**CHAPTER**

**ELEVEN**

**ERRORS**

These doc pages describe many of the error and warning message you can encounter when using LAMMPS. The common problems include conceptual issues. The messages and warnings doc pages give complete lists of all the messages the code may generate, with additional details for many of them.

# Common problems

If two LAMMPS runs do not produce the exact same answer on different machines or different numbers of processors, this is typically not a bug. In theory you should get identical answers on any number of processors and on any machine. In practice, numerical round-off can cause slight differences and eventual divergence of molecular dynamics phase space trajectories within a few 100s or few 1000s of timesteps. However, the statistical properties of the two runs (e.g. average energy or temperature) should still be the same.

If the *velocity* command is used to set initial atom velocities, a particular atom can be assigned a different velocity when the problem is run on a different number of processors or on different machines. If this happens, the phase space trajectories of the two simulations will rapidly diverge. See the discussion of the *loop* option in the *velocity* command for details and options that avoid this issue.

Similarly, the *create\_atoms* command generates a lattice of atoms. For the same physical system, the ordering and numbering of atoms by atom ID may be different depending on the number of processors.

Some commands use random number generators which may be setup to produce different random number streams on each processor and hence will produce different effects when run on different numbers of processors. A commonly-used example is the *fix langevin* command for thermostatting.

A LAMMPS simulation typically has two stages, setup and run. Most LAMMPS errors are detected at setup time; others like a bond stretching too far may not occur until the middle of a run.

LAMMPS tries to flag errors and print informative error messages so you can fix the problem. For most errors it will also print the last input script command that it was processing. Of course, LAMMPS cannot figure out your physics or numerical mistakes, like choosing too big a timestep, specifying erroneous force field coefficients, or putting 2 atoms on top of each other! If you run into errors that LAMMPS does not catch that you think it should flag, please send an email to the [developers](https://www.lammps.org/authors.html) or create an new topic on the dedicated [MatSci forum section](https://matsci.org/lammps/).

If you get an error message about an invalid command in your input script, you can determine what command is causing the problem by looking in the log.lammps file or using the *echo command* to see it on the screen. If you get an error like “Invalid . . . style”, with . . . being fix, compute, pair, etc, it means that you mistyped the style name or that the command is part of an optional package which was not compiled into your executable. The list of available styles in your executable can be listed by using *the -h command-line switch*. The installation and compilation of optional packages is explained on the *Build packages* doc page.

For a given command, LAMMPS expects certain arguments in a specified order. If you mess this up, LAMMPS will often flag the error, but it may also simply read a bogus argument and assign a value that is valid, but not what you wanted. E.g. trying to read the string “abc” as an integer value of 0. Careful reading of the associated doc page for the

command should allow you to fix these problems. In most cases, where LAMMPS expects to read a number, either integer or floating point, it performs a stringent test on whether the provided input actually is an integer or floating- point number, respectively, and reject the input with an error message (for instance, when an integer is required, but a floating-point number 1.0 is provided):

ERROR: Expected integer parameter instead of '1.0' in input script or data file

Some commands allow for using variable references in place of numeric constants so that the value can be evaluated and may change over the course of a run. This is typically done with the syntax *v\_name* for a parameter, where name is the name of the variable. On the other hand, immediate variable expansion with the syntax ${name} is performed while reading the input and before parsing commands,



**Note**

Using a variable reference (i.e. *v\_name*) is only allowed if the documentation of the corresponding command explicitly says it is. Otherwise, you will receive an error message of this kind:

ERROR: Expected floating point parameter instead of 'v\_name' in input script or data file

Generally, LAMMPS will print a message to the screen and logfile and exit gracefully when it encounters a fatal error. Sometimes it will print a WARNING to the screen and logfile and continue on; you can decide if the WARNING is important or not. A WARNING message that is generated in the middle of a run is only printed to the screen, not to the logfile, to avoid cluttering up thermodynamic output. If LAMMPS crashes or hangs without spitting out an error message first then it could be a bug (see [*this section*](#_bookmark0)) or one of the following cases:

LAMMPS runs in the available memory a processor allows to be allocated. Most reasonable MD runs are compute limited, not memory limited, so this should not be a bottleneck on most platforms. Almost all large memory allocations in the code are done via C-style malloc’s which will generate an error message if you run out of memory. Smaller chunks of memory are allocated via C++ “new” statements. If you are unlucky you could run out of memory just when one of these small requests is made, in which case the code will crash or hang (in parallel), since LAMMPS does not trap on those errors.

Illegal arithmetic can cause LAMMPS to run slow or crash. This is typically due to invalid physics and numerics that your simulation is computing. If you see wild thermodynamic values or NaN values in your LAMMPS output, something is wrong with your simulation. If you suspect this is happening, it is a good idea to print out thermodynamic info frequently (e.g. every timestep) via the *thermo* so you can monitor what is happening. Visualizing the atom movement is also a good idea to ensure your model is behaving as you expect.

In parallel, one way LAMMPS can hang is due to how different MPI implementations handle buffering of messages. If the code hangs without an error message, it may be that you need to specify an MPI setting or two (usually via an environment variable) to enable buffering or boost the sizes of messages that can be buffered.

# Error and warning details

Many errors or warnings are self-explanatory and thus straightforward to resolve. However, there are also cases, where there is no single cause and explanation, where LAMMPS can only detect symptoms of an error but not the exact cause, or where the explanation needs to be more detailed than what can be fit into a message printed by the program. The following are discussions of such cases.

* + 1. **Unknown identifier in data file**

This error happens when LAMMPS encounters a line of text with an unexpected keyword while *reading a data file*. This would be either header keywords or section header keywords. This is most commonly due to a mistyped keyword or due to a keyword that is inconsistent with the *atom style* used.

