Daniel L. Parton,¹ Patrick B. Grinaway,¹ and John D. Chodera^{1,*}

¹Computational Biology Program, Sloan Kettering Institute, Memorial Sloan Kettering Cancer Center, New York, NY 10065 (Dated: March 17, 2015)

The rapidly expanding body of available genomic and protein structural data provides a rich resource for understanding protein dynamics with biomolecular simulation. While computational infrastructure has grown rapidly, simulations on an omics scale are not yet widespread, primarily because software infrastructure to enable simulations at this scale have not kept pace. It should now be possible to study protein dynamics across entire (super)families, exploiting both 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 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 and filtering with molecular simulation to ensure stable simulation. The output of this pipeline is an ensemble of structures ready for subsequent molecular simulations using computer clusters, supercomputers, or distributed computing projects like Folding@home. Ensembler thus automates much of the timeconsuming process of preparing protein models suitable for simulation, while allowing scalability up to entire superfamilies. [JDC: Prior sentence is redundant?] 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 and open source software licensed under 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; distributed simulation

I. INTRODUCTION

Recent advances in genomics and structural biology have 8 helped generate an enormous wealth of protein data at 9 the level of amino-acid sequence and three-dimensional 10 structure. However, proteins typically exist as an ensemble of thermally accessible conformational states, and static structures provide only a snapshot of their rich dynamical behavior. Many functional properties—such as the ability to bind small molecules or interact with signaling partners-require transitions between states, encompassing anything from reorganization of sidechains at binding interfaces to domain motions to large scale folding-unfolding events. Drug discovery could also benefit from a more extensive consideration of protein dynamics, whereby small molecules might be selected based on their predicted ability to bind and trap a protein target in an inactive state [? 22

Molecular dynamics (MD) simulations have the capability, in principle, to describe the time evolution of a protein in atomistic detail, and have proven themselves to be a useful tool in the study of protein dynamics. A number of mature software packages and forcefields are available, and

However, it remains difficult for researchers to exploit the full variety of available protein sequence and structural data in simulation studies, largely due to limitations in software architecture. For example, the set up of a biomolecular simulation is typically performed manually, encompassing a sesies of fairly standard (yet time-consuming) steps such as

₂₈ much recent progress has been driven by advances in com-29 puting architecture. For example, many MD packages are 30 now able to exploit GPUs, which provide greatly improved 31 simulation efficiency per unit cost relative to CPUs, while 32 distributed computing platforms such as Folding@home [CITE], GPUGrid [CITE], and Copernicus [CITE] allow scala-34 bility on an unprecedented level. In parallel, methods for 35 building human-understandable models of protein dynam-36 ics from noisy simulation data, such as Markov state mod-37 eling (MSM) approaches, are now reaching maturity [CITE 38 MSM reviews]. MSM methods in particular have the advan-39 tage of being able to aggregate data from multiple indepen-40 dent MD trajectories, facilitating parallelization of produc-41 tion simulations and thus greatly alleviating overall compu-42 tational cost. There also exist a number of mature software ₄₃ packages for comparative modeling of protein structures, in 44 which a target protein sequence is modeled using one or 45 more structures as templates [CITE Modeller and Rosetta and a recent homology modeling review].

53 the choice of protein sequence construct and starting struc-

^{*} Corresponding author; john.chodera@choderalab.org

54 ture, addition of missing residues and atoms, solvation with 109 explicit water and salt buffer, choice of simulation parameters, and system relaxation with energy minimization and one or more short MD simulations. For this reason, simulation studies typically consider only one or a few proteins and starting configurations.

The ability to fully exploit the large base of available protein sequence and structural data in biomolecular simulation studies could open up many interesting avenues for research, enabling the study of entire protein families or superfamilies across multiple organisms. The similarity between members of a given protein family could be exploited to generate arrays of conformational models, which could be used as starting configurations to aid sampling in MD simulations. This approach would be highly beneficial for many MD methods, such as MSM construction, which require global coverage of the conformational landscape to realize their full potential, and would also be particularly useful in cases where structural data is present for only subset of the members of a protein family. It would also aid in studying protein families known to have multiple metastable conformations—such as kinases—for which the combined body of structural data for the family may cover a large range of these conformations, while the available structures for any individual member might encom-₇₉ pass only one or two distinct conformations.

