Ultra-high-resolution crystallographic evidence for protein geometry variation

# Introduction

Geometric parameters are essential for the refinement and prediction of protein structure. The original parameters (E&H 1991, 2001) were derived from the CSD. Since then there have been some reviews of these parameters (Jaskolski, 2007) based on the increasing resolution of x-ray refinement to sub-atomic resolution. As the resolution increases, there is the possibility of relaxation of refinement parameters to give a greater weight to experimental evidence. However, as E&H say “protein structures are generally solved not to build a statistically optimised protein database, but to discover biophysical functional mechanisms” (E&H 2001). We find ourselves then in an infinite spiral of looking for parameters from structures solved with the parameters.

In this study, we seek to sever this spiral by building a statistically optimised database of geometric evidence based on pure experimental evidence from the electron density of ultra-high resolution x-ray crystallographic structures solved at <=1 We decouple ourselves from the solved structures and go straight to the electron density for all atomic positions. In doing this we are freed from the problem of outliers: no cut off is required, all outliers are evidential in the electron density and thus all outliers provide us with geometric insights.

# Method

## Electron Density Topology

A method of finding: values, first and second derivatives from the electron density has been derived to enable both mathematical and visual analysis. The electron density grid points are interpolated using a multivariate interpolation method with Vandermonde matrices.

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| Values | Radiant | Hessian Eigenvalues |
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Figure 1 - 1us0 TYR A48, calculated with a 5th degree multivariate interpolating polynomial at 0.05Å samples. In the radiant image the atoms on the plane can be seen clearly by the black crosses – OH is not quite planar. The Hessian is calculated by the number of -ve or +ve eigenvalues of the Hessian. 3 negative values are a maximum and dark blue. 3 positive values, red, is a minimum. These are planar slices of 3d calculations.

The accuracy of the values can be determined by increasing the degree of the interpolated multivariate polynomial (stable by this method to a 7-degree polynomial) and the sample frequency of the interpolated points. In general, we find that a 3rd degree polynomial is sufficient mathematically, but a 5th degree polynomial is better visually. For visual images we sample at 0.05Å. The visual images can be used to verify and determine the features we find mathematically – the values show us the electron density itself, but the most striking visual images are the images of the 1st derivative. We use the L1 norm of Del of the 1st partial derivatives to produce images which we have coined the radiant of the electron density – due obviously to the 1st derivative being a gradient, but also the way the coordinate axes lines radiate out from the maxima enabling the identification of those maxima from both the circular black centres of atoms and from these lines that radiate out (see Figure 1).

## The generation of a defensible dataset

A set of pdb structures is chosen as being <=1 non-homologous at 90%, no nucleotides and >40 residues. We look at the original pdb data in these sets in 2 ways – UNRESTRICTED is all atoms, RESTRICTED is only those atoms that have single occupancy and whose bfactor is less than 1.3 x the average bfactor for the structure (CA atom of the residue).

Then, for each pdb structure, we choose a set of atoms which we wish to be evidential for geometric purposes, for the backbone geometric analysis we have chosen atoms of the types N, CA, C and O. For every atom of this type in each pdb, we look at the coordinates in the electron density and find at that position the value and the second derivative which is calculated via the 2nd partial derivatives and the eigen values of the hessian, where we choose whether the eigen values are 3 negatives to mark a maximum (Bader). If the atom is not at a maximum, we delete it from the pdb file. If the atom is a maximum, we build a cube around the point and look at all the values of the vertices – if any are bigger, we reject the point as not being a maximum, again deleted. The wider the cube the more permissive, several sets of data have been analysed at different widths.

We now have 2 files for every pdb structure – the pdb file that contains only defensible atoms, and a file of “bad” atoms. To examine whether these have given us a genuine separation of good and bad experimental evidence we need a more automated check of the electron density (manually loading every pdb in chimera is too time consuming). For this we have visualised the electron density’s first derivative via the radiant images. Using this technique with an automated tool we can generate images of all the “bad” atoms to verify they are not defensible. Additionally, once the geometry has been calculated we can look at all the outliers to verify that despite the unusual geometry the atoms are indeed where they are purported to be.

