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. 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 has 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. A particular advantage of this approach can be found in the construction of kinetic models of conformational dynamics—such as Markov state models (MSMs)—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 is compatible with 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.

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I. INTRODUCTION

Recent advances in genomics and structural biology have helped generate an enormous wealth of protein data at the level of amino-acid sequence and three-dimensional 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 [1].

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 now available, and much recent progress has been driven by advances in computing architecture. For example, many MD However, it remains difficult for researchers to exploit
the full variety of available protein sequence data (in simulating groups of related proteins) and structural data (exploiting multiple structures for each protein and its homologs/orthologs) in simulation studies in molecular simulations, largely due to limitations in software architecture.
For example, the preparation of a biomolecular simulation is typically performed manually, encompassing a series of fairly standard (yet time-consuming) steps such as
the choice of protein sequence construct and starting structure(s), addition of missing residues and atoms, solvation
with explicit water and counterions (and potentially buffer

²⁹ packages are now able to exploit GPUs [2, 3], which pro-30 vide greatly improved simulation efficiency per unit cost rel-31 ative to CPUs, while distributed computing platforms such as Folding@home [4], Copernicus [5, 6], and GPUGrid [7], al-33 low scalability on an unprecedented level. In parallel, meth-34 ods for building human-understandable models of protein 35 dynamics from noisy simulation data, such as Markov state modeling (MSM) approaches, are now reaching maturity [8– ₃₇ 10]. MSM methods in particular have the advantage of be-38 ing able to aggregate data from multiple independent MD 39 trajectories, facilitating parallelization of production simu-40 lations and thus greatly alleviating overall computational 41 cost. There also exist a number of mature software packages 42 for comparative modeling of protein structures, in which a 43 target protein sequence is modeled using one or more struc-44 tures as templates [11-14].

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57 components and cosolvents), choice of simulation param- 115 and we demonstrate that these provide wide-ranging covparameters do not yet exist), system relaxation with energy minimization, and one or more short preparatory MD simulations to equilibrate the system and relax the simufew proteins and starting configurations. Worse still, studies (or collections of studies) that do consider multiple proteins often suffer from the lack of consistent best practices in this preparation process, making comparisons between related proteins unnecessarily difficult. While notable exceptions exist—such as the Dynameomics effort of Daggett and 127 ods benefit from the ability to remove outlier MD trajecsingle initial configuration for each [15]—it is nevertheless clear that the lack of scalability of existing pipelines is a hinerance to making facile use of all available structural data.

The ability to fully exploit the large quantity of available protein sequence and structural data in biomolecular simllation studies could open up many interesting avenues for esearch, enabling the study of entire protein families or suerfamilies within a single organism or across multiple organisms. The similarity between members of a given protein family could be exploited to generate arrays of conformational models for related sequences, which could be used as starting configurations to aid sampling in MD simulations. The conformations captured in structures of related members has been shown to provide useful information about the conformations accessible to all members of the family [16, 17], though energetic differences between individuils will modify the populations and dynamics of individual conformational states. 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 a 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 150 command-line tool (ensembler) or via a flexible Python the combined body of structural data for the family may cover a large range of these conformations, while the availble structures for any individual member might encompass only one or two distinct conformations.

Here, we present the first steps toward bridging the gap between biomolecular simulation software and *omics*scale sequence and structural data: a fully automated open source framework for building simulation-ready protein models in multiple conformational substates scalable from single sequences to entire superfamilies. Ensembler provides functions for selecting target sequences and homoloous template structures, and (by interfacing with a number of external packages) performs pairwise alignments, comparative modeling of target-template pairs, and several 161

eters (or parameterization schemes for components where 116 erage of known functionally relevant conformations. By using these models as starting configurations for highly parallel MD simulations, we expect their structural diversity to ₁₁₉ greatly aid in sampling of conformational space. We further lation cell. Due to the laborious and manual nature of this 120 suggest that models with high target-template sequence process, simulation studies typically consider only one or a 121 identity are the most likely to represent native metastable 122 states, while lower sequence identity models would aid in sampling of more distant regions of accessible phase 124 space. It is also important to note that some models (especially low sequence identity models) may not represent 126 natively accessible conformations. However, MSM methcoworkers, which has simulated 100 proteins so far using a 128 tories which start from non-natively accessible conformations, and which would thus be unconnected with the phase space sampled in other trajectories. These methods essentially identify the largest subset of Markov nodes which con-132 stitute an ergodic network [18, 19].

We anticipate that **Ensembler** will prove to be useful in 134 a number of other ways. For example, the generated models could represent valuable data sets even without subsequent production simulation, allowing exploration of the conformational diversity present within the available structural data for a given protein family. Furthermore, automation of simulation preparation provides an excellent opportunity to make concrete certain "best practices", such as the 141 choice of simulation parameters, approach to the treatment of protonation states, treatment of cofactors and structural ions, and pre-simulation refinement and equilibration pro-144 cedures. While the current version of **Ensembler** only codifies some of these choices as default parameters, its modular nature allows additional stages to be easily added in the 147 future.

II. DESIGN AND IMPLEMENTATION

Ensembler is written in Python, and can be used via a 151 API to allow integration of its components into other ₁₅₂ applications. All command-line and API information in this article refers to the version 1.0.2 release of Ensembler. Up-to-date documentation can be found at ensembler.readthedocs.org.

The **Ensembler** modeling pipeline comprises a series of 157 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.

Target selection and retrieval

The first stage entails the selection of a set of target prostages of model refinement. As an example application, we 162 tein sequences—the sequences for which the user is inhave constructed models for the entire set of human tyro- 163 terested in generating simulation-ready structural models. sine kinase (TK) catalytic domains, using all available struc- 164 This may be a single sequence—such as a full-length protures of protein kinase domains (from any species) as tem- 165 tein or a construct representing a single domain—or a col-114 plates. This results in a total of almost 400,000 models, 166 lection of sequences, such as a particular domain from an

