

Ligands in Phenix

Generating & modifying for all scenarios

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User's Meeting
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What are restraints?

- Restraints are harmonic functions that provide (mostly) chemical information via residuals & gradients
- Needed because experimental data is rarely sufficient for structure determination
- Weighted with the experimental information to find the optimal result

Resolution dependence

- Ultra-hi res – Not needed
- Hi res – Can have large deviations because the experimental data dominates
- Lo res – Generally approaches the ideal values
 - If not, large scale problems

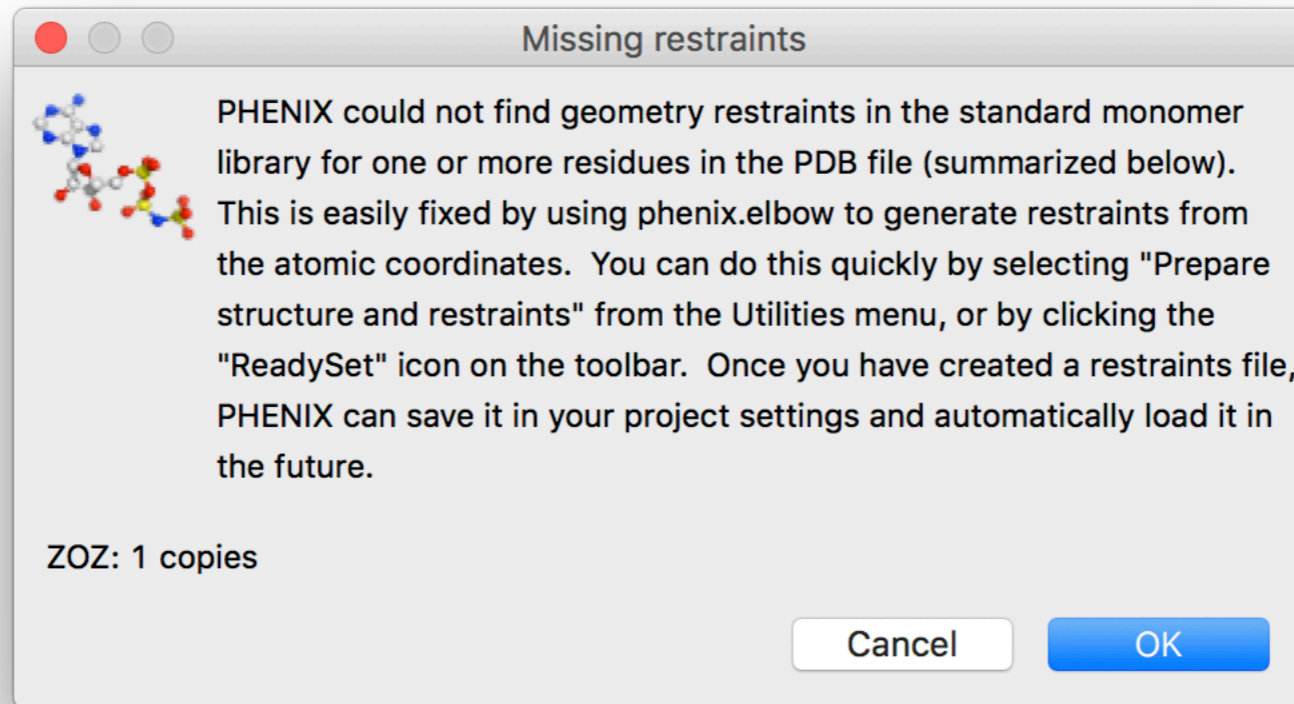
Restraints in Action

- Libraries
 - Monomer Library
 - GeoStd
- Algorithms
 - Polymer
 - Links

GeoStd

- All standard amino acids
- Current list of non-standard amino acids
- All standard RNA/DNA
- Current list of non-standard RNA/DNA
- Others – 45k Mogul validated restraints using PBEh-3c/CPCM and higher QM

What you will see



Or

Sorry: Fatal problems interpreting model file:

Number of atoms with unknown nonbonded energy type symbols: 21

Please edit the model file to resolve the problems and/or supply a CIF file with matching restraint definitions, along with `apply_cif_modification` and `apply_cif_link` parameter definitions if necessary.

CIF

- Crystallographic Information File
- mmCIF – macro-molecular CIF
- Used for
 - Model
 - Data
 - Maps
 - Ligands
 - Information
 - Restraints

Confusion

- All depositions of X-ray model use mmCIF from 1 July 2019
- “I need a CIF file.”
 - But what do you really need?

Restraints?

- Provide a reasonable geometry during refinement particularly at low resolution
 - Bonds, angles, dihedrals, chirals, planes, ...
- Must be weighted against the experimental information

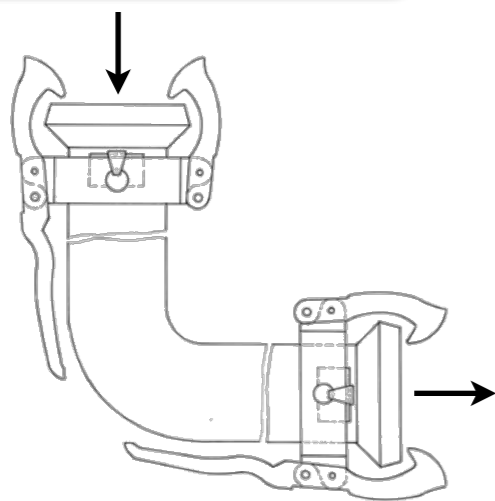
Overview

- eLBOW - electronic Ligand Builder & Optimisation Workbench
- ReadySet! - One-stop preparation for your refinement needs
- REEL - Restraints Editor Essentially Ligands

eLBOW goals

- Fast, simple and flexible procedure to include ligands
- Reduce the tedium of building 3D ligand models
- Automate generation of restraints for ligands

Chemical input



Chemical restraints (CIF)
Cartesian coordinates (PDB)

Reflection data
Protein information

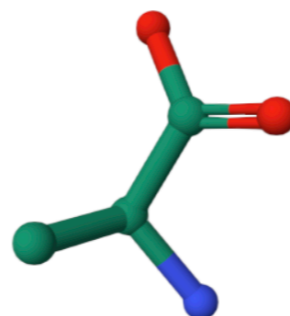
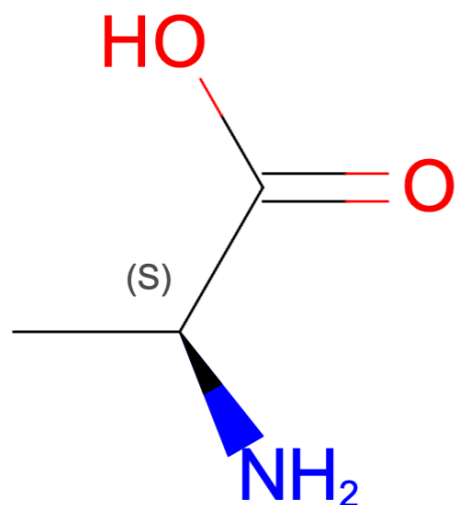
refinement

N. W. Moriarty, R. W. Grosse-Kunstleve, P. D. Adams, (2009). Acta Cryst. D 65, 1074-1080.

Chemical Component Library

- List of all the entities in the Protein Data Bank
 - Amino acids, Nucleic acids
 - Ligands, Small molecule
 - Metal clusters
- In CIF format
 - Contents chemical information
 - SMILES, atom names, bonds
 - Not restraints

Amino Acid



Toggle Hydrogen

Toggle Labels

 Display Files ▼

 Download Files ▼

 Data API

ALA

ALANINE

Find entries where: ALA

- is present as a standalone ligand in [172 entries](#)
- as a non-polymer is covalently linked to polymer or other heterogen groups [58 entries](#)
- is present in a polymer sequence [210,908 entries](#)

Find related ligands:

[Similar Ligands \(Stereospecific\)](#)

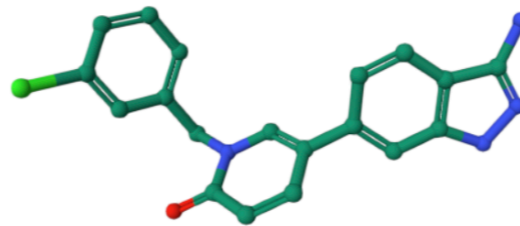
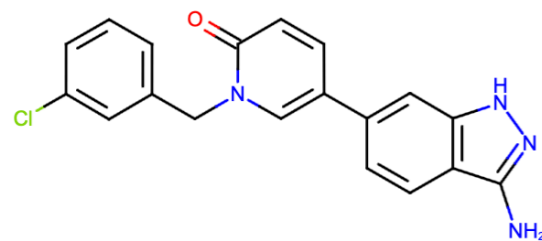
[Similar Ligands \(including Stereoisomers\)](#)

[Similar Ligands \(Quick Screen\)](#)

[Similar Ligands \(Substructure Stereospecific\)](#)

