

Tomogram reconstruction with SCIPION: from raw movies to a tomogram

Version 2 (November 2023)

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Introduction

Overview

This tutorial aims to familiarize you with the initial steps of cryo-electron tomography (cryo-ET) analysis using Scipion (de la Rosa-Trevín et al., 2016) framework. We assume you are familiar with Scipion GUI and have some experience with cryo-ET data processing. Here we will demonstrate the workflow starting from raw data up to the tomogram reconstruction, highlighting the features of our software framework. After that, you should be able to run Scipion with your data.

Software requirements

To follow this tutorial you need to have SCIPION version 3.x already installed in your system. Also, make sure you have the following plugins and their corresponding binaries installed and configured:

- scipion-em-tomo
- scipion-em-motioncorr / motioncor2
- scipion-em-cistem / ctffind4
- scipion-em-imod / imod
- scipion-em-aretomo / aretomo
- scipion-em-novactf / novactf

Test data

We will use a small subset of HIV-1 virus-like particles collected on a 300 kV TFS Titan Krios microscope at EMBL, Heidelberg, Germany. To download raw data you can use Globus or follow the instructions below. Open EMPIAR-10164 link in your browser and scroll to **Browse All Files** section. You need to select checkboxes only for the following files for downloading:

```
mdoc-files/TS_01.mrc.mdoc
mdoc-files/TS_03.mrc.mdoc
frames/TS_01_*.mrc
frames/TS_03_*.mrc
```

Once downloaded, transfer these files to a folder called **empiar-10164**. Now it should contain movies for 2 complete tilt-series in MRC format, as well as 2 MDOC files with metadata.

1 Import of tilt-series movies

We will start by launching Scipion GUI:

> scipion3

In the project window click on Create Project button and enter a project name, keeping the default location. Press Create, so that a new project window appears.



Figure 1: Create project dialog.

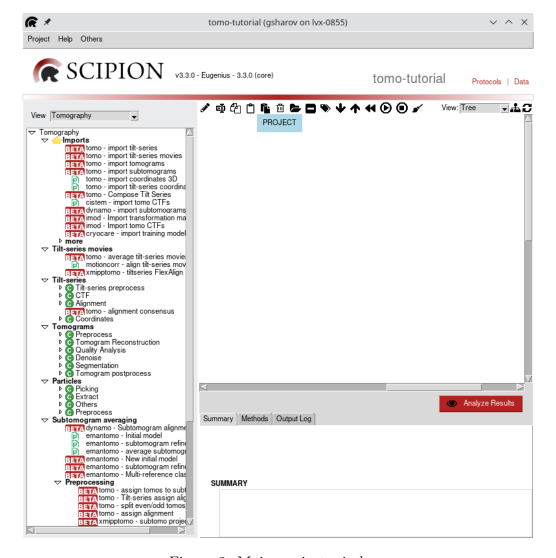


Figure 2: Main project window.

On the left panel select View Tomography and double-click on the import tilt-series movies protocol. On the *Import* tab fill in the parameters as listed below:

- Files directory: Path to empiar-10164 folder
- Pattern: *.mdoc
- Microscope voltage (kV): 300
- Pixel size (Å/px): 0.675. These movies are in the super-resolution format. The pixel size listed on the EMPIAR website is not correct.
- Dose per tilt image (e/Å²): 3.0

TIP

If you have a defects file from your detector, you will need to input it later, during motion correction. The same applies to the gain image orientation. In the import protocol, you can only input a gain reference.

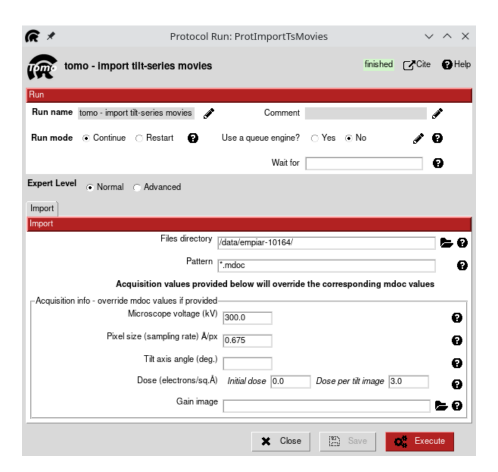


Figure 3: Import movies protocol.

This protocol (and also import tilt-series protocol) allows two import methods: via MDOC files or via filename pattern. For clarity, here we will demonstrate

both approaches. MDOC is a metadata format, designed for *IMOD* (Kremer et al., 1996) and used by SerialEM (Mastronarde, 2003) data collection software. TFS Tomography software v5.6 or newer can also produce an MDOC file for each tilt series. Using MDOC files is a recommended way of importing tilt-series data into SCIPION project because they contain all necessary information about both the microscope and data acquisition, including voltage, pixel size, acquisition order, tilt angles, and often dose rate.

If you open one of the downloaded MDOC files (e.g. $TS_03.mrc.mdoc$) in a text editor, you will see that it contains global metadata like pixel size, tilt axis angle, as well as information about each image of the tilt series (organized in sections with different Z-values). The values for dose rate and voltage are missing, that's why we had to input them manually above. Also, the pixel size in this MDOC file corresponds to a binned stack, not the raw movies. There are a few more important points to mention about MDOC files:

- For the import tilt-series movies protocol one MDOC file per tilt-series (per a group of movies that correspond to the same tilt-series) is expected. SerialEM can (if configured) produce MDOC files on a per movie basis, however, such files cannot be used for import at the moment
- For the import tilt-series protocol also one MDOC file per tilt-series is expected, each tilt-series must be in the *.mrcs stack format
- MDOC files and binary stacks/movies should be located in the same folder
- To distinguish different tilt-series Scipion will create a unique **TS_ID** based on each MDOC file basename. This key will be used to recognize the origin of all objects created from a certain tilt-series throughout Scipion workflow
- When SerialEM creates an MDOC file for each tilt series, it is associated with the <u>tilt-series stack that can have a different binning</u> compared to raw movie files. In that case, you need to specify the correct pixel size manually when importing movies into Scipion
- SerialEM has an option to sort the tilt-series stack by tilt angle immediately after the acquisition. This leads to a rearranged MDOC file. Nevertheless, SCIPION can parse *DateTime* field from MDOC to figure out the original acquisition order
- Often (e.g. due to a missing calibration), the information about dose rate is absent from MDOC file, in such case, you have to specify it manually
- SerialEM MDOC files do not contain information about spherical aberration (Cs). The default value in SCIPION import protocols is set to **2.7 mm**, which is

true for TFS Talos and Titan microscopes with C-Twin objective lens. Consult your microscope documentation to find out the correct value. If your data was acquired on a Cs-corrected microscope, input **0.001 mm**.

After checking that all input parameters are correct, press the Execute button. The protocol should complete almost immediately as it only creates symbolic links to your raw data and registers metadata in the database. The **Summary** tab should now show that 2 tilt-series have been imported, 41 movies each, 8 frames per each movie, with the pixel size of 0.68 Å/px.

