EMC: Monte Carlo Simulations

A General Simulation Package EMC version 9.4.4, 1 August, 2022

This manual is for EMC (version 9.4.4, 1 August, 2022), a simulation package that can build and simulate both atomistic and coarse-grained systems using Monte Carlo techniques. Copyright © 2007-2021 Pieter J. in 't Veld.

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1 Introduction

1.1 General Introduction

Monte Carlo simulation techniques are wide spread through branches of scientific research and financial industry. It finds its application in statistical sampling of complex functional descriptions, giving numerical solutions to theoretically unsolvable equations. Typically codes using Monte Carlo techniques are written with a specific purpose in mind. These approaches include scientific implementations where versatility and not expandability is the main design goal. The latter finds an example in Towhee, a code which offers a wide array of force fields and sampling techniques, but encounters challenges when the need arises for implementation of new techniques. The code presented here does not try to offer the same breath, but rather follows a strategy of expandability. Strict modular design isolates key features in separate objects, yet a general parser allows for easy integration of these objects in a coherent design, thus allowing for a more flexible and expandable design. The purpose of this paper is to present a design philosophy for simulation engines in general, while applying this philosophy to the field of scientific Monte Carlo simulations. This is achieved by the identification of key features, such as input/output features, force field calculation, and configurational alterations, features which are highlighted in later sections.

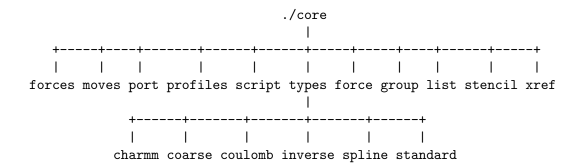
Each simulation code hinges around computational efficiency. Gain in speed can be obtained by an increase in processor speed, but more importantly, by judicious design of the program framework. This requires the identification of key factors of importance. Specifically, computational efficiency for this type of simulations is hampered by its serial nature: atoms or sites are moved one after another and not in a parallel fashion as is the case with Molecular Dynamics. Efficiency of energy calculations becomes the major concern when designing Monte Carlo codes for scientific applications. These applications hinge around the concept of an acceptance rule, in which typically is formed by a combination of system temperature with the difference in energy as the result of a change in configuration. This configuration change or move normally entails the displacement of one or a few sites. The serial nature of these moves necessitates the calculation of configurational energy both before and after the move. Optimization of energy calculations therefore plays an integral role in improving computational efficiency, for which this paper offers suggestions.

Alternatively, the speed with which a scientific field is allowed to advance rests on the foundations on which new developments can be build. Normally, code optimization results in an increase in code complexity, which in turn creates barriers for non-professional programmers to implement alterations necessary to test new scientific ideas. Strict separation of modules, yet flexible integration of concepts, creates a library-type of environment in which non-professional programmers – with the help of simple development tools – can use already existing modules as templates for their new ideas, thus creating a canvas on which only the minimal necessary changes need to be made.

1.2 Distribution Content

This distribution consist of a set of hierarchical directories, each of which represents a part of the code or documentation. The main directories are ./core, ./emc, ./examples, ./scripts, ./modules, ./lib, and ./texinfo. A general make file ./Makefile governs

compilation of separate pieces of this distribution. A 'make' in the root directory displays all compilation options. General routines are grouped under ./core, which has the following structure



Each subdirectory represents a substructure of main structures. Any module or object files placed in any of the directories are automatically compiled into each executable, provided that they conform to a required format associated with the substructure. Definition of macros based on file headers facilitates automatic compilation. This particular feature aids programming by example, which means an already existing module can be used as a template, rather than having to completely write a new model. A copy script to facilitate this process is provided by ./scripts/cpmod.pl.

Currently, the main program, which includes a scripting front-end, is located in ./emc. This directory also contains example building and running scripts in ./examples. Upon compilation through execution of 'make emc' generates a new directory ./emc/objects, which contains all .o files. Examples in ./examples make use of predefined crystal structures in ./lib.

All documentation is organized in ./texinfo. This PDF document is generated by a 'make pdf' in this directory.

2 Methodology

2.1 Lists

The presented code employs a combination of cell lists and verlet lists, where a separate object represents each list. Spatial discretization forms the cell list basis and is determined by either the largest or the smallest force field cut off, combined with a user supplied skin. The choice of cut off depends on invoking a conventional cell list, in which case the largest cut off is selected, or a multi-neighbor cell list, in which case the smallest cut off is chosen (Intveld2008). Both lists store their members in a linked list, which is accomplished by the application of a stencil or template. This template describes which cells neighbor the central cell. A distance criterion – based on the cut off and skin – then decides which sites are eventually added to a verlet list. Besides an integer vector to neighboring cells, a multineighbor template also contains the minimum distance between the cells containing the each site contributing to the pair interaction, which enables a primary selection of sites based on their cut off and therefore negating the initial need of a distance calculation. Inclusion in a verlet list only then requires a distance calculation when the neighboring particle falls within the stored minimum distance. Application of the multi-neighbor principle is optional, although initial tests do not indicate a significant drop in performance compared conventional cell list implementations.

Verlet lists describe which neighboring sites fall within a certain distance criterion of a site of interest. These lists are described by linked lists. Cross-linking between verlet lists has been added to allow for easy removal of a site when performing a Monte Carlo move. Practically, this means that each verlet list member of one site knows which verlet list member of another site it is connected to, thus creating members that contain pointers to child and parent in one particle's list and a pointer to a member of another particle's list. Additionally, the squared distance between the two sites has been added to negate the need for distance calculations during non-bonded force field evaluation.

Typically, the application of verlet lists in Molecular Dynamics makes use of the principle of a skin, which is a distance added to the force field cut off, and serves the purpose of reducing the number of verlet list rebuilds. However, an increase in distance calculations, as a result of inclusion of more sites in the verlet list, forms the downside of skin application. Hence, a balance needs to be found between performance gain as a result from less verlet list builds and performance loss as a result of an increase in distance calculations. Typical numbers used for the skin are 0.3σ in Lennard-Jones (LJ) units or 1 Å in atomistic units. An added advantage of the usage of skins is that the decision to rebuild a verlet list can be coupled to the distance with which a site has moved from when the verlet list was built last for that site. A safe criterion is to rebuild the list when the site has moved by more than half the skin. Application of this criterion enables a dynamic update of verlet lists based on particle positions only and not by an external constraint such as a preset number of passed time steps or cycles, as is commonly used in packages such as LAMMPS (*Plimpton1995*).

2.2 Force Fields

Implemented force fields include standard representations of the OPLS force field as described by Jorgensen et al. (*Jorgensen*), the CHARMM force field as described by MacKerell

et al. (MacKerell1998), and a Class 2 force field representation used by PCFF or COM-PASS force fields as described by Sun (Sun1998). A separate module describes coulombic interaction and allows for the use of either a cut off or Ewald summations to account for the long-range character (Frenkel). Coarse-grained force fields are provided by a colloidal force field as described by Everaers and Etjehadi (Everaers2003), which includes additional cross-terms for interaction between Lennard-Jones and colloidal particles (Grest2007, Veld2008), and the FENE bond potential as described by Kremer and Grest (Kremer). Pressures are calculation through standard virial expressions, which were derived when needed. All interactions can be compounded without significant effects on performance.

Generalization of virial calculations requires a solution to the gradient of a potential with respect to the cartesian coordinate system, which entails splitting partial derivatives into a product of linear parts and cartesian parts. For a nonbonded potential dependent on radius r this results in

$$\frac{\partial U(r)}{\partial \vec{x}} = \frac{\partial U(r)}{\partial r} \frac{\partial r}{\partial \vec{x}} = \frac{\partial U(r)}{\partial r} \frac{\vec{x}}{r},$$

where \vec{x} represents the constituent vector of a cartesian axis frame. Consequently, the virial tensor W_r resulting from this expression becomes

$$W_r(r_{ij}) = -\frac{\partial U(r)}{\partial r} r_{ij} \Delta \vec{r}_{ij} \Delta \vec{r}_{ij}.$$

where r_{ij} denotes the distance between site i and site j and $\Delta \vec{r}_{ij} \Delta \vec{r}_{ij}$ denotes a vector product resulting in a tensor. Bonded potentials potentials depend on bond length l_{ij} , angle θ_{ijk} , torsion ϕ_{ijkl} , and improper ψ_{ijkl} , which are defined as

$$\begin{split} \Delta \vec{r}_{ij} &= \frac{\vec{r}_j - \vec{r}_i}{|\vec{r}_j - \vec{r}_i|} \\ \vec{n}_{ijk} &= \frac{\Delta \vec{r}_{jk} \times \Delta \vec{r}_{ij}}{|\Delta \vec{r}_{kj} \times \Delta \vec{r}_{ji}|}, \\ r_{ij} &= l_{ij} = |\vec{r}_j - \vec{r}_i|, \\ \theta_{ijk} &= \Delta \vec{r}_{ij} \cdot \Delta \vec{r}_{jk}, \\ \phi_{ijkl} &= \vec{n}_{ijk} \cdot \vec{n}_{jkl}, \\ \psi_{ijkl} &= \vec{n}_{ijk} \cdot \vec{n}_{ijl}. \end{split}$$

Note that j represents the center site of impropers. The bond virial tensor W_l for potentials depending on l_{ij} is described by

$$W_l(l_{ij}) = -\frac{\partial U(l_{ij})}{\partial l_{ij}} l_{ij} \Delta \vec{r}_{ij} \Delta \vec{r}_{ij}.$$

The angular virial tensor W_{θ} for potentials depending on θ_{ijk} is described by

$$W_{\theta}(\theta_{ijk}) = -\frac{\partial U(\theta_{ijk})}{\partial \theta_{ijk}} \frac{1}{\sin(\theta_{ijk})} \left[\Delta \vec{r}_{ij} \left(\Delta \vec{r}_{jk} + \cos(\theta_{ijk}) \Delta \vec{r}_{ij} \right) + \Delta \vec{r}_{jk} \left(\Delta \vec{r}_{ij} + \cos(\theta_{ijk}) \Delta \vec{r}_{jk} \right) \right].$$

Note that vector product $\Delta \vec{r}_{ij} \Delta \vec{r}_{jk}$ is noncommunicative, i.e. $\Delta \vec{r}_{ij} \Delta \vec{r}_{jk} \neq \Delta \vec{r}_{jk} \Delta \vec{r}_{ij}$. The torsional virial tensor W_{ϕ} for potentials depending on ϕ_{ijkl} is described by

$$\begin{split} W_{\phi}(\phi_{ijkl}) &= -\frac{\partial U(\phi_{ijkl})}{\partial \phi_{ijkl}} \frac{1}{\sin(\phi_{ijkl})} \left[\\ & \left(\cos(\phi_{ijkl}) \Delta \vec{r}_{jk} \times \vec{n}_{ijk} + \Delta \vec{r}_{jk} \times \vec{n}_{jkl} \right) \Delta \vec{r}_{ij} \\ & + \left(\cos(\phi_{ijkl}) \left(\Delta \vec{r}_{kl} \times \vec{n}_{jkl} - \Delta \vec{r}_{ij} \times \vec{n}_{ijk} \right) + \Delta \vec{r}_{kl} \times \vec{n}_{ijk} - \Delta \vec{r}_{ij} \times \vec{n}_{jkl} \right) \Delta \vec{r}_{jk} \\ & - \left(\cos(\phi_{ijkl}) \Delta \vec{r}_{jk} \times \vec{n}_{jkl} + \Delta \vec{r}_{jk} \times \vec{n}_{ijk} \right) \Delta \vec{r}_{kl} \right]. \end{split}$$

The improper virial tensor W_{ψ} for potentials depending on ψ_{ijkl} is described by

$$W_{\psi}(\psi_{ijkl}) = -\frac{\partial U(\psi_{ijkl})}{\partial \psi_{ijkl}} \frac{1}{\sin(\psi_{ijkl})} \left[(\cos(\psi_{ijkl}) \Delta \vec{r}_{jk} \times \vec{n}_{ijk} + \Delta \vec{r}_{jk} \times \vec{n}_{ijl}) \Delta \vec{r}_{ij} + \cdots + (\cos(\psi_{ijkl}) \Delta \vec{r}_{jk} \times \vec{n}_{ijl} + \Delta \vec{r}_{jk} \times \vec{n}_{ijk}) \Delta \vec{r}_{kl} \right].$$

3 Program Structure

This particular Monte Carlo simulation code is modular in approach. While it uses C as its programming language foundation, it incorporates distinct object oriented features. Communication by all modules to the outside, it being storage or inter-node communication, happens through a generalized parsing module, which can be used for both textual and binary output. Binary output can be either to memory or file. Variables are stored in a tree structure of pointers with cross-links between branches when so required. The textual input/output file bears an identical but simplified structure compared to the tree structure in memory: communication only conveys essential information and cross-links are implied. The structure 'simulation' presents the top structure, consisting of pointers to the structures 'io', 'identity', 'clocks', 'output', 'units', 'channels', 'types', 'systems', 'moves', 'statistics', 'forces', 'profiles', 'clusters', 'groups', and 'sites'. These structures are grouped in the following categories:

- Molecular representation by 'clusters', 'groups', and 'sites'
- Molecular interaction by 'types' and 'forces'
- Simulation parameter definition by 'systems'
- Configuration changes by 'moves'
- Measurements by 'statistics' and 'profiles'
- Communication by 'io', 'output', and 'units', and 'channels'
- Simulation timing by 'clocks'
- Simulator identification by 'identity'

3.1 Molecular Representation

A need for generality motivates the chosen terminology of 'clusters', 'groups', and 'sites'. Clusters can represent molecules and generally define closed networks of connected sites. Groups are of the order of repeat unit or protein residue and generally define subsets of molecules, creating the possibility of group operators when applying configuration changes. Sites can resemble atoms, but could also represent a more coarse-grained model of a group of atoms lumped into one site. Sites are the backbone of all interactions. This section describes the hierarchy from the bottom up, starting with 'sites', followed by 'groups' and 'clusters'.

3.1.1 Sites

Description of sites.

3.1.2 Groups

Description of groups.

3.1.3 Clusters

Description of clusters.

3.2 Molecular Interactions

3.2.1 Types

Description of types.

3.2.2 Forces

Description of forces.

3.3 Systems

Description of systems.

3.4 Configurational Moves

Description of moves.

3.5 Measurements

Description of distributions.

4 Simulation Setup

4.1 General

EMC provides a setup script emc_setup.pl, which allows for creation of EMC build and LAMMPS input scripts. The EMC setup script serves as a wrapper around EMC itself, creating a context for abstraction of EMC scripting commands. In its simplest appearance, setup scripts only contain a few lines describing simulated chemistry and their quantities, where subsequent options are defined through the command line. More complex scripts contain several paragraphs for describing simulation conditions and chemistry. This mode will be referred to as the chemistry mode (see Section 4.6.2 [Chemistry File], page 36). The most complex mode creates an environment in which several simulations are defined in one script, allowing for looping over multiple simulation conditions. This mode will be referred to as the environment mode (see Section 4.6.1 [Environment File], page 33). EMC environment setup scripts function as a wrapper around chemistry scripts. An environment setup script can hold multiple chemistry scripts. Paragraphs use keyword ITEM as demarcation, where identifiers follow this keyword at the beginning of each paragraphs. Their ending is marked by ITEM END.

Normal parametrization uses force fields as provided by EMC (see Section 5.11 [Field], page 103) for typing. These force fields can be found in \${EMC_ROOT}/field/. Optional references.csv and parameters.csv files can be provided for DPD force fields. The latter is called from the chemistry file. Alternatively, a more general way in provided by using the field paragraph (ITEM FIELD) in an environment script. Both produce a force field file with extension .prm. Force fields are governed over types definitions (see Section 7.12 [Types], page 201). The EMC setup script can create multiphase initial configuration, but does not address the grafting capabilities of the EMC build capabilities. The EMC setup script displays all available options when called without arguments. It is, in general, called by

It can also, however, be used as an interpreter, where EMC setup scripts can function as executables when using EMC in a UNIX-type environment. An interpreter allows for including all settings and options into an EMC script, thus folding the description of one simulation or a set of simulations into one comprehensive script.

4.2 Setup Usage

Examples can be found in \${EMC_ROOT}/examples/setup/. Creating EMC and LAMMPS input with these examples occurs in steps. The first step is to convert the example setup script by means of emc_setup.pl through

```
emc_setup.pl example.esh
```

which either creates an input script for EMC, called build.emc by default, and an input script for LAMMPS, called example.in by default, or sets up an execution environment by creating subdirectories build, run, analysis, and chemistry. The first three directories contain bash scripts with the purpose of respectively building input structures, running molecular dynamics simulations, and analysis of end results. The chemistry directory can contain force fields, variation in chemistry, polymer make up, etc. Default names can be altered when desired (see \(\)\ undefined \(\)\ [Environment Options], page \(\)\ \(\)\ undefined \(\)\.

The second step involves a call of EMC itself, by for instance for LINUX,

```
emc_linux build.emc 2>&1 | tee build.out
```

The latter part of the command allows for writing output to screen and build.out simultaneously. Five extra files appear upon completion of this command, i.e. example.pdb, example.psf, and example.vmd for visualization with VMD, and example.data and example.params for use with LAMMPS. Visualization supports checking the validity of the final structure. The call to VMD, when in the user's path, is facilitated by,

```
vmd -e example.vmd
```

The example.vmd file represents a small Tcl script, which contains information about particle diameters (as defined by the chosen force field) and the type of VMD presentation to use for visualization. A molecular dynamics simulation with LAMMPS occurs with for example

```
lmp_linux <example.in 2>&1 | tee example.out
```

The example.in LAMMPS script contains all the necessary information for executing the run. Please note, that the presented examples in the \${EMC_ROOT}/examples/setup/directory are primarily meant as building examples. Strategies for managing sets of simulations are provided by the environment mode of emc_setup.pl (see Section 4.6.1 [Environment File], page 33), for which an example can be found in \${EMC_ROOT}/examples/setup/shear/.

4.3 Extensions

A comprehensive mode for EMC Setup is when using environment scripts, which creates a modeling environment consisting of build, run, and analysis scripts, complemented with a hierarchical structure for dealing with produced imulation data. The simpler EMC chemistry script creates a subset of the more comprehensive environment script, but is useful for prototyping simulations. Extensions an explanations for scripts and data files produced by both modes following from EMC Setup, EMC, LAMMPS or postprocessing analysis can be found in the following table,

Extension . data	Description EMC output containing LAMMPS data file defining coordinates, types, and topology
.csv	Postprocessed data derived from EMC or LAMMPS output resulting from execution of EMC analysis scripts; normally used for further processing by user generated scripts (using e.g. Python, Perl, Matlab, or Mathematica)
.density	Density profile produced by LAMMPS
.density3d	3D density profile produced by LAMMPS
.emc	EMC script as interpreted by EMC itself when produced by emc_setup; used to create input structures by means of energetic considerations; EMC produces a comprehensive EMC format with the same extension as output containing all topological and morphological information
.energy	Time-averaged energetic information produced by LAMMPS
.esh	EMC Setup script as interpreted by wrapper emc_setup.pl; can be either a chemistry or an environment script
.in	LAMMPS input script according to specifications as made in a EMC chemistry file
.params	EMC output following from a successful build; input needed by LAMMPS, containing all paramaters defining a full simulation
.pdb	PDB format containing initial coordinates
.pressure	Time-averaged pressure information produced by LAMMPS
.prm	EMC parameter file as produced by either EMC chemistry or environment script; used by EMC to set force field energetics

.psf PSF format containing types and (initial) topology

.vmd VMD Tcl script used to read in both .pdb and .psf, additionally defining

bead sizes following from the used force field

.volume Time-averaged volume information produced by LAMMPS

4.4 Environment Options

Option	Default	Description
analyze_archive	true	Toggle archiving of file names associated with analyzed data; file name lists can be used directly or at a later date for transferring data between computational clusters and local machines.
analyze_data	true	Toggle creation of tar archive from exchange file list; this data is used in combination with ./scripts/run_host.sh -exchange to transfer data from computational clusters to local machines; archives should not be created when analysis scripts subsequently submit analysis to queue.
analyze_last	false	Toggle inclusion of the last trajectory frame during analysis (deprecated); analysis is transferred to the analysis paragraph (see Section 4.6.1 [Environment File], page 33).
analyze_replace	true	Toggle replacement of already exisiting analysis results.
analyze_skip	0	Set the number of initial frames to skip during analysis.
analyze_source	-	Specify alternate data source directory for analysis scripts.
analyze_user	-	Set directory for user analysis scripts, which can be accessed through an analysis paragraph (see Section 4.6.1 [Environment File], page 33).
analyze_window	1	Set the number of frames over which to apply a window averaging during analysis.
modules	-	Manipulate runtime modules; format is defined by [command]=module separated by commas.
name_analyze		Set the name of job analysis scripts; analysis scripts are stored in ./analyze/; takes the base name of the chemistry script when omitted; ignored when "-" is chosen as name.

name_build		Set the name of job build scripts; build scripts are stored in ./build/; takes the base name of the chemistry script when omitted; ignored when "-" is chosen as name.
name_run		Set the name of job run scripts; run scripts are stored in ./run/; takes the base name of the chemistry script when omitted; ignored when "-" is chosen as name.
name_scripts	-	Simultaneously set analysis, build and run script names.
name_testdir	-	Set the name of a test directory in which to test the chemistry paragraph in an environment script; a setup script is created in ./test/; ignored as default.
nchains	-	Set the number of chains to be used for executing LAMMPS jobs; allows for having the next job wait on the previous (currently only with LSF).
ncores	-	Set the number of cores for execution of LAMMPS jobs.
ncorespernode	default	Set number of cores per node for execution of packed jobs; equivalent to queue_ppn.
preprocess	false	Apply C preprocessing to allow for programmatic flow when interpreting templates; C preprocessing start at the beginning of a line and are preceded by a #-mark; allowed options are define, if, elif, else, and endif.
project	-	Set the project name; derived from setup script name when not given.
queue	account=none, analyze=default, build=default, memory=default, ncores=-, ppn=default, run=default, user=none	Directly set queueing system settings, where option account sets billing account, analyze sets the analysis queue, build sets the build queue, memory sets the amount of memory needed, ncores sets the number of cores for running, ppn sets the number of processors per node, run sets the run queue, and user sets user options; the latter is dependent on the installed queueing system.

queue_account	none	Set queue account for billing.
queue_analyze	default	Set the queue used in analysis scripts; option default indicates the use of queue, which is set as system default; option local will sequentially execute all jobs on local machine.
queue_build	default	Set the queue used in build scripts; option default indicates the use of queue, which is set as system default; option local will sequentially execute all jobs on local machine.
queue_memory	default	Set memory per core in gb for executed jobs.
queue_ppn	default	Set cores per node for executed jobs.
queue_run	default	Set the queue used in run scripts; option default indicates the use of queue, which is set as system default; option local will sequentially execute all jobs on local machine.
queue_user	none	Options to be passed directly to queuing system; allowed option are dependent on the installed queueing system.
replace	false	Replace or overwrite all written script files.
time_analyze	00:30:00	Set the desired wall time for analysis scripts.
time_build	00:10:00	Set the desired wall time for build scripts.
time_run	24:00:00	Set the desired wall time for run scripts.
workdir	-	Set the work directory, which is used as a base for creating the EMC work environment.

4.5 Chemistry Options

Option	Default	Description
angle	5,180	This option can be used to set angle constants in three ways: 1. set DPD angle constants (k, theta0), 2. provide additional angle table entries through type1, type2, type3, k, theta0, 3. set force field handling for angle typing; valid options are ignore, complete, warn, empty, and error.
auto	false	Include wildcard mass entry in generated DPD .prm force field file; options are true or false.
binsize	0.01	Set the general bin size to be used in LAMMPS profiles; note, that profiles are recorded in reduced units, i.e. coordinates run from 0 to 1.
bond	25,1	Set DPD bond constants (k, 10); can also be used to provide additional bond table entries through -bond=type1,type2,k,10.
build	build	Set build script name.
build_center	false	Place first site at the origin (see also option build_origin); options are true or false.
build_dir	/build	Set location of build directory for LAMMPS script.
build_order	false	Build clusters in the order as defined by option phases; options are random or sequence.
build_origin	x=0, y=0, z=0	Set alternate origin at which to place te first site.
build_replace	false	Toggle replacement of already existing build scripts and results, which will skip build execution when a LAMMPS data file exists.
build_theta	false	Set the minimal insertion angle
charge	true	Indicator for occurence of charges in chemistry.csv.
charge_cut	9.5	Set pairwise charge interaction cut off.

chemistry	chemistry	Set chemistry input file name; extension $.\mathtt{csv}$ is implied.
chunk	true	Use chunk approach for computing profiles and samples in LAMMPS script; options are true or false.
communicate	false	Use communicate keyword instead of comm_modify in LAMMPS input script; used with DPD simulations for backwards compatibility with older versions of LAMMPS; options are true or false.
core	-1	Set particle core diameter; used with Born potentials.
cross	false	Include nonbond cross terms in LAMMPS params file; default depends on chosen force field; options are true or false.
crystal	false	Let EMC derive a crystal structure based on the offered import; options are true or false.
cut	9.5	Set pairwise interaction cutoff; options are either definite positive numbers or repulsive; the latter transfers the the parameters for a purely repulsive cut and shift potential to LAMMPS
cutoff	<pre>center=-1, charge=9.5, ghost=-1, inner=-1, outer=-1, pair=9.5, repulsive=0, rmax=-1</pre>	Set various cutoffs; options are either definite positive numbers; negative numbers refer to internal defaults; the different cutoffs can also be set with single options charge_cut, ghost_cut, inner_cut, outer_cut, cut, and rmax.
debug	-	Turn debugging information on; options are true or false.
deform	nblocks=1, ncycles=100, type=relative, xx=1, yy=1, zz=1, zy=0, zx=0, yx=0	Invoke affine deformation using Monte Carlo relaxation when ncycles is larger than zero; defomation type can either be absolute or relative.

delete	phase=1, fraction=1.0, thickness=1, type=relative, mode=include, sites=all, groups=all, clusters=all	Defines the selection criteria for deleting clusters at the edges of the box; phase sets the phase to which to apply deletion; fraction currently is not used; thickness sets the thickness of the region taken into consideration; type sets the units of the thickness (valid options are absolute and relative); mode is currently not used; sites sets the sites to include in the selection (valid options are all for all available sites or a selected list of site types separated by colons); groups sets the groups to include in the selection (valid options are all for all available groups or a selected list of group ids separated by colons); clusters sets the clusters to include in the selection (valid options are all for all available clusters build in the set phase or a selected list of cluster ids separated by colons); only full clusters are deleted, even when only one site of these clusters complies with the set selection.
density	1[,]	Set simulation density; density of separate phases can be entered through separating values for each phase by commas; units and default depend on chosen force field type.
depth	auto	Set the depth with which rings - as defined in groups - are recursively assigned; options are either auto or values larger than 2.
dielectric	0.2 or 1	Set dielectric constant of medium; default depends on chosen force field.
direction	x	Set direction in which phases in setup scripts with multiple phases are build; valid options are x , y , or z .
dtdump	100000	Frequency with which LAMMPS adds to trajectory files.
dtrestart	100000	Frequency with which LAMMPS writes restart files.
dtthermo	1000	Frequency with which LAMMPS generates thermodynamical output.
emc	true	Create EMC build script (.emc file); options are true or false.

emc_exclude	build=false	Exclude specified sections from the resulting EMC script; excluding build omits morphology generation, which can be desired when exporting SMILES.
emc_execute	-	Execute the EMC build script as created by the setup script; options false or - do not execute; option true allows EMC setup decide which EMC version to use, based on operating system, host option, and path; any other option value is interpreted as the location and name of the EMC version to use.
emc_export	smiles=false	Add export of specified format; smiles controls export of SMILES for all built clusters in both group-based and site-based representations; option false deselects export; other valid options are csv, json, and math for comma separated values, JSON, and Mathematica formats respectively.
emc_moves	displace=1	Control frequency of selected Monte Carlo move for positive values; zero or negative values unselect.
emc_output	<pre>debug=false, exit=true, info=true, warning=true</pre>	Control output as generated by EMC scripts; options for each separate keyword are true or false.
emc_progress	build=true, clusters=false	Control progress indicators resulting from executing EMC script; keyword build controls progress output during build; keyword cluster controls progress output during sequencing and cluster construction; options for each separate keyword are true or false.
emc_run	<pre>nblocks=100, ncycles=0, nequil=0, clusters=all, groups=all, sites=all</pre>	Execute a Monte Carlo simulation after building for ncycles; optionally, an equilibration phase is selected when nequil is larger than zero; nblocks selects the frequency with which output is generated; specific selections can be set when defining either clusters, groups, or sites separately or in combination; all indicates that all members of a selection are considered.
emc_test	false	Test the validity of an EMC environment script; checks up to execution of EMC build files, as generated by EMC environment; options are true or false.

emc_traject	append=true, frequency=0	Add generation of trajectory file during Monte Carlo run when frequency is larger than zero.
environment	false	Interpret EMC shell script (.esh) as an environment script; options are true and false.
ewald	true	Switch the electrostatics long-range treatment through Ewald summations on or off; default depends on the chosen force field; options are true or false.
expert	false	Invokes expert mode, which allows for overrides of certain consistency checks (e.g. nrepeat for polymers); options are true or false.
extension	.esh	Set the extension of environment scripts; extensions are derived from given environment scripts, when not defined.
extra	-	Create copies of existing nonbond interactions; format is destination:source:offset, where offset defines the DPD offset of the new interaction constant compared to the original (deprecated, replaced by replica)
field	opls	Set force field type and name based on root location; more general access to EMC provided force fields uses the type option below.
field_angle	-	Set error handling for typing of angles; valid options are complete, empty, error, ignore, and warn.
field_bond	-	Set error handling for typing of bonds; valid options are complete, empty, error, ignore, and warn.
field_dpd	<pre>auto=false, bond=false</pre>	Set options influencing the generation of a DPD force field; possible options are auto and bond with values of either true or false; auto controls the addition of wildcard additions to mass and nonbonded force field paragraphs; bond allows for verbatim transcription of pair to bond interactions.

field_debug	false	Set debug option, showing individual typing steps; valid options are full, reduced, and false; full outputs verbose debugging information, reduced generates an abridged selection, and false turns debugging information off.
field_error	true	Exit upon error (set to false when debugging a force field).
field_format	%15.10e	Sets the format of parameters in the produced field; should be format as is standard for floating point values when used in combination with printf().
field_group	-	Set field group option for applying force fields to groups as defined by their TEMPLATE section.
field_id	-	Set force field id.
field_improper	-	Set error handling for typing of impropers; valid options are complete, empty, error, ignore, and warn.
field_increment	-	Set error handling for typing of bond increments; valid options are complete, empty, error, ignore, and warn.
field_location	-	Provide force field location; intended for expert use; use field_type for EMC provided force fields.
field_name	-	Provide force field name; intended for expert use= use field_type for EMC provided force fields.
field_nbonded	0	Define the number of sites, that are bonded, but should be excluded from nonbond interactions, e.g. a value of 1 excludes 1-2 interactions, a value of 2 excludes 1-2 and 1-3 interactions, etc.; allowed values are larger or equal than 0; this option only applies to DPD force fields.
field_torsion	-	Set error handling for typing of torsions; valid options are complete, empty, error, ignore, and warn.

field_type	opls	Provide force field type; invoking this option will automatically set field, field_location, and field_name options.
field_write	true	Interpret and write the force field as defined in the ITEM FIELD paragraph (see Section 4.6.3 [Field File], page 46)
focus	-	List of clusters to focus on.
grace	0.9999, 0.9999,0	Depreciated command, use weight instead; user provided grace for building procedure; given in order of nonbonded, bonded, and focussed interactions.
ghost_cut	-	Set molecular dynamics ghost region size for property communication.
help	-	Display script help.
hexadecimal	false	Set hexadecimal output in PDB files.
host	-	Set host on which to run EMC and LAMMPS.
improper	-	Set force field handling for improper typing; valid options are ignore, complete, warn, empty, and error.
increment	-	Set force field handling for increment typing; valid options are ignore, complete, warn, empty, and error.
info	-	Turn on information during setup script execution.
inner	-	Set inner cutoff; use and default depend of chosen force field; note that inner and outer cutoffs are interpreted as fractions for colloidal force fields.
insight	true	Create InsightII CAR and MDF output files.
insight_ compress	true	Control compression of produced CAR and MDF files; options are true or false.
insight_pbc	true	Toggle application of periodic boundary conditions; options are true or false.

insight_unwrap	clusters	Set CAR unwrapping mode; options are none, clusters, or sites.
kappa	1 or 4	Set long range electrostatics Ewald summation kappa; default depends on chosen force field.
lammps	true	Create LAMMPS input script (.in file) or set lammps version using year, e.glammps=2014 (new versions start at 2015) or keywords old or new; alters the settings for -communicate and -chunk; default is new.
lammps_cutoff	false	Toggle output of pairwise cut off in parameter file; valid options are true or false.
lammps_dlimit	0.1	Set LAMMPS nve/limit distance used during equilibration.
lammps_error	false	Only restart those LAMMPS simulations for which an error occurred; valid options are true or false.
lammps_pdamp	1000	Set LAMMPS barostat damping constant for NPT simulations.
lammps_tdamp	100	Set LAMMPS thermostat damping constant for both NVT and NPT simulations.
location	<pre>analyze=., field=., include=.</pre>	Prepend path for locations of analysis scripts, force field files, and include files; always included are for 1) analyze: directories . and \${EMC_ROOT}/scripts/analyze, 2) field: directories . and \${EMC_ROOT}/field, and 3) include: directories . and \${WORKDIR}/chemistry/include when existing; multiple path entries are separated with a colon.
mass	false	Assume mass fractions in chemistry.csv input file.
memorypercore	default	Set memory per core in gigabyte for executed jobs; equivalent to queue_memory.
mol	true	Assume mol fractions in chemistry.csv input file.

momentum	100,1,1,1, angular	Control zeroing of linear and angular momentum during LAMMPS execution; order of entries (separated by commas only!), i.e. N,xflag,yflag,zflag,flag (see LAMMPS) manual; flag is omitted in the LAMMPS input script when the keyword none is used.
moves_cluster	<pre>active=false, cut=0.05, frequency=1, limit=auto:auto, max=0:0, min=auto:auto</pre>	Define cluster move settings used to optimize builds; multiple options are available, where option active switches the move on or off, cut sets a percentage of the maximum allowed displacement and rotation at which to turn the move off, frequency sets the frequency of the move, limit sets the maximum allowed displacement (in force field units) and rotation (in rad), max sets estimation for 50% acceptance of displacement and rotation (0:0 will trigger internal estimation), and min sets explicit rather than relative conditions to turn the move off.
msd	false	Add mean square displacement analysis to LAMMPS output; options are true or false.
namd	false	Create NAMD input script and parameter file.
namd_dtcoulomb	1	Set electrostatic interaction update frequency.
namd_dtdcd	10000	Set frequency with which snapshots are written to a DCD file.
namd_dtnonbond	1	Set nonbond interaction update frequency.
namd_dtrestart	100000	Set output frequency of restart files.
namd_dtthermo	1000	Set output frequency of thermodynamic quantities.
namd_dttiming	10000	Set timing frequency.
namd_dtupdate	20	Set update frequency.
namd_pres_decay	50	Set pressure ensemble decay.
namd_pres_ period	100	Set pressure ensemble period.
namd_temp_damp	3	Set temperature ensemble damping.

namd_tminimize	50000	Set number of initial minimization steps.
namd_trun	10000000	Set total number of timesteps for execution run.
nchains	-	Set the number of chains to use for execution of molecular dynamics jobs.
ncores	-1	Set the total number of cores to use for execution of one molecular dynamics job.
ncorespernode	default	Set the number of cores per node as used in packing multiple jobs on one one, when possible.
niterations	1000	Controls the number of iterations used for inserting an atom during the build process.
norestart	false	Control the capability for using run scripts – as resulting from environment scripts – for restarting; options are true or false; the latter means, that restarting with the same run script is possible.
nparallel	auto	Set the number of repeat units in the direction parallel to a surface defined in the chemistry file; values larger than 1 override the auto setting.
nrelax	100	Alter the number of relaxation cycles used during the EMC building process.
nsample	1000	Number of configurations used for averaging during execution of LAMMPS analysis routines.
ntotal	10000	Total number of created sites or atoms.
number	false	Interpret column 3 of SHORTHAND or CLUSTERS as the number of desired clusters; this number is an integer (see Section 4.6.2 [Chemistry File], page 36).
omit	false	Omit fractions from chemistry file; options are true or false.
outer	-	Set outer cutoff; note that inner and outer cutoffs are interpreted as fractions for colloidal force fields.

pair	a=25, gamma=4.5, r=1	Set DPD pair constants; only definite positive values are allowed.
parameters	parameters	Alternative parameter file name; the extension $.\mathtt{csv}$ is implied.
params	true	Create force field parameter file (.prm); currently only for DPD; options are true or false.
pdb	true	Create PDB and PSF output files.
pdb_atom	index	Set atom name behavior in PDB and PSF output; options are detect, index, and series.
pdb_compress	true	Control compression of produced PDB and PSF files; options are true or false.
pdb_connect	false	Add connectivity to PDB; options are true or false.
pdb_cut	false	Cut bonds in PSF output, which span the simulation box; options are true or false.
pdb_extend	false	Use extended format in PSF output; options are true or false.
pdb_parameters	false	Toggle output of NAMD parameter file; options are true or false.
pdb_pbc	true	Toggle application of periodic boundary conditions; options are true or false.
pdb_fixed	true	exclude flagged fixed sites when unwrapping clusters; options are true or false.
pdb_rigid	true	exclude flagged rigid sites when unwrapping clusters; options are true or false.
pdb_residue	index	Set residue name behavior in PDB and PSF output; options are detect, index, and series.
pdb_segment	index	Set segment name behavior in PDB and PSF output; options are detect, index, and series.

pdb_unwrap	clusters	Set PDB unwrapping mode; options are none, clusters, or sites.
pdb_vdw	true	Add Van der Waals representation to the VDW script created by EMC; options are true and false.
percolate	false	Treat imported structures as percolating InsightII structures; uses boundary crossing definitions as provided in the .mdf file; behavior of crystal option might be preferred; options are true or false.
phases	all	Sets which clusters to assign to each phase; each phase is separated by a +-sign; default assigns all clusters to phase 1; build order is defined by option build_order
polymer	bias=none, fraction=number, niterations=-1, order=list	Set group polymer distribution global defaults; bias helps in limiting construction time by controlling exclusion of unsuccessful subpolymers during polymer construction; valid bias options are none for no bias, binary for exclusion of subpolymers on a binary basis, and accumulative for exclusion on an accumulative basis; valid fraction options are number for using number of molecules and mass molecule mass as distribution entries; valid order options are list for sequentially and random of randomly interpreting the provided polymer distribution; the niterations keyword sets the maximum allowed number of iterations used during polymer sequence determination
precision	0.001	Provide LAMMPS kspace long range Ewald summation precision.
prefix	false	Set project name as prefix to LAMMPS output files.

pressure	<pre>false, direction= x+y+z, couple</pre>	Set the system pressure and invoke an NPT ensemble; optionally specify barostat direction and/or directional coupling; valid directions are x, y, and z; valid coupling options are couple, uncouple, true, and false, where true corresponds to coupled directions; definition of the coupling direction follows the couple keyword through separation by a colon; a + sign separates multiple directions, e.g. atomistic 1,couple=y+z sets a pressure of 1 atm in all directions with coupling in y and z directions; omission of coupling directions assumes an isotropic barostat.
profile	<pre>density=false, density3d=false, pressure=false</pre>	Generate density and/or pressure profiles of all clusters while executing LAMMPS; options are density, density3d, or pressure, followed by either true or false, separated by a colon; true is assumed when omitted; options are separated by a comma.
project	_	Set the project name; derived from setup script name when not given.
queue	account=none, analyze=default, build=default, memory=default, ncores=-1, ppn=default, run=default, user=none	Control queue settings, where account sets accounting information as needed by the queueing system, memory the amount of memory per core in gigabytes, ncores the total number of cores to be used for one MD simulation, ppn the number of cores per node; specific queues can be set for analysis, building and running through analyze, build, and run respectively; any user-defined queueing options can be passed to the queueing system directly with user; each of these commands can also be accessed separately as given by the following queue keywords.
queue_account	none	Set queue account for billing.
queue_analyze	default	Set the queue in which to execute the analysis scripts.
queue_build	default	Set the queue in which to execute the build scripts.
queue_memory	default	Set the memory per core in gigabytes.
queue_ncores	-1	Set the number of cores for execution of MD jobs.