[Similar Ligands \(Substructure including Stereoisomers\)](#)

5-letter codes



Toggle Hydrogen

Toggle Labels

Display Files

Download Files

Data API

A1LU6

5-(3-azanyl-1~{H}-indazol-6-yl)-1-[(3-chlorophenyl)methyl]pyridin-2-one

Find entries where: A1LU6

is present as a standalone ligand in 1 entries

search

Find related ligands:

[Similar Ligands \(Stereospecific\)](#)

[Similar Ligands \(including Stereoisomers\)](#)

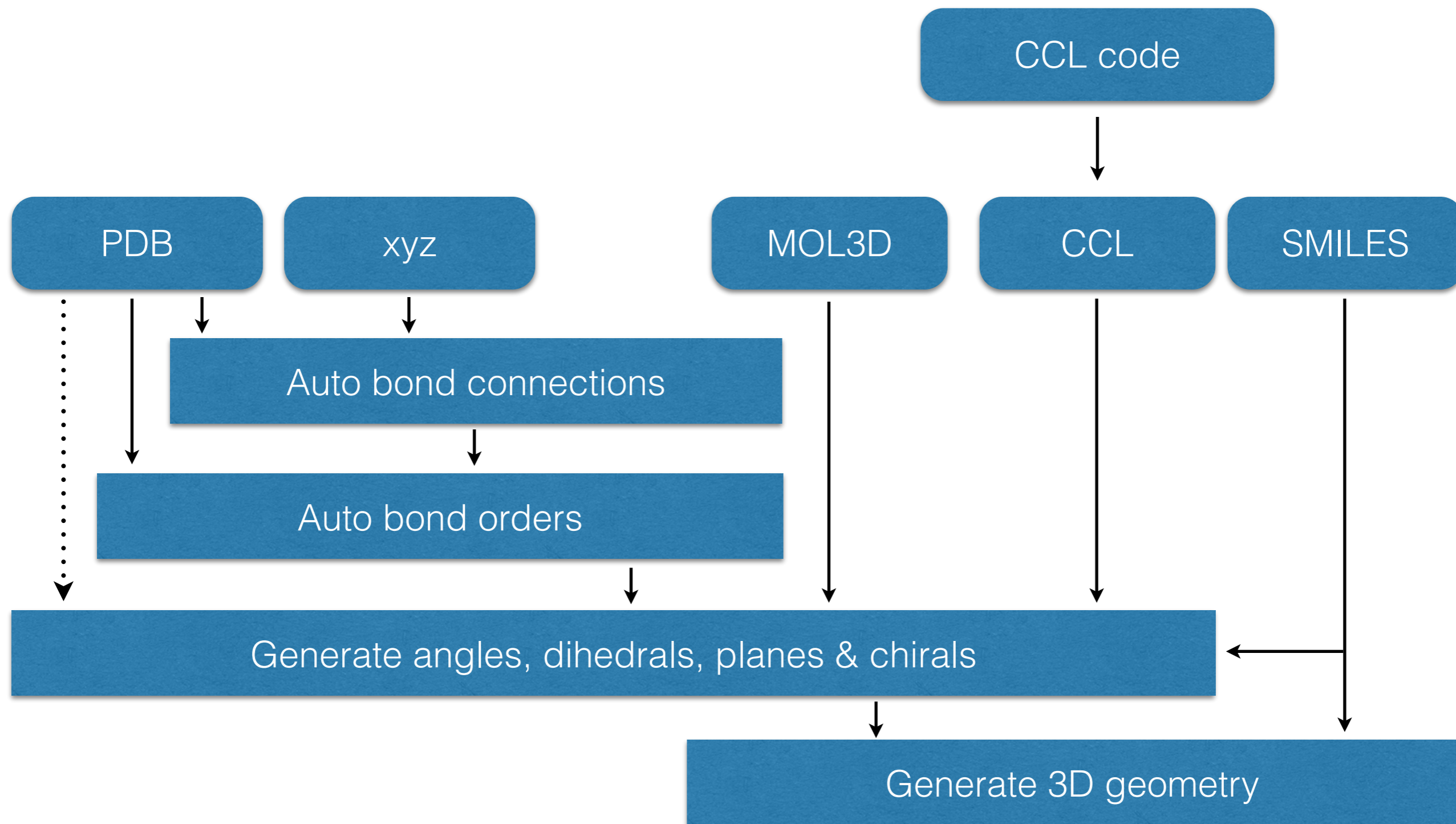
[Similar Ligands \(Quick Screen\)](#)

[Similar Ligands \(Substructure Stereospecific\)](#)

[Similar Ligands \(Substructure including Stereoisomers\)](#)

- 49k combinations for 3-letter codes

Topology

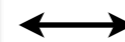


Optimisation

Topology information - Atoms, bonds, angles, ...



Simple force field geometry optimisation



Add hydrogens



Advanced geometry optimisation

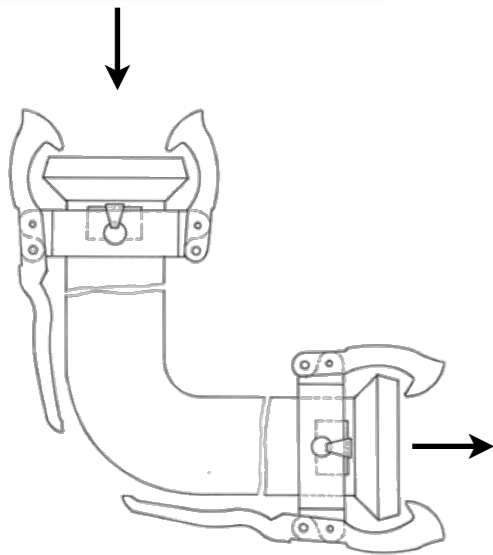


Output geometry (PDB) and restraints (CIF)

Getting ready to refine

- Many details needed to prepare for structure refinement

Chemical input



Chemical restraints (CIF)
Cartesian coordinates (PDB)

Protein Information

ReadySet!

Experimental data

refinement

ReadySet!

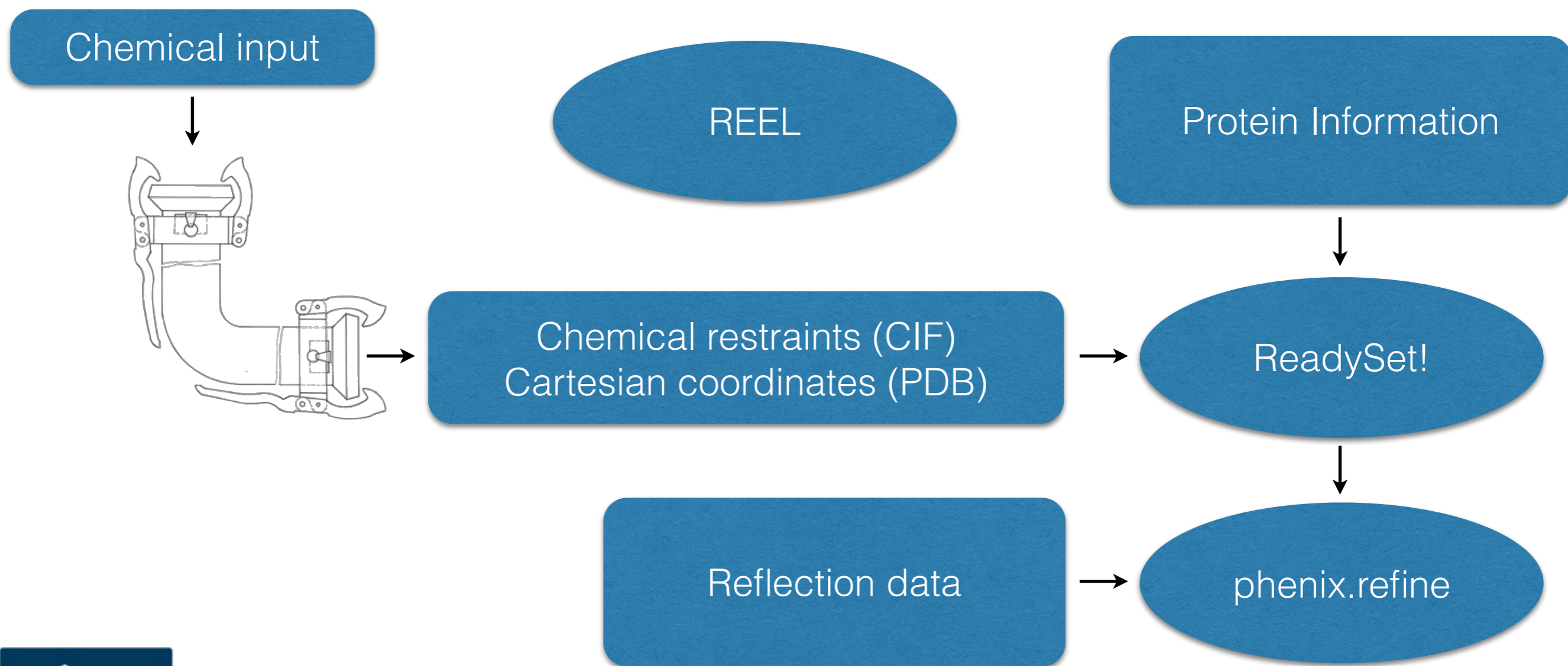
- Add hydrogens
 - Default: adds hydrogens to protein, ligands
 - Protein - Reduce
 - Ligands - eLBOW
 - Add hydrogens to water
 - Add deuteriums instead of hydrogens
 - Add hydrogen & deuteriums appropriately
- Generate restraints

ReadySet!

