Analysis Software Package

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Contents

1	Form	nat of input files for all utilities	1
	1.1	Structure file	1
	1.2	Indexed coordinate file	2
	1.3	Aggregate file	2
2	Optio	ons for all utilities	3
3	Com	mon utilities	3
_	3.1	Aggregates & Aggregates-NotSameBeads utility	3
	3.2	Average utility	4
	3.3	BondLength utility — not extensively used or tested	5
	3.4	Config utility	5
	3.5	DensityAggregates	6
	3.6	DensityMolecules	7
	3.7	DihedralMoelcules	7
	3.8	DistrAgg utility	8
	3.9	GyrationAggregates utility	10
		GyrationMolecules utility	11
		JoinAggregates utility	12
		JoinRuns utility	12
		PairCorrel utility	13
		Potential Aggregates utility	14
		SelectedVcf utility	15
		traject utility	15
		TransformVsf utility	16
	3.17		
		3.17.1 Format of output structure file	16
4	Utilit	ies for linear chains	17
	4.1	EndToEnd utility	17
5	Todo	a Liet	17

6	Data	a Structure Index	17
	6.1	Data Structures	17
7	File	Index	18
	7.1	File List	18
8	Data	a Structure Documentation	18
	8.1	Aggregate Struct Reference	18
	8.2	Bead Struct Reference	19
	8.3	BeadType Struct Reference	19
	8.4	Counts Struct Reference	20
	8.5	IntVector Struct Reference	20
	8.6	LongVector Struct Reference	21
	8.7	Molecule Struct Reference	21
	8.8	MoleculeType Struct Reference	21
	8.9	Vector Struct Reference	22
9	File	Documentation	22
	9.1	AnalysisTools.h File Reference	22
		9.1.1 Function Documentation	24
	9.2	Options.h File Reference	32
		9.2.1 Function Documentation	33
	03	Structs h File Reference	30

1 Format of input files for all utilities

All the utilities read information about studied system from vsf/vcf files (formatted as described below) and FIELD file (input file for DL_MESO simulation package). Coordinates are read from a vcf file with either indexed timesteps. Particle mass and charge is read from the FIELD file, but everything else is read from the structure, vsf, and coordinate, vcf, files. Only bead types that are in the vcf file are considered.

Aggregate file is of my own format and is used by every utility doing calculation on whole aggregates (as opposed to calculations on individual molecules).

1.1 Structure file

The software package is designed with file dl_meso.vsf in mind, which is generated by the traject utility provided in DL_MESO software (and modified by me). Generally, the utilities are tested only against files generated by traject, but other vsf files (such as the one generated by TransformVsf utility, or even the structure part of a vcf file, i.e., -i <vcf file> option can be used) should work fine, if formatted according to the following guidelines.

First line may specify default bead (or atom, as called by vsf) type, meaning that all beads not specified by their own line are of the default type. No bead of the default type can be in a molecule. If the default line is not present, the number of atom lines must be the same as the total number of beads.

All atom lines in a vsf must specify bead index number as atom < int > (starting from 0) and name as name < char[8] > . If an atom is in a molecule, the name of the molecule type is specified as segid < char[8] > and its molecule number (starting from 1) is specified as resid < int > . The following is an example of atom lines:

```
atom default 1.0 name <char[8]>
...
atom <int> name <char[8]> segid <char[8]> resid <int>
...
```

While atom keyword (or its short version, a) must be at the beginning of the line, the order of other keywords do not matter; radius 1.0 The bead indices must go from the lowest to the highest and the highest index number corresponds to the total number of beads in the systems.

Bond lines follow after atom lines. Only simple bond lines are allowed, i.e. one bond per line in the form: bond <int>:<int>.

All molecules with the same name (and different index number) have to be the same, that is, consist of the same number of beads of the same types and have the same structure (i.e., the same bonds).

Blank lines and commented lines (beginning with #) are allowed in the vsf file.

1.2 Indexed coordinate file

First line of a <code>vcf</code> file that is read is a box size line, that is, it contains <code>pbc</code> <code><double></code> <code><double></code> <code><double></code> <code><double></code> ithe three numbers correspond to the side lengths of a cuboid simulation box). Everything up to the <code>pbc</code> keywoard is ignored. Therefore, a <code>vcf</code> file with structure part can be used (as long as no atom named <code>pbc</code> is present and the box size line is after the structure part).

Each timestep starts with a comment line (i.e., line starting with #), the second line contains timestep indexed (or the short version, t i), c(oordinate) i(ndexed), or just i(ndexed) and each following line contains index and coordinates of a single bead. Not all beads from the vsf file must be present in the vcf, instead only selected bead types can be present (although all beads of the selected type(s) must be in all timesteps). The file cannot contain a tailing blank line (no check is made in any utility). Example of an indexed coordinate file:

```
pbc <float> <float> <float>
<blank line - necessary>
# 1
indexed
<int> <float> <float> <float>
```

1.3 Aggregate file 3

1.3 Aggregate file

The aggregate file with .agg ending is generated using Aggregates utility. It contains information about the number of aggregates in the system in every simulation timestep and therefore is linked to the vcf file used to calculate the aggregates. For every aggregate in each timestep there is a number and ids of molecules in that aggregate as well as a number and ids of monomeric beads near the aggregate.

The first line of an aggregate file contains the command used to generate it. The subsequent lines contain information on individual timesteps starting with Step keyword, followed by the number of aggregates in the timestep and followed by individual aggregates. Every aggregate is spread over two lines - the first one contains the number of molecules in the aggregate followed by their ids (according to a corresponding vsf structure file) and the second line contains the number of monomeric beads in the aggregate followed by its ids (again, the ids correspond to the vsf file). The line with monomeric beads is indented for easier reading. The file ends with Last step: number of timesteps>; the L marks the end of agg file for any utility using it.

Example of an aggregate file:

```
<command used to generate it>
<blank line>
Step: 1
<number of aggregates in step 1>
<blank line>
2 : 1 34
3 : 230 40000 41003
<number of molecules in the second aggregate> : <molecule ids>
<number of monomeric beads near the aggregate : <bed ids>
<blank line>
Step: 2
...
<blank line>
Last Step: <number>
```

2 Options for all utilities

Most of the utilities have several options that are the same. These options are described here, but can be used with any utility unless stated otherwise. These options must be specified after any mandatory arguments.

```
use custom .vsf structure file instead of the default dl_meso.vsf (must end
with .vsf)

-v
   verbose output providing information about the system

-V
   more detailed verbose output

-s
   run silently, that is, without any output at all (overrides verbose options)

--script
   do not rewrite terminal line (useful if output is routed to file)

-h
   print help and exit
```

3 Common utilities

Utilies that are not specific to any given system and are used for all simulations.

3.1 Aggregates & Aggregates-NotSameBeads utility

These utilities determine which molecules belong to which aggregates according to a simple criterion: two molecules belong to the same aggregate if they share at least a specified number of contact pairs. A contact pair is a pair of two beads belonging to different molecules which are closer than certain distance. Both the distance and the number of needed contact pairs are arguments of the command as well as bead types to consider. Specified molecule(s) can be excluded from aggregate calculation (both from aggregate calculation and the output .agg file).