The header section informs LAMMPS how many entries or lines are expected in the various sections (like Atoms, Masses, Pair Coeffs, *etc.*) of the data file. If there is a mismatch, LAMMPS will either keep reading beyond the end of a section or stop reading before the section has ended. In that case the next line will not contain a recognized keyword.

Such a mismatch can also happen when the first line of the data is *not* a comment as required by the format, but a line with a valid header keyword. That would result in LAMMPS expecting, for instance, 0 atoms because the “atoms” header line is the first line and thus treated as a comment.

Another possibility to trigger this error is to have a keyword in the data file that corresponds to a fix (e.g. *fix cmap*) but the *read\_data* command is missing the (optional) arguments that identify the fix and the header keyword and section keyword or those arguments are inconsistent with the keywords in the data file.

* + 1. **Incorrect format in . . . section of data file**

This error happens when LAMMPS reads the contents of a section of a *data file* and the number of parameters in the line differs from what is expected. This most commonly happens, when the atom style is different from what is expected for a specific data file since changing the atom style usually changes the format of the line.

This error can also happen when the number of entries indicated in the header of a data file (e.g. the number of atoms) is larger than the number of lines provided (e.g. in the corresponding Atoms section) and then LAMMPS will continue reading into the next section and that would have a completely different format.

# Reporting bugs

If you are confident that you have found a bug in LAMMPS, please follow the steps outlined below:

* Check the [New features and bug fixes](https://www.lammps.org/bug.html) section of the [LAMMPS WWW site](https://www.lammps.org/) or the [GitHub Releases page](https://github.com/lammps/lammps/releases) to see if the bug has already been addressed in a patch release.
* Check that your issue can be reproduced with the latest development version of LAMMPS.
* Check the manual carefully to verify that the unexpected behavior you are observing is indeed in conflict with the documentation
* Check the [GitHub Issue page](https://github.com/lammps/lammps/issues) if your issue has already been reported and if it is still open.
* Check the [GitHub Pull Requests page](https://github.com/lammps/lammps/pulls) to see if there is already a fix for your bug pending.
* Check the [LAMMPS forum at MatSci](https://matsci.org/lammps/) to see if the issue has been discussed before.

If none of these steps yields any useful information, please file a new bug report on the [GitHub Issue page](https://github.com/lammps/lammps/issues). The website will offer you to select a suitable template with explanations and then you should replace those explanations with the information that you can provide to reproduce your issue.

The most useful thing you can do to help us verify and fix a bug is to isolate the problem. Run it on the smallest number of atoms and fewest number of processors with the simplest input script that reproduces the bug. Try to identify what command or combination of commands is causing the problem and upload the complete input deck as a tar or zip archive. Please avoid using binary restart files unless the issue requires it. In the latter case you should also include an input deck to quickly generate this restart from a data file or a simple additional input. This input deck can be used with tools like a debugger or [valgrind](https://valgrind.org/) to further [*debug the crash*](#_bookmark1).

You may also post a message in the [development category of the LAMMPS forum at MatSci](https://matsci.org/c/lammps/lammps-development/) describing the problem with the same kind of information. The forum can provide a faster response, especially if the bug reported is actually expected behavior or other LAMMPS users have come across it before.

# Debugging crashes

If LAMMPS crashes with a “segmentation fault” or a “bus error” or similar message, then you can use the following two methods to further narrow down the origin of the issue. This will help the LAMMPS developers (or yourself) to understand the reason for the crash and apply a fix (either to the input script or the source code). This requires that your LAMMPS executable includes the required *debug information*. Otherwise it is not possible to look up the names of functions or variables.

The following patch will introduce a bug into the code for pair style *lj/cut* when using the examples/melt/in.melt

input. We use it to show how to identify the origin of a segmentation fault.

--- a/src/pair\_lj\_cut.cpp

+++ b/src/pair\_lj\_cut.cpp

@@ -81,6 +81,7 @@ void PairLJCut::compute(int eflag, int vflag) int nlocal = atom->nlocal;

double \*special\_lj = force->special\_lj; int newton\_pair = force->newton\_pair;

+ double comx = 0.0;

inum = list->inum; ilist = list->ilist;

@@ -134,8 +135,10 @@ void PairLJCut::compute(int eflag, int vflag)

evdwl,0.0,fpair,delx,dely,delz);

}

}

- }

+ comx += atom->rmass[i]\*x[i][0]; /\* BUG \*/

+ }

+ printf("comx = %g\n",comx);

if (vflag\_fdotr) virial\_fdotr\_compute();

}

After recompiling LAMMPS and running the input you should get something like this:

(continues on next page)

$ ./lmp -in in.melt LAMMPS (19 Mar 2020)

using 1 OpenMP thread(s) per MPI task Lattice spacing in x,y,z = 1.6796 1.6796 1.6796

Created orthogonal box = (0 0 0) to (16.796 16.796 16.796)

1 by 1 by 1 MPI processor grid

Created 4000 atoms

create\_atoms CPU = 0.000432253 secs Neighbor list info ...

update every 20 steps, delay 0 steps, check no max neighbors/atom: 2000, page size: 100000 master list distance cutoff = 2.8

ghost atom cutoff = 2.8

(continued from previous page)

binsize = 1.4, bins = 12 12 12

1 neighbor lists, perpetual/occasional/extra = 1 0 0

(1) pair lj/cut, perpetual attributes: half, newton on

pair build: half/bin/atomonly/newton stencil: half/bin/3d/newton

bin: standard Setting up Verlet run ... Unit style : lj Current step : 0

Time step : 0.005 Segmentation fault (core dumped)

* + 1. **Using the GDB debugger to get a stack trace**

There are two options to use the GDB debugger for identifying the origin of the segmentation fault or similar crash. The GDB debugger has many more features and options, as can be seen for example its [online documentation](https://www.sourceware.org/gdb/documentation/).