- Restraints (CIF) filename
- Restraints (CIF) directory
- LINKS to “edits”
- --dry-run to show ligand process pathway
- Metal coordination

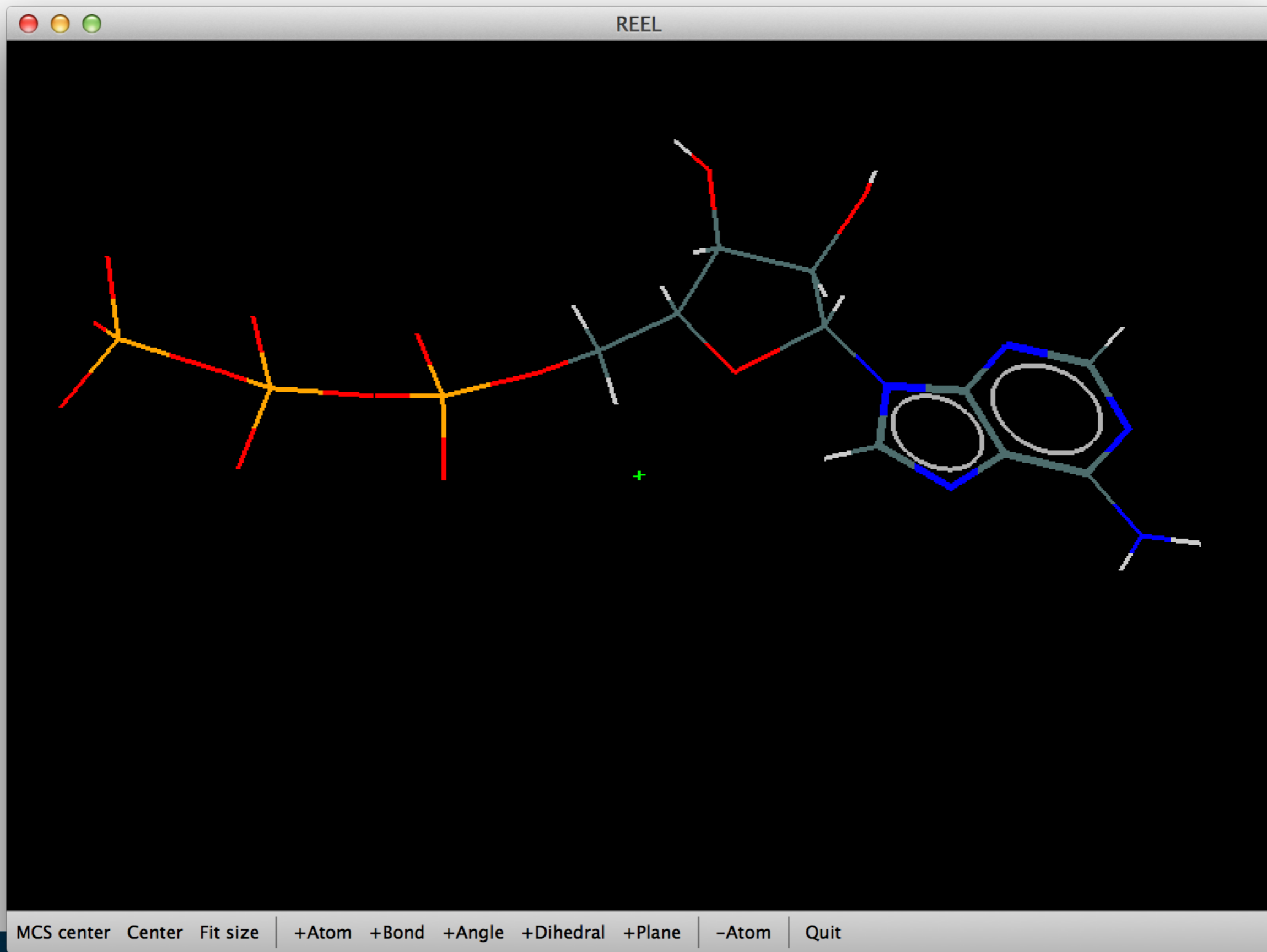
Restraints editing

- Visualisation of primary and secondary restraints
- Adjusting restraints to user preference



Restraints Editor, Essentially Ligands

- Generate a geometry for a set of restraints
- Modify restraints and generate new geometry
- Fast editing in menu items
- Highlight atom and restraints are highlighted
- Multiple ligands simultaneously
- Highlight restraint and atoms are highlighted
- Save restraints (CIF)
- Save geometry (PDB)
- Run eLBOW



Restraints Editor Especially Ligands (REEL)

Simple Optimisation eLBOW Optimisation AM1 Optimisation Search Components Find unique code

ATP

Atoms(43) Bonds(45) Angles(78) Dihedrals(30) Planes(17) Chirals(4) CisTrans Chirals Implicit(7) BoatChair

| | ? | comp_id | atom_id | type_symbol | type_energy | charge | partial_charge | x | y | z |
|----|---|---------|---------|-------------|-------------|--------|----------------|-----------|-----------|-----------|
| 1 | | ATP | PG | P | P | 0 | . | -2.009900 | -8.939900 | 0.927100 |
| 2 | | ATP | O1G | O | O | 0 | . | -0.919700 | -9.907100 | 1.322000 |
| 3 | | ATP | O2G | O | OP | -1 | . | -3.257700 | -9.235200 | 1.724400 |
| 4 | | ATP | O3G | O | OP | -1 | . | -2.306500 | -9.089900 | -0.545900 |
| 5 | | ATP | PB | P | P | 0 | . | -0.243200 | -6.679500 | 0.469800 |
| 6 | | ATP | O1B | O | O | 0 | . | 1.050300 | -7.257000 | 0.992700 |
| 7 | | ATP | O2B | O | OP | -1 | . | -0.341900 | -6.929700 | -1.016000 |
| 8 | | ATP | O3B | O | O2 | 0 | . | -1.514200 | -7.398500 | 1.233500 |
| 9 | | ATP | PA | P | P | 0 | . | 0.670000 | -3.968400 | -0.041400 |
| 10 | | ATP | O1A | O | O | 0 | . | 0.729800 | -4.332500 | -1.505600 |
| 11 | | ATP | O2A | O | OP | -1 | . | 2.063300 | -3.990800 | 0.540200 |
| 12 | | ATP | O3A | O | O2 | 0 | . | -0.276300 | -5.056600 | 0.757100 |
| 13 | | ATP | O5' | O | O2 | 0 | . | 0.029800 | -2.458300 | 0.123600 |
| 14 | | ATP | C5' | C | CH2 | 0 | . | 0.490600 | -1.443000 | -0.721300 |
| 15 | | ATP | C4' | C | CH1 | 0 | . | -0.438000 | -0.204600 | -0.596500 |
| 16 | | ATP | O4' | O | O2 | 0 | . | 0.055400 | 0.644000 | 0.263400 |
| 17 | | ATP | C3' | C | CH1 | 0 | . | -0.496500 | 0.556500 | -1.940500 |
| 18 | | ATP | O3' | O | OH1 | 0 | . | -1.718900 | 0.408900 | -2.511600 |
| 19 | | ATP | C2' | C | CH1 | 0 | . | -0.245900 | 2.038900 | -1.588600 |
| 20 | | ATP | O2' | O | OH1 | 0 | . | -1.282600 | 2.903300 | -2.232200 |
| 21 | | ATP | C1' | C | CH1 | 0 | . | -0.345300 | 2.109700 | -0.288400 |
| 22 | | ATP | N9 | N | NR5 | 0 | . | 0.564300 | 3.101200 | 0.226900 |
| 23 | | ATP | C8 | C | CR15 | 0 | . | 1.894800 | 3.031600 | 0.318500 |

View preferences loaded



Restraints Editor Especially Ligands (REEL)

Simple Optimisation

eLBOW Optimisation

AM1 Optimisation

Search Components

Find unique code

ATP

Atoms(43)

Bonds(45)

Angles(78)

Dihedrals(30)

Planes(17)

Chirals(4)

CisTrans

Chirals Implicit(7)