# Results

## Datasets

5 sets of pdb files were generated, based on electron density grid point fractions of 0.1, 0.2, 0.25, 0.3, 0.4 and 0.5. No data was calculated at 0.1 – too few atoms were found to be maxima and that tight interval to generate any geometry. At 0.5

## Radiant Images of Rejected Atoms

Some examples from 6q4g:

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## Radiant Images of Accepted Outliers

Some examples from set B03 for N:CA outliers

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| Min N:CA=1.389 | Max N:CA=1.501 | Min N:CA=1.405 | Max N:CA=1.450 |
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| Min N:CA=1.418 | Max N:CA=1.510 | Min N:CA=1.390 | Max N:CA=1.498 |
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## Statistical Analysis

We compare the datasets we get to the original data to check that it is representative as a distribution.

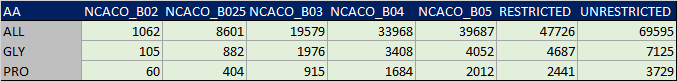
[Results]

## A comparison of sets

For each of the sets we compare the means and sd against the E&H values. We are confident that the sets give similar values.

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These are the observation counts we obtain for each of the sets.



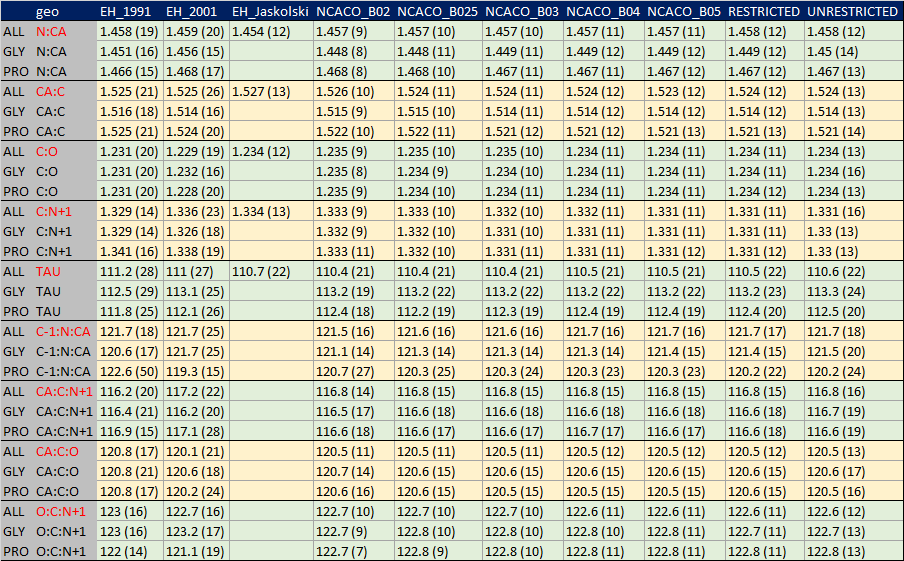
## Some histograms

For each of the E&H bond lengths and angles, a histogram is generated with the outliers specified. These are compared to the E&H recommendations and the later Jaskolski recommendations. We have good values across the datasets so we will concentrate on the B03 set as a middle ground. All the data can be found here: [Summary B03 E&H](https://rachelalcraft.github.io/PaperTau/Evidence/EhHistograms/Defensible_EH_NCACO_B03.html) and [Per Amino Acid B03 E&H](https://rachelalcraft.github.io/PaperTau/Evidence/EhHistograms/Defensible_EH_AA_NCACO_B03.html)

Some examples here:

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## A table of results



# Tau Variation

[An exploration of tau variation, start with the nice plot psi vs N:N+1 ]

# Discussion

* 1. So that's interesting - these are the new geometric recommendations.
  2. And, the outliers are interesting too. Most of them look fine.
  3. In the future we could:
     1. look at the spreads of various parameters and use the defensible data to explore what causes the spreads: we might start with tau, based on this N:N+1 rainbow pictures and the knowledge of the spreads
     2. We might also look at outliers and see whether unusual geometry is defensible or dodgy. Are there any fabled undiscovered active sites? Are there experimental problems?
     3. We could create parameters on an ad-hoc basis for anything anyone might be interested in.
     4. Re-refine the structures and look at them again. \*E&H very end of paper, last point).
  4. In conclusion, different geometric parameters are recommended here which are defensible with low sd.

# References

Engh, R A, and R Huber. ‘18.3. Structure Quality and Target Parameters’, n.d., 12.

Jaskolski, Mariusz, Miroslaw Gilski, Zbigniew Dauter, and Alexander Wlodawer. ‘Stereochemical Restraints Revisited: How Accurate Are Refinement Targets and How Much Should Protein Structures Be Allowed to Deviate from Them?’ *Acta Crystallographica Section D Biological Crystallography* 63, no. 5 (1 May 2007): 611–20. <https://doi.org/10.1107/S090744490700978X>.