

Python-based **H**ierarchical **EN**vironment for **I**ntegrated **X**tallography

Cryo fit1 FAQ

Contents

- How long does it take to run cryo fit?
- How to enlarge map box size?
- How to generate and record movie?
- How to improve initial cc?
- I see an error message at my 1 make gro step.
- <u>I see</u> "Fatal error: A charge group moved too far between two domain decomposition steps. This usually means that your system is not well equilibrated" at my 8 cryo fit step.
- <u>I see "Fatal error: Number of grid cells is zero. Probably the system and box collapsed." at my 8 cryo fit step.</u>
- I see "step 0 correlation coefficient: nan" at my 8 cryo fit step.
- I see "step 0 correlation coefficient: nan" and "Range checking error: Explanation: During neighbor searching, we assign each particle to a grid based on its coordinates. If your system contains collisions or parameter errors that give particles very high velocities you might end up with some coordinates being +-Infinity or NaN (not-a-number). Obviously, we cannot put these on a grid, so this is usually where we detect those errors. Make sure your system is properly energy-minimized and that the potential energy seems reasonable before trying again." at my 8 cryo fit step.
- I see "User's provided atomic model had 0.0 cc" in my cryo fit.overall log.

How long does it take to run cryo_fit?

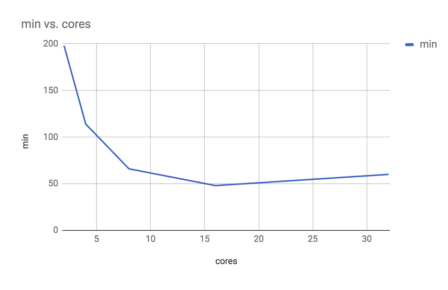
How long does it take to run cryo_fit?

- CentOS with 2 cores (2.7 Ghz)
 - tRNA (6,000 atoms): 2.5 minutes
 - Nucleosome (26,000 atoms): 1 hr
 - Beta-galactosidase (64,000 atoms): 2.5 hrs
 - ribosome (386,000 atoms): 3~7 hrs



- 24 cores (2.7 Ghz)
 - (4~16 cores would give similar/better performance)
 - Beta-galactosidase (64,000 atoms): 3.9 hrs
- 4 cores (3.1 Ghz)
 - Beta-galactosidase (64,000 atoms, 50,000 cryo_fit steps): 5.8 hrs

Benchmark with ribosome (same linux machine, 10k emsteps, 50k number_of_steps, 3x emweight)



 Based on this result, number of cores is recommended up to 16

How to enlarge map box size?

- Doo Nam uses relion image handler to change map box size.
- For example, relion_image_handler --i user.mrc --new_box 370 --o user_box_size_370.mrc
- However, relion_image_handler derived mrc file may not go through cryo_fit's automatic mrc to sit map format conversion properly.
- Therefore, please use <u>situs map2map</u> to convert your mrc format map to situs format map. You can convert by "map2map user.mrc user.sit" then enter 1 for "Convert to classic Situs (auto)*". Then, provide this user.sit file to your cryo fit. For example, phenix.cryo fit user.pdb user.sit
- When Doo Nam provided situs made sit map file, the cryo_fit ran smoothly again.

How to generate and record movie?

• Generate record movie by cryo fit1

How to improve initial cc?

• dock in map or UCSF Chimera's 'fit in map' or UCSF ChimeraX's isolde may improve initial cc.



I see an error message at my 1 make gro step.

- If a user sees "Fatal error: Atom xx in residue xx xxx was not found in rtp entry xx with xx atoms while sorting atoms." on his/her screen,
 - please remove/fix wrong atoms. Running real_spaace_refine via phenix GUI will show which atoms need to be removed/fixed.

I see "Fatal error: A charge group moved too far between two domain decomposition steps. This usually means that your system is not well equilibrated" at my 8 cryo fit step.

- Using macOS 10.13.6 helped rather than using Ubuntu 16.04. Maybe macOS has better numerical stability.
- One gromacs expert suggested to try smaller time_step_for_cryo_fit.
- However but for Doo Nam, simply using macOS solved the problem.

I see "Fatal error: Number of grid cells is zero. Probably the system and box collapsed." at my 8 cryo fit step.