**Run LAMMPS from within the debugger**

Running LAMMPS under the control of the debugger as shown below only works for a single MPI rank (for debugging a program running in parallel you usually need a parallel debugger program). A simple way to launch GDB is to prefix the LAMMPS command line with gdb --args and then type the command “run” at the GDB prompt. This will launch the debugger, load the LAMMPS executable and its debug info, and then run it. When it reaches the code causing the

segmentation fault, it will stop with a message why it stopped, print the current line of code, and drop back to the GDB prompt.

(gdb) run [...]

Setting up Verlet run ... Unit style : lj Current step : 0 Time step : 0.005

Program received signal SIGSEGV, Segmentation fault.

0x00000000006653ab in LAMMPS\_NS::PairLJCut::compute (this=0x829740, eflag=1, vflag=<optimized␣

*‹→*out>) at /home/akohlmey/compile/lammps/src/pair\_lj\_cut.cpp:139

139 comx += atom->rmass[i]\*x[i][0]; /\* BUG \*/ (gdb)

Now typing the command “where” will show the stack of functions starting from the current function back to “main()”.

(continues on next page)

(gdb) where

#0 0x00000000006653ab in LAMMPS\_NS::PairLJCut::compute (this=0x829740, eflag=1, vflag=

*‹→*<optimized out>) at /home/akohlmey/compile/lammps/src/pair\_lj\_cut.cpp:139

#1 0x00000000004cf0a2 in LAMMPS\_NS::Verlet::setup (this=0x7e6c90, flag=1) at /home/akohlmey/

*‹→*compile/lammps/src/verlet.cpp:131

#2 0x000000000049db42 in LAMMPS\_NS::Run::command (this=this@entry=0x7fffffffcca0,␣

*‹→*narg=narg@entry=1, arg=arg@entry=0x7e8750)

at /home/akohlmey/compile/lammps/src/run.cpp:177

(continued from previous page)

#3 0x000000000041258a in LAMMPS\_NS::Input::command\_creator<LAMMPS\_NS::Run> (lmp=

*‹→*<optimized out>, narg=1, arg=0x7e8750)

at /home/akohlmey/compile/lammps/src/input.cpp:878

#4 0x0000000000410ad3 in LAMMPS\_NS::Input::execute\_command (this=0x7d1410) at /home/

*‹→*akohlmey/compile/lammps/src/input.cpp:864

#5 0x00000000004111fb in LAMMPS\_NS::Input::file (this=0x7d1410) at /home/akohlmey/compile/

*‹→*lammps/src/input.cpp:229

#6 0x000000000040933a in main (argc=<optimized out>, argv=<optimized out>) at /home/akohlmey/

*‹→*compile/lammps/src/main.cpp:65

(gdb)

You can also print the value of variables and see if there is anything unexpected. Segmentation faults, for example, commonly happen when a pointer variable is not assigned and still initialized to NULL.

(gdb) print x

$1 = (double \*\*) 0x7ffff7ca1010 (gdb) print i

$2 = 0

(gdb) print x[0]

$3 = (double \*) 0x7ffff6d80010 (gdb) print x[0][0]

$4 = 0

(gdb) print x[1][0]

$5 = 0.83979809569125363

(gdb) print atom->rmass

$6 = (double \*) 0x0 (gdb)

**Inspect a core dump file with the debugger**

When an executable crashes with a “core dumped” message, it creates a file “core” or “core.<PID#>” which contains the information about the current state. This file may be located in the folder where you ran LAMMPS or in some hidden folder managed by the systemd daemon. In the latter case, you need to “extract” the core file with the coredumpctl

utility to the current folder. Example: coredumpctl -o core dump lmp. Now you can launch the debugger to load the

executable, its debug info and the core dump and drop you to a prompt like before.

$ gdb lmp core

Reading symbols from lmp... [New LWP 1928535]

[Thread debugging using libthread\_db enabled]

Using host libthread\_db library "/lib64/libthread\_db.so.1". Core was generated by `./lmp -in in.melt'.

Program terminated with signal SIGSEGV, Segmentation fault.

#0 0x00000000006653ab in LAMMPS\_NS::PairLJCut::compute (this=0x1b10740, eflag=1, vflag=

*‹→*<optimized out>)

at /home/akohlmey/compile/lammps/src/pair\_lj\_cut.cpp:139

139 comx += atom->rmass[i]\*x[i][0]; /\* BUG \*/ (gdb)

From here on, you use the same commands as shown before to get a stack trace and print current values of (pointer) variables.

* + 1. **Using valgrind to get a stack trace**

The [valgrind](https://valgrind.org/) suite of tools allows to closely inspect the behavior of a compiled program by essentially emulating a CPU and instrumenting the program while running. This slows down execution quite significantly, but can also report issues that are not resulting in a crash. The default valgrind tool is a memory checker and you can use it by prefixing the normal command line with valgrind. Unlike GDB, this will also work for parallel execution, but it is recommended

to redirect the valgrind output to a file (e.g. with --log-file=crash-%p.txt, the %p will be substituted with the process

ID) so that the messages of the multiple valgrind instances to the console are not mixed.

$ valgrind ./lmp -in in.melt

==1933642== Memcheck, a memory error detector

==1933642== Copyright (C) 2002-2017, and GNU GPL'd, by Julian Seward et al.