BoatChair

| | ? | comp_id | atom_id_1 | atom_id_2 | type | value_dist | value_dist_esd | value_dist_neutron |
|----|---|---------|-----------|-----------|--------|------------|----------------|--------------------|
| 1 | | ATP | PG | O1G | deloc | 1.510000 | 0.020000 | 1.51 |
| 2 | | ATP | PG | O2G | deloc | 1.510000 | 0.020000 | 1.51 |
| 3 | | ATP | PG | O3G | deloc | 1.510000 | 0.020000 | 1.51 |
| 4 | | ATP | PG | O3B | single | 1.648000 | 0.020000 | 1.648 |
| 5 | | ATP | PB | O1B | deloc | 1.510000 | 0.020000 | 1.51 |
| 6 | | ATP | PB | O2B | deloc | 1.510000 | 0.020000 | 1.51 |
| 7 | | ATP | PB | O3B | single | 1.648000 | 0.020000 | 1.648 |
| 8 | | ATP | PB | O3A | single | 1.648000 | 0.020000 | 1.648 |
| 9 | | ATP | PA | O1A | deloc | 1.510000 | 0.020000 | 1.51 |
| 10 | | ATP | PA | O2A | deloc | 1.510000 | 0.020000 | 1.51 |
| 11 | | ATP | PA | O3A | single | 1.648000 | 0.020000 | 1.648 |
| 12 | | ATP | PA | O5' | single | 1.648000 | 0.020000 | 1.648 |
| 13 | | ATP | O5' | C5' | single | 1.399000 | 0.020000 | 1.399 |
| 14 | | ATP | C5' | C4' | single | 1.553000 | 0.020000 | 1.553 |
| 15 | | ATP | C5' | H5'1 | single | 0.970000 | 0.020000 | 1.09 |
| 16 | | ATP | C5' | H5'2 | single | 0.970000 | 0.020000 | 1.09 |
| 17 | | ATP | C4' | O4' | single | 1.305000 | 0.020000 | 1.305 |
| 18 | | ATP | C4' | C3' | single | 1.546000 | 0.020000 | 1.546 |
| 19 | | ATP | C4' | H4' | single | 0.970000 | 0.020000 | 1.09 |
| 20 | | ATP | O4' | C1' | single | 1.617000 | 0.020000 | 1.617 |
| 21 | | ATP | C3' | O3' | single | 1.357000 | 0.020000 | 1.357 |
| 22 | | ATP | C3' | C2' | single | 1.544000 | 0.020000 | 1.544 |
| 23 | | ATP | C3' | H3' | single | 0.970000 | 0.020000 | 1.09 |

View preferences loaded



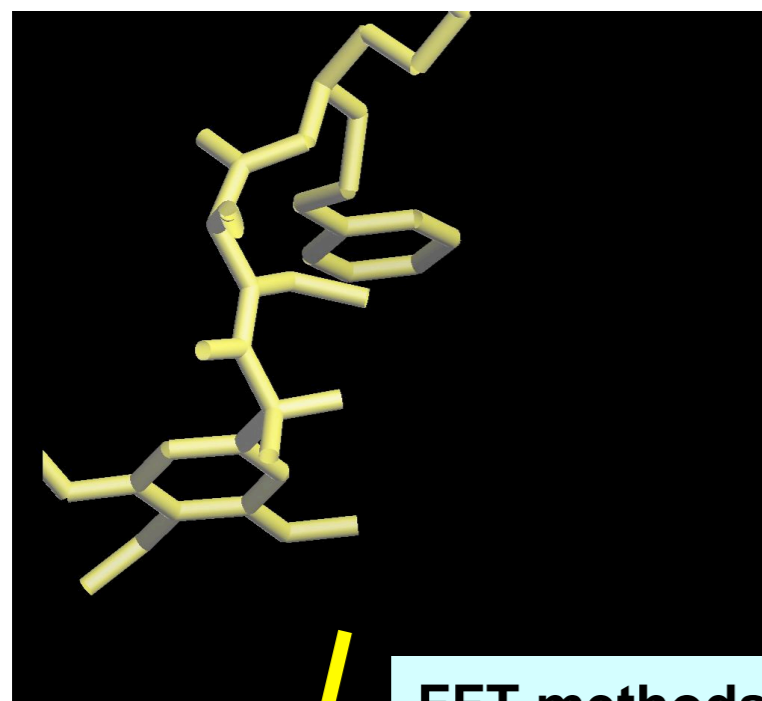
Ligand Fitting



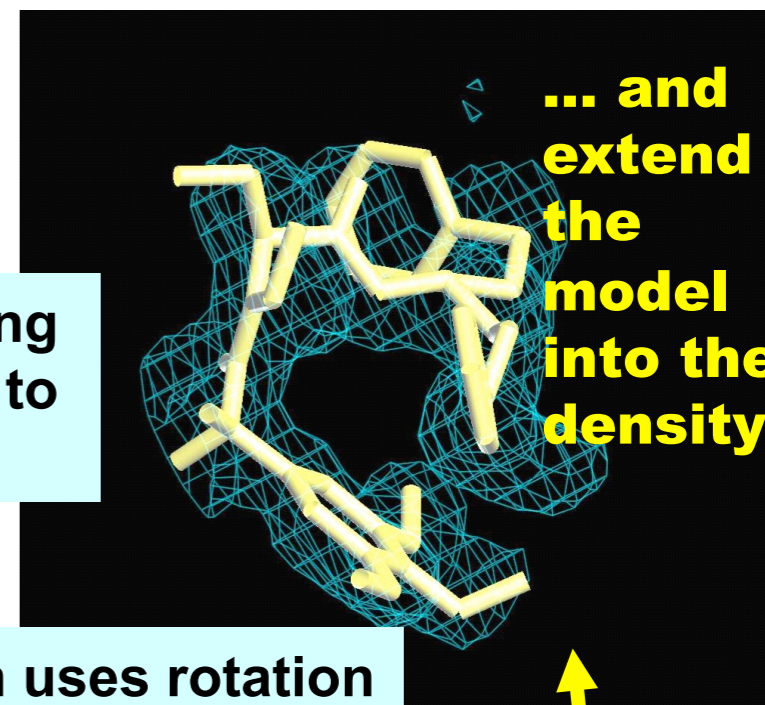
Approach

- Where is the ligand?
 - Choose the largest region of contiguous density
- What are rotatable bonds?
 - Analyze ligand for allowed rotations
- What is the orientation of the ligand?
 - Fit core of ligand
- What is the conformation of the ligand?
 - Trace the ligand out from the core

Automated Ligand Fitting



The best fit is measured using the correlation of the model to the electron density



... and extend the model into the density

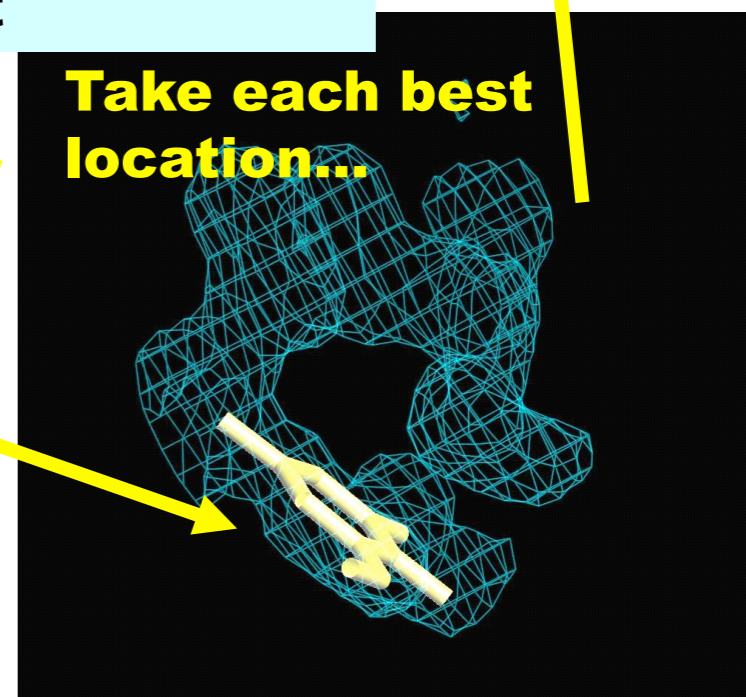
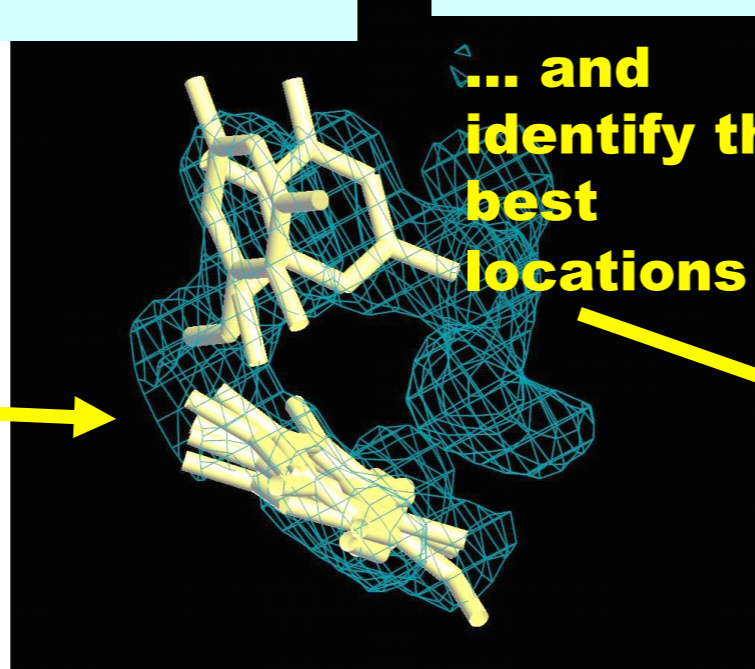
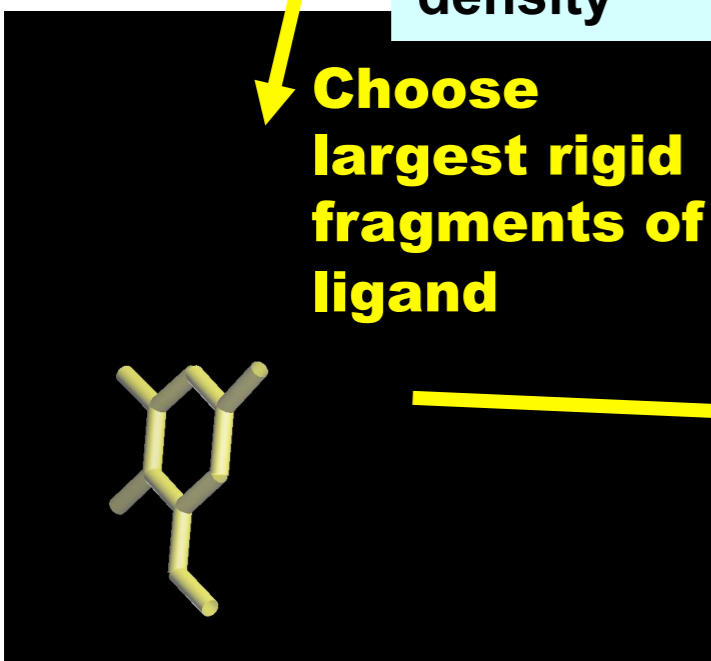
FFT methods are used to rapidly fit fragments to density