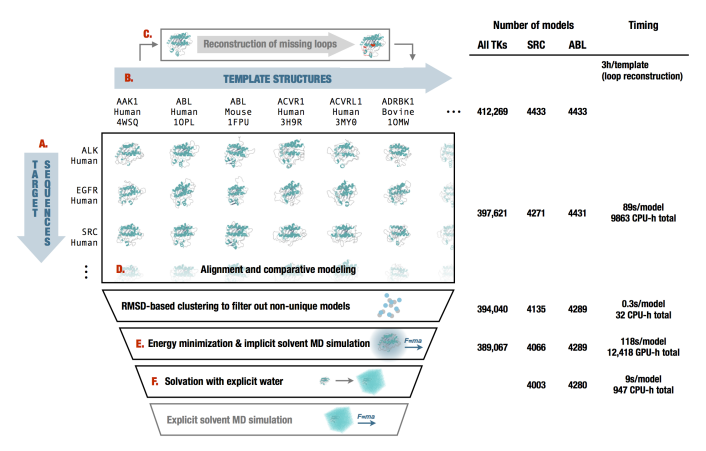


FIG. 1. Diagrammatic representation of the stages of the Ensembler pipeline and illustrative statistics for modeling all human tyrosine kinase catalytic domains. On the left, the various stages of the Ensembler pipeline are shown. The red labels indicate the corresponding text description provided for each stage in the Design and Implementation section. On the right, the number of viable models surviving each stage of the pipeline is shown for the 93 target TK domains and for two representative individual TK domains (SRC and ABL). Typical timings on a computer cluster (containing Intel Xeon E5-2665 2.4GHz hyperthreaded processors and NVIDIA GTX-680 or GTX-Titan GPUs) is reported to illustrate resource requirements per model for modeling the entire set of tyrosine kinases. Note that CPU-h denotes the number of hours consumed by the equivalent of a single CPU hyperthread and GPU-h on a single GPU—parallel execution via MPI reduces wall clock time nearly linearly.

entire family of proteins. The output of this stage is a FASTA- 188 tein. The gather_targets subcommand allows protein formatted text file containing the desired target sequences 189 domains to be selected from UniProt data by passing a reguwith corresponding arbitrary identifiers.

protein sequence and functional data (uniprot.org) [20] from UniProt, the subcommand gather_targets is used with the --query flag followed by a UniProt query string available on the UniProt website. For example, --query reviewed by a human curator. In this way, the user may se- 202 JAK1_HUMAN_D1). lect a single protein, many proteins, or an entire superfamily from UniProt. The program outputs a FASTA file, setting the UniProt mnemonic (e.g. SRC_HUMAN) as the identifier for $_{203}$ each target protein.

187 isolated protein domain, rather than the full-length pro- 206 arbitrary identifiers.

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lar expression string to the --uniprot_domain_regex flag. The ensembler command-line tool allows targets to 191 For example, the above --query flag for selecting all hube selected from UniProt—a freely accessible resource for 192 man protein kinases returns UniProt entries with domain annotations including "Protein kinase", "Protein kinase 1", via a UniProt search query. To retrieve target sequences 194 "Protein kinase 2", "Protein kinase; truncated", "Protein kinase; inactive", "SH2", "SH3", etc. The regular expression shown in Box 1 selects only domains of the first three types. conforming to the same syntax as the search function 197 If the --uniprot_domain_regex flag is used, target identi-198 fiers are set with the form [UniProt mnemonic]_D[domain mnemonic:SRC_HUMAN, would select the full-length hu- 199 index], where the latter part represents a 0-based index for man Src sequence, while the query shown in Box 1 would 200 the domain—necessary because a single target protein may select all human tyrosine protein kinases which have been 201 contain multiple domains of interest (e.g. JAK1_HUMAN_DO,

Target sequences can also be defined manually (or from 204 another program) by providing a FASTA-formatted text file In many cases, it will be desirable to build models of an 205 containing the desired target sequences with corresponding

B. Template selection and retrieval

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Ensembler uses comparative modeling to build models, and as such requires a set of structures to be used as templates. The second stage thus entails the selection of templates and storage of associated sequences, structures, and dentifiers. These templates can be specified manually, or using the ensembler gather_templates subcommand to automatically select templates based on a search of the 270 Protein Data Bank (PDB) or UniProt. A recommended approach is to select templates from UniProt which belong to 271 and templates.

The ensembler gather_templates subcommand pro- 275 vides methods for selecting template structures from either 276 UniProt or the PDB (http://www.rcsb.org/pdb), specified by the --gather_from flag. Both methods select templates at the level of PDB chains—a PDB structure containng multiple chains with identical sequence spans (e.g. for independent conformations of the protein within the asym- 282

Selection of templates from the PDB simply requires 285 residue spans are modeled in the subsequent stage. passing a list of PDB IDs as a comma-separated string, e.g. --query 2H8H,1Y57. Specific PDB chain IDs can optionally also be selected via the --chainids flag. 286 The program retrieves structures from the PDB server, as well as associated data from the SIFTS service 287 retaining only residues which are resolved and match 291 rium conditions. 240 the equivalent residue in the UniProt sequence—non- 292 domain index]_[PDB ID]_[PDB chain ID], PDB-format coordinate files.

Selection of templates from UniProt proceeds in a similar 302 fashion as for target selection; the --query flag is used to 303 own alignments automatically, a standalone function was select full-length proteins from UniProt, while the optional 304 preferable for reasons of programming convenience. As ual domains with a regular expression string (Box 1). The 306 ing the BioPython pairwise2 module [26]—which uses a returned UniProt data for each protein includes a list of as- 307 dynamic programming algorithm—with the PAM 250 scorsociated PDB chains and their residue spans, and this infor- 308 ing matrix of Gonnet et al. [27], though other choices of scortures solved by X-ray crystallography or NMR are selected, and mand, prior to the modeling step which is carried out with thus excluding computer-generated models available from 312 the build_models subcommand. The align subcommand the PDB. If the --uniprot_domain_regex flag is used, then 313 also writes a list of the sequence identities for each template 262 templates are truncated at the start and end of the domain 314 to a text file, and this can be used to select models from 263 sequence.

Templates can also be defined manually. Manual speci-₂₆₅ fication of templates simply requires storing the sequences ²⁶⁶ and arbitrary identifiers in a FASTA file, and the structures as PDB-format coordinate files with filenames matching the identifiers in the sequence file. The structure residues must also match those in the sequence file.