==1933642== Using Valgrind-3.15.0 and LibVEX; rerun with -h for copyright info

==1933642== Command: ./lmp -in in.melt

==1933642==

LAMMPS (19 Mar 2020)

OMP\_NUM\_THREADS environment is not set. Defaulting to 1 thread. (src/comm.cpp:94) using 1 OpenMP thread(s) per MPI task

Lattice spacing in x,y,z = 1.6796 1.6796 1.6796

Created orthogonal box = (0 0 0) to (16.796 16.796 16.796)

1 by 1 by 1 MPI processor grid

Created 4000 atoms

create\_atoms CPU = 0.032964 secs Neighbor list info ...

update every 20 steps, delay 0 steps, check no max neighbors/atom: 2000, page size: 100000 master list distance cutoff = 2.8

ghost atom cutoff = 2.8 binsize = 1.4, bins = 12 12 12

1 neighbor lists, perpetual/occasional/extra = 1 0 0

(1) pair lj/cut, perpetual attributes: half, newton on

pair build: half/bin/atomonly/newton stencil: half/bin/3d/newton

bin: standard Setting up Verlet run ... Unit style : lj Current step : 0

Time step : 0.005

==1933642== Invalid read of size 8

==1933642== at 0x6653AB: LAMMPS\_NS::PairLJCut::compute(int, int) (pair\_lj\_cut.cpp:139)

==1933642== by 0x4CF0A1: LAMMPS\_NS::Verlet::setup(int) (verlet.cpp:131)

==1933642== by 0x49DB41: LAMMPS\_NS::Run::command(int, char\*\*) (run.cpp:177)

==1933642== by 0x412589: void LAMMPS\_NS::Input::command\_creator<LAMMPS\_NS::Run>

*‹→*(LAMMPS\_NS::LAMMPS\*, int, char\*\*) (input.cpp:881)

==1933642== by 0x410AD2: LAMMPS\_NS::Input::execute\_command() (input.cpp:864)

==1933642== by 0x4111FA: LAMMPS\_NS::Input::file() (input.cpp:229)

==1933642== by 0x409339: main (main.cpp:65)

==1933642== Address 0x0 is not stack'd, malloc'd or (recently) free'd

==1933642==

As you can see, the stack trace information is similar to that obtained from GDB. In addition you get a more specific hint about what cause the segmentation fault, i.e. that it is a NULL pointer dereference. To find out which pointer exactly was NULL, you need to use the debugger, though.

# Error messages

This is an alphabetic list of the ERROR messages LAMMPS prints out and the reason why. If the explanation here is not sufficient, the documentation for the offending command may help. Error messages also list the source file and line number where the error was generated. For example, a message like this:

ERROR: Illegal velocity command (velocity.cpp:78)

means that line #78 in the file src/velocity.cpp generated the error. Looking in the source code may help you figure out what went wrong.

Doc page with [*WARNING messages*](#_bookmark3)

## bond count is inconsistent

An inconsistency was detected when computing the number of 1-3 neighbors for each atom. This likely means something is wrong with the bond topologies you have defined.

## bond count is inconsistent

An inconsistency was detected when computing the number of 1-4 neighbors for each atom. This likely means something is wrong with the bond topologies you have defined.

## Accelerator sharing is not currently supported on system

Multiple MPI processes cannot share the accelerator on your system. For NVIDIA GPUs, see the nvidia-smi command to change this setting.

## All angle coeffs are not set

All angle coefficients must be set in the data file or by the angle\_coeff command before running a simulation.

## All atom IDs = 0 but atom\_modify id = yes

Self-explanatory.

## All atoms of a swapped type must have same charge.

Self-explanatory.

## All atoms of a swapped type must have the same charge.

Self-explanatory.

## All bond coeffs are not set

All bond coefficients must be set in the data file or by the bond\_coeff command before running a simulation.

## All dihedral coeffs are not set

All dihedral coefficients must be set in the data file or by the dihedral\_coeff command before running a simulation.

## All improper coeffs are not set

All improper coefficients must be set in the data file or by the improper\_coeff command before running a simu- lation.

## All masses are not set

For atom styles that define masses for each atom type, all masses must be set in the data file or by the mass command before running a simulation. They must also be set before using the velocity command.

## All mol IDs should be set for fix gcmc group atoms

The molecule flag is on, yet not all molecule ids in the fix group have been set to non-zero positive values by the user. This is an error since all atoms in the fix gcmc group are eligible for deletion, rotation, and translation and therefore must have valid molecule ids.

## All pair coeffs are not set

All pair coefficients must be set in the data file or by the pair\_coeff command before running a simulation.

## All read\_dump x,y,z fields must be specified for scaled, triclinic coords

For triclinic boxes and scaled coordinates you must specify all 3 of the x,y,z fields, else LAMMPS cannot recon- struct the unscaled coordinates.

## All universe/uloop variables must have same # of values

Self-explanatory.

## All variables in next command must be same style

Self-explanatory.

## Angle atom missing in delete\_bonds

The delete\_bonds command cannot find one or more atoms in a particular angle on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid angle.

## Angle atom missing in set command

The set command cannot find one or more atoms in a particular angle on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid angle.

## Angle atoms %d %d %d missing on proc %d at step %ld

One or more of three atoms needed to compute a particular angle are missing on this processor. Typically this is because the pairwise cutoff is set too short or the angle has blown apart and an atom is too far away.

## Angle atoms missing on proc %d at step %ld

One or more of three atoms needed to compute a particular angle are missing on this processor. Typically this is because the pairwise cutoff is set too short or the angle has blown apart and an atom is too far away.

## Angle coeff for hybrid has invalid style

Angle style hybrid uses another angle style as one of its coefficients. The angle style used in the angle\_coeff command or read from a restart file is not recognized.