Here, we present the first steps toward bridging the 134 gap between biomolecular simulation software and omicsscale sequence and structural data: a fully automated open 136 This results in a total of almost 400,000 models, and we 148 'domain: "Protein kinase" AND taxonomy: 9606 AND the tool will prove to be useful in a number of other ways. 154 identifier for each target protein. For example, the generated models could represent valu- 155 108 rameters.

II. DESIGN AND IMPLEMENTATION

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

The **Ensembler** modeling pipeline comprises a series of 113 stages which are performed in a defined order. A visual overview of the pipeline is shown in Fig. 1. The various stages of this pipeline are described in detail below.

[JDC: We could really help the reader if we preface each 117 section here with a bit of an introduction of what we're trying to accomplish in each stage. Otherwise, I worry that 119 each section is a long list of things we do without reference to an overall concept of what the stage is trying to ac-121 complish or why certain decisions were made.] [DLP: Good point. I've added in brief introductions for each section.] 123 [JDC: Can you do a bit more here? I feel that the reader may need more orientation. You've essentially just added a sentence or two at the beginning of each stage that doesn't re-126 ally enlighten the user as to your terminology (tempalte and target), your motivation for why things are done each stage, or how the user is to select among the various options avail-129 able.]

Target selection

The first stage entails the selection of a set of target protein sequences, i.e. the sequences the user is interested in modeling. [JDC: Maybe explain what is meant by "target protein sequences"? These are the sequences the user is interested in modeling.] [DLP: Addressed.]

These targets can be defined manually, simply by prosource framework for building simulation-ready protein 137 viding a FASTA-formatted text file containing the desired models in multiple conformational substates scalable from 138 target sequences with arbitrary identifiers. The ensembler single sequences to entire superfamilies. Ensembler pro- 139 command-line tool also allows targets to be selected vides functions for selecting target sequences and homolo- 140 from UniProt—a freely accessible resource for protein gous template structures, and (by interfacing with a num- 141 sequence and functional data (uniprot.org) [JDC: Isn't ber of external packages) performs pairwise alignments, 142 there a real citation for UniProt?], using the subcommand comparative modeling of target-template pairs, and several [143] gather_targets. The user specifies a query string with stages of model refinement. As an example application, we 144 the --query flag, which conforms to the same syntax have constructed models for the entire set of human tyro- 145 as the search function available on the UniProt website. sine kinase catalytic domains, using all available structures 146 For example, --query 'mnemonic: SRC_HUMAN' would of protein kinase domains (from any species) as templates. ¹⁴⁷ select the full-length human Src sequence, while --query demonstrate that these provide wide-ranging coverage of 149 reviewed: yes' would select all human protein kinases known functionally relevant conformations. By using these 150 which have been reviewed by a human curator. In this models as starting configurations for highly parallel MD sim- us way, the user may select a single protein, many proteins, ulations, we expect their structural diversity to greatly aid 152 or an entire superfamily. The program outputs a FASTA in sampling of conformational space. We anticipate that 153 file, setting the UniProt mnemonic (e.g. SRC_HUMAN) as the

In many cases, it will be desirable to build models of able data sets even without subsequent production simu- 156 an isolated protein domain, rather than the full-length lation, allowing exploration of the conformational diversity 157 protein. The gather_targets subcommand allows propresent within the available structural data for a given pro- 158 tein domains to be selected from UniProt data by passtein family. Furthermore, the automation of simulation set 159 ing a regular expression string to the --domains flag. up provides an excellent opportunity to make concrete cer- 160 For example, the above --query flag for selecting all tain "best practices", such as the choice of simulation pa- 161 human protein kinases returns UniProt entries with domain annotations including "Protein kinase", "Protein ki-

