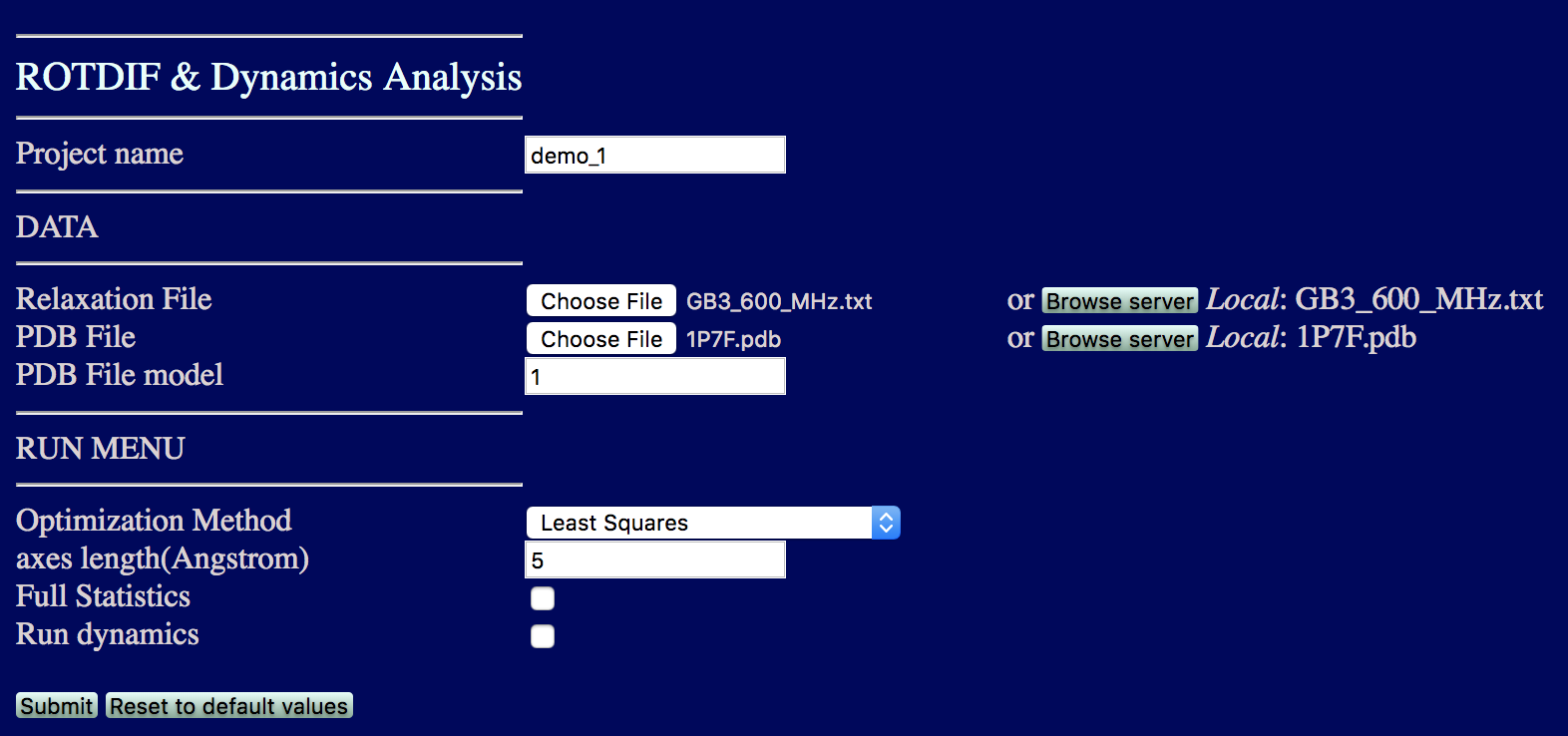
## **ROTDIF & Dynamics Analysis Module**

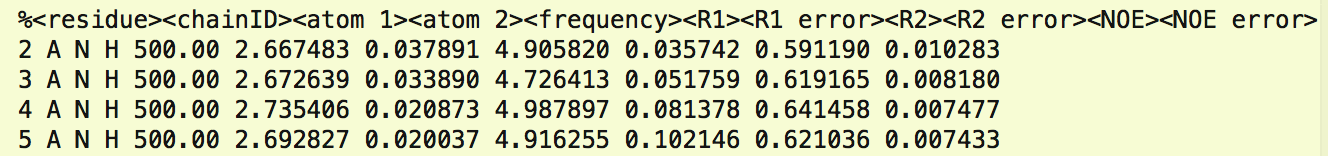
This module allows you to determine the overall rotational diffusion tensor from the experimental spin-relaxation data and to characterize local dynamics in proteins and nucleic acids.