queue_ppn	default	Set the number of cores per node as used for packing multiple jobs on one node.
queue_run	default	Set the queue in which to execute the run scripts.
queue_user	none	Set user-defined queueing options to be passed verbatim to the queueing system.
quiet	-	Supress all output generated by this setup script.
radius	5	Alter the radius of the spherical volume in which already built sites are relaxed; used during the EMC building process.
record	<pre>cut=false, frequency=1, inactive=true, name="", pbc=true, unwrap=true</pre>	Defines the record entry in the build paragraph, which records the build process by outputting a set of PDBs and PSFs; valid options are cut for cutting bonds in the last recorded frame, frequency for setting the frequency with which the relaxation process is written in each separate PDB, inactive for whether to add inactive entries to PDBs (denoted by positions $(0,0,0)$), name for the base name of the written PDBs and PSFs, pbc for mapping clusters back into the periodic box by their center of mass, and unwrap to toggle unwrapping of clusters.
references	references	Provide an alternative name for the references file; the extension $.\mathtt{csv}$ is implied.
region_epsilon	0.1	Set epsilon for excluded regions when importing structures; the given value is expected to be positive; other are ignored.
region_sigma	1	Set sigma for excluded regions when importing structures; the given value is expected to be positive; other are ignored.
replace	false	Replace or overwrite all written script files.
replica	-	Create copies of existing nonbond interactions; format is destination:source:offset, where offset defines the DPD offset of the new interaction constant compared to the original.

restart	false,	Create LAMMPS restart scripts in chemistry file mode; this option allows setting an alternative data directory using in job run scripts, from which the most recent restart file will be used as a starting point; job run scripts will always try to restart—also when this option is set to false—from the specified data directory, creating a new serial directory by adding 1 to the current highest serial; serial numbers start at 00 and currently have a maximum of 99.
rlength	-	Provide a reference length; used when selecting non-dimensional force fields.
rmass	-	Provide a reference mass; used when selecting non-dimensional force fields.
rmax	-1	Set maximum build cutoff; only applicable to field dpd and gauss, for which defaults are 1 and 1.5 respectively.
rtype	-	Provide a reference type; used when selecting non-dimensional force fields.
sample	energy=false, gyration=false, msd=false, pressure=true, volume=false, green-kubo=false	Set sampling sections in LAMMPS input script for averaging of energy for itemized energetic contributions, gyration for radii of gyration distributions, msd for mean square displacements, pressure for the pressure tensor, volume for the volume tensor, or green-kubo for Green-Kubo output concerning correlations between the off-diagonal pressure tensor contributions as used in viscosity calculations; output files will have their respective identifier as extension; options are either true or false, expect for keyword msd which has average as additional option for time averaged mean square displacement.
seed	-1	Provide initial random seed; a -1 seed will invoke the use of the number of seconds since January 1,

1970 as an initial seed.

shake		Either switch off use of shake for select force fields or set masses, types, bonds, and/or angles for which to apply the SHAKE algorithm to; allowed keywords are active, mass, type, bond, angle; contributing types are separated by a colon ':', e.g. to freeze the angle of TraPPE water in LAMMPS use bond=hw:ow, angle=hw:ow:hw; see 'fix shake' in LAMMPS manual for shake interpretation; shake additions are only added when setting active=true
shake_ iterations	20	Set the maximum number of iterations used during SHAKE.
shake_output	never	Set the output frequency with which SHAKE statistics are written to the LAMMPS log file; options are integer numbers larger than 0 and never, which is equivalent to numbers smaller than 1.
shake_tolerance	0.0001	Set the SHAKE tolerance, which defines a successful SHAKE.
shape	1	Provide the desired shape of the simulation box; the shape factor refers to the relative fration of lx/ly.
shear	false	Add shear paragraph to LAMMPS input script; options are true or false.
split	phase=1, fraction=0.5, thickness=1, type=relative, mode=random, sites=all, groups=all, clusters=all	Defines the selection criteria for applying a fractional split of clusters at the edges of the box; phase sets the phase to which to apply the split; fraction specifies the fractional split; thickness sets the thickness of the region taken into consideration; type sets the units of the thickness (valid options are absolute and relative); mode specifies the selections algorithm (valid options are distance and random); sites sets the sites to include in the selection (valid

options are all for all available sites or a selected list of site types separated by colons); groups sets the groups to include in the selection (valid options are all for all available sites or a selected list of group ids separated by colons); sites sets the sites to include in the selection (valid options are all for all available clusters build in the set phase or a selected

list of cluster ids separated by colons).

suffix	_\$HOSTNAME	Set EMC and LAMMPS suffix.
system	<pre>charge=true, geometry=true, map=true, pbc=true, id=main</pre>	Perform various system checks or set system id; charge checks if the total system charge equals to zero; geometry checks if already exisiting clusters span newly created volume upon adding of a new phase; map allows for mapping box geometry to its minimum image; pbc applies periodic boundary conditions to existing molecules when changing box shape; valid options are true and false; id sets the id used for the targetted system.
system_charge	true	Check if the total system charge equals to zero; valid options are true and false.
system_id	main	Set the EMC system ID; a string is expected.
system_geometry	true	Checks if already exisiting clusters span newly created volume upon adding of a new phase; valid options are true and false.
system_map	true	Map the active system box to its minimum shape; valid options are true and false.
system_pbc	true	Apply periodic boundary conditions after building; valid options are true and false.
temperature	300 or 1	Provide the simulation temperature; atomistic and dimensionless force fields use $300~\mathrm{K}$ and 1 respectively as default.
tequil	1000	Set the LAMMPS equilibration time
tfreq	10	Number of time steps skipped before adding a configuration to an average as sampled in LAMMPS input scripts.
tighten	false	Set margin for tightening the simulation box when importing structures; options are false or a value to represent the desired margin, e.g. 3 Angstroms for atomistic systems; note, that tightening only occurs in the chosen build direction (see direction option).
timestep		Provides the LAMMPS MD time step.

torsion	-	Set force field handling for torsion typing; valid options are ignore, complete, warn, empty, and error.
triclinic	false	Set LAMMPS triclinic mode; needed for simulation of triclinic boxes; valid options are true and false.
trun	10000000	Set LAMMPS run time; setting trun to – avoids its addition to job run scripts, thus not overriding different settings in subsequent chemistry files.
units	-	Set type of units; valid options are reduced, real, and si; alternatively, the outdated option 1j can be used instead of reduced.
units_energy	-	Set units for energetic scale.
units_length	-	Set units for length scale.
volume	false	Assume volume fractions in chemistry.csv file; options are true or false.
warn	true	Control output of warning information; options are true or false.
weight	bond=0.0001, focus=1, nonbond=0.0001	User provided energetic weight for building procedure controlling nonbonded, bonded, and focussed interactions; the latter applies to imported structures.
width	false	Sets double width for generated output scripts; options are true for 160 characters or false for 80 characters.
workdir	-	Set the work directory, which is used as a base for creating the EMC work environment.

4.6 File Formats

All entries within files are pasted into the resulting EMC scripts and force field files. Use of a chemistry file is mandatory. Use of references and parameters files is optional. All entries are assumed to be comma separated unless otherwise stated. All ITEM identifiers can be followed by an optional comment keyword, which can be set to either true or false.

ITEM COMMAND [comment=[true|false]]

The full ITEM paragraph will be ignored until the next ITEM END, when keyword comment is set to true.

4.6.1 Environment File

The environment file functions as a wrap-around to chemistry files. It allows for setting up multiple simulations within one file, including looping over variables, thus enabling setting up series of simulations scanning parameter space. The environment format allows for defining parameter scans and simulation definition in one succinct file, thus furthering compact overviews. The keyword ITEM is required to precede the identifiers listed in the following table. The identifiers are listed in the preferred order in which the are to appear in the environment script.

Identifier	Description
------------	-------------

ANALYSIS

Controls the type of analysis performed during execution of analysis scripts as generated in directory analyze; this paragraph allows to include predefined or user-defined analysis scripts into a project's analysis; sample scripts can be found in \${EMC_ROOT}/scripts/analyze; current supported types are cavity, density, energy, green-kubo, last, pressure, and volume; option active controls inclusion of the analysis type; valid values are true or false; options archive, dir, skip, and window are automatically passed to the analysis script, bearing internal settings; these options can be overridden, but is not advised; options and values are separated by an equal sign (=); different options are separated by a comma, tab or space; location of scripts is looked for in \${EMC_ROOT}/scripts/analyze/, chemistry/analyze, or the path associated with the indicated script

CLUSTERS

Sets general cluster definitions which replace the <code>Q{CLUSTERS}</code> reference in the chemistry template in the same <code>STAGE</code> and <code>TRIAL</code> section; <code>CLUSTERS</code> contents is stored in <code>chemistry/clusters/\$stage/trial.dat</code>

COMMENTS

Sets the start of a comments section, which can only be terminated by a matching END

END

Marks the end of a paragraph

ENVIRONMENT

Sets optional variables related to the environment (formerly OPTIONS); environment options are defined previously (see \(\)undefined \(\) [Environment Options], page \(\)undefined \(\))

FIELD

Allows for a direct definition of force field parameters within the environment concept; the resulting .prm parameter file and .top topology file are stored in chemistry/field/\$stage/project.{prm|top} and should be referred to by using this location (see Section 4.6.3 [Field File], page 46)

GROUPS

Sets general group definitions which replace the @{GROUPS} reference in the chemistry template in the same STAGE and TRIAL section; GROUPS contents is stored in chemistry/groups/\$stage/trial.dat

INCLUDE

Specifies a files to be included containing a subset of commands; the file name follows INCLUDE directly; INCLUDE is a single line item and is not closed with an END

LOOPS

Lists variables over which to loop; variables are expected to be lower case; variables stage, trial and copy are reserved; variables can be paired or coupled by adding:p behind the variable, which couples the current variable to the previous; add:h in order to hide or exclude the variable from the data directory structure; an:h automatically implies the variable is paired with its predecessor; add:d to the predecessor when double occurences are intended (useful in case of pairing); alternatively, permutations in variables can be accomplished by adding:2 for pairs,:3 for triplets, and:4 for quadruplets; focus on on component of the produced list can be obtained by adding a second colon followed by a number, e.g. name:2:1 would create all permutations for pairs which include element 1 for variable name; element counting starts at 0; later reference to any and all variables recorded in LOOPS are to be preceeded with @, enclosed within curly brackets {}, and written in all-caps, e.g. @{NAME} refers to variable name; these references can also be used in subsequent loop variables

PARAMETERS

Allows for direct inclusion of DPD parameters; the resulting parameter file is stored under chemistry/field/\$stage/parameters.csv (see Section 4.6.5 [Parameters File], page 51)

POLYMERS

Sets general polymer definitions which replace the <code>@{POLYMERS}</code> reference in the chemistry template in the same <code>STAGE</code> and <code>TRIAL</code> section; <code>POLYMERS</code> contents is stored in <code>chemistry/polymers/\$stage/trial.dat</code>

REFERENCES

Allows for direct inclusion of DPD references; the resulting reference file is stored under chemistry/field/\$stage/references.csv (see Section 4.6.4 [References File], page 50)

SHORTHAND Sets general shorthand definitions which replace the @{SHORTHAND} ref-

erence in the chemistry template in the same STAGE and TRIAL section; CLUSTERS contents is stored in chemistry/shorthand/\$stage/trial.dat

STRUCTURES Defines a list of structures, relating directly to trial loop entries; structures

are referred to by <code>@{STRUCTURE}</code>

STAGE Indicates the start of a section relating to loop variable stage; STAGE is

followed by a stage indicator, which should appear in the enumerated loop variable stage; all following paragraphs relate to the indicated stage; STAGE is a single line item and is not closed with an END; all identifiers following

STAGE refer to loop variable stage

TEMPLATE Defines a chemistry template; templates are stored in

chemistry/stages/stage.esh; templates are used to create
complete chemistry files upon execution of the associated run script;
references to loop variables start with Q{} and are written in all caps;

references are treated as environment variables

TRIAL Indicates the start of a section relating to loop variable trial; TRIAL is a

single line item and is not closed with an END; all identifiers following TRIAL

refer to loop variable trial

VARIABLES Lists extra environment variables; environment variables are referred to with

an at symbol @ and are enclosed by curly brackets {}; reserved variables are @{EMCROOT} for the root location of EMC and @{WORKDIR} for the directory

in which the collection of simulations takes place

WRITE Specify a message line to be written to the output; the written text directly

follows WRITE; WRITE is a single line item and is not closed with an END

EMC setup only assumes environment modus, when the environment variable is set to true in the first occurring OPTIONS paragraph. Environment mode allows for optionally setting names associated with build, analyze, and run scripts, as well as defining which queues to be used. Dashes imply undefining parameters. The ncores parameter is mandatory. For certain problems, it can be desired to simulate multiple indepedent structures in order to obtain a correct statistical sample. To this end, a loop variable copy is available, which creates multiple copies with the same conditions, but executing building with different random seeds, thus creating indepedent initial structures. Note, however, that use of this features quickly creates many simulations, which harbors the danger of overcrowding high-performance computing queueing systems and over-requesting available computational resources. An example for environment mode application can be found in \${EMC_ROOT}/examples/setup/environment/shear.

4.6.2 Chemistry File

The chemistry file – with default file name chemistry.esh – supports environment, newstyle, and legacy formats. Columns can only be separated by commas or tabs. Next line extension occurs by using an ampersand & at the end of to be extended line.

4.6.2.1 General

The new-style format allows for the definition of polymers, which uses keyword ITEM followed by an identifier to distinguish between different contributing paragraphs, as described by

Identifier	Description
CLUSTERS	Sets cluster definitions, using previously defined cluster groups
COMMENTS	Sets the start of a comments section, which can only be terminated by a matching ${\tt END}$
EMC	Defines a section to be copied verbatim into the EMC build script; optionally, the keyword can be followed by either one or two numeric values, identifying which phase and where the block verbatim text is to be added to; alternatively, identifiers phase and spot can be used, followed by an equal sign (=); valid values are the respective phase for identifier phase and a number between 0 and 2 for spot to place the defined section at specific spots in the selected phase section in the EMC build script; defaults for phase and spot are 1 and 0 respectively when value and identifiers are omitted;
END	Marks the end of a paragraph
FIELD	Allows for a direct definition of force field parameters; the resulting .prm parameter file and .top topology file are stored in the current directory (see Section 4.6.3 [Field File], page 46)
GROUPS	Sets separate group definitions, including polymeric repeat units
INCLUDE	Specifies a files to be included containing a subset of commands; the file name follows INCLUDE directly; INCLUDE is a single line item and is not closed with an END

LAMMPS

Defines a section to be copied verbatim into the LAMMPS execution script, appearing atop the simulation section; alternatively, identifiers stage and spot can be used, followed by an equal sign (=); allowed options for stage are – in order of appearance in the LAMMPS execution script – header, variables, interaction, equilibration, simulation, integrator, sampling, intermediate, run; allowed options for spot are head and tail; stage options correspond to the comments in the resulting LAMMPS execution script; defined additional variables as described above can be used when captured between \${..}.

OPTIONS

Allows for setting all command line options of $emc_setup.pl$ inside chemistry files, thus creating the option of consolidating a simulation setup within one file only; see the previous paragraph; syntax is option, value[,...], where option refers to any of the options as defined previously (see $\langle undefined \rangle$ [Chemistry Options], page $\langle undefined \rangle$)

POLYMERS

Defines polymers, using previously defined polymeric groups; this paragraph can be used for defining polymers either through the GROUPS or CLUSTERS paragraph

PARAMETERS.

Allows for direct inclusion of DPD parameters; parameters are not stored but used internally (see Section 4.6.5 [Parameters File], page 51)

PROFILES

Defines additional mass profiles, where each line generates an additional profile; syntax is name, mode[:type[:binsize]], contributor[,...]; mode can be either cluster or type; type sets the type of profile, which can either be density, density3d, or pressure; density is assumed when omitted; binsize sets the individual bin size; the general binsize is assumed when omitted; contributor refers to either a cluster name or type repectively; note, that profile coordinates are in reduced units and run from 0 to 1

REFERENCES

Allows for direct inclusion of DPD references; references are not stored but used internally (see Section 4.6.4 [References File], page 50)

SHORTHAND

Legacy format, allowing for group and cluster definitions in shorthand notation; polymeric definitions are not possible within this format

VARIABLES

Defines additional variables, which are either used by read EMC structures from the EMC structure library (\$EMC_ROOT/lib), serve as pass-through to subsequent simulation packages (e.g. LAMMPS), or are used in the generated EMC build script; the section identifier VARIABLES can be followed by keywords head for appearance at the top or tail for appearance at the bottom of the EMC script variable block

WRITE

Specify a message line to be written to the output; the written text directly follows WRITE; WRITE is a single line item and is not closed with an END

4.6.2.2 Shorthand

The SHORTHAND format can be used without a line starting with keyword ITEM, provided its lines are at the start of the chemistry file. Conversely, the shorthand format also provides quick definition of non-polymeric molecules without the need of defining groups and clusters. Its format is given by the following table.

Entry	Description
1	Cluster and Sets the group name, which optionally can be followed by f[ield]=id[,], m[ass]=#, or t[erm] – separated by colons – where characters between [] are optional and can be omitted; multiple options can be set in one line; f[ield]=id sets a specific field id when using multiple fields for typing, e.g for CHARMM (see Section 5.11 [Field], page 103), where id can be part of the full identifier; m[ass]=# overrides automated setting of group mass, where # represents the desired mass; t[erm] marks group as a terminator (see Section 5.17 [Groups], page 114)
2	Chemical representation in the form of a SMILES string (see Section 7.8 [SMILES], page 194)
3	Fraction or count; can be either molecular, mass, or volume fraction, or the number of clusters based provided option (see \(\sqrt{undefined}\)\) [Chemistry Options], page \(\sqrt{undefined}\) for options mass, mol, or number)
4	Molecular mass in g/mol
5	Molecular volume in cc/mol

Column 4 and 5 are optional.

4.6.2.3 Groups

The GROUPS paragraph defines multiple groups, including polymeric groups. Its format is given by the following table.

Entry Description

Sets the group name, which optionally can be followed by f[ield]=id[,...], m[ass]=#, or t[erm] - separated by colons - where characters between [] are optional and can be omitted; multiple options can be set in one line; f[ield]=id sets a specific field id when using multiple fields for typing, e.g for CHARMM (see Section 5.11 [Field], page 103), where id can be part of the full identifier; m[ass]=# overrides automated setting of group mass, where # represents the desired mass; t[erm] marks group as a terminator (see Section 5.17 [Groups], page 114); note, that groups with only one connection are automatically terminators

- Sets the group chemistry using SMILES strings (see Section 7.8 [SMILES], page 194) or polymer type to turn on polymer mode; in SMILES polymeric connectivity points are marked with *; valid polymer types are alternate, block, and random; column 3 and onwards are omitted when using groups in polymer mode; the polymer distribution is defined through the POLYMERS paragraph
- 3 Defines connection index for polymeric groups of current group; indices refer to * occurrences in the previously set SMILES string
- 4 Defines the connecting group name and corresponding index in the format name:index
- 5- Repeats of column 3-4 for subsequent connections

4.6.2.4 Clusters

The CLUSTERS paragraph defines multiple clusters. Options following keyword import in entry 2 can be used in any order when superceeded by their respective identifier and separated by an = sign. Column 4 and onward are optional and have different meaning depending on the mode in which the cluster option is called. In case of polymeric definitions, column 4 through 5 are used. In case of imported structures, all columns 4 and onward can be used. Identifiers instead of columns can be used when importing structures. Each identifier is described in the paragraph and table hereafter. The use of commas or tab as option separators between entries is advised when defining an import. Its format is given by the following table, where the identifier column refers to the next table.

Entry	Identifier	Description
1	-	Sets the cluster name
2	-	Sets the associated group, the type of polymer — indicated by alternate, block, or random — or whether to expect a imported structure from file as incidated by keyword import; consequently, group names are not allowed to have reserved names alternate, block, random, or import
3	- or ncells	Sets a fraction or count, which can be either a mol, mass, or volume fraction, or number of clusters, based provided command line option (see \(\text{undefined} \) [Chemistry Options], page \(\text{undefined} \), for options mass, mol, or number); in case of keyword import in entry 2, this entry contains the number of repeat units (id ncells)
4	- or name	Molecular mass in g/mol or file name (id name) in case of keyword import in entry 2

5	- or mode	Molecular volume in cc/mol or import mode (id \mathtt{mode}) when selecting keyword \mathtt{import} in entry 2
6	type	Optional entry for import type
7	flag	Optional entry for setting mobility flags of imported structures
8	density	Optional entry for setting the density treatment of imported structures
9	focus	Optional entry for marking the imported structure as focus
10	tighten	Optional entry to trim or tighten imported structures
11	ntrial	Optional entry for setting the number of trials
12	periodic	Optional entry for setting periodicity; currently not implemented
13	field	Optional entry for enforcing a specific force field for force field typing of the imported structure;
14	exclude	Optional entry for setting an exclusion region
15	depth	Optional entry for setting ring recognition recursive depth
16	percolate	Optional entry for identifying a percolating crystalline structure
17	unwrap	Optional entry for unwrapping imported structures
18	guess	Optional entry for guessing unwrap status based on input
19	charges	Optional entry for using charges from imported morphologies
20	formal	Optional entry for using formal charges from imported morphologies when available
21	translate	Optional entry for translating the all imported sites for a distance into the growth direction of the simulation box
22	map	Optional entry for mapping both sites and box shape unto their periodic minimum image

The following table describes import identifiers, which can be used without having to abide by strict column sequences. Identifiers and their values are separated by equal signs (=), omitting any spaces. Examples are mentioned after this table.

Identifier	Default	Description	
charges	false	Use charges from imported structures; allowed values ar true or false	
density	mass	Sets density treatment of imported structures, which is needed for calculating correct box dimensions when adding material; allowed values are mass or number	
depth	auto	Sets the maximum size of rings to be recognized; the provided input should be a definite positive integer or keyword auto	
exclude	true	Toggles whether to add an repulsive exclusion region around the imported structure; allowed values ar true or false	
field	-	Enforces a specific force field for force field typing of the imported structure	
flag	rigid	Sets mobility flag for imported structures; allowed flags are fixed, rigid, or mobile	
focus	true	Toggles whether to focus on the imported structure, which excludes its volume for additionally built molecules; allowed values are true or false	
formal	true	Include formal charges from imported morphologies when provided in input structure; allowed values are true or false	
guess	auto	Sets optional guessing of unwrapping based on imported structures; only functions when unwrap is set to auto; allowed values are auto, true, or false	
map	false	Map both sites and box geometry onto their periodic minimum image; allowed values are true or false	
mode	emc	Optionally enforce import mode when not using name extensions in identifier name; allowed modes are emc for EMC files, pdb for PDB files – providing both .pdb and .psf files, or insight for Insight II files – providing both .car and .mdf files	
name	-	Sets file name; allowed file name extensions are .emc for EMC files, .car or .mdf for InsightII files, and .pdb or .psf for PDB files; default is .emc; extensions override mode defaults	

ncells	1:auto:auto	Sets the number of repeated unit cells along the direction as set by option direction in the options paragraph (see \langle undefined \rangle [Chemistry Options], page \langle undefined \rangle); repeats in multiple directions can be set when dividing input by colons in order of x , y , and z direction respectively; default is 1 in the main direction and auto in the remaining directions, e.g. auto:1:auto for main direction y
ntrials	10000	Sets the number of trial iterations used for determining the volume of an import of type=line or structure (see column 6), which employs a Monte Carlo algorithm; option is a number larger than zero
percolate	auto	Sets whether imported structure reflects a percolating crystal; allowed values are auto, true, or false
periodic	-	Sets periodicity; currently not implemented
tighten	-	Set the tightening distance, which effectively shrink-wraps the box around what is imported; allowed option is a distance; default is a distance of 1 or 3 for coarse grained or atomistic force fields respectively
translate -		Translate by a length all sites into the direction of box growth; the entered value represents a scalar, which internally is multiplied

Translate by a length all sites into the direction of box growth; the entered value represents a scalar, which internally is multiplied by the unit box vector in the direction of growth; by default no translation is performed

type surface

Sets the imported structure type; allowed types are crystal when adding no additional sites, surface when adding sites in one direction as set by option direction in the options paragraph (see \(\text{undefined} \) [Chemistry Options], page \(\text{undefined} \)), tube when adding in two directions, structure when adding in three directions, or system when importing a previously built system

unwrap false

Sets whether to unwrap the imported structure; allowed values are auto, true, or false

Columns 4 is optional when defining molecules, but mandatory when importing structures. Importing options defined in columns 3 through 14 can be preceded by their respective identifiers, i.e. ncells, name, mode, type, flag, density, focus, tighten, and ntrials, separated by an equal sign. The order as given in the table above is assumed when omitting this identifier. The specified default is taken, when the an option is omitted. A formal example of an import line is given by

surface import, name=\$root+"lib/fcc", ncells=8, mode=emc, &

type=surface

which creates a surface with cluster name **surface**, using a nonbonded fcc lattice from the EMC library, consisting of 8 repeat units in the x-direction. Note, that any given order can be used, when identifiers are specified. A shorter, but equivalent definition is given by

surface import 8 \$root+"lib/fcc" emc

where now the option order matters, therefore creating the need for strictly following the column order as given in the above table. Alternatively, the import mode can be determined by EMC setup through adding the file extension, i.e.

surface import name=\$root+"lib/fcc.emc" ncells=8 type=surface

or

surface import 8 \$root+"lib/fcc.emc"

Note, that a space or comma can also be used as separator. However, this option is not advised when using a space or comma in the file name. Currently, only one structure at a time can be imported by EMC.

4.6.2.5 Polymers

In the case of a polymer, the cluster name has to appear in the POLYMERS paragraph, when defining the type of polymer in column 2. Otherwise, a group name is assumed. The POLYMERS paragraph describes the composition of one polymer at a time. Each polymer definition starts with the polymer's cluster name on a separate line. The polymer name optionally can be followed by keywords fraction, niterations, or order, separated from their value by an = sign. Defining any of these keywords overrides global definitions for the indicated polymer only (see keyword polymer under OPTIONS for globals (see \(\) undefined \(\) [Chemistry Options], page \(\) undefined \(\))). Valid fraction options are number for using number of molecules and mass molecule mass as distribution entries. Valid order options are list for sequentially and random of randomly interpreting the provided polymer distribution. The niterations keyword sets the maximum allowed number of iterations used during polymer sequence determination. Lines following the polymer name define the polymer's distribution of contributing groups. Multiple polymer definitions can be privided within this paragraph. The format of contributing groups is given by the following table.

Entry Description

Mol or mass fraction of the polymer cluster defined by this line; mass fractions are only allowed when defining polymers through groups

- 2 Group name; multiple group names can be given through separation with a ':'; weights with which groups are randomly picked follow an '=' and are separated by a ':', e.g. A:B=1:2; equal weights are assumed when omitted
- 3 Number of repeat units for the above group
- 4- Repeats of columns 2-3 for subsequent groups

Capping or end groups should be included in the above list for mass calculation purposes, although they are neglected by EMC. EMC selects end groups internally.

4.6.2.6 DPD Additions

The following identifiers are valid only, when selecting DPD as force field:

Identifier	Description
ANGLES	Defines a paragraph for additional specific angle entries, which should be provided in the order of type1, type2, type3, k, theta
BONDS	Defines a paragraph for additional specific bond entries, which should be provided in the order of $type1$, $type2$, k , l
MASSES	Defines or — in case of replicas — redefines masses, which should be provided in the order of type, mass; mass is a definite positive number and has reduced units when no reference type has been chosen; otherwise, mass is in units of the reference type (see Section 4.4 [Options], page 12)
NONBONDS	Defines a paragraph for nonbond entries, which should be provided in the order of type1, type2, a[, cutoff[, gamma]]; these entries overwrite already exisiting entries
REPLICAS	Creates duplicates of exisiting nonbond entries, which should be provided in the order of target[:factor], source[:fraction[:flag]][, source[:fraction[:flag]][,]], offset; interaction constants, mass, cutoff, and gamma are averaged when multiple sources are provided; offset defines the offset of the new DPD interaction constant a_{ij} with respect to the original; factor defines a multiplication factor with respect to the offset; factor is set to 1 when omitted; fraction defines which fraction to use of each contributing component; equal fractions are assumed when omitted; normalization of fraction is controlled by flag, which can either be true or false; by default fraction is normalized; fraction is not normalized when any one flag is set to false
TORSIONS	Defines a paragraph for additional specific torsion entries, which should be provided in the order of type1, type2, type3, type4, k, n, delta, and optionally following triplets of k , n, and delta

Both keyword and identifier must be in all caps and separated by a comma or tab. Currently, alterations or additions to force field files only function for the DPD force field. Types can be replicated in order to either create duplicates or compounded types, where

$$a_{ij}^* = f(a_{ij}^* - 25) + \Delta$$

expresses the new interaction parameter a_{ij}^* . Assume, that a new type AIR is to be created based on an alkane representation C4, where multiplication factor f=1.5 and offset $\Delta=25$. This would result in the following REPLICAS entry

For each of the parameter additions (i.e. nonbonds, bonds, angles, or torsions), entries containing a wildcard * will appear in the AUTO paragraph of the resulting force field file.

4.6.3 Field File

The field file — using the extension .define — allows for defining force field parameters and rules within one file. Translation of this file into an EMC .prm parameter file and .top topology file occurs by applying field.pl, which can be found in the EMC script directory \${EMC_ROOT}/scripts. This format can also be included in environment and chemistry files by adding the field definition between identifiers FIELD and END, both preceded by keyword ITEM.

4.6.3.1 General

The field format itself also uses keyword ITEM to identify subsequent main functionality paragraphs, as decribed by

Identifier	Description
DEFINE	Defines different force field settings as described below; this paragraph is mandatory
MASSES	Defines the types and their associated masses and subsequent definitions; the syntax is type, mass, element, number of connections, formal charge, comment; this paragraph is mandatory
COMMENTS	Allows for inclusion of comments specific to the force field's derivation; this paragraph is optional
REFERENCES	Reports the literature references used to construct the force field; the syntax is year, volume, page, journal; this paragraph is optional
PRECEDENCE	Defines the precedence table as described below; this paragraph is optional
EQUIVALENCE	Describes the equivalences of the force field; equivalences state which type can be used instead of the official type for abstracting the asociated force field parameters; the syntax is type, pair, bond, angle, torsion, improper; this paragraph is optional
NONBOND	Defines the nonbond parameters of the force field; the syntax starts with type1, type2, followed by the actual parameters (e.g. epsilon, sigma for Lennard-Jones force fields, and a, cutoff, gamme for a DPD force field); the expected number of parameters is dependent on the chosen force field mode; this paragraph is mandatory
BOND	Defines the parameters for bond length interactions; the syntax is $type1$, $type2$, k , $l0$; the interaction function depends on the the force field mode, but is in general harmonic in nature; this paragraph is optional

ANGLE	Defines the parameters for bond angle interactions; the syntax is type1, type2, type3 k, theta0; the interaction function depends on the the force field mode, but is in general harmonic in nature; this paragraph is optional
TORSION	Defines the parameters for dihedral interactions; the syntax is type1, type2, type3, type4, k, n, delta[, k, n, delta]; the allowed number of additional parameter entries depends on the chosen force field mode; the interaction function depends on the the force field mode, but is in general a Fourier series of cosines (e.g. see Section 7.12.17 [Standard], page 320, Torsion paragraph); this paragraph is optional
IMPROPER	Defines improper interactions; the syntax is $type1$, $type2$, $type3$, $type4$, k , $psi0$; the interaction function depends on the the force field mode, but is in general harmonic in nature; this paragraph is optional
RULES	Defines the rules associated with the types as defined by the MASSES paragraph; the syntax is type, partial charge, rule[, rule]; one type can have multiple rules; a rule describes a unique chemical environment, which defines the type at hand; this paragraph is optional
TEMPLATES	Defines templates, which can be used as aliases when defining group chemistry; this paragraph is optional

Each entry can be separated by either a TAB, COMMA, or SPACE. A TAB represents the preferred separator. Wildcards as represented by * can be used for types in parameter definitions. Partial wildcards are also allowed. The partial wild cards can be understood by the following example: assume c4 generally describes an SP3 carbon with no hydrogen attached, c4h with one hydrogen attached, and c4h2 with two hydrogens attached, then wildcard c4* would describe all three of these types.

4.6.3.2 Define

Specific alternative settings can be defined for force field by using the DEFINE paragraph, as decribed by the following

Identifier	Description
ANGLE	Sets the error handling for typing of angle interactions; options are <code>ignore</code> , <code>complete</code> , <code>warn</code> , <code>empty</code> , and <code>error</code>
CREATED	Sets the creation date of the force field
CUTOFF	Sets the cut off for all nonbonded interactions

TORSION

VERSION

DENSITY Sets the unit of length as used by the parameters in the force field; options are g/cc, kg/m^3, and reduced; both LENGTH and ENERGY are set to reduced when the latter option is chosen, thus resulting in a force field in reduced units Sets the unit of length as used by the parameters in the force field; options **ENERGY** are j/mol, kj/mol, cal/mol, kcal/mol, kelvin, and reduced; both LENGTH and DENSITY are set to reduced when the latter option is chosen, thus resulting in a force field in reduced units **FFDEPTH** Sets the maximum recursive with which to trace the chemical surrounding of a site during typing; this depth should equal the maximum depth of the provided rules **FFMODE** Sets the name or mode of the force field; options are born, charmm, dpd, martini, mie, opls, sdk, standard, and trappe **FFTYPE** Sets the force field type; options are atomistic, united, and coarse **IMPROP** Sets the error handling for typing of improper interactions; options are ignore, complete, warn, empty, and error INNER Sets the inner cut off when using a force field with a switching function for nonboned interactions (e.g. as in the CHARMM force field (see Section 7.12.7 [CHARMM], page 233)) LENGTH Sets the unit of length as used by the parameters in the force field; options are angstrom, nanometer, micrometer, meter, and reduced; both DENSITY and ENERGY are set to reduced when the latter option is chosen, thus resulting in a force field in reduced units MTX Sets the mixing rule; options are none, bethelot, arithmetic, geometric, and sixth_power **NBONDED** Sets the number of bonded atoms to exclude from nonbonded interactions; the number varies by force field Sets the error handling for typing of 1-4 pair interactions; options are off, PAIR14 false, exclude, on, true, and include

Sets the error handling for typing of dihedral interactions; options are

ignore, complete, warn, empty, and error

Sets the force field version

Error handling, as mentioned in the latter options of the above table, can be dealt with in a number of ways. Any errors can be either fully ignored without any output by EMC during typing by chosing option ignore. A warning will be generated, when chosing option warn. An empty parameter entry will be generated without any output to screen when chosing option empty. EMC will continue after all of these options. However, EMC will generate warnings for all missing parameters and will cease execution when chosing options error. Examples of .define files can be found for several force fields as provided in the \${EMC_ROOT}/field/ force field directory (e.g. the CHARMM, OPLS, and TraPPE force fields).

4.6.4 References File

Reference files are optional and are used to create comprehensive EMC force field files (using .prm extensions). Currently only DPD force fields are used when interpreting the reference file. Its default file name is references.csv.

Column	Description
1	Short ID as referred to in parameters.csv
2	ID as represented in produced .prm force field files
3	Bead mass in g/mol
4	Bead volume in nm^3
5	Number of connections that this bead can have
6	Effective charge of bead
7	Number of repeat units represented in this bead (e.g. 3 when 3 ethyleneoxide monomers are captured by one bead)
8	Comment describing the origin of the bead

Future emc_setup.pl versions are intended to span a wider choice of force fields when using reference files.

4.6.5 Parameters File

Parameter files are optional and are used to provide parameters. Currently they are only used in case of the DPD force field (see Section 7.12.11 [DPD], page 272). Its default file name is parameters.csv.

Column	Line	Description
1	1 2-	Mandatory empty field Alphanumeric type name of bead A (short)
2	1 2-	Mandatory empty field Alphanumeric type name of bead B (short)
3	1 2-	Temperature at which the interaction parameters in this column were determined. Numerical value refering to the interaction of bead A and B
		Identical to column 3.

Column 4 and subsequent columns can hold parameters determined at different temperatures. Their syntax is the same as for entries in column 3. Future emc_setup.pl versions are intended to include a wider choice of force field when using parameter files.

4.7 Examples

The directory \${EMC_ROOT}/examples/setup/ holds a number of examples with different complexity, all using emc_setup.pl as a base for creating EMC build and LAMMPS input scripts. The examples are subdivided in chemistry and environment related examples. This section describes ways to build bulk systems with a mixture of chemicals, typing of various force fields, construction of various types of polymers, building multiphase simulations, and ways to build material between two surfaces. A second set of examples in \${EMC_ROOT}/examples/build/ illustrate adaptation of EMC scripts directly for specific solutions. These scripts are not discussed here.

4.7.1 References

- 1. VMD Visual Molecular Dynamics, 'http://www.ks.uiuc.edu/Research/vmd/'
- 2. Humphrey, W., Dalke, A. and Schulten, K., "VMD Visual Molecular Dynamics", J. Molec. Graphics 1996, 14, 33-38
- 3. LAMMPS Molecular Dynamics Simulator, 'http://lammps.sandia.gov/'
- 4. Plimpton, S., "Fast Parallel Algorithms for Short-Range Molecular Dynamics", *J. Comput. Phys.* **1995**, *117*, 1-19

4.7.2 Chemistry Mode

Chemistry mode examples show how to set up single system simulations using a number of different common problems, which include setting up bulk systems, systems with different force fields, polymer systems, systems with multiple phases, and systems with surfaces. Chemistry examples can be found in \${EMC_ROOT}/examples/setup/chemistry/.

4.7.2.1 Bulk Mixture

Assume the desire exists to simulate a system consisting of 45% water, 40% alcohol, 5% salt, and 10% sugar by mass, where the force field of choice is PCFF and consists of about 2000 particles once built. The setup utility offers multiple ways for setting up a simulation. The quickest way is to use emc_setup.pl's legacy format. For this format, one first creates a chemistry.esh file with the following content:

Note, that the fraction indications -45, 40, 5, and 10 – in the last column are normalized internally, and can therefore be any number. Here, percentages are taken for illustrating purposes. The next step is to invoke emc_setup.pl,

```
emc_setup.pl -field=pcff -ntotal=2000 -mass -build_dir=. bulk
```

where -field selects the force field, -ntotal sets the number of particles, -mass assumes mass fractions in the chemistry.esh file, and -build_dir sets the origin of the created files to the current directory. An alternative chemistry file with the name bulk.esh uses shorthand notation in combination with options. Here all command line options are folded into the chemistry file, which reads as follows,

```
#!/usr/bin/env emc_setup.pl
```

Options

ITEM	OPTIONS
replace	true
mass	true
ntotal	2000
field	pcff
density	1
build_dir	•
ITEM	END

Shorthand

ITEM	SHORTHAND
water	0,45
alcohol	CCO,40
salt	[Na+].[Cl-],5

sugar OCC10C(0)C(0)C(0)C10,10

ITEM END

A more general way of describing the same simulations is by the use of GROUPS and CLUSTERS. EMC internally uses sites, groups, and clusters to represent atoms, repeat units, and molecules respectively. Using groups in case of small molecules makes less sense than using groups for polymers. However, as a general notation, this problem can also be defined by the following

#!/usr/bin/env emc_setup.pl

Options

ITEM	OPTIONS
replace	true
mass	true
ntotal	2000
field	pcff
density	1

ITEM END

Groups

build_dir

ITEM GROUPS

water 0 alcohol CCO

salt [Na+].[Cl-]

glucose OCC10C(0)C(0)C(0)C10

CLUSTERS

ITEM END

Clusters

ITEM

water	water,45
alcohol	alcohol,40
salt	salt,5
sugar	glucose,10

ITEM END

Execution can be performed in one of two ways,

```
emc_setup.pl bulk
```

or, in the case of a Unix-like operating system,

```
./bulk.esh
```

provided bulk.esh is executable and emc_setup.pl can be found in the shell's predefined path. This solution allows for a comprehensive representation of a simulation, without having a number of separate files describing it. Once executed, emc_setup.pl creates the EMC build script build.emc and the LAMMPS input script bulk.in. To build an input structure, the EMC build script needs to be executed using EMC. Assume, that the LINUX version emc_linux is used, which location is in your path variable \$PATH. Then,

```
emc_linux build.emc
```

will create input structures in PDB format – bulk.pdb and bulk.psf – and LAMMPS format – bulk.in. The resulting structure can be inspected by means of VMD (Visual Molecular Dynamics)^{1,2} through

```
vmd -e bulk.vmd
```

assuming, that vmd is in your path. The advantage of using bulk.vmd, is that this file contains the correct Van der Waals radii for the simulated beads. A molecular dynamics simulation with LAMMPS (Large-scale Atomic/Molecular Massively Parallel Simulator)^{3,4} can be started with

```
lmp_linux < bulk.in</pre>
```

assuming, that lmp_linux is in your path. A similar example can be found in the \${EMC_ROOT}/examples/setup/chemistry/bulk directory.