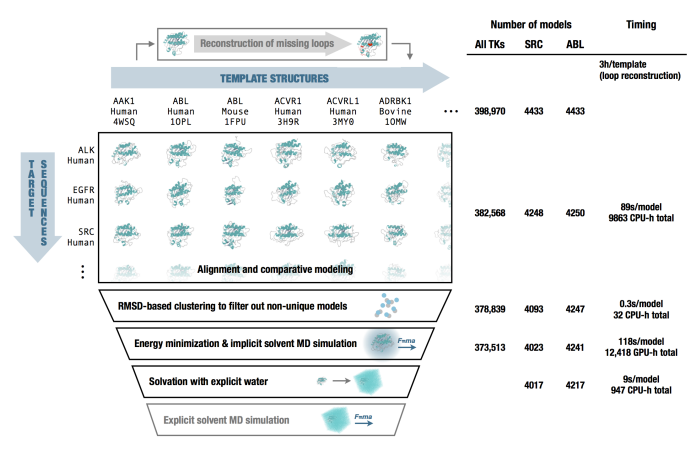


FIG. 1. Diagrammatic representation of the various stages of the Ensembler pipeline. The number of viable models surviving each stage of the pipeline are shown, either for all tyrosine kinases (All TKs) or representative individual kinases (SRC and ABL). In addition, the typical timing on a cluster (containing Intel Xeon E5-2665 2.4GHz hyperthreaded processors and NVIDIA GTX-680 or GTX-Titan GPUs) is reported to exemplify the resources required per model and for modeling the entire set of tyrosine kinases. Note that CPU-h denotes the number of hours consumed by the equivalent of a single hyperthread—parallel execution can reduce wall clock time nearly linearly.

163 nase 1", "Protein kinase 2", "Protein kinase; truncated", 184 tein family as the targets, guaranteeing some degree of hotruncated) (?!; inactive) '. In this case, target identi- 188 lect these?] [DLP: Addressed] fiers are set with the form [UniProt mnemonic]_D[domain 189 JAK1_HUMAN_D0, JAK1_HUMAN_D1. [JDC: Does it make sense 193 also match those in the sequence file. o set some of these coded examples off on their own lines?]

Template selection

174

plates. The second stage thus entails the selection of tem- 201 tures. plates and storage of associated sequences, structures and 202 using the ensembler gather_templates subcommand to 204 e.g.

"Protein kinase; inactive", "SH2", "SH3", etc. To select 185 mology between targets and templates. [JDC: Again, can only domains of the first three types, the following reg- 186 you provide more information about why this is being done? ular expression could be used: '^Protein kinase(?!; 187 What the motivation is, and how the user might expect to se-

Manual selection of templates simply requires storing the index], where the latter part represents a 0-based index for 190 sequences and identifiers in a FASTA file, and the structures the domain—necessary because a single target protein may 191 as PDB-format coordinate files with filenames matching the contain multiple domains of interest. Example identifiers: 192 identifiers in the sequence file. The structure residues must

The ensembler gather_templates subcommand provides methods for selecting template structures from either UniProt or the PDB (), specified by the --gather_from flag. 197 Both methods select templates at the level of PDB chains—a 198 PDB structure containing multiple chains with identical se-Ensembler uses comparative modeling to build models, 199 quence spans (e.g. for crystal unit cells with multiple asymand as such requires a set of structures to be used as tem- 200 metric units) would thus give rise to multiple template struc-

Selection of templates from the PDB simply requires identifiers. These templates can be specified manually, or 203 passing a list of PDB IDs as a comma-separated string, --query 2H8H,1Y57. Specific PDB chain IDs automatically select templates from the Protein Data Bank 205 can optionally also be selected via the --chainids (PDB) or from UniProt. A recommended approach is to se- 206 flag. The program retrieves structures from the PDB 183 lect templates from UniProt which belong to the same pro- 207 server, as well as associated data from the SIFTS service

