A flexible framework for multi-particle refinement in cryo-electron tomography

Alister Burt\textsuperscript{1,*}, Lorenzo Gaifas\textsuperscript{1}, Tom Dendooven\textsuperscript{2}, Irina Gutsche\textsuperscript{1,*}

\textsuperscript{1} Institut de Biologie Structurale, Universit\'e Grenoble Alpes, CEA, CNRS, IBS, Grenoble, France, \textsuperscript{2} MRC Laboratory of Molecular Biology, Cambridge, United Kingdom

* alisterburt@gmail.com (AB); irina.gutsche@ibs.fr (IG)

Abstract

Cryo-electron tomography (cryo-ET) and subtomogram averaging (STA) are increasingly used for macromolecular structure determination in situ. Here, we introduce a set of computational tools and resources designed to enable flexible approaches to STA through increased automation and simplified metadata handling. We create a bidirectional interface between the Dynamo software package and the Warp-Relion-M pipeline, providing a framework for ab initio and geometrical approaches to multiparticle refinement in M. We illustrate the power of working within this framework by applying it to EMPIAR-10164, a publicly available dataset containing immature HIV-1 virus-like particles (VLPs), and a challenging in situ dataset containing chemosensory arrays in bacterial minicells. Additionally, we provide a comprehensive, step-by-step guide to obtaining a 3.4-Å reconstruction from EMPIAR-10164. The guide is hosted on https://teamtomo.org/, a collaborative online platform we establish for sharing knowledge about cryo-ET.

Introduction

Cryo-electron tomography (cryo-ET) is an imaging technique rapidly gaining popularity for the direct visualisation of unique biological objects in 3D. Repeating structural motifs present in cryo-ET data can be reconstructed at higher resolution by subtomogram averaging (STA), allowing for the possibility of studying macromolecular structure in situ [1]. STA has developed alongside single-particle analysis (SPA) in cryo-electron microscopy (cryo-EM), a technique that has benefitted significantly in the last 10 years from advances in both electron detection hardware and image processing software [2,3].

State-of-the-art STA workflows often co-opt tools and adapt ideas from SPA for tomography and STA [4,5]. An unfortunate consequence of this side-by-side development is a somewhat fragmented software ecosystem with no standardisation of file formats or metadata conventions [6]. Despite the advent of complete or near-complete integrated solutions for cryo-ET and STA [7–9], optimal methodology for a given dataset often requires the creative combination of different approaches that may not all be present within a single integrated pipeline. For those new to cryo-ET, the burden of interfacing many different software packages in a complex workflow often represents a barrier to the testing of alternative approaches.

Image processing for STA from cryo-ET data is a complex, multistep process that can broadly be divided into 3 blocks (Fig 1). The first, “preprocessing,” generates 3D
reconstruction of the chemosensory array. Information underlying data displayed at Figs 5 and 6 is available for download at doi:10.5281/zenodo.4783129 and doi:10.5281/zenodo.4783151 respectively. Source code for all tools and resources described here is available at: • autoalign_dynamo - https://github.com/alisterburt/autoalign_dynamo • mdoscpoofer - https://github.com/alisterburt/mdoscpoofer • starfile - https://github.com/alisterburt/starfile • dynamatable - https://github.com/alisterburt/dynamatable • eulerangles - https://github.com/alisterburt/eulerangles All Python packages are made available on the Python package index (PyPI).

Benchmarking code for eulerangles can be found at https://gist.github.com/alisterburt/4a32e9c122498ac0ab482ee65ba44ba10.

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Abbreviations: API, application programming interface; cryo-EM, cryo-electron microscopy; cryo-ET, cryo-electron tomography; CSU, core signalling unit; CTF, contrast transfer function; EMDB, Electron Microscopy Data Bank; FIB, focused ion beam; FSC, Fourier shell correlation; GUI, graphical user interface; PCA, principal component analysis; RMSD, root mean square deviation; SPA, single-particle analysis; STA, subtomogram averaging; VLP, virus-like particle. reconstructions (tomograms) from experimental 2D micrographs. The second, “particle picking,” creates putative positions and orientations for objects of interest within each volume as well as initial reference(s) for subsequent refinement. Finally, the “refinement” block is concerned with the optimisation of reconstruction(s) from imaging data associated with a set of particle positions.