4.7.2.2 Force Fields

EMC allows for the use of various force fields. To illustrate this, five examples are included for commonly used force fields, which are separated into atomistic (CHARMM, OPLS-AA, PCFF), united atom (OPLS-UA, TrAPPE), and coarse grained (DPD) examples. Force field choice is indicated by the -field flag of the emc_setup.pl script, which will try to find a corresponding force field in the \${EMC_ROOT}/field directory. For the latter to work, it is assumed, that emc_setup.pl is located in the \${EMC_ROOT}/scripts directory. With exception of the coarse-grained example, all examples use a molar ratio of 1:4 for a ethanol:water mixture. The corresponding chemistry file chemistry.esh for non coarse-grained examples contains the following lines

ethanol,CCO,20 water,0,80

Fraction indications do not have to reflect percentages: 1 instead of 20, combined with 4 instead of 80 is equally valid, due to internal normalization. For a system with 1000 atoms typed with the CHARMM force field and project name solution, the call to emc_setup.pl is given by

emc_setup.pl -ntotal=1000 -field=charmm -replace solution

where -ntotal indicates the number of atoms, -field specifies the desired force field, and -replace selects overwriting of possibly existing build and input scripts. An alternative to chemistry.esh is presented by solution.esh, in which case emc_setup.pl command line options are integrated into the script by means of an options section. Execution creates an EMC build.emc and a LAMMPS^{3,4} solution.in script. A change of force field and its corresponding typing occurs by altering the -field flag to e.g. -field=trappe, when desiring TraPPE typing. Note, that each force field only includes chemistries, for which it was parameterized. This means, that not all chemistries can be typed by all force fields. Also be aware, that typing is not guaranteed to be correct. It is kindly requested to report bugs, so that improvements to force fields can be included in next versions. Next steps for executing EMC and LAMMPS are identical to the steps as mentioned in the previous section for the bulk mixture example.

4.7.2.3 Record

EMC offers the possibility to follow the building process by creating a set of PDB's and accompanying PSF's during building, which is illustrated by the example found in \${EMC_ROOT}/examples/setup/chemistry/record/. The build of a 60 carbon fullerene has been chosen to illustrate this option. EMC will produce a set of PDB's and PSF's, which are combined by cat.sh. The latter produces one PDB and PSF file, and one VMD script, after which it deletes the PDB's and PSF's produced by EMC. A exec.sh script illustrates the work flow. It uses fullerene.esh as the describing chemistry file, which set the following options,

ITEM	OPTIONS
seed	-1
ntotal	60
density	0.1
temperature	300
grace	0.999
nrelax	200
radius	9.5
center	true
field	opls-ua
record	"record",10,true
replace	true
depth	6
ITEM	END

A seed of -1 indicates for EMC to take the system clock as a seed. The total number of sites is set by ntotal, while the density is chosen to be $0.1\,\mathrm{g/cc}$. The simulation temperature is set at 300 K. The keyword grace is used to allow for overlap during initial insertion, which is corrected during relaxation. The number of relaxation moves after insertion, nrelax, is set to 200, with a site inclusion radius of 0.95 nm. The first inserted site is set at the center of the box by setting keyword center to true. The chosen force field – as set by defining field – is OPLS-UA. Recording is switched on by defining record, where PDB's and PSF's are written out for the inserting of each site. The written out files have root file name "record", where each file contains a trajectory resulting from relaxation. Here, snapshots are written every 10 cycles. Inactive sites with positions at (0,0,0) are included in PDB's. All files are opted to be replaced by setting the replace keyword to true. Ring recognition normally is automatic. However, this would take unnecessarily long for fullerenes due to the recursive nature of the underlying algorithm. EMC allows for reducing the recursion depth of this algorithm. Here, we chose to set the recursion depth to 6, since none of the rings in a fullerene have more than 6 members.

Fullerene chemistry is defined in the shorthand paragraph by

ITEM SHORTHAND

fullerene c12c3c4c5c1c6c7c8c2c9c1c3c2c3c4c4c%10c5c5c6c6c7c7c%11

c8c9c8c9c1c2c1c2c3c4c3c4c%10c5c5c6c6c7c7c%11c8c8c9c1

c1c2c3c2c4c5c6c3c7c8c1c23,1

ITEM END

Intricate connectivity is illustrated by connectivity numbering (see Section 7.8 [SMILES], page 194). The SMILES for this example as well as for many chemistries can be found in Wikipedia.¹

Execution of this example is best performed by workflow script

./exec.sh

which calls emc_setup.pl, followed by EMC. It also concatenates building trajectories as produced by EMC into one set of files, which can be viewed with VMD by

vmd -e record.vmd

assuming VMD is in the user's path.^{2,3}

References

- 1. N.N., "Buckminsterfullerene", Wikipedia, 1/1/2016.
- 2. VMD Visual Molecular Dynamics, 'http://www.ks.uiuc.edu/Research/vmd/'
- 3. Humphrey, W., Dalke, A. and Schulten, K., "VMD Visual Molecular Dynamics", J. Molec. Graphics 1996, 14, 33-38

4.7.2.4 Polymers

Polymers have a wide field of application in chemical industry. EMC offers the possibility of creating a variety polymer classes. Examples can be found in \${EMC_ROOT}/examples/setup/chemistry/polymer. The focus in this section lays on building random, block, and alternate copolymers.

Let us assume, that we would like to study a copolymer consisting of four A beads, four B beads, and two D terminators. We would like to study a coarse-grained model with 1000 beads, using a general DPD force field with a number density of 3.

The OPTIONS section is defined by (see \(\sqrt{undefined}\)\) [Chemistry Options], page \(\sqrt{undefined}\)),

ITEM	OPTIONS
project field auto ntotal density replace build_dir	<pre>dpd dpd/general true 1000 3 true .</pre>
ITEM	END

The first line sets the project name with keyword project, while the second selects the general DPD force field with keyword field. The option auto toggles the automatic addition of masses to the force field mass paragraph, in case masses are not provided. The total number of beads is set with ntotal and the system density with density. The keyword replace allows for existing files to be overwritten. The current directory is chosen as the build directory by keyword build_dir.

The generalized DPD form does not require a parameters.csv or reference.csv file, when not specifically defining pair-wise interactions. With this information, emc_setup.pl will create dpd.prm in the current directory, which is a DPD force field parameter file. This force field file is needed by EMC, in order to be able to type the different beads.

A polymer chemistry file cannot use shorthand notation as discussed in the previous paragraphs. Instead, group, cluster, and polymer specific sections need to be used (see Section 4.6.2 [Chemistry File], page 36).

First, three contributing groups are defined to describe beads A, B, and D,

ITEM	GROUPS
a	*A*,1,a:2,2,b:1
b	*B*,1,b:2
d	*D,1,a:1,1,a:2,1,b:1,1,b:2
ITEM	END

As with the bulk case, SMILES syntax is used to describe group chemistry. Additionally, connectivity between the different monomers needs to be defined. This connectivity is indicates by an asterix *. By definition, a monomer has two or more connections and a terminator only one. With the connection points defined per monomer, we have to still define where these connection points connect to. Let us consider group a as an example. We would like to connect group a to both itself and group b,

A indicates the SMILES representation, followed by connection pairs. The first pair 1,a:2 means, that the first occurring * of *A* connects to the second occurring * of group a. The first pair 2,b:1 means, that the second occurring * of *A* connects to the first occurring * of group b. The same logic applies for subsequent definitions of groups b and d. The setup script adds all redundant connections, i.e. when connecting group a to group b with 2,b:1, a connection on group b reading 1,a:2 is implied and does not have to be defined explicitly. One could define group b as

which would work, but it is not necessary. The defintion of group d through

defines all capping groups to the correct connectors. With the groups defined, we can now define the type of polymer we would like to build,

ITEM	CLUSTERS
poly	random,1
ITEM	END

The first keyword defines the polymer's name, poly here. Its name is followed by the type of polymer. The three available options are random, block, and alternate. Examples are available for each of these options in \${EMC_ROOT}/examples/polymer. Here we chose to create a random copolymer. The last entry denotes the polymer's fraction, here chosen to be 1. Different clusters and polymers can be added to this paragraph to form any mixture desired.

Finally, we define the polymer's morphology. We wanted 4 monomers A, 4 monomers B, and two terminators D,

ITEM	POLYMERS
poly	
1	a,4,b,4,d,2

ITEM END

Each polymer definition starts with its name as a header on a separate line. This name refers to the name chosen in the clusters section, here poly. The polymer's name is followed by all morphologies associated with this name. Here we only have one morphology, but we could also define a distribution of polymer morphologies. Each line contains the mol fraction of each contribution, followed by its morphological definition,

A subsequent polymer would start with its name on a separate line, followed by its definition(s), e.g. this section with two polymers would look like

ITEM	POLYMERS
poly 1	a,4,b,4,d,2
another 1 2 1	a,2,b,2,d,2 a,3,b,3,d,2 a,4,b,4,d,2
ITEM	END

Here, a polymer called another was added and assumed to have three contributing subpolymers, thus creating a distrubtion of polymer lengths within this polymer. The distribution uses mol fractions, which means, that the above example has a ditribution of 1:2:1, or 25% of the first entry, 50% of the second, and 25% of the third. The emc_setup.pl script normalizes fractions in the resulting EMC build script.

4.7.2.5 Multiphase Systems

EMC offers the possibility of building multiphase systems by building one phase after another. This allows for setting up systems, for which it is known a priori, that the equilibrated state phase separates. This then allows for studying mass distribution and interfacial tension behavior. Let us investigate this possibility by means of an example, in which we study the behavior of sodium dodecyl sulfate (SDS) in a system of water and oil. This example can be found in \${EMC_ROOT}/examples/setup/chemistry/multiphase, where the describing file multiphase.esh resides.

We assume the density of the water, SDS, and oil phase to be 1, 0.8, and 0.8 g/cc respectively. This system is at a temperature of 300K. We would like to describe the interactions with the CHARMM force field. The final simulation should have about 5000 particles. These prerequisites are captures in the OPTIONS paragraph (see \langle undefined \rangle), [Chemistry Options], page \langle undefined \rangle),

```
ITEM
             OPTIONS
             5000
ntotal
temperature 300
density
             1,0.8,0.8
phases
             water + sds + oil
field
             charmm/c32b1/all_27_prot_lipid
mass
profile
             true
replace
             true
ITEM
             END
```

Here, the keyword ntotal allows for setting the final number of sites to be simulated. The keyword temperature sets the temperature. The desnity of each phase is defined by keyword density through separation of each entry by either a comma or a tab. Chemically, each phase is defined by the keyword phases. Here, each phase is demarcated by the +-sign. Contributing clusters to each individual phase are separated by either commas, tabs, or spaces, e.g.

```
phases water sds + oil
```

would create two phases: the middle phase would consist of a random mixture of water and sds, while the outer phase would consist of just oil. The fractions entered in the SHORTHAND are assumed to be by mass, which is expressed by setting the keyword mass to true in the OPTIONS section. Furthermore, analysis mass profiles is added to the generated LAMMPS input script multiphase.in by setting the keyword profile to true. Lastly, replacement of all scripts and files produced by executing emc_setup.pl are selected to be overwritten by setting keyword replace to true. Defining the chemical composition can be entered in the SHORTHAND section (see Section 4.6.2 [Chemistry File], page 36),

ITEM SHORTHAND

Here, we select 40% by weight of both water and oil phases, while we add 20% by weight of SDS. Chemical identities are entered using the SMILES format (see Section 7.8 [SMILES], page 194).

Steps of execution are described in the usage section (see Section 4.2 [Setup Usage], page 9).

4.7.2.6 Surfaces

ITEM	OPTIONS
density	0.8 298
temperature ntotal	1000
shape	1.5
build_dir	•
field	opls-ua
charge	false
profile	true
replace	true
ITEM	END

Here, we chose the OPLS-UA force field without charges, select a mass profile section to appear in the LAMMPS input script, and allow for emc_setup.pl output scripts to be overwritten. The definition of surfaces occurs through predefined EMC input files, of which a few examples can be found in ./lib. One of these input files is a fully bonded diamond lattice called diamond_lattice.emc. EMC input files can contain variables, which can be redefined from an EMC input script. Specific variables need to be redefined, in order to turn the generic diamond lattice input file into a carbon diamond lattice. This redefintion occurs in the VARIABLES paragraph (see Section 4.6.2 [Chemistry File], page 36),

ITEM	VARIABI	LES
diamond_n diamond_a	ame	"*C(*)(*)*" "carbon" true
lbond		1.529
ITEM	END	

Four variables are predefined in the diamond lattice input file: diamond_chemistry, diamond_name, diamond_atomistic, and lbond. The above defintions overwrite the

predefined values. The value for 1bond has been taken from the OPLS-UA force field, where it represents the equilibrium bond distance between two ct types, which is 1.529 A. With options and variables set, we now need to define the group describing decane,

ITEM	GROUPS
decane	(C)10
ITEM	END

Here, an extension of SMILES was used to depict decane. Proper SMILES would require CCCCCCCC, but EMC allows for an extension to SMILES, for which a number after parenthesis is interpreted as the number of times, with which the snippet between the parenthesis is repeated. Finally, the CLUSTERS cluster section defines both decane and the surface,

ITEM	CLUSTERS			
decane surface	decane import	1 4	<pre>\$root+"lib/diamond_bonded"</pre>	emc
ITEM	END			

Here, the fraction, 1, of decane molecules is defined, followed by the surface definition. Both lines start with the name of the cluster. This name is followed by a reference to the corresponding group for the decane molecules. The second keyword for the surface molecule is a reserved keyword, which indicates, that this molecule is a surface, which uses the library file diamond_bonded as a reference file. The different phases are built in the x-direction. The number 4 indicates the number of surface cells to build into this direction, which defines the value of internal variable nx.

4.7.3 Environment Mode

The environment mode allows creating a multi-simulation environment, conveniently packaged in one clear file. One example dealing with determining shear is included. Environment examples can be found in \${EMC_ROOT}/examples/setup/environment/.

4.7.3.1 User-Defined Force Fields

This example demonstrates the concept of including a force field definition in a chemistry template, while adding loops in the environment section of the EMC Setup input script. The example can be found in ${EMC_ROOT}/examples/setup/environment/field$. EMC Setup provides the FIELD section for controlling user-defined force fields. The idea behind this example is to study the hypothetical influence of branching. To this extend, the number of branch points and the Lennard-Jones (LJ) interaction constant epsilon is varied. The work directory ${WORKDIR}$ is defined by

\${EMC_ROOT}/examples/setup/environment/field

later referred to by \${WORKDIR}. The user is advised to maintain a standardized directory structure when using the environment mode of EMC Setup. This means, that EMC Setup environment .esh files are stored in \${WORKDIR}/setup. Execution of the setup files should occur in \${WORKDIR} and not in \${WORKDIR}/setup. Once executed in \${WORKDIR}, EMC Setup creates directories analyze, build, chemistry, run, and optionally test in \${WORKDIR}. A typical convention for setup files is to name them by date, followed by a serial number, e.g. 2018070100 means, that the setup file was created on July 1, 2018 and is the first of the series of files created on that date.

Environment

All sections outside the TEMPLATE section define the environment and is considered to be the environment mode of EMC Setup. This includes the ENVIRONMENT and LOOPS sections as well as the STAGE designator. The ENVIRONMENT section defines the project name, queue settings, and high performance computing (HPC) cluster architecture parameters. Additionally, the name of a test directory can be defined by the option name_testdir (see the 'Execution' section below).

Loops

Loop variables are defined in the LOOPS section. Variables stage and trial are reserved keywords. The former indicates the stage of a project, e.g. one likes to study branched and linear structures. This could be devided in a branched stage and a linear stage. Here, only the branched variety is considered. Furthermore, the number of branches is indicated by nbranches and the settings for the LJ interaction constant by epsilon. These variables are later referred to in the TEMPLATE section by @{NBRANCHES} and @{EPSILON} respectively.

Template

The TEMPLATE section contains the definition of one simulation, here referred to as the chemistry mode of EMC Setup. In this chemistry template, one can use variables as defined in the LOOPS section of the environment. A template should at least contain an OPTIONS, GROUPS, and CLUSTERS section to fully define the simulation setup. Here, an additional POLYMERS section has been added to control the definition of the to be studied branched oligomers. Since polymers are requested, connectivity has to be defined in the GROUPS section. This connectivity is referred to with the '*' character. The numbers after the

chemical definition through a SMILES refer to the occurrence of the '*' character in the SMILES, e.g. the definition of group A

```
*a*,1,A:2,1,B:2,1,B:3,2,B:1
```

means, that the first '*' in '*a*' connects to the second of group A and the second and third of group B. The second star in '*a*' connects the first of group B.

The CLUSTERS section defines the molecule and the amount of each contributor, i.e. the solvent is defined by group S and has an 80% occurrence. The polymer is defined as a random copolymer and has a 20% occurrence.

The POLYMERS paragraph holds all definitions of polymers as referred to in the CLUSTERS paragraph. Here, only the polymer polymer exists with 10 repeat units of monomer A as defined by group A, and a variable amount of monomer B and terminator T. The amount is defined by O{NBRANCHES} which refers to nbranches in the LOOPS section.

The FIELD section defines the user-provided force fields and contains subsections related to the subsequent parameter definitions. Mandatory subsections are MASS, NONBOND, and – in case of connectivity – BOND. Additionally, in the case of this example angle terms are also defined in the ANGLE subsection. No torsions were needed. Wildcards in the form of '*' are allowed in all but the MASS section. In the case of the latter, each type can only contain one '*' character, which has to be at the end of the type. Note that the variable definition of the LJ epsilon is taken care of by <code>@{EPSILON}</code> in the <code>NONBOND</code> subsection.

Execution

Creation of the simulation environment occurs in \${WORKDIR} through

```
./setup/2018070100.esh
```

or, when the above is not executable or emc_setup.pl is not in your path,

```
${EMC_ROOT}/scripts/emc_setup.pl setup/2018070100.esh
```

Though the latter being possible, the user is adviced to add both bin and scripts directory to their path, e.g.

```
export PATH=${EMC_ROOT}/bin:${EMC_ROOT}/scripts:${PATH}
```

Once executed, several bash scripts are generated in directories analyze, build, and run with name 2018070100.sh. Subsequently, the provided test directory name appears as a subdirectory of \${WORKDIR}/test/, in which the script setup.sh is created. Execution of the latter creates an instance of the first occurrence of each loop variable, i.e. for STAGE = branched, NBRANCHES = 2, and EPSILON = 0.9. This test directory serves the purpose of testing the validity of the TEMPLATE section without having to submit to a queueing system.

Before summitting, an option which you would need to set is queue_ppn (procs or cores per node), since it is unknow how many cores can run on one of your nodes. EMC will

pack single core build jobs together on a node if it knows the ppn. Similarly, you can set the memory per core with queue_memory.

Submission to a queueing system occurs though ./build/2018070100.sh, followed by ./run/2018070100.sh once all builds as spawned by the former have finished. Chaining of builds and runs is also possible by executing the run script with build mode included, i.e.

./run/2018070100.sh -build

This spawns build and lets running of the systems LAMMPS wait for the building to finish.

4.7.3.2 Shear

The shear example, as found in \${EMC_ROOT}/examples/setup/environment/nemd, illustrates the use of the EMC setup environment mode to define a set of simulations, calculating time dependent pressure tensors for a number of n-alkanes with various shear rates. The environment mode allows for scanning through parameter space, aiding the study of material response to various conditions. The example defines 40 simulations in total. An environment script contains multiple distinct sections. A shear simulation needs an equilibrated non-sheared configuration as a starting point. First, the environment mode needs to be triggered in the first occurring OPTIONS section,

ITEM	OPTIONS		
project		pure	
replace		true	
environment		true	
name_analyze	Э	date00)
name_build		date00)
name_run		date00)
queue_build		defaul	Lt
queue_analy2	ze	defaul	Lt
shear		true,	true
restart		fal	
ncores		8	
trun		_	
ITEM	END		

Besides toggling the environment mode, this section also defines the general project name, names associated with build, run, and analysis scripts, execution queues, and the number of cores to execute LAMMPS simulations on. Build, run, and analysis names use the base name of the chemistry script by as default. Zero shear simulations do not start from a restart file, but uses the output generated by EMC instead.

Loops can be defined over project stages, trials, and user defined variables in section LOOPS. This section controls parameter space for which phenomena are to be studied.

ITEM	LOOPS						
stage		pure					
trial		c-06	c-07	c-08	c-09	c-10	&
		c-11	c-12	c-13	c-14	c-16	
shear		0.00e0					
сору		1					
ITEM	END						

Reserved variables stage, trial, and copy are internally defined. Both stage and trial relate to other sections in the the environment script. Each stage is associated with a specific chemistry file, which is defined in section TEMPLATE of the environment script. The definitions for variable trial are associated with section STRUCTURES. The reserved variable copy supports statistical sampling by creating independent structures (here, as an example, 1 structure is created). The use of the copy variable should be used with care, since it functions as a multiplier for the number of simulations, that are considered. In this example, the defined 11 trials and 4 shears comprise 44 simulations. This would increase to 440 simulations, when copy is set to 10 instead of 1.

The variable shear is a user defined variable. It is expected to be lower case. Its reference in the chemistry template is expected to be capitalized. Shell scripts are created after each LOOPS section. Build and run scripts – with names date00.sh and date10.sh, respectively, as defined in the previous OPTIONS section – are written in a newly created directory run in the current work directory. An analysis script with the name date10.sh is created in in a newly created directory analyze.

Subsequent shear simulations are defined after zero shear simulations. Here, only a few lines are needed. Non-defined parameters take the value of their definition in previous OPTIONS sections.

```
ITEM OPTIONS

name_build -
name_run date10
name_analyze date10
restart true, ../../0.00e0

ITEM END
```

These subsequent simulations will use equilibrated zero shear simulations as their starting point. A build script is not needed in this case, which is denoted by defining parameter name_build with a dash. Restart files are used as starting point, which is triggered by setting parameter restart and defining the source directory for the restart files, ../../0.00e0 in this case (the 0.00e0 value refers to the previously set zero shear).

Multiple shear rates are needed to see the shear behavior, intending to reach the Newtonian plateau at very low shear. Depending on the substance, this plateau might not be possible to be reached.

```
ITEM LOOPS

shear 1.00e-4, 1.00e-5, 1.00e-6

ITEM END
```

Shear rates are expressed in box length per time step. After the definition of this paragraph, a run shell script with name 2016103120.sh is added to directory run, and an analysis shell

script with name 2016103120.sh is added to directory analyze. Note, that the definition of stage and trial are inherited from the previous definition of section LOOPS.

With all shell scripts defined, now we can define each stage and trial separately by using sections STAGE and TRIAL respectively. After invoking STAGE and/or trial, all subsequent sections apply to the defined stage and/or trial. Here, we only need to define a section STAGE with name pure, relating back to the definition of variable stage in section LOOPS. Note, that both sections STAGE and TRIAL do not require an ITEM END designation.

ITEM STAGE pure

A template is used to define the actual simulation conditions. A template section is defined by ITEM TEMPLATE and needs an ITEM END to close. Variables appearing in section LOOPS can be referred to by starting with @ followed by the variable name in all capitals. A simulation is defined as usual withing a template section.

OPLS-UA reflects the force field chosen for our n-alkanes. A total of 10000 beads is requested by setting ntotal. The equilibration time is set to 100000 time steps with tequil. The initial density is chosen to be 0.85 g/cc. The simulation is run at equal pressure of 1 atm and temperature of 300 K by using an NPT ensemble. Note, that it is possible, that npt/sllod is not available in all LAMMPS versions, in which case the presure line should be omitted. Shear rates are set by the shear parameter. Here, @SHEAR refers back to variable shear in the LOOPS section. All exisiting scripts are overwritten by setting parameter replace to true. The usage of bond increments depend on the chosen force field. An empty bond increment is used when non-existing bond increments are encountered. Note, that OPLS-UA does not use bond increments.

ITEM	OPTIONS	
replace		true
field		opls-ua
ntotal		10000
tequil		100000
density		0.85
pressure		1.0
temperature		300
shear		@SHEAR, erate
replace		true
increment		empty
ITEM	END	

With simulation conditions set, substance definitions are in order. We chose to use a structure section to define chemical structures. As a consequence, the variable structure is defined and used to replace all occurring @STRUCTURE instances in the chemistry template.

ITEM SHORTHAND

molecule		@STRUCTURE,1
TTEM	END	

The template section is closed with an ITEM END. This template is used to define all simulations. It is stored with name pure.esh in a created directory chemistry/stages within the current work directory. Also, an adaptation shell script with name pure.sh is stored in created directory chemistry/scripts.

Finally, desired substances have to be defined. Doing so in section STRUCTURES suffices, since we are only interested in bulk properties of pure components.

ITEM	STRUCTURES	
c-06	(C)6	
c-07	(C)7	
c-08	(C)8	
c-09	(C)9	
c-10	(C)10)
c-11	(C)11	L
c-12	(C) 12	2
c-13	(C)13	3
c-14	(C) 14	1
c-15	(C) 15	5
c-16	(C)16	3
ITEM	END	

Each structure is stored in a separate file with names as occuring in the first column. These files are stored in directory chemistry/structures.

Execution of the whole simulation can be done on either the cluster on which the simulations are to run, or can be done from the machine on which the final results are to be analyzed. In the former case, each of the three scripts in directories build, run, and analysis are to be executed sequentially, i.e. build/date00.sh to build all simulations, subsequently followed by zero shear simulations through run/date00.sh when all build simulations are completed, an by finite shear simulations through run/date10.sh when all zero shear simulations are completed. Each script uses run.sh to submit jobs to the queueing system, which can be found in \${EMC_ROOT}/scripts. The run.sh supports LSF and PBS queueing systems. The run_host.sh script can be found in the same directory allows remote submission of jobs. It assumes, that the execution directory structure is identical on both local and remote host machines. run_host.sh can be used to for instance execute the build shell script remotely through execution of

run_host.sh user@remote.machine.net build/date00.sh

in the current work directory. Log files are created both locally and remotely by run_host.sh, keeping records of simulation execution. The example above would create

build/date00.log. Remote execution of subsequent run scripts would follow each other sequentially upon completion of its predecessor.

4.8 Help Output

The following output appears when invoking emc_setup.pl either without any options, with a non-exisiting option, or with the -help option.

```
EMC Setup v4.1.3 (March 30, 2022), (c) 2004-2022 Pieter J. in 't Veld
```

Usage:

```
emc_setup.pl [-command[=#[,..]]] project [phase 1 clusters [+ ...]]
```

```
Commands:
 -analyze_archive archive file names associated with analyzed data [true]
                   create tar archive from exchange file list [true]
 -analyze_data
 -analyze_last
                   include last trajectory frame (deprecated) [false]
 -analyze_replace replace already exisiting analysis results [true]
 -analyze_skip set the number of initial frames to skip [0]
 -analyze_source set data source directory for analysis scripts
 -analyze_user
                   set directory for user analysis scripts
                   set the number of frames in window average [1]
 -analyze_window
 -angle
                   set DPD angle constants k and theta or set angle field
                   option (see below) [5,180]
                   add wildcard entry to mass and nonbond sections in DPD
 -auto
                   .prm [false]
                   set bin size for LAMMPS profiles [0.01]
 -binsize
                   set bond constants k,1
  -bond
                   set build script name [build]
 -build
                   insert first site at the box center [false]
 -build_center
 -build_dir
                   set build directory for LAMMPS script [../build]
 -build_order
                   set build order of clusters [random]
 -build_origin
                   set build order of clusters [x=0, y=0, z=0]
 -build_replace
                   replace already existing build results [false]
                   set the minimal insertion angle [false]
 -build_theta
                   chemistry contains charges [true]
 -charge
                   set charge interaction cut off [9.5]
 -charge_cut
                   use chunk approach for profiles in LAMMPS script [true]
 -chunk
                   use communicate keyword in LAMMPS script [false]
 -communicate
                   set core diameter [-1]
  -core
                   include nonbond cross terms in LAMMPS params file [false]
 -cross
                   treat imported structure as a crystal [auto]
 -crystal
                   set pairwise interaction cut off [9.5]
 -cut
                   set pairwise interaction cut off [center=-1, charge=9.5,
  -cutoff
                   ghost=-1, inner=-1, outer=-1, pair=9.5, repulsive=0,
                   rmax=-1
                   control debugging information [false]
  -debug
                   deform system from given density [nblocks=1, ncycles=100,
 -deform
                   type=relative, xx=1, yx=0, yy=1, zx=0, zy=0, zz=1]
                   sets which clusters to delete; each deletion is separated
 -delete
                   by a +-sign; default assigns no clusters to delete
```

```
[clusters=all, fraction=0.5, groups=all, mode=random,
                  phase=1, sites=all, thickness=1, type=relative]
-density
                  set simulation density in g/cc for each phase [1]
                  set charge medium dielectric constant [1]
-dielectric
-direction
                  set build direction of phases [x]
                  set LAMMPS trajectory file write frequency [100000]
-dtdump
                  set LAMMPS restart file frequency [100000]
-dtrestart
                  set LAMMPS thermodynamic output frequency [1000]
-dtthermo
                  create EMC build script [true]
-emc
                  set ring recognition depth in groups paragraph [8]
-emc_depth
-emc_exclude
                  set EMC section to exclude [build=false]
                  execute EMC build script [false]
-emc_execute
-emc_export
                  set EMC section to export [smiles=]
                  set Monte Carlo moves for after build [displace=1]
-emc_moves
                  set EMC output modes [debug=false, exit=true, info=true,
-emc_output
                  warning=true]
                  set progress indicators [build=true, clusters=false]
-emc_progress
                  set Monte Carlo run conditions for after build
-emc_run
                  [clusters=all, groups=all, nblocks=100, ncycles=0,
                  nequil=0, sites=all]
-emc_test
                  test EMC build script [false]
                  settings for EMC trajectory [append=true, frequency=0]
-emc_traject
-environment
                  create project environment [false]
                  set long range ewald summations [true]
-ewald
-exclude
                  exclude previous phase during build process [true]
                  set expert mode to diminish error checking [false]
-expert
                  set environment script extension [.esh]
-extension
                  set extra interactions dest:src:offset (deprecated)
-extra
                  set force field type and name based on root location
-field
                  set angle field option (see below) [-]
-field_angle
                  set bond field option (see below) [-]
-field_bond
                  check system charge after applying force field [true]
-field_charge
                  check force field compatibility [true]
-field_check
                  set debug field option [false]
-field_debug
                  set various DPD options [auto=false, bond=false]
-field_dpd
                  override field errors (used for debugging) [true]
-field_error
                 parameter format of generated force field [%15.10e]
-field_format
                  set group field option (see below) [-]
-field_group
-field_id
                  set force field id [opls-ua]
                  set improper field option (see below) [-]
-field_improper
-field_increment set increment field option (see below) [-]
                 set force field location [~/emc/v9.4.4/field]
-field_location
                  set force field name [opls/2012/opls-ua]
-field_name
-field_nbonded
                  set number of excluded bonded interactions [0]
                 set force field reduced units flag [false]
-field_reduced
                 set torsion field option (see below) [-]
-field_torsion
                 set force field type [opls]
-field_type
```

```
-field_write
                  create field parameter file [true]
                  list of molecules to focus on [-]
-focus
-ghost_cut
                  set pairwise interaction cut off [-1]
                  (deprecated: use weight) set build relaxation grace
-grace
                  [0.9999, 0.9999, 0]
                  this message
-help
-hexadecimal
                  set hexadecimal index output in PDB [false]
-host
                  set host on which to run EMC and LAMMPS [air]
-info
                  control runtime information [true]
-inner
                  set inner cut off [-1]
-insight
                  create InsightII CAR and MDF output [false]
-insight_compress set InsightII CAR and MDF compression [false]
-insight_pbc
                  apply periodic boundary conditions [true]
-insight_unwrap
                  apply unwrapping [true]
-kappa
                  set electrostatics kappa [4]
-lammps
                  create LAMMPS input script or set LAMMPS version using
                  year, e.g. -lammps=2014 (new versions start at 2015)
                  [true]
-lammps_cutoff
                  generate output of pairwise cut off in parameter file
                  [false]
-lammps_dlimit
                  set LAMMPS nve/limit distance [0.1]
-lammps_error
                  restart LAMMPS only upon previous error [false]
-lammps_pdamp
                  set LAMMPS barostat damping constant [1000]
-lammps_tdamp
                  set LAMMPS thermostat damping constant [100]
-location
                  prepend paths for various file locations
                  [analyze=.:~/emc/v9.4.4/scripts/analyze,
                  field=::~/emc/v9.4.4/field, include=.]
                  assume mass fractions in chemistry file [false]
-mass
                  set queue memory per core in gb [default]
-memorypercore
                  manipulate runtime modules in format [command=]module
-modules
-mol
                  assume mol fractions in chemistry file [true]
                  set zero total momentum in LAMMPS [100,1,1,1,angular]
-momentum
                  define cluster move settings used to optimize build
-moves_cluster
                  [active=false, cut=0.05, frequency=1, limit=auto:auto,
                  max=0:0, min=auto:auto]
                  set LAMMPS mean square displacement output [false]
-msd
                  create NAMD input script and parameter file [false]
-namd
                  set electrostatic interaction update frequency [1]
-namd_dtcoulomb
                  set output frequency of DCD file [10000]
-namd_dtdcd
                  set nonbonded interaction update frequency [1]
-namd_dtnonbond
                  set output frequency of restart files [100000]
-namd_dtrestart
                  set output frequency of thermodynamic quantities [1000]
-namd_dtthermo
-namd_dttiming
                  set timing frequency [10000]
-namd_dtupdate
                  set update frequency [20]
-namd_pres_decay set pressure ensemble decay [50]
-namd_pres_period set pressure ensemble period [100]
                  set temperature ensemble damping [3]
-namd_temp_damp
```

```
-namd_tminimize
                  set number of minimization timesteps [50000]
                  set number of timesteps for running [10000000]
-namd_trun
-name_analyze
                  set job analyze script name [chemistry]
                  set job build script name [chemistry]
-name_build
-name_run
                  set job run script name [chemistry]
                  set analyze, job, and build script names simultaneously
-name_scripts
-name_testdir
                  set job test directory as created in ./test/ [-]
-nchains
                  set number of chains for execution of MD jobs
-ncores
                  set number of cores for execution of MD jobs [-1]
                  set queue cores per node for packing jobs [default]
-ncorespernode
-niterations
                  set number of build insertion iterations [1000]
                  control possibility of restarting when rerunning [false]
-norestart
-nparallel
                  set number of surface parallel repeat unit cells [auto]
                  set number of build relaxation cycles [100]
-nrelax
                  number of configuration in profile [1000]
-nsample
                  set total number of atoms [10000]
-ntotal
-number
                  assume number of molecules in chemistry file [false]
                  omit fractions from chemistry file [false]
-omit
-options_perl
                  export options, comments, and default values in Perl
                  syntax [false]
-options_tcl
                  export options, comments, and default values in Tcl
                  syntax [false]
                  set outer cut off [-1]
-outer
                  set pair constant defaults
-pair
                  set parameters file name [parameters]
-parameters
                  create field parameter file [true]
-params
                  create PDB and PSF output [true]
-pdb
                  set atom name behavior [index]
-pdb_atom
-pdb_compress
                  set PDB and PSF compression [true]
-pdb_connect
                  add connectivity information [false]
-pdb_cut
                  cut bonds spanning simulation box [false]
-pdb_extend
                  use extended format for PSF [false]
-pdb_fixed
                  do not unwrap fixed sites [true]
-pdb_parameters
                  generate NAMD parameter file [false]
                  apply periodic boundary conditions [true]
-pdb_pbc
-pdb_rank
                  apply rank evaluation for coarse-grained output [false]
                  set residue name behavior [index]
-pdb_residue
                  do not unwrap rigid sites [true]
-pdb_rigid
                  set segment name behavior [index]
-pdb_segment
                  apply unwrapping [clusters]
-pdb_unwrap
                  add Van der Waals representation [false]
-pdb_vdw
                  import percolating InsightII structure [false]
-percolate
                  sets which clusters to assign to each phase; each phase
-phases
                  is separated by a +-sign; default assigns all clusters to
                  phase 1 [all]
-polymer
                  default polymer settings for groups [bias=none,
```

fraction=number, niterations=-1, order=list]

port EMC setup variables to other applications -port set charge kspace precision [0.001] -precision -prefix set project name as prefix to LAMMPS output files [false] use gcc to preprocess the input script [false] -preprocess -pressure set system pressure and invoke NPT ensemble; optionally add direction and/or (un)couple for specifying directional coupling [false,direction=x+y+z,couple] -profile set LAMMPS profile output [density=false, density3d=false, pressure=false] -project set project name; slashes are used to create subdirectories queue settings [account=none, analyze=default, -queue build=default, memory=default, ncores=-1, ppn=default, run=default, user=none] set queue account for billing [none] -queue_account -queue_analyze set job analyze script queue [default] -queue_build set job build script queue [default] set queue memory per core in gb [default] -queue_memory -queue_ncores set number of cores for execution of MD jobs [-1] set queue cores per node for packing jobs [default] -queue_ppn -queue_run set job run script queue [default] options to be passed directly to queuing system [none] -queue_user turn off all information -quiet set build relaxation radius [5] -radius set record entry in build paragraph [cut=false, -record frequency=1, inactive=true, name="", pbc=true, unwrap=sites] -references set references file name [references] -region_epsilon set epsilon to use for exclusion regions [0.1] -region_sigma set sigma to use for exclusion regions [1] -replace replace all written scripts as produced by EMC setup [false] set replica interactions dest:src:offset -replica -restart create LAMMPS restart script [false,..] -rlength set reference length -rmass set reference mass set maximum build cutoff radius [-1] -rmax set reference type -rtype set sampling options for LAMMPS input script -sample [energy=false, green-kubo=false, gyration=false, msd=false, pressure=true, volume=false] set script file name [chemistry] -script set number of colums in output scripts [80] -script_ncolums set initial random seed [-1] -seed -shake set shake types -shake_iterations set maximum number of shake iterations [20] -shake_output set shake output frequency [never]

-shake_tolerance set shake tolerance [0.0001] -shape set shape factor [1] -shear add shear paragraph to LAMMPS input script [false] set LAMMPS skin [2] -skin sets which clusters to partition; each split is separated -split by a +-sign; default assigns no clusters to split [clusters=all, fraction=0.5, groups=all, mode=random, phase=1, sites=all, thickness=1, type=relative] -suffix set EMC and LAMMPS suffix [_air] system identification and checks during building -system [charge=true, geometry=true, id=main, map=true, pbc=true] check for charge neutrality after build [true] -system_charge -system_geometry check geometry sizing upon building [true] check for charge neutrality after build [main] -system_id map system box before build [true] -system_map -system_pbc apply periodic boundary conditions after build [true] -temperature set simulation temperature in K [300] set LAMMPS equilibration time [1000] -tequil set LAMMPS profile sampling frequency [10] -tfreq set LAMMPS thermo style to multi [false] -thermo_multi -tighten set tightening of simulation box for imported structures [3] set job analyze script wall time [00:30:00] -time_analyze -time_build set job build script wall time [00:10:00] set job run script wall time [24:00:00] -time_run set integration time step [2] -timestep set LAMMPS triclinic mode -triclinic -trun set LAMMPS run time [10000000] set units type [real] -units set units for energetic scale [1] -units_energy set units for length scale [1] -units_length output version information [false] -version set recalculation based on molecular volume [false] -volume control warning information [true] -warn set build relaxation energetic weights [bond=0.0001, -weight focus=1, nonbond=0.0001] use double width in scripts [false] -width set work directory [~/emc/v9.4.4/texinfo/setup] -workdir

Notes:

- * This script comes with no warrenty of any kind. It is distributed under the same terms as EMC, which are described in the LICENSE file included in the EMC distribution.
- * Queue name 'default' refers to whichever queue is default; queue name 'local' executes all jobs sequentially on local machine
- * Reference and parameter file names are assumed to have .csv extensions
- * Chemistry and environment file names are assumed to have .esh extensions

- * File names with suffixes _chem can be taken as chemistry file names wild cards
- * Chemistry file format: mol id, smiles string, fraction, mol mass[, mol vol]
- * Reserved environment loop variables are: stage, trial, and copy
- * Densities for multiple phases are separated by commas
- * Shears are defined in terms of erate; values < 0 turns shear off
- * Inner and outer cut offs are interpreted as fractions for colloidal force fields
- * Valid field options are: ignore, complete, warn, empty, or error
- * A '+' sign demarcates clusters for each phase; remaining clusters are assigned to the first empty phase

5 Scripting Commands

EMC is driven through a scripting language that accesses the different predefined routines driving building and simulation of configurations. EMC requires each script to start with the text "(* EMC: Script *)" on the first line. Absence of this line will result in an error upon start up of EMC. The following sections describe the syntax of commands in the current version. Currently, commands use a variable representation similar to Mathematica. A change to a function representation is planned in future.

