Data Production

Parameters (.info)

Plotting (.plot_rhoz)

Send to detector (.detect)

MD: Calculate correlation functions NMR: Load from file ROMANCE (frames) Selection Object Frame Object (pyDR.Frames.FrameObj) pyDR.IO.readNMR Text file with experimental All MD analyses start with specifying the All MD analyses start with specifying parameters and data atoms/bonds of interest the atoms/bonds of interest Same object as in data below Define bond frame (.tensor_frame) Define reference frames (.new frame) Define selections (.select bond) Send to data object (.frames2data) Bond selections (.sel1, .sel2) Via data fitting Send to iRED (CC) object (.frames2ired) Vectors, positions (.v.,pos) Trajectory management (.traj) (source) Data Object Default labels (.label) Same as data object below. Cross-correlation (iRED) Produces a new data object via fitting or optimization **Basic Processing** iRED Object (pyDR.iRED) (.fit,.opt2dist) pyDR.md2data Calculates reorientational modes (rank 1/2), converts to data Send to data object (.ired2data) **Project Object** Data container Storage: Batch processing Data indexing: Provides quick access to a Saves data, including data Detector optimization (.detect) [0] single index large data set, sorted via reduction (.save) Batch fitting (.fit,.opt2dist) [[0,3,4]] selection Manages data reloading various parameters. [1:5:2] slicing Display (minimize time/memory/drive Allows efficient batch ['no_opt'] keywords Plot management processing and efficientspace) ['p6.+N15'] RegEx (.plot,.plot_obj,.current_plot,.fig) storage and reloading ChimeraX manager (.chimera) **Data Object** Data: Relaxation rates / Correlation functions (.R) Metadata: Data title (.title) Standard deviation (.Rstd) Labels (.label) Order Parameters + standard deviation (.S2,.S2std) Sensitivity parameters (.info) Descriptive data history (.details) Fits: Source data (.src_data) Selection Object (.select) Fitted data (.Rc, .S2c) Statistical parameters (.chi2..chi2red..AIC..AICc) Locations in PDB corresponding to data (len(data.select)=len(data)) Display: Plot data+sensitivity (.plot) Define selections (.select bond) Plot fit quality (.plot fit) Bond selections (.sel1, .sel2) Display selections (.repr_sel) Display in chimeraX (.chimera) Default labels (.label) Display in NGL view (.nglview) Fitting: Fit data using detectors (.fit) Source Information Object (.source) Optimize fit to a distribution (.opt2dist) Organizes data, talks to Project object File locations (.filename,.original_file, .topo, etc.) Sensitivity Object (.sens) Processing info (.status,.n_det,.additional_info) Title constructor (.title) Defines data sensitivity vs. correlation time (can be a detector object) Detector Object (.detect) Correlation time (.z.,tc) Sensitivity (.rhoz) Determines fitting for data object

(child class of sensitivity class)

Same as sense (.z,.tc,rhoz,.info,.plot_rhoz)

Detector setup (.r_auto, .r_target, .r_no_opt)