The preprocessing block encompasses all steps in the generation of tomograms from experimental data. Experimental cryo-ET data are usually acquired as a set of multiframe 2D micrographs, one per tilt angle in a tilt series. Typically, this block includes per tilt interframe motion estimation and correction, contrast transfer function (CTF) estimation, tilt series alignment, and 3D reconstruction [10]. Tilt series alignment is the least automated of these steps, with significant time often spent optimising alignments in an attempt to produce more accurate reconstructions.

Particle picking is somewhat entangled with the process of initial model generation, with some particle picking methods requiring an initial model and others being reference free. Exhaustive, reference-based template matching approaches are widely accepted as an imperfect solution for particle picking due to the computational overhead and the need for subsequent dataset cleaning [11,12]. Care must also be taken to avoid model bias. Particle picking methods based on deep learning have been proposed, which may help to address some of these limitations [13,14]. Another class of particle picking methods, termed “geometrical particle picking,” derives putative particle positions and orientations from a 3D model of a supporting geometry, usually generated from minimal manual annotations [15]. These methods can be used to impose prior knowledge about how particle poses relate to the supporting geometry during refinement. Employing such a priori knowledge is advantageous, reducing both the computational burden of global searches and the likelihood of ending up in incorrect local minima during refinement. These advantages come at the expense of extra time spent on manual annotation and metadata management.

Refinement in STA has typically meant the optimisation of particle poses, considering particles as rigid bodies within fixed 3D reconstructions [16]. The potential for studying macromolecular structure at intermediate resolution (3 to 5 Å) by STA became reality with the advent of an efficient method for 3D CTF correction [17]. Inspired by progress in SPA, the refinement block may now include reference-based procedures for the a posteriori optimisation of tilt series alignment parameters, frame series alignments, and optical parameters [7–9,18]. Despite these algorithmic advances, there remain relatively few examples of reconstructions from tilt series data in this resolution regime in the Electron Microscopy Data Bank (EMDB), indicating that this work remains nontrivial (S1 Fig). We posit that a possible reason for this discrepancy is the complexity of integrating the various tools available for each step of the workflow in the fragmented cryo-ET software ecosystem.

Warp-RELION-M is a user-friendly solution for cryo-ET preprocessing, STA, and multi-particle refinement, yielding the highest-resolution reconstructions from cryo-ET data seen thus far [9]. The current version of Warp (v1.0.9) is designed to process data collected using SerialEM and does not integrate automated tilt series alignment procedures. The pipeline assumes that an initial model is available for template matching and does not interface directly with tools for geometrical particle picking. Dynamo is a flexible, extensible software environment for STA, providing a variety of powerful tools for the management, annotation, and analysis of cryo-ET data [15,19].

The combination of the tools in these software packages would yield a computational framework capable of more automated, ab initio, and geometrical approaches to multi-particle refinement in M. In this manuscript, we describe our efforts to facilitate working within this framework. In the Materials and methods section, a set of tools is presented, which automate fiducial-based tilt series alignment, simplify metadata handling, and create a bidirectional
interface between Dynamo and Warp-RELION-M. In the Results and discussion section, we demonstrate the benefits of our metadata handling tools and illustrate the power of working within this computational framework on 2 datasets. Finally, we provide a comprehensive step-by-step guide to obtaining a 3.4-Å reconstruction of the HIV-1 CA-SP1 hexamer ab initio from 5 tilt series of EMPIAR-10164. This guide is presented as a part of a collaborative, online platform (https://teamtomo.org/) that we establish as a platform for sharing knowledge about cryo-ET data processing.
Materials and methods

Integrating on-the-fly tilt series alignment into the Warp preprocessing pipeline

