MOLSIM

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Contents

1	Mols	im 1
	1.1	Introduction
	1.2	Installation
	1.3	Getting Started
2	Inpu	
	2.1	nmlSystem
	2.2	nmlScale
	2.3	nmlThermoInteg
	2.4	nmlDist
	2.5	nmlParticle
	2.6	nmlCopolymerSequence
	2.7	nmlNetworkConfiguration
	2.8	nmlRepeating
	2.9	nmlPotential
	2.10	nmlPotentialChain
		nmlPotentialExternal
		nmlPolarizationIter
		nmlSetConfiguration
		nmIMD
		nmIMC
		nmlMCAII
		nmlMCWeight
		nmlUmbrella
		nmlMCPmf
		nmlSPartSSO
		nmlBD
		nmlIntList
		nmlDump
		nmlGroup
		nmlStatic
		nmISPDF
		nmIRDF
		nmlRDFChain
		nmlRDFSph
		nmlRDFCond
		nmlG3Dist
		nmlSF
		nmlScatIntens
		nmlAngDF
		nmlAngExtDF
		nmlOriDipDF
	131	DIDIDADADODE 91

Contents Contents

	2.38	nmlKirkwoodgk	92
		nmlOriPolDF	93
		nmINNHB	94
		nmlNNDF	95
		nmlChainDF	96
		nmlChainTypeDF	96
		nmlChainTypeExtDF	96
		nmlCBPC	97
		nmlLoopTailTrain	-
		nmlCluster	
		nmlMultipoleDF	
		nmlEnergyDF	
		nmlWidom1	
		nmlWidom2	
		nmlMeanForce1	
		nmlMeanForce2	
		nmlPotMeanForce	
		nmlSurfaceArea	
		nmlTrajectory	
		nmlSubStructureDF	
		nmlNetworkDF	
		nmlNetworkRadialDF	
		nmlDynamic	
		nmlMSD	
		nmlOriXTCF, nmlOriYTCF and nmlOriZTCF	
		nmlLinVelTCF and nmlAngVelTCF	
		nmlForTCF and nmlTorTCF	
		nmllDMTCF	
		nmlUtotTCF	
		nmllmage	
		nmIVRML	
		nmIVTF	
		nmlMixed	
		nmlMixed1	
		nmlMixed2	
		nmlMixed3	
		nmlMixed4	
		nmlMixed5	
		nmlMixed6	
		nmlComplexation	
		nmlComplexDist	
	2.70	Timoompickbist	101
3	User	-provided routines	132
	3.1	PotentialUser	132
	3.2	SetParticleUser	132
	3.3	DumpUser	
	3.4	GroupUser	
	3.5	StaticUser	
	3.6	ImageUser	
4	Distr	ibuted files	134

Contents

5	Subr	outines and Functions	135
	5.1	mol.F90	136
	5.2	molsim.90	136
	5.3	particle.F90	136
	5.4	potential.F90	137
	5.5	coordinate.F90	138
	5.6	md.F90	139
	5.7	mc.F90	140
	5.8	bd.F90	142
	5.9	nlist.F90	142
	5.10	energy.F90	143
		denergy.F90	
		dump.F90	
		group.F90	
		static.F90	
		dynamic.F90	
		image.F90	
		mixed.F90	
		molaux.F90	
		mesh.F90	
		statistics.F90	
		mollib.F90	
		parallel.F90	
		sso.F90	
		gc.f90	
	5.26	celllist.F90	106
6	Exte	rnal units	159
7	Data	structures	161
•	Data	Structures	
8	Varia	bles	169
9	Addi	tional Descriptions	170
	9.1	Networks	170
	9.2	Complexation-Analysis	
	9.3	Advanced Configurations	
	9.4	Copolymer Sequence	
	9.5	Celllist	
	9.6	SSO	
	9.7	Image generation with VMD	
	0.7	mage generation with vivib	
10	Deve	eloping ·	184
		Testin	184
		Random Numbers	
		Compilation Modes	
	. 0.0		. 55
11	Appe	endix ·	188
			188
	11.2	Examples of namelists describing different objects	189
		Examples of namelists describing different objects	

1 Molsim

1.1 Introduction

MOLSIM is a modular MOLecular SIMulation software for simulation of all-atom and coarse-grained model systems with extensive static and dynamic analyses. It can be run in two different **modes**: generating configurations/trajectory and reading configurations/trajectory. The latter one is useful for subsequent analyses.

MOLSIM supports molecular dynamics (MD), Monte Carlo (MC), and Brownian dynamics (BD) simulation **methods**. Simulations can be performed the microcanonical (NVE), canonical (NVT), isobaric (NPT), and grand canonical (μ VT) **ensembles** with a number of different boundary **conditions** applied. Initial **configurations** can be generated by random, on lattice, or using user- provided routines, as well as be read from text files or from binary files of previous simulations.

More specifically, several MD integrators are provided and quaternion description is used for rigid-body rotation. Regarding MC, a large selection of different types of trial displacements is available and cluster moves are supported being useful for system with strongly interacting particles. The total number of **time steps**, in case of MD or BD, or **passes** (one attempt to move each particle), in case of MC, are divided into blocks called **macrosteps**. Complete calculations of averages are done and written for each macrostep, and grand averages are given after the simulation. For some quantities fluctuations are also given. The **precision** of grand averages (i) of scalar variables are calculated by extrapolation of block averages to infinite block length and (ii) of distribution functions from the spread of macrostep averages. Simulations can easily be restarted after an interruption or **prolonged**. Binary storage are used for saved positions and velocities, saved intermediate data, and saved data for subequent analyses for maximal accuary.

MOLSIM is able to handle four types of **objects**. They are **atom**, **particle**, **chain**, and **hierarchical structure**, and they are related to each other as follows: a hierarchical structure is built up of one or several chains, a chain of one or several particles and a particle of one or several atoms. Interactions originate from atoms and particles constitute rigid entities. A particle can, e.g., represent a molecule, a monomer in a polymer joined by a bonding potential, or a colloidal particle; a chain a linear polymer; and a hierarchical structure a dendrimer. Identical **objects** belong to the same class of **object type**. Order numbers are assigned to objects and object types as well as to **pairs of objects** and to **pairs of object types**. Whenever objects or object types, say A, B, C, appear ordered as (A, B, C), the order of the corresponding pairs becomes (AA, AB, AC, BB, BC, CC). This ordering of object and object pairs is important for the assignment of values to input variables.

The **nonbonded** pair potentials, operating between pairs of atoms, can be specified either as (i) a sum of A*/r**n terms with different prefactors and exponents, or by special routines adapted for more complicated potentials, including the possibility of user-provided potential routines. The two-body interactions are calculated through a **neighbor list** technique, and the potentials and forces are evaluated through **lookup tables**. Different routines are used whether (i) charge, (ii) charge and dipole, or (ii) charge, dipole, and induced dipole interactions are present. In the last case, atom charges, atom dipole moments, and anisotropic polarizability tensors are explicitly used in an efficient prediction method to handle the **many-body polarization**. The long-range charge and dipolar interactions are handled by **Ewald** summation, including the Smooth Particle Mesh method, or by a **reaction-field** method.

CHAPTER 1. MOLSIM 1.2. INSTALLATION

The **static analysis** comprises energy distribution functions (df), radial and angular distribution functions, and structure factors. Moreover a large selection of df's describing polarization of dipolar fluids and, the structure of water are available. Also a range of df's charactering polymer shape, structure as well as cluster formation analyses are implemented. Finally, excess chemical potential and potential of mean forces can be evaluated. Implemented **dynamic analyses** are mean square displacements, time correlation functions of velocities, orientations, angular velocities, forces, torques, induced dipole moments, and potential energy. It is possible to restrict the static and dynamic analyses to groups of molecules specified by the user.

It is easy to add **user-provided routines** to MOLSIM to enhance the capability of it. They comprise additional (i) nonbonded potentials, (ii) preparations of start configuration, (iii) selections of data to be dumped, (iv) group divisions, (v) static analyses, and (iv) protocols of exporting coordinates and other information for later image rendering. The user-provided routines are collected in a special file and do not belong to the supported software. Existing input variables control the calls of these routines.

The specification of **input variables** characterizing the model and controlling the calculation is given in an input file. That makes it easy to make sequential calculations with only minor differences in the specification. Both names and values of the variables are given in input file using the Namelist protocol, making it easy to read and prepare input files. Blocks of input data are collected into separate groups.

Text books on molecular simulation:

- Allen, M. P.; Tildelsley, D. J. 'Computer Simulation of Liquids'; Clarendon Press: Oxford, 1987 (1989 in paperback).
- Frenkel D. and Smit, B. 'Understanding Molecular Simulation: From Algorithms to Applications'; Academic Press: San Diego, 1996.
- Rapaport, D. C. 'The Art of Molecular Dynamics Simulation'; Cambridge University Press: Cambridge, 1995.
- Field, M. J. 'A practical introduction to the simulation of molecular systems'; Cambridge University Press: Cambridge, 1999.

1.2 Installation

Obtaining the Code

There are two possibilities of how to obtain the code. You can either simply download the zip of the code, or clone the whole repository.

Downloading the Zip

Download the zip from here and save it in the directory of your choice. Afterwards, navigate to that directory and extract the source code with

unzip <name of the zip file>

You might want to rename the directory to some more reasonable name.

CHAPTER 1. MOLSIM 1.2. INSTALLATION

Clone the Molsim repository

```
Just run

git clone https://github.com/joakimstenhammar/molsim.git

If you set up your ssh key at GitHub you can run

git clone git@github.com:joakimstenhammar/molsim.git
```

Installation of Molsim

Navigate into the Molsim directroy and run the configure script. This will check some dependencies. Molsim requires FFTW 3.3.4. In can be install automatically within the configure script (Note: This might take some time). The configure script will also ask you for a version name. This version name will be appended to the executables of molsim (use molsim_ser.ver instead of molsim_ser). Leave it blank for no special version name.

```
cd Molsim
./configure.sh
```

The configure script tries to locate the FFTW libary. If you want to customize the path of the libary, modify the Src/make.fftwpath file. Additionally it will select which compiler to use. To costumize the compiler which is to be used change the Src/make.arch file.

Now go to the Src directory and make Molsim:

```
cd Molsim/Src
```

and compile the code with

make all

Alternatively you can compile only the serial code with make ser, only the parallel code with make par. You might be able to speed up the compilation, by letting the compilation run in parallel (add the option -j <number of cores> to the make call e.g. make -j 4 all).

To install Molsim run

make install

If you want to uninstall Molsim run

make uninstall

After completion, (i) additional files have been created in the source directory, (ii) a subdirectory Bin has been created and contains the executables molsim_ser.exe and molsim_par.exe, and (iii) the files molsim_ser and molsim_par have been copied from the source directory to \$HOME/bin.

Note, the environmental variable PATH has to contain \$HOME/bin for the software to function (usually this is handled by the configure script)

The installation of the software is now completed.

You can check the version of molsim by passing either the -v, -V or --version to molsim, like

molsim_ser --version

1.3 Getting Started

This chapter describes how to execute the software.

- Create a working directory where input and result files will be residing

This directory will in the following be referred to as the work directory. The location of the work directory is arbitrary

Copy the input file test.in to the work directory

This input file could, e.g., be retrieved from www.physchem.lu.se/sm. At this moment we will not consider the content and the meaning of the variables in the input file. The input variables are the subject of the next chapter.

- Type 'molsim ser test'

After the execution, the files test.out and test.list (among others) should now exist in the working directory.

The content of the output file test.out , should be essentially the same as that obtained by execution trough www.physchem.lu.se/sm. If not, probably some error has occurred in the transfer of the code or during the installation.

The test calculations are normally short. The results of them are normally not applicable for the equilibrium situation, but are nevertheless useful for testing the installation.

The parallel version of MOLSIM is invoked by typing 'molsim par' instead of 'molsim ser'.

If during the installation you gave a version name while running the configure script, the version name will be appended to the executables of molsim (molsim_ser.ver instead of molsim_ser).

2 Input

All input data read from the file FIN are done with the namelist procedure. Each namelist contains input related variables. Further information on the namelist procedure is given in Appendix A.

The following namelists are available. Compulsory nameslist are checked with X.

file	namelist		variables describing
molsim.F90	nmlSystem	Х	general system variables
	nmlScale	X	scaling units
	nmlThermoInteg		thermodynamic integration
	nmlDist		distribution functions
particle.F90	nmlParticle	X	particle variables
	nmlCopolymerSequence		describe the sequence of copolymers
	nmlNetworkConfiguration		describe network configuration
	nmlRepeating		define the repeating block structure of
natantial FOO	numl Determine		copolymers
potential.F90	nmlPotential	X	potentials and forces
	nmlPotentialChain		bond and angle potential
	nmlPotentialExternal		external potential
	nmlPolarizationIter		many-body polarization calculation
coordinate.F90	nmlSetConfiguration	Х	initial start configuration
md.F90	nmlMD		molecular dynamics simulation
mc.F90	nmlMC		Monte Carlo simulation
	nmlMCAll		Monte Carlo (all) simulation
	nmlMCWeight		applying weights for pmf calculations
	nmlUmbrella		umbrella potential sampling
	nmlMCPmf		direct pmf calculation using updated weights
sso.F90	nmlSPartSSO		optimization of the single particle move
bd.F90	nmlBD		Brownian dynamics simulation
nlist.F90	nmlIntList		calculation of list of nonbonded pairs
dump.F90	nmlDump		dumping/reading of data
group.F90	nmlGroup		dividing particles into groups
static.F90	nmlStatic		call of static analysis routines
	nmlSPDF		single particle distribution functions
	nmIRDF		radial distribution functions
	nmIRDFChain		radial distribution functions for chains
	nmlRDFSph		projected radial distribution functions
	nmlRDFCond		conditional radial distribution functions
	nmlG3Dist		normalized triplet correlation functions
	nmlSF		structure factors
	nmlScatIntens		scattering intensities
	l .		I .

file	namelist	variables describing
	nmlAngDF	angular distribution functions
	nmlAngExtDF	angular distribution functions wrt ex-
		ternal frame
	nmlOriDipDF	orientation/dipole distribution functions
	nmlRadAngDF	radial-angular 2d distribution functions
	nmlKirkwoodgk	Kirkwood's Gk factor
	nmlOriPolDF	orientaitional polarization distribution functions
	nmINNHB	neighbors and hydrogen bonds
	nmINNDF	nearest neighbor distribution functions
	nmlChainDF	chain distribution functions
	nmlChainTypeDF	chain type distribution functions
	nmlChainTypeExtDF	chain type distribution functions, exter- nal frame
	nmlCBPC	probability of bead-particle contact
	nmlLoopTailTrain	loop, tail, and train characteristics
	nmlCluster	cluster size distribution functions
	nmlMultipoleDF	electrostatic multipole moment distri-
		bution functions
	nmlEnergyDF	energy distribution functions
	nmlWidom1	chemical potentials using Widom's method
	nmlWidom2	chemical potentials using Widom's method
	nmlMeanForce1	mean force between two particles
	nmlMeanForce2	mean force between two particles
	nmlPotMeanForce	potential of mean force between two particles
	nmlSurfaceArea	surface area exposed by all particles of one type
	nmlTrajectory	write trajectory on unit FLIST
	nmlSubStructureDF	distribution function of substructures
	nmlNetworkDF	network distribution functions
	nmlNetworkRadialDF	radial network distribution functions
dynamic.F90	nmlDynamic	call of static analysis routines
	nmlMSD	mean square displacement
	nmlOriXTCF, nmlOriYTCF and nmlOriZTCF	orientational tcf of particle x'/y'/z'-axis
	nmlLinVelTCF and nmlAngVelTCF	linear / angular velocity tcf
	nmlForTCF and nmlTorTCF	force and torque tcf
	nmlIDMTCF	induced dipole moment tcf
	nmlUtotTCF	energy tcf
image.F90	nmllmage	call of image data writing routines
	nmIVRML	generation of VRML coordinate files
	nmIVTF	generation of VTF coordinate files
mixed.F90	nmlMixed	general Molmix variables
	nmlMixed1	generation of potential energy curves
	TITTIIVIIACUT	generation of potential energy ourves

file	namelist	variables describing
	nmlMixed2	calculation of potential energies on a
		lattice
	nmlMixed3	calculation of global potential energy
		minimum
	nmlMixed4	generation of random coordinates
	nmlMixed5	calculation of second virial coefficients
	nmlMixed6	calculation of orientational averaged
		potential energy
moluser.F90	nmlComplexation	analysis of interparticle complexation
	nmlComplexDist	calculation of complexation distribution
		functions

The following subsections contain all input variables that are used for reading data from file FIN. The variables are grouped together and listed below their namelist name. The first line contains the name, the type, the dimension, and, if any, the default value of the variable. The following lines briefly explain the use of the variable. If the variable only can attain a limited number of values, these are listed. If description about writing data is given, the output is made on file FOUT (if nothing else is stated).

A practical point. Often there is a need of reading a variable conditionally, i.e., if some condition is true the variable is assigned a value and used later. If the condition is false, the presence of the variable in the input file does no harm; thus it has not to be deleted. In the latter case the input file contains redundant data.

2.1 nmlSystem

The namelist nmlSystem contains variables that describe the main features of the simulation and flags that controls optionally calls of routines as well as output

- Variables
 - txtitle
 - txmethod
 - txmode
 - txensemb
 - txbc
 - txstart
 - txuser
 - boxlen
 - cellside
 - sphrad
 - ellrad
 - cylrad
 - cyllen

- lenscl
- temp
- prsr
- nstep1
- nstep2
- iseed
- luseXYseed
- ixseed
- iyseed
- maxcpu
- Icont
- laver
- Iti
- Idist
- Idump
- Igroup
- Istatic
- Idynamic
- limage
- Itime
- itest
- ipart
- iatom
- iaver
- ishow
- iplot
- ilist
- Itrace
- Iblockaver

txtitle

character(90)

• User-provided title.

txmethod

character(5)

• 'md': Molecular dynamic simulation. Further specification is given in namelist nmlMD.

- 'mc': Monte Carlo simulation. Further specification is given in namelist nmlMC.
- 'mcall': Monte Carlo with simultaneous movement of all particles. Further specification is given in namelist nmlMCAll.
- 'bd': Brownian dynamic simulation (configuration space). Further specificati on is given in namelist nmlBD.

txmode

character(10) default: 'simulation'

- 'simulation': Simulation and analyses. Further specification of method, ensemble, boundary conditions, and initial configuration are specified by variables txmethod, txensemb, txbc, and txstart. On a top level, analyses are controlled by Igroup, Istatic, and Idynamic.
- 'analysis': Reading of dump data from DUMP files and analyses. On a top level, analyses are controlled by Igroup, Istatic, and Idynamic.
- 'mixed': Mixed activities controlled by namelist nmlMixed.

txensemb

character(3)

- nve: Microcanonical ensemble (only MD). This option should also be used for MD with temperature and/or volume scaling. Further specification is given in namelist nmlMD.
- 'ntv': Canonical ensemble (only MC or BD).
- 'nvt': Canonical ensemble (only MC or BD).
- 'npt': Isothermal-isobaric ensemble (only MC).
- 'ntp': Isothermal-isobaric ensemble (only MC).
- 'mvt': Grand canonical ensemble (only MC). Still not fully tested.
- 'mtv': Grand canonical ensemble (only MC). Still not fully tested.

txbc

character(3)

- 'xyz': Periodical boundary condition, x, y, and z-directions.
- 'xy': Periodical boundary condition, x and y-directions.
- 'z': Periodical boundary condition, z-direction.
- · 'rd': Periodical boundary condition, rhombic dodecahadral.
- 'to': Periodical boundary condition, truncated octahedral.
- 'sph': Hard sphere boundary (only MC).
- 'cyl': Hard cylinder boundary (only MC).
- 'ell': Hard ellipsoidal boundary (only MC).

txstart

character(8)

- 'setconf': Generation of a start configuration and accumulation variables are set to zero. This option is used to obtain a start configuration that should be equilibrated. Further specification is given in namelist nmlSetConfiguration.
- 'readfin': Read start configuration from file FIN and accumulation variables are set to zero. The format is(ro(1:3,ip),ori(1:3,1:3,ip),ip = 1,np), i.e., the same as the output of particle coordinates on FOUT.
- 'zero': Start configuration is read from FCNF and accumulation variables are set to zero. This option is used to start of a production run composed of nstep1*nstep2 steps/passes.
- 'continue': Start configuration and accumulated averages are read from FCNF. This option is used to continue an equilibration or production run if the execution was aborted due to exceeded time limit, system malfunction etc. To continue, change start to 'continue' and resubmit the job. This option may also be used to extend a completed production run. Then increase nstep1 to the new total number of macrosteps, change start to 'continue', and resubmit the job. The simulation will continue from the last run to a new total nstep1* nstep2 steps/passes.

txuser

character(80)

• Character string for engaging user-provided code in the MOLSIM core.

boxlen

real(1:3)

Box length in x-, y-, and z-directions.

cellside

real

• Side length of a rhobic dodecaheron (only txbc='rd') and truncated octahedron cell (only txbc='to').

sphrad

real

• Spherical cell radius (only txbc='sph').

ellrad

real(1:3)

• Ellipsoidal cell radii (only txbc='ell').

cylrad

real

• Cylindrical cell radius (only txbc='cyl').

cyllen

real

• Cylindrical cell length (only txbc='cyl').

lenscl

real default: 1.0

• Scaling constant which scales box, and particle coordinates (only txstart='zero'). Useful for creating a system similar to a previous one, but with different box lengths.

temp

real

• System temperature (only txensemb='npt' or 'nvt'). Desired temperature (only txensemb='nve' ensemble and temperature scaling). Temp is also used if the velocities are set according to a Maxwell distribution.

prsr

real

• Desired pressure (only txensemb='npt' or 'nve' and with volume scaling).

nstep1

integer

• Number of macrosteps. The total number of steps/passes are nstep1*nstep2. After the simulation grand averages are calculated and written for the nstep1*nstep2 steps/passes.

nstep2

integer

Number of steps (only MD or BD), or number of passes (only MC or MCALL) per macrostep.
 One pass is one attempt to move each particle in average (only MC) or one attempt to simultaneously move all particles (only MCALL). Averages are calculated and written for each set of nstep2 steps/passes. After each set the FCNF file is updated with new coordinates and accumulated averages.

iseed

integer

• Seed of random number generator. iseed > 0 is required.

luseXYseed

logical default: false

• .true.: ixseed and iyseed from the input file are used.

• .false.: Nothing.

ixseed

integer

· Initial ix used for the random number generator.

iyseed

integer

· Initial iy used for the random number generator.

maxcpu

integer default: 0

- · 0: Nothing.
- <0: The program stops before starting next macrostep if the total cpu time including the next one exceeds maxcpu (in seconds). The cpu time for the last macrostep is used as the estimate of the cpu time for the next macrostep. This is handy if batch queue installation is used. In such cases, if the job exceeds the maximum cpu time for the batch job, the job might stop abruptly (depending on system installation) without executing the remaining commands in the flink command file and possibly corrupting the file FCNF. If so, the entirely job is lost. The variable maxcpu allows the program to stop itself and promptly send the FCNF file which then can be used to continue the simulation by using the option txstart='continue'.</p>

Icont

logical default: .false.

- Quantities which use is primary to check that the simulation is properly advancing may be calculated (by driver ControlAver). The quantities are averaged over particle types and they are average square force and torque (per particle), linear and angular moments (per particle), as well as translational and rotational temperatures (only MD). Fraction accepted and rejected attempts to move (only MC). Mean square displacement per step/pass. Orientational order. Defined as the scalar product of the direction of a molecular axis and its direction at start. The value is one for a perfect alignment and approximately zero for a fluid. Useful for monitoring the relaxation of an initial lattice start. Position and orientational means. The position of the center of mass denoted $\langle r0 \rangle$, and the projection of the molecular axes on the box axes denoted $\langle x' \rangle$, $\langle y' \rangle$, and $\langle z' \rangle$.
- · . true .: Control quantities are calculated.
- false.: No calculation of control quantities.

laver

logical default: .false.

- Thermodynamic averages and their precision as well as fluctuations and their precision may be calculated (by routine MainAver). Consider the quantity Q. The precision its average $\langle Q \rangle$ and fluctuation $\sqrt{\langle Q^2 \rangle \langle Q \rangle^2}$, both given as one standard deviation are evaluated by block averaging with variable block length and extrapolation to infinite block length. The quantities considered are:
- 1. Total energy (MD)
- 2. Kinetic energy (MD)
- 3. Potential energy, total
- 4. Potential energy, total two-body contribution (only txelec='dip' or 'pol' the electrostatic interaction is excluded) Potential energy, partial two-body contributions (only txelec='dip' or 'pol' the electrostatic interaction is excluded)
- 5. Potential energy from the reciprocal space (only txelec='charge' .and. lewald)
- 6. Electrostatic energy (only txelec='dip' or 'pol'), including the reciprocal space (only lewald)

- 7. Polarization energy (only txelec='pol'), including the reciprocal space (only lewald)
- 8. Enthalpy
- 9. Heat capacity
- 10. Temperature
- 11. Pressure
- 12. Volume
- 13. Induced dipole moment: total (only txelec='pol')
- 14. Induced dipole moment: particle (only txelec='pol')
- 15. Induced dipole moment: atom (only txelec='pol')
 - . true .: Thermodynamic averages are calculated.
 - .false.: No calculation of thermodynamic averages.

lti

logical default: .false.

- . true.: Handle thermodynamic integration (by routine ThermoInteg). Further specification is given in namelist nmlThermoInteg.
- .false.: No thermodynamic integration.

Idist

logical default: .false.

- . true. Distribution functions are calculated (by routine DistFunc). Further specification is given in namelist nmlDist.
- .false. No calculation of distribution functions.

Idump

logical default: .false.

- .true. Variables are dumped or read. Further specification is given in namelist nmlDump.
- · .false. No dumping/readiing

Igroup

logical default: .false.

- .true. Particles are divided into groups. This division is used in routine static and optionally by moldyn. Further specification is given in namelist nmlGroup.
- .false. No division into groups

Istatic

logical **default:** .false.

• . true. Makes it possible to call several routines calculating static properties. Further specification is given in namelist nmlStatic (require lgroup=.true.).

• .false. No static analysis.

Idynamic

logical default: .false.

- . true. Makes it possible to call several routines calculating dynamic properties. Further specification is given in namelist nmlDynamic (require lgroup=.true.).
- · .false. No dynamic analysis.

limage

logical default: .false.

- . true. Makes it possible to call several routines preparing files for generating images. Further specification is given in namelist nmllmage.
- · .false. No image analysis.

Itime

logical default: .true.

- · . true .: Timing statistics are carried out.
- .false.: No timing statistics.

itest

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- '1': Energy, pressure, particle and atom coordinates, forces, torques, dipole moments, polarizability tensor, linear and angular accelerations, linear and angular velocities, and particle distance matrix are written after each step/configuration.
- 3: Intermediate energies (energy.F90).
- 4: Neighbour list (nlist.F90).

ipart

integer default: 0

- 0: Nothing.
- >0: Initial and final particle coordinates and velocities are written for every ipart particle.

iatom

integer default: 0

- 0: Nothing.
- >0: Initial atomic coordinates are written for every iatom atom.

iaver

integer **default:** 0

- 0: Nothing.
- >0: Cumulative (for the macrostep) thermodynamic averages are written for every iaver steps/passes.

ishow

integer default: 0

- 0: Nothing.
- >0: Calculated functions are listed at every ishow bin. Compact list.

iplot

integer **default:** 0

- 0: Nothing.
- >0: Calculated functions are plotted.

ilist

integer default: 0

- 0: Nothing.
- >0: Calculated functions are listed at every ilist bin on file FLIST. Extended list.

CHAPTER 2. INPUT 2.2. NMLSCALE

Itrace

logical **default:** .false.

• . true.: Information upon entering and exiting subroutines on three levels are written on unit 40 for the master and unit 41 for slaves.

· .false.: Nothing.

Iblockaver

logical default: .false.

• .true.: Intermediate data concerning the block averaging and its extrapolation to infinite block length are written on file blockaver.data.

· .false: Nothing.

2.2 nmlScale

The namelist nmlScale contains scaling variables that describe the relation between SI units and units used. The relation between the numerical value of quantity Q in SI units and units used is Q(SI) = S*Q, where S is the scaling factor. Below, the SI units are given in parentheses after the default scaling factors. The output list gives the scaling factor for some other derived units presented in the output. By a suitable choice of assignments, input variables could be interpreted as reduced variable. In particular, sclene = RT (ca. 2500 at ambient temperature), where R is the gas constant and T the provided temperature, energies are given units of kT.

- · Variables:
 - scllen
 - sclmas
 - scltem
 - scltim
 - sclene
 - sclhca
 - sclpre
 - scldif
 - sclang

scllen

real default: 1.0d-10 m

· Scaling factor for length.

CHAPTER 2. INPUT 2.2. NMLSCALE

sclmas

real default: 1.0d-3 kg/mol

· Scaling factor for mass.

scltem

real default: 1.0 K

· Scaling factor for temperature

scltim

real default: 1.0d-12 s

• Scaling factor for time.

sclene

real default: 1.0d+3J/mol

· Scaling factor for energy.

sclhca

real **default:** 1.0 J/(mol*K)

· Scaling factor for heat capacity.

sclpre

real default: 1.0d+6 Pa

· Scaling factor for pressure.

scldif

real **default:** 1.0d-9 m^2/s

• Scaling factor for diffusion coefficient.

sclang

real default: PI/180 1/rad

• Scaling factor for angle.

2.3 nmlThermoInteg

The namelist nmlThermoInteg contains variables that control thermodynamic integration. Presently, (i) hard-sphere, (ii) Coulomb, (iii) bond, and/or (iv) angular potentials can be changed through the coupling parameter.

