

# MDplot: Visualise Molecular Dynamics

by Christian Margreitter and Chris Oostenbrink

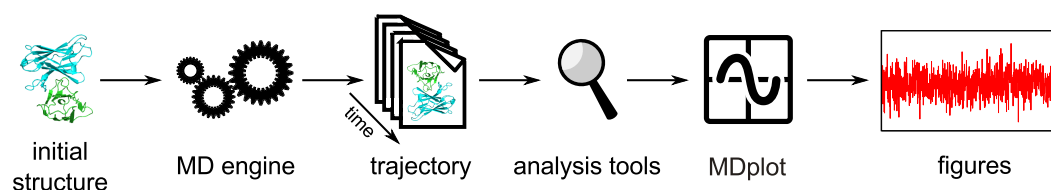
**Abstract** The MDplot package provides plotting functions to allow for automated visualisation of molecular dynamics simulation output. It is especially useful in cases where the plot generation is rather tedious due to complex file formats or when large amounts of plots are generated. The graphs that are supported range from those which are standard such as RMSD/RMSF (root-mean-square deviation and root-mean-square fluctuation, respectively) to thermodynamic integration analysis and hydrogen bond monitoring over time and address many commonly used analyses. In this article, we set out the package's functions, give examples of the function calls and show the associated plots. Plotting and data parsing is separated in all cases, i.e. the respective functions can be used independently. Thus, data manipulation and the integration of additional file formats is fairly easy. Currently, the loading functions support GROMOS, GROMACS and AMBER file formats. Moreover, we also provide a bash interface that allows simple embedding of MDplot into bash scripts as the final analysis step.

**Availability:** The package can be obtained in the latest major version from CRAN (<https://cran.r-project.org/package=MDplot>) or in the most recent version from the project's GitHub page at <https://github.com/MDplot/MDplot>, where feedback is also most welcome. **MDplot** is published under the GPL-3 license.

## Introduction

The amount of data produced by molecular dynamics (MD) engines (such as GROMOS (Schmid et al., 2012; Eichenberger et al., 2011), GROMACS (Pronk et al., 2013), NAMD (Phillips et al., 2005), AMBER (Cornell et al., 1995) and CHARMM (Brooks et al., 2009)) has been constantly increasing over recent years. This is mainly due to more powerful and cheaper hardware and as a result of this both the lengths and sheer number of MD simulations (i.e. trajectories) increased enormously. Even large sets of simulations (e.g. in the context of drug design) are attainable nowadays thus suggesting that the processing of the resulting information is undertaken automatically.

In this respect, automated yet flexible visualisation of molecular dynamics data would be highly advantageous: both in order to avoid repetitive tasks for the user and to yield the ultimately desired result instantly (see figure 1). Moreover, generating some of the graphs can be cumbersome. An example would be the plotting of a timeseries of a clustering program or hydrogen bonds. Therefore, these cases are predestined to be handled by a plotting library. There have been attempts made in that direction, for example the package **bio3d** (Grant et al., 2006; Skjærven et al., 2014) (which allows the trajectories to be processed in terms of principle component analysis (PCA), RMSD and RMSF calculations), **MDtraj** (McGibbon et al., 2015) or **Rknots** (Comoglio and Rinaldi, 2012). However, to the best of our knowledge there is currently no R package available that offers the wide range of plotting functions and engine-support that is provided by **MDplot** presented in this paper. R is the natural choice for this undertaking because of both its power in data handling and its vast plotting abilities.



**Figure 1:** Shows the overall workflow typically applied in molecular dynamics simulations beginning with a single PDB (Berman et al., 2000) structure as the input for the simulation and ending with the graphical representation of the data obtained. For large amounts of data, generating figures might become a tedious, highly repetitive task.

In the following sections we outline all of the plotting functions that are currently supported. Examples of the function calls based on the test data included in the package as well as the resulting plots, return values and argument lists for each function are described in detail. The respective code samples use the loading functions (reported below) to parse the input files located in folder 'extdata', which allows immediate testing and provides format information to users. Currently, the package supports

GROMOS, GROMACS and AMBER file formats as input.<sup>1</sup> However, extensions in both format support and plotting functionalities are planned.

## Plotting functions

The package currently offers 14 distinct plotting functions (table 1), which cover many of the graphs that are commonly required. Although the focus of the package relies on the visualisation of data, in addition to this values are calculated to characterise the underlying data when appropriate. For example `TIcurve()` calculates the thermodynamic integration free-energy values including error estimates and the hysteresis between the integration curves. In many cases, the plotting functions return useful information on the data used, e.g. range, mean and standard deviation of curves.

To provide simple access to these functions, they may be called from within a bash script. Examples are provided at the end of the manuscript.

**Table 1:** Lists all of the currently available plotting functions that have been implemented in **MDplot**. Most functions accept a boolean parameter (`barePlot`), that allows to print the plotting area only, i.e. stripped from any additional features such as axis labels.

plot function	description
<code>clusters()</code>	summary of clustering over trajectories (RMSD based)
<code>clusters_ts()</code>	timeseries of cluster populations (RMSD based)
<code>dssp()</code>	secondary structure annotation plot (DSSP based)
<code>dssp_ts()</code>	timeseries of secondary structure elements (DSSP based)
<code>hbond()</code>	hydrogen bonds summary plot
<code>hbond_ts()</code>	timeseries of hydrogen bonds
<code>noe()</code>	Nuclear-Overhauser-Effect violation plot
<code>ramachandran()</code>	dihedral angle plot
<code>rmsd()</code>	Root-Mean-Square Deviation plot
<code>rmsd_average()</code>	average Root-Mean-Square Deviation plot
<code>rmsf()</code>	Root-Mean-Square Fluctuation plot
<code>TIcurve()</code>	thermodynamic integration curves
<code>timeseries()</code>	general timeseries plot
<code>xrmsd()</code>	cross-RMSD plot (heat-map of RMSD values)

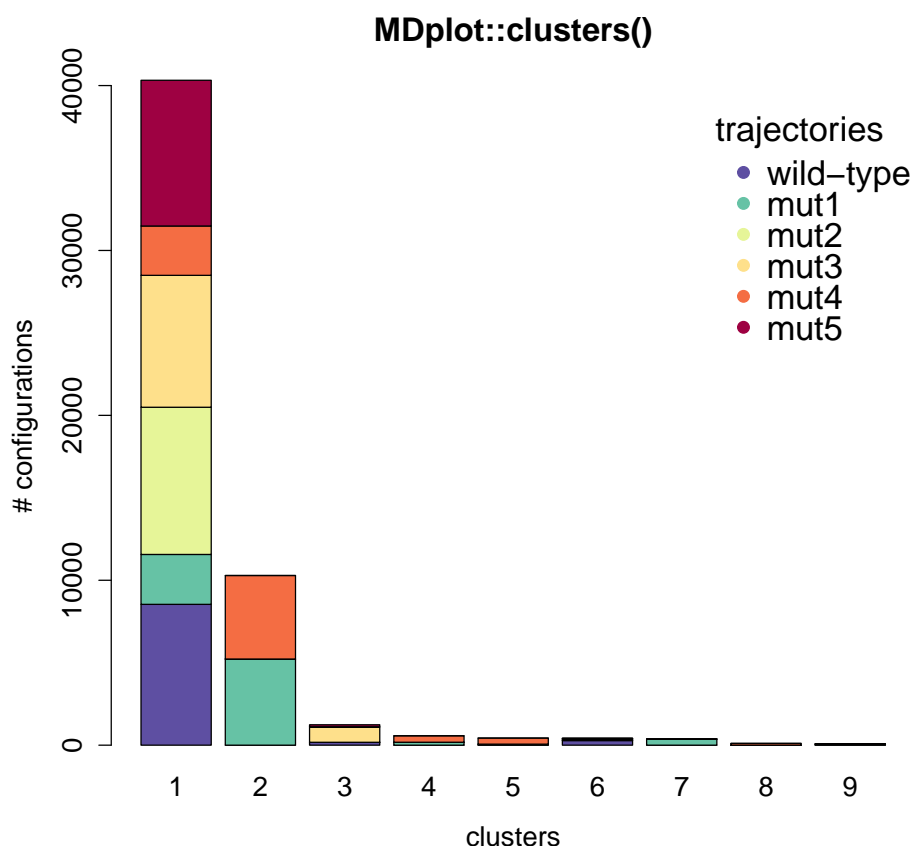
### clusters()

Molecular dynamics simulation trajectories can be considered to be a set of atom configurations along the time axis. Clustering is a method, that can be applied in order to extract common structural features from these. The configurations are classified and grouped together based on the root-mean-square deviation (RMSD). These subsets of configurations around the cluster's central member structure and their relative occurrences allow for comparisons between different and within individual simulations. `clusters()` allows to plot a summary of all of the (selected) clusters over a set of trajectories (figure 2).

```
clusters(load_clusters("inst/extdata/clusters_example.txt.gz",
                      names=c("wild-type", "mut1", "mut2",
                              "mut3", "mut4", "mut5")),
         clustersNumber=9, main="MDplot::clusters()", ylab="# configurations")
```

**Return value:** Returns a  $n \times m$ -matrix with  $n$  being the number of input trajectories and  $m$  the number of different clusters. Each element in the matrix holds the number of snapshots, in which the respective cluster occurred in the respective trajectory.

<sup>1</sup>In this manuscript, the code samples use GROMOS input (since the default value of the loading functions' parameter `mdEngine` is "GROMOS"). For information on how to load GROMACS or AMBER files, please have a look at the manual pages of the respective loading functions.