Click the button “ROTDIF & Dynamics Analysis”; the interface will look like this:

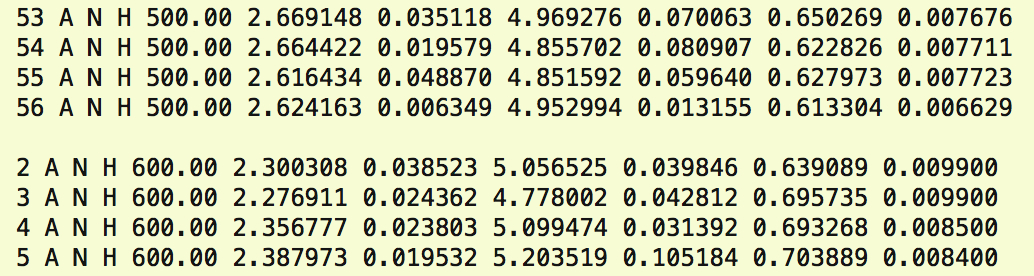


* **Project name:** The name of the folder to store all the results of the current session
* **Relaxation File:** Upload the file (e.g., GB3\_600\_MHz.txt) containing experimental relaxation data. The data file is a text (ascii) file with the following format.

**Format:**



If the input file includes relaxation data measured at multiple frequencies, please leave a blank line between the data from different fields, as shown below:



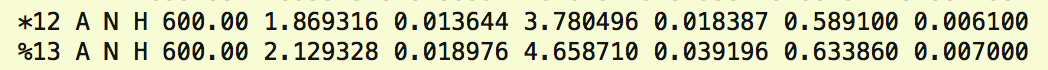
Here “A” is the “chainID” in the coordinate file, “frequency” is the 1H frequency of the spectrometer, “N” and “H” indicate the atom pair in which the relaxation of the first one (15N) was measured; R1 and R2 are the longitudinal and transverse, respectively, 15N autorelaxation rates, and NOE is the steady-state heteronuclear 15N{1H} NOE, the columns marked “R1 error” etc. correspond to the errors in the respective experimental data.

If there are unwanted residues in the data file, they can be either removed or commented out by placing % at the very beginning of the corresponding line. Two types of comments can be used:

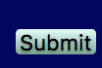
% designates residues that should be completely ignored (the same as deleting the line)  
\* designates residues that should be excluded from the determination of the diffusion tensor but will be used when determining parameters for the local dynamics.

These are typically residues affected by conformational/chemical exchange (resulting in elevated R2 values) or those residues where the molecular structure is not well defined (e.g., due to increased flexibility or other issues), as well as other outliers.

An example of comments:

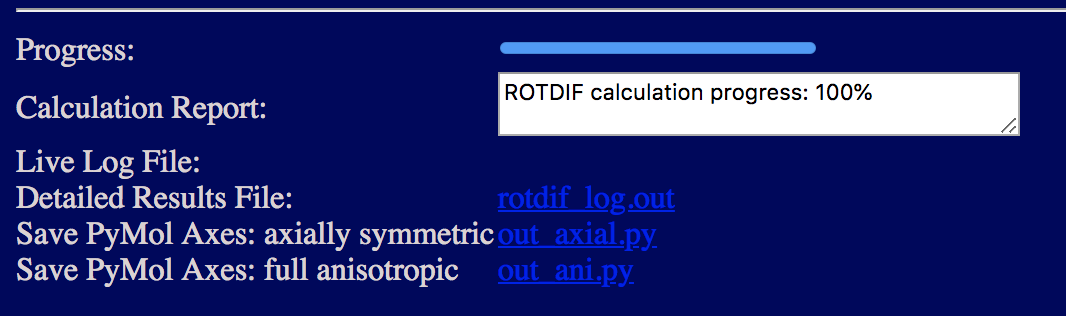


* **PDB File:** Upload the coordinate file (e.g., 1P7F.pdb).
* **PDB File model:** If the coordinate file contains more than one structural model, please specify the model you want to analyze. The default model number is 1.
* **Optimization Method:** Choose the optimization method: ”Least Squares” or “Robust Least Squares (3.0sig)”. Least Squares is the conventional minimization method, while the Robust Least Squares, introduced in ROTDIF 3, dampens the contributions of residuals that are greater than 3 (above 99.7 percentile)(see our publication [2] for more details).
* **Axes length:** Please specify according to the size of proteins. The default is 5 Angstrom. This parameter sets the desired length of the diffusion tensor axes, to be stored through the \*.py files and visualized in PyMol. If the axis length is too short (shorter than the size of the protein) it might not be clearly visible when shown together with the protein.
* **Full Statistics:** Check this box if you want to perform error analysis of the results. The program will generate synthetic (Monte-Carlo) data sets based on the experimental errors in the relaxation rates, for each set perform the same analysis, and produce a full statistics report.
* **Run Dynamics:** Check this box if you want to perform model-free analysis of internal motions in the protein.