- Variables:
 - lambda
 - powercharge
 - powerkbond
 - powerkangle

lambda

real

· Coupling parameter.

powercharge

real default: 1.0

· Power of the coupling parameter scaling the charge.

powerkbond

real default: 9.0

· Power of the coupling parameter scaling kbond.

powerkangle

real default: 6.0

Power of the coupling parameter scaling kangle.

CHAPTER 2. INPUT 2.4. NMLDIST

type	label	distribution functions	
------	-------	------------------------	--

2.4 nmlDist

The namelist nmlDist contains variables that control the calculation of distribution functions during the simulation. Any combination of the nine types of distribution functions listed below may be selected through vtype%l.

type	label	distribution functions
1	totu	total potential energy
2	paru	partial potential energy (only two-body contributions)
3	bindu	binding energy (only two-body contributions)
4	pairu	pair energy (only two-body contributions)
5	rdf	radial d.f., particle-particle (center of mass)
6	rdf	radial d.f., atom(mass > maslim)-atom(mass > maslim)
7	idm	induced dipole moment: total
8	idm	induced dipole moment: particle
9	idm	induced dipole moment: atom (polarization > pollim)
10	zdens	z-density distribution function: particle

The uncertainty of a given value of the distribution functions is given as one standard deviation, and it is evaluated by block averaging where one macrostep constitutes one block.

- · Variables:
 - vtype
 - idist
 - rcutdist
 - maslim
 - pollim
 - itestdist

vtype

static1D_var(logical, real, real, integer, logical, character, real)(1:10)

• Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.

idist

integer default: 10

Interval of sampling the distribution functions.

rcutdist

real default: rcut+racom(ipt)+racom(jpt)

• Cutoff distance for evaluating the distribution functions based on the center-ofmass of the particle. The value could be smaller but should not exceed the default one.

maslim

real default: 1d-4

· Lower mass limit for calculating atom-atom rdf.

pollim

real default: 1d-4

· Lower polarizability limit for calculating induced dipole moment distribution functions for atoms.

itestdist

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- 1: Forces and virial (routine DistFunc).

2.5 nmlParticle

The namelist nmlParticle contains variables that describe the number of particles, their geometries, masses, etc. Examples of namelist nmlParticle describing different types of particles are given in Appendix B.

- · Variables:
 - txelec
 - Iclink
 - Imultigraft
 - Ihierarchical
 - maxnbondcl
 - Ireadbondcl
 - ngen
 - ictgen
 - nbranch

- ibranchpbeg
- ibranchpinc
- nnwt
- nct
- txct
- ncct
- npptct
- txcopolymer
- nblockict
- npt
- txpt
- nppt
- natpt
- txat
- massat
- radat
- zat
- zatalpha
- sigat
- epsat
- latweakcharge
- jatweakcharge
- pK
- pH
- naatpt
- txaat
- rain
- dipain
- polain
- lintsite
- raintin
- Iradatbox
- itestpart

txelec

character(11) default: 'charge'

Describes the complexity of the electrostatic interactions. Available sources of electrostatic interactions are: (permanent) charges, weak charges, dipoles, and induced dipoles. Charges, dipole moments, and polarizability tensors employed are specified in namelist nmlParticle.

- · 'charge': Atoms possessing charges only.
- 'weakcharge': Atoms possessing charges and weak charges only. Limited to hard-sphere monoatomic particles.
- 'dip': Atoms possessing charges and static dipoles only. MD for all boundary conditions, and MC for all except Ewald summation.
- 'pol': Atoms possessing charges, static dipoles, and induced dipoles only. MD or MCALL. Ref. Chem. Phys. 191, 195 (1995). Charges, dipole moments, and polarizability tensors employed are specified in namelist nmlParticle.
- 'dipsph': Atoms possessing charges and/or dipoles in a spherical geometry, no neighbour list; special energy routines.
- 'dieldis': Atoms possessing charges with the presence of a planar or spherical dielectric discontinuity.

Iclink

logical default: .false.

- .true.: Enabling cross-links between particles and chains or between chains. Diamond-like network and bottle-brushes are currently supported.
- .false.: No cross-links.

Imultigraft

logical default: .false.

- true.: Enabling multigrafted polymers made of chains.
- .false.: No multigrafting.

Ihierarchical

logical default: .false.

- . true .: Enabling hierarchical structures made of chains.
- · .false.: No hierarchical structures.

maxnbondcl

```
integer(1:npt) default: npt*4
```

• Maximum number of cross-links of a particle (only txsetconf='hierarchical').

Ireadbondcl

logical default: .true.

- . true .: Enable reading of cross-linking information from .cnf-file when txstart='zero'
- .false.: No reading of cross-linking information when txstart='zero'

ngen

integer default: 1

Number of generations of hierarchical structure (only lhierarchical=.true. and txsetconf ='hierarchical').

ictgen

```
integer(1:ngen) default: 1
```

• Generation number -> chain type (only txsetconf='hierarchical').

nbranch

```
integer(0:ngen -1) default: 0
```

• Number of branches of a branching point of generation igen (only lhierarchical=.true. and txsetconf = 'hierarchical').

ibranchpbeg

```
integer(0:ngen -1) default: 0
```

• Chain segment of first branching point of generation igen (only lhierarchical=.true. and txsetconf = 'hierarchical').

ibranchpinc

```
integer(0:ngen -1) default: 1
```

• Segment increment between branching point of generation igen (only lhierarchical=.true. and txsetconf = 'hierarchical').

nnwt

integer default: 0

• Number of network types (only txsetconf = 'network').

nct

integer default: 0

· Number of chain types.

txct

```
character(10)(1:nct)
```

Text label for each chain type.

ncct

integer(1:nct)

· Number of chains of each chain type.

npptct

```
integer(1:npt ,1:nct )
```

• npptct (ipt,ict) is the number of particles of type ipt belonging to chain of type ict. Note, either non or all particles of a given type has to belong to chains, in the latter case sum(ncct (1:nct)*npptct (ipt,1:nct)=nppt (ipt) is required.

txcopolymer

character(30)(1:nct) default: nct *'block'

- 'block': Block copolymer.
- 'regular': Regular copolymer (alternating as possible).
- 'sequence': Copolymer with highly specific monomer distribution (the control is given in nmlCopolymerSequence).
- 'repeating': Copolymers with a defined repeating block structure (the control is given in nmlRepeating).
- · 'random': Random Copolymer.

nblockict

```
integer(1:nct) default: nct*0
```

· Number of blocks in each repeating of a chain of a chain type.

npt

integer

· Number of particle types.

txpt

```
character(10)(1:npt)
```

• Text label for each particle type.

nppt

```
integer(1:npt)
```

• Number of particles of each particle type.

natpt

```
integer(1:npt)
```

• Number of atom types of each particle type.

txat

```
character(10)(1:nat)
```

• Text label for each atom type (global atom type list).

massat

```
real(1:nat)
```

· Mass of atom type.

radat

real(1:nat)

 Hard-sphere radius of atom type. It is used to avoid an unreasonable start configuration (see namelist nmlSetConfiguration) and to check that atoms do not come too close to each other due to potential or program error. The check is performed after each macrostep by routine CheckHS
 Overlap. The value of radat should be smaller than the van der Waals radius, approximately 75% of it. In the case of a hard-core potential and MC, radat should be equal to the hard-core radius of the atom.

zat

real(1:nat) default: nat *0.0

• Charge of atom type. The charge should be given in number of elementary charges.

zatalpha

real(1:nat) default: 0.0

- >0.0: Gaussian charge distribution with width 1/(sqrt(2)*zatalpha)
- =0.0: Point charge

sigat

real(1:nat) default: nat *0.0

• Lennard-Jones σ parameter of atom type; $u(r) = 4\epsilon [(\sigma/r)^{12} - (\sigma/r)^6]$.

epsat

real(1:nat) default: (1:nat)

• Lennard-Jones ϵ parameter of atom type; $u(r) = 4\epsilon [(\sigma/r)^{12} - (\sigma/r)^6]$.

latweakcharge

logical(1:nat) default: nat*.false.

- . true .: Weak (titrating) charge of atom type (only txelec ='weakcharge').
- .false.: No weak charge.

jatweakcharge

integer(1:nat) default: nat*0

• type of atom carrying counter charge to weak charge iat (0 means no counter charge)

pΚ

```
real(1:nat) default: nat *0.0
```

pK value of the weak charge.

pН

real default: 0.0

· pH of the solution.

naatpt

```
integer(1:nat,1:npt)
```

• naatpt (ialoc,ipt) denotes the no of atoms of type ialoc (local list) on a particle of type ipt. ialoc runs from 1 to natpt (ipt), the no of atom types of particle type ipt.

txaat

```
character(10)(1:napt,1:npt)
```

• txaat (ialoc,ipt) is a text label for atom of no ialoc (local list) on particle of type ipt. ialoc runs from 1 to napt(ipt), the no of atoms of particle type ipt.

rain

```
real(1:3,1:napt,1:npt)
```

• rain (1:3,ialoc,ipt) is the x:y:z-coordinate of atom no ialoc (local list) in a particle of type ipt. ialoc runs from 1 to napt(ipt), the no of atoms of particle type ipt. rain need not necessarily be given in the principle frame axes.

dipain

```
real(1:3,1:napt,1:npt)
```

• dipain (1:3,ialoc,ipt) is the x:y:z-component of the dipole moment of atom no ialoc (local list) in a particle of type ipt. ialoc runs from 1 to napt(ipt), the no of atoms of particle type ipt. dipain has to be given in the same frame as rain.

polain

```
real(1:napt,1:npt)
```

• polain (1:6,ialoc,ipt) is the xx:yy:zz:xy:xz:yz-component of the symmetric polarizability of atom no ialoc (local list) in a particle of type ipt. ialoc runs from 1 to napt(ipt), the number of atoms of particle type ipt. polain has to be given in the same frame as rain.

lintsite

logical default: .false.

- .true.: Positions of interaction sites are given by raintin.
- .false.: Positions of interaction sites are given by rain.

raintin

```
real(1:3,1:napt,1:npt)
```

• raintin (1:3,ialoc,ipt) is the x:y:z-coordinate of interaction size no ialoc (local list) in a particle of type ipt. ialoc runs from 1 to napt(ipt), the no of atoms of particle type ipt.

Iradatbox

logical default: .false.

- . true .: Check that atoms including their hard-sphere radii are inside the box.
- .false.: No such check.

itestpart

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- · 10: Chain pointers

2.6 nmlCopolymerSequence

The namelist nmlCopolymerSequence contains variables that describe the sequence of copolymers.

- · Variables:
 - iptsegct

iptsegct

integer(maxval(npct(1:nct)),1:nct) default: npct*nct*0

· Particle type ipt of segments iseg of chain type ict

2.7 nmlNetworkConfiguration

The namelist nmlNetworkConfiguration contains variables that describe the number of networks the particles and chains, which form the networks and the network topology.

- · Variables:
 - nnwnwt
 - ncctnwt
 - txnwt
 - txtoponwt
 - iptclnwt

nnwnwt

```
integer(1:nnwt) default: nnwt*0
```

• Number of networks of network type inwt (only txsetconf = 'network').

ncctnwt

```
integer(1:nct,1:nnwt) default: nct *nnwt *0
```

• Number of chains of chain type ict of network type inwt (only txsetconf = 'network').

txnwt

```
character(10)(1:nnwt) default: nnwt *"network"
```

• Text label for each network type inwt (only txsetconf = 'network').

txtoponwt

```
character(30)(1:nnwt) default: nnwt *'default'
```

• Controls the network topology to be set (currently only 'default' possible and only txsetconf = 'network').

iptclnwt

```
integer(1:nnwt) default: nnwt*0
```

• Type of particles forming cross-links of networks of network type inwt (only txsetconf = 'network').

CHAPTER 2. INPUT 2.8. NMLREPEATING

2.8 nmlRepeating

The namelist nmlRepeating contains variables that define the repeating block structure of copolymers. The copolymers consist of repeating units, each consisting of blocks of one particle type.

- · Variables:
 - rep_block_ict

rep_block_ict

block_type(1:pt,1:np) **default:** pt*np*0

• Particle type and number of particles in in each block and chain type.

2.9 nmlPotential

The namelist nmlPotential contains variables that describe the potentials between the particles. Tabulated potential energy and force evaluation, which often gives a fast code by avoiding square root and divisions. Procedure according to Andrea, Swope, and Andersen, JCP 79, 4576 (1983).

- · Variables:
 - r2uminin
 - utoltab
 - ftoltab
 - umaxtab
 - fmaxtab
 - rcut
 - txpot
 - npot
 - ipot
 - ucoff
 - relpermitt
 - Iscrc
 - scrlen
 - Iscrhs
 - lewald
 - txewaldrec
 - iewaldopt
 - uewaldtol
 - ualpha

CHAPTER 2. INPUT 2.9. NMLPOTENTIAL

- ualphared
- ncut
- ncutregion
- Isurf
- lewald2dlc
- order
- nmesh
- Irf
- epsrf
- epsimg
- radimg
- epsi1
- epsi2
- boundaryrad
- Imaxdiel
- lbg
- Iljcut
- Ijrcut
- ljushift
- lambda_sw
- epsilon_sw
- alpha_sw
- lambda_ramp
- epsilon_ramp
- alpha_ramp
- rad_dep
- rho_dep
- factor_dep
- lellipsoid
- radellipsoid
- aellipsoid
- Isuperball
- radsuperball
- qsuperball

- txmethodsuperball
- nitersuperball
- tolsuperball
- meshdepthsuperball
- dl_damp
- dl_cut
- dr_damp
- dr_cut
- Istatsuperball
- luext
- Imonoatom
- itestpot

r2uminin

real deault: 0.1

· Square of the lower end of the tabulated potential.

utoltab

real default: 10d-5

• Energy tolerance of the tabulated potential.

ftoltab

real default: 10d-5

• Force tolerance of the tabulated potential.

umaxtab

real default: 2*10d4

• Energy at which the table is cut off at small separation.

fmaxtab

real default: 2*10d4

• Force at which the table is cut off at small separation.

rcut

real default: 0.0

Cutoff distance of the potential and forces based on the particle - particle center of mass distance. If rcut=0.0, the cutoff distance sets to

```
• (boxlen2(1)**2 + boxlen2(2)**2 + boxlen2(3)**2)**1/2 (only txbc='xyz'),
```

- (boxlen2(1)**2 + boxlen2(2)**2 + boxlen(3)**2)**1/2 (only txbc='xy'),
- (boxlen(1)**2 + boxlen(2)**2 + boxlen2(3)**2)**1/2 (only txbc='z'),
- 2*rsph (only txbc='sph'), or
- (2*rcyl)**2+lcyl**2)**1/2 (only txbc= 'cyl'),

txpot

character(20)

- Text label used for selecting potential of each pair of particle type. The potentials may either be already existing in the program or supplied by the user. The available options are:
- '(1,6,12)': Charge plus Lennard-Jones interaction $u_{ij}(r) = q_i q_j / 4\pi \epsilon_0 r + 4\epsilon_{ij} ((\sigma_{ij}/r)^{12} (\sigma_{ij}/r)^6)$. The parameters q_i , σ_{ii} , and ϵ_{ii} are given in namelist nmlParticle. Lorentz-Berthelot mixing rules are applied for cross terms.
- 'setx': Spherical Ewald potential. As the default potential, but the 1/r term is multiplied with erfc(r*ualphared/rcut).
- 'mcy': The MCY water potential, ref. JCP 64, 1351 (1976).
- 'nemo:xxx': The two-body part of the Nemo potential 'xxx'. The potential form and the coefficients are read from file molsim.lib. Present installed potentials include nemo:ww (water-water) and nemo:uw(urea-water). Note, the specification of the many-body polarization interaction is given in namelist nmlParticle.
- 'sw': Square-well potential. Parameters: lambda sw, epsilon sw and alpha sw
- 'ramp': Ramp potential. Parameters
- · 'asakura-oosawa': Asukura-Oosawa potential
- 'xxx': Search for user-provided potential labeled 'xxx' called from routine PotentialUser in file moluser.F90.
- If there is no match, the default potential form sum[ucoff (1:npot)/r**ipot(1:npot)] and the variables npot, ipot, and ucoff, which are read in this namelist, are used. If zat /= 0, the Coulomb term need not to be specified.

npot

integer(1:natat)

• npot (iatjat) denotes the number of terms of atom type pair iatjat.

ipot

```
integer(1:npot ,1:natat)
```

• ipot (m,iatjat) denotes the exponent of term m of atom type pair iatjat.

ucoff

```
real(1:npot, 1:natat)
```

• ucoff (m,iatjat) denotes the coefficient of term m of atom type pair iatjat.

relpermitt

real default: 1.0

· Relative permittivity.

Iscrc

logical default: .false.

- . true.: Screened Coulomb potential. See also Iscrhs.
- .false.: No screened Coulomb potential.

scrlen

real

Screening length for the screened Coulomb potential.

Iscrhs

logical default: .false.

- .true.: U(r) = zat (iat) * jat)/r*exp(-r/scrlen)*FAC(iat)*FAC(jat) with FAC(iat) = exp(-r*radat (iat))/(1+radat (iat)/scrlen)
- .false.: U(r) = zat (iat)*zat (jat)/r*exp(-r/scrlen). If zat all are zero ucoff (1,iatjat) is used.

lewald

logical default: .false.

- .true.: Ewald summation. Implemented for txpot='(1,6,12)', txpot=nemo... with txelec='pol', and default potential. Periodic boundary conditions are required.
- .false.: No Ewald summation.

txewaldrec

character default: 'std'

- 'std': standard Ewald summation.
- 'spm': smooth particle mesh Ewald summation (require installation of the FFTW library from www. ← fftw.org; see makefile). If only the standard Ewald summation is used, calls to subroutines fftw... in files energy.F90 and denergy.F90 can be comment away.

iewaldopt

integer

- Controls the choice of Ewald truncation analysis. ualphared = ualpha*rcut.
- For txewaldrec = 'std':
- 0: ualphared, rcut, and ncut are used.
- 1: ualphared and rcut are used to calculate uewaldtol and ncut. 1
- 2: uewaldtol and ualpha are used to calculate rcut and ncut.
- 3: uewaldtol and rcut are used to calculate ualpha and ncut. 1
- 4: uewaldtol and ncut are used to calculate ualpha and rcut.
 - ¹ According to Kolafa and Perram (charges) and Wang and Holm (dipoles).
- For txewaldrec = 'spm':
- 0: ualphared, rcut, order and nmesh are used.
- 3: uewaldtol, rcut, order and nmesh are use.

uewaldtol

real default: 0.0

Potential energy tolerance in Ewald summation (see iewaldopt).

ualpha

real

• Ewald parameter (see iewaldopt).

ualphared

real default: 3.0

 Reduced Ewald parameter used for the Ewald summation (see iewaldopt) and the Spherical Ewald potential.

ncut

integer

• Largest number of k-vectors in one direction in the reciprocal space (see iewaldopt).

ncutregion

character(6) default: 'sphere'

- · 'sphere': Spherical k-space region.
- 'cube': Cubic k-space region.

Isurf

logical default: .true.

- . true.: Inclusion of the surface term of the Ewald summation (corresponding to ϵ (surrounding) = 1).
- .false.: Exclusion of the surface term of the Ewald summation (corresponding to ϵ (surrounding) -> infinity).

lewald2dlc

logical **default**: .false.

- .true.: Correction for making a 3d-periodic system 2d-periodic according to Arnold et al. JCP 2002. Should only be applied with considerable insight on, e.g, the need of making the simulation box longer than the simulated system in the z-direction and the convergence.
- .false.: No such correction.

order

integer default: 5

• Interpolation order in the reciprocal space (see iewaldopt).

nmesh

integer default: 48

• Number of meshes used in the reciprocal space (see iewaldopt).

lrf

logical **default:** .false.

- .true.: Reaction field method (only txelec = 'pol').
- .false.: No reaction field method.

epsrf

real default: 78.0

• Relative dielectric permittivity of the surrounding beyond rcut for reaction field method. Zero means infinity.

epsimg

real

• Relative dielectric permittivity of the surrounding medium (only txelec = 'dipsphimage').

radimg

real

• Radius of the surrounding medium (only txelec = 'dipsphimage').

epsi1

real

• Dielectric constant of the sphere (only txelec = 'dipdiel').

epsi2

real

• Dielectric constant outside the sphere (only txelec = 'dipdiel').

boundaryrad

real

• Radius of dielectric boundary (only txelec = 'dipdiel').

Imaxdiel

integer

• Truncation of I-sum (only txelec = 'dipdiel').

lbg

logical

• true.: Volume charge density of inside the sphere neutralizing the system (only txelec = 'dipdiel').

Iljcut

```
logical default: .false.
```

- . true.: Apply a cutoff and shift of the Lennard-Jones potential (only txpot (ipt) = '(1,6,12)').
- .false.: No cut and shift.

Ijrcut

```
real default: 2.0**(1.0/6.0)
```

• Cutoff distance in LJ-sigma units.

ljushift

real default: 1.0

• Shift of LJ potential in LJ-epsilon units.

lambda_sw

```
real default: 1.1
```

• End of the square-well potential in units of hard-sphere diameter.

epsilon sw

real default: 1.0

· Depth of the square-well potential.

alpha_sw

```
real default: 1.0d-3
```

• Regulate the soft slope change at lambda_sw . Higher alpha implies softer change. u(r) = slope(r-r_sw)(0.5+(1/Pi)*tan(alpha_sw (r-r_sw), slope = -epsilon_sw /r1atat(lambda_sw -1), r_sw = r1atat*lambda_sw for r1atat < r1atat(lambda_sw -1).

lambda_ramp

real default: 1.1

• End of the ramp potential in units of hard-sphere diameter.

epsilon_ramp

real default: 1.0

· Depth of the ramp potential.

alpha_ramp

```
real default: 1.0d-3
```

Regulate the soft slope change at lambda_ramp. Higher alpha implies softer change. u(r) = slope(r-r_ramp)(0.5+(1/Pi)*tan(alpha_ramp(r-r_ramp), slope = -epsilon_ramp /r1atat(lambda_ramp -1), r← _ramp = r1atat*lambda_ramp for r1atat < r1atat(lambda_ramp -1).

rad_dep

real

Radius of penetrable hard sphere (Asakura-Oosawa model)

rho_dep

real

• number density of penetrable hard sphere (asakura-oosawa model)

factor_dep

real default: 1.0

· depletion-thickness factor

lellipsoid

logical default: .false.

• .true.: Hard-core ellipsoidal (prolate) particles.

• .false.: No hard-core ellipoidal particles.

radellipsoid

real default: 1.0

· Radius of degenerated axes.

aellipsoid

real default: 1.0

• Aspect ratio: >1 prolate and <1 oblate.

Isuperball

logical **default:** .false.

• .true.: particles are superballs

radsuperball

real default: One radius of superballs

qsuperball

real default: One

• q parameter of superballs

txmethodsuperball

character(4) default: "nr"

· 'nr', 'mesh', 'test'

nitersuperball

integer **default:** 10

· maximal number of nr iterations

tolsuperball

real default: 1.0d-4

· tolerance of nr iterations

meshdepthsuperball

integer **default:** 4

· depth of mesh

dl_damp

real default: One

dl_cut

real default: 1d10

dr_damp

real default: One

dr_cut

real **default**: 1d10

Istatsuperball

logical default: .false.

• . true: engage time statistics

luext

logical default: .false.

- . true.: Application of external potentials on the particles.
- .false.: No external potential.

Imonoatom

logical default: .true.

- .true.: The program checks to see if all particles have only one atom each. If so, 1) sections involving orientational integration, orientational movement, and some orientational output are omitted and 2) simpler and faster potential and force routines are used.
- . false.: Forces the program to treat the particles as polyatomic. This option, which may make the program slower, is for maintaining and checking purposes.

itestpot

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- '1': Intermediate potential variables are written
- · 2: Examination of accuracy of two-body potentials
- · 3: Plot of two-body potentials
- 4: Examination of truncation error of Ewald summation (routine PotTwoBodyTab).

2.10 nmlPotentialChain

The namelist nmlPotentialChain contains variables that describe the potentials involving particles belonging to the same chain. There is a bond potential between two consecutive particles in a chain and an angular potential between three consecutive particles in a chain. The potentials are of the type (k/p)(x-x0)*p.

- · Variables:
 - bond
 - angle
 - clink
 - itestpotchain

bond

bond_var(real, integer, real)(1:nct) default: nct*bond_var(0.0, 2, 0.0)

Force constant, power, and equilibrium separation of bond potential.

angle

```
bond_var(real, integer, real)(1:nct) default: nct *bond_var(0.0, 2, 0.0)
```

• Force constant, power, and equilibrium separation of angular potential.

clink

```
bond_var(real, integer, real) default: bond_var(0.0, 2, 0.0)
```

· Force constant, power, and equilibrium separation of crosslinks.

itestpotchain

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- 1: Bond length table and bond angle table (routines BondLengthTab and BondAngleTab).

2.11 nmlPotentialExternal

The namelist nmlPotentialExternal contains variables that describe the potentials involving atoms and an external potential.

- · Variables:
 - txuext

txuext

```
character(20)(1:npt) default: npt*''
```

- Text label used for selecting external potential. The number of external potential is growing and a number of parameters specifying these parameters are available. See routine IOPotExternal for details.
- 'wall_z': hard walls at abs(z) = wall_z-ext
- 'sw_wall_zlow': square-well wall at z = -wall z-ext
- 'ramp_wall_z': ramp walls at abs(z) = wall_z-ext
- 'lj_wall_z': LJ walls at abs(z) = wall_z-ext
- 'lj_wall_z_ts': truncated and shifted LJ walls at abs(z) = wall_z-ext
- 'lj_wall_z_mod': heterogeneous LJ walls at abs(z) = wall z-ext
- 'lj_wall_zlow': LJ + ramp wall at z = -wall_z-ext
- 'lj_wall_desorb': truncated LJ + ramp wall at z = -wall_z-ext (Niklas)

- 'estat_field_z': homogeneous electrical field in z-direction
- · 'hom_charged_walls': interaction with a charged surface
- 'i_soft_sphere': external, soft, and spherical wall
- 'Gunnar_soft_sphere': external, soft, and spherical wall (Gunnar)
- 'out_hard_ellipsoid': hard ellipsoidal wall
- 'capsid_shell'; hard spherical capsid shell
- 'uniform_shell': hard spherical capsid shell with a uniform surface charge density
- 'sphdielboundary_q': dielectric sphere (multipole expansion)
- 'sphdielboundary_p': dielectric sphere (pairwise interaction)
- 'sphdielboundary': spherical dielectric boundary (pairwise interaction)
- 'core_shell': hard inner and outer spherical walls (Steffi)
- 'insulating_sphere': penetrable uniformly charged sphere (Steffi)
- · "hollow_sphere' hollow and charged sphere (Steffi)
- · See code for further details.

2.12 nmlPolarizationIter

The namelist nmlPolarizationIter contains variables that control the calculation of the many-body polarization contribution to the potential energy and forces, ref. JPC 94, 1649 (1990).

- · Variables:
 - tpolit
 - mpolit
 - npolit
 - Idamping

tpolit

real default: 10**-4

• Relative tolerance of the induced dipole moment in the iteration.

mpolit

integer default: 15

· Maximum number of iterations.

npolit

integer default: 5

· Interval of iteration.

Idamping

logical default: .false.

- . true.: Short-distance damping of electrostatic force when calculating induced dipole moment.
- .false.: Nothing.

2.13 nmlSetConfiguration

The namelist nmlSetConfiguration contains variables that control the generation of the start configuration and is used only txstart='setconf'.

- Variables:
 - txsetconf
 - nucell
 - rclow
 - rcupp
 - roshift
 - radatset
 - Iranori
 - bondscl
 - anglemin
 - iptnode
 - ictstrand
 - rnwt
 - shiftnwt
 - txoriginnwt
 - radlimit
 - ntrydef
 - itestcoordinate

txsetconf

character(20)(1:npt)

- Text label used for selecting start coordinates. Existing routines or user provided routines are used.
- If, nmlSetConfiguration ='...lattice', particles are placed in the corner with the lowest x, y, and z-coordinate and at the surfaces with the lowest x, y, and zcoordinates, respectively, of a unit cell.
- If nmlSetConfiguration ='...random', hard-core overlap test with already existing particles is made. If coordinates could not be generated after 100*nppt (ipt) attempts with particles of type ipt, the program is stopped.
- 'origin': Set one particle in the center of the box.
- 'pclattice': Generate a primitive cubic lattice (1 particle in the unit cell).
- 'bcclattice': Generate a body centered cubic lattice (2 particles in the unit cell).
- 'fcclattice': Generate a face centered cubic lattice (4 particles in the unit cell).
- 'sm2lattice': Generate a sm2 lattice (4 particles in the unit cell).
- 'diamondlattice': Generate a cubic diamond lattice (8 particles in the unit cell).
- 'h2olattice': Generate a cubic lattice, im3m (ice VIII) (2 particles in the unit cell).
- 'n2lattice': Generate a cubic lattice, pa3 (solid N2) (4 particles in the unit cell).
- 'benzenelattice': Generate a cubic lattice, pbca (solid benzene) (4 particles in the unit cell).
- 'random': Generate random positions and orientations
- 'randomfixori': Gererate random positions and fixed orientations
- 'chainline': Generate a straight configuration with chain particles on a line (x-dir)
- 'chaincircle': Generate a straight configuration with chain particles on a circle (xy-plane)
- 'chainrandom': Generate random positions and orientations for chain particles.
- 'chainrandomintori': Generate random positions and int. fixed orientations for chain particles.
- 'sphbrushlattice': Generate a lattice brush on a spherical surface, first segment placed on a lattice.
- 'sphbrushrandom': Generate a random brush on a spherical surface, first segment randomly placed.
- 'planbrushrandom': Generate a random brush on a planar surface.
- 'hierarchicallattice': Generate a hierarchical polymer, first segment placed on a lattice.
- 'hierarchicalrandom': Generate a hierarchical polymer, first segment randomly placed.
- 'perodicnetwork': Generate a periodic network (diamond-like containing 8 nodes in a unit cell).
- · 'network': Generate a nonperiodic network (diamond-like containing 8 nodes in a unit cell).
- 'coreshell': Generate particle positions in a spherical shell.
- 'xxx': Search for user-provided routine labeled 'xxx' called from routine SetParticleUser in file moluser.F90.
- If there is no match, the program stops.

nucell

integer(1:3,1:npt) default: 3npt*0

• Number of unit cells in the x-, y-, and z-direction. The number of particles of type ipt has to be at most nucell (1,:)*nucell (2,:)*nucell (3,:)*npl, where npl is the number of particles in the unit cell (1 if PC, 2 if BCC, and 4 if FCC).

rclow

```
real(1:3,1:npt) default: 3npt*(-0.5*box(1))
```

• Lower x-, y-, and z-coordinate for the random positions or the set of unit cells. rclow may be unequal for different particle types to allow for a separation of particles of different types.

rcupp

```
real(1:3,1:npt) default: 3npt*(+0.5*box(1))
```

• Upper x-, y-, and z-coordinate for the random positions or the set of unit cells.

roshift

```
real(1:3,1:npt) default: 3npt*0.0
```

• Shift of the lattice points in a unit cell in fraction of the unit cell length.

radatset

```
real(1:npt) default: radat
```

• Hard-sphere radius used to create the start configuration.

