# AnalysisTools user manual

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# 1. Introdution

This software package is a set of utilities aimed to analyse trajectories of coarse-grained simulations. It was initially developed to work with DL\_MESO simulation package. However, it works with vtf trajectory files, so it can be used to analyse any vtf trajectories.

The emphasis is placed on assembly of molecules, e.i., self-assembly of polymers. It therefore include utilities to determine if molecules are in aggregates and to calculate various properties of aggregates. The utilities only calculate desired quantities and write them to output text files, there is no plotting or visualisation.

Examples of the resulting data can be seen in the author's thesis (Šindelka, Karel: The study of the association behavior of the amphiphilic copolymers in solutions containing low molar compounds by means of computer simulations, dissertation thesis, Charles University, 2018) as well as in papers in impacted journals (e.g., doi:10.1039/C8CP05907A, 10.1134/S1811238217010052, or 10.1007/s00396-017-4090-0).

# 2. Installation

All programs can be compiled using cmake which generates Makefile and subsequently running make. It requires C and FORTRAN compilers. The compilation should be done in a separate directory, such as build.

To create the Makefule and compile all utilities, simply run the following command (assuming build is a subdirectory of AnalysisTools root directory):

```
cmake -G "Unix Makefiles" ../
```

The binaries will be in 'bin' subdirectory of 'build'.

Cmake variable <code>-DCMAKE\_BUILD\_TYPE=Debug</code> can be used to compile the version of utilities for debugging.

To compile individual C programs using gcc, run (assuming you are in a subdirectory of AnalysisTools root directory):

gcc -03 -lm ../AnalysisTools.c ../Error.c ../Options.c ../Utility/program.c.

# 3. Format of input/output files

All utilities read information about the system from vsf/vcf files (formatted as described below) and FIELD file (input file for DL\_MESO simulation package). All system information is read from the vsf structure file (Section 3.1) and from the vcf coordinate file (Section 3.2). One vtf file containing a structure and coordinate sections can also be used. The FIELD file is only used when bead charge and/or mass is missing from the vsf structure file. The utilities consider only bead types that are present in the vcf coordinate file (i.e., bead types present in the vsf file but not in the vcf file are not seen by the utilities). A description of how the system data are read is shown in Chapter 5.

Both vsf and vcf files can be generated using a traject utility provided by the DL\_MESO simulation package (traject-v2\_5 and traject-v2\_6 provided here are modified versions from earlier DL\_MESO simulation package versions). Both structure and coordinates can be in one file (with vtf extension) – this file can be used instead of separate vsf and vcf files.

All utilities assume cuboid simulation box with dimensions from 0 to N, where N is the side length of the box (which can be different in all three dimensions).

### 3.1 vsf structure file

The structure file contains all information about all beads and bonds except for their Cartesian coordinates. The utilities are written for a vsf file created by a traject utility (for DL\_MESO versions from 2.5 to 2.7), but other vsf files should work as long as they adhere to the following format.

A vsf file is divided into two parts. The first part contains bead definitions. Each line contains the description of a single bead and follows these rules:

- the line starts with atom (or just a)
- the second string is a bead index number that starts from 0 and increases with every subsequent line (the last bead definition line therefore shows the total number of beads in the simulation)
- the line contains bead name as name <char(8)>
- the first bead definition line may contain **default** instead of the index number; every bead that is not explicitly written in the bead definition lines is has the

default name

- if the bead is in a molecule, the line contains molecule name (resname <char(8)>) and molecule id (resid <int>) that starts from 1
- mass and charge keywords are read if present (otherwise the mass and charge of beads is read from FIELD)
- other keywords are ignored

The following is an example of bead definition lines containing all required data:

```
atom default name Bead_A
atom 0 name Bead_B
atom 3 name Bead_C resname Mol_A 1
atom 4 name Bead_C resname Mol_A 1
atom 6 name Bead_C resname Mol_A 2
atom 7 name Bead_C resname Mol_A 2
atom 8 name Bead_D resname Mol_B 3
atom 9 name Bead_D resname Mol_B 3
```

In this example, there are four bead types (named Bead\_A, Bead\_B, and Bead\_C, and Bead\_D) and 10 beads in all (with indices from 0 to 9). Beads with indices 1, 2, and 5 are of the default type (Bead\_A). There are two molecule types named Mol\_A and Mol\_B with molecule indices 1 to 3. All molecules with the same name must have the same structure, i.e., the same number of beads and the same bond connectivity.

The second part of a **vsf** file contains bonds definitions and must be preceded by a blank line. Each bond definition line follows these rules:

- the line starts with bond (or just b)
- bond between two beads is specified by their indices separated by a colon (there cannot be a space between the first number and the colon)

The following is an example of bead definition lines that complement the aboveshown bead definition lines:

```
bond 3: 4
# possible comment
bond 6: 7
bond 8: 9
```

In this example, there three bonds between beads with indices 3 and 4, 6 and 7, 8 and 9.

Blank lines and comments (lines beginning with #) are allowed in both parts of the vsf file.

### 3.2 vcf coordinate file

The coordinate file contains Cartesian coordinates of the beads and the size of the cuboid simulation box. Coordinates are read from a vcf file containing either ordered timesteps (Section 3.2.1) or indexed timesteps (Section 3.2.2).

An ordered vcf file must contain all beads defined in the vsf file, while an indexed vcf file can contain only a subset of defined beads. Both indexed and ordered vcf files contain a line before every timestep specifying the file type — timestep ordered or timestep indexed (the keyword timestep can be omitted). In both ordered and indexed vcf files, the size of the simulation box is given by a line pbc <float> <float> <float> which is located before the first coordinate block. Only timestep and pbc lines are read before the first coordinates (everything else is ignored), so vtf file can be used instead of a vcf file.

The vcf file may contain comment lines (beginning with #) and blank lines between timesteps, but the coordinate block must be continuous.

#### 3.2.1 Ordered coordinate file

Coordinate lines in ordered vcf file contain only the Cartesian coordinates of the beads in the form <float> <float> <float>. The beads are written in ascending order of their indices as defined in the vsf file. The following is an example of an ordered vcf file:

```
# any number of comments or blank lines
timestep ordered
pbc 10 10 10
0.0 0.0 0.0
0.5 0.5 0.5
...
# comments between timesteps
timestep ordered
# another comment
1.0 1.0 1.0
1.5 1.5 1.5
...
```

In this example, the simulation box is cubic with side length of 10. Beginnings of two timesteps are represented by coordinates of the first two beads with indices 0 and 1 (as defined in the vsf file).

#### 3.2.2 Indexed coordinate file

Indexed coordinate files contains not only Cartesian coordinates, but also bead indices (preceding the coordinates). Therefore an indexed timestep does not have to contain all beads in the vsf structure file. Moreover, the beads do not have to be ordered according to their ascending indices. The following is an example of an ordered vcf file:

```
timestep indexed
pbc 10 10 10
2 0.5 0.5 0.5
21 0.0 0.0 0.0
...

# comments between timesteps
timestep indexed
# another comment
21 1.0 1.0 1.0
2 1.5 1.5 1.5
```

# any number of comments or blank lines

This example is similar to that for the ordered vcf file, but two beads have indices 2 and 21 instead of 0 and 1.

# 3.3 Aggregate file (agg)

The aggregate file with agg extension is generated using Aggregates utility. The file contains information about the number of aggregates in each timestep and which molecules and monomeric beads belong to which aggregate. It serves as an additional input file for utilities that calculate properties of whole aggregates — agg file is therefore linked to the vcf that was used to generate it.