Let us fill all the fields with 1P7F.pdb and GB3\_600\_MHz.txt. and click 

Caution: it may take a long time to run the calculations with “Full Statistics” checked. If this is not necessary, we suggest to leave this box unchecked.

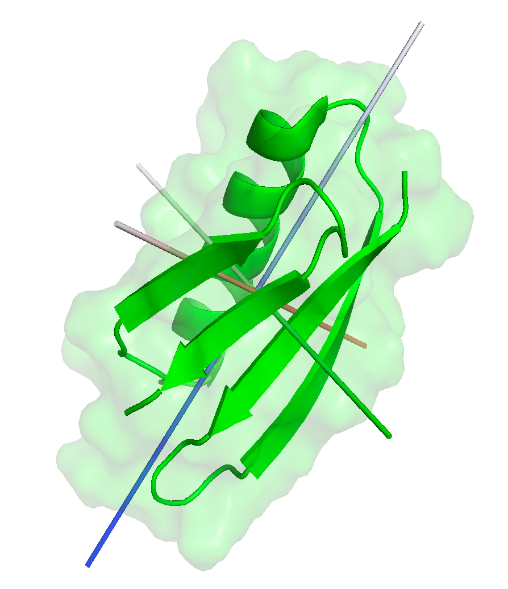
After the progress is 100%, you should be able to see all the output files and figures.



You can download the following files:

* Detailed Results File: the .out file records all the data used to generate the plots shown below
* Saved PyMol Axes: axially symmetric model: the out\_axial.py file (when run in PyMol) builds axes based on the results of the axially symmetric rotational diffusion model
* Saved PyMol Axes: fully anisotropic model: the out\_ani.py file (when run in PyMol) builds axes based on fully anisotropic rotational diffusion model

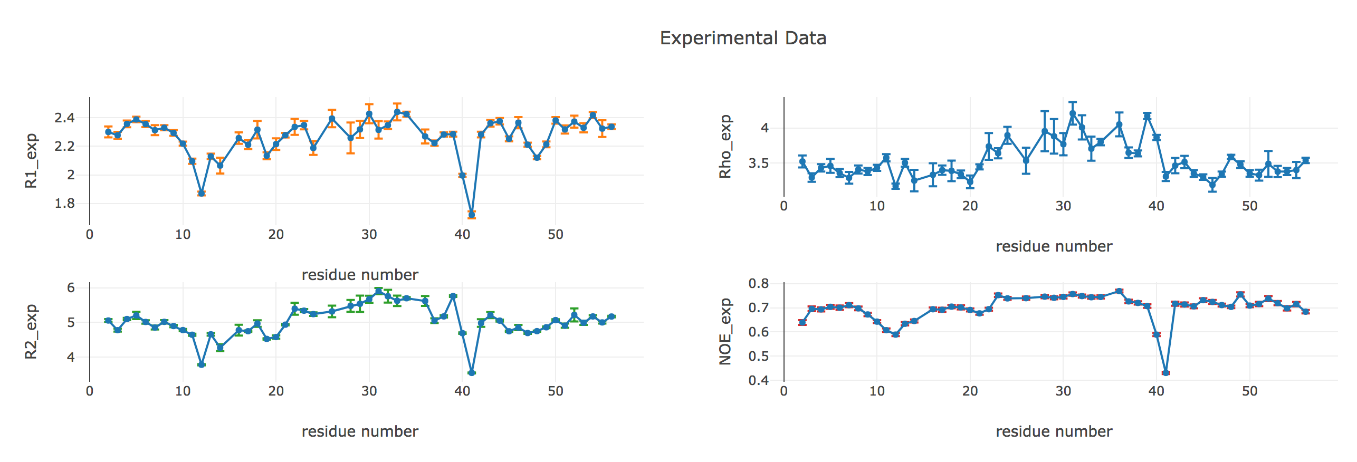
You can visualize the axes encoded in the \*.py files in PyMol together with your protein (outside ROTDIF-Web). The model below is visualized by PyMol.



Once the calculation is finished, the program will output the following plots:

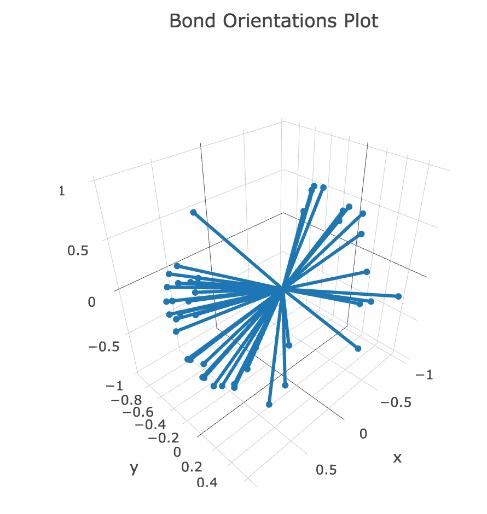
**Figure 1. Experimental Data.**

This plot shows input experimental data (with error bars) as a function of residue number. Shown are data for all residues that are not marked with %. The parameter  (Rho) represents the ratio of the modified relaxation rates – see [2]. You can use the mouse/cursor to read the values for individual residues.