Iranori

```
logical(1:npt) default: npt*.false.
```

- . true.: Random particle orientation (only txsetconf='origin' and '...lattice').
- .false.: Equal particle orientation (x' = x, y' = y, and z' = z).

bondscl

```
real(1:nct) default: nct*1.0
```

· Bond length scaling factor for chains.

anglemin

```
real(1:nct) default: nct*0.0
```

• Smallest allowed angle between consecutive particles in a chain (only nct>0).

iptnode

integer default: 0

• Type of particles of nodes (only txsetconf = 'periodicnetwork').

ictstrand

integer default: 0

Type of chain of strands (only txsetconf = 'periodicnetwork').

rnwt

```
real(1:nnwt) default: nnwt*10.0
```

• Cropping sphere radius of networks of network type inwt (only txsetconf = 'network').

shiftnwt

```
real(3,1:nnwt) default: 3*nnwt*0.0
```

• x-,y- and z-shift of the center of the cropping sphere of network type inwt in units of the unit cell

txoriginnwt

```
character(8)(1:nnwt) default: nnwt*'random'
```

- Selecting center of networks of different types (only txsetconf='network').
- 'origin': Set one network in the center of the box.
- 'random': Generate random positions of the centers of the networks.

radlimit

real(2)

· Lower and upper radial limit for placing particles (only txsetconf='coreshell').

ntrydef

integer default: 100

· number of trials of setting the configuration per particle

itestcoordinate

integer default: 0

• Flag for test output. This possibility is for maintenance purposes.

- 0: Nothing. The normal option.
- 1: Write crosslink data.

2.14 nmIMD

The namelist nmIMD contains variables that control the MD simulation.

- · Variables:
 - integ
 - nmlMDtstep
 - tvvite
 - nvvite
 - Isetvel
 - Izeromom
 - tvscl
 - tlscl
 - compre

integ

character(6)

- verlver Integration according to the velocity form of the Verlet algorithm.
- gear3 Integration according to a third-order Gear algorithm.
- gear4 Integration according to a fourth-order Gear algorithm.

nmlMDtstep

real

• Time step of the MD integration.

tvvite

real default: 0.0

• Factor that determines the initial quaternion velocity for the iteration of the quaternion velocity (only integ='velver'; 0.0 is fine).

nvvite

integer default: 2

Number of iterations for the quaternion velocities (only integ='velver'; 2 is preferred).

Isetvel

logical default: .true. (only txstart = 'setconf' .or. 'zero'); .false. (only txstart = 'continue')

- . true.: Linear and angular velocities are set according to a Maxwell distribution using the temperature temp.
- .false.: No set of linear and angular velocities.

Izeromom

logical default: .false.

- . true.: Modify linear and angular velocities (only possible) to get zero linear and angular moments while preserving the translational and rotational temperature (only lsetvel=.true.).
- · .false.: Nothing.

tvscl

real default: 0.0

• Time constant for the velocity scaling. The scaling is applied if tvscl>0.0 and is then performed every time step according to velocity(new)=velocity(old)* sqrt(1+(tstep/tvscl)*(temp /t-1)), where t is the instantaneous temperature (Berendsen et al., 1984). tvscl=tstep scales the velocities to give t=temp.

tisci

real default: 0.0

• Time constant for the length scaling. The scaling is applied if tlscl >0.0 and is then performed every time step according to length(new) = length(old)*(1+x)**(-1/3), x=(tstep/tlscl)*compre*(p-prsr), where p is the instantaneously pressure (Berendsen et al., 1984).

compre

real

· Compressibility used for the length scaling.

2.15 nmIMC

The namelist nmIMC contains variables that control the MC simulation. Each trial move involves an attempt to move one or several particles.

After each macrostep the relative difference of the total potential energy calculated from scratch and that from the updated energy is calculated and written below the heading 'check'. Normally the relative difference is less than $10^(-10)$. This might not hold if 'the linear displacement' \leq drnlist (see IONList).

- · Variables:
 - isamp
 - pspart
 - dtran
 - drot
 - lcl1spart
 - Ifixzcoord
 - Ifixxycoord
 - Ishiftzcom
 - Ifixchainstartspart
 - ispart
 - pspartcl2
 - txmembcl2
 - radcl2
 - dtrancl2
 - ppivot
 - txpivot
 - drotpivot
 - drotminpivot
 - ipivotrotmode
 - lcl1pivot
 - pchain
 - dtranchain
 - drotchain

- Icl1chain
- pslither
- pbrush
- dtranbrush
- drotbrush
- lcl1brush
- pbrushcl2
- dtranbrushcl2
- drotbrushcl2
- phierarchical
- dtranhierarchical
- pnetwork
- dtrannetwork
- pvol
- dvol
- pnpart
- chempot
- pcharge
- radcl1
- pselectcl1
- pspartsso
- Imcweight
- lautumb
- Imcpmf
- Imcsep
- itestmc

isamp

integer **default:** 1

- 0: Uniform sampling of particles, sequential selection.
- 1: Uniform sampling of particles, random selection.

pspart

```
real(1:npt) default: npt*1.0
```

Relative weight of a single-particle move. Further control is given by dtran and drot.

dtran

```
real(1:npt) default: npt*0.0
```

• 0.5*dtran is the maximal translational displacement of a particle along one box axis. Displacements are made along all box axes. If dtran>0, square region, or if dtran<0, spherical displacement region.

drot

```
real(1:npt) default: npt*0.0
```

• 0.5*drot is the maximal rotational displacement (in degrees) of one axis. If drot>0, rotation around one randomly selected box axis, or if drot<0, rotation around one randomly selected particle frame axis.

Icl1spart

logical(1:npt) default: npt*.false.

- .true.: Engage a single particle + cluster1 move involving a simultaneous trial move of (i) the selected particle (primary particle) and (ii) some or all particles located within the distance radcl1 from the selected one (secondary particles). The latter particles are selected with the probability pselectcl1.
- .false.: No single particle + cluster1 move.

Ifixzcoord

logical(1:npt) default: npt*.false.

- .true.: Fixed z-coordinate (restrict the trial move in the xy-plane) of particle. Applicable to spart and spart trial moves.
- .false.: No.

Ifixxycoord

logical(1:npt) default: npt*.false.

- .true.: Fixed xy-coordinate (restrict the trial move to the z-coordiante) of particle. Applicable to spart and spartcl2 trial moves.
- .false.: No.

Ishiftzcom

logical(1:npt) default: npt*.false.

• .true.: Shift the z-coordinate of all particles such that the center-of-mass of particles 1 and 2 is as close as possible to z = 0 (only txbc = 'cyl').

• .false.: No.

Ifixchainstartspart

logical(1:npt) default: .false.

- . true.: No trail-displacement of first particle in a chain (for grafted chains).
- · .false. No such restriction.

ispart

integer default: 0

- 0: Nothing, the normal use.
- \neq 0: For development/special use

pspartcl2

real(1:npt) default: npt*0.0

• Relative weight of a single particle + cluster2 move involving a simultaneous trial move of (i) the (original) cluster of type 1 as specified above and (ii) other similar clusters connected to the original one. The other clusters of type 1 consist of a central particle of the same type as the selected one and its neighboring particles in analogy with the original cluster. Clusters of type 1 are connected if their center-tocenter separation is directly or indirectly at most radcl2. Connected clusters of type 1 are called clusters of type 2. Further control is given by txmembcl2,dtrancl2 and radcl2.

txmembcl2

character(3)(1:npt)

- Selection of particle type of members belonging to clusters of type 2.
- =ipt: Search for members only of same particle type as that displaced.
- =all: Search for members among all particles.

radcl2

real(1:npt) default: npt*0.0

Largest separation between clusters of type 1 for belonging to same cluster of type 2.

dtrancl2

```
real(1:npt) default: npt*0.0
```

• 0.5*dtrancl2 is the maximal translational displacement of a particle along one box axis. Displacements are made along all box axes. If dtrancl2>0, square region, or if dtrancl2<0, spherical displacement region.

ppivot

```
real(1:nct) default: nct*0.0
```

• Relative weight of end-pivot rotation. The shorter end of the chain is rotated. Further control is given by drotpivot.

txpivot

```
character(5)(1:nct) default: nct*short
```

- · short: Rotation of the shorter subchain.
- lower: Rotation of the subchain containing particles with low numbers.
- upper: Rotation of the subchain containing particles with high numbers.

drotpivot

```
real(1:nct) default: nct*360
```

• Rotational displacement parameter of the end-pivot rotation.

drotminpivot

```
real(1:nct) default: nct*0.0
```

• Smallest rotational displacement of the end-pivot rotation.

ipivotrotmode

integer default: 1

- 1: Rotation around the bond joining pivot segment and the previous one. Bond angle is preserved.
- 2: Rotation round the normal to the plane formed by the pivot segments and its two neighbors.
- 3: Rotation around a random direction.

Icl1pivot

logical(1:npt) default: npt*.false.

• .true.: Engage an end-pivot + cluster1 rotation involving a simultaneous trial move of (i) the selected particle(s) (primary particle(s)) and (ii) some or all particles located within the distance radcl1 from the selected one (secondary particles). The latter particles are selected with the probability pselectcl1.

• .false.: No end-pivot + cluser1 move.

pchain

```
real(1:npt) default: npt*0.0
```

· Relative weight of chain move. Further control is given by dtranchain.

dtranchain

```
real(1:npt) default: npt*0.0
```

• Translational displacement parameter of the chain move.

drotchain

```
real(1:npt) default: npt*0.0
```

· Rotational displacement parameter of the chain move.

Icl1chain

logical(1:npt) default: npt*.false.

- .true.: Engage a chain + cluster1 move involving a simultaneous trial move of (i) the selected particles (primary particles) and (ii) some or all particles located within the distance radcl1 from the selected ones (secondary particles). The latter particles are selected with the probability pselectcl1.
- .false.: No chain + cluster1 move.

pslither

```
real(1:npt) default: npt*0.0
```

Relative weight of slithering move. All particles of a selected chain are subjected to a trial displacement (even for a homopolymer).

pbrush

```
real(1:npt) default: npt*1.0
```

• Relative weight of a brush move. Simultaneous move of a particle and its grafted chains. Requires some assumption of chain and bead labeling.

dtranbrush

```
real(1:npt) default: npt*1.0
```

· Translational parameter for brush move.

drotbrush

```
real(1:npt) default: npt*1.0
```

· Rotational parameter for brush move.

Icl1brush

```
logical(1:npt) default: npt*.false.
```

- .true.: Engage brush + cluster1 move involving a simultaneous trial move of (i) the selected particles (primary particles) and (ii) some or all particles located within the distance radcl1 from the selected ones (secondary particles). The latter particles are selected with the probability pselectcl1.
- .false.: No brush + cluster1 move.

pbrushcl2

```
real(1.npt) default: npt*0.0
```

• Relative weight of a brush + cluster2 move.

dtranbrushcl2

```
real(1:npt) default: npt*1.0 Translational parameter for brush + cluster2 move.
```

drotbrushcl2

```
real(1:npt) default: npt*1.0
```

• Rotational parameter for brush + cluster2 move.

phierarchical

```
real(1:npt) default: npt*0.0
```

· Relative weight of a hierarchical move.

dtranhierarchical

```
real(1:npt) default: npt*1.0
```

· Translational parameter for a hierarchical move.

pnetwork

```
real(1:npt) default: npt*0.0
```

· Relative weight of a network move.

dtrannetwork

```
real(1:npt) default: npt*0.0
```

• Translational parameter for network move.

pvol

```
real(1:npt) default: npt*0.0
```

• Relative weight of volume-change move (only txensemb='npt' or 'ntp').

dvol

```
real default 0.0
```

• Volume change parameter (only txensemb='npt' or 'ntp').

pnpart

```
real(1:npt) default: npt*0.0
```

• Relative weight of 'change of number of particle' move (only txensemb='mvt' or 'mtv').

chempot

```
real(1:npt) default: npt*0.0
```

• Chemical potential (only txensemb='mvt' or 'mtv').

pcharge

```
real(1:npt) default: npt*0.0
```

· Relative weight of charge-change move.

radcl1

```
real(1:npt) default: npt*0.0
```

• Largest separation between a primary and secondary particle for which the secondary particle can belong to the cluster.

pselectcl1

```
real(1:npt) default: npt*0.0
```

Probability that a particle with the separation at most radcl1 from the primary particle is selected as
a secondary particle of the cluster. 0 ≤ pselectcl1 ≤ 1.

pspartsso

```
real(1:npt) default: npt*0.0
```

Relative weight of sso move. Further control is given in nmlSPartSSO.

Imcweight

logical default: .false.

- . true.: Applying a weighting function for potential of mean force MC simulations. Further specification is given in namelist nmlMCWeight.
- · .false.: No weighting function applied.

lautumb

logical default: .false.

- . true.: Automatic umbrella sampling according to O. Engkvist and G. Karlström. Further specification is given in namelist nmlUmbrella.
- .false.: No umbrella sampling.

Imcpmf

logical default: .false.

• .true: Direct pmf calculation using updated weights. Further specification is given in namelist nmlMCPmf.

• .false.: No direct pmf calculation calculation.

Imcsep

logical default: .false.

- .true.: Either local or non-local moves are carried out during one MC-Pass, not both. When only local moves are used the neighbour list is build using drnlist as set in nmlIntlist, if non-local moves are also used drnlist is set to four times the contour length.
- .false.: Local and non-local moves are carried out during one MC-Pass.

itestmc

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- 1: Probability data of different moves (routine TestIOMCProb)
- 2: Data concerning MC trial move (routine TestMCMove)
- 3: Data concerning volume change (routine TestVolChange, TestNPartCharge1, TestNPart← Charge2, TestChargeChange1, TestChargeChange2)

2.16 nmIMCAII

The namelist nmlMCAll contains variables that control the MCALL simulation. Each trial move involves an attempt to move all particles simultaneously.

- · Variables:
 - dtranall
 - drotall
 - lautumb

dtranall

real(1:npt)

• 0.5*dtranall is the maximal translational displacement of a particle along one box axis. Displacements are made along all box axes. If dtranall>0, square region, or if dtranall<0, spherical displacement region.

CHAPTER 2. INPUT 2.17. NMLMCWEIGHT

drotall

real(1:npt)

• 0.5*drotall is the maximal rotational displacement (in degrees) of one axis. If drotall>0, rotation around one randomly selected box axis, or if drot<0, rotation around one randomly selected particle frame axis.

lautumb

logical default: .false.

- . true.: Automatic umbrella sampling according to O. Engkvist and G. Karlström. Further specification is given in namelist nmlUmbrella.
- .false.: No umbrella sampling.

2.17 nmlMCWeight

The namelist nmlMCWeight contains variables that control the application of a weighting function for potential of mean force MC simulations between two particles. Presently, only results from routine Pot← MeanForce are reweighted.

- · Variables:
 - ipmcw1
 - ipmcw2
 - txpotmcw
 - npolmcw
 - acoeffmcw

ipmcw1

integer

• Identity of the particle number 1.

ipmcw2

integer

· Identity of the particle number 2.

CHAPTER 2. INPUT 2.18. NMLUMBRELLA

txpotmcw

```
character(11)
```

- polynomial: Weighting function: a(0) + a(1)*r +
- exponential: Weighting function: a(0)*exp(-a(1)*(r-a(2))).

npolmcw

integer default: 0

· Degree of the polynomial

acoeffmcw

```
real(1:npolmcw) default: npolmcw*0.0
```

• Coefficients of the weighting function.

2.18 nmlUmbrella

The namelist nmlUmbrella contains variables that control the automatic umbrella sampling.

- · Variables:
 - typeumb
 - ipumb1
 - ipumb2
 - rminumbrella
 - delumb
 - numbgrid
 - cupdate
 - umbcoord
 - Ireadumb

typeumb

Type of umbrella potential

character(6) default: ' '

• Type of umbrella potential (e.g. particle-particle, particle-wall ...)

CHAPTER 2. INPUT 2.18. NMLUMBRELLA

ipumb1

integer default: 0

· Particle dentifier for particle-particle or atom-atom umbrella sampling.

ipumb2

integer default: 0

· Particle dentifier for particle-particle or atom-atom umbrella sampling.

rminumbrella

real default: 3.0

· Minimum particle-particle distance for particle-particle or atom-atom umbrella sampling

delumb

real default: 1.0

· Distance between two grid points in xumb

numbgrid

integer default: 0.0

· Number of grid points in xumb

cupdate

```
character(4) default: ' '
```

· Type of update of the weighting function

umbcoord

```
character(1) default: 'r'
```

• if set, particles ipumb1 and ipumb2 can only move along the coordinate set by umbcoord with fixed orientation

Ireadumb

logical default: .false.

• .true.: the initial umbrella potential is read from file.

• .false.: the potential is calculated using xumb.

2.19 nmlMCPmf

The namelist nmlMCPmf contains variables that control the calculation of potential of mean force by updating weights.

- · Variables:
 - iptmcpmf
 - nbinmcpmf
 - rlowmcpmf
 - ruppmcpmf
 - termmcpmf

iptmcpmf

integer default: 1

• Type of the two particles for which the potential of mean force should be calculated.

nbinmcpmf

integer default: 100

· Number of bins of the potential of mean force.

rlowmcpmf

rlow **default:** 0.0

· Lower limit of sampled potential of mean force.

ruppmcpmf

real **default:** 100.0

• Upper limit of sampled potential of mean force.

CHAPTER 2. INPUT 2.20. NMLSPARTSSO

termmcpmf

real

· Controls the update of weights.

2.20 nmlSPartSSO

The namelist nmlSPartSSO contains variables that handle the sso-functionality. The SSO algorithm allows for on the fly optimization of the displacement parameter of the single particle move during the equilibration run.

- · Variables:
 - dtransso
 - maxtransso
 - nstepzero
 - nstepend
 - dtranfac
 - nssobin
 - Itestsso

dtransso

real(1:npt) default: npt*1.0

• Initial displacement parameter to be used during the sso move. The displacement is in a spherical volume with diameter dtransso

maxtransso

real(1:npt)

• Maximal allowed displacement parameter the SSO can use. The default value is set according to half the system size, depending on the geometry.

nstepzero

integer default: ceiling(sqrt(real(nstep)))

• number of MC-Passes when the displacement parameter is adapted the first time. See Phys. Procedia 2011, 15, 81-86.

CHAPTER 2. INPUT 2.21. NMLBD

nstepend

```
integer default: max(nstepzero , int(0.1*nstep))
```

• duration (in MC-Passes) of the last part of the SSO. This, and only this, part will be used to give the final value for the optimal displacement parameter. Do not set it shorter than nstepzero.

dtranfac

real default: 2

• Factor by which the displacement parameter is increased if no optimal displacement parameter is found.

nssobin

integer default: 20

number of bins used by the sso move. The more bins the more accurately the displacement parameter can be determined.

Itestsso

logical default: .false.

- .true.: Print out intermediate values after each SSO-Part.
- · .false.: Nothing.

2.21 nmIBD

The namelist nmlBD contains variables that control the BD simulation. The simulation is performed in the configurational space according to Ermark, 1975.

- Variables:
 - nmlBDtstep
 - dcoeff

nmlBDtstep

real

• Time step of the BD integration.

CHAPTER 2. INPUT 2.22. NMLINTLIST

dcoeff

real

· Isotropic particle self-diffusion coefficient.

2.22 nmlIntList

The namelist nmllntList contains variables that control the calculation of lists of nonbonded pairs to be considered in two-body energy evaluations.

- · Variables:
 - txintlist
 - Ivlistllist
 - inlist
 - drnlist
 - facnneigh

txintlist

character(8) default: 'vlist'

- 'vlist': Verlet neighbour lists. Further specification is given by inlist and drnlist.
- 'llist': Linked lists.

lvlistllist

logical default: .false.

• . true.: Use of liked lists to crease Verlet neighbour lists (only txintlist='nlist').

inlist

integer default: 0

- 0: Automatic check whether a new neighbour list should be calculated or not. If the sum of the displacements of any two molecules, since last calculation, is larger than drnlist a new list is generated. The number of neighbour list calculations is written after each macrostep.
- >0: Interval of calculating a new neighbour list.

drnlist

real default: 0.0

· Distance added to the potential cutoff for calculating the neighbour list.

CHAPTER 2. INPUT 2.23. NMLDUMP

facnneigh

real default: 2.0

• 0: Multiplicative constant of radius for calculation number of interacting particles.

2.23 nmlDump

The namelist nmlDump contains variables that control the interval and variables dumped or read. Dumping is performed when txmode = 'simulation' and reading when txmode = 'analysis'.

- · Variables:
 - idump
 - txptdump
 - Idpos
 - Idori
 - Idliv
 - Idanv
 - Idfor
 - Idtor
 - Ididm
 - Idlaz
 - Idumpuser

idump

integer default: 10

Dumping/reading is performed every idump step/pass. The types of quantities dumped are controlled by the remaining variables. If txstart='continue' the dump files are properly positioned according to the step/configuration saved in file FCNF.

txptdump

character(20) default: 'all'

- 'all': Data for particles of all types are dumped.
- 'xxx': Data for particles of type ipt is dumped when 'xxx'=txpt (ipt).

CHAPTER 2. INPUT 2.23. NMLDUMP

Idpos

logical default: .false.

- .true.: Dumping/reading of particle positions.
- .false. No dumping/reading of particle positions.

Idori

logical default: .false.

- . true.: Dumping/reading of particle orientations.
- .false.: No dumping/reading of particle orientations.

Idliv

logical default: .false.

- .true.: Dumping/reading of particle linear velocities.
- .false.: No dumping/reading of particle linear velocities.

Idanv

logical default: .false.

- . true.: Dumping/reading of particle angular velocities.
- .false.: No dumping/reading of particle positions.

Idfor

logical default: .false.

- .true.: Dumping/reading of particle forces.
- .false.: No dumping/reading of particle forces.

Idtor

logical default: .false.

- . true.: Dumping/reading of particle torques.
- .false.: No dumping/reading of particle torques.

CHAPTER 2. INPUT 2.24. NMLGROUP

ldidm

logical default: .false.

• .true.: Dumping/reading of particle induced dipole moments.

• .false.: No dumping/reading of induced dipole moments.

Idlaz

logical default: .false.

• . true.: Dumping/reading of particle charge state.

• .false.: No dumping/reading of particle charge state.

Idumpuser

logical default: .false.

• . true.: Call of DumpUser, driver for user-provided dumping routines.

· .false.: No such call.

2.24 nmlGroup

The namelist nmlGroup contains variables that control the division of particles into reference and field groups. Most single particle quantities calculated in static.F90 are averaged over particles belonging to same reference group. In case of two particle quantities, the field group specifies the particles with which the reference particle interacts or is close to. More details are given below.

- · Variables:
 - ref
 - field
 - lwref

ref

character(20) default: 'type=all'

- Text label used for selecting procedure of how to divide particles into reference groups. The procedure may either be already existing in the program or supplied by the user. The available options are:
- 'type=all': Each particle type constitute one group.
- 'networkgenerations': The different chain generations of networks will be assigned to different reference groups. Requires particle type of the strand particles to be ipt = 2.
- 'type=xxx': Particles of type xxx (txpt (ipt)='xxx') constitute the only group.

xxx: Search for user-provided section labeled 'xxx' called from routine GroupUser in file moluser.
 F90.

• If there is no match, the program stops.

field

character(20) default: 'type=all'

· Describes the field groups. Same options as for ref.

lwref

logical **default:** .false.

• .true.: Reference group data are written on file FGROUP.

• .false.: No writing.

2.25 nmlStatic

The namelist nmlStatic contains variables that control the interval of the analysis and static analysis routines used.

- · Variables:
 - istatic
 - Ispdf
 - Irdf
 - Irdfchain
 - Irdfsph
 - lg3
 - Irdfcond
 - Isf
 - langdf
 - langextdf
 - loridipdf
 - Ilsphharaver
 - Iradangdf
 - lkirkwoodgk
 - loripoldf
 - Innhb
 - Inndf

- Ichaindf
- Ichaintypedf
- Ichaintypeextdf
- lcbpc
- Iltt
- Icluster
- Izerosecondmoment
- Imultipoledf
- lenergydf
- lwidom1
- lwidom2
- Imeanforce1
- Imeanforce2
- Ipotmeanforce
- Isurfacearea
- lcrystalformat
- Itrajectory
- Isubstructuredf
- Inetworkdf
- Inetworkradialdf
- Istaticuser

istatic

integer default: 1

• Conformation sampling interval. Holds both for txmode = 'simulation' and 'analysis'.

Ispdf

logical **default:** .false.

- .true.: Single particle distribution functions are calculated. Further specification is given in namelist nmISPDF.
- .false: No calculation.

Irdf

logical default: .false.

• . true.: Radial distribution functions or running coordination numbers or are calculated. Further specification is given in namelist nmIRDF.

· .false: No calculation.

Irdfchain

logical default: .false.

• . true.: Radial distribution functions or running coordination numbers are calculated between center of masses of chains. Further specification is given in namelist nmlRDFChain.

· .false: No calculation.

Irdfsph

logical default: .false.

- . true.: Radial distribution functions or running coordination numbers are calculated for particles which positions are projected on a sphere. Further specification is given in namelist nmlRDFSph.
- · .false: No calculation.

lg3

logical default: .false.

- .true.: Normalized triplet correlation functions are calculated. Further specification is given in namelist nmlG3Dist.
- · .false: No calculation.

Irdfcond

logical default: .false.

- .true.: Conditional radial distribution functions are calculated. Futher specification is given in namelist nmlRDFCond.
- .false.: No calculation.

Isf

logical default: .false.

- . true.: Partial structure factors are calculated. Further specification is given in namelist nmlSF.
- · .false.: No calculation.

langdf

logical default: .false.

• .true.: Angular distribution functions are calculated. Further specification is given in namelist nmlAngDF.

• .false.: No calculation.

langextdf

logical default: .false.

• .true.: Angular 2d distribution functions with respect to external frame are calculated. Further specification is given in namelist nmlAngExtDF.

• .false.: No calculation.

loridipdf

logical default: .false.

• .true.: Orientation/dipole distribution functions are calculated. Further specification is given in namelist nmlOriDipDF.

• .false.: No calculation.

Ilsphharaver

logical default: .false.

• .true.: Averages of unnormalized spherical harmonics are calculated.

• .false.: No calculation.

Iradangdf

logical default: .false.

• .true.: Radial-angular 2d distribution functions are calculated. Further specification is given in namelist nmlRadAngDF.

• .false.: No calculation.

Ikirkwoodgk

logical default: .false.

• . true . : Kirkwood gk-factors are calculated. Further specification is given in namelist nmlKirkwoodgk.

• .false.: No calculation.

loripoldf

logical default: .false.

• . true.: Orientation polarization distribution functions are calculated. Further specification is given in namelist nmlOriPolDF.

• .false.: No calculation.

Innhb

logical default: .false.

• . true.: Number of nearest neighbours and no of hydrogen bonds are calculated. Further specification is given in namelist nmlNNHB.

• .false.: No calculation.

Inndf

logical default: .false.

• .true.: Nearest neighbour distribution functions are calculated. Further specification is given in namelist nmlNNDF.

• .false.: No calculation.

Ichaindf

logical default: .false.

• .true.: Chain distribution functions are calculated. Further specification is given in namelist nmlChainDF.

• .false.: No calculation.

Ichaintypedf

logical default: .false.

• .true.: Chain distribution functions are calculated. Further specification is given in namelist nmlChainTypeDF.

• .false.: No calculation.

Ichaintypeextdf

logical default: .false.

• .true.: Chain type distribution functions with respect to the lab frame are calculated. Further specification is given in namelist nmlChainTypeExtDF.

• .false.: No calculation.

Icbpc

logical default: .false.

• . true.: Probabilities of chain particles to be near particles of another type are calculated. Further specification is given in namelist nmlCBPC.

• .false.: No calculation.

lltt

logical default: .false.

• .true.: Loop, tail, and train statistics are calculated. Further specification is given in namelist nmlLoopTailTrain.

• .false.: No calculation.

Icluster

logical default: .false.

• . true .: Cluster size distribution functions are calculated. Further specification is given in namelist nmlCluster.

• .false.: No calculation.

Izerosecondmoment

logical default: .false.

• . true.: Zero and second moment for an ionic, neutral system is calculated.

• .false.: No calculation.

Imultipoledf

logical default: .false.

• . true.: Electrostatic multipole moment distribution functions are calculated. Further specification is given in namelists nmlMultipoleDF.

