

AnalysisTools user manual

Karel Šindelka
k.sindelka@gmail.com

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1. Introduction

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 mainly on assembly of molecules, e.g., self-assembly of polymers. It therefore includes utilities to determine if molecules are in aggregates and to calculate various properties of aggregates. Also, throughout the utilities and this manual, the term aggregates refers to any supramolecular structure (such as, e.g., a bilayer).

However, the package also contains some utilities to generate initial configurations (or adapt existing configurations) as well as to create a data file from **vtf** trajectories for the **LAMMPS molecular dynamic simulator**.

2. Installation

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

To create the `Makefile`, simply run `cmake $ROOT_DIR` in the separate directory, where `$ROOT_DIR` is the `AnalysisTools` root directory. All utilities can be then compiled by simply running `make` without any arguments. To compile individual utilities, simply run `make <utility name>`. The binaries will be in `build/bin` directory.

Individual utilities can also be compiled without the use of `cmake` (and without `Makefile`) by using `gcc -O3 -lm $ROOT_DIR/AnalysisTools.c $ROOT_DIR/Error.c $ROOT_DIR/Options.c $ROOT_DIR/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 or for generation (or modification) of configurations. 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). Chapter 5 briefly describes how the data are read.

Both **vsf** and **vcf** files can be generated from the **DL_MESO HISTORY** file 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 of the

default type

- 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_A 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, 5, and 9 are of the default type (**Bead_A**); notice that if a bead of a default type is in a molecule, it must have its own **atom** line. 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 bond definitions. 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

The following is an example of bead definition lines that complement the above-shown bead definition lines:

```
bond 3: 4
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. Instead of a separate **vsf** file, a **vtf** file can be used (via **-i <name.vtf>** command line option).

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 **pbx** <float> <float> <float>. Because everything up to the first **timestep** or **pbx** line is ignored, **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
pbx 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 file 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; however, **AnalysisTools** utilities work with whole sets of beads, that is, when one bead of a specific type is missing, all beads of that type (i.e., beads with the same name) must be missing as well. Also, the beads do not have to be ordered according to their ascending indices. The following is an example of an ordered **vcf** file:

```
# any number of comments or blank lines

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
...
```

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** (or **Aggregates-NotSameBeads**) utility. The file contains information about the number of aggregates in each timestep and which molecules and monomeric (i.e., unbonded) beads belong to which aggregate. It serves as an additional input file for utilities that calculate aggregate properties; **agg** file is, therefore, linked to the **vcf** file 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. It follows these rules:

- each timestep starts with **Step:** <int> (only **Step** keyword is read by the utilities)
- the second line is the number of aggregates in the given timestep and is 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 (**resid** indices)
 - (2) number of monomeric beads in the aggregate followed by their indices taken from the **vsf** file (**atom** indices)
- no blank or comment lines are allowed inside the aggregate block
- not all molecules present in the **vcf** file used to generate this file must be present in every timestep

Note that the term aggregate also refers to free chains (i.e., fully dissolve dchains).

When the keyword **Last** is present instead of **Step**, it signals the end of the **agg** file; no utility will read anything beyond this keyword.

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
```

```
anything from here on is ignored
```

In this example, command **Aggregates in.vcf 1 1 out.agg A** was used to generate the file (see Section 4.2 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 an aggregate comprised of two molecules (i.e., a dimer). 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 by other utilities, the indices can be used in **vmd** to visualize, e.g., only a specific aggregate by using **resid 1 2 3** in the **Selected Atoms** box in **vmd**.

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

<code>-i <name></code>	use custom structure file instead of <code>traject.vsf</code> (<code>vsf</code> or <code>vtf</code> format)
<code>-v</code>	verbose output that provides information about all bead and molecule types
<code>-s</code>	run silently, i.e., without any output at all (overrides <code>-v</code> option)
<code>--script</code>	no progress output (useful if output is routed to a file)
<code>-h</code>	print this help and exit
<code>--version</code>	print version of the utilities 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 placing them either randomly or using provided coordinates.

The new beads replace either neutral unbonded ones with the lowest indices (as ordered in the `vsf` file) from the original system or option-specified unbonded beads. Right now, molecular beads cannot be exchanged. If molecules are added, **AddToSystem** places them at the end (for the sake of `DL_MESO` which requires molecules to be after unbonded beads). If there are not enough of the beads in the original system replace, **AddToSystem** exits with an error.

The utility has two modes: by default, it reads what is to be added from a `FIELD`-like file (see below) and places the new beads and molecules randomly; if, however, `-vtf` option is used, the utility reads provided `vsf` and `vcf` files and uses these coordinates.

If the new beads and molecules are added randomly, their coordinates are either completely random within the simulation box or ruled by constraining options; either the first bead (default behaviour) or the geometric centre (`-gc` option) of the

new molecules obey these options. The coordinates of the remaining beads in a molecule are governed by the coordinates in the **FIELD**-like file. Therefore, not all the molecular beads necessarily obey the constraining rules. Molecules are added with a random orientation.

There are two types of constraining options (that can be combined): place new beads either specified distance from other beads or in a specified interval in x, y, or z directions. The first case uses options **-ld** and **-hd** which, if present, must be accompanied by **-bt** option. Then, the new beads are placed at least **-ld <float>** and at most **-hd <float>** distance from beads specified by the **-bt** option. The second case uses options **-cx**, **-cy**, and **-cz** that basically change the box size for the added beads; for example, **-cx 5 9** would generate x coordinates only in the interval (5,9). All these options can be combined, but note that **AddToSystem** does not perform any sanity checks, that is, if any combination of the provided options is impossible to achieve, the utility will run forever.

As hinted above, 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 or unbonded beads).

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

The **molecule** keyword specifies the number of molecule types, that is the number of **finish** keywords that must be present. The **<bead name>** must be present in 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 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
A 1.0 1.0 0
B 1.0 0.0 0
CI 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.

If **-vtf** option is used, provided **vsf** and **vcf** files are read instead of a **FIELD**-like file. **AddToSystem** incorporates the beads and molecules from these files, using the provided coordinates instead of generating new ones. Therefore, above-described constraining options have no effect.