TIP

If you do not have the same number of images/movies for every tilt-series, SCIPION summary will display the number for the first imported series. Don't worry, the information is stored correctly in the database.

Now, just to practice, we will demonstrate how to import the same tilt-series movies without any MDOC files using a filename pattern. Right-click on the completed protocol and select Duplicate. Change the input parameters as listed below and execute the protocol:

• Files directory: Path to empiar-10164 folder

• Pattern: {TS}_{TO}_{TA}.mrc

• Microscope voltage (kV): 300

• Pixel size (Å/px): 0.675

• Tilt axis angle (deg): 85.3

• Dose per tilt image (e/ $Å^2$): 3.0

As you can see, we have to input more metadata manually in this case. Moreover, the file pattern becomes more complex. It consists of three parts: TS to identify individual tilt-series, TO to identify acquisition order and TA to identify tilt angle from the filename. In our case, if you look for example at file TS_01_002_-3.0.mrc,

- $TS = TS_01$
- TO = 002
- TA = -3.0

The pattern can become more complex depending on the settings of your data collection software. If you need more information about file patterns, click on ? next to the *Pattern* parameter on the *Input* protocol tab.

Now click on Analyse Results for either import protocol and in the opened dialog verify the acquisition order, tilt angles, and accumulated dose for the imported tilt-series movies. In our case, the data was collected using a dose-symmetric scheme (+/- 60 deg. starting from 0 deg.) with a 3-degree step.

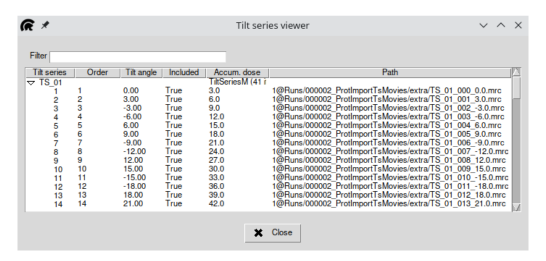


Figure 4: Tilt-series viewer. Movies are imported and sorted by <u>acquisition order</u> (second column).

2 Beam-induced motion correction

Aligning individual movie frames is necessary to correct beam-induced specimen motion and restore high-resolution information. In this practical, we will use *motion-cor2* (Zheng et al., 2017) for movie alignment. To find the corresponding protocol you can either expand the protocols tree on the left panel of the project (check the Tilt-series movies list) or use ctrl+F to search for motioncorr - align tilt-series movies. Select the movie set that you have just imported as input for this protocol. Set Binning factor: 2.0 because we have imported super-resolution movies. We will not change other default options, but you are free to explore them on your own. If you have two GPUs available on your machine, set GPU IDs to "0 1" and number of threads to 3 - this way the protocol will run on every two movies in parallel, each on a separate GPU. The third thread is required to coordinate the jobs.

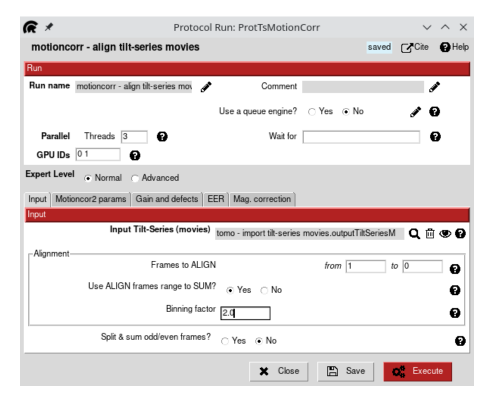


Figure 5: Movie alignment protocol for motioncor2.

After launching the protocol, we can go further and start the CTF estimation. In this case, thanks to the stream processing capability of SCIPION, we don't need to wait until the previous protocol finishes to start the next one.

ALTERNATIVES

- tomo average tilt-series movies
- xmipptomo tiltseries FlexAlign

If you would like to first check the results of the movie alignment, wait until the first output tilt-series is produced and click on Analyze Results to open the viewer. Here you have only one option to display the output. In the opened viewer verify that the output tilt-series stacks are now sorted by tilt angle. You can also right-click on the protocol output and choose *Open with ImodViewer*. Double-clicking on a tilt-series name will launch the 3dmod interface to visualize the stack file.

TIP

At the moment, Scipion does not store movie frame shifts from motion correction protocols for tomography.

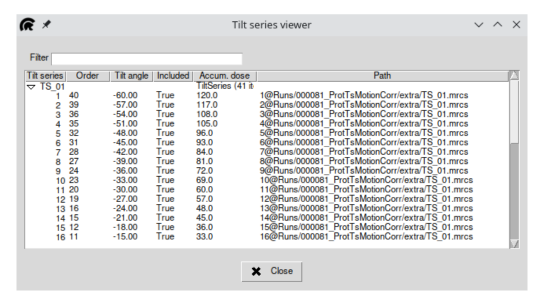


Figure 6: Tomo data viewer. Motion-corrected tilt-series are now sorted by <u>tilt angle</u> (third column) and assembled into individual *.mrcs stacks.

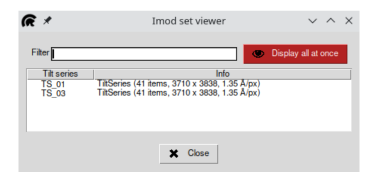


Figure 7: IMOD's tilt-series viewer. Double-click on any item to launch the 3dmod interface.

If you were to use **Split and sum odd/even frames** option of the motioncorr protocol, you would get two more output sets. This option is commonly used to later generate two identical tomograms to run denoising software (e.g. *cryoCARE*).

TIP

Dose-weighting option in motion corr protocol is disabled by default. In principle, dose-weighting can be performed at many stages of the tomo pipeline. Here we will do it after CTF estimation.

3 CTF estimation

We will now estimate the CTF parameters of the aligned images with CTFFind4 (Rohou and Grigorieff, 2015). Locate cistem - tilt-series ctffind4 protocol and select tilt-

series from the previous step as the input. Set the number of threads equal to the number of available CPU cores. Leave all options with default values. CTFFind4 will run on individual tilt-series *.mrcs stacks (that were assembled by motion correction protocol) which speeds up the execution.

TIP

Make sure you always run CTF estimation on the raw data, before any doseweighting, filtering etc.

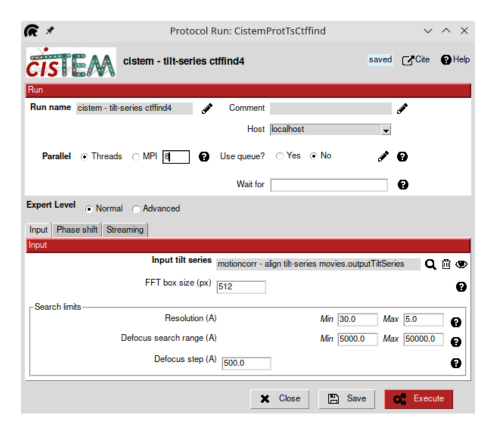


Figure 8: CTFFind4 protocol.