5.1 Build

5.1.1 Syntax

```
build
                = {
  system
                -> {id -> constant, temperature -> real,
                      density -> real, geometry -> voigt,
                      flag -> {charge -> boolean, map -> boolean}},
                -> {progress -> boolean, frequency -> long,
  select
                      message -> option, center -> boolean, origin -> vector,
                      order -> constant, check -> boolean,
                      cluster -> {constant, ...}, name -> string,
                      relax -> {ncycles -> long, radius -> real},
                      grow -> {method -> constant, check -> constant,
                        cutoff -> real, grace -> {real, real},
                        theta -> real, dphi -> real, nbonded -> long,
                        ntrials -> long, niterations -> long,
                        include -> {region, ...},
                        exclude -> {region, ...}}}
};
```

Directive system id	Parameters struct constant	Description Defines the system dimensions associated with a build. Sets identity of system to which to add the build; can be either textual or numerical.
density	real	Sets the system density; can either be set to calculate when chosing or forced when chosing values > 0.0.
geometry	voigt	Sets the system geometry; used as a fractional size when density is set, otherwise used as system dimensions from which a density is calculated (see Section 7.15 [Voigt], page 346).
flag	boolean	Control system checks and optimizations.
charge	boolean	Perform a total charge check of the system and exit on a non-zero value; options are true or false.
map	boolean	Map a stretched triclinic cell geometry into its minimal representation; options are true or false.
Directive select progress	Parameters struct boolean	Description Selects building style and clusters Shows progress indicator; options are bar, list, or none.
frequency	long	Sets the progress frequency (in percentages), where $0 \le$

frequency ≤ 100 .

message	option	Sets the style for energetic output in progress report; valid options are none, raw, nkt, or n, which corresponds to energy in no particular units, internal units, units of nkT or units of n, respectively; internal units correspond to the units of the selected force field (kcal/mol for COMPASS, OPLS, GROMACS and MARTINI).
center	boolean	Place first inserted site at origin as defined origin option; options are true or false.
origin	{0,0,0}	Define origin for first inserted site; used only when center option is set to true.
order	constant	Insert clusters in sequence or at random.
check	boolean	Check for (semi)fixed sites and abort if present; options are true or false.
cluster	array	Sets the cluster(s) participating in the build.
name	string	Sets the output name for generation of an XYZ file upon error of all active sites of the built system; no output is generated, when the output name is not set.
relax	struct	Selects intermediate structure relaxation during the building process; Monte Carlo is performed on a spherical volume with a radius around the inserted site for ncycles after each successful insertion.

Directive grow method	Parameters struct option	Description Describes growth method and settings. Sets building style; options are overlap for hard sphere excluded volume considerations, energetic for energetic considerations, and minimum for energetic minimum considerations.
check	option	Sets overlap checking; options are bonded for checking overlaps up to four bonds away, cluster for checking intramolecular, all for intra- and intermolecular overlaps, and none for no overlap checking.
cutoff	real	Sets the energetic cutoff when using method -> minimum, where decisions are based on boltzmann sampling when cutoff <= 0.
grace	{real, real}	Sets the grace fraction, which correlates to how much overlap is allowed; 0 corresponds to no overlap, while 1 corresponds to full overlap (typically < 0.75); the first value influences the nonbonded acceptance, while the second value influences the bonded acceptance; an energetical building style requires grace < 0.9995 for coherent energies.
theta	real	Sets the minimum allowed bond angle.

dphi	real	Sets the maximum allowed torsion angle for for bonded construction using a Monte Carlo scheme; needed for all
		force field definitions beyond bond length interactions
		(i.e. it is neglected in case of the existence of bonds only
		or when a lower threshold is crossed).
nbonded	integer	Sets the number of trials moves for bonded construction
	_	using a Monte Carlo scheme.
ntrials	integer	Sets the number of trial overlap or energetic calculations.
niterations	integer	Sets the number of iterative trials.
include	region	Sets inclusion regions (see Section 7.7 [Region], page 193).
exclude	region	Sets exclusion regions (see Section 7.7 [Region],
	-	page 193).

5.1.2 Usage

The build command enables the build of previously defined clusters (see Section 5.4 [Clusters], page 90) into a desired system. All building modes apply energetic sampling of bonded potentials during the bond building process. Currently a hard sphere overlap and an energetic build is available. Note that a system needs to be defined before build mode can be executed (e.g. define system before select). Note that hard and soft modes regulate insertion behavior with respect to exclusion regions. The hard mode applies the region to all sites of inserted clusters, while the soft mode only applies the region to the first site of inserted clusters.

5.1.3 Default

The default is given by

5.1.4 Examples

```
grace -> 0.5,
method -> overlap,
theta -> 0.3*pi,
cluster -> {polymer, solvent}
}
```

5.2 Cancel

5.2.1 Syntax

cancel = {style -> option};

Directive Parameters Description

style option Selects cancellation style; options are abort and exit.

5.2.2 Usage

The cancel command is used to stop a running script at the selected position. Its main application serves debugging purposes. Script debugging can use either abort or exit, while source code debugging would use abort.

5.2.3 Default

The default is given by

cancel = {style -> exit};

5.3 Carve

5.3.1 Syntax

Directive mode	Parameters option	Description Sets mode of operation; options are include and exclude.
fraction	real	Sets the fractions in each system of the groups to be carved out; the value of fraction lies between 0.0 and 1.0 ; choses nsites when fraction < 0.0 .
nsites	integer	Sets the number of sites remaining in the selected region in each system; number of sites in group sites increments.
systems	constant	Sets target systems; can be one or more systems.
inside	region	Sets the inner boundary of the region in which sites are to be deleted (see Section 7.7 [Region], page 193).
outside	region	Sets the outer boundary of the region in which sites are to be deleted (see Section 7.7 [Region], page 193).

5.3.2 Usage

The carve command carves a shape out all systems in the currently defined simulation. It will output the number of retained sites per selected system upon execution as info using the nsites directive when a fraction for the selected systems is defined. It will use nsites when the fraction for the selected systems is a number smaller than zero.

5.3.3 Default

The default is given by

5.3.4 Examples

```
An example is given by
```

5.4 Clusters

5.4.1 Syntax

```
clusters
               = {
 cluster
               -> {id -> constant, system -> constant, n -> integer,
                      [group -> constant | template -> constant]},
               -> {id -> constant, system -> constant, n -> integer,
 polymer
                     type -> option, groups -> {constant, ...},
                     nrepeat -> {integer, ...}},
                -> {core -> constant, type -> constant, polymer -> constant,
 graft
                     connect -> option},
                -> {id -> constant, system -> constant, n -> integer,
 body
                     density -> real, [surface -> constant,]
                      core -> constant, region -> region},
};
```

Directive cluster	Parameters struct	Description Defines multiple single group clusters.
polymer	struct	Defines multiple repeating group clusters.
graft	struct	Defines graft connection to a backbone cluster group.
body	struct	Defines multiple bodies constrained by a region; particles consist either one core with multiple surface sites or core sites only.
id	constant	Sets identifier, either textual or numerical.
system	constant	Sets destination system to construct in,
n	integer	Sets the number of molecules to create.
group	constant	Sets the contributing group; must be a group without branches for cluster.
template	constant	Sets the contributing template as predefined in force field files (see Section 5.11 [Field], page 103).
nrepeat	integer	Sets the number of repeat units of each contributing monomer in the polymer; one nrepeat can be given in case of multiple groups, which results in assignment of this one nrepeat to all contributing groups.
groups	constant	Sets contributing groups; must be branched and connected groups; groups will be terminated with their respective terminators as defined by the groups command (see Section 5.17 [Groups], page 114).

type	option	Sets the type of polymer; options are block for block copolymers, alternate for alternating copolymers, and random statistical copolymers.
core	constant	Sets the core cluster; must be a cluster without branches and only one contributing site; density describes core density when no surface groups are defined.
type	constant	Sets the grafting type from which to grow branch; only one grafting type per grafting core is allowed.
connect	option	Sets the growth starting point; options are head or tail.
surface	constant	Sets the surface group; must be a group without branches and only one contributing site; one core particle is assumed; density describes surface density.
region	region	Sets the region of the particle (see Section 7.7 [Region], page 193).

5.4.2 Usage

The clusters command creates clusters according to the desired schemes as outlined above, which populate the desired system as set by the system directive.

5.4.3 Default

The default is given by

5.4.4 Examples

Direct definition of molecules uses the keyword cluster, e.g. a configuration with 1000 water molecules is obtained through

Polymers can either display a block, alternating, or random constallation of its constituing monomers. Assume three monomers, combined with one terminator,

```
groups = {
  group -> {
    id -> A,
    chemistry -> "*A*",
    connects -> {
```

```
{head, {A, tail}}, {head, {B, tail}}, {head, {C, tail}},
      {tail, {A, head}}, {tail, {B, head}}, {tail, {C, head}},
      {head, {T, head}}, {tail, {T, head}}
    }
  },
                 -> {
  group
                 -> B,
    id
    chemistry
                 -> "*B*",
    connects
                 -> {
      {head, {A, tail}}, {head, {B, tail}}, {head, {C, tail}},
      {tail, {A, head}}, {tail, {B, head}}, {tail, {C, head}},
      {head, {T, head}}, {tail, {T, head}}
    }
  },
                 -> {
  group
    id
                 -> C,
    chemistry
                 -> "*C*",
    connects
                 -> {
      {head, {A, tail}}, {head, {B, tail}}, {head, {C, tail}},
      {tail, {A, head}}, {tail, {B, head}}, {tail, {C, head}},
      {head, {T, head}}, {tail, {T, head}}
    }
  },
                 -> {
  group
                 -> T,
                 -> "*T",
    chemistry
    connects
                 -> {
      {head, {A, head}}, {tail, {B, head}}, {tail, {C, head}},
      {head, {A, tail}}, {head, {B, tail}}, {head, {C, tail}}
    }
  }
};
One hundred homopolymer molecules consisting of 10 A monomers are defined by
clusters
  polymer
                 -> {id -> polymer, system -> main,
                       n -> 100, nrepeat -> 10, type -> random, groups -> A}
};
which generates 100 clusters with TAAAAAAAAAT as a resulting polymer sequence.
Alternatively, these monomers can be combined into three types of copolymers, i.e. random,
block, and alternating coplymers. A random copolymer consisting of 10 A monomers, 20 B
monomers, and 30 C monomers, is defined by
clusters
                 -> {id -> polymer, system -> main,
  polymer
                       n \rightarrow 100, repeat \rightarrow \{10, 20, 30\}, type \rightarrow random,
                       groups -> {A, B, C}}
};
```

in which all monomers are randomly distributed over the polymer backbone. Construction of a block copolymer consisting of 2 A, 3 B, and 4 C monomers, is obtained through

which generates TABCABCBCCT as a resulting polymer sequence. Note the effect as a result of the unequal numbers when defining monomers.

5.5 Crystal

5.5.1 Syntax

```
crystal = \{n \rightarrow \{x, y, z\}, \text{ miller } \rightarrow \{h, k, l\}, \text{ plane } \rightarrow \text{ option,}
periodic \rightarrow \text{ vector}\};
```

Directive	Parameters	Description
n	$\{x, y, z\}$	Sets the number of crystal repeat units in x, y, z integer increments.
miller	{h, k, 1}	Sets miller crystal plane indices specifying the desired
		tilt, where h, k, and 1 are integers.
plane	option	Specifies crystalline base plane; this plane is the plane
		which creates a crystalline cut; options are a, b, and c,
		or x , y , and z , or $m100$, $m010$, and $m001$ respectively.
periodic	vector	Specifies wether to leave or cut connections through pe-
		riodic boundaries, where vector elements \mathbf{x} , \mathbf{y} , and \mathbf{z} are
		booleans, for which true leaves and false cuts connec-
		tions in the specified direction.

5.5.2 Usage

The crystal command is used to build crystal copies of crystalline unit cells. These unit cells can either contain unconnected or connected molecules. Miller indices in combination with a base plane definition creates the possibility to rotate the final cell to any desired orientation using integer indices h, k, and 1. These indices are defined using the usual conventions. Rotation allows for cleaving a crystal by a desired plane, e.g. miller -> {2, 0, 1} in combination with plane -> m001 uses the plane defined by crystal base vectors \vec{a} and \vec{b} and rotates the cell to satisfy the miller index vector {2, 0, 1}. Box dimensions are altered to form a new unit cell which consequently can be used again as a crystalline building block. The periodic option allows for cutting connections (bonds) through periodic boundaries, which enables the creation of a free structure when using the carve command (see Section 5.3 [Carve], page 88). All vector elements are set to the given input when only one element is set, i.e. periodic -> true equals periodic -> {true, true}.

5.5.3 Default

5.5.4 Examples

```
A few examples are crystal = {
```

5.6 Cut

5.6.1 Syntax

Directive	Parameters	Description
n	integer	Defines the number clusters to be cut in each active system.
mode	option	Selects cutting mode; currently the only option is random.
nmin	integer	Specifies the minimum allowable number of mobile repeat units that remain.
region	region	Specifies in which region in the simulation cell to cut (see Section 7.7 [Region], page 193).

5.6.2 Usage

The cut command finds its main application in cutting a set of crystalline clusters during the preparation of semi-crystalline structures. It is, however, not restricted to crystalline structures alone. The cut command cuts exactly one bond on the specified number of clusters that fall within the specified region. The cutting mode governs the choice of affected clusters.

5.6.3 Default

5.6.4 Examples

```
An example is given by

cut = {
    n -> 20,
    nmin -> 3,
    region -> {shape -> cuboid, radius -> {1/2,1/2,1/4}}
};
```

5.7 Deform

5.7.1 Syntax

deform	= {mode -> constant, type -> constant, system -> constant,
	<pre>frequency -> long, density -> real, geometry -> voigt};</pre>

Directive mode	Parameters constant	Description Sets the deform mode; options are either none or affine for the site deformation mode.
type	constant	Sets the type of geometry; options are either relative for reduced units or absolute for absolute units.
system	constant	Sets identity of system to be deformed; can be either textual or numerical.
frequency	long	Sets the frequency with which to perform incremental deformations; can either be set to instant, forcing instantaneous deformation, or to a positive integer, expressing the step size (in system cycles) in which deformation is gradually applied over the course of a run.
density	real	Sets the final system density; can either be set to calculate when chosing or forced when chosing values > 0.0.
geometry	voigt	Sets the final system geometry; used as a fractional size when density is set, otherwise used as system dimensions from which a density is calculated (see Section 7.15 [Voigt], page 346).

5.7.2 Usage

The deform command is used to gradually deform a selected system to the desired target density over the cause of a run (see Section 5.29 [Run], page 136). Mobile sites are affinely deformed, rigid sites are deformed with respect of the center of mass of the rigid object, and fixed sites stay undeformed.

5.7.3 Default

```
The default is given by
```

5.7.4 Examples

Examples are given by

```
deform = {system -> main, frequency -> 2, density -> 0.7};
deform = {system -> main, frequency -> 2, geometry -> {1, 1, 3}};
```

5.8 Delete

5.8.1 Syntax

Directive mode	Parameters option	Description Sets wether to include or exclude sites in the selected region from deletion; options are include and exclude.
unwrap	boolean	Sets wether to unwrap already existing clusters (only the first instance before a build should be set to true); options are true and false.
fraction	real	Defines the distribution fraction; valid values are between 0 and 1, inclusive the extremes; currently forced to be 1.
focus	struct	Sets the selection to focus on (see see Section 7.3 [Focus], page 178).
inside	struct	Sets the inside region to consider (see see Section 7.7 [Region], page 193).
inside	struct	Sets the outside region to consider (see see Section 7.7 [Region], page 193).

5.8.2 Usage

The delete command is used to delete clusters at the growing sides of a simulation box. Specific selection criteria can be set by defining inside and outside regions in which the center of mass of targetted clusters need to reside. Inside and outside regions can be used to mimic a core-shell approach. A full region is considered when only the outside region is defined. Additionally, a focusing selection can be specified based on site types, group ids, and cluster ids. Undefined focus, inside, outside directives imply considering all active sites in the system.

5.8.3 Default

The default is given by

```
delete = {mode -> include, unwrap -> true, fraction -> 1.0};
```

5.9 Duplicate

5.9.1 Syntax

duplicate = {source -> system, destination -> {min, max}};

Directive Parameters Description

source system Defines the source system.
destination {start, end} Defines the destination systems.

5.9.2 Usage

Take source system system and copy it to destination starting at min and ending at max. Destination systems overlapping with the source system is not allowed.

5.10 Export

5.10.1 Syntax

```
export = {
  profile -> {name -> string, type -> option, style -> option},
  ...
  sample -> {name -> string, type -> option, style -> option},
  ...
};
```

Directive	Parameters	Description
profile	struct	Export settings for profiles.
sample	struct	Export settings for sampled data.
name	string	Defines the file name containing exported data.
type	option	Indicates the type of profile or sample to export; available profiles are angle, bond, bridge, density, force, loop, mass, order, and tail; the available sample is
		cavity.
style	option	Export format; options are csv, json, and math for comma separated value, JSON, and Mathematica formats respectively.

5.10.2 Usage

The export command is used to export data resulting from structure analysis into individual files, using mathematica or comma seperated value formats. Only distributions resulting from analyses are exported.

5.10.3 Default

The export command does not have any defaults.

5.10.4 Examples

Examples of the export command are

```
export = {
  profile     -> {name -> "density", type -> density, style -> math},
  profile     -> {name -> "bond", type -> bond, style -> csv},
     sample     -> {name -> "cavity", type -> cavity, style -> math}
};
```

5.10.5 Data Interpretation

export a distribution by using the export command, e.g.

```
export = {name -> "bridge", type -> bridge, style -> csv};
```

which exports the measured tail distribution to file name bridge.csv in comma separated format. An alternate style is math, which exports to bridge.m in Mathematica format. EMC output format is equivalent to Mathematica format. I typically use the latter, since it allows me to create notebooks for evaluation, which can be used within workflows.

For example, the bridge distribution under the profiles header in the output file contains of a header, followed by a distributions block:

```
profiles = {
  bridge -> {
    active -> true, convolute -> false, direction -> 0,
    nlevels -> 1, binsize -> 1, ndistributions -> 1, distributions -> {
    ncalls -> 101, offset -> 7, nbins -> 645, data -> {
        {n -> 101, accu -> 101, weight -> 101},
        ...
     }
    }
}
```

The header contains general distribution information, i.e. if the distribution is actively sampled, whether to convolute the data (when applicable), in which direction was sampled, how many levels it has, the binsize, and the number of distributions (one for each system). Levels are used to allow for 3D density sampling.

The distributions also contain a header section, which is followed by a data section. The header records for each distribution how many calls a distribution has, what offset the data is recorded at and how many bins the distribution contains. One can construct the x coordinates by combination of the offset with the bin size and the number of bins, i.e.

```
x_i = (i + offset)binsize, with 0 \le i < nbins.
```

The data section entries contain the y coordinates in three contributions: the frequency of calls to this bin, the accumulated scalar data, and the weight with which this data data was recorded. These three numbers are always the same for histograms (e.g. bridge, loop and tail distributions) and represent the frequency of a certain length or size, i.e.

```
y_i = n
```

For order profiles and other slab-wise averages, the accumulated data needs to be divided by their corresponding weight in order to get a local density or order parameter respectively, i.e.

```
y_i = accu/weight
```

For density profiles, the accumulated data needs to be divided by the number of distribution calls (ncalls) in order to get the local density, i.e.

$$y_i = accu/ncalls$$

Interpretation of cavity distributions is identical as for any histogram distribution. See the above description for brigde distributions.

5.11 Field

5.11.1 Syntax

field	= {
id	-> identifier,
mode	-> option,
name	-> {string,},
compress	-> boolean,
apply	-> boolean,
error	-> boolean,
debug	-> boolean,
angle	-> option,
torsion	-> option,
improper	-> option,
increment	-> option,
allow	-> option
} ;	

Directive	Parameters constant	Description Sets the subfield identifier, e.g. used in groups to identify
IU	Constant	the field relevant for typing.
mode	option	Sets execution, input, output, or import mode; EMC standard options are apply, get, and put; options cff, charmm, dpd, martini, and opls import native and nonnative force field formats.
name	string	Sets file names of either input or output; all import options expect a general parameter file followed by an optional topology file.
compress	boolean	Sets compression flag; options are true or false.
apply	boolean	Apply force field typing directly after reading; options are true or false.
error	boolean	Sets error flag; options are true or false; exits on error when true.
debug	boolean	Set debug flag, which controls the output of rule interpretation debugging information; options are true or false.
angle	option	Sets angle field flag; options are ignore, complete, warn, empty, or error (see see Section 7.2 [Field Flags], page 177).
torsion	option	Sets torsion field flag; options are ignore, complete, warn, empty, or error (see see Section 7.2 [Field Flags], page 177).
improper	option	Set improper field flag; options are ignore, complete, warn, empty, or error (see see Section 7.2 [Field Flags], page 177).

increment	option	Set increment field flag; options are ignore, complete,
		warn, empty, or error (see see Section 7.2 [Field Flags],
		page 177).
allow	option	Sets which selection is considered during typing; options
	_	are all or template.

5.11.2 Usage

The field command allows for force field typing. A native format governs storage through get and put modes. The apply mode executes force field typing and should be invoked after clusters assignment only. Currently field supports the Accelrys BIOSYM force field format (mode -> cff) for importing force field typing rules (.dat files) and parameter assignment tables (.frc files). CHARMM, DPD, MARTINI, and OPLS force fields (mode -> charmm, dpd, martini, or opls respectively) are imported using an EMC native textual format (.prm and .top files). A conversion script for CHARMM force fields is given by ./scripts/charmm.pl, for MARTINI force fields by ./scripts/martini.pl, and for OPLS force fields by ./scripts/opls.pl. Force fields can contain templates, which are used to define groups (see Section 5.17 [Groups], page 114) and clusters (see Section 5.4 [Clusters], page 90). Force fields are found in ./field and subsequent directories. Note, that each force field only includes chemistries, for which it was parameterized. This means, that not all chemistries can be typed by all force fields. Also be aware, that typing is not guaranteed to be correct. It is kindly requested to report bugs, so that improvements to force fields can be included in next versions.

Internally, force fields consist of a set of general and specific parameters. General parameters contain automatic parameters, which are used when specific parameters are not defined. Aside from parameters, templates can be defined for commonly used chemistries. Martini force fields use templates to preset the parametrized substances. Note, that the abovementioned Martini script can add types when doubles occur. Here, the use of templates is advised to avoid the selection of wrong types.

5.11.3 Formats

Modes get and put support an EMC-native textual format, which encompasses most possibilities, as needed by supported force fields. Support of leaner textual formats is provided as short hand for ease of use.

The CFF file format is identical to the .frc and .dat as provided by Accelrys' Materials Studio. It has been tested for PCFF and COMPASS formats.

The CHARMM, DPD, MARTINI, and OPLS file formats are created for EMC and function as intermediates between CHARMM, DPD, GROMACS, or OPLS force field families and EMC. Data blocks in the EMC .prm format are flanked by the keywords ITEM and ITEM END. The keyword ITEM is followed by an identifier to designate the following data block. This data block describes data in tab-separated columns. Note, that only exactly one tab is allowed between entries. Spaces and multiple tabs lead to erroneous behavior. The interpretation of the data depends on the ITEM identifier as described in the following table:

ITEM ID	Parameters	Description
DEFINE	See next table	Contains force field definitions and units.

MASS	type mass element nconnections charge comment	Defines masses; columns contain a textual type, a numerical mass value, a textual element, a numerical number of connections, a numerical charge, and a textual comment line respectively.
RULES	<pre>id type element {index residue atom} charge rule</pre>	Defines typing rules; columns contain a numerical id, a textual type, a textual element, a numerical index (e.g. OPLS index), a numerical charge, and a rule in SMILES format respectively; in case of the CHARMM force field, the single numerical index entry is replaced by two entries for residue and atom, respectively.
COMMENTS	id index element comment	Defines comments associated with rules; columns contain a numerical id, a numerical index (e.g. OPLS index), a textual element, and a textual comment respectively.
LITERATURE	year volume page journal	Lists literature references associated with force field; columns contain a numerical year, a textual evolume, a textual page, and a textual journal reference respectively.
EQUIVALENCE	type pair bond angle torsion improper	Tabulates equivalences between types (used to reference a derived type to its original; pair types are considered unique); all columns are in a textual format.
TEMPLATE	name smiles	Lists predefined group and cluster templates; all columns are in a textual format; SMILES format follows the Daylight Chemical Information Systems definition.
PRECEDENCE		Precendences describe the order in which rules are interpreted; parenthesis indicate precedence hierarchy: (? (c $(c1)(c2)$)) c followed by c1 and c2.
NONBOND	type1 type2 sigma epsilon	Defines the interaction parameters for Lennard-Jones 6- 12 pairwise interactions; columns contain two textual types followed by two numerical values.
BOND, BOND_AUTO	type1 type2 k 10	Defines covalent bond length interaction parameters for a harmonic spring; columns contain two textual types followed by two numerical values.
ANGLE, ANGLE_AUTO	type1 type2 type3 k theta0	Defines bond angle interaction parameters for a harmonic spring; columns contain three textual types followed by two numerical values.
TORSION, TORSION_ AUTO	<pre>type1 type2 type3 type4 k n delta [] [index]</pre>	Defines bond torsion interaction parameters for a fourier expansion; columns contain four textual types followed by up to four groups of three numerical values; the last column can contain an optional OPLS reference index.
IMPROPER, IMPROPER_ AUTO	type1 type2 type3 type4 k psi0	Defines improper interaction parameters; columns contain four textual types followed by two numerical values; each of the three possible angle combinations contribute $1/3$ to the total improper energy.

Valid DEFINE IDs are given in the following table:

DEFINE ID	Type	Description
FFNAME	string	Defines force field name.
FFTYPE	string	Defines force field type; options are ATOMISTIC, UNITED, or COARSE.
VERSION	string	Defines force field version.
CREATED	string	Defines parameter file creation date.
LENGTH	string	Sets length unit of parameters in file; options are ANGSTROM, NANOMETER, MICROMETER, METER, or REDUCED.
ENERGY	string	Sets energy unit of parameters in file; options are J/MOL, KJ/MOL, CAL/MOL, KCAL/MOL, or REDUCED.
DENSITY	string	Sets density units of parameters in file; options are G/CC, KG/M^3, or REDUCED.
MIX	string	Sets nonbonded mixing rule; options are NONE, BERTHELOT, ARTIHMETIC, GEOMETRIC, or SIXTH_POWER.
NBONDED	integer	Sets the number of bonds excluded from nonbonded pairwise calculations.
ANGLE	string	Include angle considerations during typing; options are IGNORE, COMPLETE, WARN, EMPTY or ERROR.
TORSION	string	Include torsion considerations during typing; same options as for ANLGE.
IMPROP	string	Include improper considerations during typing; same options as for ANLGE.

The distinction between general and specific force field entries is made by adding extension _AUTO to a parameter keyword. Allowed keywords for this extension are PRECEDENCE, BOND, ANGLE, TORSION, and IMPROPER.

A conversion script for the OPLS force field is provided by ./scirpts/opls.pl, which creates a .prm and .top from OPLS input formats .par and .sb. It is used through

The script assumes the existence of source.par, source.sb, and target.define. The latter is an extra file providing typing rules and possible additions or redefinitions of the OPLS source. The .define files found in ./field/opls/2012/ are currently under development.

A conversion script for the MARTINI force field in GROMACS format is provided by ./scripts/martini.pl. The script is called separately for each contribution through

```
martini.pl name[.itp]
```

The use of multiple topology files within one simulation can be obtained by using a master topology file, which consists of a concatenation of individual topology files. The martini.pl script will create extra types in case of redefinition of bonds, angles, torsions, or impropers.

MARTINI files can be found in ./field/martini/v2.0. A converted copy of the nonbonded parameter file martini.itp is stored as martini.prm in this directory.

A conversion script for the CHARMM force field is provided by ./scirpts/charmm.pl, which creates a .prm and .top from CHARMM input formats .prm and .rtf. It is used through

```
charmm.pl [-source=source] target
```

The script assumes the existence of par_source.prm, top_source.rtf, and target.define. The latter is an extra file providing typing rules and possible additions or redefinitions of the OPLS source. A conversion of all27_prot_lipid with its accompanying .define file as can be found in ./field/charmm/c32b1/. The CHARMM force field interpretation is still in beta phase.

5.11.4 Default

Unless otherwise stated, the default is given by

```
field = {
  mode -> get, compress -> false, error -> true,
  angle -> true, torsion -> true
};
```

5.12 Flag

5.12.1 Syntax

Directive	Parameters	Description
system	id	Sets system to operate on; identifiers as previously defined.
oper	mode	Sets how to operate on site flag; modes are set or unset.
select	mode	Sets selection (used for ignoring head-tail (i.e. ends) connections in crystal builds); modes are all or ends.
flag	constant	Sets site flag; input either one or more flags; options are
		fixed, semi, rigid, or branch.
site	constant	Sets the site types to include; all site types are implied when left blank.
group	constant	Sets the group to include; all groups are implied when left blank.
cluster	constant	Sets the cluster id to include; all cluster ids are implied when left blank.
region	region	Specifies in which region in the simulation cell to cut (see Section 7.7 [Region], page 193).

5.12.2 Usage

The flag command is used to define site-based functionalities through flags. Sites store these flags in a bitwise representation. Possible flag options are given in the table below.

Name	${f Bit}$	Description
fixed	0b0000000001	Prevents the site from being included in energy calcula-
		tions and from being moved during simulation.
semi	0b000000010	Prevents the site from being moved during simulation,
		however, it is included in energy calculations.
head	0b000000100	Defines a site as a head site and is normally not set in
		a global fashion, but only on a per site basis (as used in
		data files).
tail	0b0000001000	Defines a site as a tail site and is normally not set in a
		global fashion, but only on a per site basis (as used in
		data files).
rigid	0b0000010000	Defines a site as part of a rigid body and causes a collec-
		tion of sites to move as one object; only connected sites
		end up in one rigid body when the body flag is not set.

body	0ъ0000100000	Defines a site as being part of a body as defined by the region on the cluster as to which the site belongs; sites will either move inside or on the surface of this body; one cluster represents one body; all rigid sites within this body are treated as one rigid body.
surface	0ъ0001000000	Defines a site to be on the surface of a body rather than in its interior.
backbone	0ъ0010000000	Defines a site to be part of a cluster backbone; backbones are determined internally upon initialization and are able to take part in topology altering moves (e.g. endbridge and reptate moves).
branch	0b0100000000	Defines a site to be part of a cluster branch; branches are excluded from topology altering moves (e.g. endbridge and reptate moves).
unbound	0b1000000000	Defines a site to be part of a cluster branch; branches are excluded from topology altering moves (e.g. endbridge and reptate moves).

5.12.3 Default

The default is given by

5.13 Focus (Command)

5.13.1 Syntax

Directive	Parameters	Description
sites	constant	Sets site or sites to focus on; refers to mass type
		constants.
groups	constant	Sets group or groups to focus on.
clusters	constant	Sets cluster or clusters to focus on.

5.13.2 Usage

The focus command performs a translation to the origin of the accumulative center of mass of all participating sites, groups, and/or clusters. Sites, groups, and clusters are defined by constants as referred to in the simulation constant definition paragraph (see Section 7.1 [Constants], page 176) and described in the focus section (see Section 7.3 [Focus], page 178).

5.13.3 Default

Unless otherwise stated, the default is given by

```
focus = \{\};
```

5.14 Force

5.14.1 Syntax

force	= {style -:	option,	message	->	boolean};
-------	-------------	---------	---------	----	-----------

Directive	Parameters	Description
style	option	Selects force style; valid options are none, init, or list
message	option	Selects output to screen; alid options are none, raw, nkt,
		or n, which corresponds to energy in no particular units,
		internal units, units of nkT or units of n, respectively;
		internal units correspond to the units of the selected force
		field (e.g. kcal/mol for COMPASS, CHARMM, OPLS,
		GROMACS, or MARTINI).

5.14.2 Usage

The force command is used to calculate and possibly output energies and virials associated with all systems in the active simulation. It allows for an internal check of consistency of energies after a simulation by calculation of energies either directly by using the none option, or using the neighbor list resulting from a simulation using the list, or through reinitialization of the neighbor lists by using the init option. The energy output is controlled by the message directive. Energies and virials are reported in either units as defined in the units paragraph when selecting raw, or normalized by the number of active sites when selecting n, or in relative units of nk_BT when selecting nkt.

5.14.3 Default

```
The default is given by
```

```
force = {style -> init, message -> nkt};
```

5.15 Former

5.15.1 Syntax

iormer	=	{name	->	string,	mode	->	option,	forcefield	->	option};

Directive	Parameters	Description
name	string	Defines the file name to be imported; an extention of .emc
		is assumed.
mode	option	Selects i/o mode; the only currently supported option is
		get.
forcefield	option	Selects the desired force field; valid options are none, standard, and charmm.

5.15.2 Usage

The former command is used to import data files generated by previous EMC versions. Currently only the import of version 8 data files is supported.

5.15.3 Default

The default is given by

```
former = {mode -> get, forcefield -> none};
```

5.16 Get

5.16.1 Syntax

get =	{name ->	string,	compress	->	boolean};
-------	----------	---------	----------	----	-----------

Directive name	Parameters string	Description Defines the file name to be imported; an extention of .emc is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.

5.16.2 Usage

The get command is used to import data files generated by the current version of EMC, using a structured data format. Data files function as a restart file and are expected to be in a textual format.

5.16.3 Default

The default is given by

```
get = {compress -> false};
```

5.17 Groups

5.17.1 Syntax

```
groups
              = {
 group
              -> {
   id
              -> constant,
   charge -> real,
charges -> option,
   template
              -> constant,
   terminator -> boolean,
   chemistry
              -> string,
   depth
              -> integer,
   field
              -> constant,
   connects
              -> {{source -> integer,
                   destination -> {index -> constant, site -> integer}
                  }, ...}
 },
 delete
              -> {
              -> constant,
   id
   site
              -> integer
 },
};
```

Directive group id charge	Parameters struct constant double	Description Defines group parameters; repeating structure. Sets group identity; can be either numerical or textual. Sets an additional charge of the complete group, which value divided by the number of group sites is added to the site charges originating from force field or override
charges	option	charge assignments. Defines how partial charges defined in SMILES string are dealt with; valid options are forcefield for using partial charges assigned by a force field only (see Section 5.11 [Field], page 103), additive for adding specified partial charges per SMILES atom to values resulting from force field assignment, and override for only using the specified partial charges (non-specified charges are zero).
template	constant	Selects a template as predefined in force field files; can be either numerical or textual (see Section 5.11 [Field], page 103); templates replace the use of a chemistry string.
terminator	boolean	Indicates if this group can be used as a terminator, which is useful for coarse-grained polymers without termination groups; options are true or false.

chemistry	string	Sets the group's chemistry using the SMILES format (see Section 7.8 [SMILES], page 194).
depth	integer	Sets recursive depth for ring determination during SMILES interpretation; the depth should equal the maximum number of atoms participating in a ring within the given SMILES string; standard option is auto; any set depth smaller than 3 reverts to the auto setting.
field	constant	Optionally set a specific subfield identifier to use when typing this group (see Section 5.11 [Field], page 103).
connects	struct	Defines connections to other groups; both source and site correspond to the position in the SMILES of where a connection occurs; '*', '>', or '<' characters mark connection points; shortcuts head, tail, and \$end1 through \$end10 denote connections in a group withouth having to state their exact positions within the corresponding SMILES; head and tail are analogous to \$end1 and \$end2 respectively (see example below).
source	integer	Sets the source element
destination	struct	Defines the destination group and element
index	constants	Sets the destination group; can be either numerical or textual
site	integer	Sets the destination element
delete	struct	Deletes a specified group site corresponding to a chemistry entry; all corresponding simulation sites are deleted accordingly.
id	constant	Target group identity; can be either numerical or textual.
site	integer	Site number corresponding to the entry position in the SMILES string; counting starts at 0.

5.17.2 Usage

The groups command creates the necessary groups used to identify full molecules or chemical repeat units within clusters, as represented by SMILES strings. Keyword charges is used to assign extra charge to a subset of atoms in case of ionic liquids, for which case a force field might not supply charged molecules. One can also use keyword charge to increase or decrease the charge of a full group. This option assigns a partial charge to each site, which is equal to the full charge divided by the number of sites in the group.

5.17.3 Default

The default is given by

```
groups = {
  group -> {
   id -> 0,
    charge -> 0,
   charges -> forcefield,
```

```
terminator -> false
};
```

5.17.4 Examples

A few atomistic examples are

```
= {
groups
                 -> {
  group
    id
                 -> water,
    chemistry
                -> "HOH"
  },
  group
                 -> {
    id
                 -> dodecane,
                -> "(C)12"
    chemistry
  },
                 -> {
  group
    id
                 -> benzene,
    chemistry
                -> "c1ccccc1"
  group
                -> {
    id
                 -> acid,
                -> "C(=0)[0-]
    chemistry
  }
};
```

An example of deleting a group site is given by

which deletes the first site of group dodecane, effectively changing dodecane into undecane. Note, that all corresponding simulation sites are deleted accordingly.

An example of a set of monomers for a coarse-grained polymer is given by

```
= {
groups
                -> {
 group
    id
                -> monomer1,
                -> "*ab*c(d)cc",
    chemistry
    connects
                -> {
      {head, {monomer1, tail}}, {head, {monomer2, head}},
      {tail, {monomer1, head}}, {tail, {monomer2, head}},
      {head, {terminator, head}}, {tail, {terminator, head}}}
 },
                -> {
  group
```

```
id
                -> monomer2,
                -> "*c*",
    chemistry
                -> {
    connects
      {head, {monomer1, head}}, {head, {monomer1, tail}},
      {head, {monomer2, head}}, {head, {monomer2, tail}},
      {head, {terminator, head}}}
 },
                -> {
 group
    id
                -> terminator,
                -> "*t",
    chemistry
    connects
                -> {
      {head, {monomer1, head}}, {head, {monomer2, head}}}
 },
};
```

The '*' character implies the start and end or head and tail of a repeat unit of a polymer. By default, a group is not a terminator when more than one '*' character appears in the SMILES string. However, when such a group should also function as a terminator, the terminator flag can be used as indication of such. A group is automatically a terminator when a '*' character appears only once, which means, that the terminator flag need not be set. The keyword connects lists the possible connections of these start and end sites to sites in the same or other groups. The keywords head and tail represent internal short cuts and indicate the position of the '>' and '<' characters repectively.

5.18 Insight

5.18.1 Syntax

Directive	Parameters	Description
name	string	Defines the file name to be imported; an extention of .emc
		is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an
		extra extension of .gz is assumed; options are true or
		false.
system	integer	Selects system to be imported or exported
mode	option	Selects i/o mode; supported options are get or put.
forcefield	option	Selects the desired force field for writing; see Section 7.5
		[Ports], page 191, for valid options; option auto auto-selects.
atomistic	option	Select the atomistic mode normally defined by the choice
		of forcefield; options are none, united, or full.
charges	boolean	Use partial charges from imported structures; options are
		true or false.
formal	boolean	Use formal charges from imported structures; options are
		true or false.
detect	boolean	Detect and assign atom and amino acid residue types;
		options are true or false.
depth	integer	Sets recursive depth for ring determination during
		SMILES interpretation; the depth should equal the max-
		imum number of atoms participating in a ring within the
		given SMILES string; standard option is auto; any set
ou+	boolean	depth smaller than 3 reverts to the auto setting. Selects cut mode: cut bonds that cross cell boundaries;
cut	poorean	options are true or false.
pbc	boolean	Applies periodic boundary conditions; options are true
рьс	boolean	or false.
map	boolean	Selects mapping sites and box geometry according to a
		minimum image convention; options are true or false.
unwrap	boolean	Unwraps clusters to cross periodic boundaries with ap-
		plication of periodic boundary conditions to its center of
		mass when applicable; options are true or false.

crystal	boolean	Imported structure is crystalline; options are true or
		false.
percolate	boolean	Allow for percolating structures; options are true or
		false.
flag	struct	Allows for setting system flags while importing morpholo-
		gies; see see Section 7.10 [System Flags], page 198, for the
		various available flags.

5.18.2 Usage

The insight command is used to import and export InsightII data files in CAR and MDF format for exchange of configurations between EMC and Materials Studio, a product of Accelrys¹.