The **-vtf** option has priority over default behaviour, that is, if it is present, the utility ignores any **FIELD**-like file (either **FIELD** itself or a file provided via **-f** option).

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> <out.vsf> <out.vcf> <options>
```

Mandatory arguments

<input.vcf>	input coordinate file (either vcf or vtf format)
<out.vsf>	output vsf structure file for the new system
<out.vcf>	output vcf coordinate file for the new system

Non-standard options

-f <name>	FIELD -like file specifying additions to the system (default: FIELD)
-vtf <vsf> <vcf>	add ready-made system from vtf files instead of randomly placing beads read from the FIELD -like file
-st <int>	<input.vcf> timestep to add new beads to (default: 1)
-xyz <name>	also save coordinates to xyz file
-ld <float>	lowest distance from beads specified by -bt option
-hd <float>	highest distance from beads specified by -bt option
-bt <bead names>	bead types to use in conjunction with -ld and/or -hd options
-cx <num> <num2>	constrain x coordinate to interval $\langle num, num2 \rangle$
-cy <num> <num2>	constrain y coordinate to interval $\langle num, num2 \rangle$
-cz <num> <num2>	constrain z coordinate to interval $\langle num, num2 \rangle$
-gc	use molecule's geometric centre for distance check

4.2 Aggregates and Aggregates-NotSameBeads

These utilities determine which molecules belong to which aggregates (or a different compact structure) 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` text file described in Section 3.3.

Note that throughout **AnalysisTools**, an aggregate refers to any structure formed from molecules.

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 in 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

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><distance></code>	minimum distance for two beads to be in contact (thus constituting a contact pair)
<code><contacts></code>	minimum number of contact pairs between two molecules to be in one aggregate
<code><output.agg></code>	output <code>agg</code> file (must end with <code>.agg</code>) with aggregate information
<code><bead type(s)></code>	bead type name(s) to use for determining contact pairs (at least two for Aggregates-NotSameBeads)
<code><options></code>	

Non-standard options

<code>-x <mol name(s)></code>	exclude specified molecule type(s) from aggregate determination (and from the output <code>agg</code> file)
<code>-xm <mol name(s)></code>	exclude molecules that are close to specified molecule type(s)
<code>-j <output.vcf></code>	output <code>vcf</code> file with coordinates of joined aggregates (i.e., without periodic boundary conditions)

4.3 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.

Using `-n` option, the angle is specified by three bead indices taken from the bead order in the `vsf` file. These indices go 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 1 3 2` are specified, two angles will be calculated. The first angle will be between lines defined by beads with indices `1 2` and `2 3`; the second one will be between lines defined by beads with indices `1 3` and `3 2`. An 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 an `<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> <width> <output> <mol name(s)> <options>
```

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution
<code><mol name(s)></code>	molecule name(s) to calculate angles for
<code><output></code>	output file for distribution

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)
<code>-a <name></code>	write all angles for all molecules in all timesteps to <code><name></code>
<code>-n <ints></code>	multiple of three indices for angle calculation (default: 1 2 3)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)

Format of output files:

(1) `<output>` – distribution of angles

- first line: command used to generate the file
- second line: angle-specifying bead indices (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 n th 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 line: 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

4.4 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 value, statistical error, and estimate of autocorrelation time τ . Empty lines and comments (lines beginning with `#`) are skipped.

Average prints four numbers to screen: `<n_blocks>` `<average>` `<std error>` `<tau estimate>`:

<code><n_blocks></code>	number of blocks used for the binning analysis
<code><average></code>	simple arithmetic average
<code><std error></code>	one- σ statistical error
<code><tau estimate></code>	estimate of autocorrelation time τ

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

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

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, ϵ , is given by:

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

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

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

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

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

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

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

$$\epsilon^2 = \frac{\sigma_B^2}{N_B} = \frac{1}{N_B(N_B-1)} \sum_{n=1}^{N_B} (\mathcal{O}_{B,n} - \overline{\mathcal{O}})^2. \quad (4.5)$$

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

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

A way to quickly get τ estimate is to use a wide range of `<n_blocks>` values and plot the `<tau estimate>` values as a function of `<n_blocks>`. Because the number of data points in one block should be significantly larger than τ (e.g., ten times larger), plotting $f(x) = (\text{number of data lines in the file})/(10x)$ 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 considered a safe estimate of τ .

Usage:

Average `<input>` `<column>` `<discard>` `<n_blocks>`

Mandatory arguments

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

4.5 BondLength

This utility calculates a distribution of bond lengths in specified molecule type(s). It can also calculate a 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).

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 4.3). 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

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution
<code><output></code>	output file with distribution of bond lengths
<code><mol name(s)></code>	molecule name(s) to calculate bond lengths for

Non-standard options

<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-d <out> <ints></code>	write distribution of distances between specified beads in each specified molecule to <code><out></code>
<code>-w <double></code>	warn if a bond length exceeds <code><double></code> (default: half a box 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
 - 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 without any error, but it will produce an incomplete `CONFIG`. `Config` always prints first unbonded beads and then beads in molecules (required by DL_MESO).

Usage:

`Config <input.vcf> <options>`

Mandatory argument

`<input>` input coordinate file (either `vcf` or `vtf` format)

Non-standard option

`-st <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.xyz>` input xyz coordinate file

Non-standard option

`-st <int>` timestep for creating **CONFIG** file from (default: last step)

4.7 DensityAggregates

This utility calculates radial density profiles (RDPs, or radial number densities) for bead types in an aggregate with specified size (the number of molecules or aggregation number, A_S) from its centre of mass.

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 $r + dr$ (in **DensityAggregates**, dr is the `<width>` argument) divided by the volume of this shell. The utility also prints radial number profile (RNP $_i(r)$), or the number of beads in each shell without dividing it by its volume. In addition, it prints one- σ error for both RDP and RNP.

Composition of an aggregate with given size (i.e., average numbers of different types of molecules in that aggregate) is appended to the output file.