**Figure 2:** The clusters are plotted along the x-axis and the number of configurations for each trajectory for every cluster on the y-axis. The number of clusters is limited in this example to nine, which can be useful to omit scarcely populated clusters.

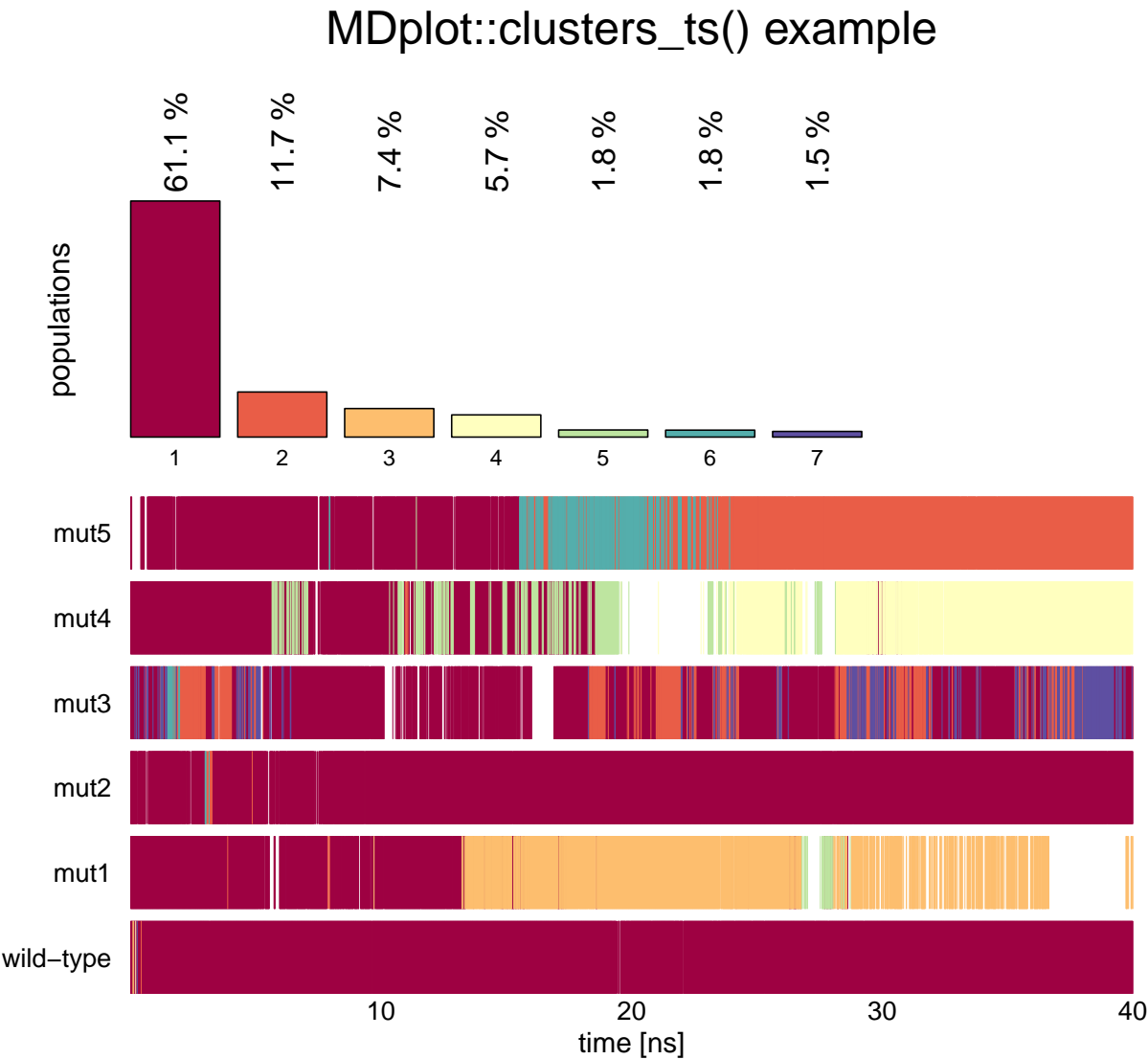
argument name	default value	description
clusters	<i>none</i>	Matrix with clusters: trajectories are given in row-wise, clusters in column-wise fashion as provided by <code>load_clusters()</code> , the associated loading function.
clustersNumber	NA	When specified, only these first clusters are shown.
legendTitle	"trajectories"	The title of the legend.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information or not.
...	<i>none</i>	Additional arguments.

### clusters\_ts()

In structural clustering, it is often instructive to have a look at the development over time rather than the overall summary. This functionality is provided by `clusters_ts()`. In the top sub-plot the overall distribution is given, while the timeseries is shown at the bottom. The clusters are sorted beginning with the most populated one, in descending order. Selections can be made and clusters that are not selected do also not appear in the timeseries plot (white areas). The time axis may be shown in nanoseconds (see figure 3 for an example).

```
clusters_ts(load_clusters_ts("inst/extdata/clusters_ts_example.txt.gz",
                           lengths=c(4000,4000,4000,4000,4000,4000),
                           names=c("wild-type","mut1","mut2",
                                   "mut3","mut4","mut5")),
           clustersNumber=7,main="MDplot::clusters_ts() example",
           timeUnit="ns",snapshotsPerTimeInt=100)
```

**Return value:** Returns a summary  $(n+1) \times m$ -matrix with  $n$  being the number of input trajectories and  $m$  the number of different clusters (which have been plotted). Each element in the matrix holds the number of snapshots, in which the respective cluster occurred in the respective trajectory. In addition, the first line is the overall summary counted over all trajectories.



**Figure 3:** The plot shows a selection of the seven most populated clusters of six trajectories. Regions that do not belong to one of the first seven clusters are shown in white.

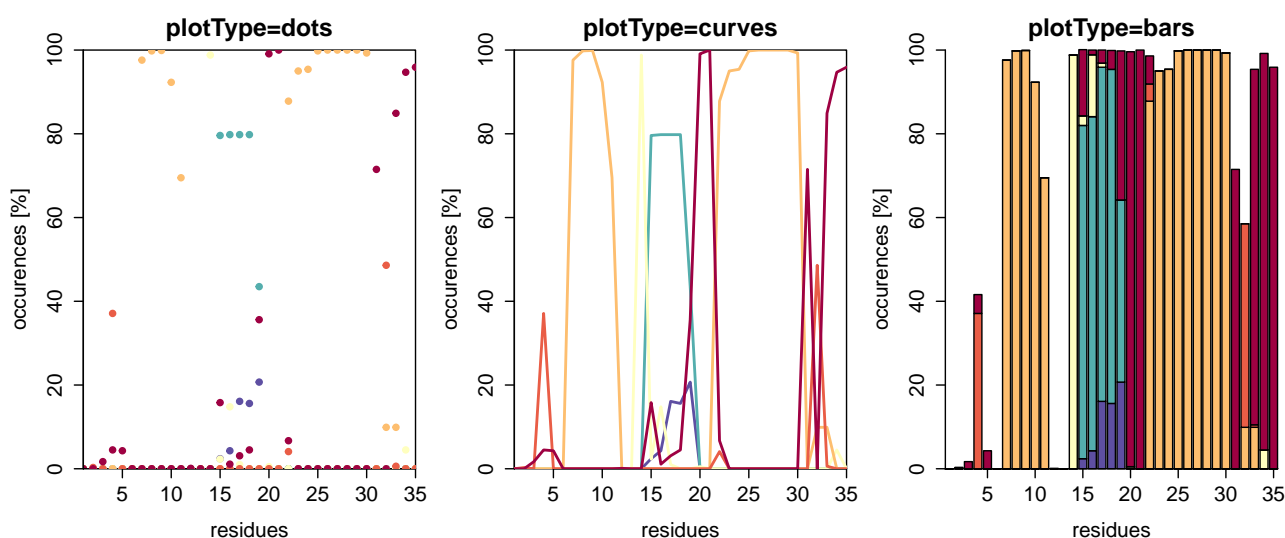
argument name	default value	description
clustersDataTS	none	List of cluster information as provided by load_clusters_ts(), the associated loading function.
clustersNumber	NA	Integer, specifying the number of clusters that is to be plotted.
selectTraj	NA	Vector of indices of trajectories that are plotted (as given in the input file).
selectTime	NA	Range of time in snapshots.
timeUnit	NA	Abbreviation of time unit.
snapshotsPerTimeInt	1000	Number of snapshots per time unit.
...	none	Additional arguments.

**dssp()**

In terms of proteins the secondary structure can be annotated by the widely used program DSSP (Definition of Secondary Structure of Proteins) (Kabsch and Sander, 1983). This algorithm uses the backbone hydrogen bond pattern in order to assign secondary structure elements such as  $\alpha$ -helices,  $\beta$ -strands, and turns to protein sequences. The plotting function `dssp()` has three different visualisation methods and plots the overall result over the trajectory and over the residues. The user can specify selections of residues and which elements should be taken into consideration (figure 4).

```
layout(matrix(1:3, nrow=1), widths=c(0.33,0.33,0.33))
dssp(load_dssp("inst/extdata/dssp_example.txt.gz"),
     main="plotType=dots",showResidues=c(1,35))
dssp(load_dssp("inst/extdata/dssp_example.txt.gz"),
     main="plotType=curves",plotType="curves",showResidues=c(1,35))
dssp(load_dssp("inst/extdata/dssp_example.txt.gz"),
     main="plotType=bars",plotType="bars",showResidues=c(1,35))
```

**Return value:** Returns a matrix, where the first column is the residue-number and the remaining ones denote secondary structure classes. Residues are given row-wise and values range from 0 to 100 percent.