Tomogram reconstruction within the Warp preprocessing pipeline currently requires the use of an external software package, IMOD [20], for tilt series alignment. In the absence of a fully integrated solution in the currently available version of Warp (1.0.9), we developed autoalign_dynamo, a package for automating fiducial-based tilt series alignment using Dynamo and IMOD. dautoalign4warp is a program for on-the-fly alignment of batches of tilt series from Warp. The program proceeds by dynamic generation of a Dynamo tilt series alignment workflow with appropriate parameters derived from minimal user input. The workflow is executed, generating a set of refined fiducial positions. Refined fiducial positions are converted using dms2mod (provided by autoalign_dynamo) and used to generate tilt series alignment parameters using the IMOD program tiltalign. Key tiltalign parameters fixed within this procedure are the following: solve for 1 tilt axis for the entire tilt series; fix tilt angles at their nominal values; and use robust fitting for parameter estimation. All tiltalign parameters can be seen and modified in the tiltalign_wrapper script. All data are subsequently organised such that alignments can be imported directly into Warp for tomogram reconstruction, requiring no further user input. The base function autoalign, which wraps Dynamo tilt series alignment functionality, is provided for those who wish to apply this procedure outside of the Warp pipeline.

Dynamo tilt series alignment workflows were released recently, and, as such, are not yet described in the literature, although details and the source code are available in the public domain [21]. To facilitate the readers understanding, we provide a brief overview of the alignment algorithm, noting that it was not implemented by us. A binary, synthetic template of a fiducial marker is generated based on user input and used to detect candidate fiducial positions within a tilt series by cross-correlation. A total of 300 subimages are extracted from the tilt series at the peaks of the resulting cross-correlation matrix and averaged to produce a template for detecting fiducial markers in the data. The template is used to detect initial fiducial positions in the tilt series by cross-correlation. Cross-correlation peaks are analysed, and those observations not meeting a minimum degree of rotational symmetry are discarded. Observations are indexed to link observations of the same fiducial marker in multiple micrographs by pairwise cross-correlation of the observations between neighbouring micrographs. The longest “trails” of linked observations are used to generate an initial 3D model of fiducial positions. The 3D model is iteratively reindexed by reprojection of fiducial positions against the tilt series, adding observations that fall within a distance threshold to the set of observations to be included for further refinement. Tilt images lacking at this point are reintegrated by a procedure comparing the reprojection of the current model with the cloud of initial observations found on that micrograph. The number of observations is maximised by the reintegration of missing fiducial markers from other images using the same procedure. The projection model is iteratively refined before the positions of fiducial markers are independently iteratively refined against an average of all observations of that marker. The final set of refined markers is pruned according to the root mean square deviation (RMSD) between measured fiducial position and the reprojected position of the 3D model.

A set of self-contained Python packages for metadata handling

In an effort to interface the primary metadata systems of RELION [22] and Dynamo [19] with the scientific Python [23] ecosystem, we provide 2 Python packages: starfile and dynamotable. These packages provide input/output functionality via a simple application programming interface (API), exposing data as pandas DataFrame objects.
Euler angles are used by many cryo-EM software packages to describe the orientation of a rigid body with respect to a fixed coordinate system. eulerangles is a Python package that provides a simple API for the batch conversion of Euler angles into rotation matrices, rotation matrices into Euler angles, and interconversion of Euler angles defined according to different conventions. Conversions can take place between all possible formulations of Euler angles in a right-handed coordinate system. A simple mechanism is provided for the definition and reuse of conventions from specific software packages. Documentation for eulerangles can be found at https://eulerangles.readthedocs.io/.

Enabling Warp preprocessing for data collected in Tomography 5

In the Warp cryo-ET preprocessing pipeline, only metadata from the SerialEM data collection program in the form of mdoc files are currently supported. Thermo Scientific Tomography 5 (Tomo 5) is the official solution provided with Thermo Scientific microscopes for electron tomography experiments [24]. We provide a small command line tool mdocspoof for the generation of mdoc files for a directory containing multiframe micrograph files of the form <basename>_<count>[<tilt_angle>]_fractions.mrc, as generated by Tomography 5. This tool enables use of the Warp preprocessing pipeline for users of Tomography 5.