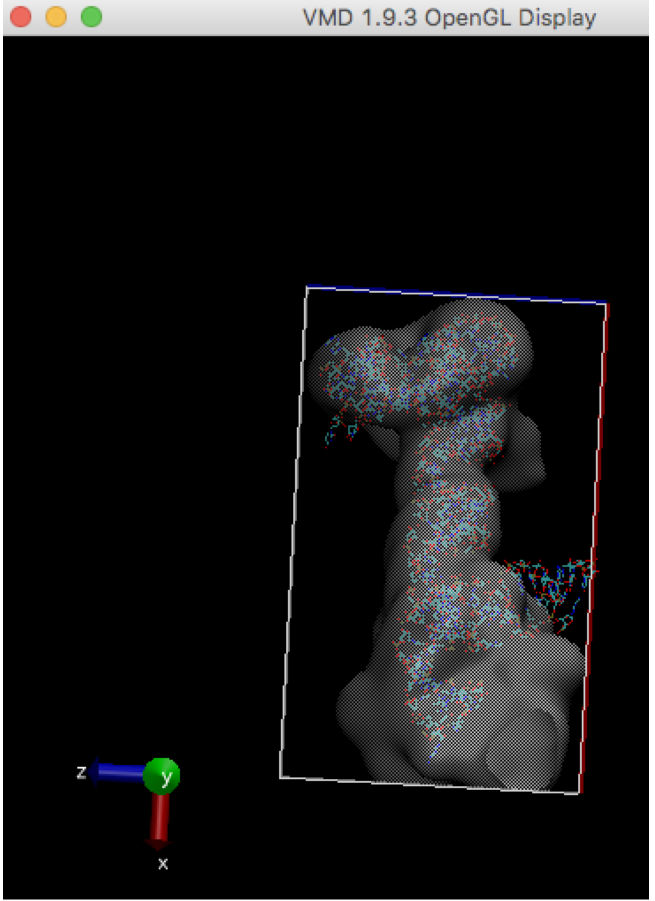
- step_8 may be full of stepxb_nx.pdb.
- Most likely, this means that initial cc is too low for MD simulation.
- When Doo Nam ran real_space_refine first, then run real_space_refined atomic model in cryo_fit, it was solved.
- Alternatively, improve initial cc by fitting initial atomic model into a map (see "How to improve initial cc?" above)
- Less likely, but still a possible case is when the map weight is too high, lowering emweight_multiply_by may help.

I see "step 0 correlation coefficient: nan" at my 8 cryo fit step.

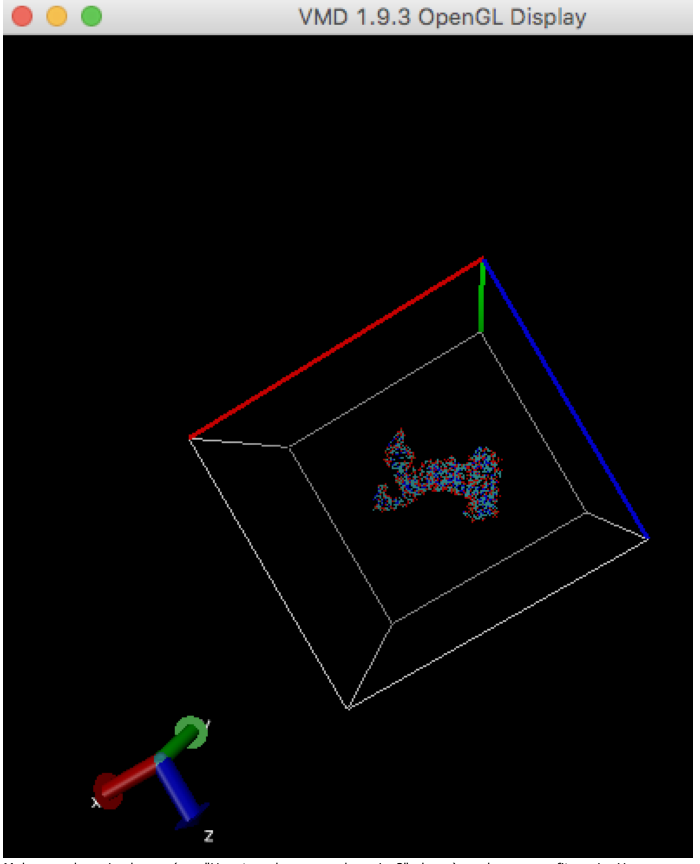
- This often indicates that the initial atomic model is not placed into a cryo-EM map.
- Improve initial cc by fitting initial atomic model into a map (see "How to improve initial cc?" above)

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- When a user used a partical map region ("boxed map") by phenix.map box or relion_image_handler, cryo_fit's automatic mrc to sit map format conversion may not work properly.
- Therefore, please use <u>situs map2map</u> to convert your mrc format map to situs format map. You can convert by "map2map user.mrc user.sit" then enter 1 for "Convert to classic Situs (auto)*". Then, provide this user.sit file to your cryo_fit. For example, phenix.cryo_fit user.pdb user.sit
- When Doo Nam provided situs made sit map file, the cryo_fit ran smoothly again.
- When a user didn't use phenix.map_box, it means that the map dimensions need to be larger.
- Like other MD simulations, gromacs need enough map box size to cover the atomic model to run (ziggle and wiggle). Refer <u>Waters seems to be out of the box</u>
- For example, stuck-out red oxygen atoms outside the right edge of the box are the problem.



• In order to run any MD simulation (including cryo_fit), a box should be large enough like



- Make map box size larger (see "How to enlarge map box size?" above), and run cryo_fit again. You can check map box size by VMD. Alternatively, remove sticking out atoms if these are unnecessary, then run cryo_fit again.
- For protein modeling, I would use <u>cryo_fit2</u> which is not limited by box size requirement. It better fits than cryo_fit1 in terms of fitting and geometry anyway.

I see "User's provided atomic model had 0.0 cc" in my cryo_fit.overall_log.

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