Acids Res 2013), which provides residue-level mappings be- 264 throughput model building. Models are output as PDBtween PDB and UniProt entries. The SIFTS data is used to ex- 265 format coordinate files. A list of all model identifiers sorted tract template sequences, retaining only residues which are 266 by sequence identity is also written to a text file. To minisequence—non-wildtype residues are thus removed from 268 gzip library to apply compression to all sizeable text files the template structures. Furthermore, PDB chains with less 269 from the modeling stage onwards. than a given percentage of resolved residues (default: 70%) 270 are filtered out. Sequences are stored in a FASTA file, with 2n plate sequence are utilized in the modeling phase, which dentifiers of the form [UniProt mnemonic]_D[UniProt domain index]_[PDB ID]_[PDB chain ID], SRC_HUMAN_DO_2H8H_A. Matching residues then extracted from the original coordinate files and stored as 275 tural similarity-based clustering. The mdtraj [CITE: mdtraj] PDB-format coordinate files.

fashion as for target selection; the --query flag is used to select full-length proteins from UniProt, while the optional 279 gorithm is then used to populate clusters. A minimum dis---domains flag allows selection of individual domains with $_{\scriptscriptstyle 280}$ tance cutoff (which defaults to 0.6 Å) is used to retain only a regular expression string. The returned UniProt data for 281 single model per cluster. each protein includes a list of associated PDB chains and their residue spans, and this information is used to select template structures, using the same method as for template 282 selection from the PDB. If the --domains flag is used, then templates are truncated at the start and end of the domain

Unresolved template residues can optionally be remodeled with the loopmodel subcommand, which employs kinematic closure algorithm [CITE] provided via the loopmodel tool of the Rosetta software suite (CITE: Rosetta and/or loopmodel). Because fewer loops need to be built during the subsequent model-building stage, prebuilding template loops tends to provide higher-quality models after completion of the **Ensembler** pipeline. Loop remodeling may fail for a small proportion of templates due to spatial constraints imposed by the original structure; the subsequent modeling step thus automatically uses the remodeled version of a template if available, but otherwise falls back to using the non-remodeled version. Furthermore, the Rosetta loopmodel program will not model missing residues at the termini of a structure—such residues spans are modeled in the subsequent stage.

Modeling

249

250

253

This stage entails the generation of models via comparative modeling of each target sequence onto each template 306 structure. Non-unique models are subsequently filtered out 307 using a RMSD-based clustering scheme.

ion [CITE: Modeller], which implements comparative structure modeling by satisfaction of spatial restraints CITE: Sali an ment/equilibration under isothermal-isobaric (NPT) condi-Blundell J Mol Biol 1993; Fiser Sali Prot Sci 9 2000]. While 312 tions. The solvation step solvates each model for a given Modeller can generate alignments automatically, we uti- $_{\scriptscriptstyle 313}$ target with the same number of waters to facilitate the intelize the BioPython pairwise2 module [CITE: BioPython]— 314 gration of data from multiple simulations, such as the conwhich uses a dynamic programming algorithm—with the 315 struction of MSMs. The target number of waters is selected PAM 250 scoring matrix of Gonnet et al. [CITE: Gaston 316 by first solvating each model with a specified padding dis-

(www.ebi.ac.uk/pdbe/docs/sifts) (CITE: Velankar Nucleic 263 to produce better quality alignments for purposes of highresolved and match the equivalent residue in the UniProt $_{267}$ mize file storage requirements, **Ensembler** uses the Python

All chains of template structures that contain the tem-272 can sometimes cause models to be nearly identical. Since the goal is to provide good coverage of conformation space, 274 **Ensembler** filters out nearly identical models using struc-Python library is used to calculate RMSD (for $C\alpha$ atoms only) Selection of templates from UniProt proceeds in a similar 277 with a fast quaternion characteristic polynomial (QCP) [Cite 278 Theobald QCP papers] implementation, and the leader al-

Refinement

This stage entails the use of molecular dynamics simulations to refine the models built in the previous step. This 285 helps to improve model quality and also prepares models for subsequent production simulation, including solvation with explicit water molecules, if desired.