Interfacing Dynamo and the Warp-RELION-M pipeline

The recommended procedure for particle picking in the Warp-RELION-M pipeline is exhaustive, reference-based template matching. Dynamo offers an interactive environment for picking particles based on supporting geometries in cryo-electron tomograms [15]. We provide warp2catalogue, a program that sets up a database called a Dynamo catalogue for tomograms reconstructed in Warp. The catalogue is set up such that all visualisation operations make use of a deconvolved reconstruction, filtered for improved visualisation, while subsequent particle extraction uses the corresponding unfiltered volume, simplifying the experience for the user.

For rigid body particle pose optimisation, STA in RELION is integrated into the existing pipeline. Alternative approaches to particle picking, pose optimisation, and classification may provide advantages over this workflow. We provide dynamo2m, a set of tools that create a bidirectional interface between Dynamo and the Warp-RELION-M pipeline. dynamo2warp allows for particle position and orientation data in Dynamo to be used for particle reconstruction in Warp, rigid body refinement in RELION, and multi-particle refinement in M. warp2dynamo provides a route to using particles reconstructed in Warp within Dynamo. An additional utility, relion_star_downgrade, is provided to enable reconstructing particles refined using RELION version 3.1 or higher in Warp 1.0.9.

Results and discussion

Tools for simplified metadata handling

The ability to test alternative approaches and rapidly iterate is key to optimising complex data analysis workflows like cryo-ET and STA. Exploratory data analysis such as checking the internal consistency of particle positions within a supramolecular assembly after a refinement is often key to understanding the limitations of a given approach. Implementing custom input/output functionality when attempting to work with a variety of nonstandard metadata presents a barrier to entry for those wishing to interactively explore their data or implement custom analysis routines. The Python packages starfile and dynamatable provide simple interfaces between the metadata systems of RELION, Dynamo, and a wealth of data analysis infrastructure, visualisation tools, and educational resources in the scientific Python ecosystem [25,26].
Converting between different Euler angle conventions is often a pain point for those wishing to interface different cryo-EM software packages due to the abundance of ambiguities in their interpretation [27]. eulerangles simplifies generating rotation matrices from Euler angles for custom analyses and facilitates building interfaces between pieces of software that interpret Euler angles according to differing conventions. Our vectorised implementation is approximately $10 \times$ faster than the popular scipy utility scipy.spatial.transform.Rotation, taking 381 ms versus 4.38 s for the conversion of a million Euler triplets, the eulerangles package thus being more suitable for working with large sets of Euler angles in Python.

As an example of the benefits derived from working within this ecosystem, we combine star-file and eulerangles with the packages mrcfile [28] (for reading and writing MRC format image files) and napari [29] (a fast, powerful multidimensional data visualisation library) to visualise tomograms, particle positions, and their oriented Z-vectors in 3D (Fig 2). Using these packages, creating such a visualisation is achieved in less than 25 lines of (human readable) code (S2 Fig).

**A flexible framework for ab initio and geometrical approaches to subtomogram averaging**

Each STA dataset is different, and analysis presents its own unique challenges. The tools provided in autoalign_dynamo and dynamo2m were designed to simplify cryo-ET data processing where possible and empower researchers with access to a vast array of alternative tooling where it may provide benefit. Combined, these tools enable working within a powerful, flexible framework for more automated, ab initio, and geometrical approaches to multi-particle refinement, linking Warp-RELION-M with Dynamo (Fig 3). In this section, we discuss and illustrate the benefits provided by our tools.