The agg file is a simple text file. The first line contains the command used to generate it – parts of this command can be necessary for subsequent analysis of aggregates. The second line is blank and from the third line the data for individual timesteps are shown. Each timestep follows these rules:

- each timestep starts with Step: <int> (only Step keyword is read by the utilities)
- the second line contains the number of aggregates in the given timestep and followed by a blank line

- there are two lines for each aggregate:
  - (1) number of molecules in the aggregate followed by their indices taken from the vsf file
  - (2) number of monomeric beads in the aggregate followed by their indices taken from the vsf file
- no blank or comment lines are allowed inside the aggregate block
- all molecules present in the vcf file used to generate this file must be present in every timestep; here, aggregate can also refer to dissolved molecules Following is an example of an agg file:

```
Aggregates in.vcf 1 1 out.agg A
```

```
Step: 1
2
2 : 1 3
3 : 10 100 1000
1 : 2
1 : 20
Step: 2
1
3 : 1 2 3
4 : 10 20 100 2000

Last Step: 2
```

In this example, command Aggregates in.vcf 1 1 out.agg A was used to generate the file (see Section 4.3 for details about this utility). There are two timesteps here – the first contains two aggregates (although one of them is a free, dissolved molecule) and the second a single aggregate. As an example, the aggregate in the second step contains three molecules with indices 1, 2, and 3 (taken from the vsf file) and four monomeric beads (i.e., solvent or counterions) with indices 10, 20, 100, and 2000 (again, taken from the vsf file).

Besides using this file for further analysis using other utilities, the indices can be used in vmd to visualize, e.g., only a specific aggregate.

# 4. Utilities

All utilities have command line help with short description when -h argument is used. Besides -h, most of the utilities have several standard command line options that are the same. The standard options can be used with any utility unless stated otherwise.

Standard options		
-i <name></name>	use custom vsf file instead of traject.vsf	
-ν	verbose output that provides information about all bead and molecule types	
-V	detailed verbose output that provides information about all individual molecules as well as about bead and molecule types	
-s	run silently, i.e., without any output at all (overrides $\neg v$ and $\neg V$ options)	
script -h	do not rewrite terminal line (useful if output is routed to a file) print help and exit	

# 4.1 AddToSystem

This utility takes an existing system specified by vcf coordinate and vsf structure files and adds new beads into it. The new beads replace neutral unbonded ones with the lowest indices (as ordered in the vsf file) from the original system. If molecules are added, AddToSystem places them at the end (for the sake of DL\_MESO which requires molecules to be after unbonded beads). The utility generates vcf and vsf files for the new system.

AddToSystem does not check whether there are enough unbonded neutral beads to be replaced by the new beads; the utility will most likely not show an error, but the resulting vcf and vsf files won't be right.

Coordinates of the new unbonded beads are ruled by the -ld, -hd, and -bt options (either the first bead or the geometric centre of the new molecules obey these options). The coordinates of the remaining beads in a molecule are governed by the provided coordinates. The molecules are added with a random orientation.

If -ld and/or -hd options are used, they must accompanied by the -bt option. The structure and number of added molecules and monomeric beads are read

from a FIELD-like file. This file must contain species section followed by molecule section as described in the DL\_MESO simulation package.

The species section contains the number of bead types and their properties:

```
species <int>
<name> <mass> <charge> <number of unbonded beads>
```

The first line must start with **species** keyword followed by the number of bead types. For each bead type a single line must contain the name of the bead, its mass and charge, and a number of these beads that are not in a molecule (i.e., monomeric beads).

The molecule section that must be behind the species section contains information about structure and numbers of molecules to be added:

```
molecule <int>
                                          Number of types of molecules
                                          Name of the first molecule type
<name>
                                          Number of these molecules
nummols <int>
                                          Number of beads in these molecules
beads <int>
                                          One line for each of the <int> beads
<bead name> <float> <float> <float>
                                            specifying bead name and its
<bead name> <float> <float> <float>
                                            Cartesian coordinates
bonds <int>
                                          Number of bonds in these molecules
                                          One line for each of the <int> bonds
<string> <int> <int>
                                            containing arbitrary string and
                                            indices connected beads
<string> <int> <int>
                                          Anything beyond here is ignored
                                          Description of a molecule is finished
finish
```

The molecule keyword specifies the number of molecule types, that is the number of finish keywords that must be present. The <beathermal the species section. The arbitrary <string> in the bonds is ignored by AddToSystem (it is a relic from the DL\_MESO simulation package, where the <string> specifies a type of bond). The indices in bond lines run from 1 to the number of beads in the molecules and are ordered according to the beads part of the section. Because molecule section in the FIELD file from DL\_MESO can also include bond angles and dihedral angles, anything beyond the last bond line is ignored (until the finish keyword is read).

If no molecules are to be added, the line molecule 0 must be still be present in the file.

The following is an example of the FIELD-like file:

```
species 3
```

```
1.0
        1.0 0
Α
    1.0 0.0
    1.0 -1.0 30
molecule 2
Dimer
nummols 10
beads 2
A 0.0 0.0 0.0
A 0.5 0.0 0.0
bonds 1
harm 1 2
finish
surfact
nummols 10
beads 3
A 0.0 0.0 0.0
B 0.5 0.0 0.0
B 1.0 0.0 0.0
bonds 2
harm 1 2
harm 2 3
angles 1
harm 1 2 3
finish
```

In this example, 30 unbonded (or monomeric) negatively charged beads called CI are added as well as 20 molecules – 10 molecules called Dimer and 10 molecules called surfact. Dimer molecules contain two A beads and one bond each; surfact molecules contain three beads and two bonds each. The part starting with angles and ending with finish is ignored. All in all, 80 beads are added – 30 CI, 30 A, and 20 B beads.

The utility creates the vcf and vsf files with the new system and can also write the coordinates into a xyz file.

Usage:

AddToSystem <input.vcf> <input add> <out.vcf> <out.vsf> <options>

#### Mandatory arguments

<input>

input coordinate file (either vcf or vtf format)

<input add=""/> <out.vcf> <out.vsf></out.vsf></out.vcf>	FIELD-like file specifying additions to the system output vcf coordinate file for the new system output vsf structure file for the new system
	T v
Non-standard option	
-st <int></int>	timestep to add new beads to (default: 1)
-xyz <name></name>	save coordinates to xyz file
-ld <float></float>	lowest distance from beads specified by -bt option
-hd <float></float>	highest distance from beads specified by -bt option
-bt <bead names=""></bead>	bead types to use in conjunction with -ld and/or -hd op-
	tions

# 4.2 AngleMolecules

This utility calculates angles between beads in each molecule of specified molecule type(s). The beads do not have to be connected, so the angle does not have to be between two bonds.

The angle is specified by three bead indices taken from the vsf file (-n option) These indices are from 1 to N, where N is the number of beads in the molecule type. Generally, the numbering of beads inside a molecule is made according to the first molecule of the given type in vsf file. For example, assume that beads of the first molecule called mol in the vsf file are ordered A (vsf index 123), B (vsf index 124), C (vsf index 200). Then, bead A is 1, bead B is 2, and C is 3.

More than one angle can be specified (i.e., a multiple of three numbers have to be supplied to the -n option.). For example, assuming indices 1 2 3 are specified (default if -n option is not used), the angle will be between lines defined by beads with indices 1 2 and 2 3. The angle is calculated in degrees and is between 0 and 180°.

The utility calculates distribution of angles for each specified trio of bead indices for each molecule type and prints overall averages at the end of <output> file. If -a option is used, it can also write all the angles for all individual molecules in each timestep (i.e., time evolution of the angle for each individual molecule).