Instead of 'true' aggregate size, a number of molecules of specific type(s) can be used (`-m` option). For example, an aggregate containing 1 **Mol_A** molecule and 2 **Mol_B** molecules (i.e., three molecules in all) can be specified in several ways:

- (1) with `<agg size(s)>` 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> <input.agg> <width> <output> <agg
size(s)> <options>
```

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><input.agg></code>	input <code>.agg</code> file
<code><width></code>	width of a single distribution bin
<code><output></code>	output file(s) (one per aggregate size) with automatic <code>#.rho</code> ending (<code>#</code> is aggregate size)
<code><agg size(s)></code>	aggregate size(s) (the number of molecules in an aggregate or the aggregation number, A_S) to calculate density for

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)
<code>-n <int></code>	number of bins to average to get smoother density (default: 1)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-m <mol name(s)></code>	instead of 'true' aggregate size, use the number of specified molecule type(s) in an aggregate
<code>-x <mol name(s)></code>	exclude specified molecule type(s) (i.e., do not calculate density for beads in molecules <code><mol name(s)></code>)

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)

- last line contains the total number of aggregates the density was calculated for
- second to last line: column headers for average numbers of different molecules in the given aggregate
- last line: average numbers of the molecules

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, that is, the box is ‘sliced’ into blocks with width `<width>` and numbers of different bead types are counted in each of the slices.

The utility does not distinguish between beads with the same name in different molecules, so if one bead type is in more than one molecule type, it’s density will be averaged over all molecule types it appears in. In this case, however, the `-x` option can be used to first run the utility and exclude one molecule type and then rerun it, excluding the other molecule type. Thus, two output files are generated, each missing densities from the specified molecule types.

Usage:

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

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution
<code><output></code>	output file with automatic <code><axis>.rho</code> ending
<code><axis></code>	direction in which to calculate density: <code>x</code> , <code>y</code> , or <code>z</code>

Non-standard options

<code>-n <int></code>	number of bins to average to get smoother density (default: 1)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-x <mol name(s)></code>	exclude specified molecule type(s) (i.e., do not calculate density for beads in molecules <code><mol name(s)></code>)

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 according to `vsf` – similarly to `-n` option in `AngleMolecules`, Section 4.3) can be used instead (`-c` option).

Usage:

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

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution
<code><output></code>	output file with automatic <code><mol_name>.rho</code> ending (one file per molecule type is generated)
<code><mol name(s)></code>	molecule name(s) to calculate density for

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)
<code>-n <int></code>	number of bins to average for smoother density (default: 1)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-c <name> <int></code>	use specified bead in a molecule <code><name></code> instead of its centre of mass

Format of output files:

(1) `<output><mol_name>.rho` – 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` is 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 provided 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 4.3). 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 can be supplied to the `-n` option.).

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

Usage:

```
DihedralMolecules <input> <width> <output> <mol name(s)> <options>
```

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution (in degrees)
<code><output></code>	output file for distribution
<code><mol name(s)></code>	molecule name(s) to calculate angles for

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)
<code>-a <file></code>	write all angles for all molecules in all timesteps to <code><file></code>
<code>-n <ints></code>	multiple of six indices for angle calculation (default: 1 2 3 2 3 4)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)

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 n th 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 three lines: arithmetic means for each calculated angle (last line), headers (second to last line) that again give range of columns in the last line for each molecule type, and calculated angles (third to last line which is the same as the second line at the file beginning)

(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 (in degrees): 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 from a supplied `agg` file (see Section 3.3 for its format). 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

$$\langle \mathcal{O} \rangle_n = \frac{\sum_i N_i \mathcal{O}_i}{N}, \quad \langle \mathcal{O} \rangle_w = \frac{\sum_i N_i m_i \mathcal{O}_i}{\sum_i N_i m_i}, \quad \text{and} \quad \langle \mathcal{O} \rangle_z = \frac{\sum_i N_i m_i^2 \mathcal{O}_i}{\sum_i N_i m_i^2}, \quad (4.7)$$

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

$$\begin{aligned} F_n &= \frac{N_{A_S}}{\sum_{A_S} N_i} = \frac{N_{A_S}}{N}, \\ F_w &= \frac{N_{A_S} m_{A_S}}{\sum_{A_S} N_i m_i} = \frac{N_{A_S} m_{A_S}}{\sum_{i=1}^N m_i} = \frac{N_{A_S} m_{A_S}}{M}, \text{ and} \\ F_z &= \frac{N_{A_S} m_{A_S}^2}{\sum_{A_S} N_i m_i^2} = \frac{N_{A_S} m_{A_S}^2}{\sum_{i=1}^N m_i^2}, \end{aligned} \quad (4.8)$$

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

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

$$\phi(A_S) = \frac{N_{A_S} m_{A_S}}{\sum_{i=1}^N n_i} = \frac{N_{A_S} m_{A_S}}{n}, \quad (4.9)$$

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 equal to the number distribution, $F_n(A_S)$. All distributions are written into the `<output distr>` file.

Lastly, **DistrAgg** can calculate number distribution of composition for aggregates with specified size(s) (`-c` option); that is, the distribution of ratios of two molecule types in aggregates with specified size, $\xi = N_{\text{Mol}_A}/N_{\text{Mol}_B}$, where N_{Mol_A} and N_{Mol_B} are the numbers of **Mol_A** and **Mol_B** molecules in the aggregate with size $A_S = N_{\text{Mol}_A} + N_{\text{Mol}_B}$. The composition distribution is defined as

$$F_n(\xi) = \frac{N_{\xi, A_S}}{N_{A_S}}, \quad (4.10)$$

where N_{ξ, A_S} is the number of aggregates with aggregate size A_S and ratio ξ ; N_{A_S} is the total number of aggregates with aggregate size A_S . If there are more than two molecule types in the simulation, `-nc` option must be used to specify which molecules should be used in the composition calculation.

The `-nc` option has no effect if used without the `-c`.

The `<avg file>` contains averages for all timesteps regardless of `-st` and `-e` options; that is, the starting step (`-st` option) and ending step (`-e` option) are ignored.

The definition of aggregate size is flexible. If none of `-m`, `-x`, or `--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
Mo1_A	Agg_1: 1 Mo1_A + 2 Mo1_B + 3 Mo1_C = 6 molecules
Mo1_B	Agg_2: 1 Mo1_A + 2 Mo1_B = 3 molecules
Mo1_C	Agg_3: 1 Mo1_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 Mo1_A Mo1_B` is used, all three aggregates are counted, but their size is the sum of only Mo1_A and Mo1_B molecules: `Agg_1 - 3`; `Agg_2 - 3`; `Agg_3 - 1`
- (3) if `-m Mo1_B Mo1_C` is used, `Agg_3` is not counted, because its size would be zero; `DistrAgg` would detect only two aggregates with sizes: `Agg_1 - 5`; `Agg_2 - 2`
- (4) if `-x Mo1_A Mo1_B` is used, `Agg_2` and `Agg_3` are not counted, because neither contains anything else than Mo1_A and/or Mo1_B; `DistrAgg` would detect only one aggregate with size: `Agg_1 - 6`
- (5) if `-x Mo1_A Mo1_B` is combined with `-m Mo1_A Mo1_B`, `DistrAgg` would again detect only `Agg_1`, but its size would be 3
- (6) if `--only Mo1_A Mo1_B` is used, `Agg_1` is not counted, because it contains a molecule not specified by `--only`; `DistrAgg` would detect two aggregates with sizes: `Agg_2 - 3`; `Agg_3 - 1`
- (7) if `--only Mo1_A Mo1_B` is combined with `-m Mo1_A`, the two detected aggregates have sizes: `Agg_2 - 1`; `Agg_3 - 1`
- (8) if `--only Mo1_A Mo1_B` is combined with `-x Mo1_A`, only `Agg_2` is detected as it is the only one composed of only Mo1_A and Mo1_B molecules and its size would be 3
- (9) if `--only Mo1_A Mo1_B` and `-x Mo1_A` are combined with `-m Mo1_A`, the size of the aggregate would be 1

Moreover, only a specified range of aggregate sizes can be taken into account (`-n <int> <int>` option). These sizes are defined by the `-m`, `-x`, and `--only` options as well.

Usage:

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

Mandatory arguments

<code><input.agg></code>	input agg file
<code><distr file></code>	output file with distribution of aggregate sizes
<code><avg file></code>	output file with per-timestep averages

Non-standard options

<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-n <int> <int></code>	use aggregate sizes in a given range
<code>-m <mol name(s)></code>	use number of specified molecule(s) as aggregate size
<code>-x <mol name(s)></code>	exclude aggregates containing only specified molecule(s)
<code>--only <mol name(s)></code>	use only aggregates composed of specified molecule(s)
<code>-c <output> <int(s)></code>	save composition distribution for specified aggregate size(s) to <code><output></code> file
<code>-nc <name> <name></code>	molecule names to use for composition calculation

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 (**<mol name>_n**) 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))
 - other column headers are the same as at the file beginning

(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 ($\langle M \rangle_n$, $\langle M \rangle_w$, and $\langle M \rangle_z$, respectively) and aggregate size ($\langle As \rangle_n$, $\langle As \rangle_w$, and $\langle As \rangle_z$, respectively) and the last column is the number of aggregates in the given step
 - 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 GenLayers

This utility generates two monolayers composed of molecules specified in a **FIELD**-like file. For now, only one type of molecule is used (the first one in the **FIELD**-like file; other molecule types are ignored). The two layers are mirror images of each other, that is, molecules in both layers are grown from box edge to box middle. The layers are placed in z direction (that is, on xy planes of the simulation box).

The first beads of the molecules are arranged on a square lattice defined either by given spacing in x and y directions (`-s` option) or by the number of molecules per layer (`-nm` option). The rest of the beads of each molecule are placed based on the coordinates in the **FIELD**-like file. By default, **GenLayers** places the two mirror layers at the edges of a simulation box; using `-g` option, a gap from box edge can be introduced. Therefore, this utility can generate, for example, polymer brushes at box edges or a double layer (such as a biological membrane) in the middle of the box.

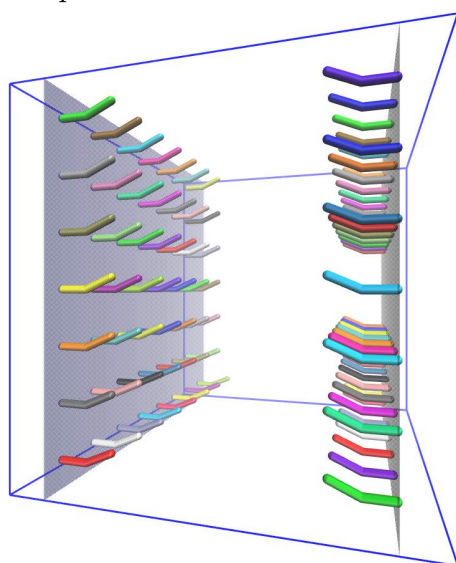
By default, the total number of beads in the generated system is equal to three times the box volume, that is, the typical number of beads in dissipative particle dynamics simulations. Beads that are not in the molecules, are put at the beginning of the output files (i.e., they have lower indices) with coordinates of the box centre, and name **None**. The idea is that once the layers are generated, **AddToSystem** utility (Section 4.1) can be used to exchange these excess beads for different species. The `-n` option changes the total number of beads. If the number is lower than the total number of beads needed to construct the two layers of molecules, it is adjusted to

exactly that number.

The input FIELD-like file must contain **species** and **molecule** sections (although, only the first molecule is considered and **nummols** is disregarded), but the **interaction** section is ignored (see DL MESO manual for details on the FIELD file or below for a simple example). The first line of the FIELD-like file, that is ignored by DL MESO, must start with box dimensions, i.e., with three numbers.

This utility does not have error checking for the provided FIELD-like file. If the file is not correct, **GenLayers** will exhibit undefined behaviour, that is, it will either freeze, crash, or run without errors, producing unexpected output files.

The following **vmd**-generated snapshot (the transparent squares are just a visual aid) is an example output using command **GenLayers out.vsf out.vcf -n 1 -nm 50 -g 1** and the provided FIELD-like file:



```
FIELD-like file
10 10 10
species 1
A 1.0 0.0 0
molecules 1
A3
nummols 1
beads 3
A 0 0.0 0.0
A 0 0.0 0.7
A 0 0.3 1.4
bonds 2
harm 1 2 100.0 0.7
harm 2 3 100.0 0.7
finish
```

The utility generates **vsf** structure and **vcf** coordinate files.

Usage (**GenLayers** does not use standard options):

GenLayers <out.vsf> <out.vcf> <options>

Mandatory arguments

<out.vsf>	output vsf structure file
<out.vcf>	output vcf coordinate file

Options

-s <x> <y>	spacing of molecules in <i>x</i> and <i>y</i> directions (default: 1 1)
-n <int>	total number of beads (default: three times the box volume)
-nm <int>	number of molecules in each layer (rewrites -s option)
-g <float>	gap between box edges and the molecules (default: 0)

<code>-f <name></code>	FIELD-like file (default: FIELD)
<code>-v</code>	verbose output that provides information about all bead and molecule types
<code>-h</code>	print this help and exit

4.13 GenSystem

This utility was not extensively tested and is not a good generator of initial configuration.

This simple utility uses modified FIELD file to create **vsf** structure file and to generate coordinates that could be used as an initial configuration. The utility assumes linear chains (no matter the connectivity in the provided FIELD file; it looks at exactly the first $n - 1$ bonds to get bond lengths, where n is the number of beads in a molecule). For each molecule type, it uses equilibrium bond length (or 0.7 if the bond length is 0) to construct a prototype molecule that is fully stretched in z -direction. 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.

This utility does not have error checking for the provided FIELD file. If the file is not correct, GenSystem will exhibit undefined behaviour, that is, it will either freeze, crash, or run without errors, producing bad output files.

Usage (GenSystem does not use standard options):

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

Mandatory arguments

<code><out.vsf></code>	output vsf structure file
<code><out.vcf></code>	output vcf coordinate file

Options

<code>-f <name></code>	FIELD-like file (default: FIELD)
<code>-v</code>	verbose output that provides information about all bead and molecule types
<code>-h</code>	print help and exit

4.14 GyrationAggregates

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

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

The radius of gyration, R_G , is defined as

$$R_G^2 = \lambda_x^2 + \lambda_y^2 + \lambda_z^2. \quad (4.11)$$

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

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

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

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

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. Per-size averages can be saved using the **-ps** option.

The gyration tensor is by default calculated for whole aggregates, but **-bt** option can be used to specify which bead types to consider.

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

The starting step (**-st** option) and ending step (**-e** option) are used only for averages (both overall averages and per-size averages with **-ps** option). Per-timestep averages in the <output> file disregard **-st** and **-e** options.

Usage:

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

Mandatory arguments

<input>	input coordinate file (either vcf or vtf format)
<input.agg>	input agg file
<output>	output file with per-timestep averages

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)
<code>-bt <bead name(s)></code>	bead type(s) used for calculation
<code>-ps <name></code>	output file with per-size averages
<code>-m <mol name(s)></code>	instead of ‘true’ aggregate size, use the number of specified molecule type(s) in an aggregate
<code>-x <mol name(s)></code>	exclude aggregates containing only specified molecule type(s)
<code>-n <int> <int></code>	use only aggregate sizes in given range
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)

(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 (R_g and R_g^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` – λ_x^2 , λ_y^2 , and λ_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
 - last column is the total number of aggregates of the given size
 - the columns in between 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, acylindricity, and asphericity, and of

all three eigenvalues (all are denoted by the above-described symbols)

4.15 GyrationMolecules

This utility calculates the gyration tensor and from it the shape descriptors similarly to `GyrationAggregates` (Section 4.14) but for individual molecules instead of for whole aggregates.

Usage:

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

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><output></code>	output file(s) (one per molecule type) with automatic <code>-<mol_name>.txt</code> ending
<code><mol name(s)></code>	molecule name(s) to calculate shape descriptors for

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for molecules do not have to be removed)
<code>-bt <bead name(s)></code>	bead type(s) to be used for calculation
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)

(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)); simple averages of relative shape anisotropy (`Anis` – Equation (4.13)), of acylindricity and asphericity (`Acyl` and `Aspher`, respectively – Equation (4.12)), and of the three eigenvalues (`eigen.x`, `eigen.y`, and `eigen.z` or λ_x^2 , λ_y^2 , and λ_z^2)
- last three lines show simple averages over the whole simulation
 - third to last line: molecule name
 - second to last line: column headers for radius of gyration and shape descriptors
 - last line: the averages

4.16 Info

This simple utility just prints information about the system read from the provided structure file. If `-c` option is used, only beads in the provided coordinate file are printed. The information is the same as when the verbose option (`-v`) is used with any other utility. Here, the `-v` option prints detailed informations about every bead and every molecule – this information also contains indices as implemented inside the **AnalysisTools** utilities (i.e., not necessarily consistent with the structure file).

Usage (**Info** does not use standard options):

Info <input> <options>

Mandatory argument	
<input>	input structure file (either <code>.vsf</code> or <code>.vtf</code> format)
Options	
<code>-c</code>	input coordinate file in either <code>.vcf</code> or <code>.vtf</code> format (default: None)
<code>-h</code>	print this help and exit

4.17 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 their 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 4.2 for details on **Aggregates** utility).

Be warned that if `-sk`, `-st`, or `-e` options are used, <output.vcf> will not be in sync with <input.agg> and, therefore, these two files cannot be used in tandem for further analysis via **AnalysisTools**.

Usage:

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

Mandatory arguments	
<input>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<input.agg>	input <code>agg</code> file
<output.vcf>	output <code>vcf</code> coordinate file with indexed coordinates

Non-standard options

<code>-sk <int></code>	number of steps skip per one used (default: 0)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)

4.18 JoinRuns

This utility joins two independent simulation runs of the same system. That is, the two systems contain 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., `trajectory.vsf` or provided by `-i` option). Only specified bead types are saved; using `-r` switch, the reverse is true, i.e., the specified bead types are excluded from the output file.

Usage:

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

Mandatory arguments

<code><1st input></code>	input coordinate file from the first simulation (either <code>vcf</code> or <code>vtf</code> format)
<code><2nd input></code>	input coordinate file from the second simulation (either <code>vcf</code> or <code>vtf</code> format)
<code><2nd vsf></code>	input structure file from the second simulation (structure file from the first simulation is <code>trajectory.vsf</code> ; changeable via <code>-i</code> option)
<code><output></code>	output <code>vcf</code> coordinate file with indexed coordinates
<code><bead type(s)></code>	bead type names to save (can be omitted if <code>-r</code> is used)

Non-standard options

<code>-r</code>	reverse function, i.e., exclude <code><bead type(s)></code> instead of including them; if no <code><bead type(s)></code> are specified, all bead types are used (requires input coordinate files with all types)
<code>--join</code>	join molecules by removing periodic boundary conditions
<code>-st1 <int></code>	starting timestep for first run (default: 1)

<code>-st2 <int></code>	starting timestep for second run (default: 1)
<code>-e1 <int></code>	ending timestep for first run (default: none)
<code>-e2 <int></code>	ending timestep for second run (default: none)
<code>-sk1 <int></code>	number of steps to skip per one used for first run (default: 0)
<code>-sk2 <int></code>	number of steps to skip per one used for second run (default: 0)

4.19 `lmp_data`

This simple utility generates `data` file for the [lammmps](#) simulation package (see [lammmps manual page](#) for details on the `data` file format).

This utility reads system composition from the provided `vsf` structure file, structure of molecules from `DL.MESO FIELD` file (numbers of bond and angle types and angles – other informations are read from the `vsf` file), and coordinates from `vcf` coordinate file. The utility ignores the `interactions` part from `FIELD`. For now, the utility also ignores dihedrals.

Option `--srp` can be used to add one more bead type into the output file (i.e., increment number of atom types and add one more line for `Masses` section with a mass of 1.0). This is intended for cases when the segmental repulsive potential (srp) is used in lammmps (srp prevents bond crossing when soft interparticle potential is used in a simulation).

Usage:

```
lmp_data <input> <out.data> <options>
```

Mandatory arguments

<code><input></code>	input <code>vcf</code> coordinate file
<code><out.data></code>	output <code>data</code> file

Options

<code>-f <name></code>	<code>FIELD</code> file (default: <code>FIELD</code>)
<code>--srp</code>	add one bead type for srp
<code>-st <int></code>	coordinate timestep for creating the <code>data</code> file

4.20 `lmp_vtf`

This utility generates `vsf` and `vcf` files from lammmps `data` file (see [lammmps manual page](#) for details on its format). Note that while lammmps uses the word ‘atom’, in

this manual, ‘bead’ is used instead.

`lmp_vtf` reads the `data` file header (`atoms`, `bonds`, `atom types`, `xlo xhi`, `ylo yhi`, and `zlo zhi` keywords) and `Masses`, `Atoms`, and `Bonds` sections; everything else is ignored. The utility requires lines in the `Atoms` section to have the following format: `<bead index> <molecule index> <bead type> <charge> <x> <y> <z>` (i.e., `atom_style full` in `lammps` terminology).

If any line of the `Masses` section ends with a comment, its first string is taken as the name of the bead type. Otherwise, the bead types are called `beadN`, where `N` is their type number in the `data` file.

Charge of every bead type is taken as the charge of the last bead of this type in the `Atoms` section.

Molecule types are determined according to the order of bead types in molecules as well as according to bead connectivity. Bead order in every molecule is considered from the lowest to the highest bead index in the `data` file. If `<molecule index>` in a bead line is 0, this bead is unbonded (i.e., not part of any molecule). If there’s only one bead in a molecule, this bead is considered unbonded as well. The types of molecules are called `molN`, where `N` goes from 1 to the total number of molecule types.

Following is an example of the `data` file:

```
7 atoms
2 bonds
3 atom types
0 10 xlo xhi
0 10 ylo yhi
0 10 zlo zhi
```

```
Masses
1 1.0
2 1.1 # some comment
3 1.2
```

```
Atoms
1 0 1 0.0 5.0 5.1 5.2
2 0 1 0.0 5.1 5.2 5.3
3 1 2 0.0 5.2 5.3 5.4
4 1 3 0.0 5.3 5.5 5.6
5 2 3 0.0 5.3 5.5 5.6
6 2 2 0.0 5.3 5.5 5.6
7 3 1 0.0 5.3 5.5 5.6
```

Bonds

```
1  1  3  4
2  1  5  6
```

In this example, there are 3 bead types called **bead1**, **some**, and **bead3**. There are 7 beads in all, 3 are unbonded (indices 1, 2, and 7; all called **bead3**), 4 are in two different molecules. The molecules are considered as different types even though they have the same numbers bonds and the same connectivity; while the first molecule has the order of beads **some**, **bead3** (according to the beads' ascending indices), the second one has the order reversed. These molecule types are called **mol1** and **mol2**.

Note that lines in both **Atoms** and **Bonds** sections do not have to be ordered in any way.

Usage (this utility does not use standard options):

```
lmp_vtf <input> <out.vsf> <out.vcf> <options>
```

Mandatory arguments

<input>	input lammps data file
<out.vsf>	output vsf structure file
<out.vcf>	output vcf coordinate file

Options

-v	verbose output
-h	print this help and exit

4.21 Orientation

This utility calculates orientation order parameter for specified molecules. It determines distribution of the values as well as average values.

Orientation order parameter s_{ij} indicates how stretched molecules in, e.g., membranes are. It is defined as

$$s_{ij} = \frac{1}{2} (3 \cos \theta - 1), \quad (4.14)$$

where

$$\cos \theta = \frac{\mathbf{r}_{ij} \cdot \mathbf{n}}{|\mathbf{r}_{ij}| |\mathbf{n}|}. \quad (4.15)$$

That is, the angle θ is between a normal vector to a plane, \mathbf{n} , and a vector connecting beads i and j , \mathbf{r}_{ij} . The value of s is between -0.5 (parallel to that plane) and 1 (perpendicular to that plane).

The default normal is z-axis, but using **-a** option, a different axis can be used. The **-a** option takes as an argument **x**, **y**, or **z**.

The two beads specifying the second vector can be defined via the `-a` option (similarly to the `-n` option in `AngleMolecules`, Section 4.3). The default is the first and second bead of the molecule, i.e., `-a 1 2`. A multiple of two indices can be provided to calculate different s_{ij} . All provided index pairs are used for all molecule types, unless both indices in a pair are higher than the number of beads in the molecule type. If only one index is higher, then this index is reduced to the highest index for the given molecule. For example, consider two molecule types to be used: `mol1` with four beads and `mol2` with two beads. Should `-a 1 2 1 3 3 4` be specified, `Orientation` would calculate $s_{1,2}$, $s_{1,3}$, and $s_{3,4}$ for `mol1` and $s_{1,2}$ and $s_{1,2}$ for `mol2` (it would literally calculate and print the same result twice). The pair `3 4` would be ignored for `mol2`.

The `<output>` file contains distributions of the orientation order parameter with averages of the parameter appended at the end.

Usage:

```
Orientation <input> <width> <output> <mol name(s)> <options>
```

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><width></code>	width of each bin of the distribution
<code><output></code>	output file with distribution of the orientation order parameter
<code><mol name(s)></code>	molecule name(s) used for calculation

Non-standard options

<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-a <axis></code>	axis to use as the normal vector (default: z)
<code>-n <ints></code>	bead indices in the molecules (multiple of two; default: 1 2)

Format of output files:

(1) `<output>` – distributions of orientation order parameters, s_{ij}

- first line: command used to generate the file
- next lines: column headers (on line per specified molecule type)
 - first column is the centre of each bin (governed by `width`); i.e., if `<width>` is 0.1, then the centre of the first bin is -0.45 (because $s_{ij} \in \langle 0.5, 1 \rangle$), centre of the second bin -0.35, etc.
 - next, each line shows the bead i and j indices in the specified molecule (according to the provided `vsf` file) to be used for calculation of s_{ij}
- next lines: calculated data
- last line: commented out (i.e., the line starts with `#`) overall averages
- commented out lines right above the last line: column headers for the averages

(same as at the beginning of the file)

4.22 PairCorrel

This utility calculates pair correlation function (pcf), or the radial distribution function, between specified bead types. All bead type pairs are used – if A and B are the specified bead types, A-A, A-B, and B-B bead pairs are used.

The utility does not differentiate beads of the same type that are in different molecules, so a pcf will be a sum of the beads from different molecule types (similar the **DensityAggregates** utility – see Section 4.7).

Usage:

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

Mandatory arguments

<input>	input coordinate file (either vcf or vtf format)
<width>	width of each bin of the pair correlation functions
<output>	output file with pair correlation functions
<bead name(s)>	bead type(s) used for calculation

Non-standard options

-n <int>	number of bins to average to get smoother pair correlation function (default: 1)
-st <int>	starting timestep for calculation (default: 1)
-e <int>	ending timestep for calculation (default: none)

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 corresponds to one bead types pair

4.23 PotentialAggregates

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

PotentialAggregates calculates electrostatic potential for aggregates of specified size(s) as a function of distance from their centre of mass. It places a virtual

particle with charge $q = 1$ at several places on the surface of an ever increasing sphere and calculates electrostatic potential acting on that virtual particle.

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

$$U_{ij}^{\text{long}} = \frac{l_B q_i q_j}{r_{ij}}, \quad (4.16)$$

where l_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,

$$U_{ij}^{\text{short}} = U_{ij}^{\text{long}} [1 - (1 + \beta r_{ij}) e^{-2\beta r_{ij}}], \quad (4.17)$$

where $\beta = \frac{5r_c}{8\lambda}$ (r_c is cut-off distance and λ 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 in the source code (around line 20), and the utility can then be recompiled.

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

Be aware that the utility does not use any Ewald sum-based algorithm, but simple ‘brute force’ approach, so the calculation is extremely slow (depending on the number of charged particles in the system).

Usage:

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

Mandatory arguments

<code><input></code>	input coordinate file (either <code>vcf</code> or <code>vtf</code> format)
<code><input.agg></code>	input <code>agg</code> file
<code><width></code>	width of each bin of the distribution
<code><output></code>	output file with electrostatic potential
<code><agg size(s)></code>	aggregate size(s) for calculation of electrostatic potential

Non-standard options

<code>--joined</code>	specify that <code><input></code> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)
<code>-st <int></code>	starting timestep for calculation (default: 1)

<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-m <mol name(s)></code>	instead of 'true' aggregate size, use the number of specified molecule type(s) in an aggregate

4.24 SelectedVcf

This utility creates a new **vcf** coordinate file containing only beads of specified types. The selected bead types are printed as comments at the beginning of the output **vcf** file. An **xyz** coordinate file can also be created from the selected bead type(s).

Using **-r** switch, the bead types can also be specified in reverse, i.e., bead types to be excluded from the output file(s).

Which timesteps are saved can be controlled using **-st** option for the first timestep to save and **-e** option for the last timestep to save; using **-sk <int>**, **<int>** steps are ignored per one step saved. If the option to save only specified timesteps (**-n**) is used, **-sk** is ignored. A maximum of 100 steps can be explicitly specified using the **-n** option.

There is also 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, no utilities can read a **vcf** file that does not contain all beads of a given type, so the **-x** option is useful only for visualization, e.g., using **vmd**.

Lastly, the **--last** saves to output file(s) only the last step from the input coordinate file; it is useful, when only the last timestep is required, but the total number of timesteps in the input file is unknown.

Usage:

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

Mandatory arguments

<input>	input coordinate file (either vcf or vtf format)
<output.vcf>	output vcf coordinate file with indexed coordinates
<bead type(s)>	bead type names to save (can be omitted if -r is used)

Non-standard options