**Automation of tilt series alignment in Warp-RELION-M with autoalign_dynamo**

Optimisation of data collection strategies has significantly increased the throughput of cryo-ET data collection in recent years, with datasets now routinely exceeding 100 tilt series. As dataset sizes increase, so does the need for accurate, automated solutions to each and every step of complex data analysis workflows. Accurate tilt series alignment is often achieved using semiautomated procedures that quickly become tedious when faced with a large dataset. Providing the automated, robust fiducial-based tilt series alignment procedures from Dynamo as an on-the-fly solution tightly integrated into the Warp preprocessing pipeline greatly simplifies the generation of accurate 3D reconstructions for downstream data analysis. Algorithmically, Dynamo tilt series alignment differs from batchruntomo in IMOD in a few significant ways [30]. In IMOD, the fiducial center position is determined on a Sobel-filtered image. Dynamo iteratively refines the center position of fiducials in each image by alignment to a per-fiducial reference image generated from images of that fiducial in the whole tilt series. Additionally, the Dynamo procedure explicitly attempts to maximise the number of observations in a tilt series by reintegration of missing fiducials, leveraging an existing projection model for detection of missing fiducials and subsequent improvement of the model. In contrast, IMOD does not attempt to reintegrate markers not included in the initial seed model unless explicitly added by the user. A quantitative comparison between different automated tilt series alignment algorithms could form the basis for interesting future work.

We provide no integrated solution for aligning tilt series lacking exogenous fiducial markers, such as those from samples prepared by focused ion beam (FIB) milling, although procedures exist in other software packages [8]. As new tilt series alignment algorithms are developed, their integration into on-the-fly data processing workflows will be important for optimal use of both microscope and researcher time.
Enabling geometrical approaches to subtomogram averaging in Warp-RELION-M

warp2catalogue and dynamo2warp enable the use of geometrical approaches to STA to users of the Warp-RELION-M pipeline (Fig 4). The catalogue system in Dynamo and the interactive
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preprocessing

multi-frame micrographs → tilts-series stacks → tomograms

particle picking

tomograms → w2c catalogue → derive particle positions and orientations from a supporting geometry → reference based template matching → initial poses and reference

refinement

initial poses and reference → particle and CTF volume reconstruction → RELION maximum-likelihood classification and pose optimisation

Dynamo powerful, flexible tools for analysis and visualisation → multi-particle refinement → optimised physical model and reconstruction(s)
3D modelling tools in the *dtmslice* viewer make it easy to generate models of supporting geometries in sets of tomograms and use them to derive particle positions and orientations for subsequent STA experiments [15]. Examples of easy-to-annotate supporting geometries from which putative particle poses can be easily derived are vesicles (Fig 4A), arbitrarily shaped membrane surfaces (Fig 4B), filaments, and crystals. *warp2catalogue* simplifies the setup of a Dynamo catalogue for *Warp-RELION-M* users: The user annotates volumes filtered to aid visualisation, whereas particle extraction operations performed from the catalogue use the corresponding unfiltered data. This automation simplifies the use of an optimal workflow without unnecessary cognitive burden for the user.

*Dynamo* also provides *dpktbl.subbox.tableOnTable* for “subboxing,” deriving particle positions and orientations, which are geometrically related to an existing set of positions and orientations (Fig 4C). This procedure is often useful in cryo-ET for focussing analysis on subunits of a large complex after an initial consensus refinement [31]. *dynamo2warp* provides a simple mechanism for the integration of these powerful tools into the *Warp-RELION-M* pipeline.

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**Fig 3.** *autoalign_dynamo* and *dynamo2m* enable a powerful, flexible framework for *ab initio* and geometrical approaches to *STA* refinement of cryo-ET data. The cryo-ET preprocessing pipeline in *Warp* (steps marked by a W, top) is extended to include on-the-fly tilt series alignment in *Dynamo* (marked by a grey cog). Users benefit from increased automation. *warp2catalogue* augments the particle picking block in *Warp-RELION-M* to facilitate the use of geometrical particle picking tools in *Dynamo*, enabling ab initio reconstruction leveraging prior knowledge of particle orientation during alignment and averaging. In the refinement block, *dynamo2m* enables users to move freely between *Dynamo* and *Warp-RELION-M*, choosing the tool most well suited to the problems posed by their data. Combined, these tools enable ab initio and geometrical approaches to multi-particle refinement of cryo-ET data. *cryo-ET*, cryo-electron tomography; *CTF*, contrast transfer function.