ALTERNATIVES

- emantomo CTF estimation
- gctf tilt-series gctf
- imod CTF estimation auto/manual
- susantomo CTF estimation

The output from CTFFind4 (or any other CTF estimation protocol) can be displayed upon clicking on the Analyze Results button (Figure 9). SCIPION CTF viewer for tomography is split into two panels. Left mouse click on a tilt-series row on the left-hand side will update the corresponding plot on the right. By default,

the plot will show defocus, estimated resolution (if available), and phase shift (if estimated) as a function of tilt angle for selected tilt-series. If you click on the arrow symbol near the tilt-series name, the list will expand with all tilt images and show the table with estimated CTF parameters (defocus, astigmatism, etc.) for each image. You can also left-click on any tilt image and the right-hand side plot will be updated with a 2D CTF fit (Figure 10). Moreover, selecting an individual tilt image and clicking the 1D fit button at the top of the viewer will open a 1D CTF fit profile.

TIP

2D and 1D fit CTF profiles are only available for CTFFind4 and gCTF.

If you want to discard some tilt-series you may tick the checkbox next to all tilt-series that you want to keep and click Generate subsets. This will create "good" and "bad" subsets with the corresponding items.

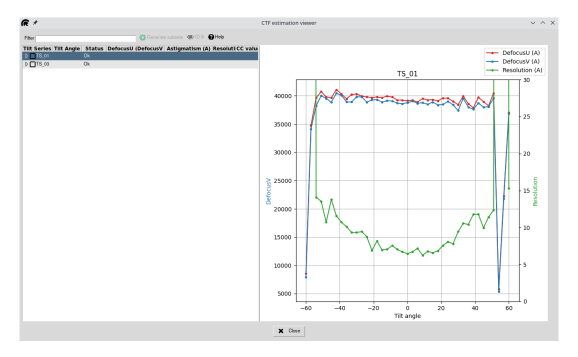


Figure 9: Visualization of CTF estimation results for each tilt-series. We can see that the defocus is consistent throughout the tilt-series, except for the high tilts where estimation has failed.

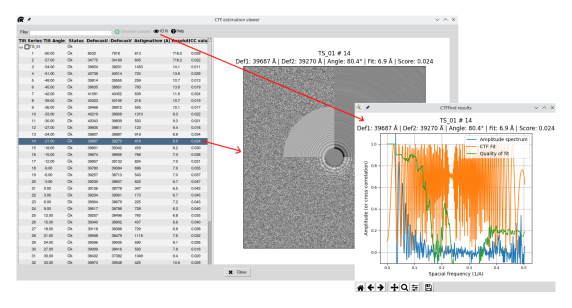


Figure 10: Individual tilt image CTF fit plots (2D and 1D) from CTFFind4.

4 Dose-weighting

As explained in (Metskas et al., 2022), dose-weighting can be done at many stages in the tomogram reconstruction pipeline, but for best effect should be done before tilt stack alignment and after CTF estimation. While the primary purpose of dose-weighting is to limit the contributions of Fourier space containing the most electron damage (Grant and Grigorieff, 2015), when combined with a dose-symmetric tilt scheme it effectively functions as a low-pass filter in the noise-ridden high tilts. This improves the overall alignment for these tilts, along with a subtle improvement in the signal-to-noise ratio in the reconstructed tomogram.

Now, locate [imod - dose filter] protocol and select tilt-series from the motion correction step as the input. Ignore other options and execute the protocol. Check the results with 3dmod viewer. The tilt-series should now have more contrast, especially at high tilts. Note that both TS_01 and TS_03 have a completely dark image at -60 degrees (section 1). We will take this into account in the following chapter.

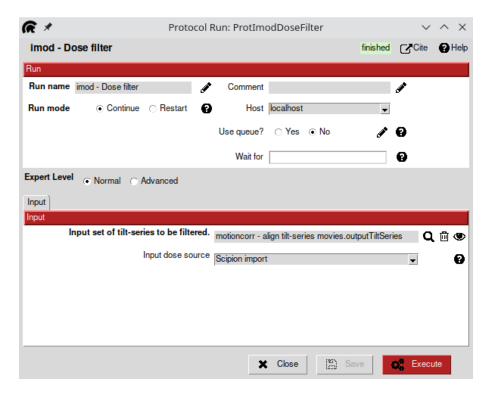


Figure 11: Dose-weighting protocol from IMOD.

5 Tilt-series alignment and reconstruction

In this chapter, we describe a few common workflows that can be used to align tilt-series and reconstruct a tomogram. Users can choose which route to take depending on their experience and data complexity. The first section goes through a manual tilt-series processing via eTomo. Here a user is presented with full capabilities of the original IMOD's graphical interface where almost any parameter can be tweaked. In this case SCIPION will simply collect and register the results. The second section describes a semi-automated approach, where the individual steps from eTomo interface have been wrapped into separate SCIPION protocols. Finally, the last section shows the power of AreTomo software that does not require fiducial markers and can align and reconstruct tomograms in a fully automated fashion. This tutorial dataset can be used with either of these three workflows, so you may choose any of them to follow.

5.1 Manual processing with eTomo GUI

At this point, we assume that you already have some experience with eTomo (Mastronarde and Held, 2017) as we will not provide a complete guide for this software, but rather illustrate how to run eTomo interactive GUI from SCIPION and how to save the results into the project. For a more comprehensive description of every step, please refer to the official eTomo tutorials [1] and [2].

Find the [imod - etomo interactive] protocol and input the dose-filtered tilt-series set. Also, input fiducial marker size of 10 nm. Set the number of threads to 4. Pixel size, tilt angles, and tilt axis position will be fetched by SCIPION automatically. After executing the protocol you will see a different tilt-series dialog showing you the steps you will need to perform within eTomo interface. Double-click on the first item in the list to launch eTomo GUI.

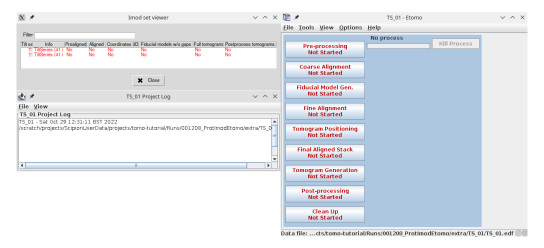


Figure 12: Tilt-series viewer of eTomo protocol (left) and eTomo interface (right).

Pre-processing

The purpose of this first step is to remove very bright or dark pixels caused by detector defects or X-rays. Such pixels can cause artifacts in a reconstructed volume so it's important to always remove them.

- 1. Click on [Pre-processing] button to open the panel for removing pixel artifacts.
- 2. Press Create Fixed Stack button.
- 3. Once completed, press on Show Min/Max for Fixed Stack and check if the min/max pixel densities distribution is uniform and there are no outliers. Deviations of 50-100 from the rest of the data will not matter in the reconstruction.
- 4. To proceed, press on Use Fixed Stack and then Done.