Usage:

AngleMolecules <input> <mol name(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<mol name(s)=""></mol>	molecule name(s) to calculate angles for	

<output></output>	output file for distribution	
Non-standard options		
joined	specify that <input/> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)	
-a <name></name>	write all angles for all molecules in all timesteps to <name></name>	
-n <ints></ints>	multiple of three indices for angle calculation (default: 1 2 3)	
-st <int></int>	starting timestep for calculation (default: 1)	

#### Format of output files:

- (1) <output> distribution of angles
  - first line: command used to generate the file
  - second line: planes to calculate angles between (the dash-separated numbers correspond to indices inside every molecule and are the same as the arguments to the -n option) with the numbers in brackets corresponding to nth column of data for each molecule type
  - third line: numbering of columns (i.e., column headers)
    - first is the centre of each bin in angles (governed by <width>); i.e., if <width> is 5°, then the centre of bin 0 to 5° is 2.5, centre of bin 5 to 10° is 7.5 and so on
    - the rest are for the calculated data: the range for each molecule type specifies which column numbers correspond to the calculated angles for that particular molecule type and the order of angles is given by the second line
  - last two lines: arithmetic means for each calculated angle (last line) and headers (second to last line) that again give range of columns in the last line for each molecule type
- (2) -a <name> all angles for all molecules in all timesteps
  - first and second lines are the same as for <output>
  - third lines: column headers
    - first is simulation timestep
    - the rest are the calculated data: the range for each molecule type corresponds to the number of molecules of the given type times the number of calculated angles; for each molecule the angles are ordered according to the second line

# 4.3 Aggregates and Aggregates-NotSameBeads

These utilities determine which molecules belong to which aggregates according to a simple criterion: two molecules belong to the same aggregate if they share at least a specified number of contact pairs. A contact pair is a pair of two beads belonging to different molecules which are closer than a specified distance. The information is written in agg format described in Section 3.3.

The number of contact pairs, the distance, and bead type(s) to use for aggregate determination are all arguments of the utilities. Any molecule type(s) can be excluded from aggregate determination (-x <mol name(s)> option); they are also excluded from the output agg file). Moreover, any molecules close to specified molecule(s) can be excluded (-xm <mol name(s)> option); here, 'close' means any of the bead types used aggregate determination is closer than <distance> to any bead of the specified molecule.

Also, periodic boundary conditions can be removed from whole aggregates and the new coordinates saved to an indexed vcf file (-j option). Therefore aggregates will not be split by simulation box boundaries when, for example, visualizing the molecules with vmd.

While the Aggregates utility uses all possible pairs of given bead types, Aggregates-NotSameBeads does not use same-type pairs. For example, if bead types A and B are given, Aggregates will use all three possible bead type pairs (i.e., A-A, A-B, and B-B), but Aggregates-NotSameBeads will use only A-B bead type pairs.

Usage:

Aggregates (or Aggregates-NotSameBeads) <input> <distance> <contacts> <output.agg> <bead type name(s)> <options>

Mandatory arguments		
<pre><input/></pre>	input coordinate file (either vcf or vtf format)	
<distance></distance>	minimum distance for two beads to be in contact (thus constituting a contact pair)	
<contacts></contacts>	minimum number of contact pairs between two molecules to be in one aggregate	
<pre><output.agg></output.agg></pre>	output agg file (must end in .agg) with aggregate information	
<pre><bead type(s)=""></bead></pre>	bead type name(s) to use for determining contact pairs (at least two for Aggregates-NotSameBeads)	
<pre><options></options></pre>	,	
Non-standard options		
-x <mol name(s)=""></mol>	exclude specified molecule type(s) from aggregate determination (and from the output agg file)	
<pre>-xm <mol name(s)=""> -j <output.vcf></output.vcf></mol></pre>	exclude molecules that are close to specified molecule(s) output vcf file with coordinates of joined aggregates (i.e., without periodic boundary conditions)	

4. Utilities 4.4. Average

# 4.4 Average

This utility uses the binning method to analyse data in a text file. It does not use any of the standard options and prints the result only to standard output (e.i., screen).

Average calculates average, statistical error, and estimate of the autocorrelation time  $\tau$ . Empty lines and comments (lines beginning with #) are skipped. Average prints to standard output (i.e., the screen) four numbers: <n\_blocks> <average> <std error> <tau estimate>:

<n_blocks></n_blocks>	number of blocks used for the binning analysis
<average></average>	simple arithmetic average
<std error=""></std>	one- $\sigma$ statistical error
<tau estimate=""></tau>	estimate of autocorrelation time $\tau$

The average of an observable  $\mathcal{O}$  is a simple arithmetic mean:

$$\langle \mathcal{O} \rangle = \frac{1}{N} \sum_{i=1}^{N} \mathcal{O}_i,$$
 (4.1)

where N is the number of measurements and the subscript i denotes individual measurements. If the measurements are independent (i.e., uncorrelated), the statistical error is given by:

$$\epsilon^2 = \frac{\sigma_{\mathcal{O}_i}^2}{N},\tag{4.2}$$

where  $\sigma_{\mathcal{O}_i}^2$  is the variance of the individual measurements,

$$\sigma_{\mathcal{O}_i}^2 = \frac{1}{N-1} \sum_{i=1}^N (\mathcal{O}_i - \langle \mathcal{O} \rangle)^2. \tag{4.3}$$

For correlated data, the autocorrelation time,  $\tau$ , representing the number of steps between two uncorrelated measurements must be determined. Every  $\tau$ -th measurement is uncorrelated, so the equation (4.2) can then be used to estimate the error.

A commonly used method to estimate the autocorrelation time is the binning (or block) method. In this method, the correlated data are divided into  $N_{\rm B}$  non-overlapping blocks of size k ( $N=kN_{\rm B}$ ) with per-block averages,  $\mathcal{O}_{{\rm B},n}$ , defined as:

$$\mathcal{O}_{B,n} = \frac{1}{k} \sum_{\substack{i=1+\\(n-1)k}}^{kn} \mathcal{O}_i.$$
 (4.4)

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If  $k \gg \tau$ , the blocks are assumed to be uncorrelated and equation (4.2) can be used:

$$\epsilon^2 = \frac{\sigma_{\rm B}^2}{N_{\rm B}} = \frac{1}{N_{\rm B}(N_{\rm B} - 1)} \sum_{n=1}^{N_{\rm B}} (\mathcal{O}_{{\rm B},n} - \overline{\mathcal{O}})^2.$$
 (4.5)

An estimate of the autocorrelation time can be obtained using the following formula:

$$\tau_{\mathcal{O}} = \frac{k\sigma_{\mathcal{B}}^2}{2\sigma_{\mathcal{O}_i}^2}.\tag{4.6}$$

A way to quickly estimate a 'real' value of  $\tau$  is to use a wide range of <n\_blocks> value and plot the <tau estimate> values as a function of <n\_blocks>. Because the number of data points in one block of the binning analysis should be significantly larger than  $\tau$  (e.g., ten times larger), plotting f(x) =(number of data lines in the file)/(10x) will produce an exponential function that intersects the <tau estimate> line. A value of <tau estimate> near the intersection (but to the left, where the exponential is above <tau average>) can be used as a good estimate of  $\tau$ .

Usage:

Average <input> <column> <discard> <n\_blocks>

Mandatory arguments		
<input/>	input text file	
<column></column>	column number in <input/> for data analysis	
<discard> <n_blocks></n_blocks></discard>	number of lines to discard from the beginning of <input/> number of blocks for binning analysis	

# 4.5 BondLength

This utility calculates a distribution of bond lengths in specified molecule type(s) and it also can calculate distribution of distances between any two beads in the molecules.

For each of the specified molecule type(s), BondLength calculates bond lengths between all different types of connected bead pairs. For example, assume two linear molecule types Mol\_1 and Mol\_2, both composed of bead types A and B. Mol\_1 is connected like this: A-A-B; Mol\_2 like this: A-B-B. If both molecule types are used, BondLength calculates for each molecule type distribution of lengths for bonds A-A, A-B, and B-B (separate for each molecule even though the molecules share the same bead types).