5.18.3 Default

```
The default is given by
```

5.18.4 References

1. Materials Studio, 'http://www.accelrys.com/'

5.19 Lammps

5.19.1 Syntax

lammps	= {name -> string, compress -> boolean,
	system -> integer, mode -> option,
	units -> option, length -> double, forcefield -> option,
	shake -> option, atomistic -> boolean, cutoff -> boolean,
	charges -> boolean, ewald -> boolean,
	bonds -> boolean, types -> boolean,
	parameters -> boolean, cross -> boolean,
	variables -> boolean, coefficients -> boolean,
	comment -> boolean, map -> boolean, unwrap -> boolean,
	<pre>version -> integer, flag -> structure};</pre>

Directive	Parameters	Description
name	string	Defines the file name to be imported; an extention of .data is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.
system	integer	Selects system to be imported or exported
mode	option	Selects i/o mode; supported options are get or put.
length	double	Defines a scaling length to superceede the simulation variable units -> angstrom; the latter is used when length is not specified (see Section 7.13 [Units], page 343).
units	option	Selects wether to represent site coordinates in reduced or real units, for which options are lj or real respectively; alternatively none is also defined.
forcefield	option	Selects the desired force field; see Section 7.5 [Ports], page 191, for valid options.
shake	option	Selects the SHAKE ^{1,2} mode for atomistic force fields; valid options are none, auto, hydrogen, water and all; auto bases the choice on the selected force field (none by default).
atomistic	boolean	Selects atomistic mode: no charge column and no bonds; options are true or false.
cutoff	boolean	adds a column containing cut offs as used for EMC force field evaluation to the LAMMPS parameter file; helpful for transferring purely repulsive potentials; options are true or false.
charges	boolean	Selects charge mode: a charge column is present in the Atoms paragraph; options are true or false.
ewald	boolean	Selects the usage of Ewald summations for long-range charge treatment; options are true or false.

bonds	boolean	Selects bonds mode: all bonded contributions are present; options are true or false.
types	boolean	Selects types mode: all types parameter definitions are present; options are true or false.
parameters	boolean	Selects the separate output of variables and (non)bonded interaction parameters in LAMMPS input script format; an extension of .params is assumed; options are true or false.
cross	boolean	Includes nonbonded cross-terms in .params file; options are true or false.
variables	boolean	Selects adding simulation variables to the separate output of variables and interaction parameters; options are true or false.
coefficients	boolean	Selects adding interaction parameters to the separate output of variables and interaction parameters; options are true or false.
comment	boolean	Includes commented references to mass names in .data and .params files; options are true or false.
map	boolean	Selects mapping sites and box geometry according to a minimum image convention; options are true or false.
unwrap	boolean	Selects unwrapping of clusters; options are true or false.
version	integer	Set LAMMPS version.
flag	structure	Set system flags.
•		-

5.19.2 Usage

The lammps command is used to import and export data files generated by LAMMPS, a Molecular Dynamics code conceived at Sandia National Laboratories.^{3,4} Textual input formats for LAMMPS are ambiguous and are not self-defining, which creates the need for predefinition of the desired format. This port allows for commonly used force field modes and site (atom) definitions.

5.19.3 Default

The default is given by

5.19.4 Examples

A few examples are given by

```
lammps = {name -> "benzene", compress -> true,
```

5.19.5 References

- 1. Ryckaert, J.-P.; Ciccotti, G. and Berendsen, H.J.C., "", *J. Comp. Phys.* **1977**, *23*, 327-341
- 2. Andersen, H. "", J. Comp. Phys. 1983, 52, 24-34
- 3. LAMMPS Molecular Dynamics Simulator, 'http://lammps.sandia.gov/'
- 4. Plimpton, S., "Fast Parallel Algorithms for Short-Range Molecular Dynamics", *J. Comput. Phys.* **1995**, *117*, 1-19

5.20 Memory

5.20.1 Syntax

```
memory = {style -> option};
```

Directive Parameters Description

style option Sets the output style; options are full or summary.

5.20.2 Usage

The memory command outputs the memory consumption of the current simulation to screen. Either a summary or a full description can be selected.

5.20.3 Default

The default is given by

```
memory = {style -> full};
```

5.20.4 Examples

An example is given by

```
memory = {style -> summary};
```

5.21 Moves

5.21.1 Syntax

```
moves = {
  ncycles -> integer,
  cycle -> integer,
  move -> integer,
  moves...
};
```

Directive	Parameters	Description
ncycles	integer	Sets the total number of simulation cycles.
cycle	integer	Sets the current simulation cycle
move	integer	Sets the current simulation move.
moves		Access to various move settings; see Section 7.4 [Moves],
		page 179, for further information.

5.21.2 Usage

This variable style describes types (see Section 7.4 [Moves], page 179, for further information) and allows direct access to all variables and parameters stored within the moves structure.

5.21.3 Default

Unless otherwise stated, the default is given by

```
Unless otherwise stated, the default is given by
@verbatim
moves = {
  ncycles    -> 0,
  cycle    -> 0,
  move    -> 0
};
```

By default, all moves are activated with zero frequency, with the exception of the displacement move (see Section 7.4.6 [Displace], page 182), which has a frequency of one.

5.22 PDB

5.22.1 Syntax

Directive	Parameters	Description
name	string	Defines the file name to be imported; an extention of .emc
		is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; unpacking and packing is added to the VMD script; options are true or false.
mode	option	Selects i/o mode; supported options are get and put
system	integer	Selects system to be imported or exported.
length	real	Set the length scale with which to scale the resulting co- ordinates (normmaly results from choice of forcefield).
forcefield	option	Selects the desired force field; see Section 7.5 [Ports], page 191, for valid options; option auto auto-selects.
atomistic	option	Select the atomistic mode normally defined by the choice
		of forcefield; options are none, united, or full.
charges	boolean	Use charges from imported structures; options are true
		or false.
depth	integer	Sets recursive depth for ring determination during SMILES interpretation; the depth should equal the maximum number of atoms participating in a ring within the given SMILES string; standard option is auto; any set depth smaller than 3 reverts to the auto setting.
detect	boolean	Detect and assign atom and amino acid residue types; options are true or false.
atom	option	Sets options for atom name representation upon output; options are detect for detection of atom types, index for use of EMC mass element ids, or series for indexing using EMC element ids followed with sequential numbering per segment. The latter is useful for CHARMM tools expecting unique atom ids.

residue	option	Sets options for residue name representation upon output; options are detect for detection of amino acid residues, index for use of EMC group ids, or series for increments starting with the letter R.
segment	option	Sets options for segment name representation upon output; options are detect for detection, index for use of EMC cluster ids, or series for increments starting with the letter M.
vdw	boolean	Include the definition for switching on Van der Waals in the resulting VMD script; options are true or false.
hexadecimal	boolean	Use hexadecimal representations for residue sequence numbers; options are true or false.
cut	boolean	Selects whether cut bonds that cross cell boundaries upon output; options are true or false.
vdw	boolean	Add Van der Waals representation to the written VMD script; options are true or false.
pbc	boolean	Applies periodic boundary conditions upon output; options are true or false.
map	boolean	Selects mapping sites and box geometry according to a minimum image convention; options are true or false.
unwrap	boolean	Unwraps clusters upon output to cross periodic boundaries with application of periodic boundary conditions to its center of mass when applicable; options are true or false.
rigid	boolean	Unwrap sites in clusters, that are flagged as rigid; options are true or false.
fixed	boolean	Unwrap sites in clusters, that are flagged as fixed; options are true or false.
connectivity	boolean	Include connectivity in the resulting PDB using keyword CONECT; options are true or false.
crystal	boolean	Selects whether a read structure is a crystal; needed for small crystal structures with box crossing bonds; options are true and false.
element	option	Selects location of element (internally used for defining masses); options are auto, element, and type; element uses elements defined in PDB files; aromatic elements are all lower case (needed when reading united atom structures); type uses types as defined in PSF files, which is needed when reading coarse-grained structures; auto internally decides which of the previous options to use based on the choice of force field (see Section 5.11 [Field], page 103).

5.22.2 Usage

The pdb command is used to export data files in PDB and PSF format for convenient visualization with VMD (Visual Molecular Dynamics), a visualization conceived and supported by the Theoretical and Computational Biophysics Group at the University of Illinois at Urbana-Champaign. Textual PDB and PSF formats are ambiguous and are not self-defining, which creates the need for predefinition of the desired format. This port allows for commonly used force field modes. Cell dimensions are defined by addition of a CRYST1 keyword to the PDB file.

5.22.3 Default

The default is given by

5.22.4 References

- 1. VMD Visual Molecular Dynamics, 'http://www.ks.uiuc.edu/Research/vmd/'
- 2. Humphrey, W., Dalke, A. and Schulten, K., "VMD Visual Molecular Dynamics", J. Molec. Graphics 1996, 14, 33-38

5.23 Put

5.23.1 Syntax

put =	{name ->	string,	compress ->	boolean,	detail ->	<pre>integer};</pre>
-------	----------	---------	-------------	----------	-----------	----------------------

Directive name	Parameters string	Description Defines the file name to be exported; an extention of .emc is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.
detail	integer	Sets level of detail in output file; a non-zero setting over- rides the already set detail.

5.23.2 Usage

The put command is used to export data files generated by the current version of EMC, using a structured data format. Data files contain all necessary information to function as a restart file and are in a textual format.

5.23.3 Default

The default is given by

```
put = {compress -> false, detail -> 0};
```

5.24 Rename

5.24.1 Syntax

Directive	Parameters	Description
site	struct	Defines multiple sites.
group	struct	Defines multiple groups.
cluster	struct	Defines multiple clusters.
system	struct	Defines multiple systems.

5.24.2 Usage

The rename command allows for renaming already existing site, group, cluster, or system ids

5.24.3 Default

No default.

5.24.4 Examples

A few examples are given by

5.25 Remove

5.25.1 Syntax

Directive	Parameters	Description
n	integer	Selects number repeat units to be removed in each active system.
mode	option	Selects removal mode; viable options are cluster, group, and site.
target	index	Specifies the targeted group to remove; additionally a group site can be specified; a value of -1 one indicates all.
nmin	integer	Specifies the minimum allowable number of mobile repeat units that remain.
region	region	Specifies in which region in the simulation cell to remove (see Section 7.7 [Region], page 193).

5.25.2 Usage

The remove command deletes a maximum of n and a minimum of nmin groups specified by group, which fall within a specified region. All sites within the specified region are deleted, when no group is provided.

5.25.3 Default

5.25.4 Examples

Remove 100 water molecules as defined by group water in the whole simulation box,

Note, that overspecification by the region radius of a factor of two ensures the inclusion of all particles in that box direction.

Remove 10 methyl groups as defined by group methyl in the center halve portion in the x direction of the simulation box,

5.26 Reset

5.26.1 Syntax

reset = {style -> option};

Directive Parameters Description

style integer Possible options are simulation, statistics,

profiles, or moves.

5.26.2 Usage

The reset command allows for resetting entries and counters of either the whole simulation structure, or of separate subsections statistics, profiles, or moves.

5.26.3 Default

The default is given by

reset = {style -> simulation};

5.27 Restart

5.27.1 Syntax

Directive name	Parameters string	Description Defines the file name to be exported; an extention of .emc is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.
format frequency	string integer	Defines counter extention format. Selects output interval frequency in units of cycles.
reset	boolean	Resets simulation after writing restart file; options are true or false.

5.27.2 Usage

The restart command allows for writing of restart files at constant intervals during simulation, which safeguards long simulations in case of compute environment failure.

5.27.3 Default

The default is given by

5.28 Retype

5.28.1 Syntax

Directive	Parameters	Description
mode	option	Sets the selection criteria for the fraction of selected sites; valid modes are: random.
charge	boolean	Selects charge transfer from mass paragraph to sites.
fraction	real	Defines the fraction of sites within the selection to be changed; values smaller than zero will be set to zero; values larger than one will be set to one; non-zero number of sites overrides fractions.
n	integer	Sets the number of sites to delete per defined targeted system; the first entry is taken for all systems when targeted systems are omited; fractions are taken when omited.
system	constant	Sets targeted system; can one or more systems; all systems are implied when omited.
source	constant	Sets source mass types; can be one or more types.
destination	constant	Sets destination mass types; can be one or more types; has to be either one element or the same number of elements as represented under the source directive.
group	index	Optional definition of group index and site for all destination types.
inside	region	Sets the inner boundary of the region in which mass types are to be changed (see Section 7.7 [Region], page 193); assumes domain center when ommited.
outside	region	Sets the outer boundary of the region in which mass types are to be changed (see Section 7.7 [Region], page 193); assumes whole domain when ommited.

5.28.2 Usage

The retype command allows for changing mass types from a source target consisting of either one or more types to a destination target consisting of one or more types. The destination target can either consist of one element or the same number of elements as the source target, in case the source target consists of more than one element. All groups, systems and regions are implied when none are supplied.

5.28.3 Default

5.28.4 Examples

Note that the new site 'graft' needs to be defined in terms of mass and force field (see Section 5.32 [Sites], page 139, and Section 5.33 [Simulation], page 140). An example is given by

5.29 Run

5.29.1 Syntax

Directive ncycles	Parameters integer	Description Defines the number of cycles for which to run the simulation.
nblocks	integer	Defines the frequency with which to output intermediate energies in intervals of cycles.
cycle	integer	Defines the cycle counter at which to start the simulation; selects the cycle as defined by the input file when negative
seed	integer	Selects a random seed; the time is used when seed is greater than or equal to zero.

5.29.2 Usage

The run command is used to start a simulation. It allows for definition of simulation length and output frequency. Starting cycle and initial random seed can also be defined.

5.29.3 Default

The default is given by

```
run = \{ \text{cycle} \rightarrow -1, \text{ seed } \rightarrow 0 \};
```

5.30 Sample

5.30.1 Syntax

Directive	Parameters	Description
cavity	struct	Descriptor for cavity size sampling as defined by CESA
		(Cavity Energetic Sizing Algorithm) ¹ (see Section 6.2
		[Cavity], page 157).
gr	struct	Descriptor for radial distribution functions sampling (see
		Section 6.3 [Gr], page 162).
gyration	struct	Descriptor for radius of gyration sampling (see Section 6.4 [Gyration], page 167).

5.30.2 Usage

This variable style describes the control of sampling options (see Chapter 6 [Sampling Tools], page 153, for applications).

5.30.3 Default

By default, all sampling is deactivated.

5.30.4 References

1. P.J. in 't Veld, M.T. Stone, T.M. Trustkett, and I.C. Sanchez, "Liquid Structure via Cavity Size Distributions", J. Phys. Chem. B 2000, 104, 12028

5.31 Shell

5.31.1 Syntax

shell = {command -> string};

Directive Parameters Description

command string Execute a shell command as specified by string.

5.31.2 Usage

The shell command is used to execute a shell command, which allows for file manuipulations during running. Any problems will be identified by a warning, thus not stopping the execution of the EMC script.

5.31.3 Default

The default is given by

shell = {command -> ""};

5.32 Sites

5.32.1 Syntax

Directive	Parameters	Description
site	struct	Identifies site parameters; repeating structure.
id	constant	Sets site identity; can be either numerical or textual.
reference	integer	Sets a numerical reference; not used internally.
name	string	Sets a longer identifying name.
mass	real	Sets the mass in units set in units.

5.32.2 Usage

The sites command creates mass entries in the mass section of the types section of simulation. Multiple sites can be entered by using multiple instances of the site directive, separated by commas.

5.32.3 Default

```
The default is given by sites = {site -> {id -> 0, reference -> 0}};
```

5.32.4 Examples

```
A few examples are
sites
                -> {id -> a, reference -> 0, name -> "colloid", mass -> 10}
  site
};
                = {
sites
                -> \{id -> c,
  site
                      reference -> 12, name -> "carbon", mass -> 12.011},
                -> \{id -> h,
  site
                       reference -> 1, name -> "hydrogen", mass -> 1.008},
                -> {id -> o,
  site
                      reference -> 14, name -> "oxygen", mass -> 15.9994}
};
```

5.33 Simulation

5.33.1 Syntax

Directive	Parameters	Description
output	{}	Description Provides access to output parameters; possible directives
•		include detail, wide, expand, math, reduced, info,
		strict, warning, message, and debug.
units	{}	Provides access to internal units; possible directives
		include mass, length, angstrom, angle, energy, kb,
		nav, charge, permittivity, and seed (see Section 7.13
		[Units], page 343).
variables	{}	Provides access to changing predefined variables; used
		in conjunction with predefined variables in input files, allowing posteriori sizing of an input structure.
tunos	{}	Provides access to all simulation-wide types; possible
types	()	directives, which define force fields and their respec-
		tive constants, include boltzmann, charmm, coarse,
		coulomb, spline, and standard (see Section 7.12
		[Types], page 201).
systems	{}	Provides access to all simulation-wide system settings;
		possible directives, which define e.g. system temperature
		or geometry, include properties; most system properties
		are derived output variables (e.g. p, v, mass, nsites,
		nclusters) rather than input variables (see Section 7.11
	r	[Systems], page 199).
moves	{}	Provides access to all moves; possible directives include displace, endbridge, rebridge, reptate, rotate, and
		temper (see Section 7.4 [Moves], page 179).
profiles	{}	Provides acces to all profile definitions; possible directives
•	- -	include density, force, mass, and order (see Section 7.6
		[Profiles], page 192).

5.33.2 Usage

The simulation command provides access to all variables and parameters defined within the simulation structure, of which the above table lists the main directives.

5.33.3 Examples

A few examples are (see data file for more suggestions)

```
simulation = {output -> {detail -> 4}};
simulation = {variables -> {lb -> 0.95*lb}};
```

5.34 Split

5.34.1 Syntax

Directive system	Parameters constant	Description Sets identity of system to which to add the build; can be either textual or numerical.
direction	option	Sets the direction in which the system is grown; options are x, y, and z.
mode	option	Sets the selection algorithm; options are distance and random.
unwrap	boolean	Sets wether to unwrap already existing clusters (only the first instance before a build should be set to true); options are true and false.
fraction	real	Defines the distribution fraction; valid values are between 0 and 1, inclusive the extremes.
focus	struct	Sets the selection to focus on (see see Section 7.3 [Focus], page 178).
region	struct	Sets the region to consider (see see Section 7.7 [Region], page 193).

5.34.2 Usage

The split command is used to redistribute clusters at the growing sides of a simulation box. Specific selection criteria can be set by defining a region in which the center of mass of targetted clusters need to reside. Additionally, a focusing selection can be specified based on site types, group ids, and cluster ids. Undefined focus and region directives imply considering all active sites in the system.

5.34.3 Default

The default is given by

5.35 Terminate

5.35.1 Syntax

terminate = {mode -> option};

Directive Parameters Description

mode option Selects termination mode; currently only the option all

is supported.

5.35.2 Usage

The terminate command is used to terminate free cluster ends created by cut and remove commands. Currently, terminators existing of only one site are supported. Future releases will include a group-based terminator.

5.35.3 Default

The default is given by

terminate = {mode -> all};

5.36 Timing

5.36.1 Syntax

timing = {style -> option};

Directive Parameters Description

style option Possible options are none, show, or reset.

5.36.2 Usage

The timing command allows for showing current timing, after which timing can be reset or allowed to accumulate further.

5.36.3 Default

The default is given by

timing = {style -> show};

5.37 Traject

5.37.1 Syntax

Directive	Parameters	Description
mode	option	Sets the operation mode; viable options are get, put, and sample.
system	integer	Selects the system to export.
name	string	Defines the file name to be exported; an extention of .traject is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.
format	string	Defines counter extention format.
start	integer	Selects the frame at which to start reading; -1 indicates the last frame.
end	integer	Selects the frame at which to end reading; -1 indicates the last frame.
frequency	integer	Selects output interval frequency in units of cycles.
append	boolean	Appends each selected cycle to the end of the trajectory file; options are true or false.
reset	boolean	Resets output file; options are true or false.
scale	boolean	Scales the output using system box dimensions; options are true or false.
unwrap	boolean	Unwraps clusters in output; options are true or false.
pbc	boolean	Applies periodic boundary conditions; options are true or false.

5.37.2 Usage

The traject command allows for reading, writing, or analyzing of LAMMPS-style trajectory files. In read mode, a configuration from a trajectory can be loaded over already existing positions. In write mode, configurations are written at constant intervals during simulation. Output can be directed to either one or separate files by setting the append option to true or false respectively. Cycle counts are added to the file name when separate files are chosen, however they are omitted when the append option is set. In analyze mode, a full trajectory is analyzed using preset sampling settings (see Section 5.30 [Sample], page 137). An example can be found in ./examples/sample/cavity/traject.emc.

5.37.3 Default

The default is given by

5.37.4 Examples

A few examples are given by

5.38 Translate

5.38.1 Syntax

Directive	Parameters	Description
sites	constant	Sets site or sites to focus on; refers to mass type
		constants.
groups clusters	constant	Sets group or groups to focus on.
delta	vector	Sets the displacement vector

5.38.2 Usage

The translate command facilitates translation of a selection of sites over a displacement vector delta. This scripting command can be helpful in translating surfaces constructed in the center of the simulation box to its edges.

5.38.3 Default

5.38.4 Examples

which translates clusters called 'surface' to the lower edge of the simulation box. Note, that \mathtt{lx} has to be provided by the user.

5.39 Types

5.39.1 Syntax

Directive	Parameters	Description
merge	boolean	Allows for merging force field constants upon input when
		true; options are true or false.
virial	boolean	Describes if virial calculations are included; options are
		true or false.
periodic	vector	Indicate periodicity with a three-element boolean vector
		with options true or false.
neighbor	constant	Describes what kind of neighbor list algorithm is used
		during pair interaction calculations; options are sector
		or pair.
stencil	constant	Describes the kind of stencil used during pair interaction
		calculations; options are standard or multi.
skin	real	Describes the skin used during pair interaction calcula-
		tions; the skin is added to the pairwise cutoff.
shake	constant	Indicates the use of the SHAKE algorithm in subsequent
		codes (e.g. LAMMPS); valid options are none, auto,
		hydrogen, water, or all.
depth	integer	Maximum depth used for construction of ring structures
		during typing; allowed values are positive, where a value
		of 8 works in most ring cases; alternatively, an auto key-
		word allows for checking rings of unknown size; please
		note, that significant slow down occurs with the latter
		options for intricate ring systems.
mass	struct	Describes the site masses.
fields		Access to various field settings; see Section 7.12 [Types],
		page 201, for further information.

5.39.2 Usage

This variable style describes types (see Section 7.12 [Types], page 201, for further information) and allows direct access to all variables and parameters stored within the types structure.

5.39.3 Default

Unless otherwise stated, the default is given by

```
= {
types
 merge
                -> false,
 virial
                -> false,
                -> {true, true, true}
 periodic
 neighbor
                -> sector,
 stencil
                -> standard,
  shake
                -> none,
 depth
                -> auto,
 skin
                -> 0
};
```

By default, all force fields are deactivated.

5.40 Variables

5.40.1 Syntax

variables = {variable -> string|integer|real, ...};

Directive	Parameters	Description
variable	${\tt string},$	Sets or redefine a variable; currently supported format
	integer, real	only include scalars of types string, integer, and real.

5.40.2 Usage

The variable command is used to define or override previously defined variables, which allows for e.g. resizing of simulation cells as defined in the original data file. Also, simulation-wide string variables can be set, which later can be used as file names, etc. Currently, only scalar variables are supported. Future releases will include structured variables as well.

5.40.3 Examples

Possible definitions include

```
variables = \{la \rightarrow 0.95*la, lb \rightarrow 1.05*lb\};
variables = \{name \rightarrow "test/polyethylene"\};
```

5.41 XYZ

5.41.1 Syntax

Directive	Parameters	Description
name	string	Defines the file name to be imported; an extention of .emc
		is assumed.
compress	boolean	Sets compression using Lempel-Ziv coding (LZ77); an extra extension of .gz is assumed; options are true or false.
	:	
system	integer	Selects system to be imported or exported
forcefield	option	Selects the desired force field; valid options are none, standard, or charmm.
cut	boolean	Selects cut mode: cut bonds that cross cell boundaries; options are true or false.
detect	boolean	Detect and assign atom and amino acid residue types;
detect	boolean	options are true or false.
segment	option	Sets options for segment representation; options are detect for detection, index for use of EMC cluster ids, or series for increments starting with the letter M.
residue	option	Sets options for residue representation; options are detect for detection of amino acid residues, index for use of EMC group ids, or series for increments starting with the letter R.
pbc	boolean	Applies periodic boundary conditions; options are true or false.
map	boolean	Selects mapping sites and box geometry according to a minimum image convention; options are true or false.
unwrap	boolean	Unwraps clusters to cross periodic boundaries with application of periodic boundary conditions to its center of mass when applicable; options are true or false.

5.41.2 Usage

The xyz command is used to export data files in XYZ format for convenient visualization with VMD (Visual Molecular Dynamics), a visualization conceived and supported by the Theoretical and Computational Biophysics Group at the University of Illinois at Urbana-Champaign. Textual XYZ formats are ambiguous and are not self-defining, which creates the need for predefinition of the desired format. This port allows for commonly used force field modes.

5.41.3 Default

The default is given by

5.41.4 References

- 1. VMD Visual Molecular Dynamics, 'http://www.ks.uiuc.edu/Research/vmd/'
- 2. Humphrey, W., Dalke, A. and Schulten, K., "VMD Visual Molecular Dynamics", J. Molec. Graphics 1996, 14, 33-38

6 Sampling Tools

EMC offers the possibility of applying sampling either on the fly or by application to pregenerated trajectories (in .traject format). Available sampling algorithms include radii of gyration, pair correlation functions, cavity size distributions, energy and density profiles. The latter profiles offer a convolution option, which promotes smoothing out results by spreading point measurables over spherical volumes. Sampling is accessed through the scripting command sample (see Section 5.30 [Sample], page 137). Examples can be found in ./examples/build/sample.

6.1 Bond

6.1.1 Syntax

```
gyration
                = {
 id
                -> integer,
                -> boolean,
  active
                -> integer,
 frequency
                -> struct,
  source
 target
                -> struct,
 binsize
                -> real,
 distributions -> struct
};
```

Directive id active	Parameters integer boolean	Description Method identifier for referencing subsequent alterations. Indicates the state of the method; options are true or false.
frequency	integer	Sets the frequency of the method, which is assumed to be a positive integer.
source, target	struct	Definition of clusters, groups, and sites as to which to focus analysis on (see Section 7.3 [Focus], page 178); both contributing types need to be included for a specific bond; all mobile and active sites are taken when not defined.
binsize	real	Sets the binsize of the resulting distributions.
distributions	struct	Distributions resulting from analysis; these distributions - only shown in .emc and exported .m files - cannot be influenced with scripting command sample.

6.1.2 Usage

The bond command is used to sample bond lengths, which are indicated by defining pairs withing focus. Resulting distributions are reported as frequency distributions, which can be exported by using the scripting command export (see Section 5.10 [Export], page 100). Averages are reported as part of the resulting distribution as shown in .emc and .m files. An example can be found in ./examples/build/sample/bond/ by running setup.sh.

6.1.3 Default

};

```
The default is given by

sample = {
    id -> 0,
    active -> false,
    frequency -> 1,
    source -> {},
    target -> {},
    binsize -> 0.01
```

6.1.4 Example

The example as provided in ./examples/sample/bond/ creates a system of octane molecules using the TraPPE-UA force field (see Section 7.12.19 [TraPPE], page 335). The EMC build script is generated by emc_setup.pl, which uses chemistry.csv to define the octane chemistry (see Chapter 4 [Simulation Setup], page 8). The final system contains a total of 1000 beads. The setup.sh calls EMC to build and sample. The sampling script sample.emc is explained in the following. First, a set of variables are set, which are used later in the script.

Next, the created structure example.emc.gz is loaded.

```
get = {name -> input, compress -> true};
```

The structure should first be equilibrated before sampling commences. To this effect, a simulation is executed for ncycles cycles.

```
run = {
  ncycles -> ncycles,
  nblocks -> nblocks
};
```

The bond lengths of bonds between CH3 and CH2 types and between CH2 types are to be sampled. To this end, sampling of bonds in defined in the sample paragraph. The TraPPE type for CH3 groups is c4h3 and for CH2 groups c4h2.

```
= {
sample
                  -> {
  bond
    {
       id
                  -> 0,
                  \rightarrow {sites \rightarrow c4h3},
       source
                  \rightarrow {sites \rightarrow c4h2},
      target
                  -> true, frequency -> frequency, binsize -> 0.001
       active
    },
    {
       id
                  -> 1,
                  -> {sites -> c4h2},
       source
      target
                  -> {sites -> c4h2},
                  -> true, frequency -> frequency, binsize -> 0.001
       active
```

```
}
};
```

The simulations is continued for another ncycles. Sampling occurs during the execution of the simulation.

```
run = {
  ncycles -> ncycles,
  nblocks -> nblocks
};
```

Trajectories can also be used for sampling analysis. To this effect, a trajectory can be loaded by using the traject keyword.

```
traject = {name -> name, frequency -> 1, mode -> analyze};
```

Sampling of trajectories occurs when the mode option analyze is selected. The resulting distributions can be exported to either math and csv formats.

```
export = {
  sample -> {name -> "bond", type -> bond, style -> math},
  sample -> {name -> "bond", type -> bond, style -> csv}
};
```

The equilibrated and resulting structure is stored in EMC format by means of

```
put = {name -> output, compress -> true};
```

6.2 Cavity

6.2.1 Syntax

cavity	= +	[
id	->	integer,
active	->	boolean,
frequency	->	integer,
focus	->	struct,
solver	->	option,
record	->	string,
separate	->	boolean,
ninserts	->	integer,
ntrials	->	integer,
npoints	->	integer,
tolerance	->	real,
cutoff	->	real,
binsize	->	real,
distributions	->	struct,
nerrors	->	struct,
zero	->	boolean,
negative	->	boolean
} ;		

Directive id active	Parameters integer boolean	Description Method identifier for referencing subsequent alterations. Indicates the state of the method; options are true or false.
frequency	integer	Sets the frequency of the method, which is assumed to be a positive integer.
focus	struct	Defines sites, groups, or clusters to focus on (see Section 7.3 [Focus], page 178).
solver	option	Selects the solver used in finding local minima or saddle points in the energy surface; available options are broyden or newton.
record	string	Sets the output file name for the positions ans sizes of found cavities; ignored when no file name is set.
separate	boolean	Toggles whether to write recorded output in one single or separate files; options are true or false.
ninserts	integer	Sets the number of initially inserted points for finding local minima or saddle points.
niterations	integer	Sets the maximum number of iteration steps allowed in finding the local minima or saddle points.
npoints	integer	Sets the number of points used to determine the cavity size by means of the root of the local energy.

tolerance	real	Sets the tolerance for the zero force at the local minimum or saddle point.
cutoff	real	Sets the cutoff in Jacobian elements used to discriminate between minima and saddle points.
binsize	real	Sets the binsize of the resulting distributions.
distribution	s struct	Distributions resulting from analysis; local minima and saddle points distributions are reported per system in alternating fashion.
nerrors	struct	Resulting bookkeeping of algorithm performance.
zero	boolean	Include negative sizes as zero size in distribution; negative sizes are ignored when false; superceedes negative when true; options are true or false.
negative	boolean	Include negative sizes in distribution; options are true or false.

6.2.2 Usage

The cavity command is used to sample cavity size distributions of all morphologies entailed by a simulation according to the algorithm as described by In 't Veld et al.[1] The algorithm uses either a Newton or Broyden iterative scheme in combination with a line search to determine local minima or saddle point in the surface of the energy landscape. This point is then used to grow a bead to such an extent, that the resulting energy equals zero. Distinction between minima and saddle points is made by requiring the Jacobian to be definate positive for minima. The definition of "definate positive" can be adjusted by means of the cutoff option. An example can be found in ./examples/build/sample/cavity/ by running build.emc. An example of trajectory analysis is given by traject.emc in the same examples directory.

6.2.3 Default

The default is given by

```
sample
  active
                 -> false,
  frequency
                 -> 1,
  focus
                 -> {},
  solver
                 -> newton,
                 -> "",
  record
  separate
                 -> false,
                 -> 1000,
 ninserts
                 -> 200,
  ntrials
                 -> 10,
 npoints
  tolerance
                 -> 1e-8,
                 -> 1e-6,
  cutoff
                 -> 0.01,
  binsize
                 -> false,
  zero
  negative
                 -> false
};
```

6.2.4 Example

The following example samples the cavity size distribution of an FCC lattice, which consists of a tetrahedral and an octahedral cavity. Its scripting equivalent can be found in ./examples/sample/cavity/build.emc. This example will be discussed in parts in the following. First, a number of variables are set,

```
variables = {
 lattice -> $root+"lib/fcc",
 output -> "lattice",
 n -> 4
};
```

Here the variable lattice describes the location of a predefined FCC lattice in EMC format. The final resulting positions are stored in filenames with starting with lattice. The variable n holds the number of replicas. An FCC lattice is obtained from the library directory by

```
get = {name -> lattice};
```

after which it is replicated n times in all directions by

```
crystal = \{n \rightarrow \{n, n, n\}\};
```

thus creating a superlattice. Subsequently, sampling of cavities using a Newton solver is turned on by

```
= {
sample
                 -> {
 cavity
    active
                 -> true,
    solver
                 -> newton,
                 -> "cavity",
    record
    binsize
                 -> 0.001,
                 -> 1000
    ninserts
 }
};
```

One thousand inserts are executed for each sampled structure. Cavity and saddle point positions are stored in file "cavity.pdb" by defining record. The resulting distribution has a bin size of 0.001. All omitted parameter definitions will internally be set to the above mentioned defaults. The selected FCC lattice does not include a force field definition, Force field parameters – which are not included in the selected FCC lattice – can be defined by using the simulation scripting command,

```
standard
                 -> {
                 -> {
      pair
        active -> true,
        cutoff \rightarrow 2.5,
        data
                 -> {
                 -> 0, i1 -> 0, sigma -> 1, epsilon -> 1}
           {i0
        }
      }
    }
  },
  moves
                 -> {
                 -> {active -> false}
    displace
  }
};
```

Here, the standard Lennard-Jones force field is chosen. Subsequently, displacement moves are switched off, thus not allowing particles to move. The seed value -1 triggers the use of the system clock as a seed value. A trajectory file is created through

```
traject = {name -> "cavity", frequency -> 1};
```

The resulting configuration is run for 0 cycles in order to sample the cavity size distribution,

```
run = {ncycles -> 0};
```

Once the cavity size distribution has been sampled, it is stored in both a Mathematica and a comma separated value format through

```
export = {
  sample -> {name -> "cavity", type -> cavity, style -> math},
  sample -> {name -> "cavity", type -> cavity, style -> csv}
};
```

The superlattice is stored in both EMC native and PDB formats by means of

```
put = {name -> output};
pdb = {name -> output};
```

The above described script is run with

```
emc_linux build.emc 2>&1 | tee build.out
```

A VMD script called cavity.vmd has been provided in the same directory. This script allows for simulataneous visualization of particles, octahedral and tetrahedral cavities, demarcated with types A and B respectively. The script is called with

vmd -e cavity.vmd

The color of the solid particles identifies the size of the cavity. Note, that due to the statistical nature of the algorithm, not all cavities necessarily will be found. Thus, possibly not all cavities will be displayed.

6.2.5 References

1. P.J. in 't Veld, M.T. Stone, T.M. Trustkett, and I.C. Sanchez, "Liquid Structure via Cavity Size Distributions", *J. Phys. Chem. B* **2000**, *104*, 12028

6.3 Gr

6.3.1 Syntax

```
= {
gr
  id
                 -> integer,
 active
                 -> boolean,
                 -> integer,
 frequency
                 -> struct,
  source
                 -> struct,
  target
 cutoff
                 -> real,
 binsize
                 -> real,
  intra
                 -> struct,
  inter
                 -> struct,
  total
                 -> struct
};
```

Directive id active	Parameters integer boolean	Description Method identifier for referencing subsequent alterations. Indicates the state of the method; options are true or false.
frequency	integer	Sets the frequency of the method, which is assumed to be a positive integer.
source, target	struct	Definition of source and target clusters, groups, and sites (see Section 7.3 [Focus], page 178); all mobile and active sites are taken when not defined.
cutoff	real	Sets the extent to which distances are evaluated; a zero value will result in initialization with the maximum cutoff resulting from active pairwise potential definitions.
binsize	real	Sets the binsize of the resulting distributions.
intra,	struct	Distributions resulting from analysis, reporting in-
inter, total		tramolecular, intermolecular, and all distances resulting from pairwise combinations as specified by source and target structures.

6.3.2 Usage

The gr command is used to sample radial distribution functions of all morphologies entailed by a simulation. Intramolecular, intermolecular, and total distributions are sampled based on the specified source and target definitions. Resulting distributions are reported as frequency distributions, which can be exported by using the scripting command export (see Section 5.10 [Export], page 100). An example can be found in ./examples/build/sample/gr/ by running setup.sh.

6.3.3 Default

```
The default is given by sample = {
```

6.3.4 Pair Correlation Functions

The following has been borrowed from Wikipedia¹. In statistical mechanics, the radial distribution function, (or pair correlation function) g(r) in a system of particles (atoms, molecules, colloids, etc.), describes how density varies as a function of distance from a reference particle.

If a given particle is taken to be at the origin O, and if $\rho = N/V$ is the average number density of particles, then the local time-averaged density at a distance r from O is $\rho g(r)$. This simplified definition holds for a homogeneous and isotropic system. A more general case will be considered below.

In simplest terms it is a measure of the probability of finding a particle at a distance of r away from a given reference particle, relative to that for an ideal gas. The general algorithm involves determining how many particles are within a distance of r and r + dr away from a particle. This general theme is depicted to the right, where the red particle is our reference particle, and blue particles are those which are within the circular shell, dotted in orange.

The RDF is usually determined by calculating the distance between all particle pairs and binning them into a histogram. The histogram is then normalized with respect to an ideal gas, where particle histograms are completely uncorrelated. For three dimensions, this normalization is the number density of the system multiplied by the volume of the spherical shell, which mathematically can be expressed as $g(r)_I = 4\pi r^2 \rho dr$, where ρ is the number density.

Given a potential energy function, the radial distribution function can be computed either via computer simulation methods like the Monte Carlo method, or via the Ornstein-Zernike equation, using approximative closure relations like the Percus-Yevick approximation or the Hypernetted Chain Theory. It can also be determined experimentally, by radiation scattering techniques or by direct visualization for large enough (micrometer-sized) particles via traditional or confocal microscopy.

The radial distribution function is of fundamental importance since it can be used, using the Kirkwood–Buff solution theory, to link the microscopic details to macroscopic properties. Moreover, by the reversion of the Kirkwood-Buff theory, it is possible to attain the microscopic details of the radial distribution function from the macroscopic properties.

6.3.5 Definition

Consider a system of N particles in a volume V (for an average number density $\rho = N/V$) and at a temperature T (let us also define $\beta = \frac{1}{kT}$). The particle coordinates are \vec{r}_i , with i = 1, ..., N. The potential energy due to the interaction between particles is $U_N(\vec{r}_1, ..., \vec{r}_N)$ and we do not consider the case of an externally applied field.