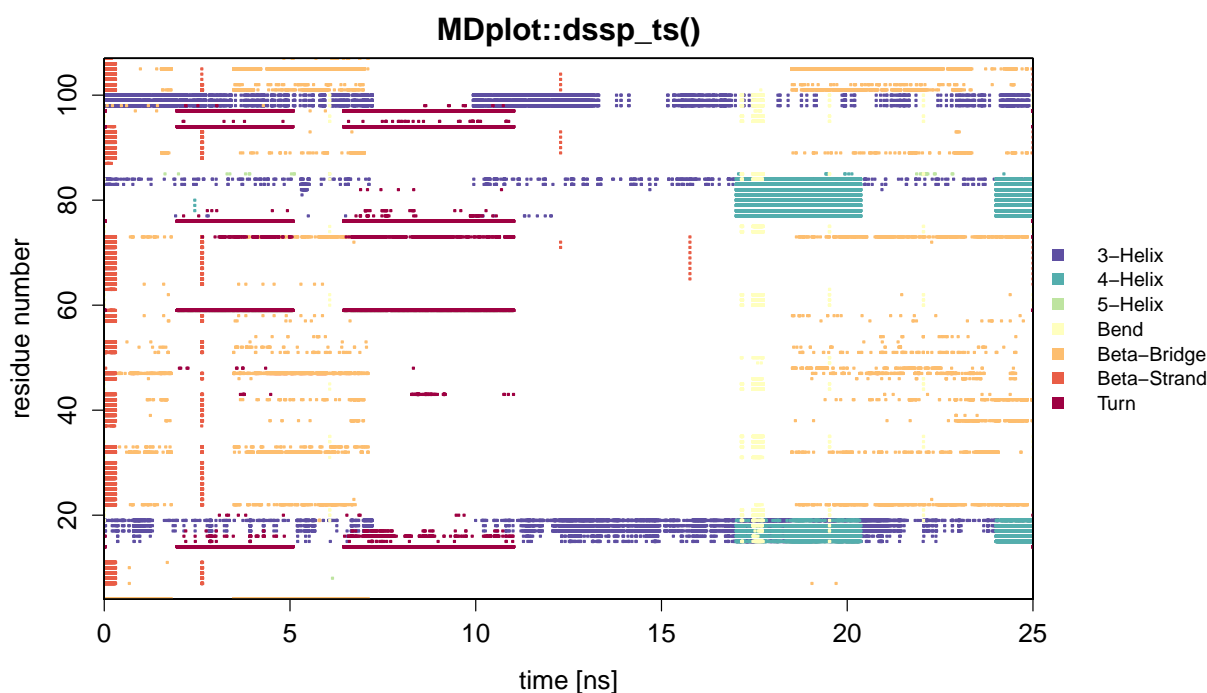
**Figure 4:** Example of `dssp()` with `plotType` set to "dots" (default), "curves" or "bars". Note that the fractions do not necessarily sum up to a hundred percent, because some residues might not be in defined secondary structure elements all the time. In this figure, there is no legend plotted due to space limitations (see figure 5 for a colour-code explanation).

argument name	default value	description
dsspData	<i>none</i>	Table containing information on the secondary structure elements. Can be generated by function <code>load_dssp()</code> .
printLegend	FALSE	If TRUE, a legend is printed on the right hand side of the plot.
useOwnLegend	FALSE	If FALSE, the names of the secondary structure elements are considered to be in default order.
elementNames	NA	Vector of names for the secondary structure elements.
colours	NA	A vector of colours that can be specified to replace the default ones.
showValues	NA	A vector of boundaries for the values.
showResidues	NA	A vector of boundaries for the residues.
plotType	"dots"	Either "dots", "curves" or "bars".
selectedElements	NA	A vector of names of the elements selected.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

### dssp\_ts()

The secondary structure information as described for the function `dssp()` can also be visualised along the time axis using function `dssp_ts()` (figure 5). The time can be annotated in snapshots or time units (e.g. nanoseconds).

```
dssp_ts(load_dssp_ts("inst/extdata/dssp_ts_example"), printLegend=TRUE,
        main="MDplot::dssp_ts()", timeUnit="ns",
        snapshotsPerTime=1000)
```



**Figure 5:** Example showing all of the defined secondary structure elements per residue over time. Note, that for this example plot a sparse data set was used to reduce the size of the data file (hence the large white areas in the middle).

argument name	default value	description
tsData	<i>none</i>	List of lists, which are composed of a name (string) and a values table (x ... snapshots, y ... residues). Can be generated by <code>load_dssp_ts()</code> .
printLegend	TRUE	If TRUE, a legend is printed on the right hand side of the plot.
timeBoundaries	NA	A vector of boundaries for the time in snapshots.
residueBoundaries	NA	A vector of boundaries for the residues.
timeUnit	NA	If set, the snapshots are transformed into the respective time (depending on parameter <code>snapshotsPerTime</code> ).
snapshotsPerTimeInt	1000	Number of snapshots per respective <code>timeUnit</code> .
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

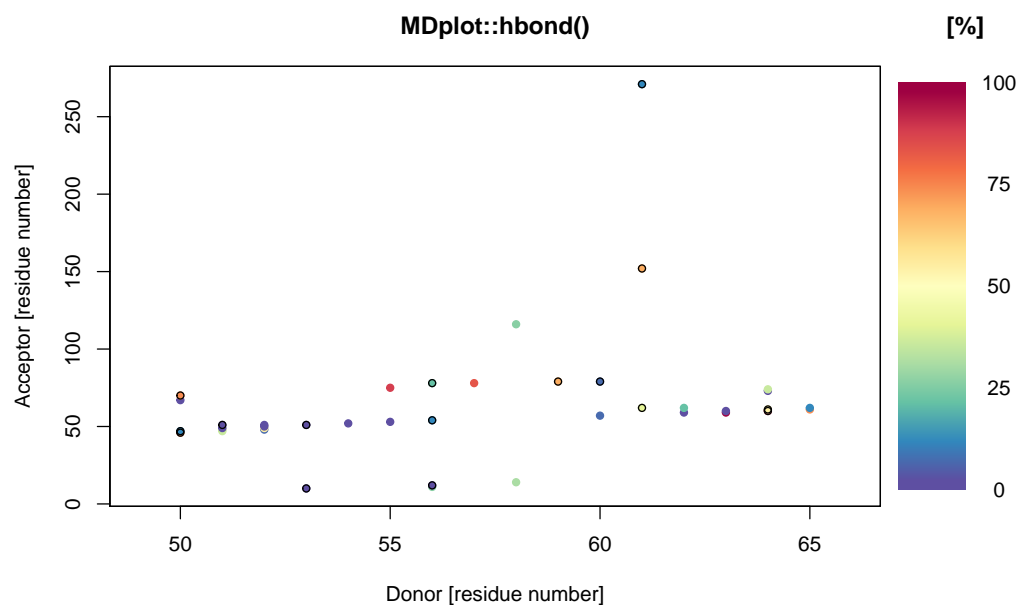
### hbond()

In the context of biomolecules, hydrogen bonds are of particular importance. These bonds take place between a donor, a hydrogen and an acceptor atom. This function plots the summary output of hydrogen bond calculations and allows selection of donor and acceptor residues. Occurrence over the whole trajectory is indicated by a colour scale. Note, that in case multiple hydrogen bond interactions between two particular residues take place (conveyed by different sets of atoms), the interaction with prevalence will be used for colour-coding (and by default, this interaction is marked with a black circle, see below). An example is given in figure 6.

```
hbond(load_hbond("inst/extdata/hbond_example.txt.gz"),
      main="MDplot::hbond()", donorRange=c(0,65))
```

**Return value:** Returns a table containing the information used for plotting in columns as follows:

- `resDonor` Residue number (donor).
- `resAcceptor` Residue number (acceptor).
- `percentage` Percentage, that has been used for colour-coding.
- `numberInteractions` Number of hydrogen bond interactions taking place between the specified donor and acceptor residues.