## Angle coeffs are not set

No angle coefficients have been assigned in the data file or via the angle\_coeff command.

## Angle extent > half of periodic box length

This error was detected by the neigh\_modify check yes setting. It is an error because the angle atoms are so far apart it is ambiguous how it should be defined.

## Angle potential must be defined for SHAKE

When shaking angles, an angle\_style potential must be used.

## Angle style hybrid cannot have hybrid as an argument

Self-explanatory.

## Angle style hybrid cannot have none as an argument

Self-explanatory.

## Angle style hybrid cannot use same angle style twice

Self-explanatory.

## Angle table must range from 0 to 180 degrees

Self-explanatory.

## Angle table parameters did not set N

List of angle table parameters must include N setting.

## Angle\_coeff command before angle\_style is defined

Coefficients cannot be set in the data file or via the angle\_coeff command until an angle\_style has been assigned.

## Angle\_coeff command before simulation box is defined

The angle\_coeff command cannot be used before a read\_data, read\_restart, or create\_box command.

## Angle\_coeff command when no angles allowed

The chosen atom style does not allow for angles to be defined.

## Angle\_style command when no angles allowed

The chosen atom style does not allow for angles to be defined.

## Angles assigned incorrectly

Angles read in from the data file were not assigned correctly to atoms. This means there is something invalid about the topology definitions.

## Angles defined but no angle types

The data file header lists angles but no angle types.

## Append boundary must be shrink/minimum

The boundary style of the face where atoms are added must be of type m (shrink/minimum).

## Arccos of invalid value in variable formula

Argument of arccos() must be between -1 and 1.

## Arcsin of invalid value in variable formula

Argument of arcsin() must be between -1 and 1.

## Assigning body parameters to non-body atom

Self-explanatory.

## Assigning ellipsoid parameters to non-ellipsoid atom

Self-explanatory.

## Assigning line parameters to non-line atom

Self-explanatory.

## Assigning quat to non-body atom

Self-explanatory.

## Assigning tri parameters to non-tri atom

Self-explanatory.

## At least one atom of each swapped type must be present to define charges.

Self-explanatory.

## Atom IDs must be consecutive for velocity create loop all

Self-explanatory.

## Atom IDs must be used for molecular systems

Atom IDs are used to identify and find partner atoms in bonds.

## Atom count changed in fix neb

This is not allowed in a NEB calculation.

## Atom count is inconsistent, cannot write data file

The sum of atoms across processors does not equal the global number of atoms. Probably some atoms have been lost.

## Atom count is inconsistent, cannot write restart file

Sum of atoms across processors does not equal initial total count. This is probably because you have lost some atoms.

## Atom in too many rigid bodies - boost MAXBODY

Fix poems has a parameter MAXBODY (in fix\_poems.cpp) which determines the maximum number of rigid bodies a single atom can belong to (i.e. a multibody joint). The bodies you have defined exceed this limit.

## Atom sort did not operate correctly

This is an internal LAMMPS error. Please report it to the developers.

## Atom style hybrid cannot have hybrid as an argument

Self-explanatory.

## Atom style hybrid cannot use same atom style twice

Self-explanatory.

## Atom style template molecule must have atom types

The defined molecule(s) does not specify atom types.

## Atom style was redefined after using fix property/atom

This is not allowed.

## Atom type must be zero in fix gcmc mol command

Self-explanatory.

## Atom vector in equal-style variable formula

Atom vectors generate one value per atom which is not allowed in an equal-style variable.

## Atom-style variable in equal-style variable formula

Atom-style variables generate one value per atom which is not allowed in an equal-style variable.

## Atom\_modify id command after simulation box is defined

The atom\_modify id command cannot be used after a read\_data, read\_restart, or create\_box command.

## Atom\_modify map command after simulation box is defined

The atom\_modify map command cannot be used after a read\_data, read\_restart, or create\_box command.

## Atom\_modify sort and first options cannot be used together

Self-explanatory.

## Atom\_style command after simulation box is defined

The atom\_style command cannot be used after a read\_data, read\_restart, or create\_box command.

## Atom\_style line can only be used in 2d simulations

Self-explanatory.

## Atom\_style tri can only be used in 3d simulations

Self-explanatory.

## Atomfile variable could not read values

Check the file assigned to the variable.

## Atomfile variable in equal-style variable formula

Self-explanatory.

## Atomfile-style variable in equal-style variable formula

Self-explanatory.

## Attempt to pop empty stack in fix box/relax

Internal LAMMPS error. Please report it to the developers.

## Attempt to push beyond stack limit in fix box/relax

Internal LAMMPS error. Please report it to the developers.

## Attempting to rescale a 0.0 temperature

Cannot rescale a temperature that is already 0.0.

## Attempting to insert more particles than available lattice points

Self-explanatory.

## Bad FENE bond

Two atoms in a FENE bond have become so far apart that the bond cannot be computed.

## Bad TIP4P angle type for PPPM/TIP4P

Specified angle type is not valid.

## Bad TIP4P angle type for PPPMDisp/TIP4P

Specified angle type is not valid.

## Bad TIP4P bond type for PPPM/TIP4P

Specified bond type is not valid.

## Bad TIP4P bond type for PPPMDisp/TIP4P

Specified bond type is not valid.

## Bad fix ID in fix append/atoms command

The value of the fix\_id for keyword spatial must start with “f\_”.

## Bad grid of processors

The 3d grid of processors defined by the processors command does not match the number of processors LAMMPS is being run on.

## Bad kspace\_modify kmax/ewald parameter

Kspace\_modify values for the kmax/ewald keyword must be integers > 0

## Bad kspace\_modify slab parameter

Kspace\_modify value for the slab/volume keyword must be >= 2.0.