The extension uses rotation around torsion angles to find the best fit

Choose largest rigid fragments of ligand

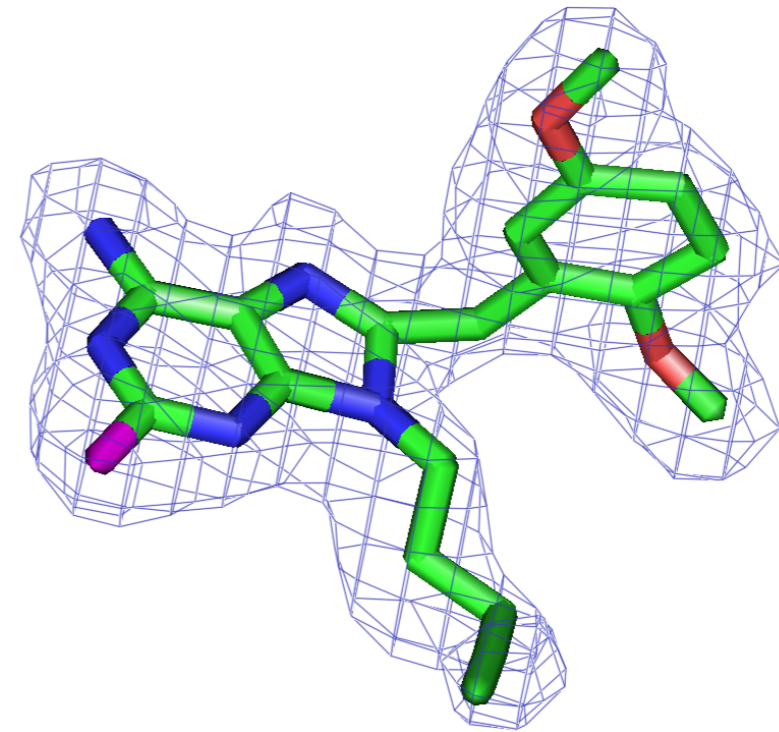
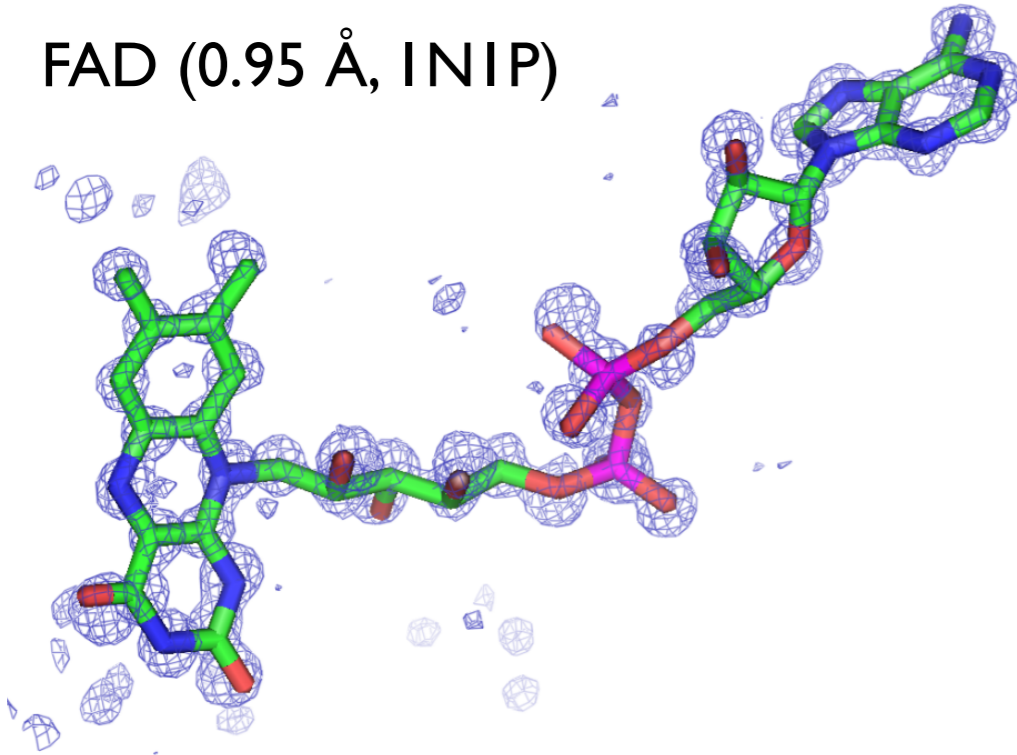
... and identify their best locations

Take each best location...



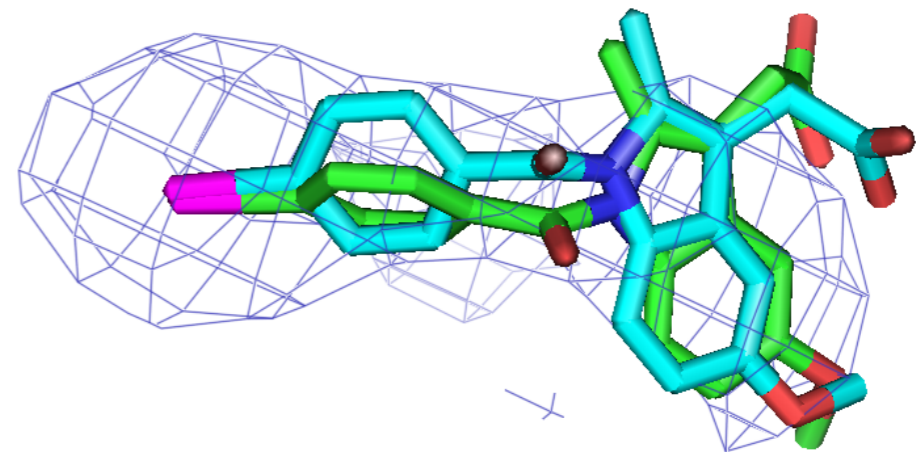
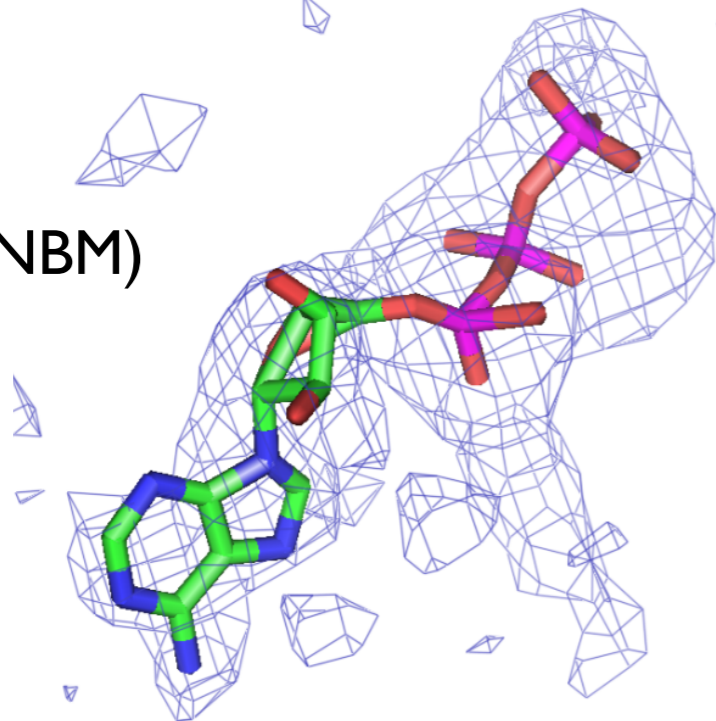
Fitting Over a Range of Resolutions