**Figure 6:** The acceptor residues are plotted on the x-axis whilst the donors are shown on the y-axis. The different colours indicate the occurrences throughout the whole trajectory.

argument name	default value	description
hbonds	<i>none</i>	Table containing the hydrogen bond information in columns "hbondID", "resDonor", "resDonorName", "resAcceptor", "resAcceptorName", "atomDonor", "atomDonorName", "atomH", "atomAcceptor", "atomAcceptorName", "percentage" (automatically generated by function load_hbond()).
plotMethod	"residue-wise"	Allows to set the detail of hydrogen bond information displayed. Options are: "residue-wise".
acceptorRange	NA	Vector, specifying the range of acceptor residues.
donorRange	NA	Vector, specifying the range of donor residues.
printLegend	TRUE	Boolean, enabling the legend.
showMultipleInteractions	TRUE	If TRUE, this option causes multiple interactions between the same residues as being represented by a black circle around the coloured dot.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

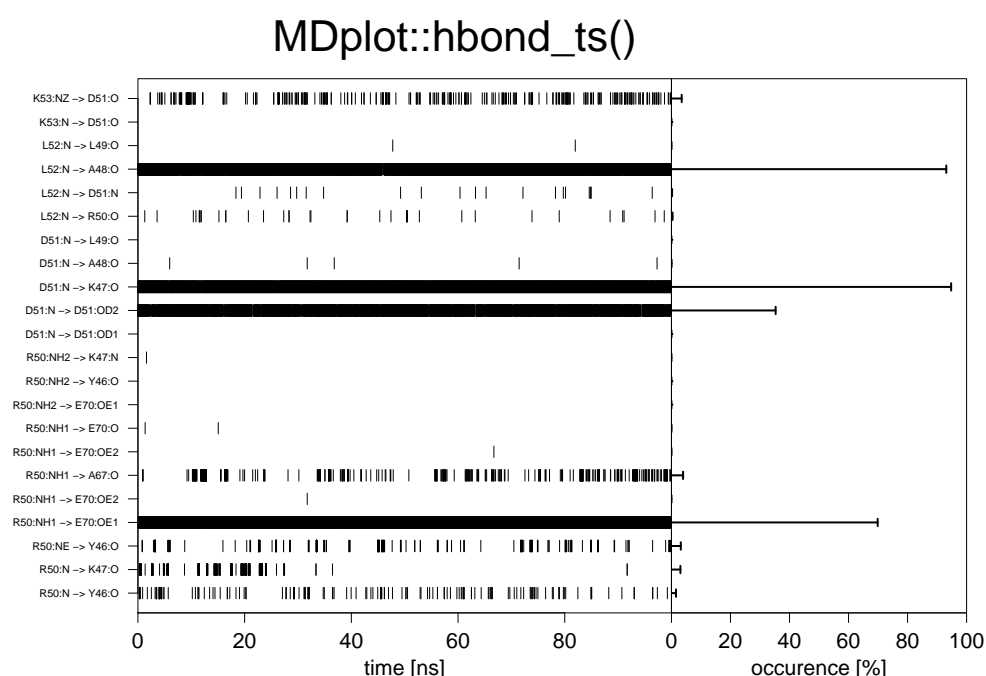


## hbond\_ts()

The timeseries of hydrogen bond occurrences can be visualised using the function `hbond_ts()`, which plots them either according to their identifiers or in a human readable form in three- or one-letter code (the participating atoms can be shown as well) on the y-axis and the time on the x-axis. If the GROMOS input format is used, this function requires two different files: the summary of the hbond program and the timeseries file. The occurrence of a hydrogen bond is represented by a black bar and the occurrence summary can be added on the right hand side as a sub-plot (figure 7). In addition to the timeseries file, depending on the MD engine format used, an additional summary file might also be necessary (see documentation of function `load_hbond_ts()` for further information).

```
hbond_ts(timeseries=load_hbond_ts("inst/extdata/hbond_ts_example.txt.gz"),
         summary=load_hbond("inst/extdata/hbond_example.txt.gz"),
         main="MDplot::hbond_ts()", acceptorRange=c(22, 75),
         hbondIndices=list(c(0, 24)), plotOccurrences=TRUE, timeUnit="ns",
         snapshotsPerTimeInt=100, printNames=TRUE, namesToSingle=TRUE,
         printAtoms=TRUE)
```

**Return value:** Returns a nx2 matrix, with the first column being the list of hydrogen bond identifiers plotted and the second one the occurrence (in percent) over the selected time range.



**Figure 7:** Example figure generated by `hbond_ts()` for both an identifier and acceptor residues' selection. The labels for the hydrogen bonds may be printed as identifiers or with names composed of residue names (in single- or three-letter code) and those of the participating atoms.

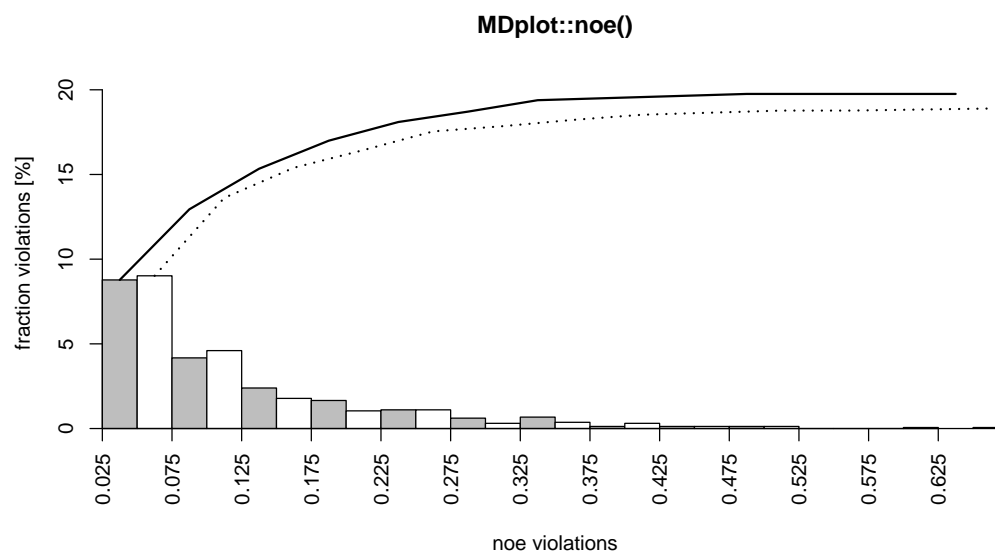
argument name	default value	description
timeseries	<i>none</i>	Table containing the timeseries information (e.g. produced by <code>load_hbond_ts()</code> ).
summary	<i>none</i>	Table containing the summary information (e.g. produced by <code>load_hbond()</code> ).
acceptorRange	NA	Vector of acceptor residues.
donorRange	NA	Vector of donor residues.
plotOccurrences	FALSE	Specifies whether the overall summary should be plotted on the right hand side.
scalingFactorPlot	NA	To manually set the scaling factor (if necessary).
printNames	FALSE	Enables human readable names rather than the hydrogen bond identifiers.
namesToSingle	FALSE	If <code>printNames</code> is TRUE, this flag enables one-letter codes instead of three-letter ones.
printAtoms	FALSE	Enables atom names in hydrogen bond identification on the y-axis.
timeUnit	NA	Specifies the time unit on the x-axis.
snapshotsPerTimeInt	1000	Specifies how many snapshots make up one time unit (see above).
timeRange	NA	Vector, selecting a certain time range.
hbondIndices	NA	A list containing vectors to select hydrogen bonds by their identifiers.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

### noe()

The Nuclear-Overhauser-Effect is one of the most important measures of structure validity in the context of molecular dynamics simulations. These interactions are transmitted through space and arise from spin-spin coupling, which can be measured by nuclear magnetic resonance (NMR) spectroscopy. These measurements provide pivotal distance restraints which should be matched on average during molecular dynamics simulations of the same system and can hence be used for parameter validation. The plotting function `noe()` allows to visualise the number of distance restraint violations and their respective spacial deviation. As shown in figure 8, multiple replicates or different protein systems are supported simultaneously. Note, that negative violations are not considered.

```
noe(load_noe(files=c("inst/extdata/noe_example_1.txt.gz",
                    "inst/extdata/noe_example_2.txt.gz")),
    main="MDplot::noe()")
```

**Return value:** Returns a matrix, in which the first column holds the bin boundaries used and the following columns represent either the percentage or absolute numbers of the violations per bin, depending on the specification.



**Figure 8:** Example plot showing two different replicates of a protein simulation (they share the same molecule, but have different initial velocities). Note, that the maximum value (x-axis) over all replicates is used for the plot. The sum over all violations from left to right is shown by an additional curve on top. The number of violations may be given as fractions (in %), as shown above, or absolute numbers (flag `printPercentages` either `TRUE` or `FALSE`).

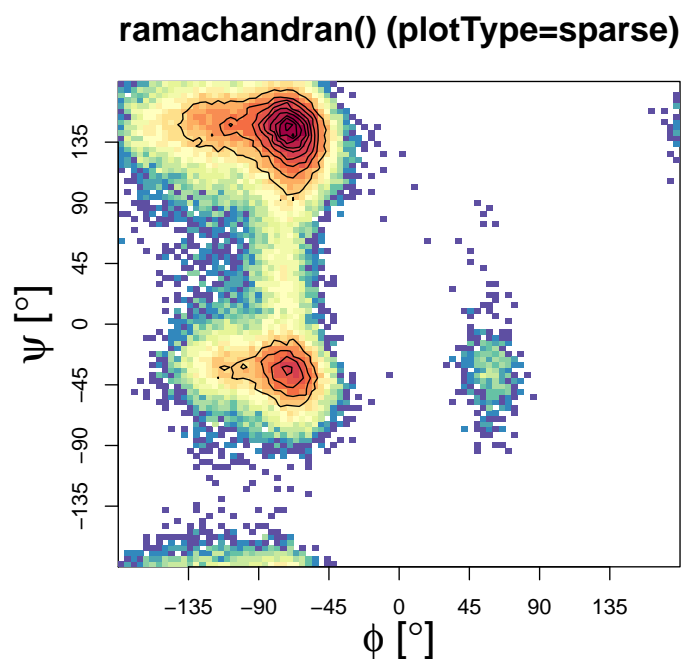
argument name	default value	description
<code>noeData</code>	<i>none</i>	Input matrix. Generated by function <code>load_noe()</code> .
<code>printPercentages</code>	<code>TRUE</code>	If <code>TRUE</code> , the violations will be reported in a relative manner (percent) rather than absolute numbers.
<code>colours</code>	<code>NA</code>	Vector of colours to be used for the bars.
<code>lineTypes</code>	<code>NA</code>	If <code>plotSumCurves</code> is <code>TRUE</code> , this vector might be used to specify the types of curves plotted.
<code>names</code>	<code>NA</code>	Vector to name the input columns (legend).
<code>plotSumCurves</code>	<code>TRUE</code>	If <code>TRUE</code> , the violations are summed up from left to right to show the overall behaviour.
<code>maxYAxis</code>	<code>NA</code>	Can be used to manually set the y-axis of the plot.
<code>printLegend</code>	<code>FALSE</code>	Boolean, which triggers plotting of the legend.
...	<i>none</i>	Additional arguments.