Template refinement

Unresolved template residues can optionally be modeled the same protein family as the targets, guaranteeing homol- 272 into template structures with the loopmodel subcommand, ogy and some degree of sequence identity between targets 273 which employs a kinematic closure algorithm provided via 274 the loopmodel tool of the Rosetta software suite [22, 23]. We expect that in certain cases, pre-building template loops with Rosetta loopmodel prior to the main modeling stage (with MODELLER) may result in improved model quality. Loop remodeling may fail for a small proportion of templates due to spatial constraints imposed by the original 280 structure; the subsequent modeling step thus automaticrystals with non-crystallographic symmetry giving rise to 281 cally uses the remodeled version of a template if available, but otherwise falls back to using the non-remodeled vermetric unit) would thus give rise to multiple template struc- 283 sion. Furthermore, the Rosetta loopmodel program will not 284 model missing residues at the termini of a structure—such

Alignment and comparative modeling

In the modeling stage, structural models of the target se-(www.ebi.ac.uk/pdbe/docs/sifts) [21], which provides 288 quence are generated from the template structures, with residue-level mappings between PDB and UniProt entries. 289 the goal of modeling the target in a variety of conforma-The SIFTS data is used to extract template sequences, 290 tions that could be significantly populated under equilib-

Modeling is performed using the automodel function of wildtype residues are thus removed from the template 293 the MODELLER software package [24, 25] to rapidly gener-242 structures. Furthermore, PDB chains with less than a 294 ate a single model of the target sequence from each temgiven percentage of resolved residues (default: 70%) are 295 plate structure. MODELLER uses simulated annealing cyfiltered out. Sequences are stored in a FASTA file, with iden- 296 cles along with a minimal forcefield and spatial restraints tifiers of the form [UniProt mnemonic]_D[UniProt 297 generally Gaussian interatomic probability densities exe.g. 298 tracted from the template structure with database-derived SRC_HUMAN_DO_2H8H_A. Matching residues then ex- 299 statistics determining the distribution width—to rapidly tracted from the original coordinate files and stored as 300 generate candidate structures of the target sequence from the provided template sequence [24, 25].

While MODELLER's automodel function can generate its -uniprot_domain_regex flag allows selection of individ- 👊 such, we implemented pairwise alignment functionality usmation is used to select template structures, using the same 309 ing matrices available within the module can be selected. method as for template selection from the PDB. Only struc- 310 The alignments are carried out with the align subcoma desired range of sequence identities. The build_models 316 subcommand and all subsequent pipeline functions have a 364 -template_seqid_cutoff flag which can be used to select only models with sequence identities greater than the given value. We also note that alternative approaches could be used for the alignment stage. For example, multiple sequence alignment algorithms [28], allow alignments to be guided using sequence data from across the entire protein family of interest, while (multiple) structural alignment algorithms such as MODELLER's salign routine [24, 25], PRO-MALS3D [29], and Expresso and 3DCoffee [30, 31], can additionally exploit structural data. **Ensembler's** modular architecture facilitates the implementation of alternative alignment approaches, and we plan to implement some of these in future versions, to allow exploration of the influence of different alignment methods on model quality.

Models are output as PDB-format coordinate files. To minimize file storage requirements, **Ensembler** uses the Python gzip library to apply compression to all sizeable text files from the modeling stage onwards. The restraints used by MODELLER could potentially be used in alternative additional refinement schemes, and **Ensembler** thus provides a flag (--write_modeller_restraints_file) for optionally saving these restraints to file. This option is turned off by default, as the restraint files are relatively large (e.g. \sim 400 kB per model for protein kinase domain targets), and are not expected to be used by the majority of users.

[Clarify how nonstandard amino acids are treated at this 385] 343 stage.]

Filtering of nearly identical models

Because **Ensembler** treats individual chains from source 397 PDB structures as individual templates, a number of mod- 398 mization, which result in an unstable simulation), as well as els may be generated with very similar structures if these 399 helping to relax model conformations. As discussed in the ter out models that are very similar in RMSD. The cluster 403 the vast majority failed within the first 1 ps of simulation. subcommand can thus be used to identify models which difspecified cutoff. Clustering is performed using the regular 406 for the refinement simulations carried out here. As such, $_{357}$ late RMSD (for C $_{lpha}$ atoms only) with a fast quaternion char- $_{409}$ along with a modified generalized Born solvent model [41] acteristic polynomial (QCP) [33–35] implementation. A min- 410 as implemented in the OpenMM package [2]. imum distance cutoff (which defaults to 0.6 Å) is used to re- 411 the other force fields or implicit water models imple-360 tain only a single model per cluster.

Filtering by MolProbity score

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363 scores, such as MolProbity or DOPE.]

Refinement of models

A number of refinement methods have been developed to help guide comparative modeling techniques toward more "native-like" and physically consistent conformations [36, 368 37], of which MD simulations are an important example. 369 While long-timescale unrestrained MD simulations (on the order of 100 μ s) have been found to be ineffective for recapit-371 ulating native-like conformations, possibly due to forcefield 372 issues [38], even relatively short simulations can be useful 373 for relaxing structural elements such as sidechain orienta-374 tion [37]. We stress that the limited refinement by molec-375 ular simulation here is primarily intended as initial relaxation and filtering stages, where implausible models might cause simulations to immediately fail, crash, or generate im-378 plausibly high energies or unstable dynamics. Exploration of conformational dynamics to derive MSMs, for example, will inevitably require orders of magnitude more simulation 381 effort—very likely tens of microseconds to milliseconds of 382 aggregate dynamics [8, 10].

Ensembler thus includes a refinement module, which 384 uses short molecular dynamics simulations to refine the models built in the previous step. As well as improving model quality, this also prepares models for subsequent production MD simulation, including solvation with explicit water molecules, if desired.

Models are first subjected to energy minimization (using the L-BFGS algorithm [39], followed by a short molecular 391 dynamics (MD) simulation with an implicit solvent representation. This is implemented using the OpenMM molecular simulation toolkit [2], chosen for its flexible Python API, and high performance GPU-acclerated simulation code. The simulation is run for a default of 100 ps, which in our example applications has been sufficient to filter out poor models (i.e. those with atomic overlaps unresolved by energy miniindividual chains are nearly identical in conformation. For 400 Results section, our example application of the Ensembler this reason, and also to allow users to select for high di- 401 pipeline to the human tyrosine kinase family indicated that ersity if they so choose, **Ensembler** provides a way to fil- 402 of the models which failed implicit solvent MD refinement,

The simulation protocol and default parameter values fer from other models in terms of RMSD distance by a user- 405 have been chosen to represent current "best practices" spatial clustering algorithm [9], as implemented in the MSM- 407 the simulation is performed using Langevin dynamics, Builder Python library [18], which uses mdtraj [32] to calcu- 408 with a default force field choice of Amber99SB-ILDN [40], mented in OpenMM can be specified using the --ff and --water_model flags respectively. The simulation length can also be controlled via the --simlength flag, and many other important simulation parameters can be controlled from either the API or CLI (via the --api_params flag). The default values are set as follows—timestep: 2 fs; temperature: 300 K; Langevin collision rate: 20 ps $^{-1}$; pH (used Insert section about optional filtering by model quality 419 by OpenMM for protonation state assignment): 7. We also 420 draw attention to a recent paper which indicates that lower 421 Langevin collision rates may result in faster phase space ex- 473 structure for production simulation on the distributed ploration [42].

residues/mutations at this stage.]