Models are first subjected to energy minimization (using 289 the L-BFGS algorithm [CITE]), followed by a short molecular 290 dynamics (MD) simulation with an implicit solvent represen-291 tation. This is implemented using the OpenMM molecular 292 simulation toolkit (link and CITE: OpenMM), chosen for its flexible Python API, and high performance GPU-acclerated 294 simulation code. By default, the Amber99SB-ILDN force field is used [CITE: amber99sbildn refs] with a modified generalized Born solvent model (GBSA-OBC) (CITE: GBSA-OBC). The **Ensembler** API allows the use of any of the other force fields implemented in OpenMM. The simulation is run for a default of 100 ps to filter out poor quality models (where atomic overlaps that cannot be resolved by energy minimization would cause the simulation to explode) and help relax models for subsequent production simulation. [JDC: What criteria were applied to filter out poor models? Do we only look for thrown exceptions or NaNs? Or do we use an energy filtering criteria too? [DLP: We currently just filter out models which throw exceptions or NaNs.]

While protein-only models may be sufficient for struc-308 tural analysis or implicit solvent simulations, Ensembler Modeling is performed with the Modeller automodel func- 309 also provides a stage for solvating models with explicit wa-310 ter and performing a round of explicit-solvent MD refine-262 Gonnet Science 1992], which we have empirically found 317 tance (default: 10 Å), then taking a percentile value from the

318 distribution (default: 68th percentile). [JDC: Would be use- 364 ful to explain why we are doing this.] [DLP: Addressed.] This helps to prevent models with particularly long, extended loops—such as those arising from template structures with unresolved termini-from imposing very large box sizes on the entire set of models. Models are resolvated with the target number of waters by first solvating with zero padding, then incrementally increasing the box size and resolvating 326 until the target is exceeded, then finally deleting sufficient waters to match the target value. The explicit solvent MD simulation is also implemented using OpenMM, using the Amber 99SB-ILDN force field and TIP3P water [JDC: CITE] by default. Other force fields or water models such as TIP4P-Ew [CITE]) can be specified via the Ensembler API. [JDC: We should allow other water models in OpenMM too, such as TIP4P-Ew?] [DLP: I forgot to mention this in the text previously - any of the OpenMM force fields can be chosen via the API. I've updated the text accordingly. Is this functionality sufficient? I guess it's ok to leave ff choice as an "advanced" feature which requires use of the API? Otherwise I could add ³³⁸ a --water_model flag to the CLI, for example.]

Packaging

Ensembler provides a packaging module which can be 340 used to compress models in preparation for data transfer, or to prepare models with the appropriate directory and file structure for subsequent production simulations on the distributed computing platform Folding@home (CITE: F@H).

Provenance

345

To aid the user in tracking the provenance of each model, 396 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 data such as hostname.

Rapidly modeling a single template

363 number of models generated.

RESULTS

IJDC: It would be useful to have some subheadings in this 366 section to give it some internal organization.]

Modeling of all human tyrosine kinase catalytic domains

As a first application of **Ensembler**, we have built mod-369 els for all 90 human tyrosine kinase (TK) domains listed in UniProt. [JDC: Is there a complete list of these somewhere? 371 Maybe reference supplementary data?] TKs (and protein 372 kinases in general) play important roles in many cellular processes and are involved in a number of types of cancer. [JDC: CITE] For example, mutations of Src are associated with colon, breast and prostate cancer [CITE: Src cancer involvement], while a translocation between the TK Abl and the pseudokinase Bcr is closely associated with chronic 378 myelogenous leukemia [CITE: Abl cancer involvement]. Protein kinase domains are thought to have multiple accessible metastable conformation states, with a single active conformation, and much effort is directed at developing kinase inhibitor drugs which bind to and stabilize inactive conformations [CITE: Lee and Craik Science 2009]. [JDC: Lee and Craik do not discuss kinases, I don't believe; you'll have to find an accurate reference on kinase conformations.] Kinases are thus a particularly interesting subject for study with MSM methods [CITE: recent kinase MSM papers], and this approach stands to benefit greatly from the ability to exploit the full body of available genomic and structural data within the kinase family, e.g. by generating large numbers of starting configurations to be used in highly parallel MD simulation.

We selected all available structures of protein kinase domains (of any species) as templates, for a total of 4433 (398,970 target-template pairs). The templates were derived from 3028 individual PDB entries and encompassed 397 23 different species, with 3634 template structures from hu-398 man kinase constructs.