FAD (0.95 Å, 1NIP)



8-(2,5-DIMETHOXY-BENZYL)-2-FLUORO-9-PENT-9H-PURIN-6-YLAMINE (2.2 Å 1UYI)

ATP (3 Å, 1NBM)



(1-(4-IODOBENZOYL)-5-METHOXY-2-METHYL-INDOLE-3-ACETIC ACID (4.5 Å, 1PGF)

Restraints in phenix.(real_space_)refine

- LINK records have no impact
- Automatically accesses the “standard” residues restraints
- Automatically links the “standard” residues
- Parameter “link_all=True” links
 - Covalent ligands
 - Carbohydrates
 - Metal ions

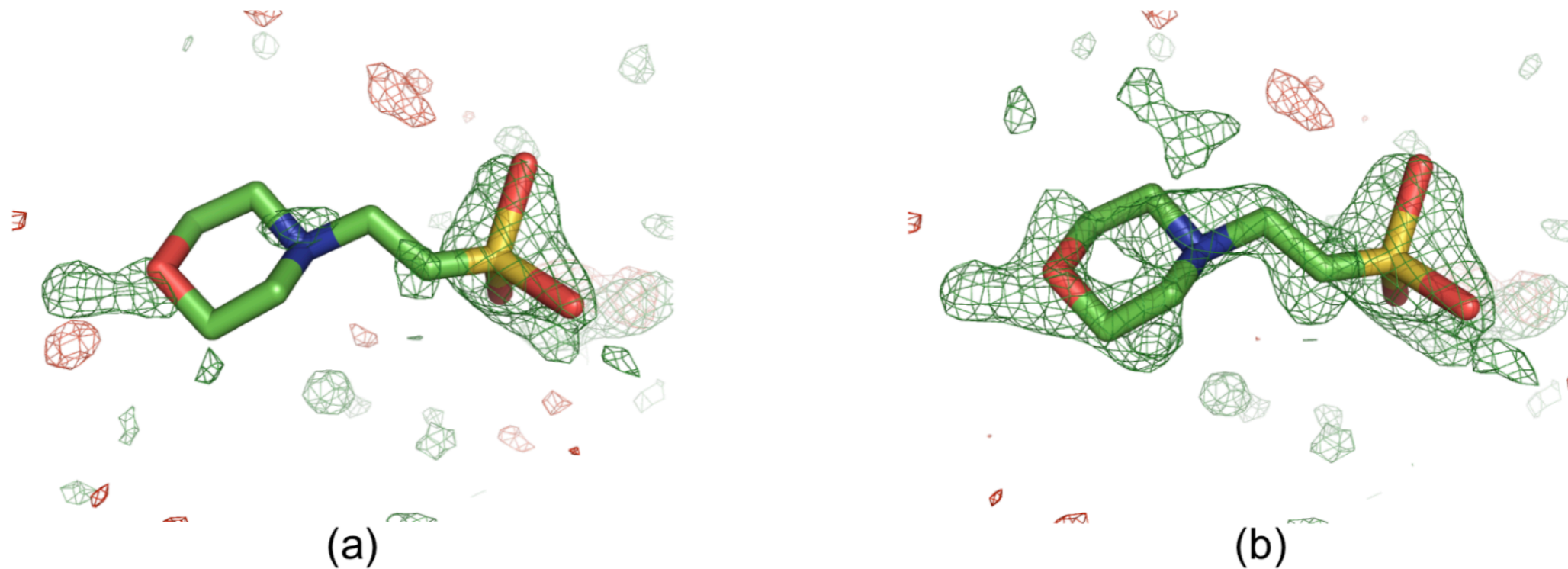
phenix.(real_space_)refine (continued)

- RNA/DNA restraints
 - Base pair hydrogen bonding
 - Base pair planarity
 - Base stacking (parallelity)
- Secondary Structure restraints
- NCS restraints
- Custom bonds & angles using edits
- All restraints used are written to .geo file including non bonded interactions

Ligands? Water?

- Be careful when inheriting a model
- Can contain ligands and water molecules
- Must be able to “see” them in the map

Polder OMIT Maps



Solvent molecule MES88 in structure 1ABA. The positive and negative mFobs-DFmodel OMIT difference density is displayed in green and red, respectively. (a) OMIT map contoured at $\pm 3\sigma$. (b) Polder map contoured at $\pm 3\sigma$. In the OMIT map, there is only some density for the O, N and S atoms. The polder map shows difference electron density for the entire molecule

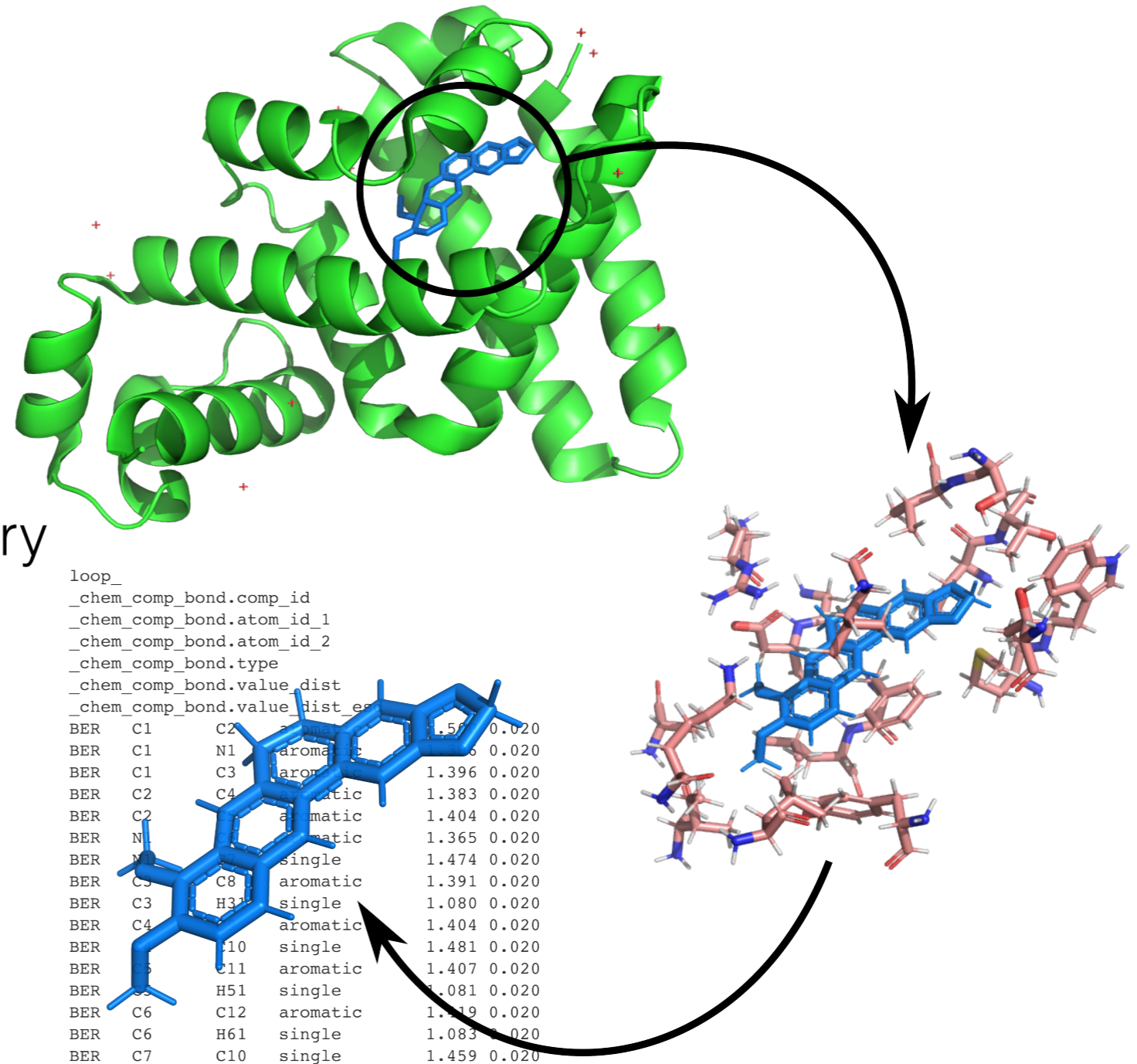
QM Restraints

- Generates restraints of ligands using Quantum Mechanics *in situ*
- Command line only
- There are two ways of using QMR
 - In *phenix.refine*
 - In a standalone program *mmtbx.quantum_interface*
- If you want to use ORCA, need to have ORCA in \$PATH (or set \$PHENIX_ORCA)