4. Utilities 4.5. BondLength

To calculate the distribution of distances between specific (possibly unconnected) beads in a molecule, use -d option which takes as arguments pairs of bead indices (according to the order of beads in the molecule in vsf - similarly to the -n option in AngleMolecules, i.e., Section 4.2). More than one pair can be specified. These indices are the same for all <mol name(s)>. If an index higher than the number of beads in the molecule is provided, the utility takes the last bead of the molecule (i.e., the highest index). For example, using -d file.txt 1 2 1 9999 would write two distributions for each <mol name(s)> into <file.txt>: distribution of distances between the first and the second bead in each <mol name(s)> and between the first bead and the last one (or the 9999th bead).

In both cases, BondLength appends at the end of the file minimum and maximum bond lengths/distances.

Usage:

BondLength <input> <width> <output> <mol name(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<width></width>	width of each bin of the distribution	
<output></output>	output file with distribution of bond lengths	
<mol name(s)=""></mol>	molecule name(s) to calculate bond lengths for	
Non-standard options		
-st <int></int>	starting timestep for calculation (default: 1)	
-d <out> <ints></ints></out>	write distribution of distances between specified beads in	
	each specified molecule to <out></out>	
-w <double></double>	warn if a bond length exceeds <double> (default: half a box</double>	
	length)	

#### Format of output files:

- (1) **<output>** distribution of bond length between all bead pairs
  - first line: command used to generate the file
  - second line: column headers
    - first is the centre of each bin (governed by <width>); i.e., if <width> is 0.1, then the centre of bin 0 to 0.1 is 0.05, centre of bin 0.1 to 0.2 is 0.15, etc.
    - the rest are for the calculated data: for each molecule type, there is a list of column numbers corresponding to all possible bead type pairs in the molecule; if no beads of the given types are connected, the data column contains nan
  - next lines: the calculated data
  - second to last line: column headers for minimal and maximal bond lengths
    - for each molecule, all the possible beads pairs are again listed

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- for each pair, the column number corresponds to the minimum, while the next column is always the maximum

- last line: commented out (i.e., the line starts with #) minimal and maximal bond lengths
- (2) -d <output> <ints> distribution of distances between specified beads
  - first line: command used to generate the file
  - second line: order of beads (given only by bead names) in each molecule
  - third line: column headers
    - first is again the centre of every bin
    - the rest are for the calculated data: for each molecule type, there is a list of column numbers corresponding to the given pairs of bead indices in the molecule (and to the -d option's arguments); the numbers also correspond to the order of beads in the previous line
  - next lines: the calculated data
  - second to last line: column headers for minimal and maximal bond lengths
    - for each molecule, all the possible beads pairs are again listed
    - for each pair, the column number corresponds to the minimum, while the next column is always the maximum
  - last line: commented out (i.e., the line starts with #) minimal and maximal bond lengths

# 4.6 Config

This utility creates DL\_MESO CONFIG file. It requires input coordinate file with all beads; otherwise the utility will still run, but will produce incomplete CONFIG. Usage:

Config <input.vcf> <options>

Mandatory argument		
<input/>	input coordinate file (either vcf or vtf format)	
Non-standard option		
-st <int></int>	timestep for creating CONFIG file from (default: last step)	

There is also utility Config\_from\_xyz which takes a xyz coordinate file and creates the DL\_MESO CONFIG file. Because xyz file does not contain information about the simulation box, the resulting CONFIG file must be modified manually — Config\_from\_xyz prints x, y, and z into the output file where box dimensions should be.

Usage:

Config\_from\_xyz <input.xyz> <options>

Mandatory argument		
<input/>	input xyz coordinate file	
Non-standard option		
-st <int></int>	timestep for creating CONFIG file from (default: last step)	

# 4.7 DensityAggregates

This utility calculates radial density profiles (RDP, or radial number densities) for bead types in an aggregate with specified size (the number of molecules or aggregation number,  $A_{\rm S}$ ) from its centre of mass.

 $\mathrm{RDP}_i(r)$  of bead type i, where r is distance from an aggregate's centre of mass, is the number of these beads in a spherical shell between the distances r and  $r+\mathrm{d}r$  (in DensityAggregates,  $\mathrm{d}r$  is the <width> argument) divided by the volume of this shell. The distance written in the output file is always  $r+0.5\mathrm{d}r$ . Besides RDP, the output file also contains radial number profiles (i.e., the number of beads in the spherical shell not divided by the shell's volume) and one-sigma errors for both RDP and radial number profiles. The output file also contains a header describing what is in each column.

Instead of 'true' aggregate size, a number of molecules of specific type(s) can be used (-m option). For example, an aggregate containg 1 Mol\_A molecule and 2 Mol\_B molecules (i.e., three molecules in all) can be specified in several ways:

- (1) with  $\langle agg size(s) \rangle$  of 3;
- (2) with <agg size(s)> of 3 and -m Mol\_A Mol\_B;
- (3) with <agg size(s)> of 1 and -m Mol\_A; or
- (4) with <agg size(s) > of 2 and -m Mol\_B.

Care must be taken when different molecule types share the same bead type. If one bead type is in more molecule types, the resulting density for that bead type will be the sum of its densities from all molecule types it appears in. The -x option can overcome this – specific molecule types can be excluded from density calculations, i.e., density of beads in the excluded molecule types will not be calculated. For example, assume two molecule types – Mol\_1 and Mol\_2. Mol\_1 contains bead types A and B; Mol\_2 contains bead types A and C. Depending on whether and how the -x option is used, the utility will calculate:

(1) densities of A, B, and C beads (density of A beads is a sum from both molecules),

if no -x is used;

- (2) densities of only A and B beads (with A beads only from Mol\_1), if -x Mol\_2 is used;
- (3) densities of only A and C beads (with A beads only from Mol\_2), if -x Mol\_1 is used; or
- (4) no densities at all if -x Mol\_1 Mol\_2 is used.

Therefore, to be able to plot density of A beads from Mol\_1 and Mol\_2 separately, (2) and (3) should be used (i.e., DensityAggregates should be run twice).

Usage:

DensityAggregates <input> <output.agg> <width> <output.rho> <agg
size(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<input.agg></input.agg>	input agg file	
<width></width>	width of each bin of the distribution	
<pre><output.rho></output.rho></pre>	output file(s) (one per aggregate size) with automatic #.rho ending (# is aggregate size)	
<agg size(s)=""></agg>	aggregate size(s) (the number of molecules in an aggregate	
	or the aggregation number, $A_{\rm S}$ ) to calculate density for	
Non-standard options		
joined	specify that <input/> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)	
-n <int></int>	number of bins to average to get smoother density (default: 1)	
-st <int></int>	starting timestep for calculation (default: 1)	
-m <mol name(s)=""></mol>	instead of 'true' aggregate size, use the number of specified molecule type(s) in an aggregate	
-x <mol name(s)=""></mol>	exclude specified molecule type(s) (i.e., do not calculate density for beads in molecules <mol name(s)="">)</mol>	

#### Format of output files:

- (1) <output.rho> bead densities for one aggregate size
  - first line: command used to generate the file
  - second line: the order of data columns for each bead type rdp is radial density profile, rnp radial number profile and stderr are one-σ errors for rdp and rnp
  - third line: column headers
    - first is the centre of each bin (governed by <width>); i.e., if <width> is 0.1,

- then the centre of bin 0 to 0.1 is 0.05, centre of bin 0.1 to 0.2 is 0.15, etc.
- the rest are for the calculated data: each number specifies the first column with data for the given bead type (i.e., rdp column)

# 4.8 DensityBox

This utility calculates number density for all bead types in the specified direction of the simulation box. The density is calculated from 0 to box length in the given direction.