• .false.: No preparation.

lenergydf

logical default: .false.

• .true.: Energy distribution functions are calculated. Further specification is given in namelist nmlEnergyDF.

• .false.: No calculation.

lwidom1

logical default: .false.

• .true.: Excess chemical potentials are prepared. Further specification is given in namelists nmlWidom1.

• .false.: No preparation.

lwidom2

logical default: .false.

• .true.: Excess chemical potentials are prepared. Further specification is given in namelists nmlWidom2.

• .false.: No preparation.

Imeanforce1

logical default: .false.

• .true.: Mean force between two particles is calculated. Further specification is given in namelist nmlMeanForce1.

• .false.: No calculation.

Imeanforce2

logical default: .false.

• .true.: Mean force between two particles is calculated. Further specification is given in namelist nmlMeanForce2.

• .false.: No calculation.

Ipotmeanforce

logical **default:** .false.

• . true.: Potential of mean force between two particles is calculated. Further specification is given in namelist nmlPotMeanForce.

· .false.: No calculation.

Isurfacearea

logical default: .false.

• . true.: Surface area available around atoms residing in particles of a specified type is calculated. Further specification is given in namelist nmlSurfaceArea.

• .false.: No calculation.

Icrystalformat

logical default: .false.

- . true.: Write atoms in the crystallographic format stating with atoms belong to molecules closest to the origin. Hard-code limitation of spatial distance and number of particles are in action.
- .false.: No calculation.

Itrajectory

logical default: .false.

- .true.: Write trajectory on FLIST. Further specification is given in namelist nmlTrajectory.
- .false.: No calculation.

Isubstructuredf

logical default: .false.

- . true.: Substructures of chains, hierarchical structures and networks are analyses. Further specification is given in namelist nmlSustructureDF.
- .false.: No calculation.

Inetworkdf

logical default: .false.

- . true .: Network distribution functions are calculated. Further specification is given in nmlNetworkDF.
- .false.: No calculation.

CHAPTER 2. INPUT 2.26. NMLSPDF

Inetworkradialdf

logical **default:** .false.

• .true.: Radial network distribution functions are calculated. Further specification is given in nmlNetworkRadialDF.

• .false.: No calculation.

Istaticuser

logical default: .false.

• . true.: StaticUser is called and from where user-provided static analysis routines are called in file moluser.F90.

• .false.: No call.

2.26 nmISPDF

The namelist nmISPDF contains variables that control the calculation of single particle distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	Quantity	
1	rrden	Reduced radial number density	
2	rren	Reduced running coordination number	
3	zden	Number density in the z-direction	
4	zden2	Number density in the z-direction (special)	
5	z'*z	Projection of molecular z'-axis on the box z-axis	
6	z'*x	Projection of molecular z'-axis on the box x-axis	
7	z'*y	Projection of molecular z'-axis on the box y-axis	
8	opt_d	Radial density projected on the z = 0 plane	
9	elpot	radial electrostaic potential evaluated at particle positions	
10	m*E	cos(m(ia)*E); m(ia) is the dipole moment of atom ia and E an external field	

· Variables:

- vtype

vtype

static1D_var(logical, real, real, integer, logical, character, real) (1:10)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- min: /0.0,0.0,-X,-X,-1.0,Y/
- X = lcyl/2 (only txbc = 'cyl'), box(3)/2 (else)

CHAPTER 2. INPUT 2.27. NMLRDF

- Y = rsph (only txbc = 'sph), rcyl (only txbc = 'cyl')
- max /10.0,10.0,X,X,1.0,0.0/

2.27 nmIRDF

The namelist nmIRDF contains variables that control the calculation of running coordination number or radial distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	com-com	particle-particle
2	com-xxx	particle-atom
3	XXX-XXX	atom-atom

- · Variables:
 - vtype
 - rmax
 - ndim
 - nbin
 - func
 - I2dtwo

vtype

static1D_var(logical, real, real, integer, logical, character, real) (1:3)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /0.0,0.0,0.0/ Max: /10.0,10.0,10.0/

rmax

real default: 10.0

• Upper distance for particle separation considered.

ndim

integer default: 3

- 2: Sample distribution function in the xy-plane.
- 3: Sample distribution function in the xyz-space.

CHAPTER 2. INPUT 2.28. NMLRDFCHAIN

nbin

integer default: 100

· Number of bins used to sample the distribution functions.

func

character(3) default: rdf

· rdf: Radial distribution functions are calculated.

• rcn: Running coordination numbers are calculated.

I2dtwo

logical default: .false.

- true.: Sampling of 2d radial distribution functions separately for z>0 and z<0. Useful for a system with txbc='xy' and two equivalent surfaces at z=±box2(3) (only ndim = 2)
- · .false.: Nothing.

2.28 nmIRDFChain

The namelist nmIRDFChain contains variables that control the calculation of running coordination number or radial distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	com-com	center of mass-center of mass

Only chains within a separation of rmax are considered.

- · Variables:
 - vtype
 - rmax
 - func

vtype

static1D_var(logical, real, real, integer, logical, character, real) (1:3)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /0.0/ Max: /10.0/

CHAPTER 2. INPUT 2.29. NMLRDFSPH

rmax

real default: 10.0

· Upper distance for particle separation considered.

func

character(3) default: rdf

· rdf: Radial distribution functions are calculated.

• rcn: Running coordination numbers are calculated.

2.29 nmIRDFSph

The namelist nmIRDFSph contains variables that control the calculation of running coordination number or radial distribution functions of particles which positions are projected on a sphere.

- · Variables:
 - ipsph
 - iptrcnsph
 - jptrcnsph
 - nbin
 - func

ipsph

integer

• The identity of the particle on which surface the projection is made.

iptrcnsph

integer

• Type of particle for which distribution function should be calculated.

jptrcnsph

integer

• Type of particle for which distribution function should be calculated.

CHAPTER 2. INPUT 2.30. NMLRDFCOND

nbin

integer default: 100

• Number of bins used to sample the distribution functions.

func

character(3) default: rdf

- · rdf: Radial distribution functions are calculated.
- rcn: Running coordination numbers are calculated.

2.30 nmIRDFCond

The namelist nmIRDFCond contains variables that control the calculation of conditional radial distribution functions. The distribution functions are made for five different bins of arccos(theta) making up -1 to 1.

- · Variables:
 - vtype
 - rmax

vtype

 $static1D_var(logical, real, real, integer, logical, character, real) Min: /0.0/ Max: /10.0/$

• Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.

rmax

real default: 10.0

• Upper distance for particle separation considered.

CHAPTER 2. INPUT 2.31. NMLG3DIST

2.31 nmlG3Dist

The namelist nmlG3Dist contains variables that control the calculation of normalized triplet correlation functions g(r12, r13, r23)/g(r12). r23 is represented by the angle theta formed by the vectors r12 and r13.

- · Variables:
 - ipt1
 - ipt2
 - ipt3
 - snbin
 - slow
 - supp
 - tnbin
 - tlow
 - tupp
 - anbin
 - alow
 - aupp
 - nskip

ipt1

integer

• Particle type of the first particle.

ipt2

integer

• Particle type of the second particle.

ipt3

integer

• Particle type of the third particle.

CHAPTER 2. INPUT 2.31. NMLG3DIST

snbin

integer

• Number of bins to sample the distance r12.

slow

real

· Lower end of r12 to be sampled.

supp

real

• Upper end of r12 to be sampled.

tnbin

integer

• Number of bins to sample the distance r13.

tlow

real

• Lower end of r13 to be sampled.

tupp

real

• Upper end of r13 to be sampled.

anbin

integer

• Number of bins to sample the angle theta.

alow

real

· Lower end of theta to be sampled.

CHAPTER 2. INPUT 2.32. NMLSF

aupp

real

· Upper end of theta to be sampled.

nskip

integer default: 1

· Interval of listing the distance r12.

2.32 nmISF

The namelist nmlSF contains variables that control the calculation of partial structure factors.

- (i.) txbc='sph' or txbc='cyl' Variables:
 - txkscale
 - klow
 - logklow
 - logkupp
 - nbin
- (ii.) Cubic box and txbc='xyz'. The largest k-vector is 2Pi/box. Variables:
 - ndim
 - nbin
 - Iqsorted
 - Isi

txkscale

character(3) default: lin

- 'lin': Linear k-scale
- 'log': Logarithmic k-scale

klow

real **default:** if sphrad \neq 0: 2Pi/(10*sphrad), else 0.01

• Lower k-vector (linear scale)

CHAPTER 2. INPUT 2.32. NMLSF

logklow

real default: -4

· Lower k-vector (logarithmic scale)

logkupp

real default: 1

• Upper k-vector (logarithmic scale)

nbin

integer default: 100

Number of bins used to sample the partial structure factors.

ndim

integer default: 3

- 2: Structure factor in the xy-plane. It is averaged over 2 100 and 2 110 directions.
- 3: Structure factor in the xyz-space. It is averaged over 3 100, 6 110 directions, and 4 110 directions.

nbin

integer default: 100

• Number of bins used to sample the partial structure factors.

Iqsorted

logical default: .true.

- .false.: List separately the structure factors of the different types directions.
- . true.: Sort the structure factors of the different types of directions and list the sorted structure factor.

Isi

logical default: .false.

- .false.: Nothing.
- . true .: Scattering intensities are calculated. Further specification is given in nmlScatIntens.

CHAPTER 2. INPUT 2.34. NMLANGDF

2.33 nmlScatIntens

The namelist nmlScatIntens contains variables that control the calculation of the scattering intensities using a multi shell profile of constant scattering properties of each particle.

- · Variables:
 - nshell
 - rshell
 - cshell

nshell

```
integer(1:npt) default: nshell(1:npt) = 1
```

· Number of shells

rshell

```
real(1:nshell,1:npt) ** default:** rshell(1,1:npt) = [ (radat (ipt), ipt = 1,npt) ]
```

· Radius of the shells.

cshell

```
real(1:nshell ,1:npt ) default: cshell(1,1:npt ) = One
```

2.34 nmlAngDF

The namelist nmlAngDF contains variables that control the calculation of angular distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	z'*z'	cos(z'(ip)*z'(jp))
2	z'*r	cos(z'(ip)*r(ijp))
3	r*z'	cos(r(ijp)*z'(jp))
4	oh∗r	cos(oh(ip)*r(ijp)) ip has to be water
5	r*oh	cos(r(ijp)*oh(jp)) jp has to be water
6	oh.o	cos(oh(ip).o(jp)) ip and jp have to be water
7	0.0.0	cos(o(jp).o(ip).o(kp))
8	m*m	cos(m(ia)*m(ja)) m(ia) is the dipole vector of atom ia = ianpn(ip) + iashift
9	m*r	cos(m(ia)*r(ijp)) (the value of iashift is currently hard coded)
10	r∗m	cos(r(ijp)*m(ja))

h(ip) denotes the z'-axis of particle ip, r(ijp) the normalized vector between particle ip and jp defined as r(jp)-r(ip), oh(ip) the direction of a oh-bond in water, oh(ip).o(jp) the largest angle of the four possible

CHAPTER 2. INPUT 2.35. NMLANGEXTDF

formed by the oh(ip) direction and the h(ip)-o(jp) vector (hydrogen bond angle), and o(jp).o(jp).o(jp) the angle formed by the location of three particles. The sampling of types 4-6 assumes that the first three sites of water are oxygen, hydrogen, and hydrogen. Generally ip refers to reference particles and jp to field particles. Only field particles within a separation of rmax from a reference particle are considered.

- · Variables:
 - vtype
 - rmin
 - rmax

vtype

static2D_var(logical, 2*real, 2*real, 2*integer, logical, character, real)(1:10)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /(-1.0, 0.0)/ Max: /(1.0, 2Pi)/

rmin

real default: 0.0

• Lower distance for particle separation considered.

rmax

real default: 10.0

• Upper distance for particle separation considered.

2.35 nmlAngExtDF

The namelist nmlAngDF contains variables that control the calculation of 2D angular distribution functions with respect to an external frame.

type	label	quantity
1	z'-axis	cos(theta)-phi 2d distributin function

- · Variables:
- vtype

vtype

```
static1D_var(logical, real, real, integer, logical, character, real)(1:3)
```

CHAPTER 2. INPUT 2.37. NMLRADANGDF

• Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.

• Min: /7*(-1.0)/ Max: /7*0.0/

2.36 nmlOriDipDF

The namelist nmlOriDipDF contains variables that control the calculation of orientation/dipole distribution functions.

type	label	distribution functions
1	dir	orientation df based on ori(1:3,1:3)
2	dir_aver	symmetry-averaged orientation df
3	dip	dipole df
4	dip_tot	total dipole df

· Variables:

- vtype

vtype

static1D_var(logical, real, real, integer, logical, character, real)(1:3)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /4*(-1)/ Max: /4*1.0/

2.37 nmlRadAngDF

The namelist nmlRadAngDF contains variables that control the calculation of radial-angular 2d distribution functions.

type	label	quantity
1	x'y'-z'-plane	two particle radial - angular r2d distributin function

- · Variables:
 - vtype
 - txCoordSys
 - txAngle

vtype

static2D_var(logical, 2*real, 2*real, 2*integer, logical, character, real)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Default values depend on txCoordSys

txCoordSys

character(5) default: "polar"

· Coordinate system of distribution functions.

'polar': r and cos(theta)

• 'cart': sqrt(x**2+y**2) and z

txAngle

character(6) default: "theta1"

· Choice of angle.

• 'theta1': theta 1

· 'theta2': tehta 2

· 'psi': psi

2.38 nmlKirkwoodgk

The namelist nmlKirkwoodgk contains variables that control the calculation of running orientation averages. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	h∗h	sum_jp h(ip)*h(jp)

- · Variables:
 - vtype
 - rmax

vtype

static1D_var(logical, real, real, integer, logical, character, real)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization (by radius**3/2), title, and number of variable of vtype.
- Min: /0.0/ Max: /10.0/

CHAPTER 2. INPUT 2.39. NMLORIPOLDF

rmax

real default: 10.0

· Upper distance for particle separation considered.

2.39 nmlOriPolDF

The namelist nmlOriPolDF contains variables that control the calculation of orientation polarization distribution functions of molecular dipole moments. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	distribution functions
1	tot	orientation polarization, total
2	par	orientation polarization, parallel to central dipole (related to Kirkwood's Gk factor)
3	per	orientation polarization, perpendicular to central dipole
4	pevpa	orientation polarization, perpendicular versus parallel

- The analysis is made for a set of different radii
- The orientation polarization is normalized by (dipole moment)**2 or radius**(3/2)*(dipole moment)**2
- · Variables:
 - vtype
 - Inorm
 - nrad
 - radius

vtype

static2D_var(logical, real, real, integer, logical, character, real)(1:4)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization (by radius**3/2), title, and number of variable of vtype.
- Min: /0.0, -10.0, 0.0, 0.0/ Max: /4*(10.0)/

Inorm

logical default: .false.

• .true.: Invoking normalization.

• .false.: No normalization invoked.

CHAPTER 2. INPUT 2.40. NMLNNHB

nrad

integer default: 2

· Number of radii to consider

radius

real(1:nrad) default: 20.0

· Radii to consider

2.40 nmINNHB

The namelist nmlNNHB contains variables that control the calculation of no of nearest neighbours and no of hydrogen bonds. Field particles within thnn are considered as nearest neighbours to a reference particle. Field particles within rmax and with an interaction energy below a threshold value are considered as hydrogen bonded to a reference particle.

- · Variables:
 - nthnn
 - thnn
 - nthhb
 - thhb
 - nnnhb
 - rmax

nthnn

integer default: 2

· Number of separations considered.

thnn

real(1:nthnn) default: [3.3, 3.5]

• Distances considered. The largest thnn should not be larger than rmax given below.

nthhb

integer default: 2

· Number of energy thresholds.

CHAPTER 2. INPUT 2.41. NMLNNDF

thhb

real(1:nthhb) default: [-10.0,-16.0]

· Energy thresholds.

nnnhb

integer **default:** 10

· Maximal number of nearest neighbours and hydrogen bonds considered in the analysis.

rmax

real default: 3.5

• Upper separation distance for hydrogen bonded particles.

2.41 nmINNDF

The namelist nmlNNDF contains variables that control the calculation of nearest neighbour distribution functions. The R-F nearest neighbour distribution function gives the distribution of distances between a reference particle R and the nearest field particle F.

- · Variables:
 - vtype

vtype

static2D_var(logical, real, real, integer, logical, character, real)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization title, and number of variable of vtype.
- Min: /0.0/ Max: /10<u>.0/</u>

type	label	quantity	
------	-------	----------	--

2.42 nmlChainDF

The namelist nmlChainDF contains variables that control the calculation of chain distribution functions. Distribution functions are calculated for each chain. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	rbb	bead-bead separation
2	ree	end-to-end separation
3	rg	radius of gyration
4	angle	angle between consequtive beads
5	cos	cos(180-angle)
6	shape	ree**2/rg**2 ratio
7	asph	asphericity
8	torp	toroidicity

- · Variables:
 - vtype

vtype

static2D_var(logical, real, real, integer, logical, character, real)(1:8)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization title, and number of variable of vtype.
- Min: /0.0, 0.0, -1.0, 0.0, 0.0, 0.0, 0.0, 0.0/ Max: /12.0, 100.0, 100.0, 180.0, 1.0, 12.0, 1.0/

2.43 nmlChainTypeDF

The namelist nmlChainTypeDF contains variables that control the calculation of chain distribution functions. Distribution functions are calculated for each type of chains. Any combination of the types of distribution functions listed below may be selected through vtype%l. Same input variables as for nmlChainDF.

2.44 nmlChainTypeExtDF

The namelist nmlChainTypeExtDF contains variables that control the calculation of chain distribution functions where the distribution functions are with respect to the lab frame. Distribution functions are calculated for each type of chains. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	com_x	com, x-direction

CHAPTER 2. INPUT 2.45. NMLCBPC

type	label	quantity
2	com_y	com, y-direction
3	com_z	com, z-direction
4	rg_z	square of radius of gyration projected on the z-axis
5	rg_xy	square of radius of gyration projected on the xy-plane

- · Variables:
 - vtype

vtype

static2D_var(logical, real, real, integer, logical, character, real)(1:5)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization title, and number of variable of vtype.
- Min: /-box2(1),-box2(2),-box2(3),0.0,0.0/
- Max: /box2(1),box2(2),box2(3),100.0,100.0/

2.45 nmlCBPC

The namelist nmlCBPC contains variables that control the calculation of probabilities that chain particles are near particles of other types. Distribution functions are calculated for each pair of chain molecules and particles of type iptpart.

- · Variables:
 - iptpart
 - rcontact

iptpart

integer

· Type of particle to be considered.

rcontact

real

• Largest separation between chain particles and particles of the other type for considering the chain particle near the other particle.

2.46 nmlLoopTailTrain

The namelist nmlLoopTailTrain contains variables that control the calculation loop, tail, and train characteristics. Properties are calculated for each type of chains separately.

- · Variables:
 - adscond

adscond

adscond_var(character, character, real)

• Adsorption condition('xy-plane'), selection of box ends, threshold distance of adsorption.

2.47 nmlCluster

The namelist nmlCluster contains variables that control the calculation of cluster size distribution functions. Objects within the distance rcluster from a given object forms a virtual bond and all objects directly or indirectly bonded form a cluster.

- · Variables:
 - txobj
 - I1d
 - I2d
 - Ipercolation
 - nobjt
 - iobjt
 - rcluster
 - txweight
 - itestcluster

txobj

character(8) default: "particle"

- · Object for which cluster analysis should be performed.
- 'particle': Particles, rcluster applies between particles.
- 'chain': Chains, rcluster applies between chain particles.
- · 'chaincom': Chains, rcluster applies between chain com's.

CHAPTER 2. INPUT 2.47. NMLCLUSTER

l1d

logical default: .true.

- . true.: Calculation of 1d cluster size distribution functions.
- .false.: No such calculation.

I2d

logical default: .false.

- . true .: Calculation of 2d cluster size distribution functions.nptcluster=2 is required.
- .false.: No such calculation.

Ipercolation

logical default: .false.

- .true.: Percolation analysis is made.
- .false.: No such analysis.

nobjt

integer default: 1

· Number of object types to be considered.

iobjt

integer(1:npt) default: npt*0

· Object types to be considered.

rcluster

real

 Upper separation for bonded objects (particles if txobj='particle' or 'chain' and chain com if txobj='chaincom').

txweight

character(6) default: "mass"

- 'mass': Mass-weighted distribution.
- 'number': Number-weighted distribution.

itestcluster

integer default: 0

- Flag for test output. This possibility is for maintenance purposes.
- 0: Nothing. The normal option.
- 1: Intermediate cluster data (several routines in static.F90). Type of particles to be inserted in the different sets.

2.48 nmlMultipoleDF

The namelist nmlMultipoleDF contains variables that control the calculation of electrostatic multipole moment distribution functions.

- · Variables:
 - Imax
 - vmin
 - vmax
 - nbin
 - nrad
 - radius

Imax

integer default: 2

• The largest multipole moment considered is 2**Imax Real and imaginary components are analyzed separately and only for nonnegative m.

vmin

real(1:lmax) default: 0.0

• Lower end of the sampled distribution function of each type.

vmax

real(1:lmax) default: 10.0

• Upper end of the sampled distribution function of each type.

CHAPTER 2. INPUT 2.49. NMLENERGYDF

nbin

integer default: 100

• Number of bins used to sample the distribution functions.

nrad

integer default: 1

Number of radii for which multipole moment df are calculated. If txbc='sph', then nrad=1 and radius=rsph are forced; otherwise multipole moment are calculated over spheres centered on all particles.

radius

real(1:nrad) default: nrad*20.0

· Radii.

2.49 nmlEnergyDF

The namelist nmlEnergyDF contains variables that control the calculation of energy distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype. Only contributions from two-body potential terms are included.

type	label	quantity
1	bindu	binding energy, total
2	bindu	binding energy with particles in group jgr
3	pairu	pair energy with particles in group jgr within rmax

- · Variables:
 - vtype
 - rmax

vtype

static1D_var(logical, real, real, integer, logical, character, real)(1:3)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /-150.0,-150.0,-25.0/ Max: /0.0,0.0,0.0/

CHAPTER 2. INPUT 2.50. NMLWIDOM1

rmax

real default: 1.0d10

• Upper distance for particle separation for pair energy distribution function.

2.50 nmlWidom1

The namelist nmlWidom1 contains variables that control the calculation of excess chemical potentials using Widom's insertion method. Only charge neutral combinations of particles should be inserted.

- · Variables:
 - ntimes
 - nset
 - nptset
 - iptset

ntimes

integer **default:** 1

· Number of samplings per occasion.

nset

integer default: 1

· Number of sets to evaluate.

nptset

integer(1:nset)

· Number of particles to be inserted in a set.

iptset

integer(1:nptset,1:nset)

· Types of particles to be inserted in a sets

CHAPTER 2. INPUT 2.51. NMLWIDOM2

2.51 nmlWidom2

The namelist nmlWidom2 contains variables that control the calculation of excess chemical potentials using Widom's insertion. Single particles are inserted and charge integration according to Svensson and Jönsson.

- · Variables:
 - ntimes
 - nset
 - iptset

ntimes

integer default: 1

· Number of samplings per occasion.

nset

integer default: 1

· Number of sets to evaluate.

iptset

integer(1:nptset,1:nset)

· Types of particles to be inserted in a sets

2.52 nmlMeanForce1

The namelist nmlMeanForce1 contains variables that control the calculation of mean force between two particles (txbc='cyl' is required) according to F = Fmean + Fhs (the surface approach).

- · Variables:
 - dr

dr

real

· Displacement for evaluation of Fhs

2.53 nmlMeanForce2

The namelist nmlMeanForce2 contains variables that control the calculation of mean force between two particles (only txbc='cyl') according to F = Fcorr + Fideal(midplane and an end) + Fhs(midplane) (midplane approach).

- · Variables:
 - thickness
 - dz

thickness

real

• Thickness of volume for sampling Fideal

dz

real

· Displacement for evaluation of fhs

2.54 nmlPotMeanForce

The namelist nmlPotMeanForce contains variables that control the calculation of the potential mean force between two particles (only txbc='cyl'). The two particles of type iptpmf are assumed to be located on the z-axis. The pmf is set to zero at the separation cyllen /2.

- · Variables:
 - iptpmf
 - rpmfzero

iptpmf

integer default: 1

• Types of particles for which the potential of mean force is calculated. nppt(iptpmf) = 2 is required.

rpmfzero

real default: cyllen /2

· Separation at which pmf is set to zero.

2.55 nmlSurfaceArea

The namelist nmlSurfaceArea contains variables that control the calculation of the surface area available for a spherical prob.

- · Variables:
 - ipt
 - rprobe
 - wradat
 - nrandom

ipt

integer

• Type of particles of interest for area determination.

rprobe

real default: 1.7

· Radius of probe particle

wradat

real(1:nat) **default:** nat*0.0

• van der Waal radius of atoms.

nrandom

integer default: 10000

· Number of random numbers.

2.56 nmlTrajectory

The namelist nmlTrajectory contains variables that control the output of a trajectory.

- · Variables:
 - iskip
 - rmin
 - rmax

iskip

integer default: 10

· Write every trajectory data for every iskip timestep.

rmin

```
real(1:3) default: -boxlen2(1:3)
```

· Lower, left, and front corner of box bounding the particles.

rmax

```
real(1:3) default: boxlen2(1:3)
```

• Upper, right, and back corner of box bounding the particles.

2.57 nmlSubStructureDF

The namelist nmlSustructureDF contains variables that control the calculation of substructures.

type	label	quantity
1	rgsub	radius of gyration of substructure
2	rden	radial number density
3	rrden	reduced radial number density
4	com-sub	distance of com of chain to center of mass of substructure
5	rg-sub	average rg of a chain a a function of the distance to the center of mass of substruc-
		ture
6	z(r)	sum of all charges as a function of the distance to the center of mass of substruc-
		ture
7	zcum(r)	cumulative charge as a function of the distance to the center of mass of substruc-
		ture
8	alpha	global degree of ionization distribution
9	a(r)	reduced radial degree of ionization

- · Variables:
 - vtype
 - Iptinsub

vtype

static1D_var(logical, real, real, integer, logical, character, real)(1:3)

- Flag for engagement, lower end, upper end, number of bins, flag for normalization, title, and number of variable of vtype.
- Min: /0.0/ Max: /100.0/

Iptinsub

logical(1:npt)

• . true . : mass of particles of type ipt are used to evaluate mass center of substructure.

2.58 nmlNetworkDF

The namelist nmlNetworkDF contains variables that control the calculation of network distribution functions. Distribution functions are calculated for each network. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	rg	radius of gyration
2	asph	asphericity
3	alpha	degree of ionization

- · Variables:
 - vtype

vtype

static1D_var(logical, real, real, integer, logical, character, real)(1:3)

- · Flag for engagement, lower end, upper end, number of bins. Other flags are not used.
- Min: /0.0,0.0,0.0/ Max: /100.0,1.0,1.0/

2.59 nmlNetworkRadialDF

The namelist nmlNetworkRadialDF contains variables that control the calculation of radial network distribution functions. Distribution functions are calculated for each network. Any combination of the types of radial distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	rpart(r)	radial particle number distribution
2	rdens(r)	radial particle density distribution
3	rgchain(r)	radial chain radius of gyration
4	q(r)	radial sum of all charges
5	qcum(r)	cumulative radial sum of all charges
6	alpha(r)	reduced degree of ionization
7	rchain(r)	radial chain number distribution

- · Variables:
 - vtype

CHAPTER 2. INPUT 2.60. NMLDYNAMIC

vtype

```
static1D_var(logical, real, real, integer, logical, character, real)(1:7)
```

• Flag for engagement, lower end, upper end, number of bins. Other flags are not used.

• Min: /0.0,0.0,0.0/ Max: /100.0,100.0,100.0/

2.60 nmlDynamic

- The namelist nmlDynamic contains variables that control the dynamic analyses. The detailed description of the use of stochastic data and evaluation of the correlation functions are given by the the data structures sf_var and cf_input_var, see chapter 8.
- · Variables:
 - sfnplow
 - sfnpupp
 - Imsd
 - lorix
 - loriy
 - loriz
 - Iliv
 - lanv
 - Ifor
 - Itor
 - lidm
 - lutot
 - itestdyn

sfnplow

integer default: 1

· Lower id of particle to be used in the dynamic analysis.

sfnpupp

integer default: np

• Upper id of particle to be used in the dynamic analysis.

CHAPTER 2. INPUT 2.60. NMLDYNAMIC

Imsd

logical **default**: .false.

• .true.: Mean square displacements are calculated. Further specification is given in namelist nmlMSD.

· .false.: Nothing.

lorix

logical default: .false.

•

logical default: .false.

- .true.: Orientational tcf for the molecular x'-axis is calculated. Further specification is given in namelist nmlOriXTCF.
- · .false.: Nothing.

loriy

logical **default**: .false.

- .true.: Orientational tcf for the molecular y'-axis is calculated. Further specification is given in namelist nmlOriYTCF.
- .false.: Nothing.

Ioriz

logical default: .false.

- .true.: Orientational tcf for the molecular z'-axis is calculated. Further specification is given in namelist nmlOriZTCF.
- · .false.: Nothing.

lliv

logical default: .false.

- . true.: Velocity tcfs for each molecular axis (molecular frame) and total velocity tcf are calculated. Further specification is given in namelist nmlLinVelCF.
- · .false.: Nothing.

CHAPTER 2. INPUT 2.60. NMLDYNAMIC

lanv

logical **default:** .false.

• .true.: Angular velocity tcfs for each molecular axis are calculated (molecular frame). Further specification is given in namelist nmlAnvVelCF.

· .false.: Nothing.

Ifor

logical default: .false.