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**Fig 4.** Geometrical approaches to STA. Particle positions and orientations are generated from minimal manual annotations. In (a), a sphere is defined by a center and edge point, and positions with an orientation normal to the surface are seeded on the sphere. In (b), an arbitrary surface is defined by annotation of points on that surface in a number of tomographic slices. A supporting mesh is derived from these points on which a set of particle positions and orientations are seeded, again with an orientation normal to the surface. (c) Different positions in a chemosensory array (EMD-10160; see also below) are annotated in blue and orange. A set of particle poses from an STA experiment (blue) is used to derive the positions and orientations of the orange particles in a “subboxing” procedure. *STA*, subtomogram averaging.

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Providing access to alternative refinement and classification procedures

RELION implements a Bayesian approach to the refinement of one or many 3D volumes from cryo-EM data called maximum a posteriori refinement [22]. Originally designed for single-particle cryo-EM data and more recently extended to work with cryo-ET data [32], RELION provides solutions for the “refinement” block of the cryo-ET pipeline. The package features an easy-to-use graphical user interface (GUI) and is already in use in a large number of cryo-EM labs. Rather than a data processing pipeline, Dynamo is a collection of powerful tools for working with STA data. With an emphasis on customisation, Dynamo often exposes the user to large numbers of parameters in fully featured GUIs.

By interfacing Dynamo with the Warp-RELION-M pipeline in the “refinement” block, we provide Warp-RELION-M users access to highly customisable STA and multireference classification procedures, principal component analysis (PCA)-based classification, and myriad data visualisation tools and analysis tools. Dynamo users gain a route to making use of the CTF estimation and particle reconstruction tools in Warp, STA and classification in RELION, and the multi-particle refinement procedure implemented in M.

As an example of a possible benefit of increased flexibility in STA workflows, we compare the options for masking during iterative refinement in Dynamo and RELION. In a 3D classification experiment using relion_refine, one mask is used for alignment, classification, and Fourier shell correlation (FSC) for the determination of regularisation parameters between iterations. In a Dynamo multireference classification project, a user may choose to provide separate masks for these 3 procedures. As an example of the possible benefits, this allows for the optimisation of particle alignment parameters during a focused classification experiment. In addition, creative combination of masks can allow for exclusion of a membrane from alignment for a target that would otherwise require using masks with extremely soft edges to avoid FSC artefacts. Such a soft mask would necessarily include the membrane or exclude membrane-proximal protein density.

Application to the HIV-1 CA-SP1 hexamer (EMPIAR-10164)

We illustrate the power of working within this framework on an example dataset by reprocessing a 5 tilt-series subset of EMPIAR-10164 containing immature HIV-1 dMACANC virus-like particles (VLPs). This dataset was contributed to the community by the Briggs group, and this subset has previously been used to benchmark NovaCTF [17] and Warp [10], resulting in 3.9 Å and 3.8 Å reconstructions, respectively.

Initial motion correction and initial CTF estimation were performed in Warp with respective spatiotemporal model resolutions of (1, 1, 8) and (2, 2, 1). Tilt series were automatically aligned using autoalign_dynamo before CTF estimation and tomogram reconstruction at 10 Å/px in Warp. VLPs were annotated in Dynamo, and initial estimates of positions and orientations were generated with an interparticle distance of 45 Å, oversampled relative to the approximately 75 Å lattice spacing seen in the tomograms. A total of 500 particles were extracted and averaged in Dynamo to produce a coarse template. The same data were subject to STA against this template in Dynamo without imposing symmetry during refinement. The resulting average contained a hexagonal lattice. To obtain an initial model, the 6-fold axis of the lattice was aligned to the z-axis, and the resulting volume was symmetrised 6-fold in Dynamo. A total of 28,516 particles were extracted from annotated VLPs and aligned against the initial model for one iteration with a limited angular search range in Dynamo. The resulting particle positions and orientations, visualised in Dynamo, formed regular lattices on the surfaces of each VLP with some less regular areas. Using simple MATLAB scripts, only the 19,810 particles with 3 or more neighbours at the expected interparticle distance were retained for subsequent analysis.
Using *dynamo2m*, metadata were converted to enable working in *Warp-RELION-M*. Particle reconstructions were carried out in Warp and 3D refinements in *RELION 3.1*, starting with local angular searches of ±15°. Resolution estimates were measured by FSC between independent reconstructions from random half-sets of the data using *relion_postprocess*. Particles were (i) reconstructed at 5 Å/px and refined (unmasked) to 10 Å resolution; (ii) reconstructed at 2.5 Å/px and refined (unmasked) to 5 Å resolution; and (iii) reconstructed at 1.7 Å/px in *Warp* and refined (masked to include only the central hexamer) to 3.8 Å resolution. This intermediate result is indicative of the accuracy of fiducial-based alignment workflows in *Dynamo* and alignments in *RELION*. Seven rounds of multi-particle refinement were performed in *M* at 1.6 Å/px in a masked region containing only the central hexamer. The first 4 rounds of multi-particle refinement, optimising only for deformation parameters, yielded a 3.6-Å reconstruction. Three further rounds including optimisation of electron-optical aberration–related parameters and tilt frame alignments refined to a resolution of 3.4 Å (Fig 5).