Modeling statistics

Unresolved template residues were first remodeled us-401 ing the loopmodel subcommand. The number of miss-402 ing residues in each template ranged from 0 to 102, with a median of 11 and a standard deviation of 13. Out of 3666 For users interested in simply using **Ensembler** to rapidly 404 templates with one or more missing residues, 3134 were generate a set of models for a single template sequence, **En-** 405 successfully remodeled, with most remodeling failures atsembler provides a command-line tool quickmodel, which 406 tributable to spatial constraints imposed by the original performs the entire pipeline for a single target with a small $_{\scriptscriptstyle 407}$ template structure. There was some correlation between renumber of templates. For larger numbers of models (such as $_{ ext{\tiny 408}}$ modeling failures and the number of missing residues; tementire protein families), modeling time is greatly reduced by 409 plates for which remodeling failed had a median of 20 missusing the main modeling pipeline, which is parallelized via 410 ing residues, compared to a median of 14 missing residues MPI, distributing computation across each model (or across 411 for templates for which remodeling was successful. The diseach template, in the case of the loop reconstruction code), 412 tributions are plotted in Fig. S1. [JDC: Can you give some and scaling (in a "pleasantly parallel" manner) up to the 413 statistics on the distribution of loop lengths modeled? Why 414 did loop modeling fail in the cases it did? Anything else you

415 can say here beyond this one sentence? [DLP: Addressed in 470 based on the sequence identity between the target and temthe text, and a SI figure.]

the most compute-intensive.

and including the implicit solvent MD refinement stage), to- 488 landscape. The high sequence identity models could be 434 talling 1.7 GB per TK target or 149 GB for all 90 TKs. The data 489 considered to be the most likely to accurately represent true from the modeling stage—with the largest contribution arising from the Modeller restraint files—and 77 kB for the im- 492 plicit solvent MD refinement stage.

Evaluation of model quality

439

The distribution of RMSDs of the final models (relative 441 to the highest sequence identity model for a given target) is shown in Fig. 3. The distributions are stratified based on the sequence identity between target and template, indicating that higher sequence identity templates result in models with lower RMSDs. The sequence identity stratifications were selected based on the sequence identity distribution plotted in Fig. 2, which suggests an intuitive division into three categories, with 307,753 models in the 0-35% sequence identity range, 69,922 models in the 35-55% range, and 4893 models in the 55-100% range.

To provide a more complete evaluation of the models generated, we have analyzed two example TKs (Src and Abl) in detail. Due to their importance in cancer, as outlined above, these kinases have been the subject of numerous studies, encompassing many different methodologies. In terms of structural data, a large number of crystal structures have been solved (with or without ligands such as nucleotide substrate or inhibitor drugs), showing the kinases n a number of different conformations. These two kinases are thus also interesting targets for MSM studies, with one recent study focusing on modeling the states which constiute the activation pathway of Src [CITE:Shukla Pande Nat Commun 2014].

Fig. 4 shows a superposition of a set of representative 519 models of Src and Abl. Models were first stratified into three 520 conda package manager for Python [?]. ranges, based on the structure of the sequence identity dis- 521 # conda install -c https://conda.binstar.org/omnia ensembler tribution (Fig. 2), then subjected to k-medoids clustering to $_{522}$ Up to date instructions can be found at https://github. 469 tity range. Each model is colored and given a transparency 524 dencies except for Modeller and Rosetta, which are not