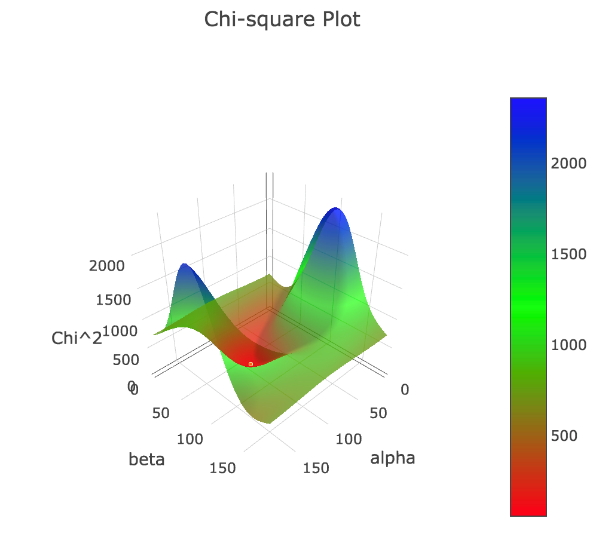
**Figure 2. Bond Orientations Plot**

This 3D plot shows the distribution of bond orientations (represented by unit vectors) for the residues included in the analysis, to allow visual inspection of how well these vectors sample the orientational space. The sampling tensor quantifying the degree of orientational sampling will be included in the Detailed Results output file. Uniform sampling of the orientations could be important for accurate analysis of the diffusion tensor – see publication [10] for more details. Use the mouse to rotate the plot or zoom.



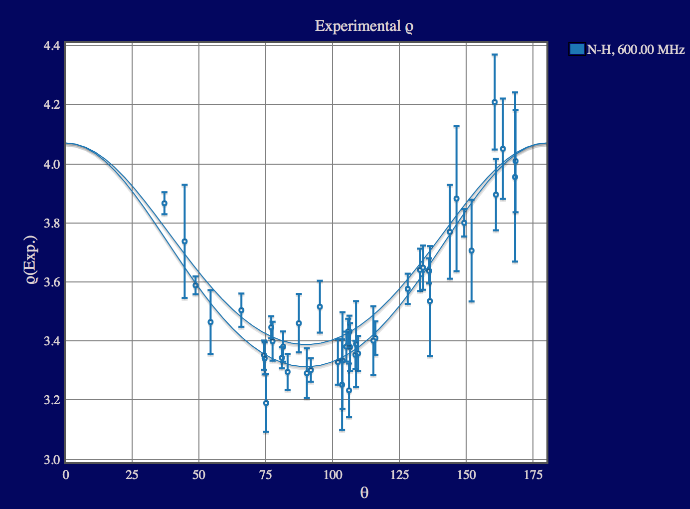
**Figure 3. Chi-square Plot**

This 3D plot shows the resulting 2 value as the function of the Euler angles alpha and beta that determine the orientation of the principal axes of the overall rotational diffusion tensor (axially symmetric model) in the coordinate frame of the protein. Use the mouse to rotate the plot or zoom.