### ramachandran()

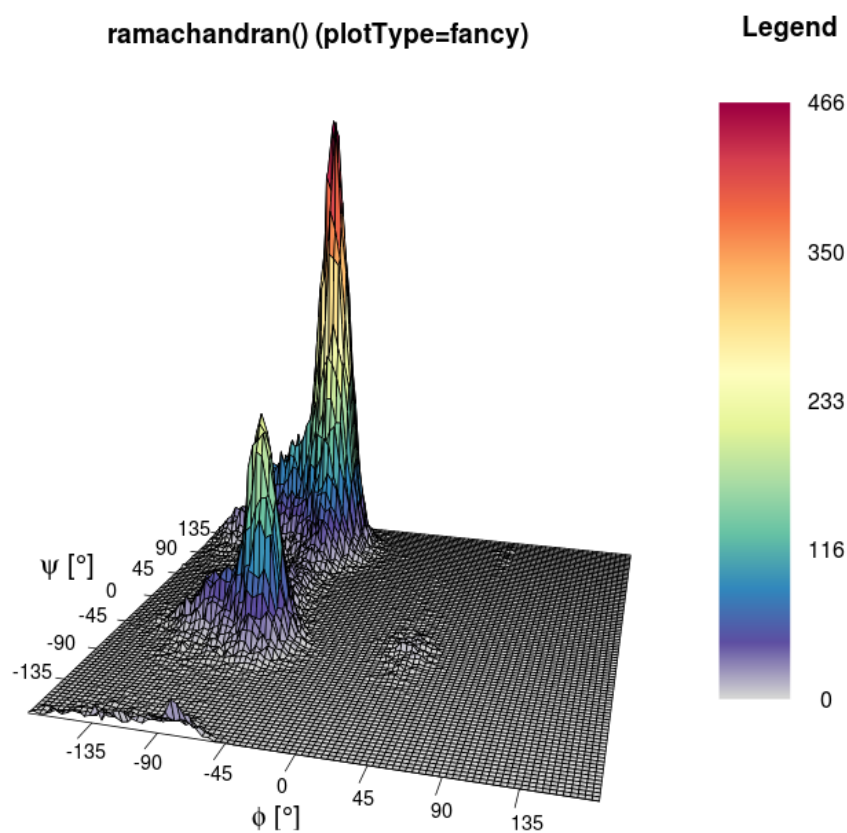
This graph type (Ramachandran et al., 1963) is often used to show the sampling of the  $\phi/\psi$  protein backbone dihedral angles in order to assign propensities of secondary structure elements to the protein of interest (so-called Ramachandran plots). These plots can provide crucial insight into energy barriers arising as required e.g. in the context of parameter validation (Margreitter and Oostenbrink, 2016). The function `ramachandran()` offers a 2D (figure 9) and 3D (figure 10) variant with the former offering the possibility to print user-defined secondary structure regions as well. The number of bins for the two axes and the colours used for the legend can be specified by the user.

```
ramachandran(load_ramachandran("inst/extdata/ramachandran_example.txt.gz"),
             heatFun="log", plotType="sparse", xBins=90, yBins=90,
             main="ramachandran() (plotType=sparse)",
             plotContour=TRUE)
ramachandran(load_ramachandran("inst/extdata/ramachandran_example.txt.gz"),
             heatFun="norm", plotType="fancy", xBins=90, yBins=90,
             main="ramachandran() (plotType=fancy)",
             printLegend=TRUE)
```

**Return value:** Returns a list of binned dihedral angle occurrences.



**Figure 9:** Two dimensional plot version "sparse" of the `ramachandran()` function with enabled contour plotting. The number of bins can be specified for both dimensions independently.



**Figure 10:** Three dimensional example of `ramachandran()`. In addition to the colour, the height (z-axis) also represents the number of dihedrals per bin.

argument name	default value	description
dihedrals	<i>none</i>	Matrix with angles (two columns). Generated by function <code>load_ramachandran()</code> .
xBins	150	Number of bins used to plot (x-axis).
yBins	150	Number of bins used to plot (y-axis).
heatFun	"norm"	Function selector for calculation of the colour. The possibilities are either: "norm" for linear calculation or "log" for logarithmic calculation.
structureAreas	<code>c()</code>	List of areas, which are plotted as black lines.
plotType	"sparse"	Type of plot to be used, either "sparse" (default, using function <code>hist2d()</code> ), "comic" (own binning, supports very few datapoints) or "fancy" (3D, using function <code>persp()</code> ).
printLegend	FALSE	Boolean, specifying whether a heat legend is to be plotted or not.
plotContour	FALSE	Boolean, specifying whether a contour should be added or not.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

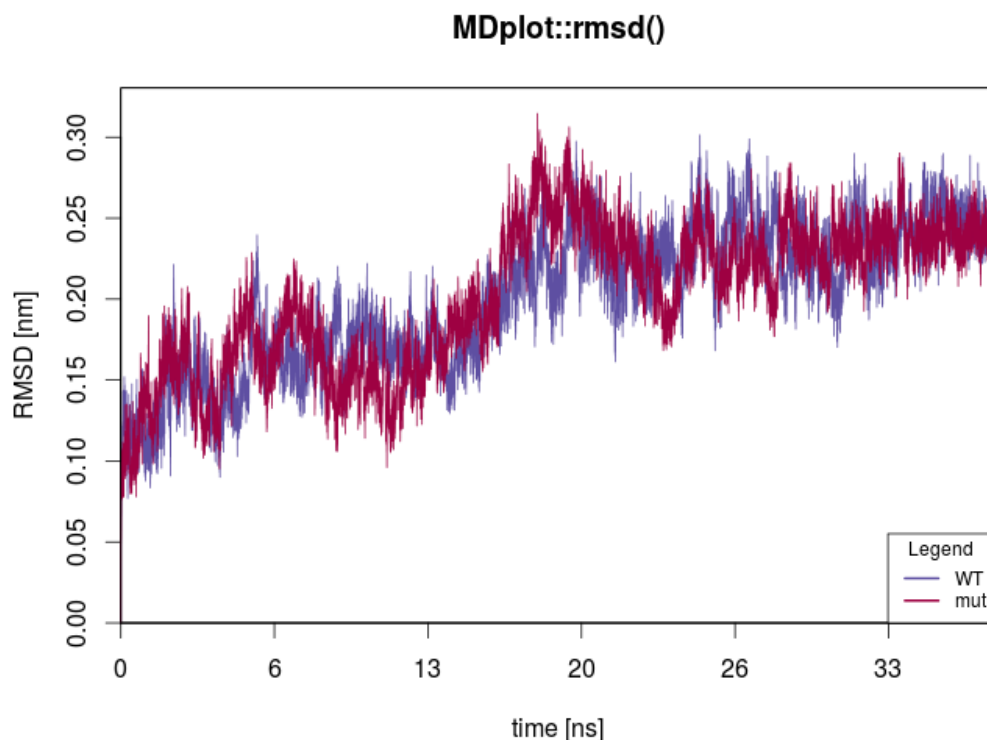
### rmsd()

The atom-positional root-mean-square deviation (RMSD) is one of the most commonly used plot types in the field of biophysical simulations. In the context of atom configurations, it is a measure for the positional divergence of one or multiple atoms. The input requires a list of alternating vectors of time indices and RMSD values. Multiple data sets can be plotted, given in separate input files. Figure 11 shows an example for two trajectories.

```
rmsd(load_rmsd(c("inst/extdata/rmsd_example_1.txt.gz",
                 "inst/extdata/rmsd_example_2.txt.gz")),
      printLegend=TRUE, names=c("WT", "mut"), main="MDplot::rmsd()")
```

**Return value:** Returns a list of lists, where each sub-list represents a RMSD curve and contains:

- `minValue` The minimum value over the whole time range.
- `maxValue` The maximum value over the whole time range.
- `meanValue` The mean value calculated over the whole time range.
- `sd` The standard deviation calculated over the whole time range.



