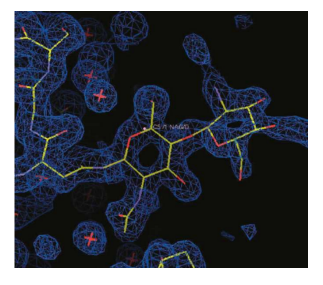
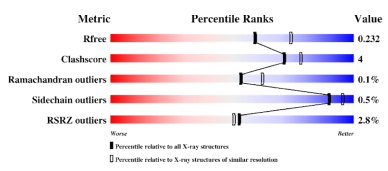
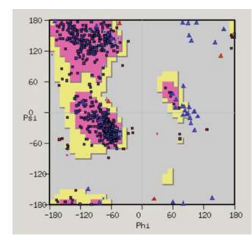
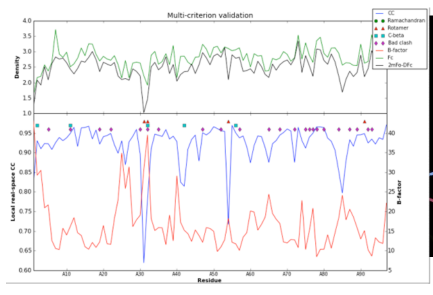
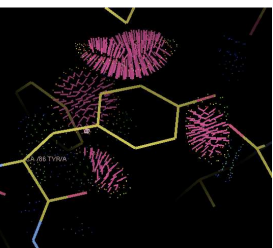
Crystal structure validation







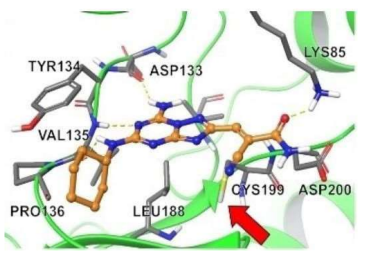




Biocrystallography and Electron Microscopy

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Importance of validation



Once deposited on the PDB, the structure can be used for a range of different studies:

- Good and correct structures are crucial for docking and structure-based drug-design
- Biochemical and mutagenesis experiments are often set up based on structural rationale



- Model bias on calculated electron density
- Low data/parameter
- Overparametrization
- Entrapment of refinement in local minima
- Misinterpretation of electron density



Analysis based on posterior probability and Bayesian statistics:

- Chemical and physical knowledge
- Expectation values for geometric parameters
- Biochemical studies

Improbable ≠ impossible

Bayesian statistics

P(A|B,C,D) Conditional probability: probability that event A occurs given the previous knowledge B,C,D...

$$P(H|E) = \frac{P(E|H)P(H)}{P(E)}$$

Bayes' theorem (inference theorem): the conditional probability of a hypothesis H given the evidence E is the probability of obtaining the evidence E given the model, multiplied by the probability of the model (hypothesis, E) and divided by the probability of the evidence (E).

Crystallographic question:

What is the probability that the protein model is correct, given

- (1) the crystallographic data and
- (2) the chemical/physical knowledge of the system?

Bayesian approach in crystallography

In crystallography:

M model, D crystallographic data,

I previous chemical/physical information

$$P(M|D,I) = \frac{P(D|M,I)P(M|I)}{P(D|I)}$$

- P(D|M,I) likelihood of the data, consistency of the data with the proposed model
- P(M|I) consistency of the model with the previous knowledge
- P(D|I) probability to obtain the dataset, considering previous knowledge (it can be trated as normalization factor and neglected)

The best model is the one for which probability is maximized.

However, rather than maximizing the probability, in the maximum likelihood approach a negative logarithm is minimized:

$$L(M|D,I) = -\log[P(D|M,I)] - \log[P(M|I)]$$

The first term has a close relation with the χ^2 parameter:

$$\chi^2 = \sum_h W_h \left[F_{obs} - F_{calc}(M) \right]^2$$

 W_h can be calculated from the variance evaluated in the previous refinement step (Bayesian approach).

Global and local analyses

Global criteria for structure evaluation:

- R-values and their difference:

$$R_{work} = \frac{\sum_{\boldsymbol{h} \notin free} |F_{obs} - kF_{calc}|}{\sum_{\boldsymbol{h} \notin free} F_{obs}}$$

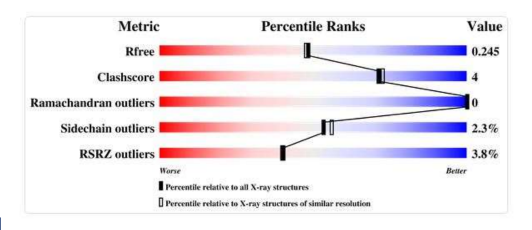
$$R_{free} = \frac{\sum_{\boldsymbol{h} \in free} |F_{obs} - kF_{calc}|}{\sum_{\boldsymbol{h} \in free} F_{obs}}$$