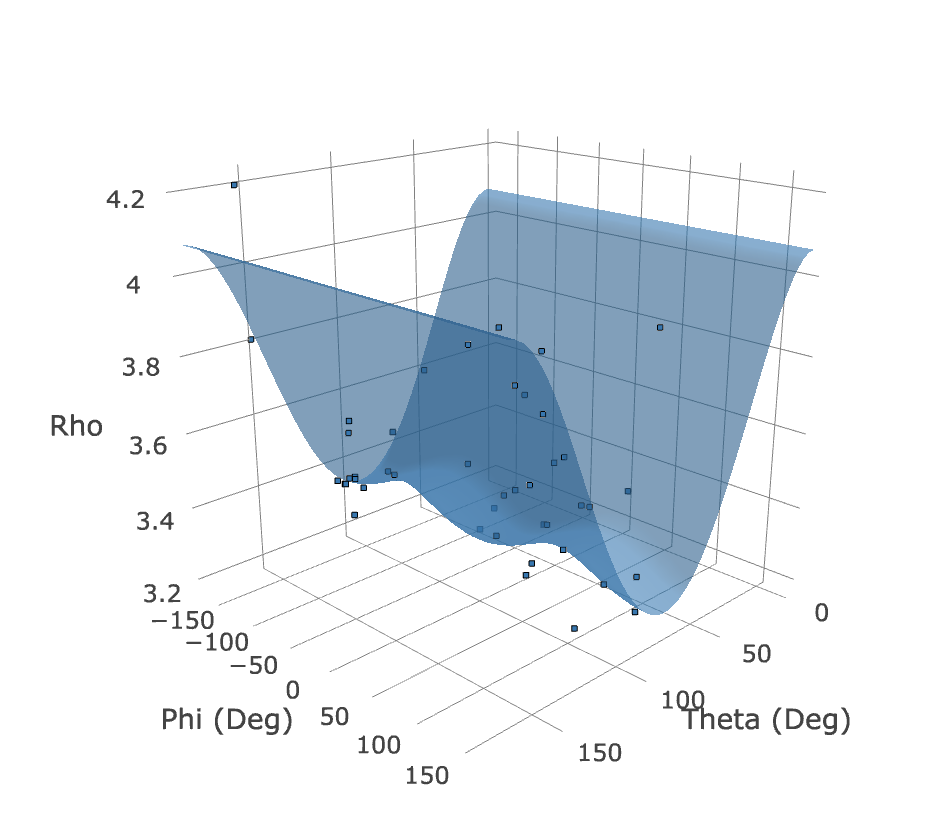
**Figure 4. 2D Model Fit**

This 2D plot shows the  values for the individual residues as the function of the angle  between the corresponding bond and the z-axis of the rotational diffusion tensor (axially symmetric model). The curves represent the upper and lower bounds of the theoretical dependence of  as a function  for the derived fully anisotropic diffusion tensor (see Figure 5 and publication [1] for details). You can use the mouse/cursor to read the values for individual residues.



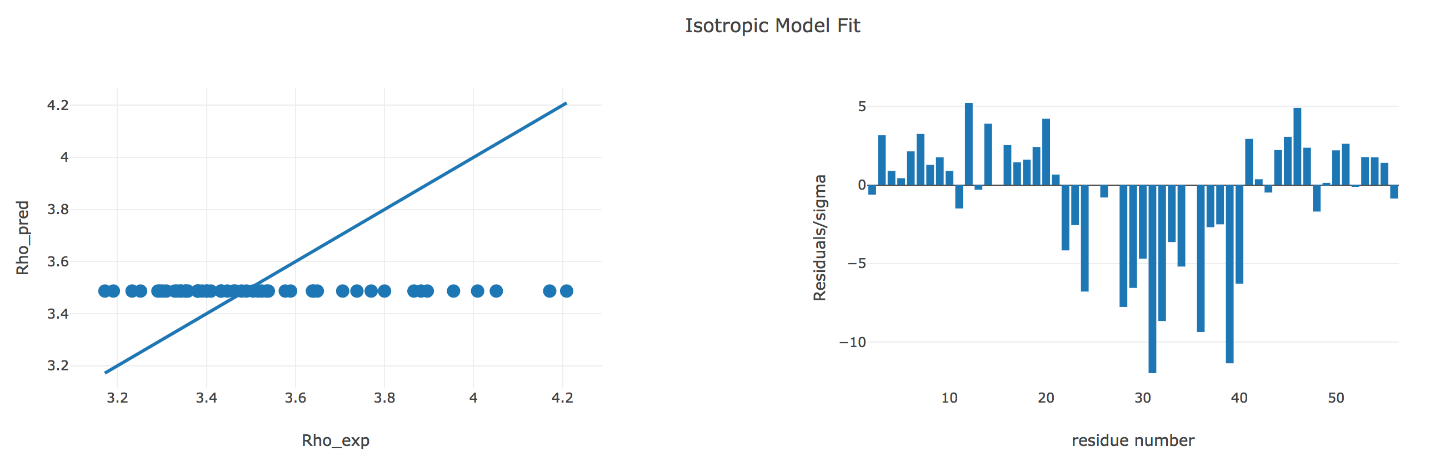
**Figure 5. 3D Model Fit**

This 3D plot shows the  values for the individual residues as the function of the angles  (theta) and  (phi) that determine the orientation of each bond with respect to the principal axes of the derived rotational diffusion tensor (fully anisotropic model). The surface represents the theoretical dependence of  as a function of  and  for the resulting diffusion tensor; the dots represent experimental data points. Use the mouse to rotate the plot or zoom. You can use the mouse/cursor to read the values for individual data points.