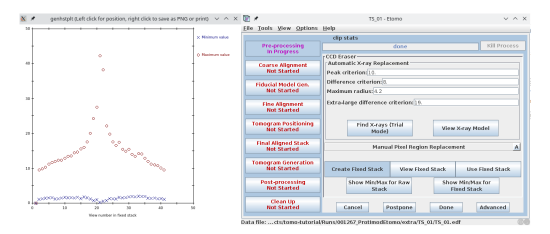


Figure 13: Removing pixel artifacts from a tilt-series stack. The plot shows minimum and maximum densities for the fixed stack. View #1 has a minimum intensity of 0 (black image), we will get rid of it in the next steps.

Coarse alignment

The goal of coarse alignment is to calculate the cross-correlation between successive tilts and apply X and Y-shifts to the tilt-series.

- 1. Press the Advanced button to display more options.
- 2. At the end of the Tiltxcorr section enter **Views to skip: 1** to ignore the first dark image of the tilt-series.
- 3. Set Coarse aligned image stack binning to 4.
- 4. Check Reduce size with antialiasing filter checkbox.
- 5. Uncheck **Convert to bytes**, otherwise grayscale depth will be degraded to 8 bits.
- 6. Check Float intensities to mean.
- 7. Press Calculate Cross-Correlation followed by Generate Coarse Aligned stack. Tilt-series images are now aligned translationally.
- 8. Check the resulting aligned stack by pressing View Aligned Stack in 3dmod. Press Done.

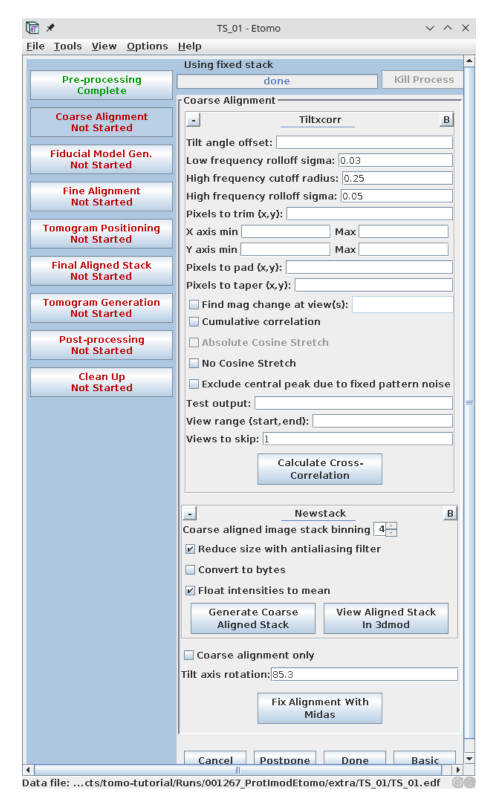


Figure 14: Coarse alignment of the tilt-series stack. We omit the first image (view) from the alignment.

Fiducial model generation

Since our data contains gold beads we will use them as fiducial markers to perform a more precise tilt-series alignment. If your data does not have any fiducials, you may try patch tracking (not described here) or use AreTomo, explained in the later section.

- 1. Press Advanced to display more parameters.
- 2. On the Seed model tab in the Beadtracker section enter View skip list: 1, check Refine center with Sobel filter, input Sobel sigma relative to bead size: 0.12 and Overall low-pass filter cutoff: 0.3 (1/nm)
- 3. In the Initial Bead Finding parameters section check **Find and adjust bead** size
- 4. In the Selection and Sorting parameters section input **Total number: 30** (the approx. number of beads in the tomogram).
- 5. Press Generate Seed Model
- 6. Switch to the *Track Beads* tab and press Track Seed Model. In the *eTomo* log window check the total number of missing points. If a large number of points are missing, press Track with Fiducial Model as Seed.

TIP

You need at least 5 beads sparsely distributed in the field of view for the tracking to work correctly.

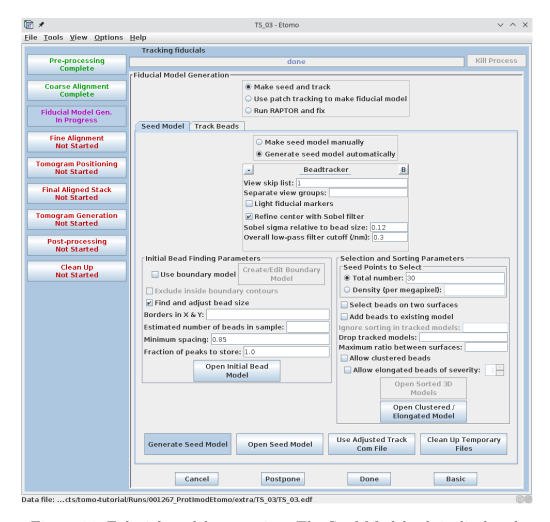


Figure 15: Fiducial model generation. The Seed Model tab is displayed.

If points are still missing, the next procedure involves an iterative process to edit this fiducial model. Otherwise, just press Done to finish this step.

- 1. It is important to make sure that the large majority of the fiducials are tracked to the two ends of the tilt-series. Press Fix fiducial model on the *Track Beads* tab, this will open the pre-aligned stack with model points overlayed along with the Bead Fixer window.
- 2. Click the Go to Next Gap button in the Bead Fixer window. This will highlight a point with a yellow circle that has a missing model point on an adjacent section (indicated by an arrow).
- 3. Use the Page Up (when an "up arrow" appears) or the Page Down (see Figure 16) to switch to the image with a missing point and use the middle mouse button to add a point in the center of the gold particle (you can use right click to adjust the current point position). It is useful to increase the magnification of the image with the + key and adjust the contrast of the images, especially at high tilt.

- 4. Keep going to the next gap and fixing missing points until the message *No more gaps found* comes up in the main 3dmod window.
- 5. Save the model file using File Save model in the main 3dmod window.
- 6. Press Track with Fiducial Model as Seed again. There should be no more missing points.
- 7. Press Done

TIP

Note that it is not necessary to fill all the gaps, particularly if there is a good excess of fiducial points. Most importantly, you should not add a point if the bead's position is not clear or it disappears from the field of view!

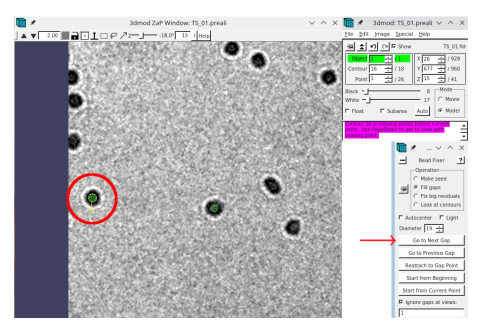


Figure 16: Filling the model gaps. Pressing Go to Next Gap (indicated by a red arrow) will highlight a bead (indicated here by a red circle) that has a missing point on the neighboring section.

Fine alignment

The goal of this step is to solve for the displacements, rotations, tilts, and magnification differences in the tilted views using the position of the gold fiducials. Usually one would aim to reduce the alignment residual error mean to 0.2 - 0.5 nm.