• . true.: Force tcfs for each molecular axis and total force tcf are calculated (box frame). Further specification is given in namelist nmlForCF.

· .false.: Nothing.

Itor

logical default: .false.

• . true.: Torque tcfs for each molecular axis and total torque tcf are calculated (box frame). Further specification is given in namelist nmlTorCF.

· .false.: Nothing.

lidm

logical default: .false.

- .true.: Induced dipole moment tcfs for each molecular axis and total induced dipole moment tcf are calculated (molecular frame). Further specification is given in namelist nmIIDPMCF.
- .false.: Nothing.

lutot

logical default: .false.

- . true .: Total energy tcf are calculated. Further specification is given in namelist nmlUtotCF.
- .false.: Nothing.

itestdyn

integer default: 0

· Flag for test output.

CHAPTER 2. INPUT 2.61. NMLMSD

2.61 nmlMSD

The namelist nmlMSD contains variables that control the calculation of the mean square displacement.

```
· Variables:
```

- sf
- cfin

sf

```
sf_var(int, int, int, int, int) default: sfnplow, sfnpupp, 3, 1, -
```

cfin

```
cf_input_var(int, int, int, logical, logical, logical) default: -, -, -, 1, -, .false., .false.
```

2.62 nmlOriXTCF, nmlOriYTCF and nmlOriZTCF

- The namelists nmlOriXTCF, nmlOriYTCF, and nmlOriZTCF contain variables which control the calculation of the three orientational tcf:s of the molecular axes.
- · Variables:
 - sf
 - cfin

sf

```
sf_var(int, int, int, int, int) default: sfnplow, sfnpupp, 3, 3, -
```

cfin

```
cf_input_var(int, int, int, int, logical, logical, logical) default: -, -, -, 1, -, -,
-
```

2.63 nmlLinVelTCF and nmlAngVelTCF

The namelists nmlLinVelTCF and nmlAngVelTCF contain variables that control the calculation of the velocity and angular velocity, respectively, tcf:s for the molecular axes.

- · Variables:
 - sf
 - cfin

sf

```
sf_var(int, int, int, int, int) default: sfnplow, sfnpupp, 3, 1, -
```

cfin

```
cf_input_var(int, int, int, int, logical, logical, logical) default: -, -, -, 1, -, -,
-
```

2.64 nmlForTCF and nmlTorTCF

The namelists nmlForTCF and nmlTorTCF contain variables that control the calculation of the force and torque, respectively, tcf:s about the box axes.

- · Variables:
 - sf
 - cfin

sf

```
sf_var(int, int, int, int, int) default: sfnplow, sfnpupp, 3, 1, -
```

cfin

```
 \texttt{cf\_input\_var(int, int, int, logical, logical, logical)} \ \textbf{default:-,-,-,-,-,-,-,-,-,--} \\ \\
```

2.65 nmlIDMTCF

The namelist nmlIDMTCF contains variables that control the calculation of the induced dipole moment tcf for the molecular axes.

- · Variables:
 - sf
 - cfin

sf

```
sf_var(int, int, int, int, int) default: sfnplow, sfnpupp, 3, 1, -
```

CHAPTER 2. INPUT 2.66. NMLUTOTTCF

cfin

```
cf_input_var(int, int, int, int, logical, logical, logical) default: -, -, -, 1, -, -,
-
```

2.66 nmlUtotTCF

The namelists nmlUtotTCF contain variables that control the calculation of tcf of the total potential energy.

- · Variables:
 - sf
 - cfin

sf

```
sf_var(int, int, int, int, int) default: 1, 1, 1, 1, -
```

cfin

```
cf_input_var(int, int, int, int, logical, logical, logical) default: -, -, -, 1, -, -,
-
```

2.67 nmllmage

The namelist nmllmage contains variables that control the interval of the calls and the calls to image date writing routines.

- · Variables:
 - iimage
 - lvrml
 - lvtf
 - limageuser
 - Igr

iimage

integer **default::** 1

CHAPTER 2. INPUT 2.68. NMLVRML

IvrmI

logical **default:** .false.

• . true . : Coordinate data file for VRML images is prepared. Further specification is given in namelist nmlVRML.

• .false.: No preparation.

lvtf

logical default: .false.

• .true.:oordinate data file for VTF images is prepared. Further specification is given in namelist nmlVTF.

• .false.: No preparation.

limageuser

logical default: .false.

- . true.: Call of ImageUser, driver for user-provided image data writing routines.
- .false.: No such call.

lgr

logical default: .false.

- .true.: coloring is following the group assignment.
- .false.: coloring is following the atom types.

2.68 nmIVRML

The namelist nmlVRML contains variables that control the preparation of coordinate data files for VRML drawing.

- · Variables:
 - txfile
 - txwhen
 - atsize
 - rgbcolor
 - blmax
 - bondr
 - tximage

CHAPTER 2. INPUT 2.68. NMLVRML

txfile

character(9) default: "merged"

- · Controls the generation of Vrml files.
- 'merged': Coordinates for each frame are merged into one Vrml file separated by a wrapper.
- 'separated': Coordinates are written on separated files.

txwhen

```
character(12) default: "after_run"
```

- · Controls the frequency of generation of Vrml files.
- 'after_run': After the run.
- 'after_macro': After each macrostep.
- 'after_iimage': At an interval of iimage steps (iimage is given in nmllmage).

atsize

```
real(1:nat) default: radat (1:nat)
```

· Size of the atoms.

rgbcolor

real(3,1:nat) **default:** some values

• Each triple defines the RGB-color of the atoms.

blmax

real default: 1.5

• Maximal separation between two atoms for drawing a bond between them.

bondr

real default: 0.3

· Radius of the bonds.

CHAPTER 2. INPUT 2.69. NMLVTF

tximage

character(20)(1:3) default: ['frame',' ',' ']

· Controls additional aspects and features of the snapshots

- (1)"frame': Make a frame around the box.
- (2)'one_plane': Draw a plane at -box(3)/2.
- (2)'two_plane': Draw planes at -box(3)/2 and box(3)/2.
- (3)'undopbc': Periodic boundary conditions are not applied when drawing chains.
- (3)'nopbc'': Same as 'undopbc'.

2.69 nmIVTF

The namelist nmIVTF contains variables that control the preparation of coordinate data files for VTF drawing.

- · Variables:
 - txfile
 - txwhen
 - tximage
 - atsize
 - rgbcolor
 - blmax
 - bondr
 - bondres
 - sphres
 - Iframezero

txfile

character(5) default: "merged"

- · Controls the generation of Vtf files. Options are:
- 'split': Coordinates are written in separated files.
- 'merge': All frames are written in one single vtf file.

CHAPTER 2. INPUT 2.69. NMLVTF

txwhen

```
character(12) default: "after_run"
```

· Controls the frequency of generation of Vrml files.

```
• 'after_run': After the run.
```

- 'after_macro': After each macrostep.
- 'after_iimage': At an interval of iimage steps (iimage is given in nmllmage).

tximage

character(20)(1:3)

- · Controls additional aspects and features of the snapshot. Options are
- (1) "frame': Make a frame around the box
- (2)'undopbc': Undo periodic boundary conditions for bonded and cross-linked systems
- (3)'center': Move center of mass of all particles to origin of the simulation cell
- (3)'xycenter'`: Move center of mass of all particles to origin of the simulation cell (consider x- and y-direction only).

atsize

```
real(1:nat) default: radat (1:nat)
```

· Size of the atoms.

rgbcolor

```
real(3,1:nat) default: some values
```

· Each triple defines the RGB-color of the atoms.

blmax

real default: 0.0

· Maximal separation between two atoms for drawing a bond between them.

bondr

real default: 0.3

· Radius of the bonds.

CHAPTER 2. INPUT 2.70. NMLMIXED

bondres

real default: 12.0

· Bond resolution: Number of prisms bonds are being drawn of

sphres

real default: 12.0

· Atom resolution: Number of prisms atoms are being drawn of

Iframezero

logical default: .true.

- .true.: Frame of the initial configuration is being stored.
- .false.: Frame of the initial configurations is not stored.
- Iframezero = .true. does not work with Igr = .true.

2.70 nmlMixed

The namelist nmlMolmix contains variables that control the choice of miscellaneous calculations In all cases two particles are involved, and their types are specified by txpt1 and txpt2.

- · Variables:
 - mode
 - txpt1
 - txpt2
 - coord1
 - coord2
 - ip

CHAPTER 2. INPUT 2.70. NMLMIXED

mode

integer(1:10) default: 0

Select type of calculation. Several calculations, at most 10, of either same or different types may
be performed by listing the corresponding values of mixmode. The correct number and order of
namelists have to follow namelist nmlMolmix.

- 1: Potential energy curve is generated. Further specification is given in namelist nmlMolmix1.
- 2: Potential energies on a lattice is calculated. Further specification is given in namelist nml

 Molmix2.
- 3: The global potential energy minimum is calculated. Further specification is given in namelist nmlMolmix3.
- 4: Random coordinates are generated, and corresponding potential energies are calculated. Further specification is given in namelist nmlMolmix4.
- 5: The second virial coefficient is calculated. Further specification is given in namelist nmlMolmix5.
- 6: Orientational averaged potential energy is calculated. Further specification is given in namelist nmlMolmix6.

txpt1

character(10)

· Particle type of particle one.

txpt2

character(10)

· Particle type of particle two.

coord1

real(1:12)

• Initial coordinate of particle of type txpt1 in the sequence ro(1), ro(2), ro(3) ori(1,1), ori(1,2), ..., ori(3,3). ro is the center-of-mass location. ori(1,1:3) is projection of the particle frame axis z' on the x, y, and z-lab axes, and similarly with ori(2,1:3) and ori(3,1:3).

coord2

real(1:12)

As for coord1 but for particle of type txpt2.

CHAPTER 2. INPUT 2.71. NMLMIXED1

ip

integer

• Particle to be moved or which coordinates should be integrated, either 1 or 2.

2.71 nmlMixed1

The namelist nmlMixed1 contains variables that control the generation of the potential energy curve.

- · Variables:
 - dxx
 - dyy
 - dzz
 - iaxis
 - dr
 - mstep
 - umax

dxx

real

• Translational step in x-direction.

dyy

real

• Translational step in y-direction.

dzz

real

• Translational step in z-direction.

iaxis

integer

- 1: Rotation about the x-axis.
- 2: Rotation about the y-axis.
- 3: Rotation about the z-axis.

CHAPTER 2. INPUT 2.72. NMLMIXED2

dr

real

· Rotational step (in degrees).

mstep

integer

· Maximum number of steps.

umax

real

• Upper potential energy limit where the generation of the potential energy curve is stopped.

2.72 nmlMixed2

The namelist nmlMixed2 contains variables that control the calculation of potential energies on a lattice. The energies are calculated on a 2D grid in position space and at each point the potential energy is minimized as a function of the orientation.

- · Variables:
 - itmax
 - udel
 - dr
 - iaxis
 - rdist
 - rlow1
 - rupp1
 - nrdiv1
 - rlow2
 - rupp2
 - nrdiv2

itmax

integer

• Maximum number of iterations of the orientation optimization.

CHAPTER 2. INPUT 2.72. NMLMIXED2

udel

real

· Potential energy tolerance for the iteration of the orientation optimization.

dr

real

• Rotation step angle for the iteration of the orientation optimization.

iaxis

integer

- 1: Grid in the yz-plane.
- 2: Grid in the zx-plane.
- 3: Grid in the xy-plane.

rdist

real

• Value of the constant coordinate, either x, y, or z depending on iaxis.

rlow1

real

• Lower value of either the y, z, or x-coordinate.

rupp1

real

• Upper value of either the y, z, or x-coordinate.

nrdiv1

integer

• Number of grid points in either the y, z, or x-coordinate.

CHAPTER 2. INPUT 2.73. NMLMIXED3

rlow2

real

• Lower value of either the y, z, or x-coordinate.

rupp2

real

• Upper value of either the y, z, or x-coordinate.

nrdiv2

integer

• Number of grid points in either the y, z, or x-coordinate.

2.73 nmlMixed3

The namelist nmlMixed3 contains variables that control the calculation of the global potential energy minimum.

- · Variables:
 - itmax
 - udel
 - dxx
 - dyy
 - dzz
 - dr

itmax

integer

· Maximum number of iterations.

udel

real

• Potential energy tolerance for the iteration.

CHAPTER 2. INPUT 2.74. NMLMIXED4

dxx

real

• Translational step in x-direction.

dyy

real

• Translational step in y-direction.

dzz

real

• Translational step in z-direction.

dr

real

• Rotational step (in degrees).

2.74 nmlMixed4

The namelist nmlMixed4 contains variables that control the calculation of random coordinates. The random coordinates are restricted to a volume, given in spherical polar coordinates, and to a potential energy range.

- · Variables:
 - iwr
 - rlow
 - rupp
 - thlow
 - thupp
 - filow
 - fiupp
 - ulow
 - uupp
 - **–** no
 - ntry

CHAPTER 2. INPUT 2.74. NMLMIXED4

iwr

integer

• 0: Evenly distributed random numbers in the r direction.

• >0: Higher probability for larger r coordinates.

rlow

real

· Lower limit in the r-direction.

rupp

real

• Upper limit in the r-direction.

thlow

real

· Lower limit in the theta-direction (in degrees).

thupp

real

• Upper limit in the theta-direction (in degrees).

filow

real

• Lower limit in the phi-direction (in degrees).

fiupp

real

• Upper limit in the phi-direction (in degrees).

ulow

real

• Lower limit of potential energy.

CHAPTER 2. INPUT 2.75. NMLMIXED5

uupp

real

· Upper limit of potential energy.

no

integer

· Number of desired random coordinates.

ntry

integer

· Maximum number of attempts.

2.75 nmlMixed5

The namelist nmlMixed5 contains variables that control the calculation of the second virial coefficient. The integration is done by generation of random coordinates restricted to a specified volume given in spherical polar coordinates. The uncertainty is given by one standard deviation obtained from ten sets of random coordinates.

- · Variables:
 - iwr
 - rlow
 - rupp
 - thlow
 - thupp
 - filow
 - fiupp
 - no
 - nblock

iwr

integer

- 0: Evenly distributed random numbers in the r direction.
- >0: Higher probability for larger r coordinates.

CHAPTER 2. INPUT 2.75. NMLMIXED5

rlow

real

· Lower limit in the r-direction.

rupp

real

• Upper limit in the r-direction.

thlow

real

• Lower limit in the theta-direction (in degrees).

thupp

real

• Upper limit in the theta-direction (in degrees).

filow

real

• Lower limit in the phi-direction (in degrees).

fiupp

real

• Upper limit in the phi-direction (in degrees).

no

integer

• Number of desired random coordinates in one set.

nblock

integer **default:** 10

CHAPTER 2. INPUT 2.76. NMLMIXED6

2.76 nmlMixed6

The namelist nmlMixed6 contains variables that control the calculation of orientation averaged potential energy at a given separation. The orientational integration is done by generation of random coordinates restricted to a specified domain, given in spherical polar coordinates. The potential energy distribution of the potential energy is plotted.

- · Variables:
 - rlow
 - rupp
 - thlow
 - thupp
 - filow
 - fiupp
 - ulow
 - uupp
 - no
 - nbin

rlow

real

• Lower limit in the r-direction.

rupp

real

• Upper limit in the r-direction.

thlow

real

• Lower limit in the theta-direction (in degrees).

thupp

real

• Upper limit in the theta-direction (in degrees).

CHAPTER 2. INPUT 2.76. NMLMIXED6

filow

real

• Lower limit in the phi-direction (in degrees).

fiupp

real

• Upper limit in the phi-direction (in degrees).

ulow

real

• Lower value of the potential energy distribution function.

uupp

real

• Upper value of the potential energy distribution function.

no

integer

· Number of random orientations.

nbin

integer

• Number of bins used to sample the distribution functions.

2.77 nmlComplexation

The namelist nmlComplexation contains variables that can be used for the analysis of interparticle complexation.

- · Variables:
 - rcut_complexation
 - IClusterDF
 - IComplexFraction
 - IComplexDist
 - ISegmentComplex

rcut_complexation

real default: 0.0

· Cut-off distance for complexation.

IClusterDF

logical default: .false.

- . true.: The cluster size distribution will be calculated.
- · .false.: Nothing.

IComplexFraction

logical default: .false.

- . true.: The fraction of complexation will be calculated.
- · .false.: Nothing.

IComplexDist

logical default: .false.

· documentation missing

ISegmentComplex

logical default: .false.

· documentation missing

2.78 nmlComplexDist

The namelist nmlComplexDist contains variables that control the calculation of complexation distribution functions. Any combination of the types of distribution functions listed below may be selected through vtype%l.

type	label	quantity
1	w	fraction of complexation

- · Variables:
 - vtype

vtype

static1D_var(logical, real, real, integer, logical, character, real)

- Flag for engagement, lower end, upper end, number of bins. Other flags are not used.
- Min: /-0.005,-0.005,-0.005/ Max: /1,005,1.005,1.005/

3 User-provided routines

The file moluser.F90 contains routines that are conditionally called and which might be modified by the user. This allows the software to be extended without having to modify the core of it.

The existing code in the distributed file moluser.F90 reflects the current use of it. The procedure to insert a new routine is as follows:

- At the appropriate position in driver moluser, extend the test of txUser and write the appropriate call to the new subroutine
- Add your new subroutine in moluser.F90
- Compile

The file moluser.F90 contains six drivers and several routines specifying different types of tasks. The six drivers are PotentialUser, SetParticleUser, DumpUser, GroupUser, StaticUser, and ImageUser.

3.1 PotentialUser

Routine PotentialUser is the driver of user-provided routines for two-body potentials. It calls routines specifying various potentials on the basis of the value of txpot(iptjpt), which is specified in namelist nml \leftarrow Potential. The procedure to install a new user-provided potential is as follows:

- Extend the test of txpot(iptjpt), and write the appropriate call to a new routine in routine PotentialUser.
- Add your new routine to file moluser.F90.

3.2 SetParticleUser

Routine SetParticleUser is the driver of user-provided routines for generating a start configuration. It calls routines specifying types of initial particle positions on the basis of the value of txsetconf(ipt), which is specified in namelist nmlSetConfiguration. The procedure to install a new user-provided start configuration is as follows:

- Extend the test of txsetconf(ipt) and write the appropriate call to a new routine in the routine Set
 — ParticleUser.
- Add your new routine to file moluser.F90.

3.3 DumpUser

Routine DumpUser is the driver of user-provided routines for dumping data. It calls routines specifying different types of dumping. Generally, all these user-provided routines are called if variable ldumpuser, specified in namelist nmlDump, is true. The procedure to install a new dump analysis is as follows:

- Write the appropriate call to the new routine in the routine DumpUser.
- Add your new routine to file moluser.F90

3.4 GroupUser

Routine GroupUser is the driver of user-provided routines for particle group division. It calls routines specifying different division of particles into groups on the basis of the values of variables ref or field [later stored in txappl(m) (m = 1 for ref and m = 2 for field particles)], which are specified in namelist nmlGroup . The procedure to install a new group division procedure is as follows:

- Extend the test of txappl(m) (m = 1 for ref and m = 2 for field particles) and write the appropriate call to the new routine in driver GroupUser.
- Add your new routine to the file moluser. F90.

Follow closely the pattern of one of the other user-provided routines for particle group division. After the statement 'if (iStage = iWriteInput) then', variables ngr , igrpt , and txgrr have to be specified. ngr denotes the number of groups, igrpt the order number of the particle type of particles that might belong to this group, and txgrr a text label of the group. After 'if (iStage = iSimStep) then', the variable igr (within the particle loop) has to be assigned the order number of the group to which the particle should belong (zero if the particle does not belong to any group). The last two lines of the particle loop should be the same as in the other cases.

3.5 StaticUser

Routine StaticUser is the driver of user-provided routines for static analysis. It calls routines specifying different types of analysis. Generally, all these user-provided routines are called if variable Istaticuser, specified in namelist nmlStatic, is true. The procedure to install a new static analysis routine is as follows:

- Write the appropriate call to the new routine in the routine StaticUser.
- Add your new routine to the file moluser. F90.

3.6 ImageUser

Routine ImageUser is the driver of user-provided image data writing routines. It calls routines specifying different types of such routines. Generally, all these user-provided routines are called if variable limageuser, specified in namelist nmlImage, is true. The procedure to install a new image data writing routine is as follows:

- Write the appropriate call to the new routine in the routine ImageUser.
- Add your new routine to the file moluser.F90.

4 Distributed files

This chapter contains a complete list of the distributed source files.

Filename	Description
molsim.F90	Routines specific for MOLSIM
particle.F90	Routines for particles
potential.F90	Routines for potential and force tables
coordinate.F90	Routines for coordinates including start configuration
md.F90	Routines for MD integration
mc.F90	Routines for MC integration
bd.F90	Routines for BD integration
nlist.F90	Routines for neighbour lists
energy.F90	Routines calculating energy and forces
dump.F90	Routines for dumping and reading variables
group.F90	Routines for particle group division
static.F90	Routines for static analysis
dynamic.F90	Routines for dynamic analysis
image.F90	Routines for preparing image data files
mixed.F90	Routines specific for mixed tasks
molaux.F90	Auxiliary routines
mesh.F90	Routines for making meshes
statistics.F90	Routines for statistical analyses
mollib.F90	General library routines
parallel.F90	Parallel interface between MOLSIM and MPI
moluser.F90	User-provided routines
molsim.lib	Potential library

Command	Description
molsim_ser	Command file for execution of serial version of MOLSIM
molsim_par	Command file for execution of parallel version of MOLSIM
makefile	Makefile for compiling and linking
archive	Command file for filing

5 Subroutines and Functions

This chapter lists the routines and functions in the MOLSIM software. The routines are listed according to the Fortran files they are located in and in the order they appear in the files.

- mol.F90
- molsim.90
- particle.F90
- potential.F90
- · coordinate.F90
- md.F90
- mc.F90
- bd.F90
- nlist.F90
- energy.F90
- denergy.F90
- dump.F90
- group.F90
- static.F90
- dynamic.F90
- image.F90
- mixed.F90
- molaux.F90
- mesh.F90
- statistics.F90
- mollib.F90
- parallel.F90
- moluser.F90
- sso.F90
- gc.f90
- celllist.F90

5.1 mol.F90

MolModule main module for the Molsim software

5.2 molsim.90

Molsimdriver driver of the Molsim program

IOMolsim(iStage) perform mixed tasks, mainly i/o

IOSystem perform i/o on general system variables

IOScale perform i/o on scaling variables

IOCnf perform i/o on the configuration file

Control Aver caluclate control quantities

MDAver calculate averages of md scific variables

DispAver calculate averages of displacements

OriOrderAver calculate averages of orientational order

PosOriAver calculate averages of positions and orientation

MainAver calculate various quantities

ThermoAver calculate averages of thermodynamic quantities

ChargeAver calculate averages of related to weak charges

IndDipMomAver calculate averages of induced dipole moment

ChainAver calculate averages of chain quantities

NetworkAver calculate averages of network quantities

HierarchicalAver calculate averages of chain quantities

ThermoInteg handle thermodynamic integration

DistFunc calculate distribution functions

PressureContact calculate contract contribution to the reduced pressure

GContact calculate rdf contract value

TestSimulation write test output

5.3 particle.F90

ParticleModule module for particle

Particle(iStage) particle variables

SetObjectParam1 setobject parameters

SetObjectParam2 set object parameters

PAxesSystem transform from input to principal axes system

5.4 potential.F90

Potential Module module for potential

Potential Driver driver of potential routines

IOPotTwoBody perform i/o on potential variables for two body interactions

PotTwoBodyTab1 set tables for two-body interactions

PotTwoBodyTab2 set tables for two-body interactions for particle pair iptjpt

PotTwoBodyTab3 set table for two-body interaction iatjat

SetUBuffer *transform to r^2 , calculate coefficients, and store them in a table*

CheckUBuffer check the accuracy of the tabulation of one interval

PotDefault default potential

Pot_1_6_12 1, 6, 12-potential

MCY MCY water potential

SphEwaldTrunc spherical ewald truncated potential

Ramp ramp potential (with soft slope change)

SquareWell Square-well potential (with a soft rise)

AsakuraOosawa Asakura-Oosawa depletion potential

LekkerkerKerTuinier Lekkerkerker-Tuinier depletion potential

calc_LT_depletion calculate Lekkerkerker-Tuinier depletion potential (in kT units)

Nemo nemo potentials

NemoType1 nemo potential of type 1

NemoType2 nemo potential of type 2

NemoType3 nemo potential of type 3

NemoType4 nemo potential of type 4

NemoType5 nemo potential of type 5

NemoType6 nemo potential of type 6

NemoType7 nemo potential of type 7

WriteUBuffer write the content of ubuf for pair iatjat

TestUBuffer calculate the accuracy of the table for pair iatjat

PlotPotTwoBodyTab plot two-body potential

TestPotTwoBodyTab test of utwobodytab and utwobody

CalcScrCUfac *calculate exp(r1/scrlen) and two related derivatives*

CalcGCDUfac calculate erfc(1/sqrt(1/a1**2+1/a2**2)*r1) and two related derivatives

CalcEwaldUfac calculate erfc(ualpha*r1) and two related derivatives

CalcRFUfac calculate modified interactions according to nymand and linse, eqs(55)

EwaldErrorUReal estimate error of real space potential energy

EwaldErrorURec estimate error of reciprocal space potential energy

TestEwald examine truncation error of ewald summation (energies in kT)

TestEwaldStd examine truncation error of standard ewald summation

TestEwaldSPM examine truncation error of the reciprocal space; smooth particle mesh ewald summation

SetImageSph Setup of image charges and dipoles according to Friedman

IOPotChain perform i/o on bond and angle interaction variables

ChainTab call routines setting up potential tables for chains

BondLengthTab make and use of lookup table to get random bond lengths

BondAngleTab make and use of lookup table to get random bond angles

IOPotExternal perform i/o on external potential variables

IOPolarizationIter perform i/o on polarization iteration variables

5.5 coordinate.F90

CoordinateModule module for coordinate

Coordinate handle coordinates and velocities

SetChargeWeakChargeCase initiate charge of for the weak-charge case

SetConfiguration generate a start configuration

SetLattice generate a lattice configuration

SetOrigin generate a point in the center of the unit cell

SetSq2D generate a 2d square lattice

SetHex2D generate a 2d hexagonal lattice

SetPC generate a primitive cubic lattice

SetBCC generate a body-centered cubic lattice

SetFCC generate a face-centered cubic lattice

SetSM2 generate a sm2 lattice

SetDiamond generate a cubic diamond lattice

SetH2O generate a cubic lattice, im3m (ice viii)

SetN2 generate a cubic lattice, pa3 (solid n2)

SetBenzene *generate a cubic lattice, pbca (solid benzene)*

SetRandom generate random positions and orientations

SetRandomFixOri generate random positions and fixed orientations

SetChainLine generate a linear configuration for chain particles (only for one chain)

SetChainCircle generate a circular configuration for chain particles (only for one chain)

SetChainRandom generate random positions and orientations for chain particles

SetChainRandomIntOri generate random positions and internally fixed orientations for chain particles

SetSphBrush generate a configuration for a spherical brush

SetPlanarBrush generate a configuration for a planar brush

SetHierarchical generate a configuration for hierarchical polymers

SetPeriodicNetwork generate a periodic network

SetNetwork generate a nonperiodic network

SetCoreShell generate a configuration with particles located in a spherical shell

SetPartPosRandom *generate a random particle position*

SetPartPosRandomN generate a random position at a given distance to a previously set particle

CheckPartOutsideBox check if a particle is outside the box

CheckTooFoldedChain check if a too folded chain

MakeCrossLink make crosslinks between particles labeled clbeg and particles labeled clend on the basis of their separation

TestCoordinate write test coordinates

StoreInteger store integer

5.6 md.F90

MDModule module for md

MDDriver molecular dynamics driver

IOMD perform i/o on molecular dynamics variables

VelVer perform one step according to the velocity form of the Verlet algorithm

Gear perform one step according to the Gear algorithm

GetLinAcc calculate new linear acceleration

GetAngAcc calculate new quaternion accelerations

ScaleLength scale lengths through a pressure coupling to an external bath

ScaleVel scale velocities and calculate new kinetic energies

ScaleLinVel scale linear velocities through a thermal coupling to an external bath

ScaleAngVel scale angular velocities through a thermal coupling to an external bath

SetVel set linear and angular velocities

SetLinVel set linear velocities according to the maxwell distribution

SetAngVel set angular velocities according to the maxwell distribution

SetZeroMom set zero linear and angular moments

SetZeroLinMom set zero linear moments

SetZeroAngMom set zero angular moments

GetLinMom calculate total linear moment

GetAngMom calculate total angular moment

WriteLinMom write total linear moment

WriteAngMom write total angular moment

GetKinEnergy calculate total kinetic energies and temperatures

GetTStep return the time step

GetTime return the time

Get|SetVel *return the value of |setve|*

GetIZeroMom *return the value of Izeromom*

TestSimulationMD write MD test output

5.7 mc.F90

MCModule module for mc

MCDriver monte carlo driver

IOMC perform i/o on monte carlo variables

MCPass perform one mc pass (np trial moves)

SPartMove perform one single-particle trial move

SPartCl2Move perform one single-particle + cluster2 trial move

PivotMove perform pivot rotation trial move

ChainMove perform one chain trial move

SlitherMove perform one chain slithering trial move

BrushMove perform one brush trial move

BrushCl2Move perform one brush + cluster2 trial move

HierarchicalMove perform one hierarchical trial move

NetworkMove perform one network trial move

VolChange perform one volume change trial move

NPartChange perform one number of particle change trial move

ChargeChange perform a charge-change trial move

SetPartPosRandomMC generate a random position

MCAllDriver mcall driver

IOMCAII performing i/o on monte carlo all variables

MCAllPass perform one mc pass (simultaneous move of all particles)