F. Solvation and NPT equilibration

While protein-only models may be sufficient for struc-432 tural analysis or implicit solvent simulations, Ensembler also provides a stage for solvating models with explicit water and performing a round of explicit-solvent MD refinement/equilibration under isothermal-isobaric (NPT) condions. The solvation step solvates each model for a given target with the same number of waters to facilitate the integration of data from multiple simulations, which is important for methods such as the construction of MSMs. The target number of waters is selected by first solvating each model with a specified padding distance (default: 10 Å), then taking a percentile value from the distribution (default: 68th percentile). This helps to prevent models with particularly long, extended loops—such as those arising from template structures with unresolved termini—from impos-446 ing very large box sizes on the entire set of models. The TIP3P water model [43] is used by default, but any of the other explicit water models available in OpenMM, such as TIP4P-Ew [44], can be specified using the --water_model flag. Models are resolvated with the target number of waters by first solvating with zero padding, then incrementally increasing the box size and resolvating 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 [40] and TIP3P water [43] by default. The force field, water model, and simulation length can again be specified using the --ff, --water_model, and --simlength flags respectively. Further simulation parameters can be controlled via the API or via the CLI --api_params flag. Pressure control is performed with a Monte Carlo barostat as implemented in OpenMM, with a default pressure of 1 atm and ₄₆₃ a period of 50 timesteps. The remaining simulation param-464 eters have default values set to the same as for the implicit 465 solvent MD refinement.

Packaging

468 can be used to prepare models for other uses. 470 tions (specified via the --package_for flag) for com- 516 a particularly interesting subject for study with MSM meth-471 pressing models in preparation for data transfer, or for 517 ods [49], and this approach stands to benefit greatly from

474 computing platform Folding@home [4]. The module could Currently, Ensembler only supports residue definitions 475 easily be extended to add methods for preparing models provided by the forcefield definition files—it does not yet 476 for other purposes. For example, production simulations have the ability to derive new forcefield parameters for un- 477 could alternatively be run using Copernicus [5, 6]—a framecommon amino acids, cofactors, or ions in a consistent way. 478 work for performing parallel adaptive MD simulations— Explain how user-specified overrides can be used to spec- 479 or GPUGrid [7]—a distributing computing platform which ify specific protonation states or alternative/non-natural 400 relies on computational power voluntarily donated by the owners of nondedicated GPU-equipped computers.

Other features

Tracking provenance information

To aid the user in tracking the provenance of each model, each pipeline function also outputs a metadata file, which 486 helps to link data to the software version used to generate it (both **Ensembler** and its dependencies), and also provides 488 timing and performance information, and other data such as hostname.

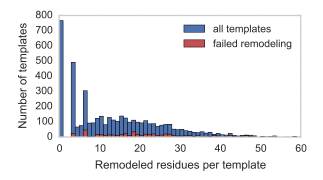
Rapidly modeling a single template

For users interested in simply using **Ensembler** to rapidly generate a set of models for a single template sequence, En-493 **sembler** provides a command-line tool quickmodel, which performs the entire pipeline for a single target with a small ₄₉₅ number of templates. For larger numbers of models (such as entire protein families), modeling time is greatly reduced by using the main modeling pipeline, which is parallelized via 498 MPI, distributing computation across each model (or across each template, in the case of the loop reconstruction code), 500 and scaling (in a "pleasantly parallel" manner) up to the number of models generated.

RESULTS

Modeling of all human tyrosine kinase catalytic domains

As a first application of **Ensembler**, we have built models for the human TK family. TKs (and protein kinases in general) play important roles in many cellular processes and are involved in a number of types of cancer [45]. For example, a translocation between the TK Abl1 and the pseudokinase Bcr is closely associated with chronic myelogenous leukemia [46], while mutations of Src are associated with colon, breast, prostate, lung, and pancreatic cancers [47]. Protein kinase domains are thought to have multiple acces-**Ensembler** provides a packaging module which 513 sible metastable conformation states, and much effort is di-The 514 rected at developing kinase inhibitor drugs which bind to package_models subcommand currently provides func- 515 and stabilize inactive conformations [48]. Kinases are thus 472 organizing them with the appropriate directory and file 518 the ability to exploit the full body of available genomic and



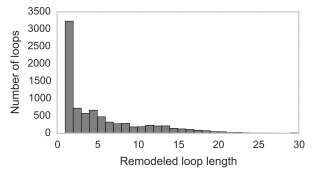


FIG. 2. Distributions for the number of missing residues in the **TK templates.** The upper histograms show the number of missing residues per template, for all templates (blue) and for only those templates for which template remodeling with the loopmodel subcommand failed (red). The lower histogram shows the number of residues in each missing loop, for all templates.

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 human TK domains annotated in UniProt as targets, and all available structures of protein kinase domains (of any species) as templates, using the commands shown in Box 1. This returned 93 target sequences and 4433 template structures, giving a total of 412,269 targetemplate pairs. The templates were derived from 3028 indiidual PDB entries and encompassed 23 different species. with 3634 template structures from human kinase con- 581 structs.

The resultant models are available as part of a supplementary dataset which can be downloaded from the Dryad Digital Repository (DOI: 10.5061/dryad.7fg32).

Ensembler modeling statistics

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Crystallographic structures of kinase catalytic domains 589

542 tion stage to reconstruct very long unresolved loops, unresolved template residues were first remodeled using the loopmodel subcommand. Out of 3666 templates with one or more missing residues, 3134 were successfully remodeled by the Rosetta loop modeling stage (with success defined simply as program termination without error); most remodeling failures were attributable to unsatisfiable spatial constraints imposed by the original template structure. There was some correlation between remodeling failures and the number of missing residues (Fig. 2, top); templates for which remodeling failed had a median of 20 missing residues, compared to a median of 14 missing residues for templates for which remodeling was successful.

Following loop remodeling, the **Ensembler** pipeline was performed up to and including the implicit solvent MD refinement stage, which completed with 389,067 (94%) surviving models across all TKs. To obtain statistics for the solvation stage without generating a sizeable amount of coordinate data (with solvated PDB coordinate files taking up about 0.9 MB each), the solvate subcommand was performed for two representative individual kinases (Src and Abl1).

