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Protocol status: Working We use this protocol and it's working

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Model building and refinement of RCKW and FL-LRRK2 bound to inhibitors

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ABSTRACT

Protocol to model and refine a PDB against a cryo-EM map.

MATERIALS

ChimeraX¹

COOT^{2,3}

Phenix⁴

Rosetta⁵

BEFORE START INSTRUCTIONS

Install the needed software (Chimera, COOT) and Phenix or Rosetta.

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- 1 Split into domains protein models using Chimera that will be used as a starting point (PDBs 6VP7 and 7LHW for RCKW and FL LRRK2, respectively). Save each domain in a PDB file separately.
- 2 Open maps with the best resolution in Chimera and fit every domain PDB into the map. Save the new position for each PDB domain file.
- 3 Open PDB files in COOT and merge all files. Save a new PDB file with all domains.
- Check all amino acids manually and refine/regularize them against the cryo-EM map. You can find a basic COOT tutorial for cryo-EM here:
 https://www2.mrc-lmb.cam.ac.uk/personal/pemsley/coot/web/tutorial/Coot-Cryo-EM-basics.html
- **4.1** You might want to model amino acids not present in your starting point PDBs if you have empty density. To do that:
 - 1. Add residue (Right side, eighth icon from the bottom)
 - 2. Click on the alpha carbon of the last amino acid.
 - 3. Mutate residue (right side, tenth icon from the bottom). It pops up a list with all amino acids. Click on the amino acid you want to add).
- 5 Once you manually checked your model, save it. Then, add your ligand. To do that:

- **5.1** Save your ligand in a PDB file. Open Phenix and use elBOW application to generate restrains for your ligand.
 - 1. On the right side, click Ligands > elBOW
 - 2. On elBOW setup, click PDB file with hydrogens and/or CONECT records > OK
 - 3. Click Use simple optimization > OK
 - 4. In geometry file, browse your PDB_ligand.pdb
 - 5. Browse your output directory
 - 6. Type Output file prefix
 - 7. Run the job.

Note

If you don't have a PDB and you have instead a SMILES notation for your ligand, you can generate a PDB and restrains for it by clicking on the step 2 SMILES string option instead

- Open the ligand PDB_Ligand.pdb in Chimera, fit it into the map, and save a new file. Open it in COOT together with your model protein and merge them. Save a new PDB with ligan plus protein.
- **5.3** Refine/Regularize your ligand against the map in COOT. Save your new PDB.
- **6** Refine your model against your map in Phenix or Rosetta:
- **6.1** In Phenix:
 - 1. Click Cryo-EM > Refinement > Real-Space Refinement
 - 2. Input/Output Tab:
 - 2.a: Type your Job Title
 - 2.b: Add file > PDB_protein+ligand.pdb
 - 2.c: Add file > map.mrc
 - 2.d: Add file > Ligand_restrains.cif
 - 3. Add map resolution
 - 4. In refinement settings Tab: Keep it as default
 - 5. Run the job
 - 6. Once the job is done, open your new PDB_protein+ligand_new.pdb in COOT and check amino acids and restrains.

7. Check Ramachandran plot. Manually correct amino acid outliers and allowed

6.2 In Rosetta:

- 1. Remove inhibitors and nucleotides from the model.
- 2. Rosseta/1.3 for refinement.
- 3. Move map and pdb into your directory
- 4. Load rosetta using module load rosetta/3.13
- 5. Once is finished go to COOT, add your ligand and remove all the hydrogens from your structure
 - 5.a: To remove hydrogens in COOT, go to Calculate > Scripting
 - 5.b: Type delete_hydrogens(#) (# is the model)
- 7 Check validation parameters and fix outliers you may have in your PDB. Run a refinement in Phenix or Rosetta (it is an iterative process, so you might have to correct/refine it several times).