1. On the *General* tab set List of views to exclude: 1 and Threshold for residual report to 3.0.

- 2. Select Do not sort fiducials into two sufraces
- 3. Check Do robust fitting with tuning factor: 1.0.
- 4. On the *Global Variables* tab set **One rotation**, **Fixed magnification at 1.0**, **Fixed tilt angles**.
- 5. Press Compute Alignment. The residual error is now reported in the Log window.
- 6. If the error is above 0.5 nm you need to optimize the alignment by removing the beads with high stdev and/or fixing bead centering manually. Depending on your microscope stage you might also adjust global variables. This is not necessary for this tutorial data, otherwise, consult eTomo guide on fixing big residuals.
- 7. Press [Done] when finished.

TIP

When playing with different parameters during final alignment always check that the residual error mean drops more significantly than the ratio of total measured values to all unknowns, otherwise, you are just overfitting the model!

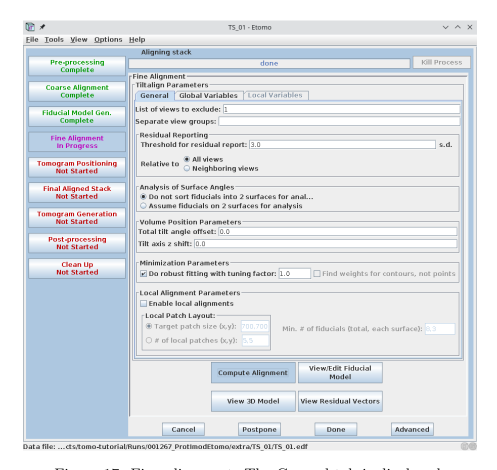


Figure 17: Fine alignment. The General tab is displayed.

Tomogram positioning

The purpose of this step is to set angles and an offset in Z so that the specimen is flat and centered in the computed volume. If you plan to do subtomogram averaging it is better to skip this step (just press Done). Note that if you correct the X-axis tilt at this point, the orientation of the missing wedge will be changed.

Final aligned stack generation

- 1. Uncheck Use linear interpolation.
- 2. Set **Aligned image stack binning to 4**. This will define the final tomogram binning (!)
- 3. Check Reduce size with antialiasing filter.
- 4. Press Create Full Aligned Stack followed by View Full Aligned Stack. Note that the stack has now been rotated 90 degrees so that the tilt axis becomes vertical. This is the default convention used by IMOD software. Check that the fiducials move horizontally on straight lines with no jumps when scrolling through the stack. Often it helps to draw a rectangle (use Shift + B) encompassing a few beads on its border, loop through Z, and check that beads do not jump vertically. If they do either improve the fine alignment or exclude such bad views.
- 5. Press Done when finished.

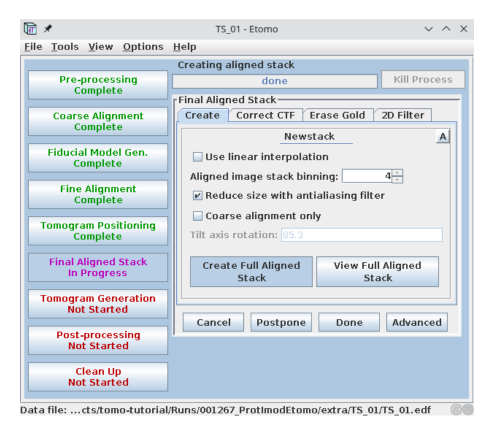


Figure 18: Making a final aligned stack. The Create tab is displayed.

TIP

If you plan to proceed later with subtomogram averaging, you would usually generate a bin 1 tomogram with the WBP reconstruction method (see below) and a binned SIRT-like filtered tomogram with high contrast for particle picking. Remember, the Final Aligned stack step is used to set the desired binning of the tomogram.

Tomogram generation

All views excluded at the **Fine Alignment** step will be automatically omitted from the final reconstruction.

- 1. Select **Back projection** method.
- 2. Check **Parallel processing** and set 4 CPUs to use in the parallel section above.
- 3. Uncheck Take logarithm of densities
- 4. Set Tomogram thickness in Z to 2000. This is in unbinned pixels (!)
- 5. Set Use SIRT-like filter equivalent to 8 iterations.

- 6. Press Generate Tomogram and wait until it's completed. Then press View Tomogram In 3dmod and open Slicer (press \(\)). In the new Slicer window set **X** rotation to 90 and **Img to 30** to increase the slicing thickness. This will display the x-z slice. Scroll through the tomogram using **View axis position** slider. Verify that (a) the thickness is correct and (b) the specimen is positioned in the middle of the slice, otherwise change **Z-shift** value (use positive unbinned pixels value to shift it upwards) and recalculate the tomogram. This step is equivalent to the *Tomogram positioning*. In the case of TS_01, you need to recalculate the tomogram using 1360 px final thickness and 300 px Z-shift.
- 7. Press Done when finished.

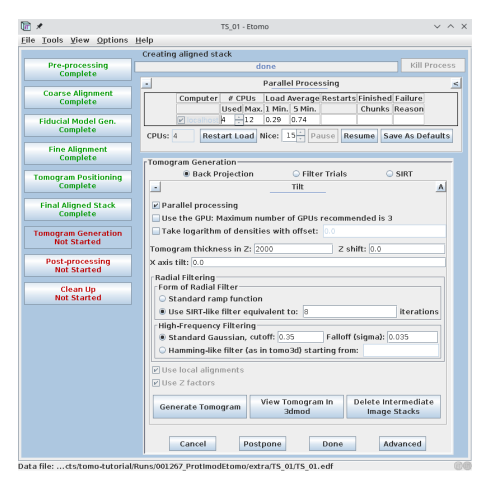


Figure 19: Creating a bin 4 tomogram with the SIRT-like filter.

Post-processing

- 1. Uncheck Convert to bytes.
- 2. Select **Rotate around X axis**. This will orient the tilt axis vertically while preserving the handedness.
- 3. Press Trim Volume. Display the trimmed volume if you like.

4. Press Done when finished.

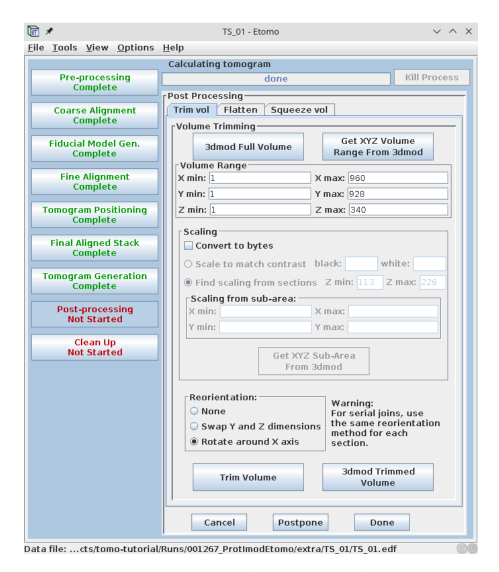


Figure 20: Post-processing the reconstructed tomogram.