Usage:

DensityBox <input> <width> <output> <axis> <options>

Mandatory arguments			
<input/> <width> <output> <axis></axis></output></width>	input coordinate file (either vcf or vtf format) width of each bin of the distribution output file with automatic <axis>.rho ending direction in which to calculate density: x, y, or z</axis>		
Non-standar	Non-standard options		
-n <int> -st <int></int></int>	number of bins to average to get smoother density (default: 1) starting timestep for calculation (default: 1)		

#### Format of output files:

- (1) <output> bead densities
  - first line: command used to generate the file
  - second line: column headers
  - first is the centre of each bin (governed by <width>); i.e., if <width> is 0.1, then the centre of bin 0 to 0.1 is 0.05, centre of bin 0.1 to 0.2 is 0.15, etc.
  - the rest are for the calculated data: each number corresponds to the density of the specified bead type

# 4.9 DensityMolecules

This utility works similarly to DensityAggregates, only instead for whole aggregates, RDPs are calculated for individual molecules. Similarly to DensityAggregates, the output file(s) also contain statistical errors and radial number profiles.

By default, the utility calculates RDPs from the molecule's centre of mass, but any bead in the molecule (with an index from vsf – similar to –n option in AngleMolecules, Section 4.2) can be used instead (-c option).

Usage:

DensityMolecules <input> <width> <output> <mol name(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<width></width>	width of each bin of the distribution	
<output></output>	output file(s) (one per molecule type) with automatic	
	<mol_name>.rho ending</mol_name>	
<mol name(s)=""></mol>	molecule name(s) to calculate density for	
Non-standard options		
joined	specify that <input/> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)	
-n <int></int>	number of bins to average for smoother density (default: 1)	
-st <int></int>	starting timestep for calculation (default: 1)	
-c <name> <int></int></name>	use specified bead in a molecule <name> instead of its centre</name>	
	of mass	

#### Format of output files:

- (1) <output> bead densities for one molecule
  - first line: command used to generate the file
  - second line: the order of data columns for each bead type rdp is radial density profile, rnp radial number profile and stderr are one-σ errors for rdp and rnp
  - third line: column headers
    - first is the centre of each bin (governed by <width>); i.e., if <width> is 0.1, then the centre of bin 0 to 0.1 is 0.05, centre of bin 0.1 to 0.2 is 0.15, etc.
    - the rest are for the calculated data: each number specifies the first column with data for the given bead type (i.e., rdp column)

#### 4.10 DihedralMolecules

This utility calculates angles between specified planes in each molecule of specified molecule type(s). The planes in a molecule are arbitrary, so they can represent true dihedral angles or improper dihedrals.

The angle is specified by six bead indices (according to the order of beads in the molecule in vsf – similarly to the –n option in AngleMolecules, Section 4.2). The first three indices specify one plane and the next three the other. For example, assuming indices 1 2 3 4 5 6, the first plane is specified by the first three beads

in the molecule; second plane by the next three beads (beads 4 5 6). The default indices (i.e., if -n option is not used) are 1 2 3 2 3 4. More than one angle can be specified (i.e., a multiple of six numbers have to be supplied to the -n option.).

The utility calculates distribution of angles for each specified trio of bead indices for each molecule type and prints overall averages at the end of <output> file. If -a option is used, it can also write all the angles for all individual molecules in each timestep (i.e., time evolution of the angle for each individual molecule).

Usage:

#### DihedralMolecules <input> <mol name(s)> <options>

Mandatory arguments		
<pre><input/> <output> <mol name(s)=""></mol></output></pre>	input coordinate file (either vcf or vtf format) output file for distribution molecule name(s) to calculcate angles for	
Non-standard options		
joined	specify that <input/> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)	
-a <file></file>	write all angles for all molecules in all timesteps to <file></file>	
-n <ints></ints>	multiple of six indices for angle calculation (default: 1 2 3 2 3 4)	
-st <int></int>	starting timestep for calculation (default: 1)	

#### Format of output files:

- (1) <output> distribution of angles
  - first line: command used to generate the file
  - second line: calculated angles (the dash-separated numbers correspond to indices inside every molecule and are the same as the arguments to the -n option) with the numbers in brackets corresponding to nth column of data for each molecule type
  - third line: numbering of columns (i.e., column headers)
    - first is the centre of each bin in angles (governed by <width>); i.e., if <width> is 5°, then the centre of bin 0 to 5° is 2.5, centre of bin 5 to 10° is 7.5 and so on
    - the rest are for the calculated data: the range for each molecule type specifies which column numbers correspond to the calculated angles for that particular molecule type and the order of angles is given by the second line
  - last two lines: arithmetic means for each calculated angle (last line) and headers (second to last line) that again give range of columns in the last line for each

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molecule type

- (2) -a <file> all angles for all molecules in all timesteps
  - first and second lines are the same as for <output>
  - third lines: column headers
    - first is simulation timestep
    - the rest are the calculated data: the range for each molecule type corresponds to the number of molecules of the given type times the number of calculated angles; for each molecule the angles are ordered according to the second line

# 4.11 DistrAgg

This utility calculates average aggregate mass and aggregation number for each timestep (i.e., time evolution) and the averages over all timesteps. It calculates number, weight, and z averages. It also calculates distribution functions of aggregation sizes and volumes.

For a quantity  $\mathcal{O}$ , the number, weight, and z averages,  $\langle \mathcal{O} \rangle_{n}$ ,  $\langle \mathcal{O} \rangle_{w}$ , and  $\langle \mathcal{O} \rangle_{z}$ , respectively, are defined as

$$\langle \mathcal{O} \rangle_{\rm n} = \frac{\sum_{i} N_i \mathcal{O}_i}{N}, \quad \langle \mathcal{O} \rangle_{\rm w} = \frac{\sum_{i} N_i m_i \mathcal{O}_i}{\sum_{i} N_i m_i}, \text{ and } \quad \langle \mathcal{O} \rangle_{\rm z} = \frac{\sum_{i} N_i m_i^2 \mathcal{O}_i}{\sum_{i} N_i m_i^2}, \quad (4.7)$$

where N is the total number of measurements, i.e., the total number of aggregates for per-aggregate averages (or molecules for per-molecule averages);  $N_i$  is the number of measurements with the value  $\mathcal{O}_i$ , and  $m_i$  is mass of an aggregate i (or a molecule i).

Per-timestep averages are written to the <output avg> and overall averages are appended as comments (with commented legend) to both <output avg> and <output distr> files.

Number, weight, and z distribution functions of aggregate sizes,  $F_n(A_S)$ ,  $F_w(A_S)$ , and  $F_z(A_S)$  respectively, are defined as

$$F_{\rm n} = \frac{N_{A_{\rm S}}}{\sum_{A_{\rm S}} N_{i}} = \frac{N_{A_{\rm S}}}{N},$$

$$F_{\rm w} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{\sum_{A_{\rm S}} N_{i} m_{i}} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{\sum_{i=1}^{N} m_{i}} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{M}, \text{ and}$$

$$F_{\rm z} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}^{2}}{\sum_{A_{\rm S}} N_{i} m_{i}^{2}} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}^{2}}{\sum_{i=1}^{N} m_{i}^{2}},$$

$$(4.8)$$

where  $N_{A_{\rm S}}$  and  $m_{A_{\rm S}}$  stand for the number and mass, respectively, of aggregates with aggregate size  $A_{\rm S}$ ; M is the total mass of all aggregates. The equations are normalised so that  $\sum F_x(A_{\rm S}) = 1$ .