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The number of models which survived each stage are shown in Fig. 1, indicating that the greatest attrition occurred during the modeling stage. The number of refined models for each target ranged from 4046 to 4289, with a median of 4185, mean of 4184, and standard deviation of 57. Fig. 1 also indicates the typical timing achieved on a cluster for each stage, showing that the build_models and refine_implicit_md stages are by far the most compute-572 intensive.

The files generated for each model (up to and including the implicit solvent MD refinement stage) totaled \sim 116 kB in size, totalling 0.5 GB per TK target or 42 GB for all 93 targets. The data generated per model breaks down as 39 kB for the output from the modeling stage (without saving MODELLER restraints files, which are about 397 kB per model) and 77 kB for the implicit solvent MD refinement stage.

Evaluation of model quality and utility

All tyrosine kinases

To evaluate the variety of template sequence similarities relative to each target sequence, we calculated sequence identity distributions, as shown in Fig. 3. This suggests an intuitive division into three categories, with 355,712 models in the 0-35% sequence identity range, 51,330 models in the 35–55% range, and 5227 models in the 55–100% range. We then computed the RMSD distributions for the models created for each target (relative to the model derived from generally contain a significant number of missing residues 590 the template with highest sequence identity) Fig. 4, to as-(median 11, mean 14, standard deviation 13, max 102) due to $_{591}$ sess the diversity of conformations captured by the modthe high mobility of several loops (Fig. 2, top), with a number 592 eling pipeline. Furthermore, to understand the influence of these missing spans being significant in length (median 5, 593 of sequence identity on the conformational similarities of mean 7, standard deviation 6, max 82; Fig. 2, bottom). To re- 594 the resulting models, the RMSD distributions were stratiduce the reliance on the MODELLER rapid model construc- 595 fied based on the three sequence identity categories de-

```
ensembler gather_targets --query 'family:"tyr protein kinase family" AND organism:"homo sapiens" AND reviewed:yes'
                         --uniprot_domain_regex '^Protein kinase(?!; truncated)(?!; inactive)
ensembler gather_templates --gather_from uniprot --query 'domain: "Protein kinase" AND reviewed: yes
                           --uniprot_domain_regex '^Protein kinase(?!; truncated)(?!; inactive)'
```

Box 1. Ensembler command-line functions used to select targets and templates. The commands retrieve target and template data by querying UniProt. The query string provided to the gather_targets command selects all human tyrosine protein kinases which have been reviewed by a curator, while the query string provided to the gather_templates command selects all reviewed protein kinases of any species. The --uniprot_domain_regex flag is used to select a subset of the domains belonging to the returned UniProt protein entries, by matching the domain annotations against a given regular expression. In this example, domains of type "Protein kinase", "Protein kinase 1", and "Protein kinase 2" were selected, while excluding many other domain types such as "Protein kinase; truncated", "Protein kinase; inactive", "SH2", "SH3", etc. Target selection simply entails the selection of sequences corresponding to each matching UniProt domain. Template selection entails the selection of the sequences and structures of any PDB entries corresponding to the matching UniProt domains.

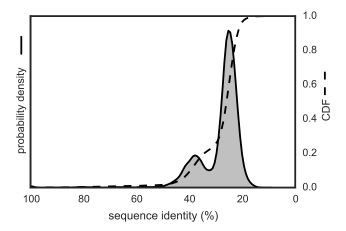
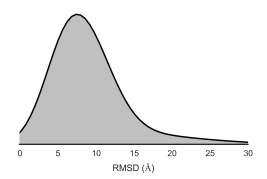


FIG. 3. Template-target sequence identity distribution for human tyrosine kinase catalytic domains. Sequence identities are calculated from all pairwise target-template alignments, where targets are human kinase catalytic domain sequences and templates are all kinase catalytic domains from any organism with structures in the PDB, as described in the text. A kernel density estimate of the target-template sequence identity probability density function is shown as a solid line with shaded region, while the corresponding cumulative distribution function is shown as a dashed line.

scribed above. This analysis indicates that higher sequence identity templates result in models with lower RMSDs, while templates with remote sequence identities result in larger RMSDs on average.

We also analyzed the potential energies of the models at the end of the implicit solvent MD refinement stage. These ranged from -14180 kT to -3160 kT, with a median of -9501 kT, mean of -9418 kT, and a standard deviation of 1198 kT (with a simulation temperature of 300 K). The 611 distributions-stratified using the same sequence identity ranges as above—are plotted in Fig. 5, indicating that higher 612 the first 1 ps of simulation.



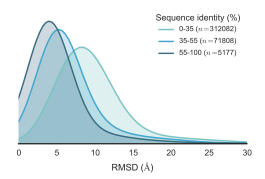


FIG. 4. Distribution of RMSDs to all TK catalytic domain models relative to the model derived from the highest sequence identity template. Distributions are built from data from all 93 TK domain targets. To better illustrate how conformational similarity depends on sequence identity, the lower plot illustrates the distributions as stratified into three sequence identity classes: high identity (55-100%), moderate identity (35-55%), and remote identity (0-35%). The plotted distributions have been smoothed using kernel density estimation.

Src and Abl1

To provide a more detailed evaluation of the variety and sequence identity templates tend to result in slightly lower 613 utility of generated models, we have analyzed two specific energy models. Of the 4973 models which failed to complete 614 TKs (Src and Abl1) in depth. Due to their importance in canthe implicit refinement MD stage, all except 9 failed within 615 cer, these kinases have been the subject of numerous de-616 tailed structural and simulation studies. In terms of struc-

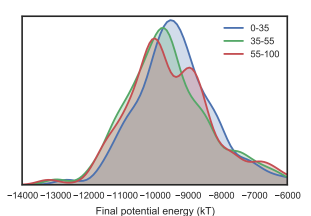


FIG. 5. Distribution of final energies from implicit solvent MD refinement of TK catalytic domain models. To illustrate how the energies are affected by sequence identity, the models are separated into three sequence identity classes: high identity (55–100%). moderate identity (35-55%), and remote identity (0-35%). The plotted distributions have been smoothed using kernel density estimation. Refinement simulations were carried out at the default 648 temperature of 300 K.

