Tutorial: Ligand Fitting with Coot

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1 Introduction

We have a protein structure (which is well-refined). However, we have not yet identified the ligand position - indeed we don't even know if the ligand is bound or not

In this tutorial, we will build a ligand and search the density for this ligand, add the ligand to main protein molecule and refine and validate the ligand.

2 Experimental Details

This protein is the catalytic domain of poly(ADP-ribose)polymerase 2 crystallised from PEG 3350 (polyethylene glycol), Tris and cryoprotected using glycerol.

3 Starting

So let's load the protein coordinates <code>ligand-fitting-no-ligand.pdb</code> and the associated MTZ file which is <code>ligand-fitting.mtz^1</code>. Use File \rightarrow AutoOpen MTZ.

4 SMILES

Now what is the ligand that we want to find? It's 3-aminobenzamide

Figure 1: 3-amino-benzamide

How do we make a 3D model of this ligand? We use a SMILES string (and LIBCHECK).

So what is the SMILES string for 3-aminobenzamide?

¹available, as usual, from http://www.biop.ox.ac.uk/coot/tutorial/

If you can't draw out the SMILES string, try this link in your browser:

http://www.molinspiration.com/cgi-bin/properties

draw the molecule there and hit the "Calculate Properties" button - the next pages gives you the SMILES string.

We need to convert this SMILES string representation to a molecule. In Coot we do that using File \rightarrow SMILES... 2 Enter you SMILES string in the second of the two entry boxes. Press "Go".

After a few moments (in which Coot runs LIBCHECK) a molecule will appear at the centre of the screen.

Is it the molecule that you wanted?

Note: check the hydrogens and planarity.

5 Fitting Ligands

We how the correct ligand now, So let's move on to search the map for entities of this kind.

Calculate \rightarrow Other Modelling Tool \rightarrow Find Ligands...

We have 2 maps from which to choose. For now, let's choose the 2mFo-DFc map and find density clusters above 1.0 sigma. Make sure that you select the correct map and protein molecule and have selected the ligand that you made with the SMILES string.

Press "Find 'Em!"

What do you see!?

You should see 4 hits.

Let's use the Fitted Ligands dialog to navigate to these hits.

Do we like all the hits? If not, why not?

Let's worry about any problematic fittings later on and for now, let's concentrate on the nicely fitting solutions.

If you like the any of the hits, merge them into the main protein molecule.

Using Calculate → Merge Molecules,

In the top pane, select the ligand hits that fit nicely by ticking them. In the lower option menu, select the protein model. Then click "Merge". Now we have 2 representations of the ligands, which can be confusing, so let's use the Display Manager to undisplay the Fitted Ligands that we merged into the main protein.

Using the Fitted Ligand dialog, navigate to the newly added monomers and optimise the fit to density - we'll use Real Space Refinement to do that. Use the blue target on right-hand toolbar and click an atom in the residue twice. Do you like the result of the refinement? What is the torsion angle of the carboxy-amine?

6 Presentation

We can emphasise the atom positions using Extensions \rightarrow Representations \rightarrow Hilight Interesting Site.

Nice?

Maybe...

 $^{^2}Note$ that the first line is the three letter code, conventionally this is 3 uppercase letters, e.g. LIG, DRG, XXX. Note that if you use several different SMILES strings, they each should have a different 3-letter-code.

7 Problem Areas

OK, now we have fitted the nicely fitting ligands, let's go back to the problematic blobs.

Validate → Find unmodelled blobs

We shall search the 2mFo-DFc as did previously using the same protein model (which now has the 3-aminobenzamide fitted). Press "Find blobs" 3

We should get a dialog with two blobs listed.

Examine those two blobs. We know that 3-aminobenzamide does not fit these blobs well. What else could these blobs be? Let's look at the crystallization and freezing conditions? Any clues?

Now, we don't necessarily know the 3-letter-codes of the molecules that might fit these blobs. We can use the searching tool in Coot to convert between a molecule name and its 3-letter-code.

File \to Search Monomer Library... add the molecule name to the entry box and press "Search"

You will see a list of molecule that include the text you typed as part of the molecule names.

If you click on any of those buttons, Coot runs LIBCHECK and generates a molecule and puts it at the centre of the screen.

You can do this for the 3 or 4 different molecules that you think this blob might be.

The centre of the screen is a bit crowded now. Undisplay those new ligands. To easily identify the best fitting ligand at the site, let's do ligand fitting here.

Calculate → Other Modelling Tools → Find Ligands

Select the 2mFo-DFc map as usual, and the main protein as before and the ligands to search for are the 3 or 4 newly created ligands.

This time we will Search "Right Here". You can turn on flexible ligands if you like, but this will slow down the search considerably.

8 Tidying Up

Real Space refinement is the tool to tidy things up. Using the blue circle icon on the right, then click click on an atom in the residue ⁴.

When we are happy with our fit, we can merge this new ligand into the main protein ligand as we did before.

9 NCS Ligands

You may have noticed by now that there are 2 protein chains related by non-crystallographic symmetry. We can exploit this NCS to help fit ligands. We have fitting a ligand that bound to the "A" chain, now let's use NCS to position a ligand in related position in the "B" chain of the protein.

Extensions \rightarrow NCS \rightarrow NCS Ligands...

The protein with the NCS is the main protein as usual, the molecule containing the ligand is also the main protein as usual (if we have done the merge mentioned above). We need to know the chain ID of the ligand that we have fitted into the "A" chain, to do that double left-click an an atom in the ligand. In the graphics you will see something like: " C1 / 1 XXX/E" - so the chain-id is "E". Also the status bar at the bottom shows information about the clicked atom.

³we could also use Validate → Difference Map Peaks to navigate to these positions.

 $^{^4}$ or (quicker) simply press the "R" key (if you have the key bindings set up).

So enter that chain-id in the "Molecule containing ligand" "Chain ${\rm ID}$ " entry. Then "Find Candidate positions".

A fitted ligands dialog box pops up and you can use the buttons in that dialog to navigate to the NCS ligand positions.

Refine this NCS ligand position if needed and merge it into the main molecule.

10 For the Adventurous

If you are feeling adventurous, you can run refmac: Calculate \to Model/Fit/Refine \to Run Refmac...

Choose an MTZ file for Refmac and use the file selector to choose the same MTZ file that we used in the beginning.

Examine the resulting molecule. Are there any problems left to fix?