**Figure 11:** This plot shows the RMSD curves for two different trajectories. The time is given in nanoseconds, which requires a properly set factor parameter.

argument name	default value	description
rmsdData	<i>none</i>	List of (alternating) indices and RMSD value vectors, as produced by <code>load_rmsd()</code> .
printLegend	TRUE	Boolean, which triggers the plotting of the legend.
factor	1000	Number, specifying how many snapshots are within one timeUnit.
timeUnit	"ns"	Specifies the time unit.
rmsdUnit	"nm"	Specifies the RMSD unit.
colours	NA	Vector of colours used for plotting.
names	NA	Vector holding the names of the trajectories.
legendPosition	"bottomright"	Indicates the position of the legend: either "bottom-right", "bottomleft", "topleft" or "topright".
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

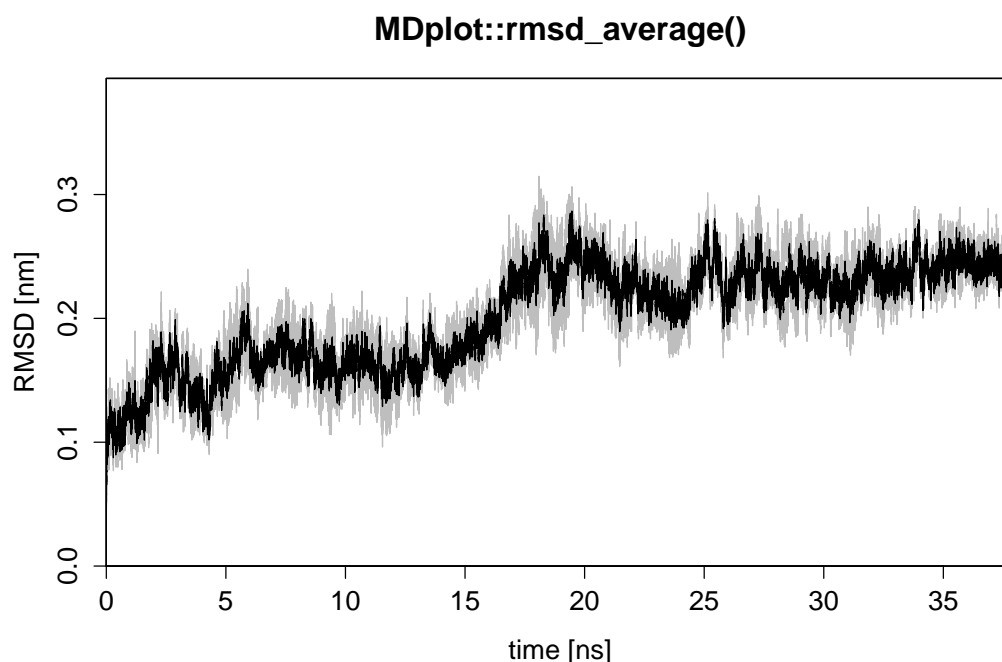
### **rmsd\_average()**

Nowadays, for many molecular systems multiple replicates of simulations are performed in order to enhance the sampling of the phase space. However, since the amount of analysis data grows accordingly, a joint representation of the results may be desirable. For the case of backbone-atom and other RMSD plots, the **MDplot** package supports average plotting. Instead of plotting every curve individually, the mean and the minimum and maximum values of all trajectories at a given time point is plotted. Thus, the spread of multiple simulations is represented as a 'corridor' over time.

```
rmsd_average(rmsdInput=list(load_rmsd("inst/extdata/rmsd_example_1.txt.gz" ),
                             load_rmsd("inst/extdata/rmsd_example_2.txt.gz")),
             maxYAxis=0.375,main="MDplot::rmsd_average()")
```

**Return value:** Returns a nx4-matrix, with the rows representing different snapshots and the columns the respective values as follows:

- snapshot Index of the snapshot.
- minimum The minimum RMSD value over all input sources at a given time.
- mean The mean RMSD value over all input sources at a given time.
- maximum The maximum RMSD value over all input sources at a given time.



**Figure 12:** In black, the mean RMSD value at a given timepoint and in grey the respective minimum and maximum values are given. In this example, two rather similar curves have been used.

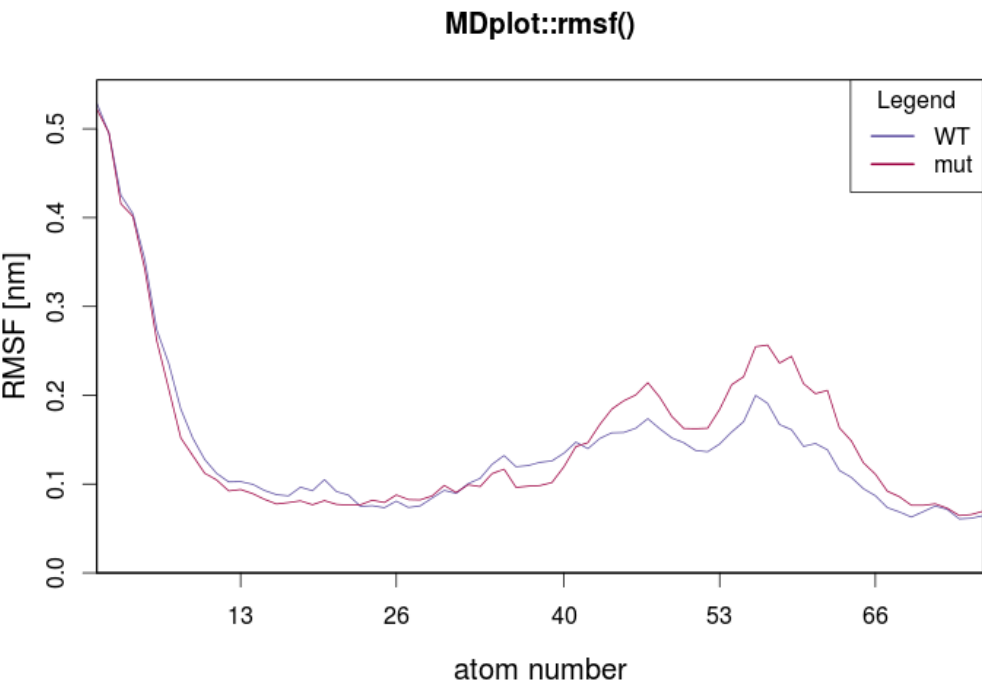
argument name	default value	description
rmsdInput	<i>none</i>	List of snapshot and RMSD value pairs, as e.g. provided by loading function <code>load_rmsd()</code> .
levelFactor	NA	If there are many datapoints, this parameter may be used to use only the <code>levelFactor</code> th datapoints to obtain a clean graph.
snapshotsPerTimeInt	1000	Number, specifying how many snapshots are comprising one <code>timeUnit</code> .
timeUnit	"ns"	Specifies the time unit.
rmsdUnit	"nm"	Specifies the RMSD unit.
maxYAxis	NA	Can be used to manually set the y-axis of the plot.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

### rmsf()

The atom-positional root-mean-square fluctuation (RMSF) represents the degree of positional variation of a given atom over time. The input requires one column with all residues or atoms and a second one holding RMSF values. Figure 13 shows, as an example, the RMSF of the first 75 atoms, calculated for two independent simulations.

```
rmsf(load_rmsf(c("inst/extdata/rmsf_example_1.txt.gz",
               "inst/extdata/rmsf_example_2.txt.gz")),
      printLegend=TRUE,names=c("WT","mut"),range=c(1,75),
      main="MDplot::rmsf()")
```

**Return value:** A list of vectors, alternately holding atom indices and their respective values.



**Figure 13:** Plot showing two different RMSF curves.

argument name	default value	description
rmsfData	<i>none</i>	List of (alternating) atom numbers and RMSF values, as e.g. produced by <code>load_rmsf()</code> .
printLegend	TRUE	Boolean, which triggers plotting of the legend.
rmsfUnit	"nm"	Specifies the RMSF unit.
colours	NA	Vector of colours used for plot.
residuewise	FALSE	Boolean, specifying whether atoms or residues are plotted on the x-axis.
atomsPerResidue	NA	If residuewise is TRUE, this parameter can be used to specify the number of atoms per residue for plotting.
names	NA	Vector of the names of the trajectories.
range	NA	Range of atoms.
legendPosition	"topright"	Indicates position of legend: either "bottomright", "bottomleft", "topleft" or "topright".
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.



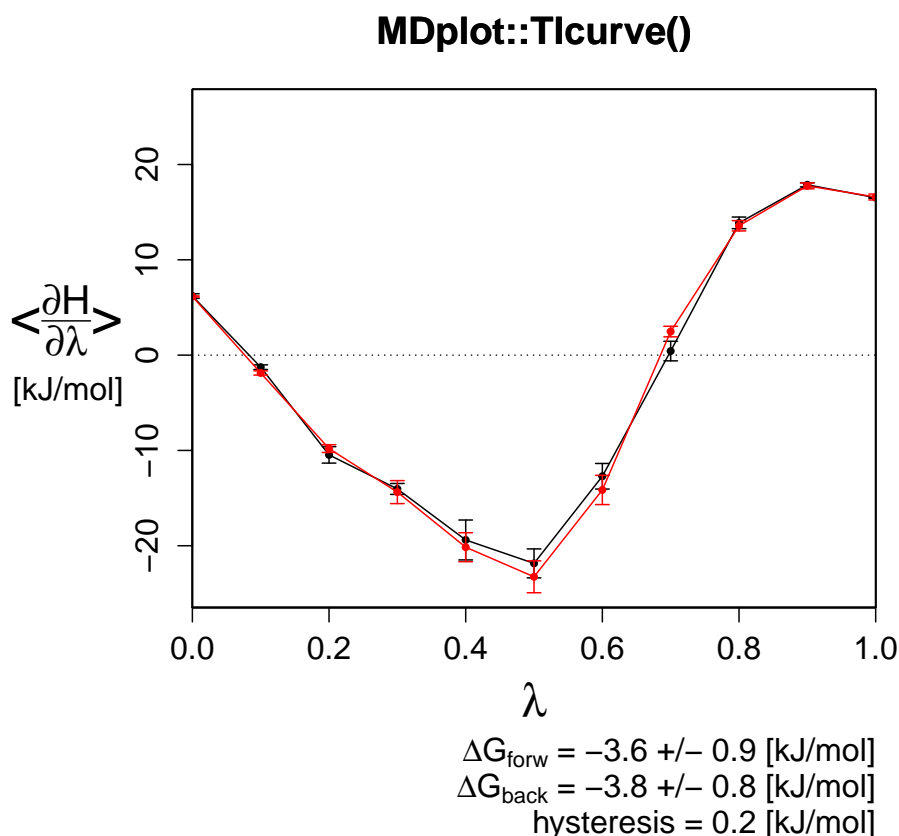
**Tlcurve()**

For calculations of the free energy difference occurring when transforming one chemical compound into another (alchemical changes) or for estimates of free energy changes upon binding, thermodynamic integration (Kirkwood, 1935) is one of the most trusted and applied approaches. The derivative of the Hamiltonian as a function of a coupling parameter  $\lambda$  is calculated over a series of  $\lambda$  state points (typically around 15). The integral of this curve is equivalent to the change in free energy (figure 14). The function `Tlcurve()` performs the integration and, if the data for both the forward and backward processes are provided, the hysteresis between them.

```
Tlcurve(load_Tlcurve(c("inst/extdata/Tlcurve_fb_forward_example.txt.gz",
                      "inst/extdata/Tlcurve_fb_backward_example.txt.gz")),
        invertedBackwards=TRUE, main="MDplot::Tlcurve()")
```

**Return value:** Returns a list with the following information:

- `lambdapoints` A list containing a (at least)  $n \times 3$ -matrix for every data input series.
- `integrationresults` A matrix containing one row of "deltaG" and "error" columns from the integration for every data input series.
- `hysteresis` If two (i.e. forward and backward) data input series are provided, the resulting hysteresis is reported (and set to be NA otherwise).