Indication on overfitting or incomplete refinement

Overall r.m.s.d. of bond distances and angles

Depending on restraints used in refinement and resolution





Global and local analyses

Global criteria for structure evaluation:

R-values and their difference:

$$R_{work} = \frac{\sum_{\boldsymbol{h} \notin free} |F_{obs} - kF_{calc}|}{\sum_{\boldsymbol{h} \notin free} F_{obs}}$$

$$R_{free} = \frac{\sum_{\boldsymbol{h} \in free} |F_{obs} - kF_{calc}|}{\sum_{\boldsymbol{h} \in free} F_{obs}}$$

Indication on overfitting or incomplete refinement

Overall r.m.s.d. of bond distances and angles

Depending on restraints used in refinement and resolution

Average B-factor

Local criteria for structure evaluation:

Residue-by-residue analysis of:

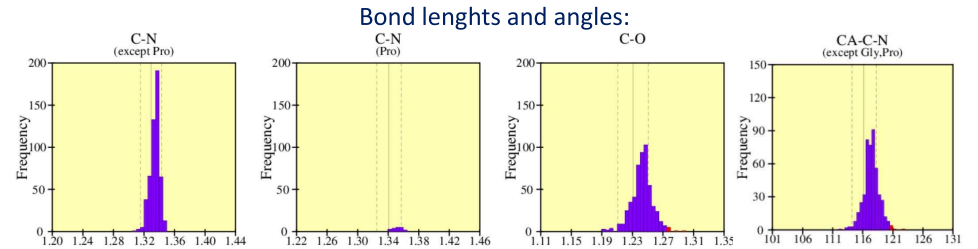
GEOMETRY ANALYSIS

- Geometric parameters: deviation from expected values
- Ramachandran plot
- Non-bonding contacts closer than the van der Waals radii of the atoms (clashes)

ELECTRON DENSITY ANALYSIS

- Real space R-value
- Fitting of the model in the electron density: Real Space Cross-correlation Coefficient
- B-factor values for each residue

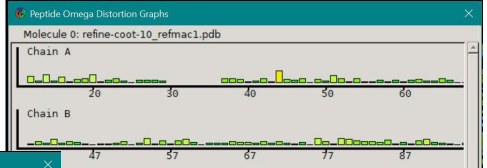
Geometry analysis: bonds and angles

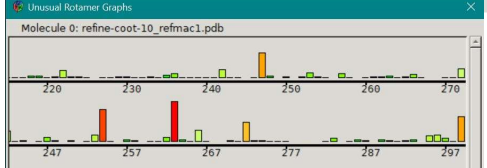


Values measured on the model compared with expectation values and their standard deviations.

In addition:

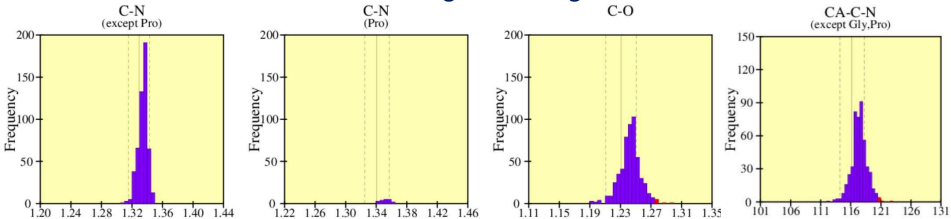
- ω angle (peptide bond dihedral angle)
- Side chain conformations





Geometry analysis: bonds and angles

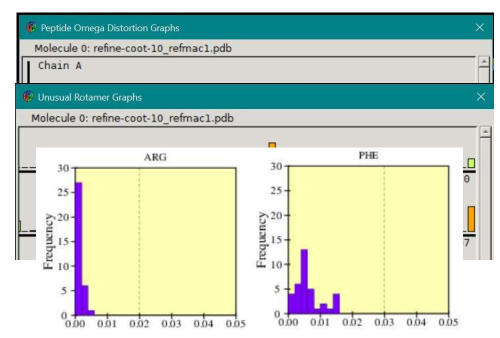




Values measured on the model compared with expectation values and their standard deviations.

In addition:

- ω angle (peptide bond dihedral angle)
- Side chain conformations
- Planarity of side chains
- •

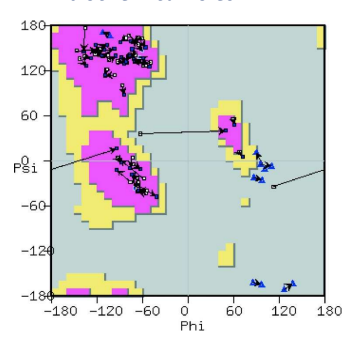


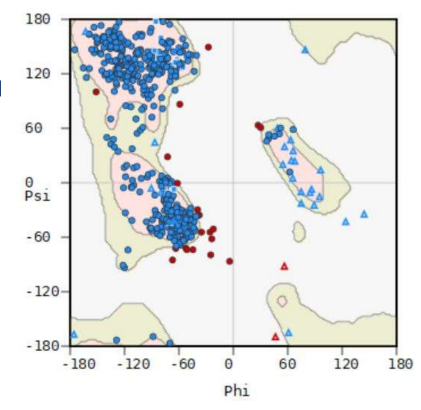
Geometry analysis: Ramachandran plots

Usually, angles ϕ and ψ are not restrained during refinement \rightarrow good for cross-validation!