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 bead type pairs, that is A-A, A-B and B-B (provided the beads are not in the same molecules), but Aggregates-NotSameBeads will not use A-A or B-B. Therefore, at least two bead types must be provided in Aggregates-NotSameBeads argument.

```
Aggregates (or Aggregates-NotSameBeads) <input.vcf> <distance> <contacts>
<output.agg> <type names> <options>
     <input.vcf>
          input coordinate filename (must end with .vcf) containing indexed timesteps
     <distance>
          minimum distance for two beads to be in contact (constituting one contact pair)
     <contacts>
          minimum number of contact pairs to consider two molecules to be in one aggregate
     <output.agg>
          output filename (must end with .agg) containing information about aggregates
     <type names>
          names of bead types to use for calculating contact pairs
     <options>
          -x < name(s) >
              exclude specified molecule(s) from calculation of aggregates
          -j <joined.vcf>
              filename for coordinates of joined aggregates (must end with .vcf)
```

3.2 Average utility 5

3.2 Average utility

Average uses binning method to analyse data stored in a supplied file. It prints average, statistical error and estimate of integrated autocorrelation time (tau). Empty lines and lines beginning with '#' are skipped. The program prints to the screen four numbers: <n_blocks> <simple average> <statistical error> <estimate of tau>.

A way to estimate a 'real' value of tau is to use a wide range of $<n_blocks>$ and then plot <tau> as a function of $<n_blocks>$. Since the number of data points in a block has to be larger than tau (say, 10 times larger), plotting <number of data lines $>/10/<n_blocks>$ vs. $<n_blocks>$ will produce an exponential function that will intersect the plotted <tau>. A value of tau near the intersection (but to the left where the exponential is above <tau>) can be considered a good estimate for tau.

Usage:

```
Average <filename> <column> <discard> <n_blocks>

<filename>
    name of data file

<column>
    column number in the file containing the data to analyze

<discard>
    number of data values considered as equilibrium

<n_blocks>
    number of blocks for binning analysis
```

3.3 BondLength utility — not extensively used or tested

BondLength utility calculates normalised distribution of bond length for specified molecule types.

```
BondLength <input.vcf> <width> <output file> <molecule names> <options>

<input.vcf>
        input coordinate filename (must end with .vcf) containing indexed timesteps

<width>
        width of each bin for the distribution

<output file>
        output filename containing distribution of bond lengths

<molecule names>
        names of molecule types to calculate the distribution for
```

3.4 Config utility

This utility takes .vcf file containing all beads (such as All.vcf created by DL_MESO traject utility which was modified by me) and creates CONFIG file (file containing initial coordinates for a simulation via DL_MESO simulation package). If a vcf file that does not contain all beads, Config will still run, but the generated file will not contain coordinates for all beads and thus will not be able to be used to start a simulation rune using DL MESO software.

Usage:

3.5 DensityAggregates

This utility calculates number bead density for aggregates of specified size from their centre of mass. During the calculation, only the current aggregate is taken into account, so there is no possibility of getting 'false' densities from adjacent aggregates. Therefore if some bead type is never present in an aggregate of specified size (but is in the .vcf file), its density will always be 0.

Care must be taken with beadtype names in various molecules types, because if one beadtype appears in more molecule types, the resulting density for that beadtype will be averaged without regard for the various types of molecule it appears in.

Instead of true aggregate size, a number of molecules of specified name can be used, i.e. an aggregate with 1 $^{\rm A}$ molecule and 2 $^{\rm B}$ molecules can be specified with <agg sizes> of 3 without -m option or 1 if -m $^{\rm A}$ is used (or 2 if -m $^{\rm B}$ is used).

Also specified molecule type(s) can be excluded via the -x option. This is useful in case of several molecules sharing the same bead type. Calculated densities take into account only name of a bead type, not in which molecule(s) it occurs. The density from the bead type in different molecule types will therefore be the sum of the densities from those molecules.

Besides densities, the output files also contain radial number profiles (i.e., the numbers of beads in a thin spherical shell as a function of distance from aggregate centre of mass) and one-sigma errors for both the density and the radial number profiles.

Usage:

```
DensityAggregates <input.vcf> <input.agg> <width> <output.rho> <agg sizes>
<options>
```

input coordinate filename (must end with .vcf) containing indexed timesteps

<input.agg>

<input.vcf>

```
input filename (must end with .agg) containing information about aggregates
<width>
    width of each bin for the distribution
<output.rho>
    output density file (automatic ending agg#.rho added)
<agg sizes>
    aggregate sizes for density calculation
<options>
     --joined
         specify that the <input.vcf> contains aggregates with joined coordinates
     -n <int>
        number of bins to average
    -st <int>
         starting timestep for calculation
     -m <molecule type name>
        instead of aggregate size, use number of molecules of specified molecule types
     -x < name(s) >
         exclude specified molecule(s)
```

Todo DensityAggregates: check if only chains in one aggregate are used – anomalies in VanDerBurgh/AddedPol/

JoinRuns: implement wholy --script common option

Completely change this - either implement -x option or remove function WriteCoorIndexed and hard code the writing to file

3.6 DensityMolecules

DensityMolecules works in similar way as the DensityAggregates, only instead of aggregates, the densities are calculated for specified molecule types.

It is possible to use specified bead instead of the centre of mass for the coordinates to calculate densities from. Care must be taken, because the order of molecule types is taken from vsf rather than from DensityMolecules arguments. For example: whether bead 1 will be connected with NameA or NameB in DensityMolecules . . . NameA NameB -c 1 2 depends on molecules' order in vsf file; that is if NameA is first in vsf, 1 will be associated with NameA and 2 with NameB, but if NameB is first, the associations are reverse, regardless of the order of names in the command's arguments. If the centre of mass should be used, x is given as argument. In the above example (assuming NameA is first in vsf) if bead 1 is intended to be used for NameB, but centre of mass for NameA, then an argument of the form -c x 1 must be used.

```
<output.rho>
  output density file (automatic ending agg#.rho added)
<mol name(s)>
  names of molecule types to calculate density for
<options>
  --joined
     specify that the <input.vcf> contains aggregates with joined coordinates
  -n <int>
     number of bins to average
  -c x/<int>
```

use specified molecule bead instead of centre of mass

width of each bin for the distribution

3.7 DihedralMoelcules

DihedralMoelcules calculated angles between specified planes in a molecule for given molecule type(s). It prints the angles for all molecules for each timestep into the output file.

For each molecule type, a multiple of four bead ids must be specified (according to the order of molecule's bead ids in an input file). If $1\ 2\ 3\ 4$ is given, the angle between planes specified by $1\ 2\ 3$ and $2\ 3\ 4$. The utility does not check bonds in the molecule, so angle between any two planes in the molecules can be specified.