Clean-up

Since deleting some intermediate files might make it impossible for Scipion to generate some of the outputs, we will skip this step. Just press $\boxed{\mathsf{Done}}$ and close eTomo window.

If you now look at the Scipion viewer dialog that stayed open while you used eTomo, you will notice that all steps for the first tilt-series have been completed and the first row has become green.



Figure 21: The Scipion tilt-series viewer registers all steps performed with eTomo GUI.

EXERCISE

Repeat the eTomo workflow for the second tilt-series. Don't forget to exclude the dark view #1 during the processing.

Now, close both eTomo GUI and the tilt-series dialog. SCIPION will now register the following outputs for each processed tilt-series:

- Prealigned tilt-series (coarse aligned stack)
- Aligned tilt-series (final aligned and rotated stack)
- Set of 3D coordinates of the fiducials
- Set of landmark models that include fiducial positions and residuals
- Raw tomogram
- Post-processed (rotated) tomogram

All these outputs may be now visualized with the installed viewers and/or connected to other Scipion protocols continuing the workflow.

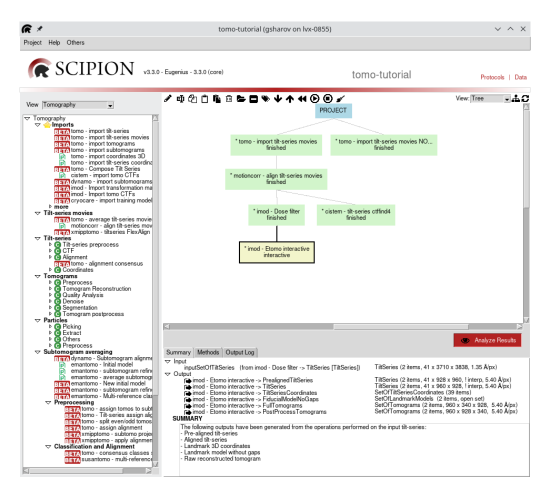


Figure 22: Outputs from *eTomo* interactive protocol.

5.2 Semi-automated processing with IMOD protocols

In this section, we describe an alternative workflow that can replace eTomo interface with a group of separate Scipion protocols. Note that not all functions available from eTomo GUI have been wrapped into individual protocols.

- 1. Find and execute [imod X-rays eraser] protocol on the dose-weighted tilt-series.
- 2. If you now display both output tilt-series with 3dmod, you may notice that the first tilt image (-60 deg.) is completely dark. Thus we need to exclude it from further processing. Find imod exclude views protocol and input the tilt-series from the previous step. Select Expert level: Advanced. Now, create a text file containing two lines:

TS_01 1

TS_03 1

The first column is tilt-series ID, while the second contains a list of views to be removed. Save and input this file as **Exclude views file** into the protocol and then execute. The output tilt-series should now contain 40 images each.

- 3. Next, let's downsample our tilt-series to speed up the processing. Find the imod tilt-series preprocess protocol and input the tilt-series from the previous step. Use a binning of 4.0.
- 4. Afterwards, run imod Xcorr prealignment protocol using the previous output. Set Generate interpolated tilt-series? Yes and input Binning of 1. This protocol will generate two output tilt-series: non-interpolated (with assigned alignment) and interpolated (aligned). You can display the interpolated output with 3dmod viewer to check if the alignment was successful.
- 5. Next, run the imod generate fiducial model protocol using the non-interpolated tilt-series from the step above, with options Find on two surfaces? No and Fiducial radius: 10 nm. Select Expert level: Advanced and also input the Number of fiducials: 30. After execution, the protocol will create a SetOfLandmarkModels which corresponds to a model containing fiducial positions and their residuals. You can display the output fiducial model with 3dmod.
- 6. Continue the workflow with the imod fiducial alignment protocol, using the set of landmark models from the previous step as input. On the *Input* tab also set Find on two surfaces? No, Generate interpolated tilt-series? Yes and Binning of 1. On the *Global variables* tab choose Solve for one rotation, leaving other parameters fixed. SCIPION will produce several outputs here, but you might want to check only the interpolated (aligned) tilt-series.
- 7. Finally, locate the imod tomo reconstruction protocol. Input a set of non-interpolated tilt-series from the step above and set **Tomogram thickness to 350 px** and **Z-shift to 75 px**. Select Expert level: Advanced and set **Iterations of a SIRT-like equivalent filter: 8**. If you have gone through the *eTomo* section above, you know that the optimal tomogram thickness for this data is about 1400 px. Here we use 1400/4 = 350 because we binned our tilt-series by a factor of 4. If you do not have any GPUs, set GPU IDs to **No**.

In the end, your workflow should look something like this:

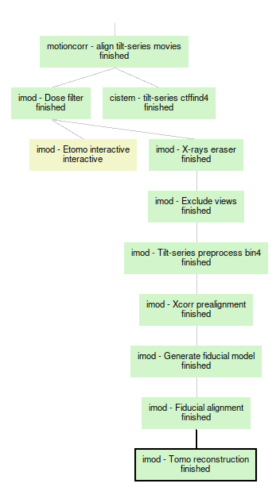


Figure 23: Project workflow for semi-automated processing with *IMOD*.

5.3 Automated processing with AreTomo

Are Tomo (Zheng et al., 2022) offers an automated marker-free tilt-series alignment and reconstruction. The program is accelerated on GPU and can produce aligned tilt-series and/or a reconstructed volume. Users can choose either weighted back projection (WBP) or simultaneous algebraic reconstruction technique (SART) to reconstruct their tomograms. Are Tomo implements both global and local alignments. The global alignment determines the tilt angle offset, translations of the tilt images, and orientations of the tilt axis as it varies throughout the tilt series. Local alignment can be used to correct local motions due to the progressive sample deformation under repeated beam exposures. Moreover, the software can apply a dose-weighting filter to aligned tilt-series or reconstructed tomograms.

Find aretomo - tilt-series align and reconstruct protocol and fill in the parameters listed below:

- Input set of tilt-series: provide the set of dose-filtered tilt-series
- Reconstruct tomogram? Yes

- Binning: 8
- Volume height (voxels): 1800. This is in unbinned voxels (!)
- Tomogram thickness (voxels): 2000. This is in unbinned voxels (!)

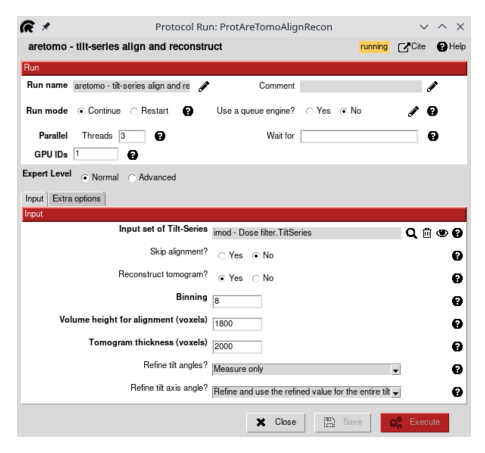


Figure 24: AreTomo protocol is used to align and reconstruct dose-filtered tilt-series.