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Distribution of volume fractions of aggregates,  $\phi(A_S)$ , is defined (assuming all beads have the same volume) as

$$\phi(A_{\rm S}) = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{\sum_{i=1}^{N} n_i} = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{n},\tag{4.9}$$

where  $n_i$  is the number of beads in aggregate i and n is the total number of beads in all aggregates. If all beads have unit mass (as is often the case in dissipative particle dynamics), the volume distribution,  $\phi(A_S)$ , is the equal to the number distribution,  $F_n(A_S)$ . These distribution are written into the **<output distr>** file.

Lastly, DistrAgg can calculate number distribution of composition for aggregates with specified size(s) (-c option). This is a number distribution of the numbers of different molecule types in the aggregate. For example, if the simulation box contains molecule types Mol\_A and Mol\_B, aggregates with the same size can contain different numbers of these molecules, or different ratios of the numbers of Mol\_A to Mol\_B molecules,  $\xi = N_{\text{Mol\_A}}/N_{\text{Mol\_B}}$ . For now, DistriAgg can calculate this distribution only for aggregates containing two molecule types. The composition distribution is defined as

$$F_{\rm n}(\xi) = \frac{N_{\xi, A_{\rm S}}}{N_{A_{\rm S}}},$$
 (4.10)

where  $N_{\xi,A_{\rm S}}$  is the number of aggregate with aggregate size  $A_{\rm S}$  and ratio  $\xi$ ;  $N_{A_{\rm S}}$  is the total number of aggregates with aggregate size  $A_{\rm S}$ .

The definition of aggregate size is flexible. If none of -m, -x, --only options is used, aggregate size is the 'true' aggregation number,  $A_S$ , i.e., the number of all molecules in the aggregate; if -m is used, aggregate size is the sum of only specified molecule type(s); if -x is used, aggregates containing only specified molecule type(s) are disregarded; if --only is used, only aggregates composed of the specified molecule type(s) are taken into account. For example, consider a system containing three aggregates composed of various numbers of three different molecule types:

Molecule types   Aggregate composition	
Mol_A	Agg_1: 1 Mol_A +2 Mol_B +3 Mol_C = 6 molecules
$Mol_B$	Agg_2: $1 \text{ Mol\_A} + 2 \text{ Mol\_B} = 3 \text{ molecules}$
$Mol_C$	Agg_3: 1 Mol_A = 1 molecule

Here is a list of some of the possibilities depending on the option(s) used:

- (1) if none of -m, -x, --only is used, all three aggregates are counted and their sizes are their aggregation numbers, i.e.,  $A_S = 6$ , 3, and 1
- (2) if -m Mol\_A Mol\_B is used, all three aggregates are counted, but their size is the sum of only Mol\_A and Mol\_B molecules: Agg\_1 3; Agg\_2 3; Agg\_3 1
- (3) if -m Mol\_B Mol\_C is used, Agg\_3 is not counted, because its size would be

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zero; DistrAgg would detect only two aggregates with sizes: Agg\_1 - 5; Agg\_2 - 2

- (4) if -x Mol\_A Mol\_B is used, Agg\_2 and Agg\_3 are not counted, because neither contains anything else than Mol\_A and/or Mol\_B; DistrAgg would detect only one aggregate with size: Agg\_1 6
- (5) if -x Mol\_A Mol\_B is combined with -m Mol\_A Mol\_B, DistrAgg would again detect only Agg\_1, but its size would be taken as 3
- (6) if --only Mol\_A Mol\_B is used, Agg\_1 is not counted, because it contains a molecule not specified by --only; DistrAgg would detect only two aggregates with sizes: Agg\_2 3; Agg\_3 1
- (7) if --only Mol\_A Mol\_B is combined with -m Mol\_A, the two detected aggregates have sizes: Agg\_2 1; Agg\_3 1
- (8) if --only Mol\_A Mol\_B is combined with -x Mol\_A, only Agg\_2 is detected as it is the only composed of only Mol\_A and Mol\_B molecules Usage:

DistrAgg <input.agg> <distr file> <avg file> <options>

Mandatory arguments	
<pre><input.agg> <distr file=""> <avg file=""></avg></distr></input.agg></pre>	input agg file output file with distribution of aggregate sizes output file with per-timestep averages
Non-standard options	
-st <int> -n <int> <int> -m <mol name(s)=""> -x <mol name(s)=""></mol></mol></int></int></int>	starting timestep for calculation (default: 1) use aggregate sizes in a given range use number of specified molecule(s) as aggregate size exclude aggregates containing only specified mole- cule(s)
<pre>only <mol name(s)=""> -c <output> <int(s)></int(s)></output></mol></pre>	use only aggregates composed of specified molecule(s) save composition distribution for specified aggregate size(s) to <output> file</output>

#### Format of output files:

- (1) <output distr> distributions of aggregate sizes
  - first line: command used to generate the file
  - second line: column headers
    - first is the aggregate size, As either true aggregation number, or the size specified by options
    - F\_n(As), F\_w(As), and F\_z(As) are number, weight, and z distribution of aggregate sizes (Equation (4.8))

- <volume distribution> is distribution according to Equation (4.9)
- next is the total number of aggregates with specified size
- the remaining columns show average numbers of every molecule type in an aggregate with the specified size
- second to last line: column headers for overall averages written in the last line
  - <M>\_n, <M>\_w, and <M>\_z are number, weight, and z averages, respectively, of aggregate masses (the averages are defined in Equation (4.7))
  - a column denoted <mol name>\_n shows an average number of molecules named
     mol name in an aggregate
- (2) <output avg> per-timestep averages
  - first line: command used to generate the file
  - second lines: column headers
    - first is simulation timestep
    - the rest are for the calculated data: number, weight, and z average aggregate mass (<M>\_n, <M>\_w, and <M>\_z, respectively) and aggregate size (<As>\_n, <As>\_w, and <As>\_z, respectively)
  - the last two lines are the same as in <output distr>
- (3) -c <name> composition distribution
  - first line: command used to generate the file
  - second lines: column headers
    - first is ratio of the two molecule types (i.e., 0 to 1)
    - the rest are aggregate sizes
    - in the data, only ratios that are non-zero for at least one aggregate size are written; in case of more than one aggregate size specified by -a option, if the ratio does not exist for some aggregate size(s), '?' is displayed instead of zero

# 4.12 GenSystem

This simple utility uses modified FIELD file to create vsf structure file and to generate coordinates that could be used as a simulation's starting point. The utility assumes linear chains and uses equilibrium bond length to construct a prototype molecule that is fully stretched in one direction for each molecule type. The utility then creates layers of molecules that are separated by layers of unbonded beads (if there are any). The utility should fill the whole box with given beads.

The input FIELD file must contain species and molecule sections, but the interaction section is ignored (see DL\_MESO manual for details on the FIELD file). The first line of FIELD that is ignored by DL\_MESO must start with box dimensions, i.e., with three numbers (the rest of the file is ignored).

Usage (GenSystem does not use standard options):

#### GenSystem <out.vsf> <out.vcf> <options>

Mandatory arguments	
<pre><out.vsf> <out.vcf></out.vcf></out.vsf></pre>	output vsf structure file output vcf coordinate file
Options	
-f <name></name>	FIELD-like file (default: FIELD) verbose output that provides information about all bead and molecule types
-h	print help and exit

# 4.13 GyrationAggregates

This utility calculates the gyration tensor and its eigenvalues (or the roots of the tensor's characteristic polynomial) for all aggregates. Using the eigenvalues, the utility calculates shape descriptors: radius of gyration, asphericity, acylindricity, and relative shape anisotropy.