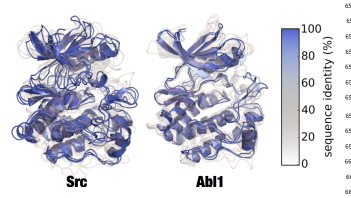


FIG. 6. Superposition of clustered models of Src and Abl1. Superposed renderings of nine models each for Src and Abl1, giving some indication the diversity of conformations generated by Ensembler. The models for each target were divided into three sequence identity ranges (as in Fig. 4), and RMSD-based k-medoids clustering was performed (using the msmbuilder clustering package [18]) 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.

tural data, a large number of crystal structures have been solved (with or without ligands such as nucleotide substrate nimetics or small-molecule inhibitors), revealing a variety of conformations accessible to these kinases. A recent largeselectivity and affinity of imatinib recognition of Abl [50].

627 Abl1. Models were first stratified into three ranges, based on the structure of the sequence identity distribution (Fig. 3), then subjected to RMSD-based k-medoids clustering (using the msmbuilder clustering package [18]) to pick three representative models from each sequence identity range. Each model is colored and given a transparency based on the sequence identity between the target and template sequence. The figure gives an idea of the variance present in the generated models. High sequence identity models (in opaque blue) tend to be quite structurally similar, with some variation in loops or changes in domain orientation.

The Abl1 renderings in Fig. 6 indicate one high sequence identity model with a long unstructured region at one of the termini, which was unresolved in the original template structure. While such models are not necessarily incorrect or undesirable, it is important to be aware of the effects they may have on production simulations performed under periodic boundary conditions, as long unstructured termini can be prone to interact with a protein's periodic image. Lower sequence identity models (in transparent white or red) indicate much greater variation in all parts of the structure. We believe the mix of high and low sequence identity models to be particularly useful for methods such as MSM building, which require thorough sampling of the conformational 651 landscape. The high sequence identity models could be considered to be the most likely to accurately represent true 653 metastable states. Conversely, the lower sequence identity 654 models could be expected to help push a simulation into re-655 gions of conformation space which might take intractably long to reach if starting a single metastable conformation.

Comparison with known biochemically relevant conformations. To evaluate the models of Src and Abl1 in the context of the published structural biology literature on 660 functionally relevant conformations, we have focused on two residue pair distances thought to be important order parameters for the regulation of protein kinase domain activity. We use the residue numbering schemes for chicken Src (commonly employed in the literature even in reference to human Src) [51, 52] and human Abl1 isoform A [53–55] respectively; the exact numbering schemes are provided in 667 Appendix 1.

Fig. 7 shows two structures of Src believed to represent inactive (PDB code: 2SRC) [51] and active (PDB code: 1Y57) [52] 670 states. One notable feature which distinguishes the two structures is the transfer of an electrostatic interaction of 672 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 [56], and a number of experimental and simulation studies have suggested that 676 this electrostatic switching process plays a role in a regulatory mechanism shared across the protein kinase fam-678 ily [49, 57, 58]. As such, we have projected the **Ensembler** scale MSM study has also studied the activation pathway of 679 models for Src and Abl1 onto a space consisting of the dis-Src [49], while a separate study employed biased sampling 🚳 tances between these two residue pairs (Fig. 8). The modtechniques to dissect the role of conformational changes in 681 els show strong coverage of regions in which either of the electrostatic interactions is fully formed (for models across Visualizing model structural diversity. Fig. 6 shows a 683 all levels of target-template sequence identity), as well as a superposition of a set of representative models of Src and 684 wide range of regions in-between (mainly models with low

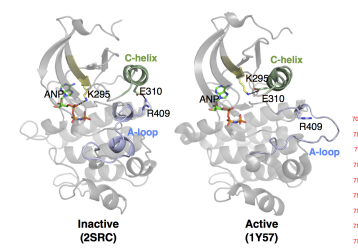


FIG. 7. Two structures of Src, indicating certain residues involved in activation. In the inactive state, E310 forms a salt bridge with R409. During activation, the α C-helix (green) moves and rotates, orienting E310 towards the ATP-binding site and allowing it to instead form a salt bridge with K295. This positions K295 in the appropriate position for catalysis. Note that ANP (phosphoaminophosphonic acid-adenylate ester; an analog of ATP) is only physically present in the 2SRC structure. To aid visualization of the active site in 1Y57, it has been included in the rendering by structurally aligning the surrounding homologous protein residues.

sequence identity). We thus expect that such a set of models, if used as starting configurations for highly parallel MD simulation, could greatly aid in sampling of functionally relevant conformational states.

AVAILABILITY AND FUTURE DIRECTIONS

Availability

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The code for **Ensembler** is hosted on the collabora-692 tive open source software development platform GitHub (github.com/choderalab/ensembler). The latest release can be installed via the conda package manager for Python (conda.pydata.org), using the two commands shown in This will install all dependencies except for MODELLER and Rosetta, which are not available through the conda package manager, and thus must be installed separately by the user. The latest source can be downloaded from the GitHub repository, which also contains up-to-date instructions for building and installing the code. Documentation can be found at ensembler.readthedocs.org. 702

A supplementary dataset can also be downloaded from the Dryad Digital Repository (DOI: 10.5061/dryad.7fg32). This contains the TK models described in the III section, general information on the targets and templates, plus a script ₇₅₅ lations combined with Monte Carlo sampling of side chain and instructions for regenerating the same dataset.

Future Directions

conda config -add channels https://conda.binstar.org/omnia conda install ensembler

Box 2. Ensembler installation using conda.

We recognize that the current version of **Ensembler** has a 710 number of limitations that limits its domain of applicability: Support for nonnatural amino acids is currently rudimentary and confined to those already appearing in the forcefield; cofactors cannot currently be automatically modeled in; ligands, cofactors, and nonnatural amino acids cannot 715 yet be automatically parameterized; protonation state as-716 signment is limited to selection of the most populated state based on the intrinsic p K_a or user-specified overrides; the modeling of missing loops is rudimentary, relying on the subsequent dynamics for relaxation; there is not yet support for modeling of distinct domains from different templates, or the use of multiple templates to model a single domain. Nevertheless, there are a great number of use cases for this ₇₂₃ first version of an automated tool for simulation preparation at the superfamily scale. To expand this domain of applicability, there are a number of obvious additions and improvements which we plan to implement in future versions of **En-**727 sembler.

Comparative modeling. Comparative protein modeling can be approached in a number of different ways, with varying degrees of complexity. The comparative modeling stage of **Ensembler** currently uses MODELLER, but a number of 732 excellent alternatives—such as RosettaCM [13] and the I-TASSER Suite [14]—can be added as user-selectable alter-₇₃₄ native choices. Additional options could be added to allow more expensive loop-modeling approaches to be employed to handle long insertions.