## Bad matrix inversion in mldivide3

This error should not occur unless the matrix is badly formed.

## Bad principal moments

Fix rigid did not compute the principal moments of inertia of a rigid group of atoms correctly.

## Bad quadratic solve for particle/line collision

This is an internal error. It should normally not occur.

## Bad quadratic solve for particle/tri collision

This is an internal error. It should normally not occur.

## Bad real space Coulombic cutoff in fix tune/kspace

Fix tune/kspace tried to find the optimal real space Coulombic cutoff using the Newton-Rhaphson method, but found a non-positive or NaN cutoff

## Balance command before simulation box is defined

The balance command cannot be used before a read\_data, read\_restart, or create\_box command.

## Balance produced bad splits

This should not occur. It means two or more cutting plane locations are on top of each other or out of order. Report the problem to the developers.

## Balance rcb cannot be used with comm\_style brick

Comm\_style tiled must be used instead.

## Balance shift string is invalid

The string can only contain the characters “x”, “y”, or “z”.

## Bias compute does not calculate a velocity bias

The specified compute must compute a bias for temperature.

## Bias compute does not calculate temperature

The specified compute must compute temperature.

## Bias compute group does not match compute group

The specified compute must operate on the same group as the parent compute.

## Big particle in fix srd cannot be point particle

Big particles must be extended spheroids or ellipsoids.

## Bigint setting in lmptype.h is invalid

Size of bigint is less than size of tagint.

## Bigint setting in lmptype.h is not compatible

Format of bigint stored in restart file is not consistent with LAMMPS version you are running. See the settings in src/lmptype.h

## Bitmapped lookup tables require int/float be same size

Cannot use pair tables on this machine, because of word sizes. Use the pair\_modify command with table 0 instead.

## Bitmapped table in file does not match requested table

Setting for bitmapped table in pair\_coeff command must match table in file exactly.

## Bitmapped table is incorrect length in table file

Number of table entries is not a correct power of 2.

## Bond and angle potentials must be defined for TIP4P

Cannot use TIP4P pair potential unless bond and angle potentials are defined.

## Bond atom missing in box size check

The second atom needed to compute a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond atom missing in delete\_bonds

The delete\_bonds command cannot find one or more atoms in a particular bond on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid bond.

## Bond atom missing in image check

The second atom in a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond atom missing in set command

The set command cannot find one or more atoms in a particular bond on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid bond.

## Bond atoms %d %d missing on proc %d at step %ld

The second atom needed to compute a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond atoms missing on proc %d at step %ld

The second atom needed to compute a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond coeff for hybrid has invalid style

Bond style hybrid uses another bond style as one of its coefficients. The bond style used in the bond\_coeff command or read from a restart file is not recognized.

## Bond coeffs are not set

No bond coefficients have been assigned in the data file or via the bond\_coeff command.

## Bond extent > half of periodic box length

This error was detected by the neigh\_modify check yes setting. It is an error because the bond atoms are so far apart it is ambiguous how it should be defined.

## Bond potential must be defined for SHAKE

Cannot use fix shake unless bond potential is defined.

## Bond style hybrid cannot have hybrid as an argument

Self-explanatory.

## Bond style hybrid cannot have none as an argument

Self-explanatory.

## Bond style hybrid cannot use same bond style twice

Self-explanatory.

## Bond style quartic cannot be used with 3,4-body interactions

No angle, dihedral, or improper styles can be defined when using bond style quartic.

## Bond style quartic cannot be used with atom style template

This bond style can change the bond topology which is not allowed with this atom style.

## Bond style quartic requires special\_bonds = 1,1,1

This is a restriction of the current bond quartic implementation.

## Bond table parameters did not set N

List of bond table parameters must include N setting.

## Bond table values are not increasing

The values in the tabulated file must be monotonically increasing.

## BondAngle coeff for hybrid angle has invalid format

No “ba” field should appear in data file entry.

## BondBond coeff for hybrid angle has invalid format

No “bb” field should appear in data file entry.

## Bond\_coeff command before bond\_style is defined

Coefficients cannot be set in the data file or via the bond\_coeff command until an bond\_style has been assigned.

## Bond\_coeff command before simulation box is defined

The bond\_coeff command cannot be used before a read\_data, read\_restart, or create\_box command.

## Bond\_coeff command when no bonds allowed

The chosen atom style does not allow for bonds to be defined.

## Bond\_style command when no bonds allowed

The chosen atom style does not allow for bonds to be defined.

## Bonds assigned incorrectly

Bonds read in from the data file were not assigned correctly to atoms. This means there is something invalid about the topology definitions.

## Bonds defined but no bond types

The data file header lists bonds but no bond types.

## Both restart files must use % or neither

Self-explanatory.

## Both restart files must use MPI-IO or neither

Self-explanatory.

## Both sides of boundary must be periodic

Cannot specify a boundary as periodic only on the lo or hi side. Must be periodic on both sides.

## Boundary command after simulation box is defined

The boundary command cannot be used after a read\_data, read\_restart, or create\_box command.

## Box bounds are invalid

The box boundaries specified in the read\_data file are invalid. The lo value must be less than the hi value for all 3 dimensions.

## Box command after simulation box is defined

The box command cannot be used after a read\_data, read\_restart, or create\_box command.

## CPU neighbor lists must be used for ellipsoid/sphere mix.

When using Gay-Berne or RE-squared pair styles with both ellipsoidal and spherical particles, the neighbor list must be built on the CPU

## Can not specify Pxy/Pxz/Pyz in fix box/relax with non-triclinic box

Only triclinic boxes can be used with off-diagonal pressure components. See the region prism command for details.