Starting timestep can also be specified.

```
Usage:
```

```
DihedralMolecules <input.vcf> <molecule(s)> <options>

<input.vcf>
    input coordinate filename (must end with .vcf) containing indexed timesteps

<mol name(s)>
    names of molecule types to calculate density for

<options>
    --joined
        specify that the <input.vcf> contains aggregates with joined coordinates
        -a <name> <ints>
            output filename and a series of four bead ids for angle calculation

-st <int>
            starting timestep for the calculation
```

3.8 DistrAgg utility 9

3.8 DistrAgg utility

DistrAgg calculates number and weight average aggregation masses for each timestep (i.e. their time evolution) as well as their average over the whole simulation. The number average and weight average aggregation mass, $\langle M \rangle_{\rm n}$ and $\langle M \rangle_{\rm w}$ respectively, are defined as:

$$\langle M \rangle_{\rm n} = \frac{\sum_{A_{\rm S}} m_i N_i}{N}$$
, and $\langle M \rangle_{\rm w} = \frac{\sum_{A_{\rm S}} m_i^2 N_i}{\sum_{A_{\rm S}} m_i N_i}$, (1)

where the sums go over all aggregate sizes. m_i is the mass of an aggregate with aggregation number $A_{\rm S}=i$, N_i is the number of such aggregates and N is total number of aggregates. The equations can be written more conveniently (for programming purposes) as sums over aggregates themselves instead of their sizes:

$$\langle M \rangle_{\rm n} = \frac{\sum_{i=1}^N m_i}{N} , \text{ and } \langle M \rangle_{\rm w} = \frac{\sum_{i=1}^N m_i^2}{\sum_{i=1}^N m_i}. \tag{2}$$

The utility also calculates number, weight and z distribution function of aggregation numbers. The distributions, $F_{\rm n}(A_{\rm S})$, $F_{\rm w}(A_{\rm S})$, and $F_{\rm z}(A_{\rm S})$ respectively, are defined as:

$$F_{\rm n}(A_{\rm S}) = \frac{N_{A_{\rm S}}}{\sum_{A_{\rm S}} N_i} = \frac{N_{A_{\rm S}}}{N} \,, \quad F_{\rm w}(A_{\rm S}) = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{\sum_{A_{\rm S}} N_i m_i} \,, \quad \text{ and } \quad F_{\rm z}(A_{\rm S}) = \frac{N_{A_{\rm S}} m_{A_{\rm S}}^2}{\sum_{A_{\rm S}} N_i m_i^2}, \tag{3}$$

where $N_{A_{\rm S}}$ and $m_{A_{\rm S}}$ stand for the number and mass, respectively, of aggregates with aggregation number $A_{\rm S}$. The equations are normalized so that $\sum F_x(A_{\rm S})=1$. Equations for $F_{\rm w}$ and $F_{\rm z}$ can again be transformed to contain sums over aggregates, not their sizes:

$$F_{\rm w}(A_{\rm S}) = \frac{N_{A_{\rm S}} m_{A_{\rm S}}}{\sum_{i=1}^{N} m_i} \quad \text{and} \quad F_{\rm z}(A_{\rm S}) = \frac{N_{A_{\rm S}} m_{A_{\rm S}}^2}{\sum_{i=1}^{N} m_i^2}, \tag{4}$$

Lastly, the utility calculates distribution of volume fractions of aggregates under the assumption that all beads have the same volume, which is essential in dissipative particle dynamics simulation method. Since in DPD the volumes of all beads are identical, volume fraction of an aggregate with aggregation number $A_{\rm s}$ is calculated as:

$$\phi(A_{\rm S}) = \frac{N_{A_{\rm S}} n_{A_{\rm S}}}{\sum_{i=1}^{N} n_i},\tag{5}$$

where n_i is the number of beads in an aggregate with $A_s = i$.

The definition of aggregation number , $A_{\rm S}$, is somewhat flexible. The -m <name> option can be used to specify that the aggregation number is not the number of all molecules in an aggregate, but rather only the number molecules of the specified type(s). Two values are then taken as the mass and aggregation number for calculations of distributions and averages – the 'true' value (noted as whole agg mass in output files) and the '-m option value' (noted as options mass). For example, let's assume an aggregate contains A and B molecules. Using -m A will count the aggregate size as the number of As, but its mass as both the mass of only A molecules and as the sum of A and B molecules' masses. The resulting distributions will therefore be the functions of aggregation number specified by the -m option, but for every distribution function, there will be two data columns in the output file (and two sets of overall averages).

Also using the --only <name> uses only aggregates composed exclusively of a specified molecule type. On the contrary, -x <name(s) > option discounts aggregates containing only the specified molecules.

Next, the calculations can be made only for a given range of aggregation numbers (specified by the -m option if present) if -n option is used.

Lastly, it can calculate composition distribution, that is, if an aggregate of specified size can be composed of two molecule types A and B (or two types specified by -m option), the aggregate can contain various A/B molecule ratios (or ratios $\xi = N_{\rm A}/N_{\rm B}$). The option -c <int(s)> <name> calculates number distribution of these

ratios for every given size (or <int (s) >) and the distribution is written to file <name>. This number distribution is defined as

 $F_{\rm n}(\xi) = \frac{N_{\xi, A_{\rm S}}}{N_{A_{\rm S}}},$ (6)

where $N_{\xi,A_{\rm S}}$ is the number of aggregates with size $A_{\rm S}$ (possibly specified by the -m option) and ratio ξ and $N_{A_{\rm S}}$ is the total number of aggregates with the given $A_{\rm S}$.

The utility reads information about aggregate from input file with Aggregate format. This file can be generated using Aggregates utility.

Usage:

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

<input>

input filename with information about aggregates

<distr file>

output filename with weight and number distribution functions

<avg file>

output filename with weight and number average aggregation number in each timestep

<options>

-st <int>

starting timestep for calculation (does not affect calculation of time evolution)

-n < int > < int >

range of aggregation numbers to calculate distributions and averages for

-m < name(s) >

instead of aggregate size, use number of molecules of specified molecule types

-x < name(s) >

exclude aggregates containing only specified molecule(s)

--only <molecule type name>

use only aggregates composed of specified molecule type

-c < name > < int(s) >

calculate composition distribution for specified aggregate size(s) and write it to file $<\!\texttt{name}\!>$

3.9 GyrationAggregates utility

This utility calculates a gyration tensor and its eigenvalues (as roots of the tensor's characteristic polynomial) for all aggregates. Using the eigenvalues, various shape descriptors are determined.