ClusterMember calculate secondary cluster members using Verlet neighbour list

ClusterMemberLList calculate secondary cluster members using linked lists

GetRandomTrialPos calculate translational displacement and trial position

GetRandomTrialOri calculate a trail orientation

GetFixedTrialOri get fixed trial orientation

GetRotatedTrialOri get rotated trial orientation

RotateSetPart rigid-body rotation of a set of particles

FizedZCoord no displacement in z-direction

FizedXYCoord no displacement in the xy-plane

FixedChainStart put rotm = ro and drotm to zero if particle is the first segment in a chain

ShiftZDirection center the system at z = 0 as much as possible (only for lbccyl)

CheckPartBCTM check boundary conditions of trial particle coordinate

AddNeighbours adding neighbours

UpdateOri get fixed trial orientation

SetTrialAtomProp set trial atom properties from trial particles properties

CheckAtomBCTM check boundary conditions of trial atom coordinate

Metropolis *Metropolis test*

MCUpdate update energies and coordinates after accepted mc trial move

Restorelptm restore lptm

MCAver calculate means of mc specific variables

MCAllAver calculate means of mc specific variables for mcall

MCWeightIO perform i/o for an umbrella sampling with a distance dependent weight

MCWeight calculate weight based on trial and current position for mcweight

MCWeightInverse calculate the inverse weight based on current position for mcweight

UmbrellalO perform i/o of the umbrella potential sampling

UmbrellaWeight calculate weight for umbrella sampling

UmbrellaBin calculate bin for umbrella potential

UmbrellaUpdate update weight function for updated umbrella potential

MCPmfIO perform i/o of the mc pmf sampling

MCPmfWeight calculate weight for mcpmf

MCPmfBin calculate bin for umbrella potential

MCPmfUpdate update weight function for mc pmf

TestMCMove test output for MC move

5.8 bd.F90

BDDriver Brownian dynamics driver

IOBD perform i/o on Brownian dynamics variables

BDStep perform one bd step (configuration space)

5.9 nlist.F90

NListModule module for neighbour (verlet and liked) list

NListDriver driver of calls for making lists of interacting particles

IONList perform i/o on neighbour list variables

LoadBalance control the calls of LoadBalanceRealSpace and LoadBalanceRecSpace

LoadBalanceRealSpace set iobjmyid for load balancing of single loop with equal load

LoadBalanceRecSpace set vbounds for load balancing of reciprocal space loop over nz and ny

NList control the calls of SetVList, VListAver, SetLList, and LListAver

SetVList set neighbour (verlet och liked) lists

SetVListMD set verlet lists for md

SetVListMDLList set verlet lists for md using linked lists

SetVListMC set verlet lists for mc

SetVListMCLList set verlet lists for mc using linked lists

VListAver calculate statistics involving verlet lists and particle partitioning

TestVList write verlet list and node decomposition test output

SetLList set linked lists

SetNCell set list of cells to be used for a position

LListAver calculate statistics involving linked cell lists and particle partitioning

TestLList write linked list

CalcTotNeighbourPair calculate number of neigbouring pairs

Getdrnlist *return the value of drnlist*

Getnpmyid return the value of npmyid

Getdrnlist return the value of ncellllist

5.10 energy.F90

EnergyModule *module for energy*

UTotal calculate energies, forces, torques, virial, and pressure

UTwoBodyA calculate two-body potential energy; only monoatomic particles

UTwoBodyALList calculate two-body potential energy; only monoatomic particles

UTwoBodyACellList calculate two-body potential energy

UTwoBodyP calculate two-body potential energy; general particles

UEwald calculate potential energy, forces, and virial from charges; k-space

EwaldSetArray calculate eikx, eiky, and eikz arrays used for ewald summation; k-space

EwaldSetArray2d calculate sinkx, coskx, sinky, and cosky arrays used for ewald summation; k-space

UDipole calculate potential energy from charges and dipoles

UDipoleP calculate potential, field, and field gradient from charges and dipoles

UDipoleEwald calculate potential, field, and field gradient from charges and dipoles; k-space

UManyBodyP calculate many-body potential energy; general particles

FieldStat calculate potential and field from charges and static dipoles

FieldStatEwald calculate potential and field from charges and static dipoles; k-space

IterIdm iterate induced dipole moments self-consistently

FieldIdm calculate field from induced dipoles

FieldIdmEwald calculate field from induced dipoles; k-space

FieldTot calculate field, field gradient, and virial from charges and total dipoles

FieldTotEwald calculate field, field gradient, and virial from charges and total dipoles; k-space

UlntraReac calculate intramolecular contribution to energy and force in rf

PackReduceU pack the different energy contributions and perform all reduce

UDipoleSph calculate charge and dipole potential energy using image charge approximation

UDielDis calculate energy for charge and in a system with dielectric discontinuities

UDielDisPlane calculate coulomb energy in a system with dielectric discontinuity at z = 0

UDielDisSph calculate coulomb energy in a system with spherical dielectric discontinuity

ImageIntSph get the image interaction for spherical geometry

UTwoBodyPair calculate two-body potential energy and force between two particles

UBond calculate bond potential energy, forces, and virial

UAngle calculate angle potential energy, forces, and virial

UCrossLink calculate crosslink potential energy, forces, and virial

UExternal calculate external potential energy

UreactionSphere calculate the reaction energy of an ion and a dielectric sphere

UExternalUpdate update arrays for external energy

xCCLM return the spherical harmonics

xPLM return the associate legendre polynomial p(l,m) at x

Longrangecontr calculate long-range contribution of the electrostatic energy

SPMFFTRec make Fourier transformation, reciprocal space operations, and back FFT

EwaldSetup setup for ewald summation

Getnkvec return the value of nkvec

Getnkvec2d return the value of nkvec2d

Gettime ewald return the value of ncut2d

UWeakChargeA calculate two-body potential energy; only monoatomic particles

UWeakChargeP calculate two-body potential energy; general particles

5.11 denergy.F90

DUTotal calculate energy difference between two configurations

DUTwoBody calculate two-body potential energy difference

UTwoBodyANew calculate two-body potential energy for new configuration

UTwoBodyAOId calculate two-body potential energy for old configuration

UTwoBodyANewLList calculate two-body potential energy for new configuration

UTwoBodyAOIdLList calculate two-body potential energy for old configuration

UTwoBodyANewCellList calculate two-body potential energy for old configuration

UTwoBodyAOldCellList calculate two-body potential energy for old configuration

UTwoBodyPNew calculate two-body potential energy for new configuration

UTwoBodyPOld calculate two-body potential energy for old configuration

DUTwoBodyEwald calculate two-body potential energy difference; k-space

DUWeakChargeEwald calculate two-body potential energy difference for titrating systems; k-space

EwaldSetArrayTM calculate eikxtm, eikytm, and eikztm arrays for moving particles

EwaldSetArray2dTM calculate sinkxtm, coskxtm, sinkytm, and sinkytm arrays for moving particles

EwaldUpdateArray update eikx, eiky, eikz, and sumeikr for moving particles

DUDipole calculate potential energy difference from charges and dipoles

DUDipolePNew calculate potential energy from charges and dipoles for new configuration

DUDipolePOId calculate potential energy from charges and dipoles for old configuration

DUDipoleEwald calculate potential energy change; k-space

DUDipoleSph calculate charge and dipole potential energy difference

DUDipoleSphNew calculate charge and dipole potential energy for new configuration

DUDipoleSphOld calculate charge and dipole potential energy for old configuration

DUDielDis calculate coulomb energy in a system with dielectric discontinuities

DUDielDisPlane calculate coulomb energy in a system with dielectric discontinuity at z = 0

DUDielDisSph calculate coulomb energy in a system with a radial dielectric discontinuity

DUBond calculate bond potential energy difference

DUAngle calculate angle potential energy difference

DUCrossLink calculate crosslink potential energy difference

DUExternal calculate external potential difference

UWeakChargeANew calculate two-body potential energy for new configuration

UWeakChargeAOld calculate two-body potential energy for old configuration

UWeakChargePNew calculate two-body potential energy for new configuration

UWeakChargePOld calculate two-body potential energy for old configuration

5.12 dump.F90

DumpModule module for dumping

DumpDriver dump driver

IODump performing i/o on dump variables

DoDump performing dumping matters

5.13 group.F90

Group classify reference and field particles into groups and make average

GroupAll group division: each particle type forms one group

GroupType group division: select particle type forms one group

5.14 static.F90

StaticDriver driver of static analysis routines

SPDF calculate single particle distribution functions

RDF calculate running coordination number or radial distribution function

RDFChain calculate running coordination number or radial distribution function

RDFSph calculate rcn or rdf of particles which positions are projected on a sphere

RDFCond calculate conditional radial distribution function

G3Dist *calculate g*(*r*12,*r*13,*r*23)/*g*(*r*12)

SFDriver calculate partial structure factors

SFPBC calculate partial structure factors

ScatIntens calculate scattering intensity, form factor, and structure factor

SFNoPBC calculate partial structure factors

AngDF calculate angular distribution functions

AngExtDF calculate angular 2d distribution functions with respect to external frame

OriDipDF calculate orientation/dipole distribution function

SphHarAver calculate averages of unnormalized spherical harmonics

RadAngDF calculate radial-angular 2d distribution functions

Kirkwoodgk calculate Kirkwood gk-factor

OriPoIDF calculate orientation polarization distribution functions

NNHB calculate number of nearest neighbours and number of hydrogen bonds

NNDF calculate the nearest neighbour distribution

ChainDF calculate chain distribution functions

ChainTypeDF calculate chain type distribution functions

ChainTypeExtDF calculate chain type distribution function relative external frame

ChainBeadPartContact calculate probability of chain bead-particle contact

LoopTailTrain calculate loop, tail, and tail characteristics for chains

CheckAdsChainSeg check if a chain and its segments are adsorbed

ClusterSD calculate cluster size distribution

ClusterSD1D calculate cluster size distribution

ClusterSD2D calculate 2d cluster size distribution

ZeroSecondMoment calculate zero and second moment for an ionic, neutral system

MultipoleDF calculate multipole moment distribution function and mean square average

CalcMultipole calculate multipole moments of spherical volumes arising from dipoles

EnergyDF calculate energy distribution functions

Widom1 calculate excess chemical potential using Widom's method (neutral set)

Widom2 calculate excess chemical potential using Widom's method (charge integration)

DUWidomx calculate the change of the potential energy of the insertion

UTwoBodyWidom calculate two-body potential between two particles; monoatomic particles

UTwoBodyWidomP calculate two-body potential between two particles, polyatom particles

MeanForce1 calculate mean energies and forces between two particles

MeanForce2 calculate mean energies and forces between two particles

UBondAcrossZ calculate potential energy from bonds across z = 0

PotMeanForce calculate potential of mean force

SurfaceArea calculate surface area exposed by all particles of one type

Crystalformat write atoms in the crystalographic format starting with atoms

Trajectory write the trajectory on flist using every iskip timestep

ExcessAmount calculate excess amount

SubStructureDF calculate distribution functions of properties of a SubStructure

NetworkDF calculate network distribution functions

NetworkRadiaIDF calculate radial distribution functions of properties of networks

5.15 dynamic.F90

DynamicModule module for dynamic

DynamicDriver driver of dynamic analysis routines

MSD mean square displacement

OrixTCF orientation time correlation function of paricle x'-axis

OriyTCF orientation time correlation function of paricle y'-axis

OrizTCF orientation time correlation function of paricle z'-axis

LinVeITCF *linear velocity time correlation function*

AngVeITCF angular velocity time correlation function

ForTCF force time correlation function

TorTCF torque time correlation function

IDMTCF induced dipole moment time correlation function

UtotTCF total potential time correlation function

PrepareDynamic prepare for dynamic analyse

MemoryDynamic allocate memory

CFCalc calculate correlation function

CFCalcSub calculate time correlation function, subroutine

CFWrite write time correlation function

5.16 image.F90

ImageDriver driver of image preparation routines

ImageVRML generate input files for vrml 97 viewers

VRMLSub writes a vrml-97 file

ImageVTF generate vtf file(s) and tcl-script for VMD

UpdateVTFFileName update vtf file name with respect to a given frame iframe

WriteVTFHeader write header of the vtf file

WriteVTFCoordinates writes current atom coordinates to vtf-file

WriteTCLScript TCL: "tool command language"

UndoPBCModule collection of subroutines and functions needed for subroutine UndoPBC

UndoPBC undo periodic boundary conditions using an iterative strategy

MakeBondList make bond list

5.17 mixed.F90

MixedDriver mixed task driver

Mixed1 generate a potential energy curve

Mixed2 calculate potential energies on a 2d lattice

Mixed3 calculate the global potential energy minimum

Mixed4 generate random coordinates

Mixed5 calculate the second virial coefficient

Mixed6 calculate orientational averaged potential energy

LineMove perform one movement, rotation or translation, to a new minimum

NewCoord translate and rotate a particle and update its atom coordinates

RandomPos generate a random position

CopyCoord copy coordinates

5.18 molaux.F90

MolauxModule molaux module

CalcCOM Calculate the center of mass of given coordinates

PBC apply periodic boundary conditions

PBCr2 apply periodic boundary conditions and calculate r**2

PBC2 calculate periodic boundary condition displacement

FileOpen open external files

FileFlush flush external files (contains a system call)

SetBoxParam set box related parameters

SetPartOriRandom set particle orientation matrix corresponding to a random orientation

SetPartOriLab set particle orientation matrix to match lab frame orientation

SetPartOriBond set particle orientation matrix to match bond directions of the particle

SetAtomPos set atom positions in the laboratory frame

SetAtomDipMom set atom dipole moments in the laboratory frame

SetAtomPolTens set atom polarization tensors in the laboratory frame

CalcIndDipMom calculate induced dipole moment: total and particle

CalcSysDipMom calculate system dipole moment from charges, static, and induced dipoles

CalcPartDipMom calculate particle dipole moment from charges, static, and induced dipoles

dvol return the volume between two concentric shells, 4*pi/3 is not included

darea return the area between two concentric spheres, pi is not included

angcos return the angle cosine between two 3d vectors

angle_rad return the angle between two 3d vectors

angle_deg return the angle between two 3d vectors

WritePartAtomProp write particle and atom properties

IWarnHCOverlap warn if hard-core overlap between particle ip and other particles

WarnHCOverlap warn if hard-core overlap among atoms

WarnAtomOutsideBox warn if atoms are outside box

WarnPartOutsideBox warn if particles are outside box

UndoPBCChain undo the periodical boundary conditions for a chain molecule

CalcChainProperty calculate properties of a single chain

CalcNetworkProperty calculate properties of network number inw

Asphericity calculate the asphericity based on the 2:nd moments in the principal frame

CalcLTT calculate loop, tail, and tail characteristics for a single chain

CalcChainPairListGeneral calculate a list of chain pairs based on particle-particle separation

CalcChainComPairListGeneral calculate a list of chain pairs based on com-com separation

CalcPartPairListAll calculate a list of particle pairs based on their separations

CalcPartPairListGeneral calculate a list of particle pairs

CalcPartPairListIpt calculate pair list for particles

Perculation determine if a system of bonded objects is perculated

CorrAnalysis make correlation analysis of a single variable

BlockAverAnalysis read data and calculate average and precision for different block lengths

WriteTrace write trace information

WriteVecAppend write real array at the end of an external unit

EllipsoidOverlap check overlap between two ellipsoids with equal semi-axes

SuperballOverlap superball overlap check

SuperballOverlapOF superball overlap check, return overlap function

SuperballOverlap_NR check for overlap between two superballs

SuperballStatNR superball check statistics, NR overlap method

SuperballStatMesh superball check statistics, mesh overlap check

SuperballAver superball overlap-check average

SuperballDF superball overlap-check distribution functions

5.19 mesh.F90

MeshModule module for mesh

BuildSuperball mesh a superball

BuildDopTree build a DOP tree

BuildNode build a node

TriTri check if two triangles intersect

Cint tries to find interval on the line intersection between the triangle planes

Coplanar check for overlap in coplanar triangles

PointInTriangle check if point in a triangle

Cross2 cross product of two 2d vectors

Cross Cross product of two 3d vectors

Sort2 sort two elements in ascending order

MidPoint find midpoint of two 3d arrays, midpoint may be modified

12i change a logic variable to an integer

findExtrema documentation_missing

union documentation missing

volume documentation_missing

idSort documentation missing

swap documentation_missing

transformation documentation_missing

rotation documentation missing

TestOverlap documentation missing

OverlapMesh translate and rotate a mesh and check overlap of two trees

OverlapTree check overlap of two trees

OverlapDop check overlap between two bounding boxes

OverlapLeaf check overlap of two leafs

5.20 statistics.F90

StatisticsModule module for statistics

SetIBlockAver *set variable lblockaver*

ScalarSample sample scalar quantities

ScalarSampleBlock sample scalar quantities; fixed block length

ScalarSampleExtrap sample scalar quantities; extrapolate to large blocklen

ScalarNorm normalize scalar quantities

ScalarWrite write scalar quantities

Scalar All Reduce make all_reduce of scalar quantities

DistFuncSample sample distribution functions

DistFuncNorm normalize distribution functions

DistFuncHead write heading of distribution functions

DistFuncWrite control the calls of DistFuncShow, DistFuncPlot, and DistFuncList

DistFuncShow list bin numbers, x, y, and dy of distribution functions

DistFuncPlot plot distribution functions

DistFuncList list x, y, and dy values of distribution functions

DistFuncAverValue calculate and write average of distribution functions

DistFuncAverDist calculate average and spread among distribution functions

DistFuncAllReduce make all_reduce of df quantities

DistFunc2DSample sample two-dimensional distribution functions

DistFunc2DNorm normalize two-dimensinal distribution functions

DistFunc2DHead write heading of two-dimensional distribution functions

DistFunc2DShow *list bin numbers and z as well as bin numbers and dz of two-dimensional distribution functions*

DistFunc2DList *list bin numbers and z as well as bin numbers and dz of two-dimensional distribution functions*

5.21 mollib.F90

MollibModule mollib module

Invint return the inverse of an integer

Center center a string

SpaceOut space out a string

ReplaceText Replace text in a string

ErfLocal return the error function, erf(x)

Erf1 return the error function, erf(x)

Erf2 return the error function, erf(x)

GammaLn return the logarithm of the gamma function

PL return the legendre polynomial p(l) at x

PLM return the associate legendre polynomial p(l,m) at x

CCLM return the spherical harmonics

DDJKM return one wigner's rotation matrix element

SetW3J precalculate wigner's 3j-symbols

WW3J return one wigner 3j-symbol

Trap integration by the trapezoidal rule

TrapNe integration by the trapezoidal rule with varying steplength

IntegCirPoints get points and weight for numerical integration over a circle

PolFit calculate least square fit to a polynomial

PolVal calculate the value of the polynomial for a given x-value

MatVecMul multiply a matrix with a vector, c = a*b

VecMatMul multiply a vector with a matrix, c = b*a

MatInv2 matrix inversion of a 2x2 matrix

MatInv3 matrix inversion of a 3x3 matrix

Matinv matrix inversion with solution of linear equations

Diag diagonalize a real matrix and calculate eigenvectors

Eigensort Given the eigenvalues d and eigenvectors v as output from Diag this

Smooth smooth by using spline functions argonne routine e350s

CsEval return the value and the first two derivatives of a spline func

VInter interpolate in a 1d array

LinInter linear interpolation between (x0,y0) and (x1,y1)

CardinalBSpline calculate values and derivatives of the n:th order cardinal B-spline

HeapSort sort an array into ascending order, Heapsort algorithm

HeapSortIndex generate an index array such that vec(index(j)) is ascending, Heapsort algorithm

MakeCluster make clusters

MakeCluster1 make clusters

MakeCluster2 make clusters

CalcClusterMember get members of the cluster to which object iobj belongs to

CarToSph transform a vector from a cartensian to a spherical polar coordinate system

SphToCar transform a vector from a spherical polar to a cartesian coordinate system

CarToStd1 transform a vector from a cartensian coordinate system to standard form

Std1ToCar transform a vector from standard form to a cartensian coordinate system

OriToEuler transforming orientation matrix ori to Euler angles a, b, c

EulerToQua transform Euler angles a, b, c to quaternions

OriToQua transform orientation matrix to quaternions

QuaToOri transform quaternions qua to orientation matrix

AxisAngToOri transform rotation axis and rotation angle to a rotation matrix

LabToPri transform a vector from lab to principal frame

PriToLab transform a vector from principal to laboratory frame

EulerRot rotation of a column vector (x,y,z) using the Euler angles

EulerRotStd rotate tensor of standard form using Euler angles (NOTE CONVENTION NOT CLEAR)

OrthoOri orthogonalize particle frame

CheckOriOrtho check that a matrix is orthogonal

QuaNorm normalize quaternions

AngVelToQuaVel calculate quaternion velocities from quaternions and angular velocities (principal frame)

QuaVelToAngVel calculate angular velocities (principal frame) from quaternions and quaternion velocities

Random return a random number in the range of 0 < ran < 1

GauRandom return a normally distributed stochastic variable

CirRandom return two random numbers x,y on a unit circle

SphRandom return three random numbers x,y,z on a unit sphere

InvFlt return the inverse of a real variable

GetReIDiff calculate relative differences

CpuAdd add and write total cpu time elapsed since start

CpuLeft check to see if the execution should be stopped

CpuTot write total cpu time elapsed since start

SecondsSinceStart return cpu time used since start in seconds

WriteVec write a vector

WriteMat write a matrix

WriteStd write a variable of standard form

WriteFront write a front

WriteHead write a heading

WriteDateTime write date and time

WritelOStat write value of iostat and take appropriate action

Warn write a warning message

Stop write a stop message and stop process

LowerCase change a string to lower case

UpperCase change a string to upper case

SubStr determine the number of substrings of a string

SignMagn get sign and magnitude of a real number

Advance read from an external unit until a string match is found

Plot plot a function

BrentMod get the minimum of a function

KnuthShuffle Shuffle 1-Dimensional List if Integers !Pascal Hebbeker

5.22 parallel.F90

ParallelModule module to be used in routines invoking mpi

par_initialize initialize parallel matter

par_finalize finalize parallel matter

par comm size get number of processes

par_comm_rank get my id and set master and slave

par_barrier barrier synchronisation

par_bc_characters broadcast character variables

par_bc_logicals broadcast logicals

par_bc_logical broadcast logical

par bc ints broadcast integers

par_bc_int broadcast of scalar integer par bc ints8 broadcast integers of kind 8 par bc reals broadcast double precision reals par bc real broadcast double precision real par bc comps broadcast double precision complex variables par_allreduce_logicals perform global logical or and redistribution of logical variables par_allreduce_logical perform global logical or and redistribution of logical variables par_allreduce_ints perform global sum and redistribution of integer variables par allreduce int perform global sum and redistribution of integer variable par allreduce reals perform global sum and redistribution of double precision variables par_allreduce_real perform global sum and redistribution of double precision variable par allreduce comps perform global sum and redistribution of double precision complex variables par max ints find maximum value and redistribute integer variables par min reals find minimum value and redistribute double precision variables par_max_reals find maximum value and redistribute double precision variables par_reduce_reals perform global sum and redistribution of double precision variables par allgather shortints perform a distribution of array of integer(2) par_allgather_ints perform a distribution of array of integer(4) par_scatter_reals scatter double precision data from root to the other processes par_gather_reals gather double precision data from all processes except root to root par handshake hand shaking between master and slaves par timing parallel timing par error error handler for par routines

5.23 moluser.F90

PotentialUser driver of user-provided routines for potentials

SodiumChloride na-na, na-cl, and cl-cl potentials

EwaldSquareWell real space ewald plus square-well potential (with a soft rise)

ElstatScreen potential where the first term is screened if it is a 1/r term

SetConfigurationUser driver of user-provided routines for generating a start configuration

SetCNF generate a configuration by reading ro and ori from the begining of fcnf

SetOneHelix generate a helical conformation

SetCubic2D1Surf generate a cubic configuration with particles at z = -(boxlenz/2 + delta)

SetCubic2D2Surf generate a cubic configuration with particles at z = +-(boxlenz/2 + delta)

SetChainRandomTrans generate a random start configurations for chain particles + chain translation

SetCapsid generate a uniform distribution on the surface of a sphere of rcap+2

SetCapsid 250 generate a random configuration with 250 particles on a spherical surface

SetCapsidRandom generate a random start configurations for paticles of type ipt

SetInsideCapsid generate a random configuration inside a capsid

SetChainInsideCapsid generate a random chain configuration inside a capsid

SetSpool generate a spool-like structure inside a capsid

SetSpoolm generate a multilayered spool-like structure inside a capsid

SetSpoolm1 generate a multilayered spool-like structure inside a capsid

setsheet set bcc 'sheet' or 'tube'

SetRandomCylinderShell generate a random configuration inside a cylinder with a hard cylinder within

SetChainLinePMA generate a line configuration for PMA chain particles

DumpUser driver of user-provided routines for dumping

BBDump dumps three angles and two energies

ChainCOMDump dump center of mass of chains

ChainReeDump dump end-to-end separations of chains

DipMomTotDump dump the total dipole moment

xy_coordinates_Jasper read x,y coordinates

GroupUser driver of user-provided routines for particle group division

GroupBW1 group division for one benzene molecule in water

GroupBW2 group division for one benzene molecule in water

GroupBW3 group division for one benzene molecule in water

GroupBBW1 group division for two benzene molecules in water

GroupBBW2 group division for two benzene molecules in water

GroupBBW3 group division for two benzene molecules in water

GroupBBW4 group division for two benzene molecules in water

GroupSurface group division for particles of type ipt = 2 at surface and in bulk

groupsurface2 group division for particles only at surface

GroupAds1 group division for particles belonging to adsorbed chains

GroupAds layer1 ramp group division for adsorbed (layer 1) particles

GroupNetworkGenerations group division according to generations of non-periodic network

GroupWeakCharge group division according to titratable species - respective division in

StaticUser driver of user-provided routines for static analysis

ScalarDemo1 calculate an average of a scalar quantity

Scalar Demo2 calculate average of scalar quantities

BondOrder calculate an average of a scalar quantity

TabulationQl calculate an average of a scalar quantity

ChainBeadCylContact calculate probability of chain bead-cylinder contact

Min calculate minimum energy configuration of a bcc "sheet" or "tube"

S1 calculate orientational order parameter S1 in all dimensions

S2 calculate orientational order parameter S2 in all dimensions

C2 calculate C2 nematic order parameter of projection on xy-plane

Q calculate nematic order parameter of projection on xy-plane with x

SFPBC2D documentation_missing

SFPBC2DNoAv Does not average over x and y, compatible with boxlength in x = y

PosMeanSph calculate means of positions

MacroionOneSph calculate various df for system of one macroion + ions, sph geometry

MacroionTwoCyl calculate various df for system of two macroions + ions, cyl geometry

MeanElFieldZCyl calculate mean electrostatic field in the z-direction

DomainModule module for domain analysis

DomainIO domain analyse based on Kirkwood gk-factor

CalcDomain calculate dipole domains based on Kirkwoods gk-factor (G Karlstrom)

DomainDominating sample gk for the dominating domain of each configuration

DomainDistFunc Domain distribution functions

UDomain domain-domain dipole-dipole interaction energy

TCFDipDomain sample gk for the dominating domain of each configuration

SCDF calculate single chain distribution functions as a function of z coordinate

AdsRadGyr calculate radius of gyration distribution functions of adsorbed and adsorbing chains

AdsBondOrder calculate bond order distribution function for adsorbed chains

AdsPropDyn write chain adsorption dynamic data on external unit

CalcAvBondOrder calculation of the average bond order among provided chains

CalcBondOrder calculation of the bond order among bonds in chains of type ict

AdsEventDyn write time and occurrence of adsorption events on FUSER

AdsModule module for adsorption module

AdsExam call of AdsExam routines

AdsExam1 generate curvz file using the primary adsorption data

AdsExam2 analysis of primary adsorption data; adsorption events

AdsExam3 analysis of primary adsorption data; adsorption length

AdsExam4 analysis of primary adsorption data; adsorption length

ReadPrimAdsData read user-provided primary adsorption data

WritePrimAdsData write user-provided primary adsorption data

Z DF Slit distribution function based on *z*-coordinates, planar geometry

ElMom electrostatic moment of a particle

XYProjectDF calculate on 'z' = 0 projected normalized density df

SPDF_COMB calculate single particle distribution functions for comb polymers (coarse_grained)

COMB_DF calculate type distribution functions of comb polymer type (coarse model)

COMBAver calculate averages of comb chain quantities

SFPBC_COMB calculate partial structure factors

OCF calculate bond-bond orientational correlation function and where particle i belongs to group igr

OCF_DF calculate type distribution functions of comb polymer type (coarse model)

ChainBeadBeadContact documentation missing

ElPot calculate electrostatic potential as a function of distance from the center of lab frame

ImageUser driver of user-provided routines for generating image input files

JosUser auxillary routine for syncronizing input data for Jos' project

ComplexationModule Module for analysing the Complexation

DoComplexation documentation missing

5.24 sso.F90

SSOModule Module for the SSO simulation

5.25 gc.f90

NPartChange perform one number of particle change trial move

5.26 celllist.F90

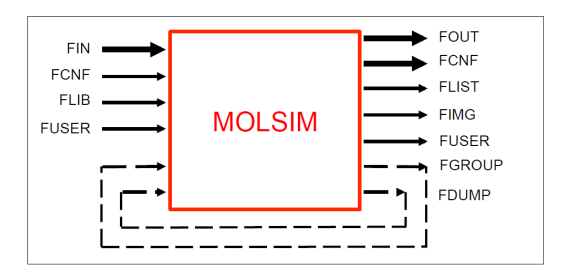
CellListModule a new improved implementation of the linked list

6 External units

A file is specified by its name and type, which are separated by a dot. The file types are used in a predetermined way to describe the use of the file and leaves file name to label the project. Within a project the file name is independent of program.