**Figure 14:** A forward and backward thermodynamic integration curve with the resulting hysteresis between them (precision as permitted by the error).

argument name	default value	description
lambdas	<i>none</i>	List of matrices (automatically generated by <code>load_TIcurve()</code> ) holding the thermodynamic integration information.
invertedBackwards	FALSE	If a forward and backward TI are provided and the lambda points are enumerated reversely (i.e. 0.3 of one TI is equivalent to 0.7 of the other), this flag can be set to be TRUE in order to automatically mirror the values appropriately.
energyUnit	"kJ/mol"	Defines the energy unit used for the plot.
printValues	TRUE	If TRUE, the free energy values are printed.
printErrors	TRUE	Boolean, indicating whether error bars are to be plotted.
errorBarThreshold	0	If the error at a given lambda point is below this threshold, it is not plotted.
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

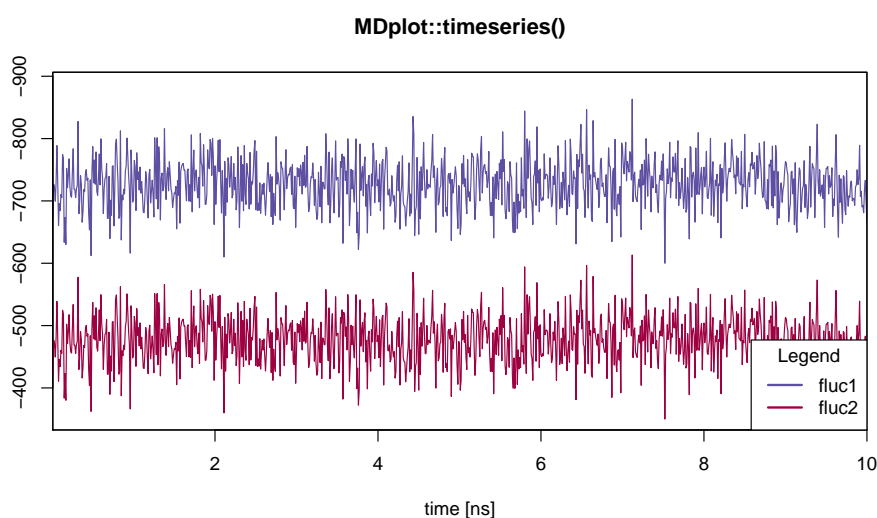
### timeseries()

This function provides a general interface for any timeseries given as a time-value pair (figure 15).

```
timeseries(load_timeseries(c("inst/extdata/timeseries_example_1.txt.gz",
                             "inst/extdata/timeseries_example_2.txt.gz")),
           main="MDplot::timeseries()",
           names=c("fluc1", "fluc2"),
           snapshotsPerTimeInt=100)
```

**Return value:** Returns a list of lists, each of the latter holding for every data input series:

- `minValue` The minimum value over the whole set.
- `maxValue` The maximum value over the whole set.
- `meanValue` The mean value over the whole set.
- `sd` The standard deviation over the whole set.



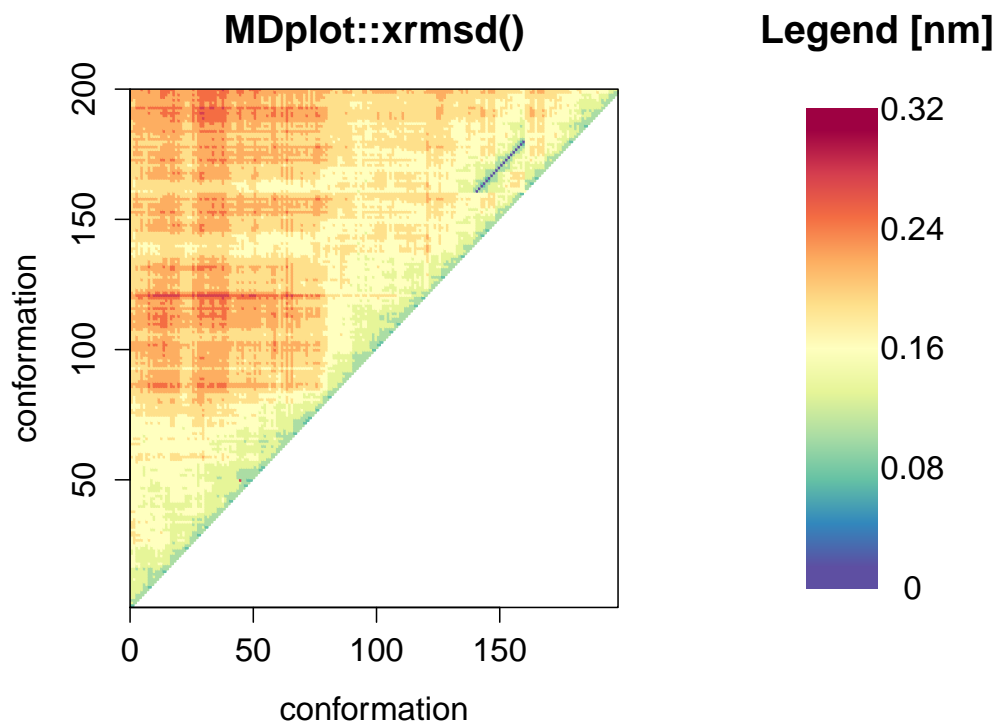
**Figure 15:** Shows timeseries with parameter `snapshotsPerTimeInt` set in a way such, that the proper time in nanoseconds is plotted. In addition, the legend has been moved to the bottom-right position.

argument name	default value	description
tsData	<i>none</i>	List of (alternating) indices and response values, as produced by <code>load_timeseries()</code> .
printLegend	TRUE	Parameter enabling the plotting of the legend.
snapshotsPerTimeInt	1000	Number specifying how many snapshots make up one timeUnit.
timeUnit	"ns"	Specifies the time unit.
valueName	NA	Name of response variable.
valueUnit	NA	Specifies the response variable's unit.
colours	NA	Vector of colours used for plotting.
names	NA	Vector of names of the trajectories.
legendPosition	"bottomright"	Indicates position of legend: either "bottom-right", "bottomleft", "topleft" or "topright".
barePlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

### xrmsd()

This function generates a plot which shows a heat-map of the atom positional root-mean-square differences between snapshots (figure 16). The structures are listed on the x- and y-axes. The heat-map shows the difference between one structure and another using a coloured bin. The legend is adapted in accordance to the size of the values.

```
xrmsd(load_xrmsd("inst/extdata/xrmsd_example.txt.gz"),
      printLegend=TRUE,main="MDplot::xrmsd()")
```



**Figure 16:** An example `xrmsd()` plot showing only the upper half because of the mirroring of the values.

argument name	default value	description
xrmsdValues	<i>none</i>	Input matrix (three rows: x-values, y-values, RMSD-values). Can be generated by function <code>load_xrmsd()</code> .
printLegend	TRUE	If TRUE, a legend is printed on the right hand side.
xaxisRange	NA	A vector of boundaries for the x-snapshots.
yaxisRange	NA	A vector of boundaries for the y-snapshots.
colours	NA	User specified vector of colours to be used for plotting.
rmsdUnit	"nm"	Specifies, in which unit the RMSD values are given.
barPlot	FALSE	Boolean, indicating whether the plot is to be made without any additional information.
...	<i>none</i>	Additional arguments.

## Additional functions and the bash interface

Given that the plotting functions expect input to be stored in a defined data structure, the step of loading and parsing data from the text input files has been implemented in separate loading functions. Currently, they support GROMOS, GROMACS and AMBER file formats and further developments are planned to cover additional ones as well.