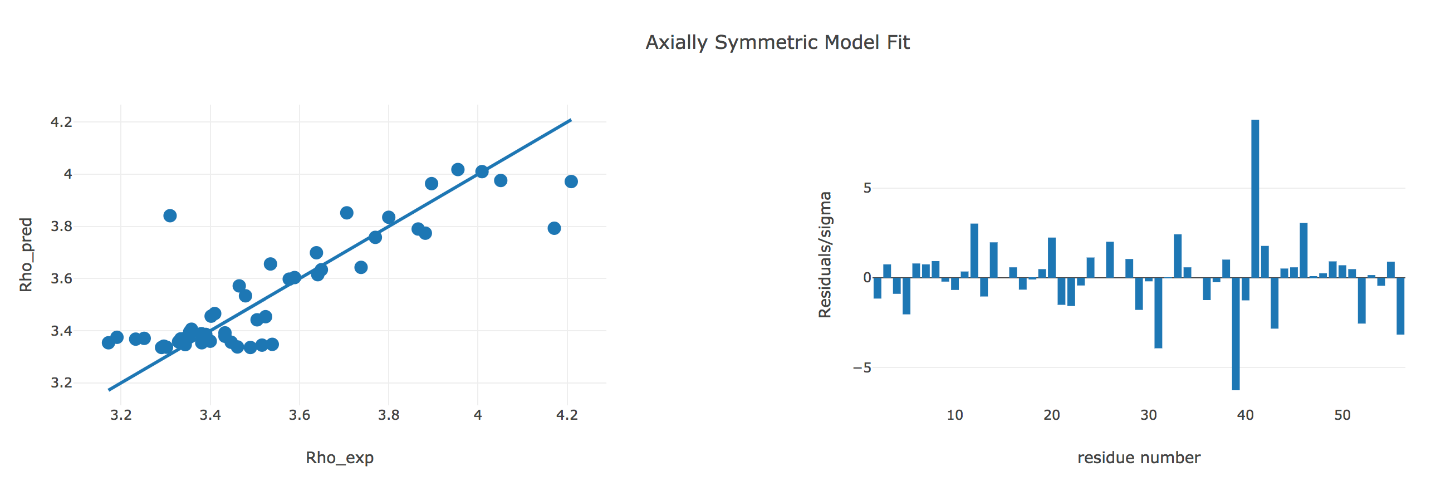
**Figure 6. Isotropic Model Fit**

These plots illustrate how well or not the experimental data agree with the isotropic rotational diffusion tensor model. Shown on the left is the agreement between the experimental and back-calculated/predicted values of ; the line corresponds to absolute agreement. Shown on the right are the normalized residuals of fit for each residue. You can use the mouse/cursor to read the values for individual residues.



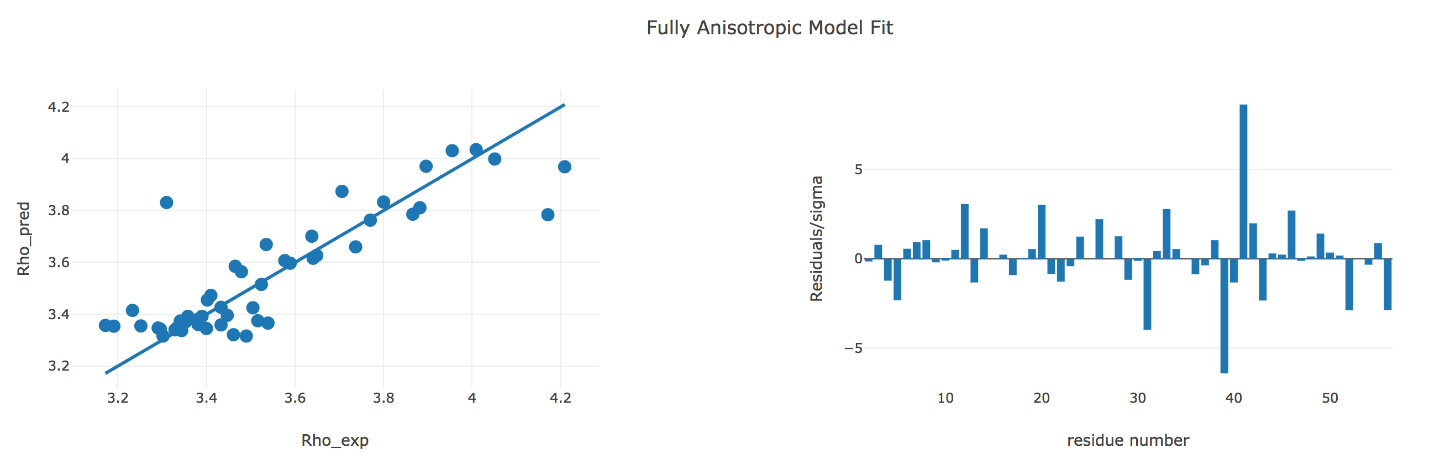
**Figure 7. Axially Symmetric Model Fit**

These plots illustrate how well or not the experimental data agree with the axially symmetric rotational diffusion tensor model. Shown on the left is the agreement between the experimental and back-calculated/predicted values of ; the line corresponds to absolute agreement. Shown on the right are the normalized residuals of fit for each residue. You can use the mouse/cursor to read the values for individual residues.



