

## Refmac tutorial

Download tutorial file from the website:

[www.yybl.york.ac.uk/refmac/refmac\\_tutorial.tar.gz](http://www.yybl.york.ac.uk/refmac/refmac_tutorial.tar.gz)

Create a subdirectory where you usually work and copy tutorial file to this directory.

```
mkdir refmac
cd refmac
mv wherever_tutorial_file/refmac_tutorial.tar.gz .
```

Or copy tutorial materials from the \$CEXAM directory

```
cp $CEXAM/refmac_tutorial.tar.gz .
```

Then untar and go to the refmac\_tutorial directory.

```
tar xzvf refmac_tutorial.tar.gz
cd refmac_tutorial
```

There are several files to play around. These files can only give you some idea about refinement in general and refmac5 in particular.

Start ccp4i. For each case you should create a project in ccp4i using “Directories&ProjectDir” on the top right of the interface. It will help you to trace back what you have done and find files you need. There are five subdirectories. Each subdirectory is for one tutorial. They are: **rnase** – a simple refinement, **1n5b** - tls refinement, **twin** – twin refinement, **sad** – sad refinement, **lowres** – low resolution refinement and automatic NCS restraints.

### I) Simple refinement.

On the top of the ccp4i select “Refinement”. Click “Run Refmac5”. It will bring up refmac5 interface. Define input mtz and coordinate files. They are rnase115\_unique1.mtz and rnase.pdb. Press Run/Run now on the left bottom side.

Current version of the program uses automatic weighting. For many cases it works sufficiently well. Sometimes (especially at high resolution) default weighting may give a relaxed geometry. In this case you can play with the weighting of X-ray and geometry terms. If you want to change weights then click “Refinement Parameters”, under this unclick “Use automatic weighting” and add an appropriate number into the field “Use weighting term”. For low resolution you may want to use very small values - 0.01 or even smaller. For higher resolution this number may need to be as high as 10. You may need to run refinement with different values to get it right. If after refinement run rms bond distances are more than 0.02 then you may want to reduce weighting, if rms bond value is less than 0.01 then you may need to increase. Smaller weighting value means tighter geometry.

Now click on the “run” button. Program should run.

While it is running you can have a look at the log file. When reftmac finishes this job you should use coot (or another program) to view coordinates, maps and make corrections if you need to.

```
coot -pdb rnase_refmac1.pdb -data rnase115_refmac1.mtz
```

and analyse the electron density.

## II) TLS refinement

This example of TLS refinement in Refmac5 uses 1n5b, a molecular chaperone. The asymmetric unit contains 2 dimers, chains A + B and chains C + D. Data is in P212121 to 2.0Å

To run Refmac5, we need to specify what TLS groups we wish to use. CCP4 wiki site has a page about different ways of creating TLS groups.

To create TLS group using ccp4i:

1. Go to ccp4i -> Refinement -> Model preparation -> Create/Edit TLS file.
3. Give "TLS out" a name such as 1n5b\_in.tls
4. Define 4 groups (use "Add another TLS group" to create additional entries):
  - Chain A, residues 3 to 130
  - Chain B, residues 3 to 130
  - Chain C, residues 3 to 129
  - Chain D, residues 3 to 130
5. Click "Run now" to create the file.

We are now ready to do TLS refinement

6. Select the task Run Refmac5
7. Select mode "TLS & restrained refinement" from the protocol folder (top left button after the "Job title").
8. Set the files:
  - HKLIN 1n5b.mtz
  - XYZIN 1n5b.pdb
  - TLSIN 1n5b\_in.tls (from above)

Note 1: If you do not give input tls file then each chain will be taken as a tls group.

Note 2: Waters close to tls groups will be added to those groups (it is a new feature and available from ccp4 6.0.99c onwards)

Note 2: If you do not give tls file then reftmac will choose each chain as single tls group. So you can try just step 7 (Select mode "TLS & restrained refinement")

9. In the folder "TLS parameters" select "Set initial Bfactors"
10. For a quick run, use 5 TLS cycles and 5 restrained cycles

When Refmac5 has finished, check the logfile for the usual things: Rfree, geometry, warning messages. TLS section shows the raw TLS parameters at each cycle. These don't mean much on their own, but you can see if the TLS refinement

is converging. Now we need to analyse the TLS parameters. It is available from ccp4 6.1.1 onwards.