The eigenvalues,  $\lambda_x^2$ ,  $\lambda_y^2$ , and  $\lambda_z^2$ , (sorted so that  $\lambda_x^2 \le \lambda_y^2 \le \lambda_z^2$ ) are also written to output file(s), because their square roots represent half-axes of an equivalent ellipsoid.

The radius of gyration,  $R_{\rm G}$ , is defined as

$$R_{\rm G}^2 = \lambda_x^2 + \lambda_y^2 + \lambda_z^2. \tag{4.11}$$

The asphericity, b, and the acylindricity, c, are defined, respectively, as

$$b = \lambda_z^2 - \frac{1}{2}(\lambda_x^2 + \lambda_y^2) = \frac{3}{2}\lambda_z^2 - \frac{R_G^2}{2}$$
 and  $c = \lambda_y^2 - \lambda_x^2$ . (4.12)

The relative shape anisotropy is defined in terms of the other descriptors as

$$\kappa^2 = \frac{b^2 + 0.75c^2}{R_G^4} \tag{4.13}$$

Number average of all properties and, additionally, weight and z averages for the radius of gyration are calculated (see Equation (4.7) in Section 4.11 for general definitions of averages). Per-timestep averages (i.e., time evolution) are written to the <output> file. To save averages for aggregate sizes, -ps option can be used.

The shape descriptors are by default calculated for all beads in the aggregates, but -bt option can be used to specify which bead types to use.

Similarly to DistrAgg, the definition of aggregate size is flexible – see Section 4.11 for explanation of the -m and -x options.

Usage:

GyrationAggregates <input> <input.agg> <output> <options>

Mandatory arguments	
<input/>	input coordinate file (either vcf or vtf format)
<input.agg></input.agg>	input agg file
<output></output>	output file with per-timestep averages
Non-standard options	
joined	specify that <input/> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)
-bt <bead name(s)=""></bead>	bead type(s) used for calculation
-ps <name></name>	output file with per-size averages
-m <mol name(s)=""></mol>	instead of 'true' aggregate size, use the number of speci-
	fied molecule type(s) in an aggregate
-x < mol name(s) >	exclude aggregates containing only specified molecule
	type(s)
-n <int> <int></int></int>	use only aggregate sizes in given range
-st <int></int>	starting timestep for calculation (default: 1)

#### (1) <output> – per-timestep averages

- first line: command used to generate the file
- second line: column headers
  - first is timestep
  - rest are for calculated data: number, weight, and z averages (denoted by \_n, \_w, and \_z respectively) of radius of gyration and square of radius of gyration (Rg and Rg^2 respectively Equation (4.11)); number averages of relative shape anisotropy (Anis Equation (4.13)), acylindricity and asphericity (Acyl and Aspher, respectively Equation (4.12)), and all three eigenvalues (eigen.x, eigen.y, and eigen.z  $\lambda_x^2$ ,  $\lambda_y^2$ , and  $\lambda_z^2$ )
- second to last line: column headers for overall averages written in the last line
  - <M>\_n and <M>\_w are number and weight, respectively, averages of aggregate masses (the averages are defined in Equation (4.7)); aggregate mass here is the mass of all beads of the chosen type(s) in the aggregate
  - average numbers of molecules of each type in an aggregate are shown in columns denoted <mol names>
  - the remaining column represent overall averages of the per-timestep quantities

described above

- (2) -ps <name> per-size averages
  - first line: command used to generate the file
  - second line: column headers
    - first is aggregate size, As
    - last column is the total number of aggregates of the given size
    - the rest are for the calculated data: simple averages of the numbers of molecules of each type in the aggregate, of radius of gyration and its square, of relative shape anisotropy, of acylindricity, of asphericity, and of all three eigenvalues (all denoted by the above-described symbols)

# 4.14 GyrationMolecules

This utility calculates shape descriptors similarly to GyrationAggregates (Section 4.13), but for individual molecules instead for whole aggregates.

Usage:

GyrationMolecules <input> <output> <mol name(s)> <options>

Mandatory arguments	
<input/>	input coordinate file (either vcf or vtf format)
<output></output>	output file(s) (one per molecule type) with automatic
	- <mol_name>.txt ending</mol_name>
<mol name(s)=""></mol>	molecule name(s) to calculate shape descriptors for
Non-standard options	
joined	specify that <input/> contains joined coordinates (i.e., pe-
	riodic boundary conditions for molecules do not have to
	be removed)
-bt <bead name(s)=""></bead>	bead type(s) to be used for calculation
-st <int></int>	starting timestep for calculation (default: 1)

- (1) <output> per-timestep averages (one file per molecule type)
  - first line: command used to generate the file
  - second line: name of molecule type
  - third line: column headers
    - first is timestep
    - rest are for calculated data: number, weight, and z averages (denoted by \_n, \_w, and \_z respectively) of radius of gyration (Rg Equation (4.11)); number averages of relative shape anisotropy (Anis Equation (4.13)), acylindricity

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and asphericity (Acyl and Aspher, respectively – Equation (4.12))

# 4.15 JoinAggregates

This utility is meant for cases when non-standard option -j is omitted from Aggregates (or Aggregates-NotSameBeads) command. JoinAggregates uses the provided vcf and agg files to join aggregates, i.e., to remove periodic boundary conditions and save the new coordinates into a vcf file. The utility reads Aggregates command from the agg file to determine distance and number of contact pairs for aggregate check (see Section 4.3 for details on Aggregates utility).

The output file is a vcf coordinate file. Usage:

JoinAggregates <input> <input.agg> <output.vcf> <options>

Mandatory arguments	
<input/> <input.agg> <output.vcf></output.vcf></input.agg>	input coordinate file (either vcf or vtf format) input agg file output vcf coordinate file with indexed coordinates
Non-standard options	
-st <int></int>	starting timestep for calculation (default: 1)

### 4.16 JoinRuns

This utility probably does not work correctly.

This utility joins two independent simulation runs of the same system. That is, the system contains identical beads and molecules, but these beads and molecules are numbered differently in the vsf and vcf files from different simulations. The two input vcf files must contain the same timestep type (i.e., both indexed or both ordered) and the same number of beads (i.e., if one bead type is absent from one vcf file, it must be absent from the second one as well).

The output is a vcf coordinate file with beads indexed according to the first vsf structure file (i.e., traject.vsf or provided by -i option).

Usage:

JoinRuns <1st input> <2nd input> <2nd vsf> <output> <bead type(s)>
<options>

4. Utilities 4.17. lmp\_data

Mandatory arguments		
<1st input>	input coordinate file from the first simulation (either vcf or vtf format)	
<2nd input>	input coordinate file from the second simulation (either vcf or vtf format)	
<2nd vsf>	input structure file from the second simulation (structure file from the first simulation is traject.vsf; changeable via -i option)	
<output></output>	output vcf coordinate file with indexed coordinates	
<pre><bead type(s)=""></bead></pre>	bead type names to save	
Non-standard options		
join -st1 <int></int>	join molecules by removing periodic boundary conditions starting timestep first run (default: 1)	
202 2220		
-st2 <int></int>	starting timestep second run (default: 1)	
-sk1 <int></int>	number of steps skip per one used for first run (default: 0)	
-sk2 <int></int>	number of steps skip per one used for second run (default: 0)	

# 4.17 lmp\_data

This utility generate data file for the lammps simulation package (see lammps manual page for details on the data file format).

The utility reads information on system composition from DL\_MESO file (information on all beads and structure of molecules – although it does not read dihedrals) and coordinates from vcf coordinate file. The utility ignores the interactions part from FIELD. The utility also ignore dihedrals.

Usage:

lmp\_data <input> <out.data> <options>

Mandatory arguments	
<input/> <out.data></out.data>	input vcf coordinate file output data file
Options	
-f <name></name>	FIELD file (default: FIELD) coordinate timestep for creating the data file

## 4.18 PairCorrel

This utility calculates pair correlation function (pcf) between specified bead types. All bead type pairs are used – if A and B bead types, A-A, A-B, and B-B bead type pairs are used. Right now, the pcfs are not correctly normalised.