Do not change other options and execute the protocol. At this point, we intentionally chose a large thickness value and high binning factor to estimate the correct tomogram thickness. Upon completion, the protocol produced two outputs: a non-interpolated tilt-series (the original ones with alignment metadata) and a set of tomograms. Right-click on the *outputSetOfTomograms* output and choose *Open with ImodViewer*. Double-click on the first tomogram (TS_01) to launch the *3dmod* interface.

To check the tomogram thickness you need to run 3dmod's Slicer: press key or choose Slicer in the 3dmod menu. In the new Slicer window set X rotation to 90 and Img to 30 to increase the slicing thickness. This will display the x-z slice. Scroll through the tomogram using View axis position slider. The current binned tomogram thickness is 2000/8=250 px. Draw a rectangle (use Shift + B) encompassing the sample and all beads to estimate the specimen thickness. The rectangle dimensions will be displayed instead of the Lo/Hi buttons in the Slicer

window when drawing. Remember that the pixels are 8x binned! Our estimate is 140 px * 8 = 1120 px.

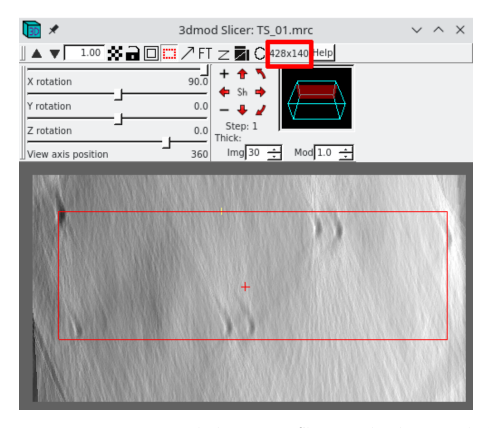


Figure 25: Estimating tomogram thickness using Slicer in 3dmod. Rectangle dimensions are indicated with a red box.

Now let's re-run the protocol using the newly estimated thickness value and then verify tilt-series alignment. Right-click on the completed protocol box and select **Edit**. This time input **binning 4**, volume height of **1120 px**, and tomogram thickness of **1300 px**. Once the run is finished, double-check the output tomogram again with *3dmod*.

To verify the tilt-series alignment we can use the non-interpolated tilt-series output (bin 1) that has alignment metadata. Find imod - apply transformation protocol and input the set of tilt-series from the *AreTomo* protocol. Set **Binning to 4** and press Execute.

TIP

Are Tomo protocol can generate aligned interpolated tilt-series when you choose not to reconstruct a tomogram. This can also be used to quickly check the accuracy of alignment. You might notice that interpolated tilt-series have fewer tilt images than the input, this is because Are Tomo automatically finds and discards dark tilt images. The parameter that controls this behavior (Dark tolerance) can be found on the Extra options tab of the protocol.

Once the *IMOD*'s protocol run is completed, let's check the tilt-series alignment. Right-click on the *InterpolatedTiltSeries* output and choose *Open with ImodViewer*. Double-click on the first item (TS_01) to launch the *3dmod* interface. Middle-click on the image (ZaP) window to scroll through the aligned tilt-series. The aligned stack is displayed as an x-y view with the vertical tilt axis. If the alignment has been successful, the gold beads should only move horizontally (perpendicular to the tilt axis) without any vertical jumps. You might also notice that the first dark tilt image has been removed from these tilt-series since it was detected by *AreTomo* and marked as disabled in the non-interpolated set.

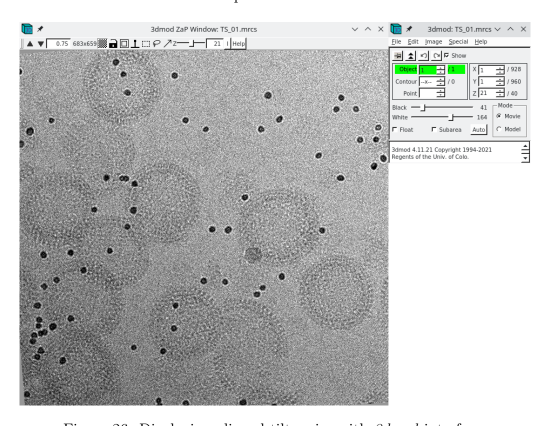


Figure 26: Displaying aligned tilt-series with 3dmod interface.

ALTERNATIVES

- emantomo align tilt series
- emantomo tomo reconstruction
- imod tomogram reconstruction
- tomo3d tomo3D

6 CTF correction

CTF correction in cryo-ET is an active research area. Conventionally, CTF estimation and correction are performed on tilt-series before tomogram reconstruction.

In the **2D** strip-based approaches, implemented e.g. in *IMOD*, each tilt image is divided into strips parallel to the tilt axis. Defocus for the central strip is estimated by averaging over overlapping tiles and then extrapolating to the other strips using a linear relationship depending on the distance from the tilt axis and the tilt angle. Each strip is then CTF-corrected by phase flipping. A corrected aligned stack of tilt-series is then used for tomogram reconstruction. This method only considers the defocus gradient caused by tilting the sample.

3D CTF-correction approaches additionally consider the sample thickness, which creates another defocus gradient along the direction of the electron beam. In the case of *NovaCTF* (Turoňová et al., 2017) during the reconstruction of a tomogram by back-projection, each voxel is calculated from tilt images that were CTF-corrected with defocus values corresponding to the position (given by x and z coordinates) of that voxel at each tilt. To achieve this, each image in the tilt-series is CTF-corrected multiple times with different defocus values. The number of different CTF corrections performed per image depends upon how finely the defocus gradient should be sampled and is a user-defined parameter.

CTF correction can also be performed at **subtomograms level**, taking into account their known 3D positions inside a tomogram volume. Such methods have been implemented in *Relion* and *EMAN2*. However, this imposes additional challenges of dealing with the missing wedge, extra interpolations, etc. *Relion* 4.0 has now implemented a different concept of pseudo-subtomograms (Zivanov et al., 2022) comprised of 3D arrays of CTF pre-multiplied 2D tilt-series images and other auxiliary data.

More recently, several high-resolution subtomogram averaging pipelines have been derived using **per-particle per-tilt averaging** algorithms (e.g. EMAN2, emClarity or M). In the case of EMAN2, the tomograms are only used for particle picking and once the coordinates are known, subtilt series are extracted from the original tilt series, one for each particle. These subtilt series come with CTF information based on where each particle is located in a 3D volume, different for each tilt in the subtilt series. This information is used for CTF correction during subtomogram averaging which essentially becomes subtilt averaging.

It's important to note that all CTF correction methods mentioned above depend on the accuracy of the CTF estimation. In many cases CTFFind4 or gCTF are used, that were originally designed for single-particle analysis. The low signal-to-noise ratio of tomographic tilt images makes CTF estimation less robust and precise than with single particle data, as the power spectra of higher tilt images show fewer Thon rings for fitting. An error in the defocus estimation of 250 nm limits the resolution to 10 Å, and an error of 63 nm limits the resolution to 5 Å (Kudryashev, 2017).