471 plate sequence. The figure gives an idea of the variance Following loop remodeling, the Ensembler pipeline was 472 present in the generated models. High sequence identity performed up to and including the implicit solvent MD re- 473 models (in opaque blue) tend to be quite structurally similar, finement stage, which completed with 373,513 surviving 474 with some variation in loops or changes in domain orientamodels. To obtain statistics for the solvation stage with- 475 tion. The Abl1 renderings indicate one high sequence idenout generating a sizeable amount of coordinate data, the 476 tity model with a long unstructured region at one of the tersolvate subcommand was performed for just two repre- 477 mini, which was unresolved in the original template strucsentative individual kinases (Src and Abl). The number of 478 ture. While such models are not necessarily incorrect or unnodels which survived each stage are shown in Fig. 1, indi- $_{479}$ dersirable, it is important to be aware of the effects they cating that the greatest attrition occurred during the mod- 400 may have on production simulations performed under perieling stage. The number of refined models for each target 481 odic boundary conditions, as long unstructured termini can ranged from 4005 to 4248, with a median of 4160 and stan- 👑 be prone to interact with a protein's periodic image. Lower dard deviation of 60. Fig. 1 also indicates the typical tim- 483 sequence identity models (in transparent white or red) ining achieved on a cluster for each stage, showing that the 484 dicate much greater variation in all parts of the structure. build_models and refine_implicit_md stages are by far 485 We believe the mix of high and low sequence identity models to be particularly useful for methods such as MSM build-Each model generated about 513 KB of file data (up to 487 ing, which require thorough sampling of the conformational generated per model breaks down as 436 kB for the output 490 metastable states. Conversely, the lower sequence identity models could be expected to help push a simulation into regions of conformation space which might take intractably 493 long to reach if starting a single metastable conformation.

> To evaluate the models of Src and Abl1 in the context of the published literature, we have focused on two residue pair distances thought to be important for the regulation of protein kinase domains. Fig. 5 shows two structures of Src believed to represent inactive (PDB code: 2SRC) [CITE: 2SRC] and active (PDB code: 1Y57) [CITE: 1Y57] states. One notable feature which distinguishes the two structures is the transfer of an electrostatic interaction of E310 from R409 (in the inactive state) to K295 (in the active state), brought about by a rotation of the α C-helix. These three residues are also well conserved [CITE Kannan Neuwald JMB 2005] and a num-505 ber of experimental and simulation studies have suggested that this electrostatic switching process plays a role in a reg-507 ulatory mechanism shared across the protein kinase fam-508 ily [CITE Foda Shan Seeliger Src Nat Commun 2015; Shukla Pande Nat Commun 2014; Ozkirimli Post Prot Sci 2008]. As such, we have projected the **Ensembler** models for *Src* and ⁵¹¹ Abl1 onto a space consisting of the distances between these two residue pairs (Fig. 6). The models show strong coverage of regions in which either of the electrostatic interactions is ₅₁₄ formed, as well as a wide range of regions inbetween. We thus expect that such a set of models, if used as starting con-516 figurations for a highly parallel MD simulation, could greatly ⁵¹⁷ aid in sampling of the activation process.

AVAILABILITY AND FUTURE DIRECTIONS

The latest release of **Ensembler** can be installed via the

pick three representative models from each sequence iden- 523 com/choderalab/ensembler. This will install all depen-

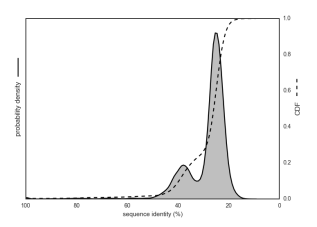


FIG. 2. Sequence identity distribution for human TK models. Distribution of sequence identities for all 373,513 models generated for the human tyrosine kinases. Sequence identities are calculated from pairwise target-template alignments. The cumulative distribution function is shown by the dashed line. The plotted distributions have been smoothed using kernel density estimation. [JDC: Font size too small.]

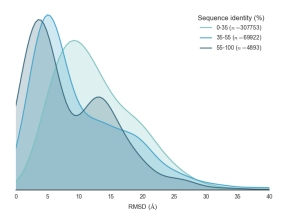


FIG. 3. RMSD distribution by sequence identity. RMSD distributions for all 373,513 human TK models, divided into three sequence identity ranges. For a given target, model RMSDs are calculated relative to the highest sequence identity model for that target. The plotted distributions have been smoothed using kernel density estimation. [JDC: Font size too small.]

525 available through the conda package manager, and thus ₅₂₆ must be installed separately by the user. The latest source 527 can be downloaded from the above GitHub repository, which also contains instructions for building and installing 529 the code.