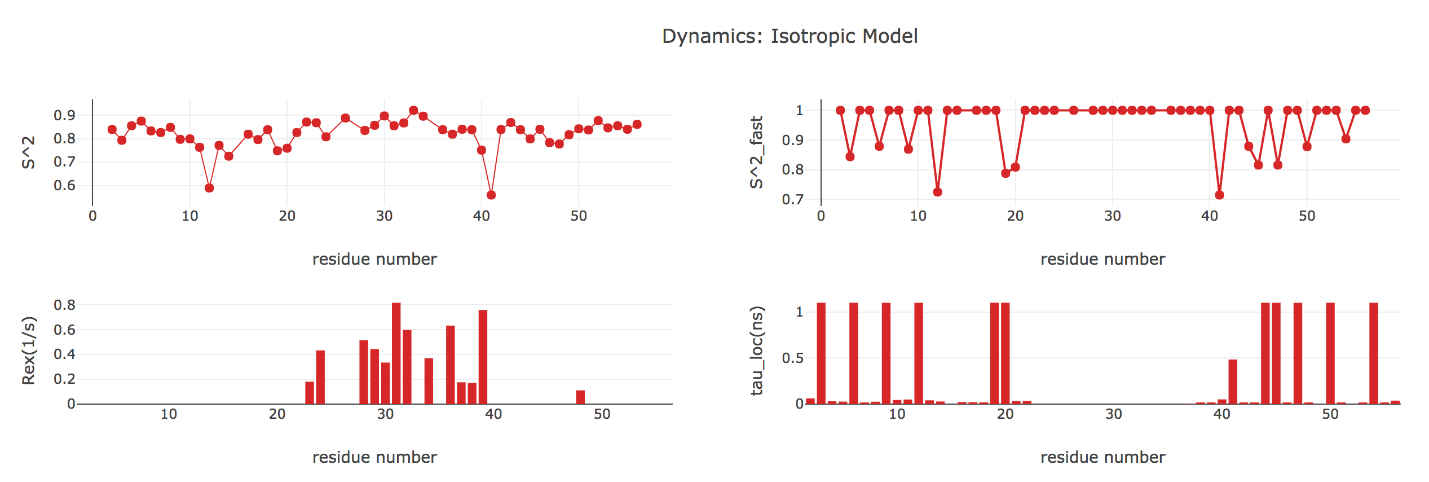
**Figure 8. Fully Anisotropic Model Fit**

These plots illustrate how well or not the experimental data agree with the fully anisotropic rotational diffusion tensor model. Shown on the left is the agreement between the experimental and back-calculated/predicted values of ; the line corresponds to absolute agreement. Shown on the right are the normalized residuals of fit for each residue. You can use the mouse/cursor to read the values for individual residues.