It calculates radius of gyration, $R_{\rm G}$, as:

$$R_{\rm G}^2 = \lambda_x^2 + \lambda_y^2 + \lambda_z^2,\tag{7}$$

where λ_i^2 is the i-th eigenvalue of the gyration tensor (or its i-th principle moment). The eigenvalues are sorted so that $\lambda_x^2 \leq \lambda_y^2 \leq \lambda_z^2$. Then it calculates the asphericity, b:

$$b = \lambda_z^2 - \frac{1}{2} \left(\lambda_x^2 + \lambda_y^2 \right) = \frac{3}{2} \lambda_z^2 - \frac{R_G^2}{2},\tag{8}$$

the acylindricity, c:

$$c = \lambda_y^2 - \lambda_x^2 \tag{9}$$

and the relative shape anisotropy, κ :

$$\kappa^2 = \frac{b^2 + 0.75c^2}{R_G^4} = \frac{3}{2} \frac{\lambda_x^4 + \lambda_y^4 + \lambda_z^4}{\left(\lambda_x^2 + \lambda_y^2 + \lambda_z^2\right)^2} \tag{10}$$

Number averages of all the properties and weight and z averages for radius of gyration are calculated. The number average for a quantity \mathcal{O} is defined as:

$$\langle \mathcal{O} \rangle_{\rm n} = \frac{\sum_i \mathcal{O}_i}{N},$$
 (11)

where N is the total number of aggregates. The weight average is:

$$\langle \mathcal{O} \rangle_{\mathbf{w}} = \frac{\sum_{i} m_{i} \mathcal{O}_{i}}{\sum_{i} m_{i}},$$
 (12)

where m_i is mass of an aggregate i. Lastly, the z-average is

$$\langle \mathcal{O} \rangle_{\mathbf{z}} = \frac{\sum_{i} m_{i}^{2} \mathcal{O}_{i}}{\sum_{i} m_{i}^{2}}.$$
 (13)

All the sums go over individual aggregates. Both number and weight average according to these equations were already used in the DistrAgg utility.

Averages of the shape descriptors during the simulation (or their time evolution) are written to an output file and overall averages are appended to that file.

Sums dependent on aggregation number can be written to output file with -ps <name> option. The aggregation number can be either the number of all molecules in an aggregate or a number of molecules of the type specified by -m <name> option.

Usage:

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

<input.vcf>

input coordinate filename (must end with .vcf) containing indexed timesteps

<input.agg>

input filename (must end with .agg) containing information about aggregates

<output.vcf>

output filename with shape descriptors for chosen sizes throughout simulation

<agg sizes>

aggregate sizes for gyration calculation

<options>

--joined

specify that the <input.vcf> contains aggregates with joined coordinates

-bt

specify bead types to be used for calculation (default is all)

-m < name >

take as an aggregate size the number of <name> molecules in aggregates instead of the number of all molecules

-ps <name>

output filename with per-size (or per-aggregation-number) averages

-n <int> <int>

range of aggregation numbers to calculate distributions and averages

3.10 GyrationMolecules utility

This utility function in the same way as GyrationAggregates, but it calculates radii of gyration for specified molecule types instead of aggregate sizes.

Right now it calculates gyration for all beads in the specified molecule types.

```
Usage:
```

```
GyrationMolecules <input.vcf> <input.agg> <output> <molecule names> <options>

<input.vcf>
        input coordinate filename (must end with .vcf) containing indexed timesteps

<input.agg>
        input filename (must end with .agg) containing information about aggregates

<output.vcf>
        output.vcf>
        output filename with radii of gyration throughout simulation (automatic ending #.txt)

<molecule names>
        molecule types for gyration calculation

<options>
        -bt
            specify bead types to be used for calculation (default is all)

--joined
            specify that the <input.vcf> contains joined coordinates
```

3.11 JoinAggregates utility

This utility reads input .vcf and .agg files and removes periodic boundary conditions from aggregates - e.i. it joins the aggregates. The distance and the bead types for closeness check are read from the first line of .agg file with contains full Aggregates command used to generate the file. JoinAggregates is meant for cases, where -j flag was omitted in Aggregates utility.

```
Aggregates <input.vcf> <input.agg> <output.vcf> <options> <input.vcf> input coordinate filename (must end with .vcf) containing indexed timesteps <input.agg> input filename (must end with .agg) containing information about aggregates <output.vcf> output filename (must end with .vcf) with joined coordinates
```

3.12 JoinRuns utility 13

3.12 JoinRuns utility

MOST LIKELY NOT WORKING - IT'S NOT USED.

This program is to be used if two simulation runs with different initial seeds (that is, two simulations with different bead id numbers .vsf files, but identical FIELD files) should be joined. Two .vcf files that contain the same bead types must be provided as well as the .vsf structure file for the second simulation. The output .vcf coordinate files has bead ids according to the structure file of the first simulation. The program is, however, extremely inefficient with unbonded beads, while bonded beads are always sorted in the same way by DL_MESO simulation software. The usefullness of such utility is confined to cases with more then one type of unbonded beads and under those conditions the utility may take around 1 minute per step (of the second simulation run) for system in box of side length 40.

Usage:

```
JoinRuns <1st input.vcf> <2nd input.vcf> <2nd input.vsf> <output.vcf>
<type names> <options>
     <1st input.vcf>
          input coordinate filename (must end with .vcf) containing indexed timesteps for the first
          simulation
     <2nd input.vcf>
          input coordinate filename for the second sumation in the same format as the first coordinate
          file
     <2nd input.vsf>
           .vsf structure file for the second simulation (must end with .vsf)
     <output.vcf>
          output filename with indexed coordinates (must end with .vcf)
     <type names>
          names of bead types to save
     <options>
          --joined
              join individual molecules by removing periodic boundary conditions
          -n1 <int>
              number of timestep to start the first simulation from
          -n2 < int >
              number of timestep to start the second simulation from
          -u1 <int>
              leave out every skip steps in the first simulation
          -u2 < int >
```

Todo JoinRuns: base reindexing of beads in the second simulation on comparison between the two .vsf files

leave out every skip steps in the second simulation

3.13 PairCorrel utility

This utility calculates pair correlation function (pcf) between specified bead types. All pairs of bead types (including same pair) are calculated - given $\mathbb A$ and $\mathbb B$ types, pcf between $\mathbb A-\mathbb A$, $\mathbb A-\mathbb B$ and $\mathbb B-\mathbb B$ are calculated. The pcfs are not normalised.

Currently, the utility cannot recognise between beads of the same type that are in different molecule types - i.e. if bead type A is both in molecule type 1 and molecule type 2, only one pcf will be calculated regardless of the molecule type A is in. Use SelectedVcf with -x option to resolve the problem.

Usage:

3.14 Potential Aggregates utility

This utility calculates electrostatic potential as a function of distance from the centre of mass of specified aggregate size(s). It calculates the electrostatic potential at every charged bead in the aggregate (both molecular beads and monomer beads). It then sums up all calculated potentials in the bins of specified widths at specified distance from the aggregate centre of mass.

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

$$U_{ij}^{\text{long}} = \frac{l_{\text{B}} q_i q_j}{r_{ij}},\tag{14}$$

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

$$U_{ij}^{\text{short}} = U_{ij}^{\text{long}} \left[1 - (1 + \beta r_{ij}) \exp(-2\beta r_{ij}) \right],$$
 (15)

where $\beta=\frac{5r_c}{8\lambda}$ (r_c is cut-off distance and λ is smearing constant). It also calculates contributions from periodic images of the simulation box. Parameters of the potential are (for now) hard-coded in the sourcode: Bjerrum length bjerrum=1.1 (aqueous conditions), cut-off distance r_c=3, charge smearing constant lambda=0.2, and number of periodic images of the simulation box images=5.

The size of aggregate can be modified using -m and -x options similarly to DensityAggregates.