The appropriate averages are taken in the canonical ensemble (N, V, T), with $Z_N = \int \dots \int e^{-\beta U_N} d\vec{r}_1 \dots d\vec{r}_N$ the configurational integral, taken over all possible combinations of particle positions. The probability of an elementary configuration, namely finding particle 1 in $d\vec{r}_1$, particle 2 in $d\vec{r}_2$, etc. is given by

$$P^{(N)}\left(\vec{r}_1,\;...,\vec{r}_N\right)\;d\vec{r}_1\;...\;d\vec{r}_N = \tfrac{e^{-\beta U_N}}{Z_N}\;d\vec{r}_1\;...\;d\vec{r}_N\;.$$

The total number of particles is huge, so that $P^{(N)}$ in itself is not very useful. However, one can also obtain the probability of a reduced configuration, where the positions of only n < N particles are fixed, in \vec{r}_1 ..., \vec{r}_n , with no constraints on the remaining N-n particles. To this end, one has to integrate the above equation over the remaining coordinates \vec{r}_{n+1} , ..., \vec{r}_N :

$$P^{(n)}(\vec{r}_1, ..., \vec{r}_n) = \frac{1}{Z_N} \int ... \int e^{-\beta U_N} d\vec{r}_{n+1} ... d\vec{r}_N.$$

The particles being identical, it is more relevant to consider the probability that any n of them occupy positions $\vec{r}_1, ..., \vec{r}_n$ in any permutation, thus defining the n-particle density

$$\rho^{(n)}(\vec{r}_1, ..., \vec{r}_n) = \frac{N!}{(N-n)!} P^{(n)}(\vec{r}_1, ..., \vec{r}_n).$$

For n = 1, this equation gives the one-particle density which, for a crystal, is a periodic function with sharp maxima at the lattice sites. For a (homogeneous) liquid, it is independent of the position \vec{r}_1 and equal to the overall density of the system:

$$\frac{1}{V} \int \rho^{(1)}(\vec{r}_1) d\vec{r}_1 = \rho^{(1)} = \frac{N}{V} = \rho.$$

It is now time to introduce a correlation function $g^{(n)}$ by

$$\rho^{(n)}(\vec{r}_1, ..., \vec{r}_n) = \rho^n g^{(n)}(\vec{r}_1, ..., \vec{r}_n).$$

 $g^{(n)}$ is called a correlation function, since if the atoms are independent from each other $\rho^{(n)}$ would simply equal ρ^n and therefore $g^{(n)}$ corrects for the correlation between atoms. From the latter two equations it follows that

$$g^{(n)}(\vec{r}_1, \ \ldots, \ \vec{r}_n) = \tfrac{V^n N!}{N^n (N-n)!} \cdot \tfrac{1}{Z_N} \ \int \ldots \int e^{-\beta U_N} \ d\vec{r}_{n+1} \ \ldots \ d\vec{r}_N.$$

6.3.6 Relations Involving g(r)

6.3.6.1 Structure Factor

The second-order correlation function $g^{(2)}(\vec{r}_1, \vec{r}_2)$ is of special importance, as it is directly related (via a Fourier transform) to the structure factor of the system and can thus be determined experimentally using X-ray diffraction or neutron diffraction. If the system consists of spherically symmetric particles, $g^{(2)}(\vec{r}_1, \vec{r}_2)$ depends only on the relative distance between them, $\vec{r}_{12} = \vec{r}_2 - \vec{r}_1$. We will drop the sub- and superscript: $g(\vec{r}) \equiv g^{(2)}(\vec{r}_{12})$. Taking particle 0 as fixed at the origin of the coordinates, $\rho g(\vec{r}) dr = dn(\vec{r})$ is the number

of particles (among the remaining N-1) to be found in the volume $d\vec{r}$ around the position \vec{r} . We can formally count these particles as $dn(\vec{r}) = \langle \sum_{i \neq 0} \delta(\vec{r} - \vec{r}_i) \rangle d\vec{r}$, with $\langle ... \rangle$ the ensemble average, yielding

$$g(\vec{r}) = \frac{1}{\rho} \langle \sum_{i \neq 0} \delta(\vec{r} - \vec{r}_i) \rangle = V \frac{N-1}{N} \, \langle \delta(\vec{r} - \vec{r}_1) \rangle,$$

where the second equality requires the equivalence of particles 1, ..., N-1. The formula above is useful for relating $g(\vec{r})$ to the static structure factor $S(\vec{q})$, defined by $S(\vec{q}) = 1/N\langle \sum_{ij} e^{-i\vec{q}(\vec{r}_i - \vec{r}_j)} \rangle$, since we have

$$\begin{split} S(\vec{q}) &= 1 + \frac{1}{N} \langle \sum_{i \neq j} e^{-i\vec{q}(\vec{r}_i - \vec{r}_j)} \rangle = \\ &= 1 + \frac{1}{N} \left\langle \int_V e^{-i\vec{q}\vec{r}} \sum_{i \neq j} \delta \left[\vec{r} - (\vec{r}_i - \vec{r}_j) \right] \, d\vec{r} \right\rangle = \\ &= 1 + \frac{N(N-1)}{N} \int_V e^{-i\vec{q}\vec{r}} \left\langle \delta(\vec{r} - \vec{r}_1) \right\rangle \, d\vec{r} \,, \end{split}$$

and thus

$$S(\vec{q}) = 1 + \rho \int_V e^{-i\vec{q}\vec{r}} g(\vec{r}) d\vec{r},$$

proving the Fourier relation alluded to above. This equation is only valid in the sense of distributions, since $g(\vec{r})$ is not normalized: $\lim_{r\to\infty}g(\vec{r})=1$, so that $\int_V d\vec{r}g(\vec{r})$ diverges as the volume V, leading to a Dirac peak at the origin for the structure factor. Since this contribution is inaccessible experimentally we can subtract it from the equation above and redefine the structure factor as a regular function,

$$S'(\vec{q}) = S(\vec{q}) - \rho \delta(\vec{q}) = 1 + \rho \int_V e^{-i\vec{q}\vec{r}} [g(\vec{r}) - 1] d\vec{r}.$$

Finally, we rename $S(\vec{q}) \equiv S'(\vec{q})$ and, if the system is a liquid, we can invoke its isotropy,

$$S(q) = 1 + \rho \int_V e^{-i\vec{q}\vec{r}} [g(r) - 1] d\vec{r} = 1 + 4\pi \rho \frac{1}{a} \int r \sin(qr) [g(r) - 1] dr$$
.

6.3.6.2 Compressibility Equation

Evaluating the latter equation for q = 0 while using the relation between the isothermal compressibility χ_T and the structure factor at the origin yields the compressibility equation:

$$\rho kT\chi_T = kT\left(\frac{\partial\rho}{\partial p}\right) = 1 + \rho\int_V [g(r)-1]\,d\vec{r}.$$

6.3.6.3 Potential of Mean Force

It can be shown that the radial distribution function is related to the two-particle potential of mean force $w^{(2)}(r)$ by²

$$g(r) = \exp\left[-\frac{w^{(2)}(r)}{kT}\right].$$

6.3.6.4 Energy Equation

If the particles interact via identical pairwise potentials³,

$$U_N = \sum_{i>j=1}^{N} u(|\vec{r}_i - \vec{r}_j|),$$

the average internal energy per particle is

$$\frac{\langle E \rangle}{N} = \frac{3}{2}kT + \frac{\langle U_N \rangle}{N} = \frac{3}{2}kT + \frac{1}{2}\rho \int_V d\vec{r} \ u(r)g(r,\rho,T).$$

6.3.6.5 Pressure Equation of State

Developing the virial equation yields the pressure equation of state,

$$p = \rho kT - \frac{1}{6}\rho^2 \int_V r g(r, \rho, T) \frac{du(r)}{dr} d\vec{r}.$$

6.3.6.6 Thermodynamic Properties

The radial distribution function is an important measure because several key thermodynamic properties, such as potential energy and pressure can be calculated from it. For a 3-D system where particles interact via pairwise potentials, the potential energy of the system can be calculated as follows⁴

$$PE = 2\pi\rho N \int_0^\infty r^2 u(r)g(r)dr,$$

where N is the number of particles in the system, ρ is the number density, u(r) is the pair potential. The pressure of the system can also be calculated by relating the 2nd virial coefficient to g(r). The pressure can be calculated as follows

$$P = \rho k_B T - \frac{2}{3} \pi \rho^2 \int_0^\infty r^3 g(r) \frac{du(r)}{dr} dr,$$

where T is the temperature and k_B is Boltzmann's constant. Note that the results of potential and pressure will not be as accurate as directly calculating these properties because of the averaging involved with the calculation of g(r).

6.3.7 References

- 1. N. N., Radial Distribution Function, Wikipedia (1/1/2016).
- Chandler, D., Introduction to Modern Statistical Mechanics. Chapter 7.3, Oxford University Press (1987).
- 3. Hansen, J. P. and McDonald, I. R., Theory of Simple Liquids (3rd ed.), Academic Press (2005).
- 4. Frenkel, D. and Smit, B., Understanding Molecular Simulation from Algorithms to Applications (2nd ed.), Academic Press (2002).

6.4 Gyration

6.4.1 Syntax

```
gyration
                = {
 id
                -> integer,
 active
                -> boolean,
 frequency
                -> integer,
 focus
                -> struct,
  cutoff
                -> real,
 binsize
                -> real,
  distributions -> struct
};
```

Directive	Parameters	Description
id	integer	Method identifier for referencing subsequent alterations.
active	boolean	Indicates the state of the method; options are true or
		false.
frequency	integer	Sets the frequency of the method, which is assumed to
		be a positive integer.
focus	struct	Definition of clusters, groups, and sites as to which to
		focus analysis on (see Section 7.3 [Focus], page 178); all
		mobile and active sites are taken when not defined.
cutoff	real	Sets the extent to which distances are evaluated; a zero
		value will result in initialization with the maximum cutoff
		resulting from active pairwise potential definitions.
binsize	real	Sets the binsize of the resulting distributions.
distributions struct		Distributions resulting from analysis; these distributions
		- only shown in .emc and exported .m files - cannot be
		influenced with scripting command sample.

6.4.2 Usage

The gyration command is used to sample radius of gyration distributions of all morphologies entailed by a simulation. Resulting distributions are reported as frequency distributions, which can be exported by using the scripting command export (see Section 5.10 [Export], page 100). Averages are reported as part of the resulting distribution as shown in .emc and .m files. An example can be found in ./examples/build/sample/gyration/ by running setup.sh.

6.4.3 Default

6.4.4 Theory

The following has been borrowed from Wikipedia¹. In polymer physics, the radius of gyration is used to describe the dimensions of a polymer chain. The radius of gyration of a particular molecule at a given time is defined as:

$$R_g^2 = \frac{1}{N^2} \sum_{i=1,j>i}^{N} (\vec{r}_i - \vec{r}_j)^2,$$

where N represents the number of particles in the molecule and vectors \vec{r}_i and \vec{r}_j denote particle positions. Since the chain conformations of a polymer sample are quasi infinite in number and constantly change over time, the "radius of gyration" discussed in polymer physics must usually be understood as a mean over all polymer molecules of the sample and over time. That is, the radius of gyration which is measured as an average over time or ensemble:

$$\langle R_q^2 \rangle = \frac{1}{N^2} \langle \sum_{i=1,j>i}^N (\vec{r}_i - \vec{r}_j)^2 \rangle,$$

where the angular brackets $\langle ... \rangle$ denote the ensemble average. An entropically governed polymer chain (i.e. in so called theta conditions) follows a random walk in three dimensions. The radius of gyration for this case is given by

$$R_g = \frac{1}{\sqrt{6}} \sqrt{N} \ a$$

Note that although aN represents the contour length of the polymer, a is strongly dependent of polymer stiffness and can vary over orders of magnitude. N is reduced accordingly. One reason that the radius of gyration is an interesting property is that it can be determined experimentally with static light scattering as well as with small angle neutron- and x-ray scattering. This allows theoretical polymer physicists to check their models against reality. The hydrodynamic radius is numerically similar, and can be measured with Dynamic Light Scattering (DLS).

6.4.5 References

1. N. N., Radius of Gyration, Wikipedia (1/1/2016).

6.5 Interaction

6.5.1 Syntax

interaction	= {	
id	->	integer,
active	->	boolean,
frequency	->	integer,
cutoff	->	real,
mode	->	option,
coulomb	->	option,
bias	->	option,
weight	->	option,
ntrials	->	integer,
nvolumes	->	integer,
ninits	->	integer,
nwidth	->	integer,
source	->	struct,
target	->	struct,
mass	->	real,
dvolume	->	real,
volume1	->	struct,
volume2	->	struct,
denergy	->	real,
energy	->	struct
} ;		

Directive	Parameters	Description
id	integer	Method identifier for referencing subsequent alterations.
active	boolean	Indicates the state of the method; options are true or false.
frequency	integer	Sets the frequency of the method, which is assumed to be a positive integer.
mode	global	Sets the desired pairwise potential cut off mode; options are global, individual, and repulsive
coulomb	none	Sets the desired type of coulombic interactions; options are none, cut, switch, and long.
bias	none	Sets the bias used during interaction sampling; options are none, energy, and frequency.
weight	none	Sets the weighting used for populating the resulting interaction distribution; options are none, boltzmann, and montecarlo.
ntrials	0	Sets the number of energetic interaction sampling trials.
nvolumes	100	Sets the number of volume sampling trials.

ninits	1000	Sets the number of trials for populating the initial distribution as used by energy and frequency biased sampling.
nwidth	1	Sets the spread used during sampling for accessing non- initialized bins resulting from the previous option.
source,	struct	Definition of source and target clusters, groups, and sites
target		(see Section 7.3 [Focus], page 178) setting the two interaction species; all mobile and active sites are taken when not defined.
cutoff	real	Sets the extent to which distances are evaluated; a zero value will result in initialization with the maximum cutoff resulting from active pairwise potential definitions.
dvolume	real	Sets the binsize of the resulting distributions.
volume1,	real	Resulting distribution of source and target volume anal-
volume2		ysis; produces the volume distribution of the two interacting species.
denergy	real	Sets the binsize of the resulting energy distributions.
energy	struct	Distributions resulting from analysis, reporting pairwise interaction as specified by source and target structures.

6.5.2 Usage

The interaction command is used to sample the pairwise potential of mean force, which can either be calculated from provided or from internally generated structures. Internally generated structures are sampled over distances within the specified cutoff and over randomly chosen orientations. Resulting pairwise interactions can be weighted by several weighting techniques. Resulting samples per bin can be controlled by different biasing schemes. Volumetric and energetic distributions are sampled based on the specified source and target definitions. Resulting distributions are reported as frequency distributions, which can be exported by using the scripting command export (see Section 5.10 [Export], page 100).

6.5.3 Default

The default is given by

```
= {
sample
  id
                 -> O,
                 -> false,
  active
  frequency
                 -> 1,
                 -> option,
 mode
                 -> option,
  coulomb
 bias
                 -> none,
  weight
                 -> none,
 ntrials
                 -> 0,
                 -> 100,
  nvolumes
                 -> 1000,
 ninits
  nwidth
                 -> 1,
  source
                 -> {},
```

```
target -> {},
mass -> 0,
cutoff -> 0,
dvolume -> 0.01,
denergy -> 0.01
};
```

6.6 Profiles

Density, pressure, and energy profile explanation. $\,$

6.7 Examples

Choice examples.

7 Variable Descriptions

EMC uses hierarchical variables, which can be recursive in character. This chapter describes the syntax of variables that are used often within the context of the EMC scripting language. Currently, variables use a representation similar to Mathematica.

Mathematical operations supported for scalar variable types integer and real are

Operator	Application	Description
+	a + b	Addition
_	a - b	Subtraction
*	a * b	Multiplication
/	a / b	Division
^	a^b	Power
?	q ? a : b	Logical; select a when q is true, b otherwise

Mathematical functions supported for scalar variable types integer and real are

Operator	Application	Description
acos	acos(x)	Arccosine using a radial basis
asin	asin(x)	Arcsine using a radial basis
atan	atan(x)	Arctangent using a radial basis
cos	cos(x)	Cosine using a radial basis
eval	eval(expr)	Evaluate a string expression into a value
exp	exp(x)	Exponential
int	int(x)	Floor of a real number
log	log(x)	Natural log
sin	sin(x)	Sine using a radial basis
sqrt	sqrt(x)	Square root
tan	tan(x)	Tangent using a radial basis
mass	mass(index)	Mass of a predefined group index
mtotal	<pre>mtotal(focus)</pre>	Mass of sites in selection as defined by focus (See
		Section 7.3 [Focus], page 178)
nclusters	nclusters(Number of clusters in selection as defined by focus
	focus)	(See Section 7.3 [Focus], page 178)
nsites	nsites(index)	Number of sites of a predefined group index
ntotal	ntotal(focus)	Number of sites in selection as defined by focus (See
		Section 7.3 [Focus], page 178)
type	type(index)	Determine type number of site index (used for trans-
• •	• •	fer to e.g. LAMMPS input scripts)
vsites	vsites(focus)	Volume captured by selection as defined by focus (See
		Section 7.3 [Focus], page 178)
vtotal	vtotal(focus)	Total system volume as defined by system selection
		(See Section 7.3 [Focus], page 178)

Note, that groups must have been defined (see Section 5.17 [Groups], page 114) and a force field must have been applied (see Section 5.11 [Field], page 103) in order for mass(index) and nsites(index) to return correct values. See also scripts/emc_setup.pl.

Reverved variable names are represented by the following constants

Constant	Type	Description
\$root	string	EMC root directory
\$arg#	varies	command line argument
pi	real	number: 3.14159265358979323846264338327950288
е	real	number: 2.71828182845904523536028747135266250
true	integer	number: 0
false	integer	number: 1
null	integer	number: -1

Internally defined constants are booleans true and false, mathematical constants pi and e, internal variable null, and the program's root directory \$root. For the latter, it is assumed, that the emc executable resides in \$root/bin. All other locations will result in erroneous behavior of \$root. Command line arguments not preceded by a "-" are accessed through the prefix \$arg followed by a number, starting at 0. For example

```
variables = {name -> $arg0};
```

sets the variable name to the first command line argument. In case of the command

```
emc_${HOST} build.emc atoms
```

the variable name in the above example would be set to the text "atoms". Note, that — in this example — calling \$arg1 and subsequent non-exisiting arguments as a variable will result in an error due to the fact, that \$arg1 is not defined on the command line and therefore also not internally. Addition operations using a + b, where either a or b is of variable type string, will result in a string. For example

```
"text" + 1 + 1 := "text11"
but
    "text" + (1 + 1) := "text2"
Previously defined variables can also be included
    a = 2^2
    "text" + a := "text4"
or
    a = 2*3
    dir = "/home/user/text"
    dir + "_" + a := "/home/user/text_6"
```

7.1 Constants

7.1.1 Syntax

Directive	Parameters	Description
systems	constant	Adds a constant to the systems category; constants are expected to be alpha-numerical.
clusters	constant	Adds a constant to the clusters category; constants are expected to be alpha-numerical.
groups	constant	Adds a constant to the groups category; constants are expected to be alpha-numerical.
sites	constant	Adds a constant to the sites category; constants are expected to be alpha-numerical.

7.1.2 Usage

This variable style describes constants. The style is additive when used in combinations with see Section 5.40 [Variables], page 150.

7.1.3 Default

Unless otherwise stated, the default is given by

```
constants = {
  systems -> {},
  clusters -> {},
  groups -> {},
  sites -> {}
```

7.2 System Flags

7.2.1 Syntax

flag -> option
Valid options are, when set,

Directive Parameters

ignore Ignore any missing contribution and do not generate any output

complete Complete any missing contribution

warn Generate a warning when a contribution is missing

empty Create an empty entry when a contribution is missing

error Exit on errors resulting from missing contributions; all missing contributions will be listed before an exit on error occurs

7.2.2 Default

Unless otherwise stated, the default is given by

flag -> error

7.3 Focus

7.3.1 Syntax

Directive clusters	Parameters constant	Description Refers to cluster constants as defined by the constants table.
groups	constant	Refers to group constants as defined by the constants table.
ntrials	integer	Defines the number of trials used to determine volume; volume is determined by means of Monte Carlo integration.
sites	constant	Refers to site constants as defined by the constants table.
systems	constant	Refers to system constants as defined by the constants table.

7.3.2 Usage

Defines a system sites subset by means of their constants identifiers (see Section 7.1 [Constants], page 176).

7.3.3 Default

The default is described by an empty list for all three contributors,

```
focus = {
  clusters -> {},
  groups -> {},
  sites -> {},
  systems -> {},
  ntrials -> 10000
};
```

7.4 Moves

7.4.1 Syntax

```
moves
               = {
 ncycles
               -> integer,
 cycle
               -> integer,
 move
               -> integer,
 body
               -> struct,
 deform
               -> struct,
 displace
               -> struct,
 endbridge
               -> struct,
 migrate
               -> struct,
 rebridge
               -> struct,
 reptate
               -> struct,
 rotate
               -> struct,
 surface
               -> struct,
 temper
               -> struct
};
```

Directive	Parameters	Description
ncycles	integer	Sets the total number of simulation cycles.
cycle	integer	Sets the current simulation cycle
move	integer	Sets the current simulation move.
body	struct	Settings for displacing sites inside a body.
deform	struct	Settings for system box deformation (see Section 7.4.6
		[Displace], page 182).
displace	struct	Settings for displacing sites in a system box.
endbridge	struct	Settings for recombining cluster ends in a system box.
migrate	struct	Settings for migrating short branches.
rebridge	struct	Settings for rebridging clusters in a system box.
reptate	struct	Settings for reptating cluster ends in a system box.
rotate	struct	Settings for rotating cluster ends in a system box.
surface	struct	Settings for displacing sites on a body surface.
temper	struct	Settings for parallel tempering between systems.

7.4.2 Usage

This variable style describes moves. These moves encompass standard and advanced Monte Carlo moves. 1,2

7.4.3 Default

Unless otherwise stated, the default is given by

```
moves = {
  ncycles -> 0,
  cycle -> 0,
  move -> 0
```

};

By default, all moves are activated with zero frequency, with the exception of the displacement move (see Section 7.4.6 [Displace], page 182), which has a frequency of one.

7.4.4 References

- 1. P.J. in 't Veld, M. Hütter, and G.C. Rutledge, "Temperature-Dependent Thermal and Elastic Properties of the Interlamellar Phase of Semicrystalline Polyethylene by Molecular Simulation", *Macromolecules* **2006**, *39*, 439
- 2. V. Kumar, C.R. Locker, P.J. in 't Veld, G.C. Rutledge, "Effect of Short Chain Branching on the Interlamellar Structure of Semicrystalline Polyethylene", *Macromolecules* **2017**, *50*, 1206

7.4.5 Deform Move

7.4.5.1 Syntax

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
mode	option	Set the mode of deformation; valid options are isotropic, shape, full, xx, yx, yy, zx, zy, and zz.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
n	integer	Number of masses available in the simulation; controlled internally: output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.5.2 Usage

The deform move alters the box geometry based the mode and the selected pressure as set in systems (see Section 7.11 [Systems], page 199).

7.4.5.3 Default

7.4.6 Displace

7.4.6.1 Syntax

```
= {
displace
 active
               -> boolean,
 frequency
               -> integer,
 dlimit
               -> real,
 nsites
               -> integer,
               -> boolean,
 couple
               -> integer,
 dmax
               -> real,
               -> struct
 accept
};
```

Directive	Parameters	Description
active	boolean	Indicates the state of the move; options are true or
		false.
frequency	integer	Sets the frequency of the move, which is assumed to be
		a positive integer.
dlimit	real	Set the maximum displacement limit.
couple	boolean	Couple all individual acceptances when internally deter-
		mining displacement limits.
nsites	integer	Set the number of sites to be displaced within one move.
n	integer	Number of masses available in the simulation; controlled
		internally.
dmax	real	Array of adapted displacement limits based on simulation
		progress; internally determined using a feedback loop;
		output only.
accept	struct	Array of accumulative accepted and total trials.

7.4.6.2 Usage

The displace move is used to displace sites by means of perturbation. The target acceptance is set to 50%.

7.4.6.3 Default

The default is given by

7.4.7 Endbridge

7.4.7.1 Syntax

```
= {
endbridge
 active
               -> boolean,
               -> integer,
 frequency
 target
               -> constant,
 nmin
               -> integer,
 nmax -> integ
tolerance -> real,
               -> integer,
 dmin
               -> real,
 dmax
n
accept
               -> real,
               -> integer,
               -> struct
};
```

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
target	constant	Target group.
nmin	integer	Minimum chain length after end-bridging; valid values are values larger or equal than three.
nmax	integer	Maximum chain length after end-bridging; chain length is unlimited when zero.
tolerance	real	Tolerance; currently not in use.
dmin	real	Minimum distance between parent chain end and child end-bridging candidate chain.
dmax	real	Maximum distance between parent chain end and child end-bridging candidate chain.
n	integer	Number of masses available in the simulation; controlled internally: output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.7.2 Usage

The endbridge move allows for recombination of chain ends with other chains; currently, only linear non-branched chains are allowed.

7.4.7.3 Default

The default is given by

```
sample = {
  active -> false,
```

```
frequency -> 1,
  nmin -> 3
};
```

7.4.8 Migrate

7.4.8.1 Syntax

```
migrate = {
  active -> boolean,
  frequency -> integer,
  target -> constant,
  radius -> real,
  n -> integer,
  accept -> struct
};
```

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
target	constant	Target type for moving the branch to.
radius	real	Select sites with target types within radius from type on backbone; selects all available sites in the system when the radius equals zero.
n	integer	Number of masses available in the simulation; controlled internally: output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.8.2 Usage

The migrate move is used to migrate branches on the backbone of a chain. Branches should be created by using grafting short side clusters. Branches should not exceed a length of three sites. Target candidates are chosen at random from the candidate list following from the chosen radius.

7.4.8.3 Default

```
The default is given by
```

7.4.9 Rebridge

7.4.9.1 Syntax

```
rebridge = {
  active -> boolean,
  frequency -> integer,
  drivers -> integer,
  tolerance -> real,
  n -> integer,
  accept -> struct
};
```

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
drivers tolerance	integer real	Number of driver sites; valid options are 0, 1, and 2. Move tolerance; currently not in use.
n	integer	Number of masses available in the simulation; controlled internally: output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.9.2 Usage

The rebridge move performs concerted rotations on single chains to promote accelerated equilibration of long chain polymers.¹

7.4.9.3 Default

```
The default is given by
```

1. V.G. Mavrantzas, T.D. Boone, E. Zervopoulou, and D.N. Theodorou, "End-Bridging Monte Carlo: A Fast Algorithm for Atomistic Simulation of Condensed Phases of Long Polymer Chains", *Macromolecules* **1999**, *32*, 5072.

7.4.10 Reptate

7.4.10.1 Syntax

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
target	constant	Optional target end group and site; considers all available chain ends when not defined.
nmin	integer	Minimum length of the resulting chain; valid values are values larger or equal than three.
n	integer	Number of masses available in the simulation; controlled internally: output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.10.2 Usage

The reptate move promotes chain reptation by taking the end of one chain and moving it to the end of another.

7.4.10.3 Default

```
The default is given by
```

7.4.11 Rotate

7.4.11.1 Syntax

Directive active	Parameters boolean	Description Indicates the state of the move; options are true or false.
frequency	integer	Sets the frequency of the move, which is assumed to be a positive integer.
nmax	integer	Currently not in use.
n	integer	Number of masses available in the simulation; controlled internally: output only.
amax	real	Array of adapted rotation limits based on simulation progress; internally determined using a feedback loop; output only.
accept	struct	Array of accumulative accepted and total trials; output only.

7.4.11.2 Usage

The rotate move selects and rotate chain ends at random. One chain is considered per attempted move. Per attempted on to three sites are included in a random rotation.

7.4.11.3 Default

7.4.12 Surface

7.4.12.1 Syntax

Directive	Parameters	Description
n	integer	Number of masses available in the simulation; controlled
		internally: output only.
dmax	real	Array of adapted displacement limits based on simulation
		progress; internally determined using a feedback loop;
		output only.
accept	struct	Array of accumulative accepted and total trials; output
		only.

7.4.12.2 Usage

The **surface** move represents a subclass of the displacement move and operates on sites that are part of a body representing a surface. The move is automatically invoked when the latter applies. User settings are controlled via the displacement move.

7.4.12.3 Default

Defaults are given by the displacement move.

7.4.13 Temper

7.4.13.1 Syntax

Directive	Parameters	Description
active	boolean	Indicates the state of the move; options are true or
		false.
frequency	integer	Sets the frequency of the move, which is assumed to be
		a positive integer.
n	integer	Number of masses available in the simulation; controlled
		internally: output only.
accept	struct	Array of accumulative accepted and total trials; output
		only.

7.4.13.2 Usage

The temper move controls the exchange between multiple replicas as represented by existing systems. The exchange is governed by the per system chosen temperatures.

7.4.13.3 Default

7.5 Port

7.5.1 Syntax

port = {forcefield -> option};

Directive forcefield	Parameters option	Description Sets the ported force field type.
Option	Description	
none	No force field into ported format.	erpretation; bonded contributions are not included in the
auto	_	s of ported format; based on which force field family is only works correctly when just one force field family is
boltzmann	Assumes Boltzma	ann force fields (see Section 7.12.4 [Boltzmann], page 204).
cff	Assumes CFF for	cce fields (see Section 7.12.6 [CFF], page 217).
charmm	Assumes CHARM	IM force fields (see Section 7.12.7 [CHARMM], page 233).
dpd	Assumes DPD o	coarse-grained force fields (see Section 7.12.11 [DPD],
_	page 272).	
martini	Assumes MARTI TINI], page 290).	NI coarse-grained force fields (see Section 7.12.13 [MAR-
opls	• · · · · · /	all-atom and united-atom force fields (see Section 7.12.14
1	[OPLS], page 298	· · · · · · · · · · · · · · · · · · ·
sdk		coarse-graine force fields (see Section 7.12.15 [SDK],
standard	,	d force fields (see Section 7.12.17 [Standard], page 320).
table		ed force fields (see Section 7.12.18 [Table], page 329).
trappe		all-atom and united-atom force fields (see Section 7.12.19
огаррс	[TraPPE], page 3	· ·
coarse	• •	rained force fields.
colloid	Assumes colloida	
fene	Assumes FENE f	

7.5.2 Default

7.6 Profiles

7.6.1 Syntax

Directive bond	Parameters struct	Description Settings for bond length distributions of all bonds within all systems; no discrimination is made between bond types.
density	struct	Settings for mass density profiles; no discrimination is made between site masses.
force	struct	Settings for energy and virial density profiles; resulting binned contributions reflect the interaction of that bin with its surroundings.
mass	struct	Settings for mass profiles; distinction is made between contributing site masses.
order	struct	Settings for order profiles; no distinction is made between contributing bond types.

7.6.2 Usage

This variable style describes profiles.

7.6.3 Default

By default, all profiles are deactivated and all entries are zeroed out.

7.7 Region

7.7.1 Syntax

Directive	Parameters	Description
shape	option	Sets region shape; possibilities are cuboid or spheroid.
type	option	Sets region type; possibilities are relative or absolute.
mode	option	Sets region mode; possibilities are hard or soft; used in
		conjunction with sites growth.
center	vector	Defines the center of a region.
h	voigt	Defines the extent of a region as a voigt notation shape
		(see Section 7.15 [Voigt], page 346).
radius	vector	Defines the extent of a region as a vector (see Section 7.14
		[Vector], page 345); the radius is a derived alternative
		to h and translated into h upon execution.

7.7.2 Usage

Specifies a region of a certain shape within system simulation cell. Units of the shape center and radius can either be expressed in a relative or an absolute fashion. Relative units express both shape center and radius in units of base vectors \vec{a} , \vec{b} , and \vec{c} and lengths thereof. Absolute units make use of the simulation-wide settings as expressed under in the 'Units' section (see Section 7.13 [Units], page 343) of the simulation structure. The region's mode influences the growth of sites by either hard or soft exclusion or inclusion. The hard mode applies the region to all sites of inserted clusters, while the soft mode only applies the region to the first site of inserted clusters.

7.7.3 Default

Unless otherwise stated, the default is given by

```
region = {shape -> spheroid, type -> relative, mode -> hard, center -> {0,0,0}, h -> {0,0,0,0,0,0}};
```

7.8 SMILES

7.8.1 Syntax

chemistry = string;

7.8.2 Usage

The Simplified Molecular Input Line Entry System (or SMILES for short) is used to describe chemistry as reflected in the definitions of groups (see Section 5.17 [Groups], page 114). It follows the syntax as described Daylight Chemical Information Systems, Inc., but is extended to also capture non-periodic system representations as are used in defining coarse-grained systems. The distinction between periodic and non-periodic representations is identified by keywords atomistic and coarse respectively. In atomistic mode, hydrogen completion is implicit and lower character case signifies aromaticity, where allowed characters are c, n, o, p, and s. A short summary is listed in the table below. Extensions to the original SMILES format are marked with (extension).

Modifier	Example	Description
>X	>C	First branch point, connecting to chains in the down direction (extension).
Х<	C<	Second branch point, connecting to chains in the up direction (extension).
*	*	General branch point, connecting chains (extension).
~X	~C	Represents any bond (extension).
-X	-C	Represents a single bond (extension).
: X	:C	Represents a partial double bond (extension).
= X	=C	Represents a double bond.
#X	#N	Represents a triple bond.
Х	С	Lower case indicate aromatics in atomistic mode; al-
F3	F 7	lowed characters are c, n, o, p, and s.
[XX]	[He]	Needed for usage of symbols with more than one character.
()	CC(C)C	Indicates branching from backbone.
(X)1	(C)3	Indicates repeating sequences (here $(C)3 = CCC$); note, that $CC(C)3C = CCCCCC$ and $CC((C)3)C = CC(CCC)C$; multi-digits are allowed (extension).
1X	2H	Assignes an aberrant mass to a symbol (here 2).
X1	c1ccccc1	Indicates a link number to create ring structures (here a benzene ring); duplicate link numbers are allowed.
%	c%11ccccc%11	Used for multi-digit link numbers.
\X or \X	C/C=C/C	Identifies cis $(C/C=C\setminus C)$ or trans $(C/C=C/C)$ isomers.
@	[C@H]	Identifies L- ([C@@H]) or R- ([C@H]) stereoisomers.
+1	[Na+1]	Indicates an assigned positive charge; charges can be partial (e.g. 0.5); a + with omitted digits indicates an increase by +1, e.g. [Na+] equals [Na+1] or [Ca++] represents a calcium atom with charge +2.

-1 [Cl-1] Indicates an assigned negative charge; charges can be partial (e.g. 0.5); a - with omitted digits indicates an decrease by -1.

7.8.3 Examples

The following table shows a few example chemistry and their corresponding SMILES strings.

Chemistry **SMILES** water HOH cyclohexanone O=C1CCCCC1 dodecane (C)12CCCCCCCCCC iso-octane CC(C)CC(C)(C)Cdiphenylmethane c1ccccc1Cc1ccccc1 Methylene diphenyl 4,4'-diisocyanate O=C=Nc1ccc(cc1)Cc2ccc(N=C=O)cc2 trans-2-butene C/C=C/Cc1ccc2c(c1)c(c[nH]2)C[C@@H](C(=0)0)NL-tryptophan

7.9 Splines

7.9.1 Introduction¹

In mathematics, a spline is a sufficiently smooth polynomial function that is piecewise-defined, and possesses a high degree of smoothness at the places where the polynomial pieces connect (which are known as knots).^{2,3}

In interpolating problems, spline interpolation is often referred to as polynomial interpolation because it yields similar results, even when using low-degree splines, to interpolating with higher degree polynomials while avoiding instability due to Runge's phenomenon. In computer graphics splines are popular curves because of the simplicity of their construction, their ease and accuracy of evaluation, and their capacity to approximate complex shapes through curve fitting and interactive curve design.

The most commonly used splines are cubic spline, i.e., of order 3 – particular, cubic B-spline and cubic Bezier spline. They are common, in particular, in spline interpolation simulating the function of flat splines. The term spline is adopted from the name of a flexible strip of metal commonly used by draftsmen to assist in drawing curved lines.⁴

Splines are curves, which are usually required to be continuous and smooth. Splines are usually defined as piecewise polynomials of degree n with function values and first n-1 derivatives that agree at the points where they join. The abscissa values of the join points are called knots. The term "spline" is also used for polynomials (splines with no knots) and piecewise polynomials with more than one discontinuous derivative. As such, splines with no knots are generally smoother than splines with multiple discontinuous derivatives. Splines with few knots are generally smoother than splines with many knots; however, increasing the number of knots usually increases the fit of the spline function to the data. Knots give the curve freedom to bend to more closely follow the data.

It is commonly accepted that the first mathematical reference to splines is the 1946 paper by Schoenberg,⁵ which is probably the first place that the word "spline" is used in connection with smooth, piecewise polynomial approximation. However, the ideas have their roots in the aircraft and shipbuilding industries.

7.9.2 Linear Spline

The linear splined applied by EMC follows

$$\Delta x_i = x - x_i,$$

$$S_i(x) = \sum_{j=0}^{1} k_{j,i} \Delta x_i^j,$$

$$k_{0,i} = y_i,$$

$$k_{1,i} = (y_i - y_{i-1})/(x_i - x_{i-1}).$$

Linear splines find their application in cases of sudden large gradients or steps in y data with respect to x.

7.9.3 Cubic Spline

Internally, EMC applies a natural cubic spline by default. The algorithm behind a natural cubic spline is given by

$$\Delta x_i = x - x_i,$$

$$S_i(x) = \sum_{j=0}^3 k_{j,i} \Delta x_i^j,$$

$$S_i(x_i) = S_{i-1}(x_i) = y_i,$$

$$S_i'(x_i) = S_{i-1}'(x_i),$$

$$S_i''(x_i) = S_{i-1}''(x_i),$$

$$S_i''(x_0) = S_{n-1}''(x_n) = 0,$$

for which the above set of equations defines all constants k needed to describe the full spline function. Cubic splines are useful for capturing the behavior of reasonably well-behaved data.

7.9.4 References

- 1. Wikipedia on "Cubic Splines"
- 2. K. L. Judd, "Numerical Methods in Economics", MIT Press 1998, 225 (ISBN 978-0-262-10071-7).
- 3. W.-K. Chen, Wai-Kai (2009 "Feedback, Nonlinear, and Distributed Circuits", CRC Press 2009, 9-20 (ISBN 978-1-4200-5881-9).
- 4. M. H. Katz, "Multivariable Analysis: A Practical Guide for Clinicians and Public Health Researchers", Cambridge University Press 2011, 82 (ISBN 978-0-521-14107-9).
- 5. Schoenberg, "Contributions to the problem of approximation of equidistant data by analytic functions", Quart. Appl. Math. 1946, 4, 45-99 and 112-141.

7.10 System Flags

7.10.1 Syntax

flag = {charge -> boolean, map -> boolean, pbc -> boolean};

Directive	Parameters	Description
charge	boolean	Check for charge neutrality; options are true or false.
map	boolean	Map configurations into the system box; options are true or false.
pbc	boolean	Apply periodic conditions to system geometry; options
•		are true or false.

7.10.2 Default

Unless otherwise stated, the default is given by

flag = {charge -> true, map -> true, pbc -> true};

7.11 Systems

7.11.1 Syntax

```
systems
              = {
 n
              -> integer,
              -> {
 properties
   {
     id
              -> constant,
              -> real,
     v
              -> real,
     t
              -> real,
     mass
              -> real,
     nclusters -> integer,
     nsites -> struct,
     plane -> integer,
     geometry -> voigt
   },
};
```

Directive	Parameters	Description
n	integer	Sets the number of contributing systems to the total simulation.
properties		Describes the properties of each individual system.
id	constant	Identifies the system; refers to the systems paragraph in constants (see Section 7.1 [Constants], page 176).
p	real	Defines the system pressure; only used when deformation moves are active.
v	real	Defines the system volume.
t	real	Defines the system temperature.
mass	real	Reflects the total system mass; cannot be altered.
nclusters	integer	Reflects the number of clusters in this system; cannot be altered.
nsites	struct	Reflects the number of sites in this system; cannot be altered.
plane	integer	Reflects the original crystal plane the system was constructed with.
geometry	voigt	Defines the system geometry (see Section 7.15 [Voigt], page 346).

7.11.2 Usage

This variable style describes the definition of systems.

7.11.3 Default

Unless otherwise stated, the default is given by systems = $\{n \rightarrow 0\}$;

7.12 Types

7.12.1 Syntax

types	= {	
merge	-> boolean,	
virial	-> boolean,	
periodic	-> vector,	
neighbor	-> constant,	
stencil	-> constant,	
skin	-> real,	
shake	-> constant	
depth	-> integer,	
mass	-> struct,	
boltzmann	-> struct,	
charmm	-> struct,	
cff	-> struct,	
coarse	-> struct,	
colloid	-> struct,	
coulomb	-> struct,	
dpd	-> struct,	
gromacs	-> struct,	
inverse	-> struct,	
martini	-> struct,	
opls	-> struct,	
sdk	-> struct,	
spline	-> struct,	
standard	-> struct,	
table	-> struct,	
trappe	-> struct	
};		

Directive	Parameters	Description
merge	boolean	Allows for merging force field constants upon input when
		true; options are true or false.
virial	boolean	Describes if virial calculations are included; options are
		true or false.
periodic	vector	Indicate periodicity with a three-element boolean vector
		with options true or false.
neighbor	constant	Describes what kind of neighbor list algorithm is used
		during pair interaction calculations; options are sector
		or pair.
stencil	constant	Describes the kind of stencil used during pair interaction
		calculations; options are standard or multi.
skin	real	Describes the skin used during pair interaction calcula-
		tions; the skin is added to the pairwise cutoff.

shake	constant	Indicates the use of the SHAKE algorithm in subsequent codes (e.g. LAMMPS); valid options are none, auto, hydrogen, water, or all.
depth	integer	Maximum depth used for construction of ring structures during typing; allowed values are positive, where a value of 8 works in most ring cases; alternatively, an auto keyword allows for checking rings of unknown size; please note, that significant slow down occurs with the latter options for intricate ring systems.
mass	struct	Describes the site masses.
boltzmann	struct	Describes Boltzmann force fields (see Section 7.12.4 [Boltzmann], page 204).
born	struct	Describes Born force fields (see Section 7.12.5 [Born], page 209).
cff	struct	Describes CFF (Class2) force field families (see Section 7.12.6 [CFF], page 217).
charmm	struct	Describes CHARMM force fields (see Section 7.12.7 [CHARMM], page 233).
coarse	struct	Describes multiple coarse-grained force field definitions (see Section 7.12.8 [Coarse], page 244).
colloid	struct	Describes colloidal force fields (see Section 7.12.9 [Colloid], page 260).
coulomb	struct	Describes coulombic contribution definitions (see Section 7.12.10 [Coulomb], page 268).
martini	struct	Describes MARTINI coarse-grained force fields (see Section 7.12.13 [MARTINI], page 290).
gromacs	struct	Describes GROMACS force fields (see Section 7.12.12 [GROMACS], page 281).
inverse	struct	Describes settings used in applications of inverse bonded interactions.
dpd	struct	Describes DPD coarse-grained force fields (see Section 7.12.11 [DPD], page 272).
opls	struct	Describes OPLS force fields (see Section 7.12.14 [OPLS], page 298).
sdk	struct	Describes SDK force fields (see Section 7.12.15 [SDK], page 307).
spline	struct	Describes spline-based force fields (see Section 7.12.16 [Spline], page 316).
standard	struct	Describes standard force fields (see Section 7.12.17 [Standard], page 320).
table	struct	Describes tabular force fields (see Section 7.12.18 [Table], page 329).
trappe	struct	Describes tabular force fields (see Section 7.12.19 [TraPPE], page 335).