In order to allow for direct calls from bash scripts, users might use the Rscript interface located in the folder 'bash' which serves as a wrapper shell. Pictures in the file formats PNG, TIFF or PDF can be used provided that the users' R installation supports them. If `help=TRUE` is set, all the other options are ignored and a full list of options for every command is printed. In general, the names of the arguments of the functions are the same for calls by script. The syntax for these calls is `Rscript MDplot_bash.R {function name} [argument1=...] [argument2=...]`, which can be combined with bash variables (see below). The file path can be given in an absolute manner or relative to the Rscript folder path. The package holds a file called 'bash/test.sh' which contains several examples.

```
#!/bin/bash
# clusters
Rscript MDplot_bash.R clusters files=../extdata/clusters_example.txt.gz \
                             title="Cluster analysis" size=900,900 \
                             outformat=tiff outfile=clusters.tiff \
                             clustersNumber=7 \
                             names=WT,varA,varB,varC2,varD3,varE4

# xrmsd
Rscript MDplot_bash.R xrmsd files=../extdata/xrmsd_example.txt.gz title="XRMSD" \
                             size=1100,900 outformat=pdf outfile=XRMSD.pdf \
                             xaxisRange=75,145

# ramachandran
Rscript MDplot_bash.R ramachandran files=../extdata/ramachandran_example.txt.gz \
                             title="Ramachandran plot" size=1400,1400 resolution=175 \
                             outformat=tiff outfile=ramachandran.tiff angleColumns=1,2 \
                             bins=75,75 heatFun=norm printLegend=TRUE plotType=fancy
```

## Loading functions

In order to ease data preparation, loading functions have been devised which are currently able to load the output of standard GROMOS, GROMACS and AMBER analysis tools and store these data such, that they can be interpreted by the **MDplot** plotting functions.<sup>2</sup> Loading functions are named after their associated plotting function with 'load\_' as prefix. For other molecular dynamics engines than the aforementioned ones, the user has to specify how their output should be read. However, in case other file formats are requested we appreciate suggestions, requests and contributions (to be made on our GitHub page). For detailed descriptions of the data structures used, we refer to the manual pages of the loading functions and the respective examples. For storage reasons the example input files have been compressed using gzip with R being able to load both compressed and uncompressed files.

<sup>2</sup>Functions `load_timeseries()` and `load_TICurve()` do not require engine-specific loading and function `noe()` is only available for GROMOS because no input files for the other engines could be retrieved.

## Conclusions

In this paper we have presented the package **MDplot** and described its application in the context of molecular dynamics simulation analysis. Automated figure generation is likely to aid in the understanding of results at the first glance and may be used in presentations and publications. Planned extensions include both the integration of new functionalities such as a DISICL (secondary structure classification (Nagy and Oostenbrink, 2014a,b)) as well as the provision of loading interfaces for additional molecular dynamics engines. Further developments will be published on the projects' GitHub page and on CRAN.

## Acknowledgements

The authors would like to thank Prof. Friedrich Leisch for his useful comments and guidance in the package development process, Markus Fleck for providing GROMACS analysis files, Silvia Bonomo for her help in dealing with AMBER, Sophie Krecht for her assistance in typesetting and Jamie McDonald for critical reading of the manuscript.

## Funding

This work was supported by the European Research Council (ERC; grant number 260408), the Austrian Science Fund (FWF; grant number P 25056) and the Vienna Science and Technology Fund (WWTF; grant number LS08-QM03).

## Bibliography

- H. M. Berman, J. Westbrook, Z. Feng, G. Gilliland, T. N. Bhat, H. Weissig, I. N. Shindyalov, and P. E. Bourne. The Protein Data Bank. *Nucleic Acids Research*, 28(1):235, 2000. doi: 10.1093/nar/28.1.235. [p1]
- B. Brooks, C. Brooks, A. MacKerell, L. Nilsson, R. Petrella, B. Roux, Y. Won, G. Archontis, C. Bartels, S. Boresch, A. Caffisch, L. Caves, Q. Cui, A. Dinner, M. Feig, S. Fischer, J. Gao, M. Hodoseck, W. Im, K. Kuczera, T. Lazaridis, J. Ma, V. Ovchinnikov, E. Paci, R. Pastor, C. Post, J. Pu, M. Schaefer, B. Tidor, R. M. Venable, H. L. Woodcock, X. Wu, W. Yang, D. York, and M. Karplus. CHARMM: The Biomolecular Simulation Program. *Journal of Computational Chemistry*, 30(10):1545–1614, July 2009. ISSN 0192-8651. doi: 10.1002/jcc.21287. [p1]
- F. Comoglio and M. Rinaldi. Rknots: topological analysis of knotted biopolymers with R. *Bioinformatics*, 28(10):1400, 2012. doi: 10.1093/bioinformatics/bts160. [p1]
- W. D. Cornell, P. Cieplak, C. I. Bayly, I. R. Gould, K. M. Merz, D. M. Ferguson, D. C. Spellmeyer, T. Fox, J. W. Caldwell, and P. A. Kollman. A Second Generation Force Field for the Simulation of Proteins, Nucleic Acids, and Organic Molecules. *Journal of the American Chemical Society*, 117(19):5179–5197, May 1995. ISSN 0002-7863. doi: 10.1021/ja00124a002. [p1]
- A. P. Eichenberger, J. R. Allison, J. Dolenc, D. P. Geerke, B. A. C. Horta, K. Meier, C. Oostenbrink, N. Schmid, D. Steiner, D. Wang, and W. F. van Gunsteren. GROMOS++ Software for the Analysis of Biomolecular Simulation Trajectories. *Journal of Chemical Theory and Computation*, 7(10):3379–3390, Oct. 2011. ISSN 1549-9618. doi: 10.1021/ct2003622. [p1]
- B. J. Grant, A. P. C. Rodrigues, K. M. ElSawy, J. A. McCammon, and L. S. D. Caves. Bio3d: an R package for the comparative analysis of protein structures. *Bioinformatics*, 22(21):2695–2696, 2006. ISSN 1367-4803, 1460-2059. doi: 10.1093/bioinformatics/btl461. [p1]
- W. Kabsch and C. Sander. Dictionary of protein secondary structure: Pattern recognition of hydrogen-bonded and geometrical features. *Biopolymers*, 22(12):2577–2637, Dec. 1983. ISSN 1097-0282. doi: 10.1002/bip.360221211. [p5]
- J. Kirkwood. Statistical mechanics of fluid mixtures. *Journal of Chemical Physics*, pages 300–313, 1935. [p17]
- C. Margreitter and C. Oostenbrink. Optimization of Protein Backbone Dihedral Angles by Means of Hamiltonian Reweighting. *Journal of Chemical Information and Modeling*, 56(9):1823–1834, sep 2016. doi: 10.1021/acs.jcim.6b00399. [p11]

- R. T. McGibbon, K. A. Beauchamp, M. P. Harrigan, C. Klein, J. M. Swails, C. X. Hernández, C. R. Schwantes, L.-P. Wang, T. J. Lane, and V. S. Pande. MDTraj: A Modern Open Library for the Analysis of Molecular Dynamics Trajectories. *Biophysical Journal*, 109(8):1528 – 1532, 2015. doi: 10.1016/j.bpj.2015.08.015. [p1]
- G. Nagy and C. Oostenbrink. Dihedral-Based Segment Identification and Classification of Biopolymers II: Polynucleotides. *Journal of Chemical Information and Modeling*, 54(1):278–288, jan 2014a. doi: 10.1021/ci400542n. [p21]
- G. Nagy and C. Oostenbrink. Dihedral-based segment identification and classification of biopolymers I: proteins. *Journal of Chemical Information and Modeling*, 54(1):266–277, Jan. 2014b. ISSN 1549-960X. doi: 10.1021/ci400541d. [p21]
- J. C. Phillips, R. Braun, W. Wang, J. Gumbart, E. Tajkhorshid, E. Villa, C. Chipot, R. D. Skeel, L. Kalé, and K. Schulten. Scalable molecular dynamics with NAMD. *Journal of Computational Chemistry*, 26(16):1781–1802, Dec. 2005. ISSN 1096-987X. doi: 10.1002/jcc.20289. [p1]
- S. Pronk, S. Páll, R. Schulz, P. Larsson, P. Bjelkmar, R. Apostolov, M. R. Shirts, J. C. Smith, P. M. Kasson, D. v. d. Spoel, B. Hess, and E. Lindahl. GROMACS 4.5: a high-throughput and highly parallel open source molecular simulation toolkit. *Bioinformatics*, 29(7):845–854, Apr. 2013. ISSN 1367-4803, 1460-2059. doi: 10.1093/bioinformatics/btt055. [p1]
- G. N. Ramachandran, C. Ramakrishnan, and V. Sasisekharan. Stereochemistry of polypeptide chain configurations. *Journal of Molecular Biology*, 7:95–99, July 1963. ISSN 0022-2836. [p11]
- N. Schmid, C. D. Christ, M. Christen, A. P. Eichenberger, and W. F. van Gunsteren. Architecture, implementation and parallelisation of the GROMOS software for biomolecular simulation. *Computer Physics Communications*, 183(4):890–903, Apr. 2012. ISSN 0010-4655. doi: 10.1016/j.cpc.2011.12.014. [p1]
- L. Skjærven, X.-Q. Yao, G. Scarabelli, and B. J. Grant. Integrating protein structural dynamics and evolutionary analysis with Bio3D. *BMC Bioinformatics*, 15(1), dec 2014. doi: 10.1186/s12859-014-0399-6. [p1]

Christian Margreitter  
Institute of Molecular Modeling and Simulation  
University of Natural Resources and Life Sciences (BOKU)  
Austria  
[christian.margreitter@gmail.com](mailto:christian.margreitter@gmail.com)

Chris Oostenbrink  
Institute of Molecular Modeling and Simulation  
University of Natural Resources and Life Sciences (BOKU)  
Austria  
[chris.oostenbrink@boku.ac.at](mailto:chris.oostenbrink@boku.ac.at)