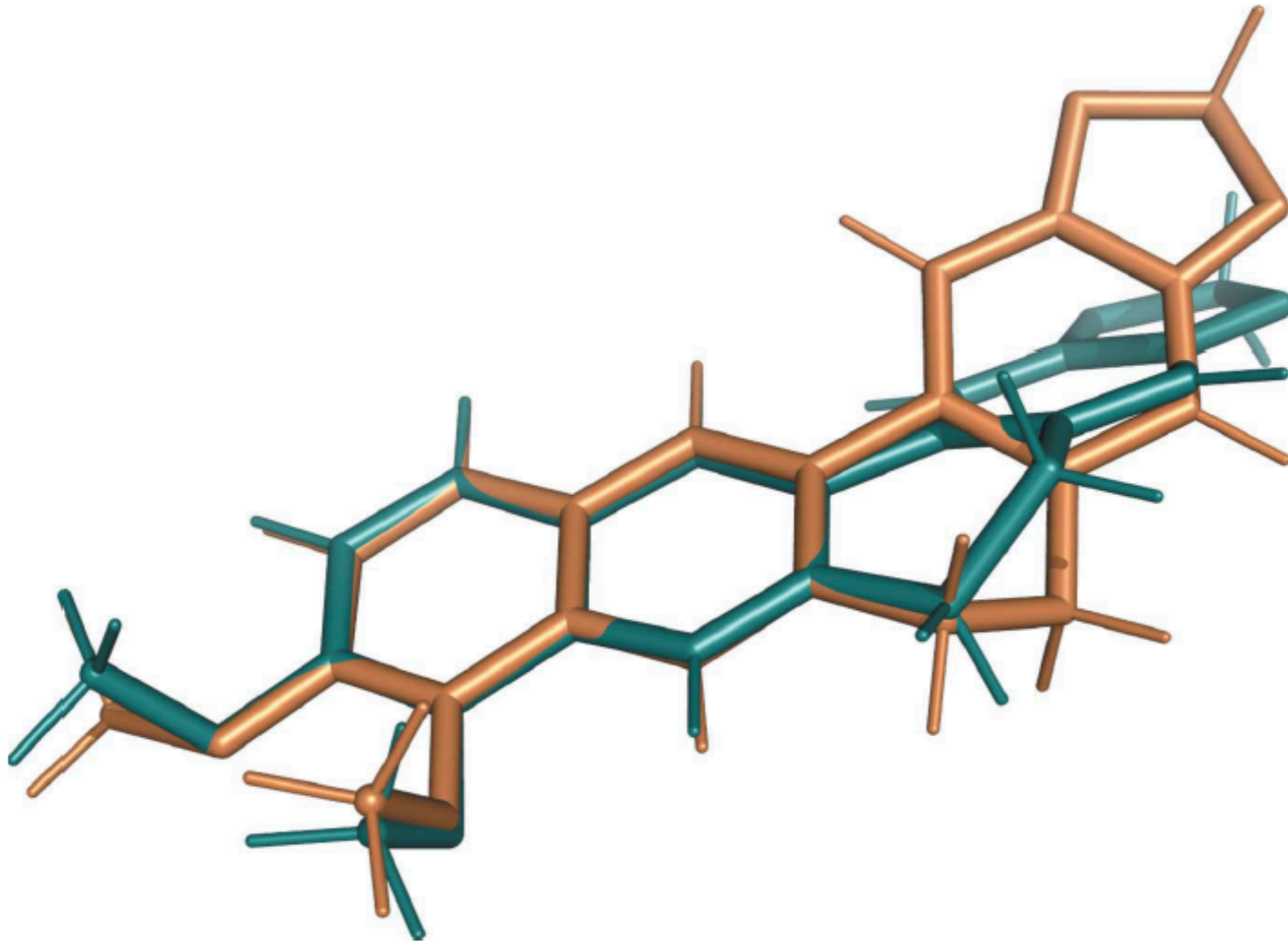
QMR

In situ restraints generation

- Carve out the ligand environment
- Minimise the ligand geometry *in situ*
- Transfer geometry values to restraints (and write to disk)
- Refinement with modified restraints



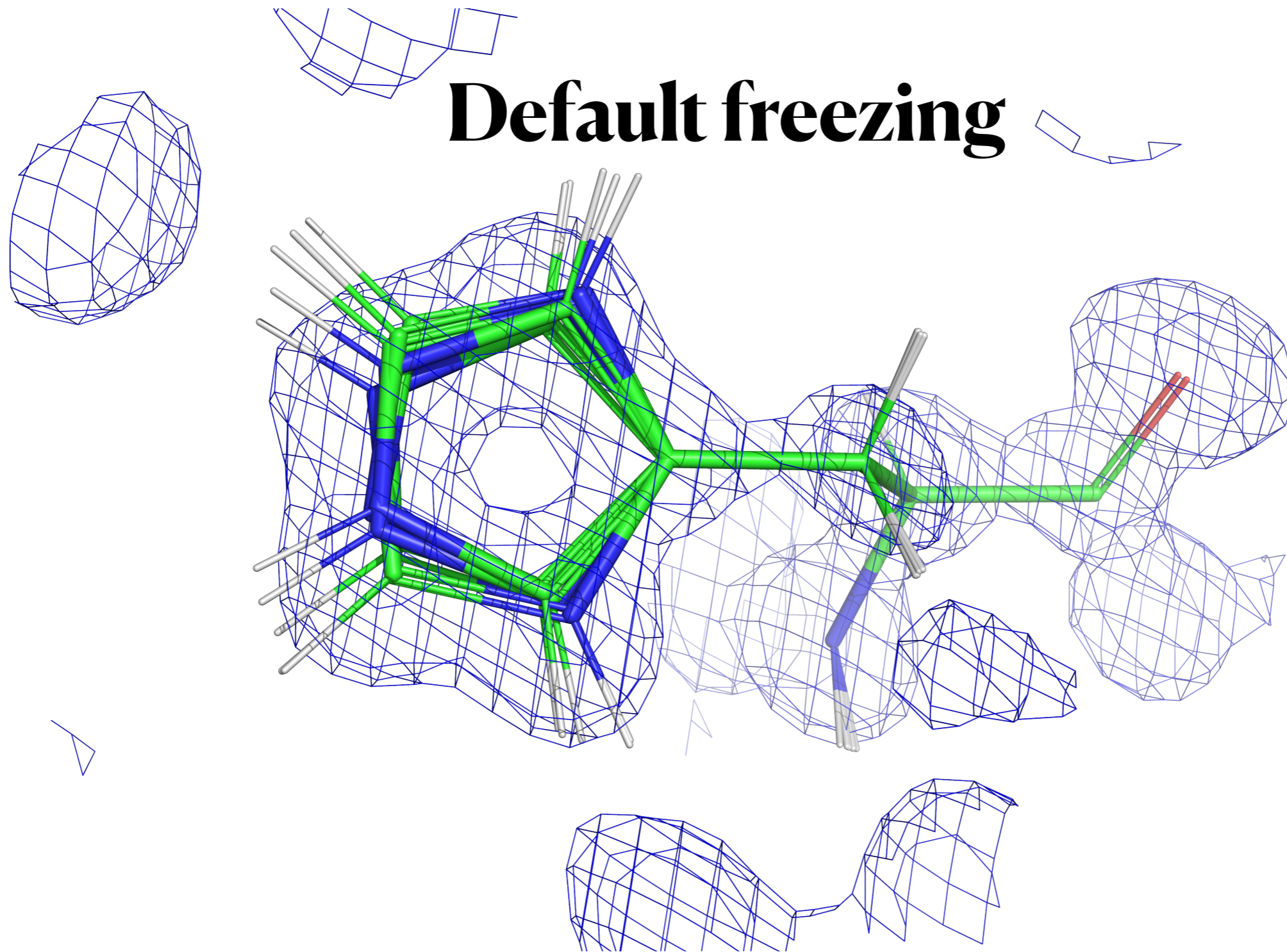
BER in 3vw2



QM Flipping

- Generate the three pronation states of HIS
- Flip χ_2 180 for total of six configurations
- Perform a QM geometry minimisation of the side-chain while freezing the heavy atoms (non-H) for rest of model
- Compare metrics
 - Energy
 - H-bonds
 - RMSD