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

```
<input.vcf>
    input coordinate filename (must end with .vcf) containing indexed timesteps
<input.agg>
    input filename (must end with .agg) containing information about aggregates
<width>
    width of each bin for the distribution
<output.txt>
    output file with electrostatic potential (automatic ending agg#.txt added)
<agg size(s)>
    aggregate sizes for calculation of electrostatic potential
<options>
     --joined
         specify that the <input.vcf> contains aggregates with joined coordinates
     -st <int>
         starting timestep for calculation
     -m <molecule type name>
         instead of aggregate size, use number of molecules of specified molecule types
    -x < name(s) >
         exclude specified molecule(s)
```

3.15 SelectedVcf utility

This utility takes .vcf file containing either indexed timesteps and creates a new .vcf coordinate file containing only beads of selected types with an option of removing periodic boundary condition and thus joining molecules. The otput .vcf file contains indexed timesteps.

Specified molecules can be excluded which is useful when the same bead type is shared between more molecule types.

The selected <type names > are printed at the beginning of the output file.

```
--join
   join individual molecules by removing periodic boundary conditions
-st <int>
        starting timestep for calculation
-sk <int>
        number of steps to skip per one used
-x <name(s) >
        exclude specified molecule(s)
```

3.16 traject utility

This utility is from the DL_MESO simulation package. While originally it creates a .vtf file containing both structure and coordinates, I have changed it to create a separate dl_meso.vsf structure file and All.vcf coordinate file containing indexed timesteps.

There are two versions from two versions of the DL_MESO simulation package, namely versions 2.5 and 2.6.

number of computer cores used for the simulation run (or the number of HISTORY file)

Usage:

```
traject-v2_5 <cores> or traject-v2_6 <cores> <cores>
```

The standard options cannot be used with this utility.> }}}

3.17 TransformVsf utility

This utility takes .vsf structure file and DL_MESO input file FIELD and transforms them into a different .vsf structure file that is better suited for visualisation using VMD software.

4 Utilities for linear chains

3.17.1 Format of output structure file

Every atom line in the generated structure file contains bead's index number, mass, charge and name. Atom lines for beads in molecules also contain molecule's id number and the name of the type of molecule. The bond section of output.vsf lists all bonds one by one (i.e. no chains of bonds in the format <id1>:: <id2> are used). Information about which bonds belong to which molecule is provided as a comment. The file has the following format:

```
atom default name <name> mass <m> charge <q>
...
atom <id> name <name> mass <m> charge <q>
...
atom <id> name <name> mass <m> charge <q> segid <name> resid
<id>
...
# resid <id>
<bonded bead id1>: <bonded bead id2>
...
```

For VMD atom selection:

```
segid <name>
    selects all molecules with given name(s)

resid <id>
    selects molecule(s) with given index number(s)

charge <q>
    selects all beads with given charge(s) (double quotes are required for negative charge)

mass <m>
    selects all beads with given mass(es)
```

4 Utilities for linear chains

This section provides information about utilities with calculations that are sensible to do only on linear polymer chains. No check whether the molecules are linear is done.

4.1 EndToEnd utility

This utility calculates end-to-end distance of specified molecules. End-to-end distance makes sense only for linear chains, therefore it is assumed that the provided molecule names are linear chains. No check is performed. The distance is calculated between the first and the last bead of the molecule; that is, between the first and the last bead in the FIELD entry for the given molecule. Also the use of joined coordinates (that is, without periodic boundary condition) is required, because the utility does not remove periodic boundary conditions.

The output is a file containing average end-to-end distance for every molecule type for each timestep.

Usage:

```
EndToEnd <input.vcf> <output.vcf> <molecule names> <options>

<input.vcf>
    input coordinate filename (must end with .vcf) containing either ordered or indexed timesteps (with joined coordinates)

<output.vcf>
    output.vcf>
    output filename with indexed coordinates (must end with .vcf)

<molecule names>
    names of molecule types (linear chains) to use
```

5 Todo List

Page Common utilities

DensityAggregates: check if only chains in one aggregate are used - anomalies in VanDerBurgh/AddedPol/

JoinRuns: implement wholy --script common option

Completely change this - either implement -x option or remove function WriteCoorIndexed and hard code the writing to file

JoinRuns: base reindexing of beads in the second simulation on comparison between the two .vsf files

6 Data Structure Index

6.1 Data Structures

Here are the data structures with brief descriptions:

Aggregate Information about every aggregate Bead Information about every bead 19 BeadType Information about bead types 19

7 File Index

Counts Total numbers of various things	20
IntVector 3D vector of integers	20
LongVector 3D vector of floats	21
Molecule Information about every molecule	21
MoleculeType Information about molecule types	21
Vector 3D vector of floats	22
7 File Index	
7.1 File List	
Here is a list of all documented files with brief descriptions:	
AnalysisTools.h Functions common to all analysis utilities	22
Options.h Options usable in utilities	32
Structs.h Structures for utilities	39
Common/Aggregates.h	??
8 Data Structure Documentation	
8.1 Aggregate Struct Reference	
Information about every aggregate.	
#include <structs.h></structs.h>	
Data Fields	
• int nMolecules	
number of molecules in aggregate • int * Molecule	
ids of molecules in aggregate	
• int nBeads	
number of bonded beads in aggregate	

· int * Bead

ids of bonded beads in aggregate

· int nMonomers

number of monomeric beads in aggregate

int * Monomer

ids of monomeric beads in aggregate

· double Mass

total mass of the aggregate

bool Use

should aggregate be used for calculation?

8.2 Bead Struct Reference

Information about every bead.

```
#include <Structs.h>
```

Data Fields

· int Type

type of bead corresponding to index in BeadType struct

· int Molecule

index number of molecule corresponding to Molecule struct (-1 for monomeric bead)

· int nAggregates

number of aggregates the bead is in (only monomeric beads can be in more aggregates - allocated memory for 10)

int * Aggregate

index numbers of aggregates corresponding to Aggregate struct (-1 for bead in no aggregate)

int Index

index of the bead according to .vsf file (needed for indexed timesteps)

Vector Position

cartesian coordinates of the bead

8.3 BeadType Struct Reference

Information about bead types.

```
#include <Structs.h>
```

Data Fields

• char Name [16]

name of given bead type

int Number

number of beads of given type

bool Use

should bead type in .vcf file be used for calculation?

· bool Write

should bead type in .vcf file be written to output .vcf?

· double Charge

charge of every bead of given type

· double Mass

mass of every bead of given type

8.4 Counts Struct Reference

Total numbers of various things.

```
#include <Structs.h>
```

Data Fields

· int TypesOfBeads

number of bead types

int TypesOfMolecules

number of molecule types

· int Beads

total number of beads in all molecules

· int Bonded

total number of beads in all molecules (TO BE REMOVED)

· int Unbonded

total number of monomeric beads (TO BE REMOVED)

· int BeadsInVsf

total number of all beads in .vsf file (not necessarily in .vcf)

· int Molecules

total number of molecules

• int Aggregates

total number of aggregates

8.5 IntVector Struct Reference

3D vector of integers.

```
#include <Structs.h>
```

Data Fields

- int x
- int y
- int z

8.6 LongVector Struct Reference

3D vector of floats.

```
#include <Structs.h>
```

Data Fields

- long double **x**
- · long double y
- long double z

8.7 Molecule Struct Reference

Information about every molecule.

```
#include <Structs.h>
```

Data Fields

· int Type

type of molecule corresponding to index in Molecule Type struct

int * Bead

ids of beads in the molecule

· int Aggregate

id of aggregate molecule is in (corresponding to index in Aggregate struct)

8.8 MoleculeType Struct Reference

Information about molecule types.

```
#include <Structs.h>
```

Data Fields

• char Name [16]

name of given molecule type

· int Number

number of molecules of given type

· int nBeads

number of beads in every molecule of given type

• int nBonds

number of bonds in every molecule of given type

int ** Bond

pair of ids for every bond (with relative bead numbers from 0 to nBeads)

• int nBTypes

number of bead types in every molecule of given type

int * BType

ids of bead types in every molecule of given type (corresponds to indices in BeadType struct)

· double Mass

total mass of every molecule of given type

bool InVcf

is molecule type in vcf file?