<code>-r</code>	reverse function, i.e., exclude <code><bead type(s)></code> instead of including them; if no <code><bead type(s)></code> are specified, all bead types are used (requires <code><input></code> with all bead types)
<code>--join</code>	join molecules by removing periodic boundary conditions
<code>-w</code>	wrap simulation box (i.e., apply periodic boundary conditions)
<code>-st <int></code>	starting timestep for calculation (default: 1)
<code>-e <int></code>	ending timestep for calculation (default: none)
<code>-sk <int></code>	number of steps skip per one used (default: 0)
<code>-n <int(s)></code>	save only specified timesteps
<code>-x <name(s)></code>	exclude molecules of specified name(s) – do not use if <code><output.vcf></code> is further analysed
<code>-xyz <name></code>	save coordinates to xyz file – does not take into account <code>-x</code> option
<code>--last</code>	save only the last step

4.25 Surface

Surface utility determines the average surface of structures such as membranes or polymer brushes.

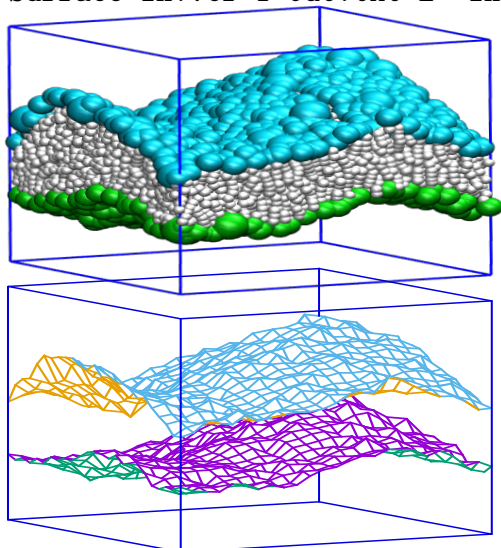
The utility cuts the plane perpendicular to the chosen axis into squares of size `<width>*<width>` and then finds the beads with the highest and lowest coordinates along the chosen axis whose other two coordinates fall into the square for each square on the plane. By default, it searches the chosen axis for the beads with the highest coordinate in the interval $\langle 0, (boxlength)/2 \rangle$ and with the lowest coordinate in the interval $\langle (boxlength)/2; (boxlength) \rangle$, i.e., it assumes something such as a polymer brush on each wall. If `-in` option is used, it searches for a layer structure inside the box, i.e., it searches for the bead with the lowest and highest coordinates in the whole box, $\langle 0; (boxlength) \rangle$ (this mode finds surfaces for something like a bilayer).

Note that the utility does not determines any structures for now, therefore any molecules outside the brush/membrane can be recognised as a part of the surface (`-bt` and `-m` can be sometimes used to eliminate this problem).

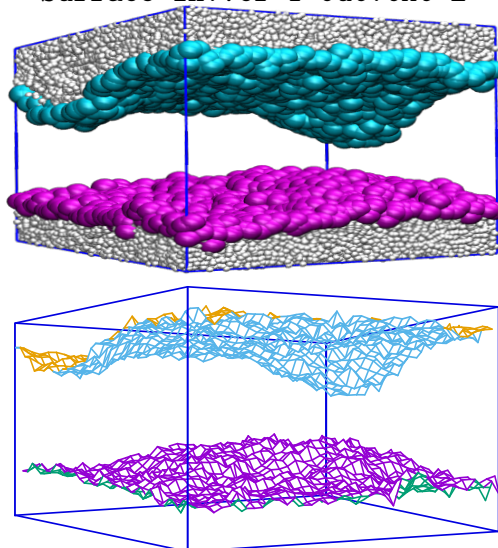
By default, all bead types and all molecule types are used, but using `-bt` and `-m` options, only specified bead and/or molecule types can be used. This is particularly useful when, e.g., other molecules solubilise inside the brush/membrane, leaving some molecules in the solution.

Following are two examples of usage with and without `-in` option (on top are snapshots with coloured balls representing the detected surface and on the bottom are graphs of the surface):

Surface in.vcf 1 out.txt z -in



Surface in.vcf 1 out.txt z



Usage:

Surface <input> <width> <output> <axis> <options>

Mandatory arguments

<input>	input coordinate file (either vcf or vtf format)
<width>	side length of each square
<output>	output file
<axis>	direction in which to determine the surface: x , y , or z

Non-standard options

-in	start from the box centre instead of from its edges
-m <name(s)>	molecule type(s) to use (default: all)
-bt <name(s)>	bead type(s) to use (default: all)
-st <int>	starting timestep for calculation (default: 1)
-e <int>	ending timestep for calculation (default: none)

Format of output files:

(1) <output> – 3D coordinates

- first line: command used to generate the file
- second line: column headers
 - first two numbers represent the centre of each square (being the coordinates on the sliced up plane); i.e., if <width> is 1, then the centre of the first square is 0.5 0.5, the centre of the second one is 0.5 1.5, etc.
 - the second two numbers are coordinates of the surface in the third dimension, i.e., along the chosen axis

4.26 **traject**

This utility is from the DL_MESO simulation package and comes in three versions 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.exe** 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 argument

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

4.27 **TransformVsf**

This utility reads **FIELD** and **vsf** files to create a new structure file. Generally, this is useful only if **traject-v2.5** or **traject-v2.6** were used to generate the **dl_meso.vsf** file. This file does not contain mass and charge of particles, whereas the one generated with **TransformVsf** (or with **traject.exe** from the **dl_meso** version 2.7) does and therefore the original **FIELD** file is not needed for further analysis.

Usage (**TransformVsf** does not use standard options):

TransformVsf <output.vsf> <options>

Mandatory argument

<output.vsf> output structure file

Options

-i <name> use custom structure file instead of **traject.vsf** (**vsf** or **vtf** format)

-v verbose output

-h print this help and exit

4.28 VisualizeAgg

This utility prints aggregates of given size to output **vcf** files (one file per aggregate size). The output files contain one aggregate per timestep regardless of how many aggregates of the given size are in the initial coordinate file.

The definition of aggregate size is fairly flexible (similarly to **DensityAggregates** utility – Section 4.7).

As the output file(s) do not necessarily contain all beads of a bead type, these **vcf** files cannot be used for further analysis via **AnalysisTools**.

Usage:

```
VisualizeAgg <input> <input.agg> <output> <agg
size(s)> <options>
```

Mandatory arguments

<input>	input coordinate file (either vcf or vtf format)
<input.agg>	input .agg file
<output>	output file(s) (one per aggregate size) with automatic #.vcf ending (# is aggregate size)
<agg size(s)>	aggregate size(s) to save

Non-standard options

--joined	specify that <input> contains joined coordinates (i.e., periodic boundary conditions for aggregates do not have to be removed)
-st <int>	starting timestep for calculation (default: 1)
-e <int>	ending timestep for calculation (default: none)
-m <mol name(s)>	instead of ‘true’ aggregate size, use the number of specified molecule type(s) in an aggregate
-x <mol name(s)>	exclude aggregates containing only specified molecule type(s)

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.