FPOS, FORI, FLIV, FANV, FFOR, FTOR, and FIDM are collectively called dump files.

File type	Generic name	Content	binary
in	FIN	Input data	
out	FOUT	Output data	
cnf	FCNF	Configuration and average data	binary
lib	FLIB	Potential library	
list	FLIST	List data	
img	FIMG	Image data	
user	FUSER	At the disposal to the user	
group	FGROUP	Particle group data	
pos	FPOS	Box lengths and positions of particles	binary
ori	FORI	Orientations of particles	binary
liv	FLIV	Linear velocities (box frame) of particles	binary
anv	FANV	Angular velocities (principal frame) of particles	binary
for	FFOR	Forces (box frame) of particles	binary
tor	FTOR	Torques (box frame) of particles	binary
idm	FIDM	Induced dipole moments (box frame) of particles	binary



Flow chart of the use of external files. Bold arrows denote files that are always required or generated, while thin arrows denote files that might be required or generated, depending on input variables.

7 Data structures

The following data structures are used for input variables:

bonds etc

```
type bond_var
  real(8) :: k ! force constant
  integer(4) :: p ! power
  real(8) :: eq ! equilibrium value
end type bond_var
```

adsorption conditions

```
type adscond_var
   character(8) :: txplane !'xy-plane' is yet the only option
   character(3) :: txend ! '-','+','-|+' for negative, positive, both ends
   real(8) :: offset ! threshold distance of adsorption for the com of a particle
end type adscond_var
```

stochastic data

```
type sf_var
  integer(4) :: nplow ! lower particle number
  integer(4) :: npupp ! upper particle number
  integer(4) :: ndim ! number of dimension of the stochastic variable
  integer(4) :: fac ! factor describing the separation of stoch. data
  integer(4) :: ngr ! logical flag if for single sf value
end type sf_var
```

correlation functions (input)

```
type cf_input_var
  integer(4) :: nmean ! length (number of values) forming a mean
  integer(4) :: nlevel ! length (number of values) forming a level
  integer(4) :: nolevel ! number of levels
  integer(4) :: legendre ! order of Legendre polynomial (1, 2, or 3)
  logical :: lsvalue ! logical flag if for single sf value
  logical :: lsubmean ! logical flag if subtraction of mean of cf
  logical :: lnorm ! logical flag if normalization of cf
end type cf_input_var
```

correlation functions

```
type cf_var
  integer(4)
                                            ! length (number of values) forming a mean
                :: nmean
                                            ! length (number of values) forming a level
  integer(4)
                :: nlevel
   integer(4)
                  :: ratio
                                                        ! nlevel/nmean
                                                        ! number of levels
   integer(4)
                  :: nolevel
  integer(4)
                :: legendre
                                            ! order of Legendre polynomial (1, 2, or 3)
                                                 ! logical flag if for single sf value
  logical
                :: lsvalue
  logical
               :: lsubmean
                                            ! logical flag if subtraction of mean of cf
                :: lnorm
                                                 ! logical flag if normalisation of cf
  logical
   real(8), allocatable
                                                        ! stochastic function
                            :: sf(:,:,:,:)
  real(8), allocatable :: sf_aver(:,:,:)
                                                 ! stochastic function, local average (of particle)
  real(8), allocatable
                                                    ! stochastic function, gobal mean
                          :: sf_mean(:,:)
                                                        ! time correlation function
  real(8), allocatable
                           :: cf(:,:,:)
  real(8), allocatable
                          :: cf2(:,:,:)
                                                   ! time correlation function squared
  integer(4), allocatable :: Np(:,:,:)
                                                 ! counter of correlation sampling (of particle)
  integer(4), allocatable :: Ngr(:,:,:)
                                                 ! counter of correlation sampling (of group)
  integer(8), allocatable :: Nlev(:,:)
                                                     ! counter of level (of particle)
end type cf_var
```

1D static variables

```
type static1d_var
  logical :: l ! logical flag for engagement
  real(8) :: min ! minimum value
  real(8) :: max ! maximum value
  integer(4) :: nbin ! number of bins
  logical :: lnorm ! logical flag for normalization
  character(10):: label ! title
  real(8) :: nvar ! expanded into nvar variables (not read)
end type static1D_var
```

2D static variables

```
type static2d_var
  logical :: l ! logical flag for engeagement
  real(8) :: min(2) ! minimum value
  real(8) :: max(2) ! maximum value
  integer(4) :: nbin(2) ! number of bins
  logical :: lnorm ! logical flag for normalization
  character(10):: label ! title
  real(8) :: nvar ! expanded into nvar variables (not read)
end type static2D_var
```

potential energy

```
type potenergy_var
real(8), allocatable :: twob(:)
                                    ! two-body contribution (excluding ewald contribution)
  real(8), allocatable :: oneb(:)
                                      ! one-body contribution (dielectric discontinuity)
   real(8)
                                           ! total
                         :: tot
                                 ! reciprocal electrostatic space contribution (UEwald)
  real(8)
                   :: rec
  real(8)
                                        ! static electrostatic contribution (umbodyp)
                       :: stat
                    :: pol
  real(8)
                                   ! polarization electrostatic contribution (umbodyp)
                         :: bond
   real(8)
   real(8)
                         :: angle
   real(8)
                         :: crosslink
                                           ! external contribution
   real(8)
                         :: external
end type potenergy_var
```

chain properties

```
type chainprop_var
   real(8)
                                          ! center of mass
              :: ro(3)
   real(8)
              :: rbb2
                                          ! bead-to-bead distance squared
   real(8)
              :: angle
                                          ! angle between consecutive beads
  real(8)
              :: cos
                                         ! cos(180 - angle between consecutive beads)
   real(8)
              :: ree(3)
                                           ! end-to-end vector
                                          ! end-to-end distance squared
   real(8)
              :: ree2
                                          ! radius of gyration squared
   real(8)
              :: rg2
                                    ! square extention along principal axes (smallest)
  real(8)
            :: rg2s
  real(8)
            :: rg2m
                                     ! square extention along principal axes (middle)
                                     ! square extention along principal axes (largest)
  real(8)
            :: rg2l
                                   ! radius of gyration squared projected on the z-axis
  real(8)
            :: rg2z
                                 ! radius of gyration squared projected on the xy-plane
  real(8)
            :: rg2xy
                                   ! persistence length based on end-to-end separation
            :: lpree
  real(8)
  real(8)
            :: lprg
                                      ! persistence length based on radius of gyration
                                 ! square end-to-end distance / square of radius of gyration
  real(8)
            :: shape
   real(8)
              :: asph
                                           ! asphericity (JCP 100, 636 (1994))
   real(8)
              :: torp
                                          ! toroidicity
end type chainprop_var
```

network properties

```
type networkprop_var
                                          ! center of mass
   real(8)
              :: ro(3)
                                          ! radius of gyration squared
   real(8)
              :: rg2
  real(8)
           :: rg2x
                                  ! radius of gyration squared projected on the x-axis
                                  ! radius of gyration squared projected on the y-axis
  real(8) :: rg2y
  real(8)
                                  ! radius of gyration squared projected on the z-axis
           :: rg2z
                                   ! square extention along principal axes (smallest)
  real(8)
          :: rg2s
  real(8)
           :: rg2m
                                     ! square extention along principal axes (middle)
  real(8)
                                    ! square extention along principal axes (largest)
           :: rg2l
  real(8)
           :: eivr(3,3)
                                     ! normalized eigenvectors of the principal frame
            :: theta(3)
                                  ! angles of axes of largest extension and x-, y-, and z-axes of
  real(8)
       main frame
                                          ! asphericity (JCP 100, 636 (1994))
   real(8)
              :: asph
  real(8)
             :: alpha
                                       ! degree of ionization (for titrating systems)
end type networkprop_var
```

simple averaging

```
type aver_var
  real(8) :: s2 ! summation/averaging over steps
  real(8) :: s1 ! summation/averaging over macrosteps
end type aver_var
```

complexation

```
type cluster_var
  integer(4), allocatable :: ip
  integer(4), allocatable :: np
end type cluster_var
```

blocks in chains

```
type :: block_type
  integer(4) :: pt !particle type
  integer(4) :: np !number of particles
end type block_type
```

scalar quantities

```
type scalar_var
                                                        ! label
   character(40) :: label
                                                        ! normalization factor
   real(8)
                 :: norm
                                                        ! number of macrosteps sampled
   integer(4)
                 :: nsamp1
                                               ! number of values sampled per macrostep
  integer(4)
                :: nsamp2
                                                         ! average of the run
   real(8)
                 :: avs1
  real(8)
                                                     ! precision of average of the run
                 :: avsd
                                                        ! average of a macrostep
   real(8)
                 :: avs2
   real(8)
                 :: fls1
                                                         ! fluctuation of the run
                                                 ! precision of fluctuation of the run
  real(8)
                :: flsd
                 :: fls2
                                                        ! fluctuation of a macrostep
   real(8)
   real(8)
                 :: value
                                                        ! value of a configuration
                                                        ! number of samplings
                 :: nsamp
   integer(4)
  integer(4)
                :: nblocklen
                                                 ! for sampling with variable blocklen
  integer(4)
               :: nblock(mnblocklen)
                                                 ! for sampling with variable blocklen
               :: av_s1(mnblocklen)
                                                 ! for sampling with variable blocklen
  real(8)
  real(8)
               :: av_sd(mnblocklen)
                                                 ! for sampling with variable blocklen
                                                 ! for sampling with variable blocklen
  real(8)
               :: av_s2(mnblocklen)
                                                 ! for sampling with variable blocklen
  real(8)
               :: fl_s1(mnblocklen)
  real(8)
               :: fl_sd(mnblocklen)
                                                 ! for sampling with variable blocklen
  real(8)
               :: fl_s2(mnblocklen)
                                                 ! for sampling with variable blocklen
               :: av_sd_extrap
                                                 ! for sampling with variable blocklen
  real(8)
  real(8)
               :: av_sd_stateff
                                                 ! for sampling with variable blocklen
               :: fl_sd_extrap
                                                 ! for sampling with variable blocklen
  real(8)
  real(8)
               :: fl_sd_stateff
                                                 ! for sampling with variable blocklen
end type scalar_var
```

1D distribution functions

```
type df_var
   character(27) :: label
                                                        ! label
                 :: norm
                                                        ! normalization factor
   real(8)
                                                        ! number of macrosteps sampled
   integer(4)
                 :: nsamp1
  integer(4)
                :: nsamp2
                                               ! number of values sampled per macrostep
                                                        ! minimum value of df
   real(8)
                 :: min
                                                        ! maximum value of df
   real(8)
                  :: max
                 :: nbin
                                                        ! number of grid points
   integer(4)
   real(8)
                 :: bin
                                                        ! grid length of df
   real(8)
                  :: bini
                                                        ! inverse of bin
                                                ! number of values sampled in each bin during
  real(8)
              :: nsampbin(-1:mnbin_df)
       macrostep
   real(8)
                  :: avs1(-1:mnbin_df)
                                                         ! average of the run
                :: avsd(-1:mnbin_df)
                                                     ! precision of average of the run
  real(8)
   real(8)
                 :: avs2(-1:mnbin_df)
                                                        ! average of a macrostep
end type df_var
```

2D distribution function

```
type df2d_var
                                                       ! label of df
   character(27) :: label
                                                       ! normalization factor
   real(8)
                 :: norm
   integer(4)
                 :: nsamp1
                                                      ! number of macrosteps sampled
                                              ! number of values sampled per macrostep
  integer(4)
               :: nsamp2
                 :: min(2)
   real(8)
                                                       ! minimum value of df
                                                       ! maximum value of df
   real(8)
                 :: max(2)
   integer(4)
                 :: nbin(2)
                                                       ! number of grid points
                                                       ! grid length of df
   real(8)
                 :: bin(2)
   real(8)
                 :: bini(2)
                                                       ! 1/bini
                 :: avs1(-1:mnbin_df2d,-1:mnbin_df2d) ! average of the run
   real(8)
  real(8)
               :: avsd(-1:mnbin_df2d,-1:mnbin_df2d) ! precision of average of the run
                 :: avs2(-1:mnbin_df2d,-1:mnbin_df2d) ! average of a macrostep
   real(8)
end type df2d_var
```

cluster1 trial move

```
type cluster1_tm_var
                                         ! flag for cluster1 tm
   logical
               :: 1
   real(8)
                                         ! relative probability of cluster1 move
                :: p
                                         ! radius of region for cluster members
   real(8)
                :: rad
  real(8)
                :: psel
                                       ! probability to select a particle within rad
   real(8)
                :: dtran
                                         ! maximal translational trial displacement
                :: drot
                                         ! maximal rotational trial displacement
   real(8)
end type cluster1_tm_var
```

cluster 2 trial move

```
type cluster2_tm_var
   logical
               :: 1
                                         ! flag for cluster1 tm
   real(8)
                :: p
                                         ! relative probability of cluster2 move
   real(8)
                :: dtran
                                         ! maximal translational trial displacement
                                         ! maximal rotational trial displacement
   real(8)
                :: drot
                                         ! =0 : search members only of type iptmove
   integer(4)
                :: mode
                                     ! =1 : search members across all particle types
end type cluster2_tm_var
```

trial move

```
type trialmove_var
  logical
               :: 1
                                        ! flag for type of trial move
                                       ! relative probability of type of trial move
   real(8)
                :: p
                :: dtran
                                        ! maximal translational trial displacement
   real(8)
   real(8)
                :: drot
                                        ! maximal rotational trial displacement
                                        ! specific for type of trial displacment
   integer(4)
                :: mode
   logical
                :: lcl1
                                        ! flag for cluster1 trial move
  real(8)
                :: radcl1
                                        ! radius of region for cluster members
  real(8)
               :: pselcl1
                                      ! probability to select a particle within rad
end type trialmove_var
```

node in the DOP-tree

triangle mesh, with DOP-tree

```
type trimesh ! triangle mesh, with DOP-tree
  !private
  real(8), allocatable :: c(:,:) ! c(3,np) coordinates of triangle verticies
  integer(4), allocatable :: t(:,:) ! t(3,nt) triangles as index as verticies into c
  type(node), allocatable :: n(:) ! nodes in tree
  integer(4) :: levels ! levels of subdivisions of triangles
end type
```

affine transformation

sso step

```
type :: step
  integer(8) :: n !number of steps
  real(8) :: d2 !squared displacement
  real(8) :: d4 !displacement**4
end type step
```

SSOPart

```
type :: ssopart_var
   real(8)
               :: fac
                           !increment of part length
   integer(4) :: nextstep !step at which next part starts
   integer(4) :: i
                           !current part
                           !number of parts
   integer(4) :: n
end type ssopart_var
type(ssopart_var), save :: SSOPart
SSOParameters
type :: ssoparam_var
                           ! used dtran
   real(8)
               :: used
   real(8)
               :: opt
                           ! dtran with the highest mobility
   real(8)
               :: err
                        ! accuracy of opt
end type ssoparam_var
type(ssoparam_var), save, allocatable :: SSOParameters(:,:)
sso mobility
type :: mobility_var
   real(8)
               :: val
                              !value
               :: error
   real(8)
                              !error
   real(8)
              :: smooth
                              !smooth
end type mobility_var
type(mobility_var), allocatable, save :: Mobility(:)
celllist cell-pointer-array
type cell_pointer_array
                                                      ! pointer to a cell, usefull to create an
  type(cell_type), pointer
                                   :: p => null()
       array of pointers
end type cell_pointer_array
celllist cell-type
type cell_type
  integer(4)
                                         :: id
                                                               ! for easy recognition
  integer(4)
                                   :: npart
                                                       ! number of particles per cell
                                                       ! number of neighbouring cells
  integer(4)
                                   :: nneighcell
  type(cell_pointer_array), allocatable :: neighcell(:)
                                                            ! pointer to the neighbouring cells
  integer(4)
                                :: iphead
                                                   ! first particle in the linked list
end type cell_type
```

8 Variables

A brief explanation of most global variables is found in file mol.F90. Beside that, the following convention of indices is used.

Indices	Description
ic, jc	Chain number
ip, jp	Particle number
ia, ja	Atom number
ict, jct	Number of chain type
ipt, jpt	Number of particle type
iat, jat	Number of atom type
ictjct	Chain type-chain type pair number
iptjpt	Particle type-particle type pair number
iatjat	Atom type-atom type pair number
iploc	Particle number (local)
ialoc	Atom number (local)

9 Additional Descriptions

9.1 Networks

Overview

Non-periodic network structures in Molsim

The aim of this branch is the "proper" implementation of polymer networks as an integral part of the simulation software Molsim. These networks are called non-periodic networks in order to distinguish them from their macroscopic counter parts (i.e. macrogels). This follows Molsim's philosophy of having different objects in a hierarchy of different levels.

Object type	Hierarchy Level	Object type	Hierarchy Level
Atoms	1		
Particles	2		
Chains	3		
Hierarchical Structures	4	Non-periodic Network Structures	4

Within this scheme, atoms are part of particles, which in turn may be connected to form a chain. These can be interconnected by cross-links to form more complex structures, as e.g. hierarchical structures or network-like structures. Note, that for all object types on the left hand side of the table, a programming infrastructure in form of various characteristic parameters and (pseudo-)pointers is established already, thus, leading to a high programming flexibility, whenever one wishes to expand the functionality of the software with respect to one of these object types. On the other hand, non-periodic network structures have been introduced to Molsim later and therefore a programming infrastructure is not given. Besides the mentioned characteristic parameters and (pseudo-)pointers, a series of subroutines in Molsim handles the statistical treatment of the left hand side object types. Within the scope of this branch the corresponding parameters and functionalities shall be established.

Model description

The aim of the implemented network model is to simulate the properties of non-periodic polymer networks of spherical shape (i.e. microgels). The model network comprises cross-linking particles (so called nodes) and chain particles. The nodes are set on the carbon positions of the diamond lattice (space group Fd3m, no. 227). The cubic lattice is cropped to obtain a spherical shape by defining a cutoff radius - all nodes within the cutoff radius will be kept; all nodes outside of the cutoff radius will be discarded. The remaining nodes are then connected via polymer chains, which consist of a number of chain particles per chain. Within the model the so called dangling chains are included. They are connected to only one cross-link and thus they form the outer periphery of the modelled gel. Via the input parameter nnwt it is possible to define the number of network types present in the system. Different network types may for example differ in their number (nnwnwt(inwt)) and size (rnwt(inwt)). Further, different types of particles may be used as building blocks and hence, polymer networks with different physical properties may be obtained.

Within this scheme it is important to note, that always one particle type forms the nodes of one certain network type. Analogues, one chain type forms the chains of a certain network type. This is controlled via the parameters iptclnwt(inwt) (i.e. the particle type forming the cross-links of network type inwt) and ncctnwt(ict,inwt) (number of chains of a chain type ict belonging to one network of network type inwt). The size of the networks cannot be chosen in a continous manner, as the nodes are set on discrete diamond lattice positions, hence the generated networks are available with discrete numbers of nodes.

Details of implementation

Network Configuration

General

The possibility to set finite networks has already been possible prior to the here undertaken implementations by means of the subroutine SetNetwork. Within the branch Networks the already existing routine SetNetwork has been adapted to the new nomenclature regarding network related parameters (see below). Both, in the subroutine SetNetwork and in the general output about the system parameters, the output about networks has been refined.

Shift of cropping sphere

The parameter shiftnwt(1:3,1:nnwt) was introduced. It enables one to obtain topology-modified networks, which mainly differ in how the structure (*i.e.* dangling chains and interconnectivity of the outer layer) of their periphery looks like. In the process of generating the network structure a cubic diamond lattice will be span and cropped by a spherical cut-off. The default network will be set having the (0,0,0) position of the diamond lattice unit cell in the center of the cropping sphere. This corresponds to $shiftnwt(1 \leftarrow :3,inwt) = 0.0$ of a certain network type inwt. By setting 'shiftnwt(1:3,inwt)' the center of the cropping sphere may be shifted to any point of the unit cell. Note, that shiftnwt accepts its values in the range of 0.0 to 1.0, respectively. Higher values will work aswell, but due to the underlying periodic boundary conditions which the diamond lattice is subject to, shiftnwt(1:3,inwt) = 0.0 is equivalent to shiftnwt(1:3,inwt) = 1.0 and so on.

Network Properties

Similar to the possibility to evaluate characteristic properties of chain objects in Molsim, a new subroutine has been implemented, namely CalcNetworkProperty. Within the subroutine CalcNetworkProperty the following properties are evaluated:

- ro(3): center of mass
- · rg2: radius of gyration squared
- rg2x: radius of gyration squared projection on x-axis
- rg2y: radius of gyration squared projection on y-axis
- · rg2z: radius of gyration squared projection on z-axis
- rg2s: square extention along principal axes (smallest)
- rg2m: square extention along principal axes (middle)

- rg21: square extention along principal axes (largest)
- asph: asphericity (JCP 100, 636 (1994))
- theta(3): angle between x-, y- and z-axis, respectively, and the axis of the network's largest extension
- alpha: degree of ionization (for titrating systems)
- eivr(3,3): normalized eigenvectors of the principal frame

Averages of Network Properties

Within the subroutine NetworkAver the averages and precisions with regards to the properties available by CalcNetworkProperties are formed (except for ro(3) and eivr(3,3)). The obtained network quantities will be written to the Output-File when networks are present in the system. Example output:

* network quantities						*		

quantity		average	precision	fluctuation	precision	stat eff		
<r(g)**2>**0.5</r(g)**2>	=	75.32408	0.00915	1.77217	0.06023	2.		
<r(g)**2_x>**0.5</r(g)**2_x>	=	43.51678	0.00452	1.36440	0.09226	0.		
<r(g)**2_y>**0.5</r(g)**2_y>	=	43.49056	0.00458	1.37475	0.14198	0.		
<r(g)**2_z>**0.5</r(g)**2_z>	=	43.45776	0.01588	1.66017	0.01477	3.		
smallest rms mom. p.a.	=	43.39747	0.01547	1.77924	0.10399	2.		
intermediate rms mom. p.a.	=	43.47212	0.01324	1.65132	0.11078	2.		
largest rms mom. p.a.	=	43.59531	0.01354	1.74615	0.07283	1.		
<asphericity></asphericity>	=	0.00001	0.00000	0.00000	0.00000	1.		
<xtheta></xtheta>	=	72.47060	8.72678	44.88033	3.26752	0.		
<ytheta></ytheta>	=	101.41857	3.97221	38.12409	4.66430	0.		
<ztheta></ztheta>	=	78.86201	2.54458	19.66774	2.88301	0.		

asphericity (<2:nd moments>) = 0.00001

Here, the asphericity is being calculated according to Zifferer, G. and Olaj, O. F., Journal of Chemical Physics 1994, 100, 636. The second displayed asphericity (asphericity ($<2\leftarrow$:nd moments>)) is being directly calculated from the respective average of the principal moments of a macrostep or the whole simulation.

Network Distribution Functions

The functionality of finite network-related simulations has been extended by allowing for statistical analysis of network properties by means of distribution functions.

Network Property Distribution Functions

The following network property distribution functions are available by means of the setting in the nml← NetworkDF.

type	label	description
1	rg	network radius of gyration distribution
2	asph	network asphericity distribution
3	alpha	network degree of ionization distribution
4	rgchain	chain radius of gyration distribution of network chains
5	reechain	chain end-to-end distance distribution of network chains

The described type i refers to the array index to be used in the respective vtype(i) construct.

Radial Network Distribution Functions

The following radial network distribution functions are available by means of the setting in the nml← NetworkRadialDF.

type	label	description
1	rpart	radial particle number distribution
2	rdens	radial particle density distribution
3	rgchain	radial chain radius of gyration
4	q	radial sum of all charges
5	qcum	radial cumulated sum of all charges
6	alpha	radial degree of ionization
7	rchain	radial chain number distribution

The described type i refers to the array index to be used in the respective vtype(i) construct.

Network Generation Groups

A new way of forming groups within networks has been established. Within this grouping scheme the chains are numbered starting from the dangling chains at the periphery of the network and taking the shortest way to the core chains of the network structure. A schematic representation of the numbering is presented below.

Note, that this scheme works only for structures set with 'txsetconf = 'network'. The cross-links of the network will be assigned to have no group. In order to obtain this grouping set eitherreforfieldin thenmlGrouptonetworkgenerations`.

Testin

fnw.nwt2.mc.in

The testin-file was added to the in-directory. The corresponding out-file was stored in the Save-directory. The documents "todiff.txt" and "goall.sh" were expanded by the new input-file.

fnw.nwgen.df.mc.in

The testin-file was added to the in-directory. The corresponding out-file was stored in the Save-directory. The documents "todiff.txt" and "goall.sh" were expanded by the new input-file.

Network variables

Internal Variables

Besides of the external network related parameters (i.e. input variables) a number of internal variables has been introduced in order to set up the programming infrastucture of non-periodic networks. All variables are declared in mol . F90.

Keyword	Туре	Description
nnw	integer	Total number of networks
nctnwt(1:nnwt)	integer	Number of different chain types in network type inwt
ncnwt(1:nnwt)	integer	Number of chains per network of network type inwt
npnwt(1:nnwt)	integer	Number of particles per network of network type inwt
nclnwt(1:nnwt)	integer	Number of cross-links per network of network type inwt
nnwtnwt	integer	Number of different network type pairs
txnwtnwt(1:nnwtnwt)	character(21)	Name of network type pair inwtnwt
massnwt(1:nnwt)	real	Mass per network of network type inwt
massinwt(1:nnwt)	real	Inverse mass per network of network type inwt
lnetwork	logical	. true . if networks are used
<pre>lptnwt(1:mnpt,1:nnwt)</pre>	logical	.true. if particle type ipt is part of networks of network type inwt
lpnnwn(1:np,1:nnw)	logical	.true. if particle ip is part of network inw
npweakchargenwt(1:nnwt)	integer	Number of titratable charges per network of network type inwt

Network pointer

Keyword	Type	Description
inwtnwn(1:nnw)	integer	In: network number inw, Out: its network type inwt
inwtct(1:nct)	integer	In: chain type ict, Out: its network type inwt
inwtcn(1:nc)	integer	In: chain number ic, Out: its network type inwt

Keyword	Туре	Description
inwncn(1:nc)	integer	In: chain number ic, Out: its network numberinw
inwnnwt(1:nnwt)	integer	In: network type inwt, Out: its first network inw
<pre>inwtnwt(1:nnwt,1:nnwt)</pre>	integer	In: two network types inwt/jnwt, Out: its network
		type pair number inwtnwt
icnclocnwn(1:ncnwt,1:nnw)	integer	In: local chain number icloc and newtwork number
		inw, Out: its chain number ic
<pre>ipncllocnwn(1:nclnwt,1:nnw)</pre>	integer	In: local cross-link number iclloc and network num-
		ber inw, Out: its particle number ip
ipnplocnwn(1:npnwt,1:nnw)	integer	In: local particle number iploc and network number
		inw, Out: its particle number ip

9.2 Complexation-Analysis

Complexation Analysis

Interparticle complexation can be analyzed using Molsim. To use it set

```
lstaticuser = .true.
```

in nmlStatic. Additionally the first 4 characters of txuser in nmlSystem have to be comp. The control of the complex-statistics is given by the namelist nmlComplexation:

```
&nmlcomplexation
  rcut_complexation = ...,
  lcomplexfraction = ...,
  lclusterdf = ...,
/
```

control over the ComplexDF routine is given by the namelist nmlComplexDF:

```
&nmlcomplexdf
  vtype = ...
```

where rcut_complexation defines the maximum distance for two particles to form a complex; $1 \leftarrow \text{ComplexFraction (logical)}$ switches on the calculation of the fraction of complexed particles; $1 \leftarrow \text{ClusterDF (logical)}$ switches on the calculation of the size distribution of complexed formed by the complexation.

• Testfile: complexation.mc.in

Fraction of Complexation

This routine return the fraction of complexed particles of each particle-particle commbination. It return the fraction of X beads which are closer than $rcut_complexation$ to a Y bead as w(cmplx): X - Y. E.g.:

quantity	average	precision	fluctuation	precision	stat eff
w(cmplx): a - a	0.30000	0.00000	0.00000	0.00000	0.
w(cmplx): a - b	0.20000	0.00000	0.00000	0.00000	0.
w(cmplx): b - a	0.40000	0.00000	0.00000	0.00000	0.
w(cmplx): b - b	0.50000	0.00000	0.00000	0.00000	0.

means that 30% of the A beads are complexed to other A beads; 20% of the A beads are complexed to a B bead; 40% of the B beads form a complex with an A bead and 50% of the B beads form a Complex with another B bead.

Complexation Distribution

This routine return the distribution of the fraction of complexed particles of each particle-particle commbination. It return the fraction of X beads which are closer than rcut_complexation to a Y bead. To control of the distribution function the nmlComplexDist is used. Additionally also the distribution of the complexation rate of the individual chains can be calculated.

Segment Complexation

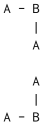
This routine return the fraction of complexation of each segment along the chains of the system (averaged over all chains of one type).

Size distribution of the binary clusters

Here the size of clusters formed by neighbouring complexed particles of two different types are measured. A cluster is formed only by connections between particles of different particle type. E.g.



forms one cluster of size 5 and one of size 1 (single uncomplexed A particle). In contrast



forms two clusters of size 3.

As an output the size distribution (average number of cluster of a specific size) are given for all combinations of two different particle types.

9.3 Advanced Configurations

The support for branched, random and repeating structures is improved:

Branched Structures

When setting a hierarchical structure using the subroutine SetHierarchical the inability to set one
particle will not lead to an abortion of the whole program. Instead another it is attempted ntrydef
(default 100) times to set the whole structure. If one particle cannot be set the whole structure is
reset.