· bool Use

should molecule type be used for calculation?

· bool Write

should molecule type be used for calculation?

8.9 Vector Struct Reference

3D vector of floats.

```
#include <Structs.h>
```

Data Fields

- double x
- double y
- double z

9 File Documentation

9.1 AnalysisTools.h File Reference

Functions common to all analysis utilities.

```
#include "Structs.h"
```

Functions

• void CommonHelp (bool error)

Function printing help for common options.

 void VerboseOutput (bool Verbose2, char *input_vcf, char *bonds_file, Counts Counts, BeadType *BeadType, Bead *Bead, MoleculeType *MoleculeType, Molecule *Molecule)

Function printing basic information about system if $\neg v$ or $\neg V$ option is provided.

• bool ReadStructure (char *vsf_file, char *vcf_file, char *bonds_file, Counts *Counts, BeadType **BeadType, Bead **Bead, MoleculeType **MoleculeType, Molecule **Molecule)

Function reading information from dl_meso FIELD and vsf structure files.

int ReadCoorOrdered (FILE *vcf_file, Counts Counts, Bead **Bead, char **stuff)

Function reading ordered coordinates from .vcf coordinate file.

• int ReadCoorIndexed (FILE *vcf_file, Counts Counts, Bead **Bead, char **stuff)

Function reading ordered coordinates from .vcf coordinate file.

bool SkipCoor (FILE *vcf_file, Counts Counts, char **stuff)

Function to skip one timestep in coordinates file.

 bool ReadAggregates (FILE *agg_file, Counts *Counts, Aggregate **Aggregate, BeadType *BeadType, Bead **Bead, MoleculeType *MoleculeType, Molecule **Molecule)

Function reading information about aggregates from .agg file.

 void WriteCoorIndexed (FILE *vcf_file, Counts Counts, BeadType *BeadType, Bead *Bead, MoleculeType *MoleculeType, Molecule *Molecule, char *stuff)

Function writing indexed coordinates to a .vcf file.

int FindBeadType (char *name, Counts Counts, BeadType *BeadType)

Function to identify type of bead from its name.

• int FindMoleculeType (char *name, Counts Counts, MoleculeType *MoleculeType)

Function to identify type of molecule from its name.

• Vector Distance (Vector id1, Vector id2, Vector BoxLength)

Function to calculate distance vector between two beads.

 void RemovePBCMolecules (Counts Counts, Vector BoxLength, BeadType *BeadType, Bead **Bead, MoleculeType *MoleculeType, Molecule *Molecule)

Function to join all molecules.

• void RemovePBCAggregates (double distance, Aggregate *Aggregate, Counts Counts, Vector BoxLength, BeadType *BeadType, Bead **Bead, MoleculeType *MoleculeType, Molecule *Molecule)

Funcion to join all aggregates.

void RestorePBC (Counts Counts, Vector BoxLength, Bead **Bead)

Function to restore pbc.

Vector CentreOfMass (int n, int *list, Bead *Bead, BeadType *BeadType)

Function to calculate centre of mass for a collection of beads.

- Vector Gyration (int n, int *list, Counts Counts, Vector BoxLength, BeadType *BeadType, Bead **Bead)
- double Min3 (double x, double y, double z)

Function returning the lowest number from three floats.

Vector Sort3 (Vector in)

Function returning sorted numbers x < y < z.

• void FreeBead (Counts Counts, Bead **Bead)

Free memory allocated for Bead struct array.

• void FreeMolecule (Counts Counts, Molecule **Molecule)

Free memory allocated for Molecule struct array.

void FreeMoleculeType (Counts Counts, MoleculeType **MoleculeType)

Free memory allocated for Molecule Type struct array.

void FreeAggregate (Counts Counts, Aggregate **Aggregate)

Free memory allocated for Molecule Type struct array.

9.1.1 Function Documentation

9.1.1.1 CentreOfMass()

```
Vector CentreOfMass (
                int n,
                int * list,
                Bead * Bead,
                BeadType * BeadType )
```

Parameters

in	n	number of beads	
in	list	list of bead ids (corresponding to indices in Bead struct)	
in	Bead	information about individual beads (coordinates)	
in	BeadType	information about beadtypes (masses)	

Returns

coordinates of centre of mass of a given aggregate

Function to calculate centre of mass for a given list of beads.

9.1.1.2 CommonHelp()

```
void CommonHelp (
          bool error )
```

Parameters

in	error	true or false whether to use stderr or stdout	
----	-------	---	--

Function to print help for common options, either for -h help option or program error.

9.1.1.3 Distance()

Parameters

in	id1	first coordinate vector
in	id2	second coordinate vector
in	BoxLength	dimensions of simulation box

Returns

distance vector between the two provided beads (without pbc)

Function calculating distance vector between two beads. It removes periodic boundary conditions and returns x, y, and z distances in the range <0, BoxLength/2).

9.1.1.4 FindBeadType()

Parameters

in	name	bead name
in	Counts	numbers of beads, residues, etc.
in	BeadType	information about bead types

Returns

bead type id corresponding to index in BeadType struct (or -1 if non-existent bead name)

9.1.1.5 FindMoleculeType()

Parameters

	in	name	molecule name
	in	Counts	numbers of beads, residues, etc.
ſ	in	MoleculeType	information about bead types

Returns

molecule type id corresponding to index in BeadType struct (or -1 for non-existent molecule)

9.1.1.6 FreeAggregate()

Parameters

in	Counts	number of beads, molecu.es, etc.
out	Aggregate	information about individual molecules

Free memory allocated for Aggregate struct array. This function makes it easier other arrays to the Aggregate struct in the future

9.1.1.7 FreeBead()

Parameters

in	Counts	number of beads, molecu.es, etc.
out	Bead	information about individual beads

Free memory allocated for Bead struct array. This function makes it easier to add other arrays to the Bead struct in the future

9.1.1.8 FreeMolecule()

Parameters

in	Counts	number of beads, molecu.es, etc.
out	Molecule	information about individual molecules

Free memory allocated for Molecule struct array. This function makes it easier other arrays to the Molecule struct in the future

9.1.1.9 FreeMoleculeType()

Parameters

in	Counts	number of beads, molecu.es, etc.
out	MoleculeType	information about individual molecules

Free memory allocated for MoleculeType struct array. This function makes it easier other arrays to the MoleculeType struct in the future

9.1.1.10 Gyration()

```
Vector Gyration (
                int n,
                 int * list,
                Counts Counts,
                Vector BoxLength,
                BeadType * BeadType,
                Bead ** Bead )
```

Function to calculate the principle moments of the gyration tensor.