It should be noted that working within this framework readily yielded a 3.4-Å reconstruction from 19,810 particles in 5 tomograms, without use of an external reference and leveraging the geometry of the system evidenced in the tomograms during alignments. The combination of *Dynamo* tools and the *Warp-RELION-M* pipeline, enabled by *autoalign_dynamo* and *dynamo2m*, rendered this ab initio, geometrical approach a smooth, efficient process. An ab initio approach is not strictly necessary for this dataset, as demonstrated by *emClarity* [7] and *Warp-RELION-M* [9]. We, however, note that obtaining an initial model is often a challenging step in the first stages of a project. Thus, access to ab initio and geometrical approaches is provided as an alternative, should the integrated template matching procedure prove inadequate. The final resolution of the reconstruction serves only to demonstrate the ability of *M* to further optimise image alignment and electron-optical parameters.

### Application to the *Escherichia coli* chemosensory array in situ

We are currently using an optimised minicell system [33] to investigate structural bases of the chemotactic response in motile bacteria. To illustrate the advantages of working within this framework, we choose to present intermediate results from a set of 109 minicell tomograms (Fig 6).

![Figure 5](https://doi.org/10.1371/journal.pbio.3001319.g005)
The chemotaxis-mediating molecular machinery forms large supramolecular arrays, a highly cooperative network of core signalling units (CSUs), in the bacterial inner membrane at the cell poles. The elongated nature of the CSU, the crowded cellular milieu, and increased sample thickness complicate template matching approaches with this system, increasing the false-positive rate and impeding appropriate peak extraction. Instead, we opt for a seed-over-sampling approach in which initial positions and orientations are distributed on Dynamo surface models, oversampling with respect to the expected interparticle distance of approximately 120 Å. Particle positions were generated from the surface model with an interparticle distance of 30 Å using Dynamo. Particles were extracted and subject to one round of STA in Dynamo with local angular searches of ±30˚. Constraining the angles of particles relative to initial estimates from a supporting geometry ensures that particles within each array have orientational parameters coherent with our understanding of the system. Particles were extracted at 7.5 Å/px in Warp and imported into Dynamo using warp2dynamo.

A consensus refinement of these particles, centred on the 3-fold symmetry axis of a p6-symmetric assembly, was performed in Dynamo, allowing local deviations (±30˚) in out-of-plane orientation, complete search of in-plane orientation, and enforcing C3 symmetry during refinement. In the resulting average, the expected 3-fold symmetric arrangement of P4 domains in the structure was not readily observed. The resulting reconstruction had a nominal resolution of 18 Å, measured by gold-standard FSC, with the FSC curve indicating the presence of significant heterogeneity. The Dynamo tool dpktbl.subbox.tableOnTable was used to derive the positions and orientations of particles centred on 6 intertrimer-of-dimer axes of the array. Duplicate particles were removed using dpktbl.exclusionPerVolume prior to a consensus refinement of these particles enforcing C2 symmetry. The resulting reconstruction had a nominal resolution of 19 Å. Attempts to disentangle structural heterogeneity using multireference approaches to 3D classification in a variety of masks using both Dynamo and RELION failed to yield sensible results at this stage.