Default freezing

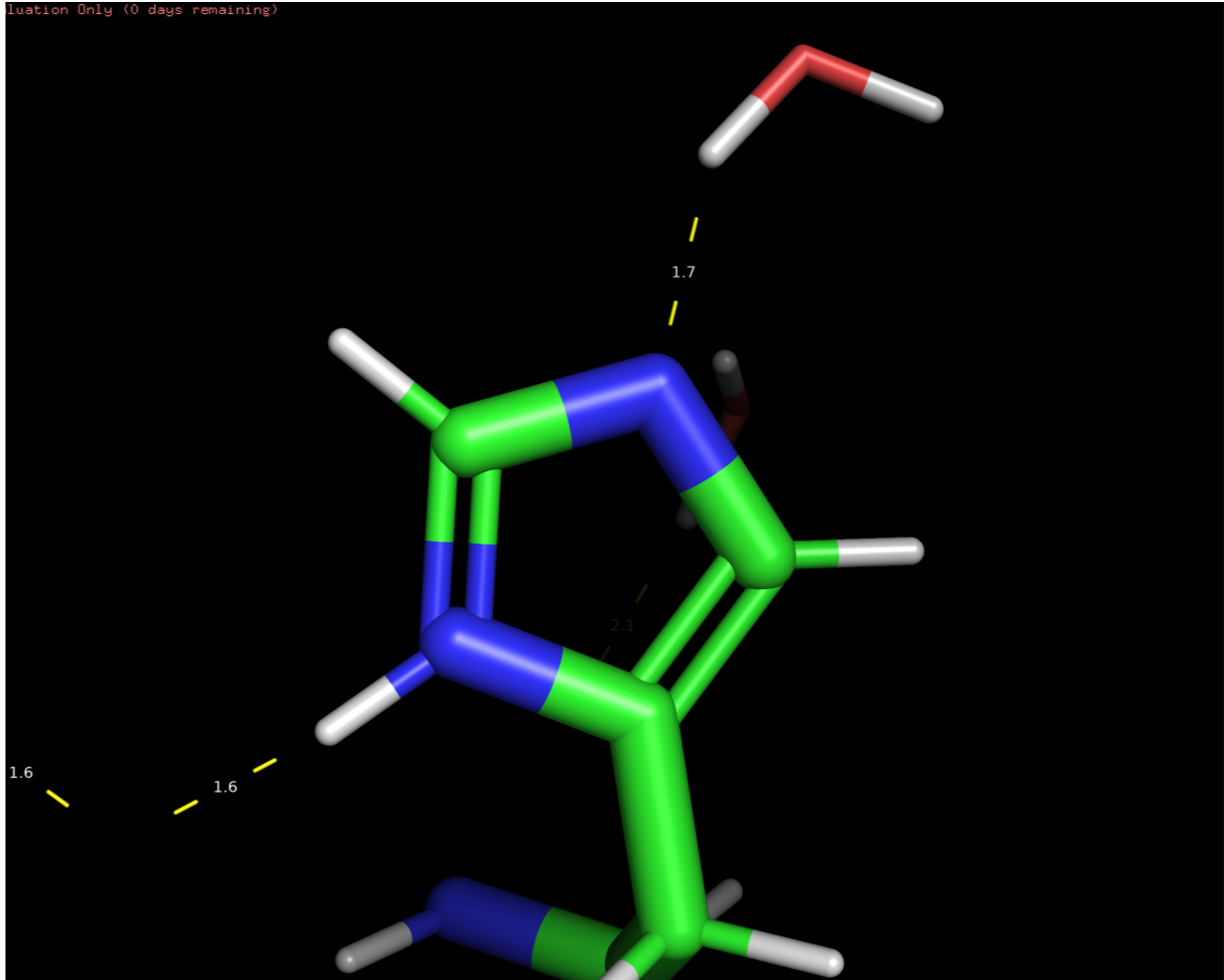
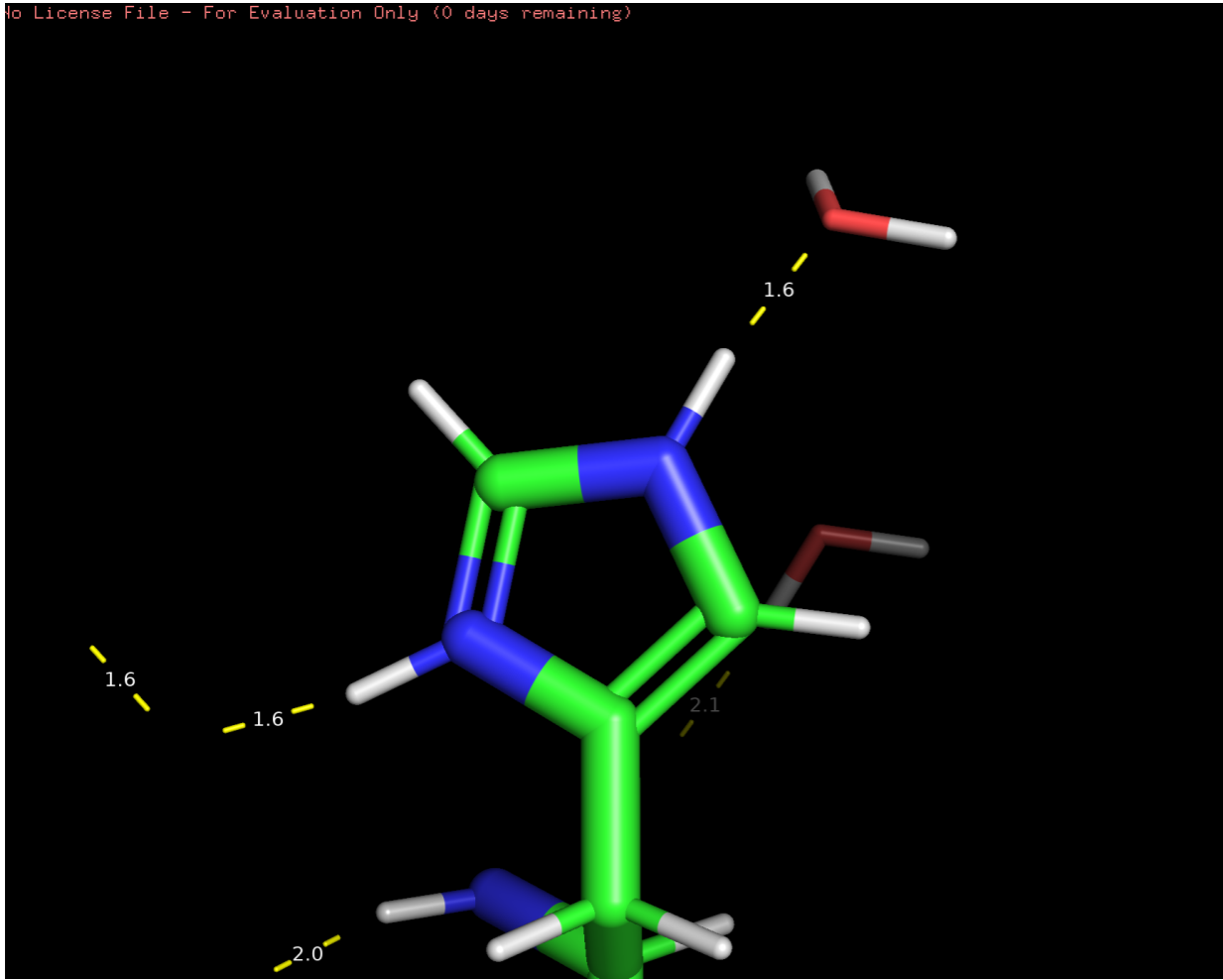


Metrics

| | Configuratio | Energy (kcal/ | ΔE | H bonds | r.m.s.d. | Rotamer |
|---|--------------|---------------|------------|---------|----------|---------|
| 0 | HD1, HE2 | | | | | m90 |
| 1 | HD1, HE2 | -1019,6 | 2,6 | 14 | 0,04 | m90 |
| 2 | HD1 only | -1022,2 | 0,0 | 14 | 0,04 | m90 |
| 3 | HE2 only | -1003,0 | 19,3 | 13 | 0,05 | m90 |
| 4 | HD1, HE2 | -1004,0 | 18,3 | 12 | 0,29 | m-70 |
| 5 | HD1 only | -1004,4 | 17,9 | 12 | 0,38 | m-70 |
| 6 | HE2 only | -1009,4 | 12,9 | 11 | 0,32 | m-70 |

The QMF results for the histidine resseq 4 in chain A of PDB 4rj2

Minimised geometry of two histidine configurations of histidine in chain A and resid 4 of PDB 4rj2.



Summary

- eLBOW & ReadySet! perform better when provided with better input. (GIGO)
- Need to know something about the ligand
 - Hierarchy of input file value
- Check your .geo file for confirmation of restraints