530

[JDC: In the Discussion, let's be sure to talk about the limitations and what could be improved or added in the future. For example, we don't yet handle counterions (e.g. 540

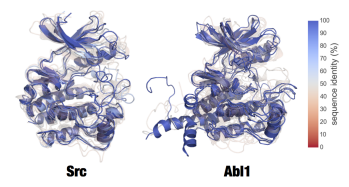


FIG. 4. Superposition of clustered models of Src and Abl. Superposed renderings of nine models each for Src and Abl, [JDC: Src and Abl, or Src and Abl1? The description should match the captions above.] giving some indication the diversity of conformations generated by Ensembler. The models for each target were divided into three sequence identity ranges (as in Fig. 3), and RMSDbased k-medoids clustering was performed to select three clusters from each. The models shown are the centroids of each cluster. Models are colored and given transparency based on their sequence identity, so that high sequence identity models are blue and opaque, while lower sequence identity models are transparent and red. [JDC: Font size too small.]

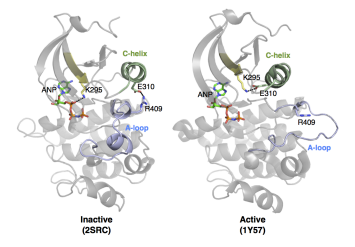


FIG. 5. Two structures of Src, indicating certain residues involved in activation. In the inactive state, E310 forms a salt bridge with R409. During activation, the α C-helix (green) moves and rotates, orienting E310 towards the ATP-binding site and allowing it to instead form a salt bridge with K295. This positions K295 in the appropriate position for catalysis.

scale modeling, but there's a lot of work yet to be done.]

ACKNOWLEDGMENTS

The authors are grateful to Kyle A. Beauchamp (MSKCC), structural Zn²⁺), prosthetic groups (e.g. heme), or cofactors s41 Robert McGibbon (Stanford), Arien S. Rustenburg (MSKCC) (e.g. ATP) yet. We don't handle post-translational modifica- 542 for many excellent software engineering suggestions. The tions either (such as phosphorylation, methylation, glycosy- 543 authors thank Sonya M. Hanson (MSKCC), Nicholas M. Levin-556 lation, etc.). It's a good idea to suggest that this is an impor- 544 son (?), Markus A. Seeliger (Stony Brook), Diwakar Shukla tant first step toward enabling superfamily- and genomics- 545 (Stanford), and Avner Schlessinger (Mount Sinai) for help-

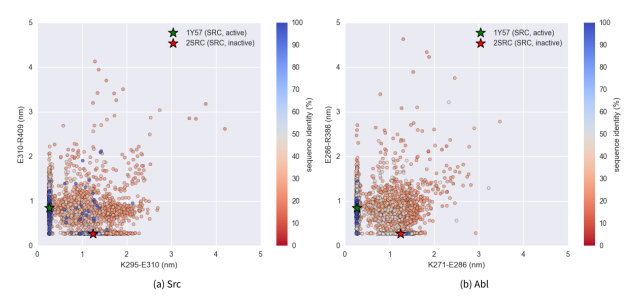


FIG. 6. Src and Abl1 models projected onto the distances between two conserved residue pairs, colored by sequence identity. Two Src structures (PDB entries 1Y57 [CITE] and 2SRC [CITE]) are projected onto the plots for reference, representing active and inactive states respectively. These structures and the residue pairs analyzed here are depicted in Fig. 5. Distances are measured between the center of masses of the three terminal sidechain heavy atoms of each residue. The atom names for these atoms, according to the PDB coordinate files for both reference structures, are—Lys: NZ, CD, CE (ethylamine); Glu: OE1, CD, OE2 (carboxylate); Arg: NH1, CZ, NH2 (part of guanidine). [JDC: Font size too small if single-column, so I made this double-column. Not sure if that's what you want.] [JDC: Fill in rest of caption.]

546 ful scientific feedback on modeling kinases. The authors 550 ilyn Gunner (CCNY) for assistance with MCCE2. DLP and this are grateful to Benjamin Webb and Andrej Sali (UCSF) for ssi work was supported in part by the generous support of a 548 help with the MODELLER package, Peter Eastman and Vi- 552 Louis V. Gerstner Young Investigator Award. jay Pande (Stanford) for assistance with OpenMM, and Mar-