## Can not specify Pxy/Pxz/Pyz in fix nvt/npt/nph with non-triclinic box

Only triclinic boxes can be used with off-diagonal pressure components. See the region prism command for details.

## Can only use -plog with multiple partitions

Self-explanatory. See page discussion of command-line switches.

## Can only use -pscreen with multiple partitions

Self-explanatory. See page discussion of command-line switches.

## Can only use Kokkos supported regions with Kokkos package

Self-explanatory.

## Can only use NEB with 1-processor replicas

This is current restriction for NEB as implemented in LAMMPS.

## Can only use TAD with 1-processor replicas for NEB

This is current restriction for NEB as implemented in LAMMPS.

## Cannot (yet) do analytic differentiation with pppm/gpu

This is a current restriction of this command.

## Cannot (yet) request ghost atoms with Kokkos half neighbor list

This feature is not yet supported.

## Cannot (yet) use ‘electron’ units with dipoles

This feature is not yet supported.

## Cannot (yet) use Ewald with triclinic box and slab correction

This feature is not yet supported.

## Cannot (yet) use K-space slab correction with compute group/group for triclinic systems

This option is not yet supported.

## Cannot (yet) use MSM with 2d simulation

This feature is not yet supported.

## Cannot (yet) use PPPM with triclinic box and TIP4P

This feature is not yet supported.

## Cannot (yet) use PPPM with triclinic box and kspace\_modify diff ad

This feature is not yet supported.

## Cannot (yet) use PPPM with triclinic box and slab correction

This feature is not yet supported.

## Cannot (yet) use kspace slab correction with long-range dipoles and non-neutral systems or per-atom energy

This feature is not yet supported.

## Cannot (yet) use kspace\_modify diff ad with compute group/group

This option is not yet supported.

## Cannot (yet) use kspace\_style pppm/stagger with triclinic systems

This feature is not yet supported.

## Cannot (yet) use molecular templates with Kokkos

Self-explanatory.

## Cannot (yet) use respa with Kokkos

Self-explanatory.

## Cannot (yet) use rigid bodies with fix deform and Kokkos

Self-explanatory.

## Cannot (yet) use rigid bodies with fix nh and Kokkos

Self-explanatory.

## Cannot (yet) use single precision with MSM (remove -DFFT\_SINGLE from Makefile and re-compile)

Single precision cannot be used with MSM.

## Cannot add atoms to fix move variable

Atoms can not be added afterwards to this fix option.

## Cannot append atoms to a triclinic box

The simulation box must be defined with edges aligned with the Cartesian axes.

## Cannot balance in z dimension for 2d simulation

Self-explanatory.

## Cannot change box ortho/triclinic with certain fixes defined

This is because those fixes store the shape of the box. You need to use unfix to discard the fix, change the box, then redefine a new fix.

## Cannot change box ortho/triclinic with dumps defined

This is because some dumps store the shape of the box. You need to use undump to discard the dump, change the box, then redefine a new dump.

## Cannot change box tilt factors for orthogonal box

Cannot use tilt factors unless the simulation box is non-orthogonal.

## Cannot change box to orthogonal when tilt is non-zero

Self-explanatory.

## Cannot change box z boundary to non-periodic for a 2d simulation

Self-explanatory.

## Cannot change dump\_modify every for dump dcd

The frequency of writing dump dcd snapshots cannot be changed.

## Cannot change dump\_modify every for dump xtc

The frequency of writing dump xtc snapshots cannot be changed.

## Cannot change timestep once fix srd is setup

This is because various SRD properties depend on the timestep size.

## Cannot change timestep with fix pour

This is because fix pour pre-computes the time delay for particles to fall out of the insertion volume due to gravity.

## Cannot change to comm\_style brick from tiled layout

Self-explanatory.

## Cannot change\_box after reading restart file with per-atom info

This is because the restart file info cannot be migrated with the atoms. You can get around this by performing a 0-timestep run which will assign the restart file info to actual atoms.

## Cannot change\_box in xz or yz for 2d simulation

Self-explanatory.

## Cannot change\_box in z dimension for 2d simulation

Self-explanatory.

## Cannot clear group all

This operation is not allowed.

## Cannot close restart file - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot compute initial g\_ewald\_disp

LAMMPS failed to compute an initial guess for the PPPM\_disp g\_ewald\_6 factor that partitions the computation between real space and k-space for Dispersion interactions.

## Cannot create an atom map unless atoms have IDs

The simulation requires a mapping from global atom IDs to local atoms, but the atoms that have been defined have no IDs.

## Cannot create atoms with undefined lattice

Must use the lattice command before using the create\_atoms command.

## Cannot create/grow a vector/array of pointers for %s

LAMMPS code is making an illegal call to the templated memory allocators, to create a vector or array of pointers.

## Cannot create\_atoms after reading restart file with per-atom info

The per-atom info was stored to be used when by a fix that you may re-define. If you add atoms before re-defining the fix, then there will not be a correct amount of per-atom info.

## Cannot create\_box after simulation box is defined

A simulation box can only be defined once.

## Cannot currently use pair reax with pair hybrid

This is not yet supported.

## Cannot currently use pppm/gpu with fix balance.

Self-explanatory.

## Cannot delete group all

Self-explanatory.

## Cannot delete group currently used by a compute

Self-explanatory.

## Cannot delete group currently used by a dump

Self-explanatory.

## Cannot delete group currently used by a fix

Self-explanatory.

## Cannot delete group currently used by atom\_modify first

Self-explanatory.

## Cannot delete\_atoms bond yes for non-molecular systems

Self-explanatory.

## Cannot displace\_atoms after reading restart file with per-atom info

This is because the restart file info cannot be migrated with the atoms. You can get around this by performing a 0-timestep run which will assign the restart file info to actual atoms.