7.12.2 Usage

This variable style describes types.

7.12.3 Default

Unless otherwise stated, the default is given by

By default, all force fields are deactivated.

7.12.4 Boltzmann

7.12.4.1 Syntax

```
boltzmann = {
  bond -> struct,
  angle -> struct,
  pair -> struct
};
```

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
pair	struct	Pair interaction descriptors

The boltzmann force field uses exponential functions for bonded interactions and tabulated functions for nonbonded interactions. Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{boltzmann} = E_{bond} + E_{angle} + E_{pair}
```

for angle, bond and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/nanoparticle/coarse/ for an application.

7.12.4.2 Example

```
variables
                      -> 1*<sup>-</sup>10,
  a
                      -> "~/emc/version/force/polystyrene/smoothed_"
  prefix
};
simulation
                      = {
                      -> {
  types
     merge
                      -> true,
     boltzmann
                      -> {
        bond
                      -> {
           active
                      -> true,
                      -> {
           data
              {i0 \rightarrow m, i1 \rightarrow m, coefficients \rightarrow {
                 a \rightarrow 0.015e-10, w \rightarrow 0.09e-10, length \rightarrow 2.46e-10},
           }
        },
        angle
                      -> {
           active -> true,
                      -> {
           data
              \{i0 \rightarrow m, i1 \rightarrow m, i2 \rightarrow m, coefficients \rightarrow \{i\}\}
                    \{a \rightarrow 0.140, w \rightarrow 14.2, theta \rightarrow 147.3\},\
                    \{a \rightarrow 0.030, w \rightarrow 15.5, theta \rightarrow 158.0\}\}
```

```
}
},
pair -> {
   active -> true,
   nbonded -> 2,
   data -> {
      {i0 -> m, i1 -> m, order -> 2, name -> prefix+"m-m.m"}
   }
}
}
```

7.12.4.3 References

1. T. Spyriouni, C. Tzoumanekas, D. Theodorou, F. Mueller-Plathe, and G. Milano, "Coarse-Grained and Reverse-Mapped United-Atom Simulations of Long-Chain Atactic Polystyrene Melts: Structure, Thermodynamic Properties, Chain Conformation, and Entanglements", Macromolecules 2007, 40, 3876

7.12.4.4 Pair

```
= {
bond
  active
                 -> boolean,
 nbonded
                 -> integer,
                 -> integer,
  data
  {
    {
      i0
                 -> id,
                 -> id,
      i1
                 -> real,
      cutoff
      order
                 -> integer,
      name
                 -> string
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
cutoff	real	Defines the cutoff of the potential; takes the last data file entry when cutoff <= 0.
order	integer	Interpolation order; either 1 for linear interpolation or 2 for cubic splines.
name	string	Defines the data file name; ./force/polystyrene/ shows examples of the data file format (i.e. $\{\{x, y\},\}$).

The energetic functional form of the pair contributions to the total potential is formed by a linear interpolation or a cubic spline through the provided data.

7.12.4.5 Bond

```
= {
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      coefficients ->
      {
        {
                 -> real,
                 -> real,
          length-> real
      }
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
coef	struct	Describes individual coefficient contributions
a, w	real	Force constants
length	real	Equilibrium length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = \sum_{i=1}^{n} a_i / (2\pi w_i) \exp[-2(l - l_{0,i})^2 / w_i^2],$$

where l represents the bond length.

7.12.4.6 Angle

```
angle
  active
                -> boolean
                -> integer,
 n
 data
  {
    {
      i0
                 -> id,
      i1
                -> id,
      i2
                -> id,
      coefficients ->
      {
        {
                 -> real,
                 -> real,
          theta -> real
        },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
coef	struct	Describes individual coefficient contributions
a, w	real	Force constants
theta	real	Equilibrium angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = \sum_{i=1}^{n} a_i / (2\pi w_i) \exp[-2(\theta - \theta_{0,i})^2 / w_i^2],$$

where θ represents the bond angle.

7.12.5 Born

7.12.5.1 Syntax

```
born = {
  bond -> struct,
  angle -> struct,
  torsion -> struct,
  improper -> struct,
  pair14 -> struct,
  pair -> struct
};
```

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The OPLS force field is a compounded force field based on the Born force field.^{1–2}. Bonded terms are equivalent to OPLS force fields³. Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{opls} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}
```

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/born/ for applications.

7.12.5.2 Example

```
simulation
  types
                       -> {
                       -> true,
     merge
                       -> {
     born
                       -> {
        bond
           active
                      -> true,
                       -> {
           data
                       \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
           }
        },
                       -> {
        angle
           active
                      -> true,
           data
                       \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, k \rightarrow 100, theta \rightarrow 110}
              {i0
           }
        },
```

```
torsion
                      -> {
           active -> true,
                      -> {
           data
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
              {i0
                    \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                    \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
                 }
              }
           }
        },
        improper
                     -> {
           active
                      -> true,
           data
                      -> {
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
              {i0
           }
        },
                       -> {
        pair
           active -> true,
           nbonded -> 3,
           mode
                       -> global,
           cutoff -> 10,
                       -> berthelot,
           mix
                       -> {
           data
                       \rightarrow a, i1 \rightarrow a, sigma \rightarrow 4.0, epsilon \rightarrow 0.4},
              {i0
              {i0
                      \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
              {i0
                      -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
              {i0
                       -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
        }
     }
  }
};
```

7.12.5.3 References

- 1. F.G. Fumi and M.P. Tosi, Ionic sizes and born repulsive parameters in the NaCl-type alkali halides—I: The Huggins-Mayer and Pauling forms, J. Phys. Chem. Solids 1964, 25, 31-44.
- 2. F.G. Fumi and M.P. Tosi, Ionic sizes and born repulsive parameters in the NaCl-type alkali halides—II: The generalized Huggins-Mayer form, J. Phys. Chem. Solids 1964, 25, 31-44.
- 3. W. L. Jorgensen and J. Tirado-Rives, "The OPLS Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.5.4 Pair

```
pair
               = {
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
  shift
               -> boolean,
  coulomb
               -> option,
  cutoff
               -> real,
 mode
               -> option,
 n
               -> integer,
 data
               ->
  {
    {
      i0
               -> id,
      i1
               -> id,
      sigma
               -> real,
     rho
               -> real,
               -> real,
      a
               -> real,
      С
               -> real,
               -> real,
      core
               -> real
      cutoff
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in- teractions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants

i0, i1	id	Site id, referring to types as defined by the mass
		paragraph.
sigma	real	Interaction dependent parameter.
rho	real	Ionic pair dependent paramater.
a	real	Exponential prefactor.
С	real	Attractive parameter.
d	real	Repulsive parameter.
core	real	Distance at which the potential is treated as a purely
		repulsive potential; needed to avoid singular behavior at
		small distances.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = A \exp\left(\frac{\sigma - r_{ij}}{\rho}\right) - \frac{C}{r_{ij}^6} + \frac{D}{r_{ij}^8},$$

for $r_{ij} > r_{core}$, where r_{ij} represents the distance between site i and site j.

7.12.5.5 Bond

```
= {
bond
  active
                 -> boolean
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = k_{bond}(l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.5.6 Angle

```
= {
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = k_{angle}(\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.5.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active n	Parameters boolean integer	Description Interaction activator; either true or false. Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine
		function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} 1/2 \ k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.5.8 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive active n	Parameters boolean integer	Description Interaction activator; either true or false. Number of data entries; can be ommitted when type = {merge -> true}.
data i0, i1, i2, i3	struct id	Summary of interaction constants Site id, referring to types as defined by the mass paragraph in types
k psi	real real	Force constant Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.6 CFF

7.12.6.1 Syntax

```
= {
 pair
                         -> struct,
 bond
                         -> struct,
 angle
                         -> struct,
 bond_bond
                         -> struct,
 bond_angle
                         -> struct,
  torsion
                         -> struct,
  end_bond_torsion
                         -> struct,
 middle_bond_torsion
                        -> struct,
 bond_bond_13
                        -> struct,
  angle_torsion
                        -> struct,
  angle_angle_torsion
                        -> struct,
  improper
                        -> struct,
  angle_angle
                        -> struct
};
```

Directive	Parameters	Description
pair	struct	Pair interaction descriptors
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
bond_bond	struct	Bond-bond interaction descriptors
bond_angle	struct	Bond-angle interaction descriptors
torsion	struct	Torsion interaction decriptors
end_bond_	struct	End bond torsion interaction decriptors
torsion		
middle_	struct	Middle bond torsion interaction decriptors
bond_		
torsion		
bond_bond_	struct	Bond-bond-13 interaction decriptors
13		
angle_	struct	Angle-torsion interaction decriptors
torsion		<u>.</u>
angle_	struct	Angle-angle-torsion interaction decriptors
angle_		
torsion		
improper	struct	Improper interaction decriptors
angle_angle	struct	Angle-angle interaction decriptors
9 0		• •

The CFF force field, as described by Sun,¹ uses a combination of standard and cross-coupled bonded interactions combined with Lennard-Jones 6-9 nonbonded interactions. Its functional form is a summation of bonded and nonbonded interactions given by

```
\begin{split} E_{cff} &= E_{bond} + E_{angles} + E_{torsions} + E_{impropers} + E_{pair} \\ E_{angles} &= E_{angle} + E_{bond-bond} + E_{bond-angle} \\ E_{torsions} &= E_{torsion} + E_{end-bond-torsion} + E_{middle-bond-torsion} + E_{bond-bond-13} + E_{angle-torsion} \\ &\quad + E_{angle-angle-torsion} \\ E_{impropers} &= E_{improper} + E_{angle-angle} \end{split}
```

for bond, angle, torsion, improper, and pair contributions respectively. The following paragraphs describe each contribution in detail. Note, that improper contributions are calculated differently (see Improper). Force field typing is available for the PCFF force field.

7.12.6.2 Examples

The following example creates topology for water and types these sites with the PCFF force field as included with EMC,

```
= {
field
                -> cff,
 mode
                -> {$root+"pcff/pcff_templates.dat", $root+"pcff/pcff.frc"
 name
};
groups
                = {
                -> {id -> water, chemistry -> "0"}
  group
clusters
                -> {id -> water, system -> main, group -> water, n -> 1000}
  cluster
};
field
                = {
 mode
                -> apply
};
```

Application of the field scripting command saves the need for manual force field typing (see Section 5.11 [Field], page 103). However, CFF-type force field alterations or additions can also be entered directly through

```
active \rightarrow true, n \rightarrow 0},
        angle_torsion -> {
           active \rightarrow true, n \rightarrow 0},
                      -> {
        bond
           active \rightarrow true, n \rightarrow 1, data \rightarrow
                      \rightarrow o*, i1 \rightarrow hw, 1 \rightarrow 0.97, k \rightarrow {563.28, -1428.22,
                 1902.12}}},
        bond_angle -> {
           active \rightarrow true, n \rightarrow 1, data \rightarrow
              \{i0 \rightarrow hw, i1 \rightarrow o*, i2 \rightarrow hw, k \rightarrow \{22.35, 22.35\}\}\},
        bond_bond -> {
           active \rightarrow true, n \rightarrow 1, data \rightarrow
              \{i0 \rightarrow hw, i1 \rightarrow o*, i2 \rightarrow hw, k \rightarrow -9.5\}\},
        bond_bond_13 -> {
           active \rightarrow true, n \rightarrow 0},
        end_bond_torsion -> {
           active \rightarrow true, n \rightarrow 0},
        improper -> {
           active \rightarrow true, n \rightarrow 0},
        middle_bond_torsion -> {
           active \rightarrow true, n \rightarrow 0},
        pair -> {
           active -> true, mix -> sixth, shift -> false, coulomb -> cut,
           nbonded \rightarrow 2, mode \rightarrow repulsive, cutoff \rightarrow 9.5, n \rightarrow 3, data \rightarrow {
                      -> o*, i1 -> o*, epsilon -> 0.274, sigma -> 3.608},
                      \rightarrow o*, i1 \rightarrow hw, epsilon \rightarrow 0.00336154972409,
                 sigma -> 3.21478799199},
              {i0
                      -> hw, i1 -> hw, epsilon -> 0.013, sigma -> 1.098}}},
        torsion
                      -> {
           active \rightarrow true, n \rightarrow 0}},
  }
};
```

The above definition would result from application of the PCFF force field.

7.12.6.3 References

1. H. Sun, "COMPASS: An ab Initio Force-Field Optimized for Condensed-Phase Applications - Overview with Details on Alkane and Benzene Compounds", J. Phys. Chem. B 1998, 102, 7338.

7.12.6.4 Pair

```
= {
pair
 active
             -> boolean,
 nbonded
              -> integer,
 mix
               -> option,
 shift
               -> boolean,
 coulomb
              -> option,
 cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
   {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
   },
   . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants.
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-9 potential.
sigma	real	Site size; the location of the potential well.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-9 Lennard-Jones potential,

$$E_{pair} = \epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[2\left(\frac{\sigma}{r_{ij}}\right)^3 - 3\right],$$

where r_{ij} represents the distance between site i and site j. The nonbond interactions, which include Lennard-Jones and coulombic interactions, encompass all pairs between atoms which are separated by two or more bonded atoms, i.e. nbonded = 2.

7.12.6.5 Bond

```
= {
bond
  active
                 -> boolean
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      1
                 -> real,
                 -> {real, real, real}
      k
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
1	real	Equilibrium length
k	real	Defines up to three bond constants

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = \sum_{i=1}^{3} k_i (l - l_{0,i})^{i+1},$$

where l represents the bond length of bond $\{i_0, i_1\}$.

7.12.6.6 Angle

```
= {
angle
  active
                 -> boolean
                 -> integer,
  n
  data
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id
      theta
                 -> real,
                 -> {real, real, real}
    },
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
theta	real	Equilibrium angle
k	real	Defines up to three angle constants

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = \sum_{i=1}^{3} k_i (\theta - \theta_{0,i})^{i+1},$$

where θ represents the angle between bonds $\{i_0,i_1\}$ and $\{i_1,i_2\}$.

7.12.6.7 Bond-Bond

```
bond_bond
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id
                 -> real
      k
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
k	real	Defines the bond-bond cross term constant

The energetic functional form of the bond-bond cross-coupling contributions to the total potential is described by

$$E_{bond-bond} = k(l_1 - l_{0.1})(l_2 - l_{0.2}),$$

where l_1 represents the length of bond $\{i_0, i_1\}$ with equilibrium value $l_{0,1}$ and l_2 the length of bond $\{i_1, i_2\}$ with equilibrium value $l_{0,2}$. Equilibrium values reference the bond contributions.

7.12.6.8 Bond-Angle

```
= {
bond_angle
                 -> boolean
  active
  n
                 -> integer,
  data
                 ->
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id
                 -> {real, real}
      k
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
k	real	Defines the bond-angle cross term constants

The energetic functional form of the bond-angle cross-coupling contributions to the total potential is described by

$$E_{bond-angle} = (k_1(l_1 - l_{0,1}) + k_2(l_2 - l_{0,2}))(\theta - \theta_{0,i}),$$

where l_1 represents the length of bond $\{i_0, i_1\}$ with equilibrium value $l_{0,1}$, l_2 the length of bond $\{i_1, i_2\}$ with equilibrium value $l_{0,2}$, and θ the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$. Equilibrium values reference both bond and angle contributions.

7.12.6.9 Torsion

```
torsion
                -> boolean
  active
                -> integer,
 n
 data
  {
    {
      i0
                -> id,
      i1
                -> id,
      i2
                -> id,
      i3
                -> id,
      coefficients ->
      {
        {k
                -> real, n -> integer, delta -> real},
                -> real, n -> integer, delta -> real},
        {k
                -> real, n -> integer, delta -> real}
        {k
      }
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to mass paragraph in types
i3		
coefficients	struct	List of up to three sets of coefficients
k	real	Torsion constants
n	integer	Torsion prefactor
delta	real	Torsion offset

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{3} k_i (1 - cos[n_i \phi - \delta]),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}, \{i_1, i_2\},$ and $\{i_2, i_3\}.$

7.12.6.10 End-Bond-Torsion

```
end_bond_torsion = {
                -> boolean
  active
 n
                -> integer,
 data
  {
    {
      i0
                -> id,
      i1
                -> id,
                -> id,
      i2
                -> id,
      i3
                -> {real, real, real, real, real}
      k
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to mass paragraph in types
i 3		
k	real	Three left $(k_1 \text{ through } k_3)$ and three right $(k_4 \text{ through } k_3)$
		k_6) interaction constants

The energetic functional form of the end-bond-torsion cross-coupling contributions to the total potential is described by

$$E_{end-bond-torsion} = \sum_{i=1}^{3} (k_i(l_1 - l_{0,1}) + k_{i+3}(l_2 + l_{0,2})) \cos[i\phi],$$

where l_1 represents the length of bond $\{i_0, i_1\}$ with equilibrium value $l_{0,1}$, l_2 represents the length of bond $\{i_2, i_3\}$ with equilibrium value $l_{0,2}$, and ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$.

7.12.6.11 Middle-Bond-Torsion

```
middle_bond_torsion = {
  active
                 -> boolean
                 -> integer,
  n
  data
  {
    {
                 -> id,
      i0
      i1
                 -> id,
                 -> id,
      i2
                 -> id,
      i3
                 -> {real, real, real}
      k
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
	_	{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to mass paragraph in types
i3		
k	real	Three $(k_1 \text{ through } k_3)$ interaction constants

The energetic functional form of the middle-bond-torsion cross-coupling contributions to the total potential is described by

$$E_{end-bond-torsion} = \sum_{i=1}^{3} k_i (l_1 - l_{0,1}) \cos[i\phi],$$

where l_1 represents the length of bond $\{i_1,i_2\}$ with equilibrium value $l_{0,1}$ and ϕ represents the bond torsion constructed by bonds $\{i_0,i_1\}$, $\{i_1,i_2\}$, and $\{i_2,i_3\}$.

7.12.6.12 Bond-Bond-13

```
= {
bond_bond_13
 active
                 -> boolean
                 -> integer,
 n
 data
                 ->
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants.
i0, i1, i2,	id	Site id, referring to mass paragraph in types.
i 3		
k	real	Bond-Bond-13 interaction constant.

The energetic functional form of the bond-bond-13 cross-coupling contributions to the total potential is described by

$$E_{bond-bond-13} = k_{bond-bond-13}(l_1 - l_{0,1})(l_2 - l_{0,2}),$$

where l_1 represents the length of bond $\{i_0, i_1\}$ with equilibrium value $l_{0,1}$, l_2 represents the length of bond $\{i_2, i_3\}$ with equilibrium value $l_{0,2}$.

7.12.6.13 Angle-Torsion

```
angle_torsion
 active
                -> boolean
 n
                -> integer,
 data
                ->
  {
    {
                -> id,
      i0
      i1
                -> id,
                -> id,
      i2
                -> id,
      i3
                -> {real, real, real, real, real}
      k
   },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants.
i0, i1, i2,	id	Site id, referring to mass paragraph in types.
i3		
k	real	Six $(k_1 \text{ through } k_6)$ interaction constants.

The energetic functional form of the angle-torsion cross-coupling contributions to the total potential is described by

$$E_{angle-torsion} = \sum_{i=1}^{2} (\theta_i - \theta_{0,i}) \sum_{j=1}^{3} k_{3(i-1)+j} \cos[j\phi],$$

where θ_1 the angle between bonds $\{i_0,i_1\}$ and $\{i_1,i_2\}$, θ_2 the angle between bonds $\{i_1,i_2\}$ and $\{i_2,i_3\}$, and ϕ the bond torsion constructed by bonds $\{i_0,i_1\}$, $\{i_1,i_2\}$, and $\{i_2,i_3\}$.

7.12.6.14 Angle-Angle-Torsion

```
angle_torsion
  active
                 -> boolean
  n
                 -> integer,
  data
                 ->
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      iЗ
                 -> id,
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to mass paragraph in types
i3		

The energetic functional form of the angle-angle-torsion cross-coupling contributions to the total potential is described by

```
E_{angle-angle-torsion} = k_{angle-angle-torsion}((\theta_1 - \theta_{0,1})((\theta_2 - \theta_{0,2})\cos[\phi],
```

where θ_1 the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, θ_2 the angle between bonds $\{i_1, i_2\}$ and $\{i_2, i_3\}$, and ϕ the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$.

7.12.6.15 Improper

```
angle
  active
                  -> boolean
                  -> integer,
  n
  data
                  ->
  {
    {
      i0
                  -> id,
      i1
                  -> id,
      i2
                  -> id,
      i3
                  -> id,
      k
                  -> real,
                  -> real
      psi
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}, \{i_2, i_0\}, \text{ and } \{i_3, i_0\}$. EMC represents site i_0 as the central site, as opposed to some representations, for which i_1 is the central site. Note, that each improper contributes to the total improper energy individually. Note, that Discover and LAMMPS use the average of the three contributing angles ψ and calculate the improper energy accordingly. EMC, however, uses the average of the energy for all three contributing factors.

7.12.7 CHARMM

7.12.7.1 Syntax

```
= {
charmm
 pair
                 -> struct,
 bond
                 -> struct,
  angle
                 -> struct,
  urey
                 -> struct,
  torsion
                 -> struct,
 pair14
                 -> struct,
 improper
                 -> struct
};
```

Directive	Parameters	Description
pair	struct	Pair interaction descriptors
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
urey	struct	Urey 1-3 interaction decriptors
torsion	struct	Torsion interaction decriptors
pair14	struct	Pair 1-4 interaction descriptors
improper	struct	Improper interaction descriptors

The CHARMM force field as included in EMC is based on the CHARMM description as defined by MacKerrell et al. EMC provides force field files, which are derived from the original CHARMM force field files, but are governed by typing rules, rather than residue templates, as is the case in the original definition. An included script (./scripts/charmm.pl) derives typing rules from the original CHARMM force field files in an automated fashion. The force field files are found in ./field/charmm. Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{charmm} = E_{bond} + E_{angle} + E_{urey} + E_{torsion} + E_{improper} + E_{pair14} + E_{pair}$$

for bond, angle, urey, torsion, pair14, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/charmm/ for applications.

7.12.7.2 Examples

The following example creates topology for water and types these sites with the CHARMM force field as included with EMC,

Application of the field scripting command saves the need for manual force field typing (see Section 5.11 [Field], page 103). However, CHARMM force field alterations or additions can also be entered directly through

```
= {
simulation
                    -> {
  types
                    -> true,
     merge
     charmm
                    -> {active -> true,
       angle
                    -> {active -> true, n -> 1, data ->
          {i0
                    \rightarrow HT, i1 \rightarrow OT, i2 \rightarrow HT, k \rightarrow 55, theta \rightarrow 104.52}},
                    -> {active -> true, n -> 1, data ->
       bond
          {i0
                    \rightarrow OT, i1 \rightarrow HT, k \rightarrow 450, 1 \rightarrow 0.9572}},
       improper \rightarrow {active \rightarrow true, n \rightarrow 0},
                    -> {active -> true, mix -> berthelot, shift -> false,
       pair
          coulomb -> none, nbonded -> 3, mode -> individual, inner -> 0,
          cutoff -> 9.5, n -> 3, data -> \{
             {i0
                    -> OT, i1 -> OT, epsilon -> 0.1521, sigma -> 3.15057422683},
                    \rightarrow OT, i1 \rightarrow HT, epsilon \rightarrow 0.0836456812992,
             {i0
               sigma -> 1.77529387564},
                  -> HT, i1 -> HT, epsilon -> 0.046,
               sigma -> 0.400013524445}}},
                    \rightarrow {active \rightarrow true, n \rightarrow 3, data \rightarrow {
       pair14
                    \rightarrow OT, i1 \rightarrow OT, epsilon \rightarrow 0.1521, sigma \rightarrow 3.15057422683},
             {i0
                  -> OT, i1 -> HT, epsilon -> 0.0836456812992,
               sigma -> 1.77529387564},
             \{i0 \rightarrow HT, i1 \rightarrow HT, epsilon \rightarrow 0.046,
               sigma -> 0.400013524445}}},
       torsion \rightarrow {active \rightarrow true, n \rightarrow 0},
                    -> {active -> true, n -> 1, data ->
       urey
          {i0
                    \rightarrow HT, i1 \rightarrow OT, i2 \rightarrow HT, k \rightarrow 0, 1 \rightarrow 0}},
  }
};
```

7.12.7.3 References

1. A. D. MacKerrell, Jr. et al., "All-Atom Empirical Potential for Molecular Modeling

and Dynamics Studies of Proteins", J. Phys. Chem. B 1998, 102, 3586-3616.

7.12.7.4 Pair

```
= {
pair
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
               -> boolean,
  shift
  coulomb
               -> option,
  inner
               -> real,
  cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
  data
  {
    {
      i0
               -> id,
      i1
               -> id,
      epsilon
               -> real,
      sigma
               -> real,
      inner
               -> real,
      cutoff
               -> real
    },
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in- teractions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
inner	real	Sets global inner start of cut off for global mode.
cutoff	real	Sets global cut off for global mode.
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs
		per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type =
11	integer	{merge -> true}in which case the entered entries are merged with already existing ones.
data	struct	Summary of interaction constants.

i0, i1	id	Site id, referring to types as defined by the mass
		paragraph.
epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
inner	real	Sets pairwise inner start of cut off; can be omitted upon
		global inner definition.
cutoff	real	Sets pairwise cut off; can be omitted upon global $cutoff$
		definition.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between sites i and j.

7.12.7.5 Bond

```
= {
bond
  active
                 -> boolean
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = k_{bond}(l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.7.6 Angle

```
= {
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = k_{angle}(\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.7.7 Urey

```
= {
urey
                 -> boolean
  active
                 -> integer,
 n
  data
                 ->
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Urey bond spring constants
1	real	Equilibrium urey bond length

The energetic functional form of the Urey-Bradley contributions to the total potential is described by

$$E_{urey} = k_{urey}(l - l_0)^2,$$

where k_{urey} represents a spring constant, l the urey bond length of bond $\{i_0, i_2\}$, and l_0 the equilibrium bond length.

7.12.7.8 Torsion

```
torsion
                 -> boolean
  active
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
           delta -> real
        }
      }
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to mass paragraph in types

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \cos(n_i \phi - \delta_i)),$$

where k_i represents a set of torsion constants, δ_i a torsion offset, n_i the torsion pre-factor, and ϕ the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. The maximum allowed value of n_i is 6.

7.12.7.9 Pair14

```
pair
  {\tt active}
                   -> boolean,
  {\tt mix}
                   -> option,
                   -> integer,
  data
  {
    {
       i0
                   -> id,
                   -> id,
       i1
                  -> real,
       epsilon
                   -> real
       sigma
    },
  }
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}in which case the entered entries are merged with already existing ones.
data	struct	Summary of interaction constants.
i0, i1	id	Site id, referring to types as defined by the mass paragraph.
epsilon	real	Force interaction constant; identical to the well depth for a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{l}\right)^6 \left[\left(\frac{\sigma}{l}\right)^6 - 1\right],$$

where l represents the distance between sites i0 and i1.

7.12.7.10 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.8 Coarse

7.12.8.1 Syntax

```
coarse
  1j
                 -> struct,
  repulsive
                 -> struct,
  sphere
                 -> struct,
  colloid
                 -> struct,
  dpd
                 -> struct,
  charge
                 -> struct,
  fene
                 -> struct,
                 -> struct
  angle
};
```

Directive	Parameters	Description
lj	struct	Hard sphere core with Lennard-Jones shell interaction
		descriptors
repulsive	struct	Hard sphere core with r^-6 repulsive shell interaction
		descriptors
sphere	struct	Coarse-grained sphere interaction descriptors (Girifalco ¹)
colloid	struct	Colloidal interaction descriptors (Everaers and
		Ejtehadi ²)
dpd	struct	DPD interaction descriptors (Groot and Warren ³)
charge	struct	Coarse-grained charge interaction decriptors (internal
		form)
fene	struct	FENE interaction descriptors (Kremer and Grest ⁴)
angle	struct	Angle interaction descriptors

The coarse force field represents a set of coarse-grained interations functions. Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{coarse} = E_{LJ} + E_{repulsive} + E_{sphere} + E_{colloid} + E_{DPD} + E_{charge} + E_{FENE} + E_{angle}$$

for hard core Lennard-Jones, hard core repulsive, colloidal, coarse-grained sphere angle, DPD, charge, FENE, and angle contributions respectively. The following paragraphs describe each contribution in detail. Typically, all conributions are not used simulataneously. Note, that the DPD contributions are included for backwards compatibility. Use the DPD force field when typing and transferring to e.g. LAMMPS. See ./examples/nanoparticle/coarse/ for applications.

7.12.8.2 Examples

```
variables = {
};
simulation = {
```

```
-> {
  types
    merge
                 -> true,
    coarse
                 -> {
                 -> {
      dpd
                -> true,
        active
        data
                 -> {
    }
 }
};
simulation
                 = {
                 -> {
  types
                 -> true,
    merge
    coarse
                 -> {
      colloid
                 -> {
        active
                -> true,
        data
                 -> {
        }
      }
    }
 }
};
```

7.12.8.3 References

- 1. L. A. Girifalco, "Molecular Properties of C₆₀ in the Gas and Solid Phases" J. Chem. Phys. **1992**, 96, 858-861.
- 2. R. Everaers and M. R. Ejtehadi, "Interaction potentials for soft and hard ellipsoids", Phys. Rev. E. 2003, 67, 041710.
- 3. R. D. Groot and P. B. Warren, "Dissipative particle dynamics: Bridging the gap between atomistic and mesoscopic simulation", J. Chem. Phys. 1997, 107, 4423-4435.
- 4. K. Kremer and G. S. Grest, J. Chem. Phys. 1990, 92, 5057.

7.12.8.4 LJ

```
= {
1j
 active
              -> boolean,
               -> integer,
 nbonded
 mix
               -> option,
 shift
               -> boolean,
 cutoff
               -> real,
 mode
               -> option,
 n
               -> integer,
 data
  {
   {
     i0
               -> id,
               -> id,
     i1
     epsilon
               -> real,
     sigma
               -> real,
     core
               -> real,
     cutoff
               -> real
   },
   . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.
epsilon	real	Force interaction constant; identical to the well depth for a Lennard-Jones 6-12 potential
sigma	real	Site size; the point where the potential equals zero

core	real	Site hard core size
cutoff	real	Sets the local pairwise cut off

The energetic functional form of the pair contributions to the total potential is described by a hard core, extended with a 6-12 Lennard-Jones potential,

$$E_{LJ} = 4\epsilon \left(\frac{\sigma}{r_{ij} - d_{core}}\right)^6 \left[\left(\frac{\sigma}{r_{ij} - d_{core}}\right)^6 - 1\right] \text{ for } r_{ij} \ge d_{core},$$

where r_{ij} represents the distance between site i and site j, and d_{core} its core size.

7.12.8.5 Repulsive

```
repulsive
  active
                 -> boolean,
  nbonded -> integer,
mix -> option,
shift -> boolean,
  shift
                 -> boolean,
  cutoff
                -> real,
  mode
                 -> option,
  n
                 -> integer,
  data
                 ->
  {
    {
      i0
                 -> id,
                 -> id,
      i1
                 -> real,
      epsilon
                 -> real,
      sigma
      core
                 -> real
      cutoff
                 -> real
    },
    . . .
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.
epsilon	real	Force interaction constant; identical to the well depth for a Lennard-Jones 6-12 potential
sigma	real	Site size; the point where the potential equals zero

core real Site hard core size

The energetic functional form of the pair contributions to the total potential is described by a hard core, extended with a repulsive potential,

$$E_{repulsive} = 4\epsilon \left(\frac{\sigma}{r_{ij} - d_{core}}\right)^{12} \text{ for } r_{ij} \ge d_{core},$$

where r_{ij} represents the distance between site i and site j, and d_{core} its core size.

7.12.8.6 Sphere

```
1j
 active
               -> boolean,
 active -> boolean,
nbonded -> integer,
mix -> option,
 cutoff
                -> real,
 mode
                -> option,
                -> integer,
 n
 data
  {
    {
                -> id,
      i0
      i1
                -> id,
      epsilon
                 -> real,
      sigma
                -> real,
      d
                 -> real,
                -> real,
      n
                -> real
      cutoff
    },
    . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
mix	option	Sets mixing rule; options are none, berthelot,
		arithmetic, geometric, and sixth.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of
		cutoff for all contributions, individual defining cut offs
		per contribution, and repulsive defining the well loca-
		tion for each separate contribution as the cut off distance
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass
		paragraph.
epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential
d	real	Sets the diameter of both particles
n	real	Sets the number of contributing Lennard-Jones beads to
		each spherical FCC particle

cutoff real Sets the local pairwise cut off

The energetic functional form of an integration of two interacting spherical FCC lattices consisting of n Lennard-Jones particles,

$$E_{LJ} = 4 \epsilon n^2 \left(\frac{\sigma}{d}\right)^6 \left[\frac{1}{12} \left(\frac{1}{s(s-1)^3} + \frac{1}{s(s+1)^3} - \frac{2}{s^4}\right) + \frac{1}{90} \left(\frac{\sigma}{d}\right)^6 \left(\frac{1}{s(s-1)^9} + \frac{1}{s(s+1)^9} - \frac{2}{s^{10}}\right)\right],$$

where ϵ and σ are the standard Lennard-Jones constants, $s = r_{ij}/d$, and r_{ij} represents the distance between the particle centers.

7.12.8.7 Colloid

```
colloid
  active
                 -> boolean,
 nbonded
                 -> integer,
                 -> integer,
 data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      Α
      sigma
                 -> real,
                 -> real,
      d2
                 -> real,
                 -> real
      cutoff
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
A	real	Pairwise Hamaker constants
sigma	real	Pairwise Lennard-Jones sigma
d1, d2	real	Respective colloidal particle diameter
cutoff	real	Pairwise potential cutoff

The energetic functional form of the pair contributions to the total potential, as described by Everaers and Ejtehadi, is formed by the sum of attractive and repulsive energy:

$$E = E_A + E_R$$

Attractive energy is expressed by

$$E_A = -\frac{A}{6} \left[\frac{2d_i d_j}{r_{ij}^2 - (d_i + d_j)^2} + \frac{2d_i d_j}{r_{ij}^2 - (d_i - d_j)^2} + \ln \left(\frac{r_{ij}^2 - (d_i + d_j)^2}{r_{ij}^2 - (d_i - d_j)^2} \right) \right],$$

where A is referred to as Hamaker's constant, d_i and d_j represent the diameter sites i and j, and r_{ij} represents the distance between sites. Repulsive energy is expressed by

$$E_{R} = \frac{A}{37800} \frac{\sigma^{6}}{r_{ij}} \left[\frac{r_{ij}^{2} - 7r_{ij}(d_{i} + d_{j}) + 6\left(d_{i}^{2} + 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} - d_{i} - d_{j})^{7}} + \frac{r_{ij}^{2} + 7r_{ij}(d_{i} + d_{j}) + 6\left(d_{i}^{2} + 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} + d_{i} + d_{j})^{7}} - \frac{r_{ij}^{2} - 7r_{ij}(d_{i} - d_{j}) + 6\left(d_{i}^{2} - 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} - d_{i} + d_{j})^{7}} \right],$$

where σ represents the Lennard-Jones (LJ) constant sigma. Hamaker's constant A in LJ units is given by $A = 4\pi\epsilon_{LJ} (\rho\sigma^3)^2$, with ρ representing reduced density.

7.12.8.8 DPD

```
= {
pair
  active
                  -> boolean,
  {\tt mix}
                  -> option,
  nbonded
                 -> integer,
  mode
                  -> option,
  cutoff
                  -> real,
  n
                  -> integer,
  data
  {
    {
      i0
                  -> id,
      i1
                  -> id,
                  -> real,
                  -> real,
      gamma
      {\tt cutoff}
                  -> real
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
mix	option	Sets mixing rule; options are none, berthelot,
		arithmetic, geometric, and sixth.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
mode	option	Sets cut off mode; options are global using the value of
		cutoff for all contributions, individual defining cut offs
		per contribution, and repulsive defining the well loca-
		tion for each separate contribution as the cut off distance.
cutoff	real	Sets global cut off for global mode
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass
		paragraph.
a	real	Force interaction constant.
gamma	real	Force damping constant.
cutoff	real	Site size; the point where the potential equals zero.

The energetic functional form of pair contributions to the total soft potential is described by a repulsive Hertzian spring,

$$E_{pair} = 1/2 \ a \ r_{ij,cutoff} \left(1 - r_{ij}/r_{ij,cutoff}\right)^2, \ r_{ij} < r_{ij,cutoff}$$

where a represents the pairwise interaction paramater, r_{ij} the distance between site i and site j, and $r_{ij,cutoff}$ the cutoff distance between these two sites.

7.12.8.9 Charge

```
charge
  active
                 -> boolean,
 nbonded
                 -> integer,
 mix
                 -> option,
  shift
                 -> boolean,
                 -> integer,
 n
  data
  {
    {
                 -> id,
      i0
                 -> id,
      i1
      a
                 -> real,
                 -> real,
      kappa
      d1
                 -> real,
      d2
                 -> real,
      cutoff
                 -> real
    },
};
```

Directive active nbonded	Parameters boolean integer	Description Interaction activator; either true or false. Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
a	real	Model constant
kappa	real	Electrostatic screening factor
d1	real	Sets the diameter of particle i0
d2	real	Sets the diameter of particle i1
cutoff	real	Sets the local pairwise cut off

The energetic functional form of the charge contributions to the total potential, which is based on a Yukawa potential. It is described by

$$E_{charge} = A \ln (1 + e^{-\kappa H}) (4 - 2H/d_1 - 2H/d_2),$$

where d_1 and d_2 represent particle diameters, $H = r_{ij} - \frac{1}{2} (d_1 + d_2)$, and A the electrostatic constant, which can be expressed in terms of surface potentials ψ_1 and ψ_2 ,

$$A = \pi \epsilon_0 \epsilon_r (\psi_1 + \psi_2)^2 d_1 d_2 / (d_1 + d_2) / 8,$$

assuming, that both surface potentials are approximately the same.

7.12.8.10 FENE

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = -\tfrac{1}{2} \ k \ l_0^2 \ \ln \left[1 - (l/l_0)^2 \right],$$

where k represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.8.11 Angle

```
angle
 active
               -> boolean
 n
                -> integer,
 data
                ->
 {
   {
      i0
               -> id,
     i1
               -> id,
      i2
                -> id
    },
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types

The energetic functional form of the bond contributions to the total potential is described by

$$E_{angle} =$$

where

7.12.9 Colloid

7.12.9.1 Syntax

Parameters	Description
struct	Pairwise colloidal interaction descriptors
struct	Pairwise charge interaction descriptors
struct	Bond interaction descriptors
struct	Angle interaction decriptors
	struct struct struct

The colloid force field is a compounded force field, that uses Hamaker and charge screening interactions. Its functional form is a summation of nonbonded and bonded interactions given by

```
E_{colloid} = E_{pair} + E_{charge} + E_{bond} + E_{angle}
```

for pair, charge, bond, and angle contributions respectively. The following paragraphs describe each contribution in detail.