Protonation states. Some amino acids can exist in different protonation states, depending on pH and on their local environment. These protonation states can have im-740 portant effects on biological processes. For example, long 141 timescale MD simulations have suggested that the conformation of the DFG motif of the TK Abl1—believed to be an important regulatory mechanism [59]—is controlled by protonation of the aspartate [60]. Currently, protonation states 745 are assigned simply based on pH (a user-controllable parameter). At neutral pH, histidines have two protonation states which are approximately equally likely, and in this sit-148 uation the selection is therefore made based on which state 749 results in a better hydrogen bond. It would be highly de-750 sirable to instead use a method which assigns amino acid 751 protonation states based on a rigorous assessment of the ₇₅₂ local environment. We thus plan to implement an interface and command-line function for assigning protonation 754 states with MCCE2 [61-63], which uses electrostatics calcuconformers to calculate pKa values.

Cofactors, structural ions, and ligands. Many proteins require the presence of various types of non-protein atoms and molecules for proper function, such as metal ions $_{760}$ (e.g. Mg $^{+2}$), cofactors (e.g. ATP) or post-translational modi-

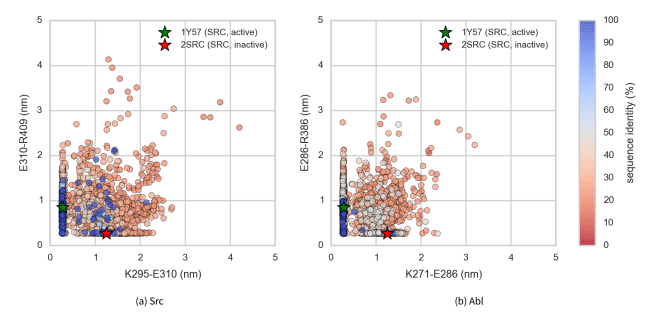


FIG. 8. Src and Abl1 models projected onto the distances between two conserved residue pairs, colored by sequence identity. Two Src structures (PDB entries 1Y57 [52] and 2SRC [51]) are projected onto the plots for reference, representing active and inactive states respectively. These structures and the residue pairs analyzed here are depicted in Fig. 7. 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).

761 fications (e.g. phosphorylation, methylation, glycosylation, 791 yet the minimum and maximum lengths are 102 and 801 remetal cations, and display significantly increased activity in 797 questions would arise as to how best to approach this. the presence of Mg^{2+} [64], the divalent cation with highest oncentration in mammalian cells. Metal ions are often not esolved in experimental structures of proteins, but by takng into account the full range of available structural data, it should be possible in many cases to include metal ions based on the structures of homologous proteins. We are careful to point out, however, that metal ion parameters in classical MD force fields have significant limitations, particularly in their interactions with proteins [65]. Cofactors and post-translational modifications are also often not fully resolved in experimental structures, and endogenous cofactors are frequently substituted with other molecules to facilitate experimental structural analysis. Again, Ensembler could exploit structural data from a set of homologous proeins to model in these molecules, although there will likely be a number of challenges to overcome in the design and 184 implementation of such functionality.

Long insertions and deletions. Another limitation with 809 the present version of Ensembler involves the treatment of 810 and Arien S. Rustenburg (MSKCC) for many excellent softmembers of a protein family with especially long residue in- sn ware engineering suggestions. The authors thank Nicholas sertions or deletions. For example, the set of all human pro- 812 M. Levinson (University of Minnesota), Markus A. Seeliger tein kinase domains listed in UniProt have a median length 813 (Stony Brook), Diwakar Shukla (Stanford), and Avner Schof 265 residues (mean 277) and a standard deviation of 45, 814 lessinger (Mount Sinai) for helpful scientific feedback on

etc.), and we thus plan for **Ensembler** to eventually have 792 spectively. The latter value corresponds to the protein kithe capability to include such entities in the generated mod- 793 nase domain of serine/threonine-kinase *greatwall*, which els. Binding sites for metal ions are frequently found in pro- 794 includes a long insertion between the two main lobes of teins, often playing a role in catalysis. For example, pro- $_{795}$ the catalytic domain. In principle, such insertions could be tein kinase domains contain two binding sites for divalent 796 excluded from the generated models, though a number of

Conclusion

We believe **Ensembler** to be an important first step toward enabling computational modeling and simulation of proteins on the scale of entire protein families, and suggest 802 that it could likely prove useful for tasks beyond its original aim of providing diverse starting configurations for MD simulations. The code is open source and has been developed with extensibility in mind, in order to facilitate its customization for a wide range of potential uses by the wider scientific 807 community.

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Appendix 1: Sequences and residue numbering schemes for Src and Abl1

⁹⁸⁰ Kinase catalytic domains are highlighted in red, and the conserved residues analyzed in the main text (Figs. 7 and 8) are ⁹⁸¹ highlighted with yellow background.