9.1.1.11 Min3()

Parameters

in	X	first double precision number
in	У	second double precision number
in	Z	third double precision number

Returns

lowest of the supplied numbers

Function returning the lowest number from three floats.

9.1.1.12 ReadAggregates()

```
bool ReadAggregates (
    FILE * agg_file,
    Counts * Counts,
    Aggregate ** Aggregate,
    BeadType * BeadType,
    Bead ** Bead,
    MoleculeType * MoleculeType,
    Molecule ** Molecule
```

Parameters

in	agg_file	name of input aggregate file
in	Counts	numbers of beads, molecules, etc.
in	BeadType	information about bead types
out	Bead	information about individual beads
out	Aggregate	information about aggregates
in	MoleculeType	information about molecule types
out	Molecule	information about individual molecules

Returns

1 if 'Last Step' detected or 0 for no error

Function reading information about aggregates from .agg file (Aggregate file) generated by Aggregates utility.

9.1.1.13 ReadCoorIndexed()

```
int ReadCoorIndexed (
    FILE * vcf_file,
    Counts Counts,
    Bead ** Bead,
    char ** stuff )
```

Parameters

in	vcf_file	name of input .vcf coordinate file
in	Counts	numbers of beads, molecules, etc.
out	Bead	coordinates of individual beads
out	stuff	first line of a timestep

Returns

0 for no errors or index number of bead (starting from 1) for which coordinates cannot be read

Function reading coordinates from .vcf file with indexed timesteps (Indexed coordinate file).

9.1.1.14 ReadCoorOrdered()

```
Counts Counts,
Bead ** Bead,
char ** stuff )
```

Parameters

in	vcf_file	name of input .vcf coordinate file
in	Counts	numbers of beads, molecules, etc.
out	Bead	coordinates of individual beads
out	stuff	first line of a timestep

Returns

0 for no errors or index number of bead (starting from 1) for which coordinates cannot be read

Function reading coordinates from .vcf file with ordered timesteps. Not used anymore!

9.1.1.15 ReadStructure()

Parameters

in	vsf_file	.vsf structure file
in	vcf_file	.vcf coordinate file
in	bonds_file	filename with bonds
out	Counts	numbers of beads, molecules, etc.
out	BeadType	information about bead types
out	Bead	informationn about individual beads
out	MoleculeType	information about molecule types
out	Molecule	information about individual molecules

Returns

'true' or 'false' for .vcf file with indexed or ordered timesteps, respectively

Function reading information about beads and molecules from DL_MESO FIELD file, .vsf structure file, and .vcf coordinate file. Charge and mass of beads is read from FIELD file, but all other information is read from vsf structure file. If vcf coordinate is passed to ReadStructure (), bead types not present in the vcf file get ignored.

Overly complicated, some things done more than once, needs complete overhaul. Thankfully, it works.

9.1.1.16 RemovePBCAggregates()

Parameters

in	distance	distance for closeness check (taken from agg file)
in	Aggregate	information about aggregates
in	Counts	number of beads, molecu.es, etc.
in	BoxLength	dimensions of the simulation box
in	BeadType	information about bead types
out	Bead	information about individual beads (coordinates)
in	MoleculeType	information about molecule types
in	Molecule	information about individual molecules

Function to remove periodic boundary conditions from all aggregates, thus joining them.

9.1.1.17 RemovePBCMolecules()

Parameters

in	Counts	numbers of beads, molecules, etc.
in	BoxLength	dimension of the simulation box
in	BeadType	information about bead types
out	Bead	information about individual beads (coordinates)
in	MoleculeType	information about molecule types
in	Molecule	information about individual molecules

Function to remove periodic boundary conditions from all individual molecules, thus joining them

9.1.1.18 RestorePBC()

Parameters

in	Counts	numbers of beads, molecules, etc.
in	BoxLength	dimension of the simulation box
out	Bead	information about individual beads (coordinates)

Function to restore removed periodic boundary conditions. Used in case of cell linked list, because it needs coordinates <0, BoxLength>.

9.1.1.19 SkipCoor()

Parameters

in	vcf_file	file with vcf coordinates
in	Counts	number of beads in vcf file
out	stuff	first line of a timestep

Returns

1 if premature end of file or 0 for no error

Function to skip one timestep in coordinates file. It works with both indexed and ordered vcf files.

9.1.1.20 Sort3()

```
Vector Sort3 ( \begin{tabular}{ll} Vector in \end{tabular}
```

Parameters

in in first double precision numb	 ⊵r
-----------------------------------	--------

Returns

sorted vector

Function returning sorted numbers x < y < z.

9.1.1.21 VerboseOutput()

```
BeadType * BeadType,
Bead * Bead,
MoleculeType * MoleculeType,
Molecule * Molecule )
```

Parameters

in	Verbose2	print extra information if 'true'
in	input_vcf	.vcf structure file
in	bonds_file	filename with bonds
in	Counts	numbers of beads, molecules, etc.
in	BeadType	information about bead types
in	Bead	informationn about individual beads
in	MoleculeType	information about molecule types
in	Molecule	information about individual molecules

Function providing standard verbose output (for cases when verbose option is used). It prints most of the information about used system.

9.1.1.22 WriteCoorIndexed()

```
void WriteCoorIndexed (
    FILE * vcf_file,
    Counts Counts,
    BeadType * BeadType,
    Bead * Bead,
    MoleculeType * MoleculeType,
    Molecule * Molecule,
    char * stuff )
```

Parameters

in	vcf_file	name of output .vcf coordinate file
in	Counts	numbers of beads, molecules, etc.
in	BeadType	information about bead types
in	Bead	coordinates of individual beads
in	MoleculeType	information about molecule types
in	Molecule	coordinates of individual molecules
in	stuff	array of chars containing comment line to place at the beginning

Function writing coordinates to a .vcf file. According to the Use flag in BeadType structure only certain bead types will be saved into the indexed timestep in .vcf file (Indexed coordinate file).

9.2 Options.h File Reference

Options usable in utilities.

```
#include "Structs.h"
```

Functions

bool VsfFileOption (int argc, char **argv, char **vsf_file)

Option whether to use .vsf file different from $dl_{meso.vsf}$ (-i < name.vsf>).

bool BondsFileOption (int argc, char **argv, char **bonds_file)

Option whether to use bonds file (-b < name>).

void VerboseLongOption (int argc, char **argv, bool *verbose, bool *verbose2)

Option whether to use long verbose output (overrides VerboseShortOutput) (-V).