Multi-particle refinement in M yielded improvements when performed in a masked region encompassing a large array region of 400 Å diameter, resulting in a reconstruction with a nominal resolution of 11 Å. No improvement was seen when refinement was performed on smaller regions encompassing 1 to 4 CSUs, presumably due to insufficient signal for accurate alignments. Work is underway to improve upon this intermediate result by classification and
further refinement. However, these initial results are already quite promising, showing features such as a hole in the center of the 4-helical bundle of a receptor dimer, not yet seen in situ.

**A comprehensive guide to obtaining a 3.4-Å reconstruction of the HIV-1 CA-SP1 hexamer**

Information about best practices and approaches to problems presented by cryo-ET data analysis is constantly evolving and currently divided between the literature, mailing lists, and documentation specific to various pieces of software. We provide, in the form of a rich, living document, a step-by-step walk-through for the procedure used to obtain the 3.4-Å reconstruction of the HIV-1 CA-SP1 hexamer (https://teamtomo.org/walkthroughs/EMPIAR-10164/introduction.html). This guide takes a user through downloading a 5 tilt series subset of EMPIAR-10164, preprocessing multiframe micrographs in Warp, aligning tilt series automatically using autoalign_dynamo, generating tomograms in Warp, setting up a Dynamo catalogue for geometrical particle picking using warp2catalogue, picking particles based on supporting geometries in Dynamo, initial model generation in Dynamo, isolating an optimal subset of particles using custom scripts, reconstruction of particles in Warp via dynamo2m, rigid body STA in RELION, and multi-particle refinement in M. Working through this guide is designed to expose a user to the theory and practice of obtaining a reconstruction from cryo-ET data. They will learn to use a variety of computational tools to address problems they will likely encounter when analysing their own data. The guide is provided as part of https://teamtomo.org/, an open collaborative platform we established for sharing knowledge in the growing cryo-ET community. Those wishing to share their expertise for the benefit of the wider community are invited to contribute.

**Conclusions**

The tools and resources presented in this manuscript increase automation within the Warp preprocessing pipeline, simplify metadata handling, and interface two complementary cryo-ET data processing packages. The interface provided by dynamo2m enables working within a powerful, flexible framework for geometrical and ab initio approaches to STA, which can benefit from multi-particle refinement in M. Our tools increase the functionality available to users of both Dynamo and Warp-RELION-M, lowering the barrier to employing more customisable workflows for cryo-ET data processing, which are often key to success. Finally, we have attempted to make structural cryo-ET more accessible by providing a comprehensive, step-by-step walk-through to obtaining state-of-the-art STA results. It is our hope that this online guide can serve as a starting point for newcomers to the growing cryo-ET community, enabling them to more quickly understand and use cryo-ET to solve their specific biological questions. Furthermore, we anticipate that the existence of a collaborative, community-driven resource for sharing cryo-ET knowledge and tools will encourage open science and contribute to the development of cryo-ET as a valuable tool for structural cell biology.

**Supporting information**

S1 Fig. The number of STA reconstructions with different reported resolution values in the EMDB. The data used to produce this figure can be found at https://www.ebi.ac.uk/pdbe/emdb/statistics_main.html. EMDB, Electron Microscopy Data Bank; STA, subtomogram averaging. (TIF)
S2 Fig. The source code for generating the scene depicted in Fig 2, combining starfile and eulerangles with existing tools in the scientific Python ecosystem.

(TIF)

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Author Contributions

Conceptualization: Alister Burt, Irina Gutsche.
Data curation: Alister Burt.
Funding acquisition: Irina Gutsche.
Investigation: Alister Burt, Lorenzo Gaifas, Tom Dendooven, Irina Gutsche.
Methodology: Alister Burt, Irina Gutsche.
Project administration: Irina Gutsche.
Resources: Irina Gutsche.
Software: Alister Burt.
Supervision: Irina Gutsche.
Validation: Alister Burt, Lorenzo Gaifas, Tom Dendooven, Irina Gutsche.
Visualization: Alister Burt.
Writing – original draft: Alister Burt, Irina Gutsche.
Writing – review & editing: Alister Burt, Lorenzo Gaifas, Irina Gutsche.

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