7.12.9.2 Example

```
simulation
                     -> {
  types
                     -> true,
     merge
     standard
                     -> {
                     -> {
       bond
          active -> true,
          data
                     \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
          }
       },
                     -> {
       angle
          active -> true,
                     -> {
          data
                     \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, k \rightarrow 400, theta \rightarrow 110}
          }
       },
                     -> {
       pair
          active -> true,
          inner
                     -> 1.0001,
          outer
                     -> 1.25,
```

```
nbonded -> 2,
                  -> {
         data
           {i0
                  -> a, i1 -> a,
                  \rightarrow 1e-21, d1 \rightarrow 1e-7, d2 \rightarrow 1e-7, cutoff \rightarrow 2.5e-7},
             Α
           {i0
                 -> a, i1 -> b,
                  -> 1.1e-21, d1 -> 1e-7, d2 -> 2e-7, cutoff -> 3.75e-7},
             Α
           {i0
                  -> a, i1 -> c,
                  -> 1.2e-21, d1 -> 1e-7, d2 -> 3e-7, cutoff -> 5e-7},
             Α
           {i0
                  -> b, i1 -> b,
                  -> 1.2e-21, d1 -> 2e-7, d2 -> 2e-7, cutoff -> 5e-7},
             Α
           {i0
                  -> b, i1 -> c,
                  \rightarrow 1.3e-21, d1 \rightarrow 2e-7, d2 \rightarrow 3e-7, cutoff \rightarrow 6.25e-7},
             Α
           {i0
                 -> c, i1 -> c,
                  -> 1.4e-21, d1 -> 3e-7, d2 -> 3e-7, cutoff -> 7.5e-7}
      }
    }
 }
};
```

7.12.9.3 References

1. R. Everaers and M. R. Ejtehadi, "Interaction potentials for soft and hard ellipsoids", Phys. Rev. E. 2003, 67, 041710.

7.12.9.4 Pair

```
= {
pair
  active
                 -> boolean,
  nbonded
                 -> integer,
                 -> integer,
  inner
                 -> real,
  outer
                 -> real,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      Α
                 -> real,
                 -> real,
      sigma
      d1
                 -> real,
      d2
                 -> real,
      cutoff
                 -> real
    },
};
```

Directive active nbonded	Parameters boolean integer	Description Interaction activator; either true or false. Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
inner	real	Fraction of diameter defining the inner interaction rim; must be a number larger than 1.
outer	real	Fraction of diameter defining the outer interaction rim; must be a number larger than 1.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
A	real	Pairwise Hamaker constants
sigma	real	Pairwise Lennard-Jones sigma
d1, d2	real	Respective colloidal particle diameter
cutoff	real	Pairwise potential cutoff

The energetic functional form of the pair contributions to the total potential, as described by Everaers and Ejtehadi, is formed by the sum of attractive and repulsive energy:

$$E = E_A + E_R$$

Attractive energy is expressed by

$$E_A = -\frac{A}{6} \left[\frac{2d_i d_j}{r_{ij}^2 - (d_i + d_j)^2} + \frac{2d_i d_j}{r_{ij}^2 - (d_i - d_j)^2} + \ln \left(\frac{r_{ij}^2 - (d_i + d_j)^2}{r_{ij}^2 - (d_i - d_j)^2} \right) \right],$$

where A is referred to as Hamaker's constant, d_i and d_j represent the diameter sites i and j, and r_{ij} represents the distance between sites. Repulsive energy is expressed by

$$E_{R} = \frac{A}{37800} \frac{\sigma^{6}}{r_{ij}} \left[\frac{r_{ij}^{2} - 7r_{ij}(d_{i} + d_{j}) + 6\left(d_{i}^{2} + 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} - d_{i} - d_{j})^{7}} + \frac{r_{ij}^{2} + 7r_{ij}(d_{i} + d_{j}) + 6\left(d_{i}^{2} + 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} + d_{i} + d_{j})^{7}} - \frac{r_{ij}^{2} - 7r_{ij}(d_{i} - d_{j}) + 6\left(d_{i}^{2} - 7d_{i}d_{j} + d_{j}^{2}\right)}{(r_{ij} - d_{i} + d_{j})^{7}} \right],$$

where σ represents the Lennard-Jones (LJ) constant sigma. Hamaker's constant A in LJ units is given by $A = 4\pi\epsilon_{LJ} (\rho\sigma^3)^2$, with ρ representing reduced density.

7.12.9.5 Charge

```
charge
  active
                 -> boolean,
 nbonded
                 -> integer,
 mix
                 -> option,
  shift
                 -> boolean,
                 -> integer,
 n
  data
  {
    {
                 -> id,
      i0
                 -> id,
      i1
      a
                 -> real,
                 -> real,
      kappa
      d1
                 -> real,
      d2
                 -> real,
      cutoff
                 -> real
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
a	real	Model constant
kappa	real	Electrostatic screening factor
d1	real	Sets the diameter of particle i0
d2	real	Sets the diameter of particle i1
cutoff	real	Sets the local pairwise cut off

The energetic functional form of the charge contributions to the total potential, which is based on a Yukawa potential. It is described by

$$E_{charge} = A \ln (1 + e^{-\kappa H}) (4 - 2H/d_1 - 2H/d_2),$$

where d_1 and d_2 represent particle diameters, $H = r_{ij} - \frac{1}{2} (d_1 + d_2)$, and A the electrostatic constant, which can be expressed in terms of surface potentials ψ_1 and ψ_2 ,

$$A = \pi \epsilon_0 \epsilon_r (\psi_1 + \psi_2)^2 d_1 d_2 / (d_1 + d_2) / 8,$$

assuming, that both surface potentials are approximately the same.

7.12.9.6 Bond

```
= {
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2 \ k_{bond} (l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.9.7 Angle

```
= {
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2 \ k_{angle} (\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.10 Coulomb

7.12.10.1 Syntax

DirectiveParametersDescriptionchargestructDiffuse coarse-grained charge interaction descriptorspairstructPair interaction descriptors

The coulomb force field is a collection of electrostatic interactions. Its functional form is a summation of nonbonded interactions given by

$$E_{coulomb} = E_{pair} + E_{charge}$$

for charge and pair contributions respectively. The following paragraphs describe each contribution in detail.

7.12.10.2 Examples

```
variables
};
                = {
simulation
  types
                -> {
   merge
                -> true,
    coulomb
                -> {
      charge
                -> {
        active -> true,
                -> {
        data
        }
      }
    }
};
simulation
                = {
                -> {
 types
   merge
                -> true,
    coulomb
                -> {
                -> {
      pair
        active -> true,
        data
                -> {
      }
```

```
}
};
```

7.12.10.3 References

- 1. R. D. Groot, "Electrostatic interactions in dissipative particle dynamics simulation of polyelectrolytes and anionic surfactants", J. Chem. Phys. 2003, 118, 11265-11277.
- 2. M. Gonzalez-Melchor, E. Mayoral, M. E. Valsquez, J. Alejandre, "Electrostatic interactions in dissipative particle dynamics using the Ewald sums", J. Chem. Phys. 2006, 125, 224107.

7.12.10.4 Pair

```
bond = {
  active -> boolean,
  nbonded -> integer,
  cutoff -> real
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types

The energetic functional form of the bond contributions to the total potential is described by

$$E_{pair} = \frac{q_i q_j}{4\pi\epsilon_0 \epsilon_r r_{ij}},$$

where q_i and q_j represent the point charges of sites i and j respectively, ϵ_0 represents the dielectric permittivity, ϵ_r the dielectric constant, and r_{ij} the distance between site i and site j. Charges are set per site.

7.12.10.5 Charge

```
charge = {
  active -> boolean,
  nbonded -> integer,
  k -> real,
  cutoff -> real
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types

The distribution of charges in a coarse-grained system cannot be described by simple point charges only. Instead, a spatial distribution following from the Poisson-Boltzmann equation (PBE) should be used, as described by Groot [Groot 2003]. A first approximation of a two-body approach for the solution to the PBE was given by Gonzalez-Melchor et al. [Gonzalez-Melchor 2006] as a function of exponentials. This solution, however, prooved to inadequately describe the two-body solution. We propose s better approximation, which is based on an error function and is given by

$$E_{charge} = \frac{\operatorname{erf}[kr_{ij}]}{r_{ij}},$$

where k represents a force constant and r_{ij} represents the distance between site i and site j. Charges are set per site. The advantage of this function is its easy integration in already existing standard point-charge descriptions, thus allowing for representation of long-range contributions by Ewald summations without change of already existing implementations.

7.12.11 DPD

7.12.11.1 Syntax

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair	struct	Pair interaction descriptors

The Dissipative Particle Dynamics (DPD) force field - developed by Groot and Warren - is a compounded force field, that finds its application in mesoscale simulations. ¹ Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{dpd} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}$$

for bond, angle, torsion, improper, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/dpd/ for applications.

7.12.11.2 Example

```
simulation
  types
                      -> {
                      -> true,
     merge
     dpd -> {
        bond
                      -> {
          active -> true,
                      -> {
          data
                      \rightarrow a, i1 \rightarrow b, k \rightarrow 20, 1 \rightarrow 1}
              {i0
          }
        },
                      -> {
        angle
          active -> true,
          data
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, k \rightarrow 5, theta \rightarrow 180}
          }
        },
        torsion
          active -> true,
```

```
data
              {i0
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
                    \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                    \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
                 }
              }
           }
        },
                       -> {
        pair
           active -> true,
           nbonded -> 3,
           mode
                       -> global,
           cutoff -> 1,
           data
                       -> {
              {i0
                       -> a, i1 -> a, a -> 25.0, gamma -> 4.5, cutoff -> 1},
              {i0
                       -> a, i1 -> b, a -> 26.2, gamma -> 4.5, cutoff -> 1},
              {i0
                      \rightarrow a, i1 \rightarrow c, a \rightarrow 22.1, gamma \rightarrow 4.5, cutoff \rightarrow 1},
              {i0
                      \rightarrow a, i1 \rightarrow d, a \rightarrow 20.0, gamma \rightarrow 4.5, cutoff \rightarrow 1},
              {i0
                      \rightarrow b, i1 \rightarrow b, a \rightarrow 25.0, gamma \rightarrow 4.5, cutoff \rightarrow 1},
                       \rightarrow b, i1 \rightarrow c, a \rightarrow 28.0, gamma \rightarrow 4.5, cutoff \rightarrow 1},
              {i0
              {i0
                      -> b, i1 -> d, a -> 29.0, gamma -> 4.5, cutoff -> 1},
                      \rightarrow c, i1 \rightarrow c, a \rightarrow 25.0, gamma \rightarrow 4.5, cutoff \rightarrow 1},
              {i0
              {i0
                      -> c, i1 -> d, a -> 21.3, gamma -> 4.5, cutoff -> 1},
              {i0
                      -> d, i1 -> d, a -> 25.0, gamma -> 4.5, cutoff -> 1}
        }
     }
  }
};
```

7.12.11.3 References

1. R.D. Groot and P.B. Warren "Dissipative particle dynamics: Bridging the gap between atomistic and mesoscopic simulation", J. Chem. Phys. 1997, 107, 4423-4435.

7.12.11.4 Pair

```
= {
pair
  active
                  -> boolean,
  {\tt mix}
                  -> option,
  nbonded
                  -> integer,
  mode
                  -> option,
  cutoff
                  -> real,
  n
                  -> integer,
  data
  {
    {
      i0
                  -> id,
      i1
                  -> id,
                  -> real,
                  -> real,
      gamma
      {\tt cutoff}
                  -> real
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
mix	option	Sets mixing rule; options are none, berthelot,
		arithmetic, geometric, and sixth.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
mode	option	Sets cut off mode; options are global using the value of
		cutoff for all contributions, individual defining cut offs
		per contribution, and repulsive defining the well loca-
		tion for each separate contribution as the cut off distance.
cutoff	real	Sets global cut off for global mode
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass
		paragraph.
a	real	Force interaction constant.
gamma	real	Force damping constant.
cutoff	real	Site size; the point where the potential equals zero.

The energetic functional form of pair contributions to the total soft potential is described by a repulsive Hertzian spring,

$$E_{pair} = 1/2 \ a \left(1 - r_{ij}/r_{ij,cutoff}\right)^2, \quad r_{ij} < r_{ij,cutoff}$$

where a represents the pairwise interaction paramater, r_{ij} the distance between site i and site j, and $r_{ij,cutoff}$ the cutoff distance between these two sites.

7.12.11.5 Bond

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2 \ k_{bond} (l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.11.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2 \ k_{angle}(\cos(\theta) - \cos(\theta_0))^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.11.7 Torsion

```
torsion
                 -> boolean
  active
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
           delta -> real
        }
      }
    },
};
```

Directive active n	Parameters boolean integer	Description Interaction activator; either true or false. Number of data entries; can be ommitted when type =
data	struct	{merge -> true}. Summary of interaction constants
uata	Struct	· ·
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 6
delta	real	Offset angle, mainly used to change the sign of the cosine
		function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \cos(n_i \phi - \delta_i)),$$

where k_i represents a set of torsion constants, δ_i a torsion offset, n_i the torsion pre-factor, and ϕ the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. The maximum

allowed value of n_i is 6. Note, that the functional form presented here differs from the original OPLS definition.

7.12.11.8 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
                 -> id,
      i1
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.12 GROMACS

7.12.12.1 Syntax

```
gromacs = {
  bond -> struct,
  angle -> struct,
  torsion -> struct,
  improper -> struct,
  pair14 -> struct,
  pair -> struct
};
```

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The gromacs force field is a compounded force field based on the GROMACS force field.
Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{gromacs} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}
```

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/nanoparticle/coarse/ for an application.

7.12.12.2 Example

```
simulation
                   -> {
  types
    merge
                  -> true,
    gromacs
                   -> {
      bond
                  -> {
                  -> true,
         active
                  -> {
         data
                  \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
         }
      },
       angle
                  -> {
         active -> true,
                  -> {
         data
                   -> a, i1 -> b, i2 -> c, k -> 100, theta -> 110}
         }
      },
                  -> {
      torsion
```

```
active -> true,
                     -> {
          data
                    \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
                  \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                  \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
               }
             }
          }
       },
        improper -> {
          active -> true,
          data
                     -> {
             {i0
                     \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
          }
       },
       pair14
                     -> {
          active -> true
       },
       pair
                     -> {
          active -> true,
          nbonded -> 3,
                     -> global,
          mode
          cutoff -> 10,
          mix
                     -> berthelot,
          data
                     -> {
             {i0
                     -> a, i1 -> a, sigma -> 4.0, epsilon -> 0.4},
             {i0
                     \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
             {i0
                     -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
             {i0
                     -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
          }
       }
     }
  }
};
```

7.12.12.3 References

1. W. L. Jorgensen and J. Tirado-Rives, "The OPLS Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.12.4 Pair

```
= {
pair
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
  shift
               -> boolean,
  coulomb
               -> option,
  cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
    {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
    },
    . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.12.5 Bond

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2 \ k_{bond} (l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.12.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2 \ k_{angle} (\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.12.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2, i3	id	Site id, referring to types as defined by the mass paragraph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.12.8 Pair14

```
pair14 = {
  active -> boolean,
};
```

Directive Parameters Description
active boolean Interaction activator; either true or false.

Representation of the 1-4 intra-molecular pair-wise contributions, which have the same functional form as the pair contributions in the following paragraph. The pre-factor, however is 0.5, as described in Jorgensen et al. [Jorgensen 1988].

7.12.12.9 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.13 MARTINI

7.12.13.1 Syntax

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The martini force field is a compounded force field based on the MARTINI force field.¹ Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{martini} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}
```

for bond, angle, torsion, improper, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/martini/ for an application.

7.12.13.2 Example

```
simulation
  types
                      -> {
                      -> true,
     merge
                      -> {
     martini
                      -> {
        bond
           active -> true,
                      -> {
           data
                      \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
           }
        },
                      -> {
        angle
           active -> true,
           data
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, k \rightarrow 100, theta \rightarrow 110}
              {i0
        },
```

```
torsion
                      -> {
           active -> true,
           data
                      -> {
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
              {i0
                   \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                   \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
                }
             }
           }
        },
        improper -> {
           active -> true,
           data
                      -> {
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
              {i0
           }
        },
        pair14
                      -> {
           active -> true
        },
                      -> {
        pair
           active -> true,
           nbonded -> 3,
                      -> global,
           mode
           cutoff -> 10,
           mix
                      -> berthelot,
           data
              {i0
                      \rightarrow a, i1 \rightarrow a, sigma \rightarrow 4.0, epsilon \rightarrow 0.4},
              {i0
                      \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
              {i0
                      -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
              {i0
                      -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
           }
        }
     }
  }
};
```

7.12.13.3 References

1. W. L. Jorgensen and J. Tirado-Rives, "The OPLS Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.13.4 Pair

```
= {
pair
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
  shift
               -> boolean,
  coulomb
               -> option,
  cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
   {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
    },
    . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.13.5 Bond

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2 \ k_{bond} (l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.13.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2 \ k_{angle} (\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.13.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active n	Parameters boolean integer	Description Interaction activator; either true or false. Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine
		function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.13.8 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i 3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.14 OPLS

7.12.14.1 Syntax

```
opls = {
  bond -> struct,
  angle -> struct,
  torsion -> struct,
  improper -> struct,
  pair14 -> struct,
  pair -> struct
};
```

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The OPLS force field is a compounded force field based on the OPLS force field.¹. Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{opls} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}
```

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/opls/ for applications.

7.12.14.2 Example

```
simulation
                   -> {
  types
                   -> true,
    merge
    opls
                   -> {
       bond
                   -> {
                  -> true,
         active
                  -> {
         data
                   \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
         }
       },
                   -> {
       angle
         active -> true,
                   -> {
         data
                   -> a, i1 -> b, i2 -> c, k -> 100, theta -> 110}
         }
       },
                  -> {
       torsion
```

```
active -> true,
                     -> {
          data
                    \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
                  \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                  \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
               }
             }
          }
       },
        improper -> {
          active -> true,
          data
                     -> {
             {i0
                     \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
          }
       },
       pair14
                     -> {
          active -> true
       },
       pair
                     -> {
          active -> true,
          nbonded -> 3,
                     -> global,
          mode
          cutoff -> 10,
          mix
                     -> berthelot,
          data
                     -> {
             {i0
                     -> a, i1 -> a, sigma -> 4.0, epsilon -> 0.4},
             {i0
                     \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
             {i0
                     -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
             {i0
                     -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
          }
       }
     }
  }
};
```

7.12.14.3 References

1. W. L. Jorgensen and J. Tirado-Rives, "The OPLS Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.14.4 Pair

```
= {
pair
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
  shift
               -> boolean,
  coulomb
               -> option,
  cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
    {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
    },
    . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.14.5 Bond

```
bond
  active
                 -> boolean
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = k_{bond}(l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.14.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = k_{angle}(\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.14.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active n	Parameters boolean integer	Description Interaction activator; either true or false. Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine
		function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} 1/2 \ k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.14.8 Pair14

```
pair14 = {
  active -> boolean,
};
```

Directive Parameters Description
active boolean Interaction activator; either true or false.

Representation of the 1-4 intra-molecular pair-wise contributions, which have the same functional form as the pair contributions in the following paragraph. The pre-factor, however is 0.5, as described in Jorgensen et al. [Jorgensen 1988].

7.12.14.9 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.15 SDK

7.12.15.1 Syntax

```
sdk = {
  bond -> struct,
  angle -> struct,
  torsion -> struct,
  improper -> struct,
  pair14 -> struct,
  pair -> struct
};
```

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction descriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The SDK force field is a compounded force field based on the SDK force field.¹. Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{sdk} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}$$

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/sdk/ for applications.

7.12.15.2 Example

```
simulation
                  -> {
  types
    merge
                  -> true,
    sdk -> {
                  -> {
      bond
         active -> true,
                  -> {
         data
                  \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
         }
      },
                  -> {
       angle
         active -> true,
                  -> {
         data
                  -> a, i1 -> b, i2 -> c, k -> 100, theta -> 110}
         }
      },
                  -> {
      torsion
```

```
active -> true,
                     -> {
          data
                    \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
                   \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                   \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
               }
             }
          }
       },
        improper -> {
          active -> true,
          data
                     -> {
             {i0
                     \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
          }
       },
       pair14
                     -> {
          active -> true
       },
       pair
                     -> {
          active -> true,
          nbonded -> 3,
                     -> global,
          mode
          cutoff -> 10,
          mix
                     -> berthelot,
          data
                     -> {
             {i0
                     -> a, i1 -> a, sigma -> 4.0, epsilon -> 0.4},
             {i0
                     \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
             {i0
                     -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
             {i0
                     -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
          }
       }
     }
  }
};
```

7.12.15.3 References

- 1. Shinoda, Devane, Klein papers
- 2. W. L. Jorgensen and J. Tirado-Rives, "The SDK Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.15.4 Pair

```
= {
pair
 active
             -> boolean,
 nbonded
              -> integer,
 mix
               -> option,
 shift
               -> boolean,
 coulomb
              -> option,
 cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
   {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
   },
   . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.15.5 Bond

```
bond
  active
                 -> boolean
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = k_{bond}(l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.15.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = k_{angle}(\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.15.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2, i3	id	Site id, referring to types as defined by the mass paragraph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} 1/2 \ k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.15.8 Pair14

```
pair14 = {
  active -> boolean,
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.

Representation of the 1-4 intra-molecular pair-wise contributions, which have the same functional form as the pair contributions in the following paragraph. The pre-factor, however is 0.5, as described in Jorgensen et al. [Jorgensen 1988].

7.12.15.9 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.16 Spline

7.12.16.1 Syntax

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
pair	struct	Pair interaction descriptors

The spline force field uses a set of splined functions for nonbonded and bonded interactions (see Section 7.9 [Splines], page 196). Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{spline} = E_{pair} + E_{bond} + E_{angle} + E_{torsion}$$

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/spline/ for an application.

7.12.16.2 Bond

```
bond
                 = {
 active
                 -> boolean
                 -> integer,
 data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      spline
                 -> struct
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants

The energetic functional form of the bond contributions to the total potential is described by

```
E_{bond} =,
```

where

7.12.16.3 Angle

```
= {
angle
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
                 -> id,
      i0
                 -> id,
      i1
      i2
                 -> id,
      spline
                 -> struct
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
spline	struct	Spline function structure.

The energetic functional form of the angle contributions to the total potential is described by

```
E_{angle} =,
```

where θ represents the bond angle.

7.12.16.4 Torsion

```
torsion = {
  active -> boolean
  n -> integer,
```

```
data
                ->
  {
    {
      i0
                -> id,
                -> id,
      i1
      i2
                -> id,
      i3
                 -> id,
      spline
                 -> struct
    },
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2, i3	id	Site id, referring to mass paragraph in types
spline	struct	Spline function structure.

The energetic functional form of the torsion contributions to the total potential is described by

```
E_{torsion} =,
```

where ϕ represents the bond torsion.

7.12.16.5 Pair

```
bond
                = {
  active
               -> boolean,
 nbonded
                -> integer,
                -> integer,
  data
                ->
  {
    {
      i0
                -> id,
      i1
                -> id,
      spline
                -> struct
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded in-
		teractions; the given number is expected to be larger or
		equal to zero.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
spline	struct	Spline function structure.

The energetic functional form of the pair contributions to the total potential is formed by a linear interpolation or a cubic spline through the provided data.

7.12.16.6 Example

```
variables
};
simulation
                -> {
  types
                -> true,
    merge
    boltzmann
                -> {
      bond
                -> {
        active -> true,
        data
                -> {
        }
     },
      angle
                -> {
        active -> true,
                -> {
        data
        }
      },
      pair
                -> {
        active -> true,
        nbonded -> 3,
        data
                -> {
      }
    }
  }
};
```

7.12.16.7 References

1. Spyriouni et al., Macromolecules 2007, 40, 3876

7.12.17 Standard

7.12.17.1 Syntax

```
standard = {
  pair -> struct,
  bond -> struct,
  angle -> struct,
  torsion -> struct,
  improper -> struct
};
```

Directive	Parameters	Description
pair	struct	Pair interaction descriptors
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors

The standard force field is a compounded force field, that is OPLS-like in its appearance. Its functional form is a summation of nonbonded and bonded interactions given by

```
E_{standard} = E_{pair} + E_{bond} + E_{angle} + E_{torsion} + E_{improper}
```

for pair, bond, angle, torsion, and improper contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/lj/ for an application. Note, that no standardized typing is provided for the standard force field.

7.12.17.2 Example

```
= {
simulation
  types
                   -> {
                   -> true,
    merge
    standard
                   -> {
      bond
                   -> {
         active
                  -> true,
         data
                  \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
            {i0
         }
      },
                   -> {
       angle
         active -> true,
         data
                   -> a, i1 -> b, i2 -> c, k -> 100, theta -> 110}
         }
      },
       torsion
                   -> {
         active -> true,
```

```
data
                      \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
              {i0
                    \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                    \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
                 }
              }
           }
        },
        improper
                     -> {
           active
                      -> true,
           data
                       -> {
                       \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
              {i0
           }
        },
                       -> {
        pair
           active -> true,
           nbonded -> 3,
           mode
                       -> global,
           cutoff -> 10,
                       -> berthelot,
           mix
           data
                       -> {
              {i0
                       \rightarrow a, i1 \rightarrow a, sigma \rightarrow 4.0, epsilon \rightarrow 0.4},
                      \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
              {i0
              {i0
                       -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
              {i0
                      -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09}
           }
        }
     }
  }
};
```

7.12.17.3 References

1. W. L. Jorgensen and J. Tirado-Rives, "The OPLS Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.17.4 Pair

```
= {
pair
 active
             -> boolean,
 nbonded
              -> integer,
 mix
              -> option,
 shift
               -> boolean,
 coulomb
              -> option,
 cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
   {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
   },
   . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
_		a Lennard-Jones 6-12 potential
sigma	real	Site size: the point where the potential equals zero

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.17.5 Bond

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2 \ k_{bond} (l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.17.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
 n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2 \ k_{angle} (\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.17.7 Torsion

```
torsion
                 -> boolean
  active
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
           delta -> real
        }
      }
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 6
delta	real	Offset angle, mainly used to change the sign of the cosine
		function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \cos(n_i \phi - \delta_i)),$$

where k_i represents a set of torsion constants, δ_i a torsion offset, n_i the torsion pre-factor, and ϕ the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. The maximum

allowed value of n_i is 6. Note, that the functional form presented here differs from the original OPLS definition.

7.12.17.8 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.12.18 Table

7.12.18.1 Syntax

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
pair	struct	Pair interaction descriptors

The tabular force field uses tabulated functions for nonbonded and bonded interactions. Its functional form is a summation of bonded and nonbonded interactions given by

$$E_{table} = E_{pair} + E_{bond} + E_{angle} + E_{torsion}$$

for bond, angle, torsion, and pair contributions respectively. The default interpolation order is set to cubic (see Section 7.9 [Splines], page 196). The following paragraphs describe each contribution in detail. See ./examples/field/table/ for an application.

7.12.18.2 Table Syntax

```
data = {
    {
        x -> real,
        energy -> real
    },
    ...
};
```

Directive	Parameters	Description
х	real	Location; either interparticle distance for pair, bond dis-
		tance for bond, bond angle of angle, or bond torsion for
		torsion interactions.
energy	real	Energy at the specified location.

Table entries are given in pairs, where the x coordinate represents the function variable and the y coordinate the resulting energy. The x interval between each entry must be equidistant. Non-equidistant entries result in poor interpolation.

7.12.18.3 Example

```
simulation
                = {
                -> {
 types
   merge
                -> true,
   boltzmann
                -> {
     bond
               -> {
       active -> true,
       order
               -> cubic,
       data
                -> {
          {i0
               -> a, i1 -> a, name -> "bond_aa.dat"},
          {i0
               -> a, i1 -> b, name -> "bond_ab.dat"},
          {i0
               -> b, i1 -> b, name -> "bond_bb.dat"}
       }
     },
                -> {
     angle
       active -> true,
       order
                -> cubic,
       data
               -> {
               -> a, i1 -> a, i2 -> a, name -> "angle_aaa.dat"},
          {i0
               -> a, i1 -> a, i2 -> b, name -> "angle_aab.dat"},
          {i0
                -> a, i1 -> b, i2 -> b, name -> "angle_abb.dat"},
          {i0
          {i0
               -> b, i1 -> b, i2 -> b, name -> "angle_bbb.dat"}
       }
     },
                -> {
     pair
       active -> true,
       order
               -> cubic,
       nbonded -> 2,
                -> {
       data
          {i0
               -> a, i1 -> a, name -> "pair_aa.dat"},
               -> a, i1 -> b, name -> "pair_ab.dat"},
          {i0
          {i0
               -> b, i1 -> b, name -> "pair_bb.dat"}
     }
   }
 }
};
```

7.12.18.4 Pair

```
= {
bond
  active
                -> boolean,
  order
                -> option,
 nbonded
                -> integer,
                -> integer,
 n
 data
  {
    {
      i0
                -> id,
      i1
                -> id,
                -> string
      name
    },
 }
};
```

Directive active order nbonded	Parameters boolean option integer	Description Interaction activator; either true or false. Spline interpolation order; either linear or cubic. Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data i0, i1 name	struct id string	Summary of interaction constants Site id, referring to mass paragraph in types Table file name; for table format see [Table Syntax] below.

The energetic functional form of the pair contributions to the total potential is formed by a linear interpolation or a cubic spline through the provided data.

7.12.18.5 Bond

```
bond
  active
                -> boolean,
  order
                -> option,
                -> integer,
  data
  {
    {
      i0
                -> id,
                -> id,
      i1
     name
                -> string
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
order	option	Spline interpolation order; either linear or cubic.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to mass paragraph in types
name	string	Table file name; for table format see [Table Syntax]
		below.

The energetic functional form of the bond contributions to the total potential is described by

```
E_{bond} = Interpolation(l),
```

where l represents the bond distance. Either a linear or a cubic spline through the provided data defines the interpolation function.

7.12.18.6 Angle

```
angle
 active
                 -> boolean,
 order
                 -> option,
                 -> integer,
 data
  {
    {
      i0
                 -> id,
                 -> id,
      i1
      i2
                 -> id,
                 -> string
      name
    },
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
order	option	Spline interpolation order; either linear or cubic.
n	integer	Number of data entries; can be ommitted when type =
		<pre>{merge -> true}.</pre>
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to mass paragraph in types
name	string	Table file name; for table format see [Table Syntax]
		below.

The energetic functional form of the angle contributions to the total potential is described by

```
E_{angle} = Interpolation(\theta),
```

where θ represents the bond angle. Either a linear or a cubic spline through the provided data defines the interpolation function.

7.12.18.7 Torsion

```
torsion
  active
                -> boolean,
  order
                -> option,
                -> integer,
  data
  {
    {
      i0
                -> id,
                -> id,
      i1
      i2
                -> id,
                -> id,
      i3
                -> string
      name
    },
 }
};
```

Directive active order n	Parameters boolean option integer	Description Interaction activator; either true or false. Spline interpolation order; either linear or cubic. Number of data entries; can be ommitted when type = {merge -> true}.
data i0, i1, i2, i3	struct id	Summary of interaction constants Site id, referring to mass paragraph in types
name	string	Table file name; for table format see [Table Syntax] below.

The energetic functional form of the torsion contributions to the total potential is described by

```
E_{torsion} = Interpolation(\phi),
```

where ϕ represents the bond torsion. Either a linear or a cubic spline through the provided data defines the interpolation function.

7.12.19 TraPPE

7.12.19.1 Syntax

Directive	Parameters	Description
bond	struct	Bond interaction descriptors
angle	struct	Angle interaction decriptors
torsion	struct	Torsion interaction decriptors
improper	struct	Improper interaction decriptors
pair14	struct	1-4 intra-molecular pair interaction activator
pair	struct	Pair interaction descriptors

The TraPPE force field is a compounded force field based on the TraPPE force field.¹. Its functional form is a summation of bonded and nonbonded interactions given by

```
E_{trappe} = E_{bond} + E_{angle} + E_{torsion} + E_{improper} + E_{pair}
```

for bond, angle, torsion, and pair contributions respectively. The following paragraphs describe each contribution in detail. See ./examples/field/trappe/ for applications.

7.12.19.2 Example

```
simulation
                   -> {
  types
    merge
                  -> true,
    trappe
                   -> {
       bond
                  -> {
         active -> true,
                  -> {
         data
                  \rightarrow a, i1 \rightarrow b, k \rightarrow 100, 1 \rightarrow 2}
         }
       },
       angle
                  -> {
         active -> true,
                  -> {
         data
                   -> a, i1 -> b, i2 -> c, k -> 100, theta -> 110}
            {i0
         }
       },
                  -> {
       torsion
```

```
active -> true,
                     -> {
          data
                    \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, coefficients \rightarrow {
                  \{k \rightarrow 1.5, n \rightarrow 1, delta \rightarrow 0\},\
                  \{k \rightarrow 2.5, n \rightarrow 2, delta \rightarrow 0\}
               }
             }
          }
       },
        improper -> {
          active -> true,
          data
                     -> {
             {i0
                     \rightarrow a, i1 \rightarrow b, i2 \rightarrow c, i3 \rightarrow d, k \rightarrow 100, psi \rightarrow 0}
          }
       },
       pair14
                     -> {
          active -> true
       },
       pair
                     -> {
          active -> true,
          nbonded -> 3,
                     -> global,
          mode
          cutoff -> 10,
          mix
                     -> berthelot,
          data
                     -> {
             {i0
                     -> a, i1 -> a, sigma -> 4.0, epsilon -> 0.4},
             {i0
                     \rightarrow b, i1 \rightarrow b, sigma \rightarrow 3.7, epsilon \rightarrow 0.5},
             {i0
                     -> c, i1 -> c, sigma -> 4.1, epsilon -> 0.22},
             {i0
                     -> d, i1 -> d, sigma -> 2.4, epsilon -> 0.09},
          }
       }
     }
  }
};
```

7.12.19.3 References

- 1. Siepmann papers
- 2. W. L. Jorgensen and J. Tirado-Rives, "The TraPPE Potential Functions for Proteins. Energy Minimizations for Crystals of Cyclic Peptides and Crambin", J. Am. Chem. Soc. 1988, 110, 1657-1666.

7.12.19.4 Pair

```
= {
pair
  active
               -> boolean,
 nbonded
               -> integer,
 mix
               -> option,
  shift
               -> boolean,
  coulomb
               -> option,
  cutoff
               -> real,
 mode
               -> option,
               -> integer,
 n
 data
               ->
  {
   {
     i0
               -> id,
     i1
               -> id,
     epsilon
               -> real,
     sigma
               -> real,
     cutoff
               -> real
    },
    . . .
 }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
nbonded	integer	Number of bonded sites to exclude from nonbonded interactions; the given number is expected to be larger or equal to zero.
mix	option	Sets mixing rule; options are none, berthelot, arithmetic, geometric, and sixth.
shift	boolean	Sets shifting of potential at cutoff; options are true and false.
coulomb	option	Set type of coulomb treatment; options are none, cut, and long.
cutoff	real	Sets global cut off for global mode
mode	option	Sets cut off mode; options are global using the value of cutoff for all contributions, individual defining cut offs per contribution, and repulsive defining the well location for each separate contribution as the cut off distance.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass paragraph.

epsilon	real	Force interaction constant; identical to the well depth for
		a Lennard-Jones 6-12 potential.
sigma	real	Site size; the point where the potential equals zero.
cutoff	real	Cut off; distance at which the pair-wise contribution is
		set to zero; can be omitted.

The energetic functional form of the pair contributions to the total potential is described by a 6-12 Lennard-Jones potential,

$$E_{pair} = 4\epsilon \left(\frac{\sigma}{r_{ij}}\right)^6 \left[\left(\frac{\sigma}{r_{ij}}\right)^6 - 1\right],$$

where r_{ij} represents the distance between site i and site j.

7.12.19.5 Bond

```
bond
                 -> boolean
  active
                 -> integer,
 n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
                 -> real,
      k
      1
                 -> real
    },
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Bond spring constants
1	real	Equilibrium bond length

The energetic functional form of the bond contributions to the total potential is described by

$$E_{bond} = 1/2k_{bond}(l - l_0)^2,$$

where k_{bond} represents a spring constant, l the bond length of bond $\{i_0, i_1\}$, and l_0 the equilibrium bond length.

7.12.19.6 Angle

```
angle
  active
                 -> boolean
                 -> integer,
  n
  data
                 ->
  {
    {
                 -> id,
      i0
      i1
                 -> id,
      i2
                 -> id,
                 -> real,
      k
                 -> real
      theta
    },
  }
};
```

Directive	Parameters	Description
active	boolean	Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type =
		{merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2	id	Site id, referring to types as defined by the mass para-
		graph in types
k	real	Angle spring constants
theta	real	Equilibrium bond angle

The energetic functional form of the angle contributions to the total potential is described by

$$E_{angle} = 1/2k_{angle}(\theta - \theta_0)^2,$$

where k_{angle} represents a spring constant, θ the the angle between bonds $\{i_0, i_1\}$ and $\{i_1, i_2\}$, and θ_0 the equilibrium angle.

7.12.19.7 Torsion

```
torsion
                 -> boolean
  active
                 -> integer,
  n
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      coefficients ->
      {
        {
          k
                 -> real,
                 -> integer,
          delta -> real
        }
      }
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2, i3	id	Site id, referring to types as defined by the mass paragraph in types
k	real	Force constant
n	integer	Torsional angle prefactor; values range between 1 and 4
delta	real	Offset angle, mainly used to change the sign of the cosine function

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{torsion} = \sum_{i=1}^{m} k_i (1 + \operatorname{sign}(n_i) \cos(n_i \phi - \delta_i)),$$

where ϕ represents the bond torsion constructed by bonds $\{i_0, i_1\}$, $\{i_1, i_2\}$, and $\{i_2, i_3\}$. Function $sign(n_i)$ is +1 for odd and -1 for even values of n_i .

7.12.19.8 Improper

```
improper
  active
                 -> boolean
  n
                 -> integer,
  data
  {
    {
      i0
                 -> id,
      i1
                 -> id,
      i2
                 -> id,
      i3
                 -> id,
      k
                 -> real,
                 -> real
      psi
    },
};
```

Directive active	Parameters boolean	Description Interaction activator; either true or false.
n	integer	Number of data entries; can be ommitted when type = {merge -> true}.
data	struct	Summary of interaction constants
i0, i1, i2,	id	Site id, referring to types as defined by the mass para-
i3		graph in types
k	real	Force constant
psi	real	Improper equilibrium angle

The energetic functional form of the torsion contributions to the total potential is described by

$$E_{improper} = k_{improper}(\psi - \psi_0)^2,$$

where ψ represents the improper angle, which is given by the average of torsion angles formed by planes out of permutations of any of the following two bonds: $\{i_1, i_0\}$, $\{i_2, i_0\}$, and $\{i_3, i_0\}$. Note, that site i_0 is the central site, as opposed to some representations, where i_1 is the central site.

7.13 Units

7.13.1 Syntax

```
= {
units
 type
                 -> constant,
 mass
                 -> real,
 length
                 -> real,
 angstrom
                 -> real,
 angle
                 -> real,
  energy
                 -> real,
 kb
                 -> real,
 nav
                 -> real,
                 -> real,
 charge
 permittivity
                -> real,
 seed
                 -> real,
 reduced
                 -> boolean
};
```

Directive	Parameters	Description
type	constant	Defines the type of units used (needed upon first initial-
		ization); options are reduced or si.
mass	real	Defines the global mass scaling constant.
length	real	Defines the global length scaling constant.
angstrom	real	Defines the length scaling for ported formats (e.g. PDB
		or LAMMPS).
angle	real	Defines the global angle scaling constant.
energy	real	Defines the global energetic scaling constant.
kb	real	Defines Boltzmann's constant with respect to the chosen scaling.
nav	real	Defines Avogadro's constant with respect to the chosen scaling.
charge	real	Defines the global scaling of a unit charge.
permittivity	real	Defines the global permittivity scaling.
seed	real	Defines the random seed.
reduced	boolean	Reflects the output state (normally false); options are
		true or false.

7.13.2 Usage

This variable style describes units. Units are used to internally scale all variables with respect to their units. The units can also be used to define an appropriate relation between force field energies, angles, bead sizes, etc. Upon initialization, a suitable scaling will be deduced from force field or internal natural constant definitions when the contributor is zero (see Section 7.12 [Types], page 201).

7.13.3 Default

```
Unless otherwise stated, the default is given by
```

By default, all other contributors are set to zero and deduced upon initialization.

7.14 Vector

7.14.1 Syntax

vector = $\{x \rightarrow real, y \rightarrow real, z \rightarrow real\};$

Directive Parameters Description

x, y, z real Sets vector components.

7.14.2 Usage

This variable style describes a vector. Vectors always consist of three components, which can be given in real numbers. Interpretation can either be real or integer, dependent on the command in which vector occurs.

7.14.3 Default

Unless otherwise stated, the default is given by

vector = $\{x \rightarrow 0, y \rightarrow 0, z \rightarrow 0\};$

7.15 Voigt

7.15.1 Syntax

Directive Parameters Description
xx, yy, zz, real Sets voigt components.
zy, zx, yx

7.15.2 Usage

This variable style describes a voigt. Voigts always consist of six components, which can be given in real numbers. Interpretation can either be real or integer, dependent on the command in which voigt occurs.

7.15.3 Default

Unless otherwise stated, the default is given by

voigt =
$$\{xx \rightarrow 0, yy \rightarrow 0, zz \rightarrow 0, zy \rightarrow 0, zx \rightarrow 0, yx \rightarrow 0\};$$

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