module, which 117 can be used to compress models in preparation for data 118 transfer, or to prepare models with the appropriate di-119 rectory and file structure for subsequent production sim-120 ulations on the distributed computing platform Folding@Home (CITE: F@H).

Other features

The command-line tool also provides a quickmodel function, which performs the entire Ensembler pipeline for a sin-125 gle target with a small number of templates. For larger numbers of models (such as entire protein families), the main 127 pipeline functions should be used. The modeling and refinement functions use MPI to trivially parallelize computation across each model (or across each template, in the case of the loop reconstruction code).

Each pipeline function also outputs a metadata file, which helps to link data to the software version used to generate it (both Ensembler and its dependencies), and also provides timing and performance information, and other 135 data such as hostname.

III. RESULTS

IV. AVAILABILITY AND FUTURE DIRECTIONS

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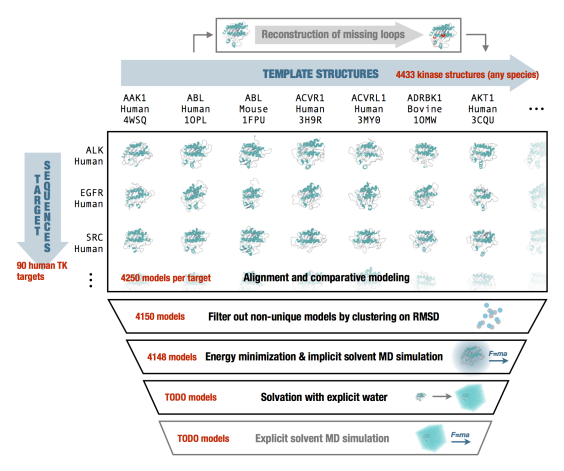


FIG. 1. Ensembler pipeline