11. In the "Model Completion & Analysis" section select "Analyse TLS parameters" task.
12. Set the files:
  - TLSIN is TLSOUT from previous Refmac job
  - XYZIN is XYZOUT from previous Refmac job
13. Click "Output file containing the axis .." and give the AXES file a name  
1n5b\_tls.vector
14. In the Axes folder, select mmCIF format
15. In Other Options, select "Analyse derived atomic ...."

Check the log file for "NON-POSITIVE DEFINITE WARNINGS" (I think there will be none!)

Output coordinates contain ANISOU records derived from the TLS plus residual B factors. Use coot to view them. Axis can be viewed using ccp4mg (I think?)

### III) Twin

Run refmac5 as usual. Click Twin refinement button on the interface. It will activate twin refinement.

In some versions of ccp4i there was a bug and twin refinement was not activated. To check if it is activated check the log file. If it has TWIN keyword then everything is fine. If twin does not run automatically then use "Include keyword file". Click on "Browse" and select file keyw.dat. This file contains a single keyword - TWIN. Adding this should activate twin refinement.

Keyword file could be used to add options that are not available on the current interface.

*Note* that in this case Rfactor and Rfree are diverging. It is because 1) This data set is from pdb and there was no twin flag there 2) Twin relates two (or more) set of reflections. Usual free R selection does not take this fact into account. Refmac5 puts twin related reflections to the same set (i.e. either all related reflections belong to "free" or to "working" set). In general free reflection selection should be done before starting refinement accounting for potential twin operators. One way of achieving this is to select free reflections at higher group and expand them. It will ensure that throughout refinement and all other treatment of data all related reflections will belong to the same set.

Look at the maps.

### IV) SAD

Refinement against F+ and F- directly (SAD refinement).

Start refmac5 as usual. On the top right side of the interface choose “SAD data directly” instead of “no prior phase information”. Type in the “Anomalous atom” field SE, f’ field -8.0 and f’’ field 4.0. You can also use wavelength. Then all atoms that have significant f’’ will be used in anomalous refinement.

After refinement finished you should look at the maps. There are coefficients for anomalous difference map also (corresponding mtz labels are FAN and PHAN for amplitudes and phases of anomalous difference map).

## V) Low resolution refinement

If you have older version of ccp4 then you can follow the instructions described in the presentation:

[www.yesbl.york.ac.uk/refmac/Presentations/Refmac\\_Erice\\_workshop.ppt](http://www.yesbl.york.ac.uk/refmac/Presentations/Refmac_Erice_workshop.ppt)

slides 45-50

Full description of new features are in:

[www.yesbl.york.ac.uk/refmac/data/refmac\\_news.html](http://www.yesbl.york.ac.uk/refmac/data/refmac_news.html)

In the new version of ccp4 there are options for jelly body, automatic NCS and map sharpening.

- a) Jelly body is under “Refinement Parameters”. You need to click “Use jelly-body refinement with sigma”. Change sigma to 0.01 or 0.02. This value defines “jelliness”. Smaller value means tighter restraints
- b) Automatic NCS restraints are under “Setup Non-Crystallographic Symmetry”. You need to click “use automatically generated local NCS restraints”. You can also use global NCS
- c) Map sharpening is under “Monitoring and Output Options”. You need to click “Perform map sharpening with B value 20.0”. B value should a little bit smaller than Wilson’s B value.

## Dictionary

Tutorials for dictionary are located on the web page:

<http://www.yesbl.york.ac.uk/mxstat/JLigand/index.html>

To run jligand you need to have java 1.5 or later versions. There are three tutorials on this site: 1) How to create ligand description; 2) Link description and 3) ligands with metal.