• Testfile: hierarchical.mc.in

9.4 Copolymer Sequence

Overview

In general chains consisting of more than one particle type (i.e. copolymers) may be simulated using Molsim. Different ways to generate such chains are triggered by the parameter txcopolymer. Existing modes comprise alternating, and block copolymers. Within the scope of this branch the functionality to simulate copolymers with random, repeating or irregular or highly specific monomer distribution has been added (sequence).

Usage

Sequence

The input of the copolymer sequence has been realized by introducing the parameter iptsegct(iseg,ict), which specifies the particle type ipt of chain segments iseg of chains of type ict. First of all the parameter txcopolymer(ict) has to be set to sequence for the chain type in question in the namelist nmlParticle:

```
&nmlParticle
...
txcopolymer(ict) = 'sequence' ,
...
/
```

where ict should be replaced by the corresponding chain type number. The parameter iptseg(iseg,ict) is then being set in the namelist nmlCopolymerSequence. For example

```
&nmlCopolymerSequence
  iptsegct(1,ict) = ipt , iptsegct(2,ict) = ipt , ... , iptsegct(npct,ict) = ipt ,
   ... /
```

where again ict should be substituted by the corresponding chain type number, ipt should be replaced by the intended particle type number and npct corresponds to the number of particles per chain of chain type ict.

Repeating Copolymers

Copolymers with a defined repeating block structure can be generated. It consists of repeating units, each consisting of blocks of one particle type.

• in nmlParticle set the parameters of the chain type ict which is to be repeating

```
txcopolymer(ict) = "repeating",
and set
nblockict(ict)
```

to the number of blocks within a repeating unit to the intended number.

• The detailed repeating structure is given by nmlRepeating:

```
&nmlrepeating
rep_iblock_ict( iblock , ict)%pt = ... ,
rep_iblock_ict(iblock , ict)%np = ... ,
/
```

where the particle type pt and number of particles np of each iblock for eachict is defined.

If not enough particles of a certain type are given for a chain to fulfill the regular structure, the repeating structure is continued, but only filling the blocks with the remaining particles.

Example: One chain with the following input (excerpt)

```
&nmlparticle
  nct = 1,
  ncct = 1,
  npptct(1:3,1) = 4,5,10,
  nblockict(1) = 3,
  txcopolymer='repeating',
/
&nmlrepeating
  rep_iblock_ict(1:3,1)%pt = 2, 3, 1
  rep_iblock_ict(1:3,1)%np = 2, 3, 1
```

will generate such a repeating structure:

Seg-	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
ment																			
Par-	5	6	10	11	12	1	7	8	13	14	15	2	9	16	17	18	3	19	4
ticle																			
Num-																			
ber																			
Par-	2	2	3	3	3	1	2	2	3	3	3	1	2	3	3	3	1	3	1
ticle																			
Type																			
Re-	1		2			3	1		2			3	1	2		•	3	2	3
peat-																			
ing																			
Block																			
Re-	1						2						3					4	
peat-																			
ing																			
Unit																			

Random Copolymers

Random copolymers can be generated. The random sequence is generated by first creating a list where the particle type of each segment is given as for a block-like copolymer. This list is then shuffled using a Knuth Shuffle. Now the particles are sequentially assigned to segments of its type. This way the particle types are randomly distributed, but the particles of one type are of increasing index along the chain. Note that if you want the exact sequence printed in your output file you need to set itestpart = 10 in the nmlParticle.

• in nmlParticle set the parameters of the chain type ict which is to be random:

```
txcopolymer(ict) = "random",
```

Testin

The Testin-file covering these functionalities are chain. sequence. fnw.mc.in and copoly-seq.mc. \leftarrow in.

9.5 Celllist

Overview

The cell list is implemented in molsim acts a a neighbor list, so that fewer particle pairs with a distance larger than $r_{\rm cut}$ are considered. This allows for much quicker simulations, especially for large systems with short ranged interactions. The simulation duration scales with $\mathcal{O}(n)$. The advantages are compared to the other neighbor lists are the following:

- The cell list is updated after every accepted step, so that is is always up to date. This allows for non-local moves to be carried out without needing to artificially enlarge the cell size. This is not the case for vlist and llist which are currently implemented in molsim.
- The cell structure is generated only once at the beginning of the simulation. The time for the generation of the cell list scales with the number of cells, but the execution speed should be independent of the number of cells. In llist the number of cells were capped to a small number (1000), so that large systems had to use large cells. Increasing the number of possible cells did not help either, as the time to find the neighbors in each step increased with the number of cells, making the llist very slow when having many cells ($\approx 10^6$).

Details of Implementation

Working Procedure

The usage of a neighbor list within Molsim is controlled by the namelist nmlIntList. Usage if the cell list is achieved by setting txintlist = 'cellList'. The cell size is the chosen to be as small as possible to fill the whole space evenly with cells, while still remaining larger than rcut.

One can create cells of different size, by changing the value of drnlist, which will be added to rcut for setting up the cell list. Most notably one can set drnlist to a negative value to even further reduce the cell size. This will lead to less particles with a distance larger than rcut, but more cells will be considered. Whether or not this approach increases the speed of the simulation may depend on the system.

All parameters except txintlist and drnlist of nmlIntList are not used for the cell list. A typical input file could be the following:

```
&nmlintlist txintlist = 'cellList', drnlist = 0.0, /
```

Testin

An example for the usage of the cell list can be found the Testin directory.

```
• lj.mc.clist.in
```

Further Reading

• cell List https://en.wikipedia.org/wiki/Cell_lists

9.6 SSO

Overview

SSO

In this section the sso-functionality is added to Molsim. For a detailed description of the algorith see the master-thesis of pascal or the corresponding publication. The SSO algorithm allows for on the fly optimization of the displacement parameter of the single particle move during the equilibration run.

Sperarate Local from Non-Local

In addition the 1mcsep parameter was introduced. This separates local (Single-Particle and SSO Move) from non-local moves (all the others) during the simulation. During each MC-Pass it is evaluated, wheter non-local moves are to be carried out (accoring to their probability) and only if they will be carried out the neihbour list ist updated with a large drnlist (4* contou length of the longest chain). This allows for using a tigh neighbour-list during the simulation with local-moves.

Testin

The following test-projects are added to the Testin directory:

- · sso.sso.in: Test of the sso-move
- sso.lmcsep.in: Test of the Imcsep-setting

9.7 Image generation with VMD

Overview

In order to generate images for publications, presentations and for visual insight in simulated systems, Molsim offers the possibility to generate image files in the VTF format ("VTF Trajectory Format"). The VTF format has specifically been designed for people working with their own simulation code with the aim of visualization via the image software VMD ("Visual Molecular Dynamics"). VMD offers a broad spectrum of functionality with regards to the visualization of molecular systems with the possibility of rendering high quality images.

Details of Implementation

File Types

The visualization of simulation snapshots requires two different file types.

- vtf: The vtf file contains atom, bond and box length definitions, followed by an arbitrary number of coordinate blocks.
- tc1: The tcl file forms the interface between Molsim and VMD. The whole settings of VMD may be controlled via tcl scripts. Here, color specifications, additional geometric objects (*e. g.* simulation cell) and the loading of the vtf file is stored.

Working Procedure

The generation of images with Molsim is controlled by the namelists nmlSystem, nmlImage and namelist nmlVTF/nmlVRML. Note, that this manual covers only the image generation using the vtf format for the visualization with VMD. For the generation of images in the vrml format, please confer the MOLSIM manual. In the following the work flow for the image generation is described.

Request the generation of images by setting limage = .true. in the namelist nmlSystem

```
&nmlsystem limage = .true.,/
```

 Request the generation of images using the vtf format by setting lvtf = .true. in the namelist nmlImage

```
&nmlimage lvtf = .true.,/
```

- Further specifications may be given in the namelist nm1VTF. With just the default settings, images of the initial and of the final configuration will be stored, respectively.
- Run the simulation
- Open the vtf image file by executing the accompanying tcl script in VMD

```
vmd -e {name-of-project}.tcl
```

· Apply individual modifications to representation

Render a snapshot

Here are some tipps on how to generate nice images using VMD.

- use the AOShiny material
- turn on the lights, often the scene generated looks less bright when rendered as presented one the screen in VMD

light 2 on light 3 on

- turn on the shadows with display shadows on
- turn on the ambientocclusion display ambientocclusion on
- use the GLSL rendermode display rendermode GLSL

Then render the scene (with the path to you Tachyon executable adapted

```
render Tachyon scene.dat exec "<path to you Tachyon executable>" -aasamples 48 scene.dat -res 2096 1048 -format BMP -o scene.b
```

You might want to adapt the name of the image (scene.*), the resolution and the the aasamples.

Testin

An example for the usage of the image generation in the vtf format may be found in the Testin directory.

· vtfimage.fnw.mc.in

Further Reading

- VTF https://www.github.com/olenz/vtfplugin/wiki/VTF-format
- TCL https://www.tcl.tk
- VMD http://www.ks.uiuc.edu/Research/vmd

10 Developing

When developing Molsim, please adhere to the contributing guide, which is shipped in the MOLSIM repository.

10.1 Testin

In the Molsim directory you will find a directory called Testin. The Testin offers an efficient way to check whether changes of the Molsim source code lead to any malfunction of the program.

General idea

The general idea of the Testin-directory is to account for the sustainment of the functionality of Molsim. Whenever the source code of Molsim is changed, one has to ensure that none of the functionalities of Molsim have been broken. Despite of a few aspects (e.g. format changes) the program execution should yield consistent results. By comparing the results of test simulations of the current with the previous version it is easy to detect whatever kind of malfunctions.

The Testin-directory provides a method to compare the output of the current version to the last stable version.

Usage

- First you need to generate the output from the stable version. The stable version should be the latest master branch from which your branch diverged. If you are unsure, check when the file Testin/stable.md5sum was last changed. The repository at this stage is the latest stable version. When you have checked out the stable version you want to compare with, run make stable in the Testin directory. This will compile the current source code with the mode = test flag and create the output in the Testin/out_stable directory.
- at any time you can run make out to run the same input files and store the output in the Testin/out.
- to compare the output of the out and out_stable directories, run make diff. Note that if the out files are not present, make diff will create them accordingly. The collected differences are written into the file diff.out

The diff should, in general, yield no differences. If differences were found, one needs to carefully inspect each of these instances for whether one deals with expected changes or unexpected changes.

- Unexpected changes: Investigate where the difference is coming from and fix the bug!
- Expected changes: When you have only expected changes, add comments to the diff.out file, which describe the changes. This commented version of the diff.out files should be attached to the pull request when you want to merge into the stable version. Note that the diff.out file will be overwritten when running make so you might want to add you comments to a copy of diff.out

- After your merge request has been positively reviewed, run make declarestable to declare your
 current version of the code as the stable one. The files included in the check for the stable version
 are all .F90 files, the makefile and the molsim_ser file in the Src directory. Additionally also the
 configure.sh file in the molsim root is checked.
- to delete all output of the out runs (and the diff.out file!) run make clean, to delete the out_ stable dir run make cleanstable. To delete both run make cleanall.

Further notes:

- You can speed up the process, by running make in parallel (e.g. use make -j 4, to run on 4 cores).
- When running make all needed files are generated automatically, so you do not need to run make out before running make diff. A single call of make is enough.
- The out_stable files can not be created automatically, when you are in the directory of a modified version of molsim. Checkout the stable version, run make stable, and go back to your branch. The out_stable dir should stay, and can be used to compare with.

List of elements contained in the Testin-directory

element	description
in	input files covering all functionalities of Molsim
todiff.txt	all output files to diff
Makefile	Makefile to run all tests
scripts/diff1.sh	script to compare the output files
scripts/molsim_stable.sh	script to generate outputs from the stable version
stable.md5sum	file defining the current stable version

After executing make you will additionally find:

element	description
out	output files resulting from the Molsim version under examination
out_stable	output files corresponding to the input file in in, created with the latest stable version.
diff.out	Output of the diff between the current and the stable versions

10.2 Random Numbers

The random number generator used in Molsim is rand function as described by Press et al. in Numerical Recipes in Fortran 77, *The Art of Scientific Computing*, Second Edition, 1997. It has a a period of $\approx 3.1 \times 10^{18}$ and returns a uniform random number in the range (0.0, 1.0).

The routine uses a pair of of integers (ixseed and iyseed) to generate random numbers. The iseed you give in the nmlSystem is used to generate the first integer number pair. When iseed is negative the current time is used to set iseed. In most cases it is sufficient to set only iseed. The case where you also want to set ixseed and iyseed are when you want to use a specific random number sequence from a previous simulation.

- You want to set ixseed and iyseed to some specific values:
 - In nmlSystem set iseed, iyseed and ixseed to the desired values and set luseXYseed to
 .t..
 - Note that if you set iseed to a negative value, then the seed will not be set by the cpu
 time. A negative seed will reinitialize the random number generator and overwrite ixseed and
 iyseed.
- Given an output file where you want to recreate the random numbers you have three possibilities:
 - If you want to generate a new simulation with the same initial seed: set i seed to the value of the seed given at the beginning of the input file
 - If you want to generate a new simulation with the same initial seed of the previous simulation
 which was started by setting iyseed and ixseed explicitly: set iseed, iyseed and ixseed to
 the ones given in the beginning of the output file set luseXYseed in nmlSystem to .t..
 - If you want to run a simulation, continuing with the same chain of random numbers where a
 the previous simulation left of: set iseed, iyseed and ixseed to the ones given in the end of
 the output file set luseXYseed in nmlSystem to .t..

10.3 Compilation Modes

To make the development easier, several compilation modes are provided. They are used as following:

when making Molsim a second argument can be massed to make: mode=<mode>. Example:

make ser mode=test

to compile Molsim in the test mode. The following modes are supported:

- · normal: compile with normal options to achieve high computational efficiency.
- test: compile in such a way, that the results are reproducible. Use this mode when performing tests in the Testin directory.
- debug: During run-time the compiler checks for out-of-bounds arrays and uninitialized variables. Only use for debug purposes.
- quick: Fast compilation without optimization. Use only for testing purposes.
- · warn: The compiler gives all warnings during compile time.
- gprof: For profiling the execution with gprof (only for the gfortran compiler)

Gprof

To profile you program with gprof, do the following (only for the gfortran compiler):

• Compile with mode=gprof:

make ser mode=gprof

• run Molsim on you project in the directory of you input file. A file gmon.out should be generated

molsim_ser <Projectname>

• run gprof

gprof <path to the molsim exe> gmon.out

11 Appendix

11.1 Namelist

- A namelist consists of (i) one start statement, (ii), one end statement and (iii) a list (possibly empty) between the start and end statements.
- The start statement consists of an ampersand (&) followed by the name of the namelist.
- The end statement consists of a slash (/).
- The list is composed of variables and their values separated with either by commas.
- The order of the variables and their assignments is normally irrelevant.
- However, if a variable is assigned values more than once, the last one takes place.
- Arrays are allocated either element by element, consecutively by listing values separated by commas, or by a combination of these to ways.
- A value, say 5, appearing r times may be expressed as r∗5.
- Note the order Fortran stores arrays elements: the left most index is running fastest and the right most slowest.
- Comments are allowed and should proceed by an exclamation sign (!).
- Generally, if a program reads several namelists, they have to occur in the inputfile in the same order as they are read from the program. Here, the order of the namelist is arbitrary.
- A namelist with no list (empty namelist) has to be specified.
- Namelists that are not read by the program do no harm.

Example: Consider the following declarations

```
character(4) :: title
integer(4) :: m, n
real(8) :: arr1(1:3), arr2(1:2,1:2), arr3(1:2,1:2)
```

The namelist nm1System below illustrates an element-by-element array assignment

```
&nmlSystem
title = 'head',
n = 10, m = 10,
n = 20,
arr1(1) = 1.5, arr1(2) = 2.5, arr1(3) = 2.5,
arr2(1,1) = 1.0, arr2(2,1) = 2.0, arr2(1,2) = 3.0, arr2(2,2) = 4.0,
arr3(1,1) = 1.0, arr3(2,1) = 2.0, arr3(1,2) = 3.0, arr3(2,2) = 4.0,
```

which could, for example, be shortened to

```
&nmlSystem
title = 'head',
n = 20,
m = 10,
arr1 = 1.5, 2*2.5,
arr2 = 1.0, 2.0, 3.0, 4.0,
arr3(1:2,1) = 1.0, 2.0, arr2(1:2,2) = 3.0, 4.0,
/
```

11.2 Examples of namelists describing different objects

Examples of namelist &nmlParticle describing different types of objects are given in this appendix.

Hard-sphere particles

```
&nmlParticle
          beginning of the namelist
npt = 1,
          number of particle types
txpt = 'hs',
         particle's label
nppt = 100,
          number of particles of this type
natpt = 1,
          number of atom types which this particle is composed of
txat = 'hs',
          atom's label
massat= 1.0,
         molar mass of the atom (in grams per mole)
radat = 1.0,
         hard-sphere radius of the atom
naatpt(1,1) = 1,
         number of 'hs' atoms in the 'hs' particle
txaat(1,1) = 'hs',
         label of 'hs' atom in the 'hs' particle
          end of the namelist
```

Lennard-Jones particles

```
txpt = 'lj',
          particle's label
nppt = 500,
          number of particles of this type
natpt = 1,
          number of atom types which this particle is composed of
txat = 'center',
          atom's label
massat= 40.0,
         molar mass of the atom
radat = 1.0,
          hard-sphere radius of the atom
sigat = 3.405,
         Lennard-Jones parameter 'sigma'
epsat = 0.99606555,
          Lennard-Jones parameter 'epsilon'
naatpt(1,1) = 1,
          number of 'center' atoms in the 'lj' particle
txaat(1,1) = 'lj',
          label of 'center' atom in the 'lj' particle
          end of the namelist
```

Lennard-Jones particles with an embedded dipole

```
&nmlParticle
         beginning of the namelist
txelec= 'dip',
         enable atoms to possess a static dipole
npt = 1,
         number of particle types
txpt = 'LJ',
         particle's label
nppt = 100,
         number of particles of this type
natpt = 1,
         number of atom types which this particle is composed of
txat = 'LJ',
         atom's label
massat= 18.0,
         molar mass of the atom
radat = 0.5,
         hard-sphere radius of the atom
zat = 0.0,
         atom's valency
sigat = 2.88630,
         Lennard-Jones parameter 'sigma'
epsat = 1.97023,
         Lennard-Jones parameter 'epsilon'
naatpt(1,1) = 1,
```

```
number of 'LJ' atoms in the 'LJ' particle
txaat(1,1) = 'LJ',
         label of 'LJ' atom in the 'LJ' particle
dipain(1,1,1) = 0.0000,
         x, y, and z components of the static dipole of the first atom type of the first particle type
0.0000, 0.10584,
         end of the namelist
Note: Here, dipain(1,1,1) is a simplified version of dipain(1:3,1,1).
Nemo water (3-site water model with dipoles and polarizabilities)
&nmlParticle
         beginning of the namelist
txelec= 'pol',
         atoms possess charges, static and induced dipoles
npt = 1,
         number of particle types
txpt = 'water',
         particle's label
nppt = 2,
         number of particles of this type
natpt = 2,
         number of atom types which 'water' particles are composed of
txat = 'o ','h ',
         atom's labels; 1st atom type is oxygen
massat= 16.0, 1.0,
         molar mass of the atom
radat = 0.0, 0.0,
         hard-sphere radius of both atom types
zat = -0.80100, 0.400500,
         valency of oxygen and hydrogen atom
naatpt(1,1) = 1, 2,
         number of oxygen and hydrogen atom in water
txaat(1,1) = 'o', 'h', 'h',
         molecule atom's label
rain(1,1,1) = 0.0, 0.0, -0.0656,
         x, y, and z coordinates of oxygen
rain(1,2,1) = 0.7572, 0.0, 0.5205,
         x, y, and z coordinates of 1st hydrogen
rain(1,3,1) = -0.7572, 0.0, 0.5205,
         x, y, and z coordinates of 2nd hydrogen
dipain(1,1,1) = 0.0000, 0.0000, -0.1299,
         x, y, and z components of the static dipole of the oxygen atom
dipain(1,2,1) = 0.0784, 0.0000, 0.0422,
         x, y, and z components of the static dipole of the 1st hydrogen atom
dipain(1,3,1) = -0.0784, 0.0000, 0.0422,
         x, y, and z components of the static dipole of the 2nd hydrogen atom
polain(1,1,1) = 0.6715, 0.6133, 0.7002, 0.0000, 0.0000, 0.0000,
```

xx, yy, zz, xy, xz, and zz components of the symmetric polarizability of the oxygen atom

```
polain(1,2,1) = 0.2199, 0.0756, 0.1441, 0.0000, 0.1005, 0.0000, xx, yy, zz, xy, xz, and zz components of the symmetric polarizability of the 1st hydrogen atom polain(1,3,1) = 0.2199, 0.0756, 0.1441, 0.0000, -0.1005, 0.0000, xx, yy, zz, xy, xz, and zz components of the symmetric polarizability of the 2nd hydrogen atom / end of the namelist
```

Simple 1:1 elecrolyte

```
&nmlParticle
          beginning of the namelist
npt = 2,
          number of particle types
txpt = 'ion1', 'ion2',
          labels of both particle types
nppt = 108, 108,
          number of both particle types
natpt = 1, 1,
          number of atom types which particles are composed of
txat = 'site1','site2',
         labels of atom types
massat= 23.0, 35.4,
         molar mass of the atom (in grams per mole)
radat = 2.0, 2.0,
          hard-sphere radius of both atom types
zat = 1.0, -1.0,
          valency of both atom types
naatpt(1,1) = 1,
          number of 'site1' atoms in 'ion1' particle
txaat(1,1) = 'site1',
          label of 'site1' atom in 'ion1' particle
naatpt(1,2) = 1,
          number of 'site2' atoms in 'ion2' particle
txaat(1,2) = 'site2',
         label of 'site2' atom in 'ion2' particle
          end of the namelist
```

Macroion + counterions

```
natpt = 1, 1,
          number of atom types which particles are composed of
txat = 'mic', 'ion',
          labels of atom types
massat= 460.0, 23.0,
          molar mass of the atom (in grams per mole)
zat = -60, 1,
          valency of both atom types
radat = 20.0, 2.0,
         hard-sphere radius of both atom types
naatpt(1,1) = 1,
          number of 'mic' atoms in 'macroion' particle
txaat(1,1) = 'macroion',
          label of 'macroion' atom in 'macroion' particle
naatpt(1,2) = 1,
          number of 'ion' atoms in 'ion' particle
txaat(1,2) = 'ion',
          label of 'ion' atom in 'ion' particle
          end of the namelist
```

Polyions + counterions

```
&nmlParticle
         beginning of the namelist
nct = 1,
         number of chain types
txct = '100-mer',
         label of the chain type
ncct = 10,
         number of chains in the system
npptct(1,1) = 100, 0
         number of particles in the '100-mer' is equal to 100 of type 'pe' and zero of type ion'
npt = 2,
         number of particle types
txpt = 'pe', 'ion',
         labels of both particle types
nppt = 1000, 1000,
         number of particles of each particle types
natpt = 1, 1
         number of atom types which particles are composed of
txat = 'bead', 'ion',
         labels of atom types
massat = 10.0, 10,
         molar mass of the atom (in grams per mole)
radat = 2.0, 2.0,
         hard-sphere radius of both atom types
zat = 1.0, -1.0,
         valency of both atom types
naatpt(1,1) = 1
```

Diamond-like polyelectrolyte gel + counterions

```
&nmlParticle
          beginning of the namelist
lclink=.true.,
          flag to enable crosslinking
maxnbondcl= 4, 2, 0,
          maximum number of crosslinks for types of particles
nct = 1,
          number of chain types
txct ='strand'
          label of chain type
ncct = 16,
          number of chains of type 'strand'
npptct(2,1) = 10,
          number of particles of type 'strand' in the chain type 'strand'
npt = 3,
          number of particle types
txpt='node','strand','countion',
          labels of all particle types
nppt = 8, 160, 168,
          number of particles of each particle types
natpt = 1, 1, 1
          number of atom types which particles are composed of
txat='bead1','bead2','countion'
          labels of atom types
radat = 2.0, 2.0, 2.0,
          hard-sphere radius of both atom types
zat = 1.0, 1.0, -1.0,
          valency of all three atom types
naatpt(1,1) = 1,
          number of 'bead' atoms in 'node' particle
txaat(1,1) = 'bead',
          label of 'bead' atom in 'node' particle
naatpt(1,2) = 1,
          number of 'bead2' atoms in 'strand' particle
txaat(1,2) = 'bead2',
          label of 'bead2' atom in 'strand' particle
```

Bottle-brushes polymers with uneven side-chain distribution

```
&nmlParticle
         beginning of the namelist
lclink =.true.,
         flag to enable crosslinking
maxnbondcl = 1, 1,
         maximum number of crosslinks for types of particles
ngen = 1,
         number of generations of hierarchichal structure
ictgen(0) = 1,
         chain type of 0th generation (main chain)
ictgen(1) = 2,
         chain type of 1st generation (side chain)
nbranch = 5,
         number of branches on the main chain
ibranchpbeg = 1,
         main chain particle number of the first branch point
ibranchpinc = 1,
         segment increment for branch points on the main chain
nct = 2,
         number of chain types
txct ='10-mer1','5-mer2',
         label of each chain type
ncct = 2, 10,
         number of chains of each type
npptct(1,1) = 10, 0,
         number of particles of each type in the 1st chain
npptct(1,2) = 0, 5,
         number of particles of each type in the 2nd chain
npt = 2,
         number of particle types
txpt = 'bead1', 'bead2',
         labels of both particle types
nppt = 20, 50,
         number of particles of each particle types
natpt = 1, 1,
         number of atom types which particles are composed of
txat = 'site1', 'site2',
         labels of atom types
massat= 10.0, 10.0,
```

CHAPTER 11. APPENDIX 11.2. EXAMPLES OF NAMELISTS DESCRIBING DIFFERENT OBJECTS

Note: Here npptct(1,1) is a simplified version of npptct(1:2,1) etc.

```
particle id
           10 11 15
                         20
 21 0 0 26 0 41
              46 0 0 51 0 66
 0 0
     0
              0 0
                  0
 0 0 ... 0
              0 0 .. 0
 0 0
     0
              0 0
 0 0
      0
              0 0
                  0
```

number of particle types

```
Third generation dendrimers
&nmlParticle
          beginning of the namelist
lclink =.true.,
          flag to enable crosslinking
maxnbondcl = 2, 2, 2, 1,
          maximum number of crosslinks for types of particles
ngen = 3
          number of generations of hierarchichal structure
ictgen(0) = 1,
          chain type of 0th generation
ictgen(1) = 2, 3, 4,
          chain type of 1st, 2nd, and 3rd generation
nbranch = 2, 2, 2,
          number of branches of 1st, 2nd and 3rd generation
ibranchpbeg = 1, 4, 4,
          segment number of the first branch point of 1st, 2nd, and 3rd generation
ibranchpinc = 0, 0, 0,
          segment increment for branch points of 1st, 2nd, and 3rd generation
nct = 4,
          number of chain types
txct ='1-mer1', '4-mer2', '4-mer3', '4-mer4'
          label of each chain type
ncct = 1, 2, 4, 8,
          number of chains of each type
npptct(1,1) = 1, 0, 0, 0,
          number of particles of type 'bead1' in the chain type '1-mer1'. Chain '1-mer1' is composed only
of 'bead1' particles.
npptct(1,2) = 0, 4, 0, 0,
          number of particles of type 'bead2' in the chain type '4-mer2'. Chain '4-mer2' is composed only
of 'bead2' particles.
npptct(1,3) = 0, 0, 4, 0,
          number of particles of type 'bead3' in the chain type '4-mer3'. Chain '4-mer3' is composed only
of 'bead3' particles.
npptct(1,4) = 0, 0, 0, 4,
          number of particles of type 'bead4' in the chain type '4-mer4'. Chain '4-mer4' is composed only
of 'bead4' particles.
npt = 4,
```

```
txpt = 'bead1', 'bead2', 'bead3', 'bead4',
         labels of all particle types
nppt = 1, 8, 16, 32,
         number of particles of each particle types
natpt = 1, 1, 1,
          number of atom types which particles are composed of
txat = 'site1', 'site2', 'site3', 'site4'
          labels of atom types
massat= 10., 10., 10., 10.,
         molar mass of the atom (in grams per mole)
radat = 2.0, 2.0, 2.0, 2.0,
          hard-sphere radius of both atom types
naatpt(1,1) = 1,
          number of 'site1' atoms in 'bead1' particle
txaat(1,1) = 'site1',
          label of 'site1' atoms in 'site1' particle
naatpt(1,2) = 1,
          number of 'site2' atoms in 'bead2' particle
txaat(1,2) = 'site2',
          label of 'site2' atoms in 'site2' particle
naatpt(1,3) = 1,
          number of 'site3' atoms in 'bead3' particle
txaat(1,3) = 'site3',
          label of 'site3' atoms in 'site3' particle
naatpt(1,4) = 1,
          number of 'site4' atoms in 'bead4' particle
txaat(1,4) = 'site4',
          label of 'site4' atoms in 'site4' particle
itestpart = 10,
          test output of chain pointers
          end of the namelist
```

Note: Here nptct(1,1) is a simplified version of nptct(1:2,1) etc.

```
particle id
    30 33 49 46
  13+ # # # # # # # +21
 26# +
                 + #42
  # +
               + #
29# # 10+5 2 1 6 9+18 # #45
    0 0 0 0 x 0 0 0 0
37# # 14+
               +22 # #53
   # +
             + #
  34# +
              + #50
   17+ #### #### +25
    38 41 57 54
```

11.3 Suggestion or Problems?

If you have suggestions or problems with Molsim, open an issue online. Please add the following information to help us help you:

- MOLSIM version number (check by running molsim --version)
- · Host computer (name and configuration)
- Suggestion or problem in as much detail as possible
- · Any Input/Cnf files needed to reproduce the error