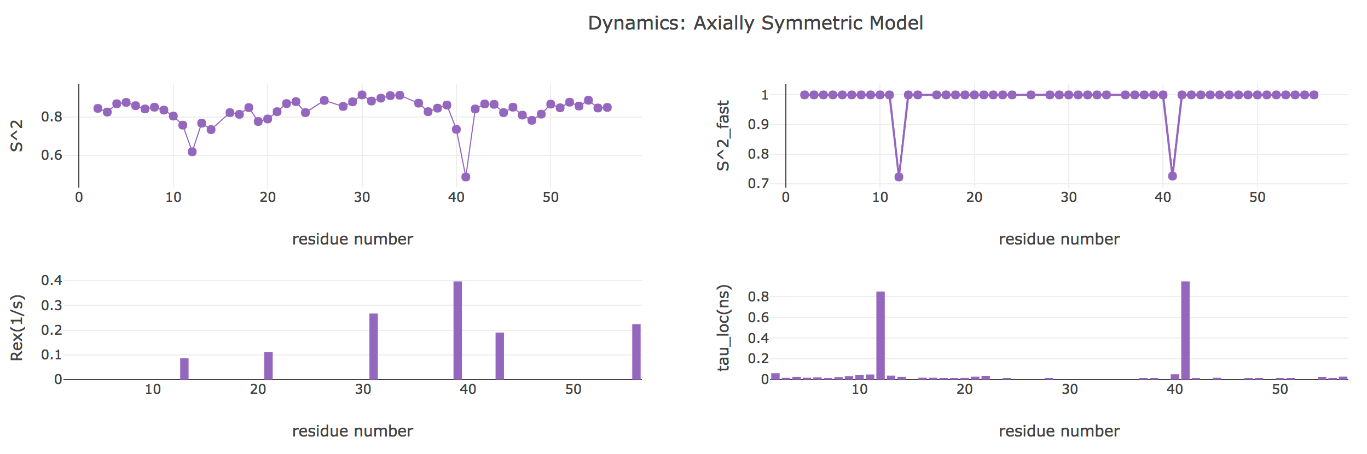
**Figure 9. Dynamics: Isotropic Model**

These plots depict the model-free parameters of local motions derived from the experimental data assuming isotropic rotational diffusion tensor model: the squared order parameter (S^2) and the related local correlation time (tau\_loc), as well as, when applicable, the squared order parameter for fast motion (S^2\_fast) and the conformational exchange contribution (Rex) to R2. See publications [3, 11] for a detailed description of these parameters. You can use the mouse/cursor to read the values for individual residues.



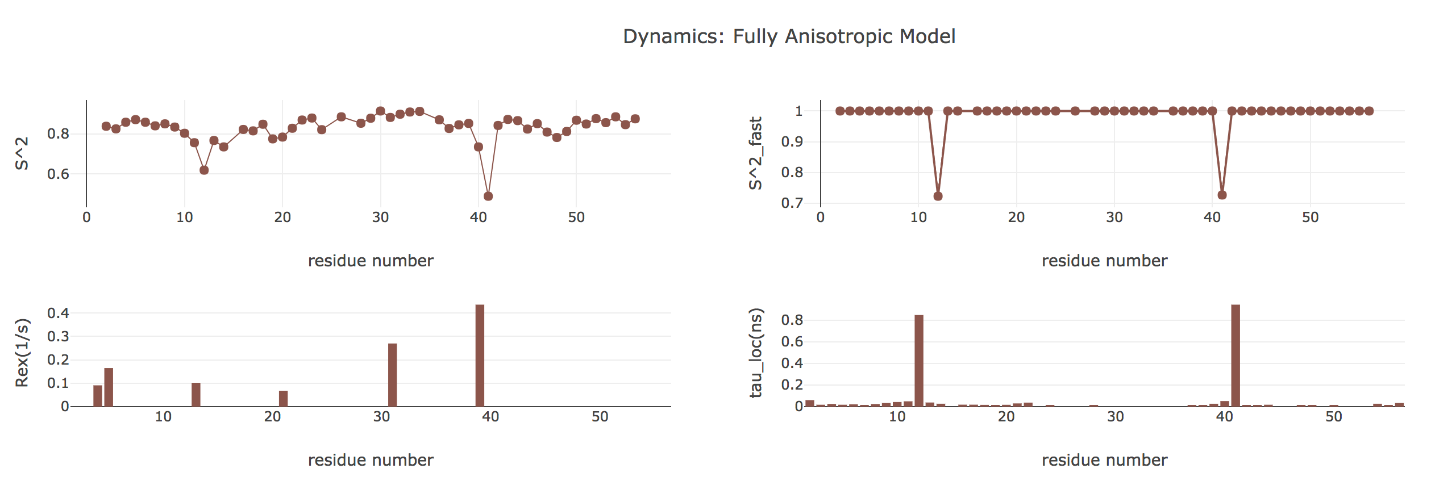
**Figure 10. Dynamics: Axially Symmetric Model**

These plots depict the model-free parameters of local motions derived from the experimental data assuming axially symmetric rotational diffusion tensor model: the squared order parameter (S^2) and the related local correlation time (tau\_loc), as well as, when applicable, the squared order parameter for fast motion (S^2\_fast) and the conformational exchange contribution (Rex) to R2. See publications [3, 11] for the details of these parameters. You can use the mouse/cursor to read the values for individual residues.



**Figure 11. Dynamics: Fully Anisotropic Model**

These plots depict the model-free parameters of local motions derived from the experimental data assuming fully anisotropic rotational diffusion tensor model: the squared order parameter (S^2) and the related local correlation time (tau\_loc), as well as, when applicable, the squared order parameter for fast motion (S^2\_fast) and the conformational exchange contribution (Rex) to R2. See publications [3, 11] for the details of these parameters. You can use the mouse/cursor to read the values for individual residues.



# Exercise Examples

This exercise includes analysis of 15N relaxation data at three different fields (1H resonance frequencies: 500 MHz, 600 MHz, and 800 MHz). Please use files “1P7F.pdb” and “GB3\_3Fields.txt” to run ROTDIF-Web in all applicable modules. You should be able to see plots like these:



# References:

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2. K. Berlin, A. Longhini, T. K. Dayie, D. Fushman, “Deriving Quantitative Dynamics Information for Proteins and RNAs using ROTDIF with a Graphical User Interface”, J Biomol NMR (2013) 57, 333-352.

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