6.1 2D CTF correction with IMOD

In this tutorial, we will explore both 2D and 3D CTF correction methods. Let's start with the first one. If we look at the *CTFFind4* estimation that we executed at the beginning of the analysis, we can notice two issues: a) high tilts have wrong defocus estimates, and b) we did not remove the first dark image of each tilt-series before running the estimation. In principle, one could use imod - exclude views protocol to remove bad images and then re-run CTF estimation. Let's do this and also explore *IMOD* protocol for CTF estimation.

- 1. Find and execute imod exclude views protocol. Input motion-corrected tilt-series from motioncor2 and the text file with excluded view numbers that we have already used before.
- 2. Create a text file containing two lines as below. We will use this as defoci list for the two tilt-series:

```
TS_01 4000.0
TS_03 1500.0
```

3. Locate imod - Automatic CTF estimation protocol and change its input parameters:
a) input motion-corrected tilt-series with dark views excluded, b) Expected defocus file: the file created above, c) autorefinement Angle step: 0, d) Search astigmatism? Yes. Using a value of zero for the angular step will force IMOD to fit CTF to each single image separately.

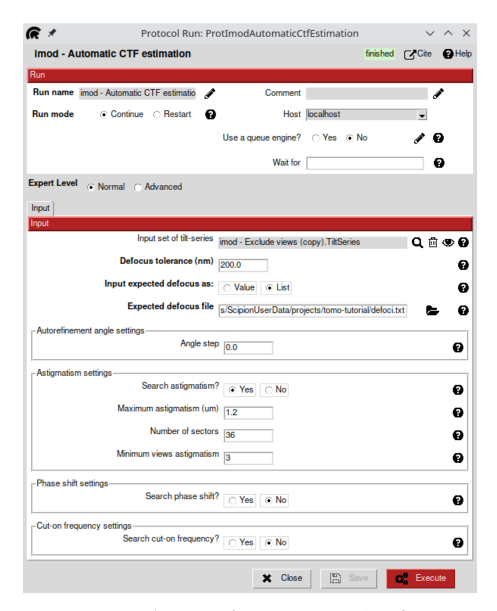


Figure 27: Automatic CTF estimation with IMOD.

- 4. If we now analyze the results of this CTF estimation, we can notice that hightilt views no longer deviate from the rest of tilt-series. We will now continue with these values, though you could run manual CTF estimation with *IMOD* instead to verify the 2D CTF fit for each image.
- 5. Next, we will execute imod CTF correction protocol using the CTF estimation above and non-interpolated tilt-series (bin 4) from imod fiducial alignment protocol. Non-interpolated tilt-series have alignment information and the estimated astigmatism will be rotated to match the tilt axis alignment automatically by *IMOD* protocol. We also ensured that the excluded views were the same in our workflow so that the tilt-series could be matched with the CTF estimation.

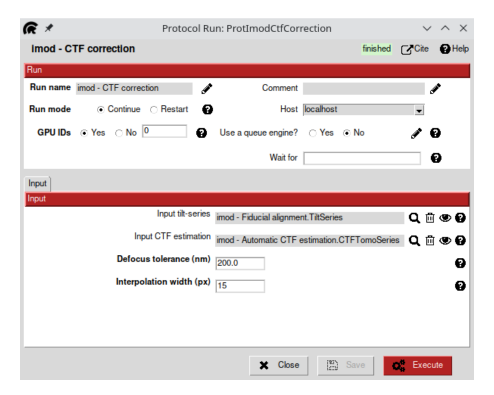


Figure 28: 2D strip-based CTF correction in IMOD.

6. Finally, CTF corrected and aligned tilt-series can be reconstructed with imod - tomo reconstruction protocol. In the end, your project workflow should look something like below.

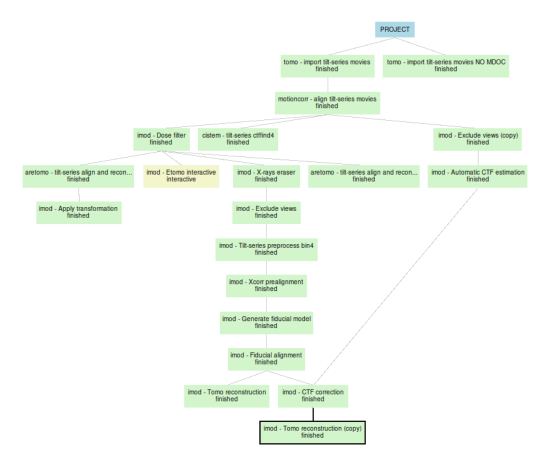


Figure 29: Project workflow after CTF correction.

6.2 3D CTF correction with NovaCTF

3D CTF correction workflow with *NovaCTF* consists of two SCIPION protocols: compute defocus array and 3D CTF correction and reconstruction. Find the first one and input there the non-interpolated tilt-series (bin 4) from [imod - fiducial alignment] protocol. Also, input CTF estimation from the last *IMOD* run. For tomogram thickness use **350** px with Z-shift of **75** px. Instead of the default recommended defocus step of 15 nm you can use **30** nm to speed up processing. We will use phase flipping as a CTF correction method, though multiplication can be used as well, albeit at the cost of reducing the tomogram <u>visual</u> quality. The protocol should finish very quickly as it only generates multiple defocus files used for CTF correction later.

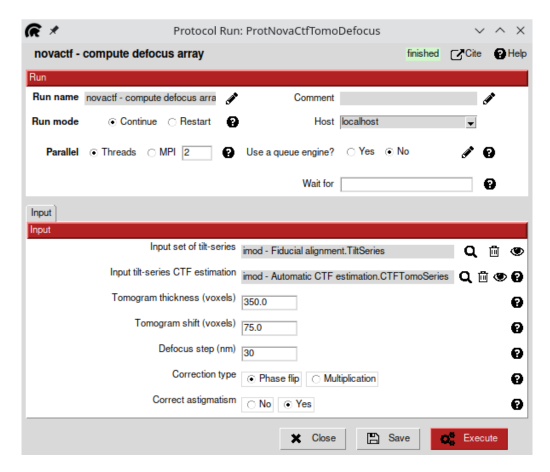


Figure 30: NovaCTF - compute defocus array protocol.

Now locate novactf - 3D CTF correction and reconstruction protocol and use the previous protocol as input. Set **Apply tilt-series alignment? Yes** and leave the other options with default values. Set **Number of threads** equal to the number of available CPU cores to parallelize steps execution. This protocol will perform CTF correction, apply alignment, and filter input tilt-series N times, where N is the number of defocus files generated in the previous protocol. Afterward, it will combine all stacks into a final CTF-corrected tomogram.

7 Wrapping up

That's it! If you have completed the tutorial and read and understood the guide up to this point, you should be ready for some serious tomo data processing using the SCIPION framework! If you found any mistakes in this guide or have more questions, get in touch with us. Below are the resources for further reading:

- Our paper on using SCIPION for tomography (Jiménez de la Morena et al., 2022).
- Other Scipion tutorials for tomography

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