Human Abl1 sequence

983	1	${\tt MLEICLKLVG}$	CKSKKGLSSS	SSCYLEEALQ	RPVASDFEPQ	GLSEAARWNS	KENLLAGPSE	60
984	61	${\tt NDPNLFVALY}$	${\tt DFVASGDNTL}$	SITKGEKLRV	LGYNHNGEWC	EAQTKNGQGW	VPSNYITPVN	120
985	121	SLEKHSWYHG	PVSRNAAEYL	LSSGINGSFL	VRESESSPGQ	RSISLRYEGR	VYHYRINTAS	180
986	181	DGKLYVSSES	RFNTLAELVH	HHSTVADGLI	TTLHYPAPKR	${\tt NKPTVYGVSP}$	NYDKWEMERT	240
987	241	DITMKHKLGG	GQYGEVYEGV	WKKYSLTVAV	K TLKEDTMEV	EEFLK E AAVM	KEIKHPNLVQ	300
988	301	LLGVCTREPP	FYIITEFMTY	GNLLDYLREC	NRQEVNAVVL	LYMATQISSA	MEYLEKKNFI	360
989	361	${\tt HRDLAARNCL}$	VGENHLVKVA	$DFGLS^{\mathbf{R}}LMTG$	DTYTAHAGAK	FPIKWTAPES	LAYNKFSIKS	420
990	421	DVWAFGVLLW	EIATYGMSPY	PGIDLSQVYE	LLEKDYRMER	PEGCPEKVYE	LMRACWQWNP	480
991	481	SDRPSFAEIH	QAF ETMFQES	SISDEVEKEL	GKQGVRGAVS	TLLQAPELPT	KTRTSRRAAE	540
992	541	${\tt HRDTTDVPEM}$	${\tt PHSKGQGESD}$	PLDHEPAVSP	LLPRKERGPP	${\tt EGGLNEDERL}$	LPKDKKTNLF	600
993	601	$\mathtt{SALIKKKKKT}$	APTPPKRSSS	FREMDGQPER	${\tt RGAGEEEGRD}$	ISNGALAFTP	LDTADPAKSP	660
994	661	${\tt KPSNGAGVPN}$	GALRESGGSG	FRSPHLWKKS	STLTSSRLAT	${\tt GEEEGGGSSS}$	KRFLRSCSAS	720
995	721	${\tt CVPHGAKDTE}$	WRSVTLPRDL	QSTGRQFDSS	TFGGHKSEKP	ALPRKRAGEN	RSDQVTRGTV	780
996	781	${\tt TPPPRLVKKN}$	EEAADEVFKD	IMESSPGSSP	${\tt PNLTPKPLRR}$	QVTVAPASGL	PHKEEAGKGS	840
997	841	${\tt ALGTPAAAEP}$	VTPTSKAGSG	${\tt APGGTSKGPA}$	EESRVRRHKH	${\tt SSESPGRDKG}$	KLSRLKPAPP	900
998	901	${\tt PPPAASAGKA}$	GGKPSQSPSQ	EAAGEAVLGA	KTKATSLVDA	VNSDAAKPSQ	PGEGLKKPVL	960
999	961	${\tt PATPKPQSAK}$	PSGTPISPAP	VPSTLPSASS	ALAGDQPSST	AFIPLISTRV	SLRKTRQPPE	1020
1000	1021	${\tt RIASGAITKG}$	VVLDSTEALC	LAISRNSEQM	ASHSAVLEAG	KNLYTFCVSY	VDSIQQMRNK	1080
1001	1081	FAFREAINKL	ENNLRELQIC	PATAGSGPAA	TQDFSKLLSS	VKEISDIVQR		1130

979

982

1002

Sequences for human and chicken Src, aligned using Clustal Omega

1003	SRC_HUMAN	1	MGSNKSKPKD	ASQRRRSLEP	AENVHGAGGG	AFPASQTPSK	PASADGHRGP	SAAFAPAAAE	60
1004	SRC_CHICK	1	${\tt MGSSKSKPKD}$	${\tt PSQRRRSLEP}$	PDSTHHG	GFPASQTPNK	${\tt TAAPDTHRTP}$	SRSFGTVATE	57
1005	i		***.*****	******	:* *	.******	*: * ** *	* :**:*	
1006	SRC_HUMAN	61	PKLFGGFNSS	${\tt DTVTSPQRAG}$	${\tt PLAGGVTTFV}$	ALYDYESRTE	TDLSFKKGER	LQIVNNTEGD	120
1007	SRC_CHICK	58	PKLFGGFNTS	${\tt DTVTSPQRAG}$	${\tt ALAGGVTTFV}$	ALYDYESRTE	TDLSFKKGER	LQIVNNTEGD	117
1008	1		******	******	******	******	******	*****	
1009	SRC_HUMAN	121	WWLAHSLSTG	QTGYIPSNYV	APSDSIQAEE	WYFGKITRRE	SERLLLNAEN	PRGTFLVRES	180
1010	SRC_CHICK	118	WWLAHSLTTG	QTGYIPSNYV	APSDSIQAEE	WYFGKITRRE	SERLLLNPEN	PRGTFLVRES	177
101	l		******:**	******	******	******	***** **	******	
1012	SRC_HUMAN	181	ETTKGAYCLS	VSDFDNAKGL	${\tt NVKHYKIRKL}$	DSGGFYITSR	TQFNSLQQLV	AYYSKHADGL	240
1013	SRC_CHICK	178	ETTKGAYCLS	${\tt VSDFDNAKGL}$	${\tt NVKHYKIRKL}$	DSGGFYITSR	TQFSSLQQLV	AYYSKHADGL	237
1014	ı		******	******	******	******	***.*****	******	
1015	SRC_HUMAN	241	CHRLTTVCPT	${\tt SKPQTQGLAK}$	${\tt DAWEIPRESL}$	RLEVKLGQGC	FGEVWMGTWN	GTTRVAIKTL	300
1016	SRC_CHICK	238	CHRLTNVCPT	${\tt SKPQTQGLAK}$	${\tt DAWEIPRESL}$	RLEVKLGQGC	FGEVWMGTWN	GTTRVAIKTL	297
1017	,		*****	******	******	******	******	******	
1018	SRC_HUMAN	301	KPGTMSPEAF	LQEAQVMKKL	RHEKLVQLYA	VVSEEPIYIV	TEYMSKGSLL	DFLKGETGKY	360
1019	SRC_CHICK	298	KPGTMSPEAF	$LQ\textcolor{red}{\mathbf{E}} AQVMKKL$	RHEKLVQLYA	VVSEEPIYIV	TEYMSKGSLL	DFLKGEMGKY	357
1020	1		******	******	******	******	******	***** ***	
102	SRC_HUMAN	361	LRLPQLVDMA	AQIASGMAYV	ERMNYVHRDL	RAANILVGEN	LVCKVADFGL	AR LIEDNEYT	420
1022	SRC_CHICK	358	LRLPQLVDMA	AQIASGMAYV	ERMNYVHRDL	RAANILVGEN	LVCKVADFGL	ARLIEDNEYT	417
1023	1		******	******	******	******	******	******	
1024	SRC_HUMAN	421	ARQGAKFPIK	${\tt WTAPEAALYG}$	RFTIKSDVWS	FGILLTELTT	${\tt KGRVPYPGMV}$	NREVLDQVER	480
1025	SRC_CHICK	418	ARQGAKFPIK	${\tt WTAPEAALYG}$	RFTIKSDVWS	FGILLTELTT	KGRVPYPGMV	NREVLDQVER	477
1026	i		******	******	******	******	******	******	
1027	SRC_HUMAN	481	GYRMPCPPEC	${\tt PESLHDLMCQ}$	${\tt CWRKEPEERP}$	TFEYLQAFLE	DYFTSTEPQY	QPGENL	536
1028	SRC_CHICK	478	GYRMPCPPEC	${\tt PESLHDLMCQ}$	${\tt CWRKDPEERP}$	TFEYLQAFLE	DYFTSTEPQY	QPGENL	533
1029)		******	******	****:****	******	******	*****	