• void SilentOption (int argc, char **argv, bool *verbose, bool *verbose2, bool *silent)

Option whether not to print to stdout (overrides Verbose options) (-s).

bool ExcludeOption (int argc, char **argv, Counts Counts, MoleculeType **MoleculeType)

Option whether to exclude molecule types (-x < name(s) >).

bool JoinCoorOption (int argc, char **argv, char *joined_vcf)

Option whether to write joined aggregate coordinates to file (-j < joined.vcf>).

bool BeadTypeOption (int argc, char **argv, Counts Counts, BeadType **BeadType)

Option to choose which bead types to use in calculations (-bt < name(s) >).

bool BoolOption (int argc, char **argv, char *opt)

Option whether not to print rewrite stdout line (--script).

bool IntegerOption (int argc, char **argv, char *opt, int *value)

Function for any option with integer argument.

bool TwoIntegerOption (int argc, char **argv, char *opt, int *values)

Function for any option with two integer arguments.

• bool HundredIntegerOption (int argc, char **argv, char *opt, int *values, int *count, char *file)

Function for any option with two integer arguments.

bool FileOption (int argc, char **argv, char *opt, char **name)

Function for any option with filename.

bool MoleculeTypeOption (int argc, char **argv, char *opt, int *moltype, Counts Counts, MoleculeType
 **MoleculeType)

Function for any option with molecule type name.

bool MoleculeTypeOption2 (int argc, char **argv, char *opt, int **moltype, Counts Counts, MoleculeType
 **MoleculeType)

Function for any option with molecule type name(s).

9.2.1 Function Documentation

9.2.1.1 BeadTypeOption()

```
bool BeadTypeOption (
          int argc,
          char ** argv,
          Counts Counts,
          BeadType ** BeadType )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	Counts	numbers of beads, molecules, etc.
out	BeadType	information about bead types

Returns

```
true or false error or not error
```

Option to choose which bead types to use for calculation. If the option is absent, all bead types are switched to Use = true. Argument: -bt < name(s) >

9.2.1.2 BondsFileOption()

```
bool BondsFileOption (
                int argc,
                char ** argv,
                 char ** bonds_file )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
out	bonds_file	filename with bonds

Returns

true or false error or not error

Option whether to use bonds file with alternative bond definitions.

9.2.1.3 BoolOption()

```
bool BoolOption (
                int argc,
                char ** argv,
                char * opt )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)

Returns

```
true if opt present, false otherwise
```

Function for any boolean option (i.e. without argument). The option (e.g. --script) is an argument of this function.

9.2.1.4 ExcludeOption()

```
bool ExcludeOption (
                int argc,
                char ** argv,
                Counts Counts,
                 MoleculeType ** MoleculeType )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	Counts	numbers of beads, molecules, etc.
out	MoleculeType	information about molecule types

Returns

true or false error or not error

Option to exclude specified molecule types from calculations. Gives specified molecule types use = false and the rest use = true. Arguments: -x < name(s) >

9.2.1.5 FileOption()

```
bool FileOption (
                int argc,
                char ** argv,
                char * opt,
                char ** name )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)
out	name	array containing the filename

Returns

true or false for error

Generic option for file name. The option is an argument of this function.

9.2.1.6 HundredIntegerOption()

```
bool HundredIntegerOption (
    int argc,
    char ** argv,
    char * opt,
    int * values,
    int * count,
    char * file )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -c)
out	values	array of two integer values of given option
out	n	number of numeric arguments

Returns

```
true or false for error
```

Function for any option with up to 100 integer arguments. The option is an argument of this function.

9.2.1.7 IntegerOption()

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)
out	value	integer value of given option

Returns

```
true or false for error
```

Function for any option with integer argument. The option (e.g. -n) is an argument of this function.

9.2.1.8 JoinCoorOption()

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	Counts	numbers of beads, molecules, etc.
out	MoleculeType	information about molecule types

Returns

```
true or false error or not error
```

Option whether to join aggregates and save joined coordinates into a specified file. Arguments: $-j < joined. \leftarrow vcf >$

9.2.1.9 MoleculeTypeOption()

```
char ** argv,
char * opt,
int * moltype,
Counts Counts,
MoleculeType ** MoleculeType )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)
in	Counts	numbers of beads, molecules, etc.
out	moltype	id of the molecule type
in	MoleculeType	information about molecule types

Returns

true or false for error

Generic option for molecule type that can take more than one argument. The option is an argument of this function.

9.2.1.10 MoleculeTypeOption2()

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)
in	Counts	numbers of beads, molecules, etc.
out	moltype	array for the molecule type
in	MoleculeType	information about molecule types

Returns

true or false for error

Generic option for molecule types. The option is an argument of this function.

9.2.1.11 SilentOption()

```
void SilentOption (
    int argc,
```

```
char ** argv,
bool * verbose,
bool * verbose2,
bool * silent )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
out	verbose	bool for -v option (verbose output)
out	verbose2	bool for -V option (detailed verbose output)
out	silent	bool for this option

Returns

true or false for error on common options

Option to not print anything to stdout (or at least no system definitions and no Step: #). Overrides VerboseShort ← Option and VerboseLongOption. Argument: ¬s

9.2.1.12 TwoIntegerOption()

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
in	opt	option switch (e.g. array containing -n)
out	values	array of two integer values of given option

Returns

```
true or false for error
```

Function for any option with two integer arguments. The option (e.g. -n) is an argument of this function.

9.2.1.13 VerboseLongOption()

```
void VerboseLongOption (
    int argc,
    char ** argv,
    bool * verbose,
    bool * verbose2 )
```

Parameters

in	argc	number of program's arguments
in	argv	program's arguments
out	verbose	bool for -v option (verbose output)
out	verbose2	bool for -V option (detailed verbose output)

Returns

true or false for error on common options

Option whether to print detailed data to stdout. Data are printed via VerboseOutput() function (and possibly some in-program code). Argument: $-\nabla$

9.2.1.14 VsfFileOption()

```
bool VsfFileOption (
                int argc,
                char ** argv,
                char ** vsf_file )
```

Parameters

in	argc	number of utility's arguments
in	argv	utility's arguments
out	vsf_file	filename with structure information

Returns

true or false error or not error

Option whether to use .vsf file different from the default $dl_meso.vsf$.

9.3 Structs.h File Reference

Structures for utilities.

Data Structures

• struct Vector

3D vector of floats.

struct LongVector

3D vector of floats.

struct IntVector

3D vector of integers.

struct Counts

Total numbers of various things.

struct BeadType

Information about bead types.

struct MoleculeType

Information about molecule types.

struct Bead

Information about every bead.

• struct Molecule

Information about every molecule.

• struct Aggregate

Information about every aggregate.

Macros

• #define PI 3.141593

value of pi

#define SQR(x) ((x)*(x))

macro for algebraic square

#define CUBE(x) ((x)*(x)*(x))

macro for algebraic cube

Typedefs

· typedef struct Vector Vector

3D vector of floats.

typedef struct LongVector LongVector

3D vector of floats.

• typedef struct IntVector IntVector

3D vector of integers.

typedef struct Counts Counts

Total numbers of various things.

typedef struct BeadType BeadType

Information about bead types.

typedef struct MoleculeType MoleculeType

Information about molecule types.

• typedef struct Bead Bead

Information about every bead.

• typedef struct Molecule Molecule

Information about every molecule.

typedef struct Aggregate Aggregate

Information about every aggregate.