Ramachandran violations are possible, but should be carefully analyzed:

- 1. Evaluate electron density of the residue (often Ramachandran violations in disordered loops poorly modeled)
- 2. Analyze similar structures: strained conformations may have important biochemical roles





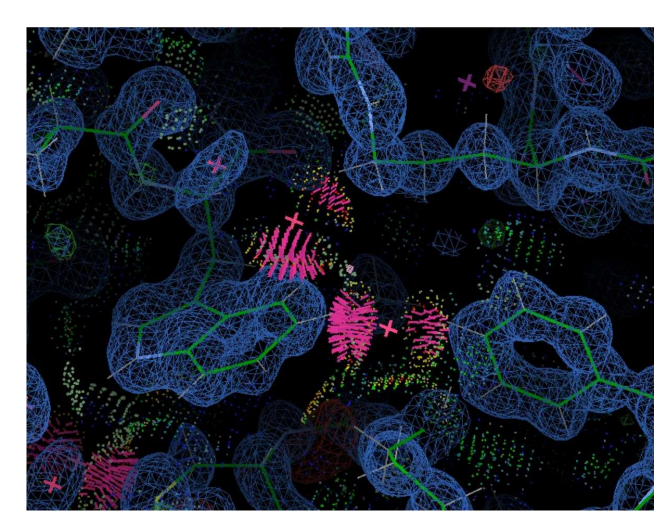
Kleywegt plots: analysis of difference in backbone dihedral angles for residues of NCS related molecules.

Geometry analysis: clashes

Non-bonding contacts shorter than van der Waals radii of atoms.

Software can recognize hydrogen bonding networks and identify hydrophobic close contacts.

Particularly dangerous contacts with symmetry related molecules.



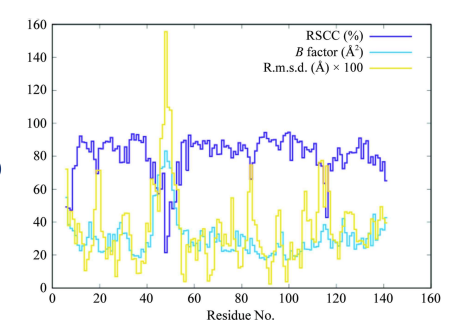
Electron density validation

Real Space R-value (RSR):

$$RSR = \frac{\sum_{r} |\rho_{obs}(r) - \rho_{calc}(r)|}{\sum_{r} |\rho_{obs}(r) + \rho_{calc}(r)|}$$

with $\rho_{obs} = \sum_{r} (2F_{obs} - F_{calc}) \cdot \exp(-i\varphi_{calc})$ and $\rho_{calc} = \sum_{r} F_{calc} \cdot \exp(-i\varphi_{calc})$ for each voxel of the real space (r)

Values are plotted residue by residue, often together with the B-factor value.



Real Space Cross-correlation Coefficient (RSCC):

$$RSCC = \frac{\sum_{\boldsymbol{r}} \left[\rho_{obs}(\boldsymbol{r}) - \overline{\rho_{obs}(\boldsymbol{r})} \right] \cdot \left[\rho_{calc}(\boldsymbol{r}) - \overline{\rho_{calc}(\boldsymbol{r})} \right]}{\left(\sum_{\boldsymbol{r}} \left[\rho_{obs}(\boldsymbol{r}) - \overline{\rho_{obs}(\boldsymbol{r})} \right]^{2} \cdot \sum_{\boldsymbol{r}} \left[\rho_{calc}(\boldsymbol{r}) - \overline{\rho_{calc}(\boldsymbol{r})} \right]^{2} \right)^{1/2}}$$

Visual analysis of maps: bias-minimized maps, omit maps (using calculated phases and amplitudes from a model where a specific portion was removed).

Ligands

For known ligands:

Geometric restraint file are available for the main ligands

For new ligands:

Geometric restraints should be generated considering chemical knowledge (e.g. ibridization state, bond leght and angles...)

Omit maps and difference maps are extremely useful tools to confirm the presence of ligands.

Often, ligands occupy the binding site only in a percentage of proteins. **Partial occupancy** should be considered together with **B-factor values**.

Careful analysis of ligand contacts (**LIGPLOT**):