The utility do not recognise between beads of the same type that are in different molecules, so a pcf will be a sum of the beads from different molecule types.

Usage:

PairCorrel <input> <width> <output> <bead name(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<width> <output></output></width>	width of each bin of the pair correlation functions output file with pair correlation functions	
<pre><bead name(s)=""></bead></pre>	bead type(s) used for calculation	
Non-standard options		
-n <int></int>	number of bins to average to get smoother pair correlation function (default: 1)	
-st <int></int>	starting timestep for calculation (default: 1)	

#### Format of output files:

- (1) **<output>** pair correlation functions between all bead types
  - first line: command used to generate the file
  - second line: column headers
  - first is the centre of each bin (governed by <width>); i.e., if <width> is 0.1, then the centre of bin 0 to 0.1 is 0.05, centre of bin 0.1 to 0.2 is 0.15, etc.
  - the rest are for the calculated data: each column correspond to one pair of bead types

# 4.19 Potential Aggregates

This utility should be working, but it needs more testing.

This utility calculates electrostatic potential as a function of distance from the centre of mass of specified aggregate size(s). It places a virtual particle with charge q = 1 at several places on the surface of ever increasing sphere and calculates electrostatic potential acting on that virtual particle.

At long range, the potential is calculated using Coulomb potential,

$$U_{ij}^{\text{long}} = \frac{l_{\text{B}}q_iq_j}{r_{ij}},\tag{4.14}$$

where  $l_{\rm B}$  is the Bjerrum length,  $q_i$  and  $q_j$  are charges of particles i and j, and  $r_{ij}$  is interparticle distance. At short range, the potential is for now calculated using potential between two charges with exponentially decreasing charge density,

$$U_{ij}^{\text{short}} = U_{ij}^{\text{long}} \left[ 1 - (1 + \beta r_{ij}) e^{-2\beta r_{ij}} \right],$$
 (4.15)

where  $\beta = \frac{5r_c}{8\lambda}$  ( $r_c$  is cut off distance and  $\lambda$  is smearing constant). The utility takes into account periodic images of the simulation box.

For now, parameters for the potential are hard coded in the source code: the Bjerrum length is bjerrum=1.1 (aqueous solution), cut-off is r\_c=3, charge smearing constant lambda=0.2, and number of periodic images of the simulation box is images=5. The parameters can be changed but the utility must then be recompiled.

The aggregate size can be modified using -m options similarly to Density-Aggregates (Section 4.7).

Usage:

PotentialAggregates <input> <input.agg> <width> <output> <agg size(s)> <options>

Mandatory arguments		
<input/>	input coordinate file (either vcf or vtf format)	
<input.agg></input.agg>	input agg file	
<width></width>	width of each bin of the distribution	
<output></output>	output file(s) (one per aggregate size) with automatic	
	agg#.txt ending (# is aggregate size)	
<agg size(s)=""></agg>	aggregate size(s) for calculation of electrostatic potential	
Non-standard options		
joined	specify that <input/> contains joined coordinates (i.e., pe-	
	riodic boundary conditions for aggregates do not have to	
	be removed)	
-st <int></int>	starting timestep for calculation (default: 1)	
-m <mol name(s)=""></mol>	instead of 'true' aggregate size, use the number of specified molecule type(s) in an aggregate	

4. Utilities 4.20. SelectedVcf

## 4.20 SelectedVcf

This utility creates a new vcf coordinate file containing only beads of specified types with output vcf file. The selected bead types are printed as comments at the beginning of the output vcf file.

There is an option to remove periodic boundary conditions (i.e., to join molecules). Conversely, the simulation box can be wrapped (i.e., the periodic boundary conditions applied). If both --join and -w options are used, the simulation box is first wrapped and then the molecules are joined.

Also, specified molecules can be excluded which is useful when the same bead type is shared between more molecule types. However, as of now, no utilities can read a vcf file that does not contain all beads of a given type, so this can be used only for vmd visualization.

Lastly, xyz coordinate file can also be created from the selected bead type(s). Usage:

SelectedVcf <input> <output> <bead type(s)> <options>

Mandatory arguments		
<pre><input/> <output.vcf> <bead type(s)=""></bead></output.vcf></pre>	input coordinate file (either vcf or vtf format) output vcf coordinate file with indexed coordinates bead type names to save (can be omitted if -r is used)	
Non-standard options		
-r	reverse function, i.e., exclude <bead type(s)=""> instead of including them; if no <bead type(s)=""> are specified, all bead types are used (requires <input/> with all bead types)</bead></bead>	
join	join molecules by removing periodic boundary conditions	
<b>-</b> ₩	wrap simulation box (i.e., apply periodic boundary conditions)	
-st <int></int>	starting timestep for calculation (default: 1)	
-e <int></int>	ending timestep for calculation (default: none)	
-sk <int></int>	number of steps skip per one used (default: 0)	
-n <int(s)></int(s)>	save only specified timesteps	
-x <name(s)></name(s)>	exclude molecules of specified $name(s)$ – do not use if $output.vcf$ is further analysed	
-xyz <name></name>	save coordinates to $\mathbf{x}\mathbf{y}\mathbf{z}$ file – does not take into account $-\mathbf{x}$ option	

4. Utilities 4.22. TransformVsf

# 4.21 traject

This utility is from the DL\_MESO simulation package and comes in three version for 2.5, 2.6, and 2.7 versions of DL\_MESO. For the latest DL\_MESO version, the utility is unmodified and therefore is not included here. The utilities traject-v2\_5 and traject-v2\_6 shipped with DL\_MESO 2.5 and 2.6 were modified to generate separate vsf and vcf file (the traject for DL\_MESO 2.7 does this natively with -sc command line option).

Usage of 2.5 and 2.6 versions (see DL\_MESO manual for the description of latest version):

traject-v2\_5 <int> or traject-v2\_6 <int>

Mandatory arg	gument
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<int> number of computer cores used for the simulation run (equals the number of HISTORY files)

### 4.22 TransformVsf

This not-very-useful utility just rewrites vsf file for better visualization with vmd. The output vsf file contains not only bead name and index, but its charge and mass as well.

Usage:

TransformVsf <output.vsf> <options>

Mandatory a	argument
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<output.vsf> output structure file

# 5. Computational details

This chapter will contain some information about how things are coded in the utilities.

# 5.1 Read system data

ReadStructure() function reads all system information from vsf and vcf files. FIELD file is used only to get mass and charge of beads if the information is not in vsf file. Provided vsf file is used to get all information about beads and molecules – names and numbers of bead and molecules, bonds in molecules. The first timestep of the vcf file is used to determine numbers and ids of beads in that vcf file which means that all timesteps must contain the same beads.

The procedure in ReadStructure() is as follows:

- (1) Go through atom section of the vsf file to identify default bead type (if atom default line is present), to find highest bead and molecule ids, and to find bead type names (and charges and masses if present).
- (2) Go again through the atom section to read names and ids of beads and molecules as well as numbers of all beads and molecules for each type.
- (3) Go through bond section of the vsf file to calculate number of bonds in each molecule type.
- (4) Go again through the bond section to read bonds for each molecule type.
- (5) Go through the atom section (for the third time) to assign bead ids to molecules.
- (6) Go through the first timestep of vcf file to find which beads are in that vcf file (if no vcf file is provided e.g., for DistrAgg utility assume all beads are used).
- (7) Modify all arrays to accommodate only the beads that are present in the vcf file.
- (8) Read charge and mass from the FIELD file, if not already known from vsf file.