## Cannot do GCMC on atoms in atom\_modify first group

This is a restriction due to the way atoms are organized in a list to enable the atom\_modify first command.

## Cannot do atom/swap on atoms in atom\_modify first group

This is a restriction due to the way atoms are organized in a list to enable the atom\_modify first command.

## Cannot dump sort on atom IDs with no atom IDs defined

Self-explanatory.

## Cannot dump sort when multiple dump files are written

In this mode, each processor dumps its atoms to a file, so no sorting is allowed.

## Cannot embed Python when also extending Python with LAMMPS

When running LAMMPS via Python through the LAMMPS library interface you cannot also user the input script python command.

## Cannot evaporate atoms in atom\_modify first group

This is a restriction due to the way atoms are organized in a list to enable the atom\_modify first command.

## Cannot find create\_bonds group ID

Self-explanatory.

## Cannot find delete\_bonds group ID

Group ID used in the delete\_bonds command does not exist.

## Cannot find specified group ID for core particles

Self-explanatory.

## Cannot find specified group ID for shell particles

Self-explanatory.

## Cannot have both pair\_modify shift and tail set to yes

These 2 options are contradictory.

## Cannot intersect groups using a dynamic group

This operation is not allowed.

## Cannot mix molecular and molecule template atom styles

Self-explanatory.

## Cannot open -reorder file

Self-explanatory.

## Cannot open ADP potential file %s

The specified ADP potential file cannot be opened. Check that the path and name are correct.

## Cannot open AIREBO potential file %s

The specified AIREBO potential file cannot be opened. Check that the path and name are correct.

## Cannot open BOP potential file %s

The specified BOP potential file cannot be opened. Check that the path and name are correct.

## Cannot open COMB potential file %s

The specified COMB potential file cannot be opened. Check that the path and name are correct.

## Cannot open COMB3 lib.comb3 file

The COMB3 library file cannot be opened. Check that the path and name are correct.

## Cannot open COMB3 potential file %s

The specified COMB3 potential file cannot be opened. Check that the path and name are correct.

## Cannot open EAM potential file %s

The specified EAM potential file cannot be opened. Check that the path and name are correct.

## Cannot open EIM potential file %s

The specified EIM potential file cannot be opened. Check that the path and name are correct.

## Cannot open LCBOP potential file %s

The specified LCBOP potential file cannot be opened. Check that the path and name are correct.

## Cannot open MEAM potential file %s

The specified MEAM potential file cannot be opened. Check that the path and name are correct.

## Cannot open SNAP coefficient file %s

The specified SNAP coefficient file cannot be opened. Check that the path and name are correct.

## Cannot open SNAP parameter file %s

The specified SNAP parameter file cannot be opened. Check that the path and name are correct.

## Cannot open Stillinger-Weber potential file %s

The specified SW potential file cannot be opened. Check that the path and name are correct.

## Cannot open Tersoff potential file %s

The specified potential file cannot be opened. Check that the path and name are correct.

## Cannot open Vashishta potential file %s

The specified Vashishta potential file cannot be opened. Check that the path and name are correct.

## Cannot open balance output file

Self-explanatory.

## Cannot open coul/streitz potential file %s

The specified coul/streitz potential file cannot be opened. Check that the path and name are correct.

## Cannot open custom file

Self-explanatory.

## Cannot open data file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open dir to search for restart file

Using a “\*” in the name of the restart file will open the current directory to search for matching file names.

## Cannot open dump file

Self-explanatory.

## Cannot open dump file %s

The output file for the dump command cannot be opened. Check that the path and name are correct.

## Cannot open file %s

The specified file cannot be opened. Check that the path and name are correct. If the file is a compressed file, also check that the gzip executable can be found and run.

## Cannot open file variable file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix ave/chunk file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix ave/correlate file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix ave/histo file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix ave/time file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix balance output file

Self-explanatory.

## Cannot open fix poems file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix print file %s

The output file generated by the fix print command cannot be opened

## Cannot open fix qeq parameter file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix qeq/comb file %s

The output file for the fix qeq/combs command cannot be opened. Check that the path and name are correct.

## Cannot open fix reax/bonds file %s

The output file for the fix reax/bonds command cannot be opened. Check that the path and name are correct.

## Cannot open fix rigid infile %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix rigid restart file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix rigid/small infile %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open fix tmd file %s

The output file for the fix tmd command cannot be opened. Check that the path and name are correct.

## Cannot open fix ttm file %s

The output file for the fix ttm command cannot be opened. Check that the path and name are correct.

## Cannot open gzipped file

LAMMPS was compiled without support for reading and writing gzipped files through a pipeline to the gzip program with -DLAMMPS\_GZIP.

## Cannot open input script %s

Self-explanatory.

## Cannot open log.cite file

This file is created when you use some LAMMPS features, to indicate what paper you should cite on behalf of those who implemented the feature. Check that you have write privileges into the directory you are running in.

## Cannot open log.lammps for writing

The default LAMMPS log file cannot be opened. Check that the directory you are running in allows for files to be created.

## Cannot open logfile

The LAMMPS log file named in a command-line argument cannot be opened. Check that the path and name are correct.

## Cannot open logfile %s

The LAMMPS log file specified in the input script cannot be opened. Check that the path and name are correct.

## Cannot open molecule file %s

The specified file cannot be opened. Check that the path and name are correct.

## Cannot open nb3b/harmonic potential file %s

The specified potential file cannot be opened. Check that the path and name are correct.

## Cannot open pair\_write file

The specified output file for pair energies and forces cannot be opened. Check that the path and name are correct.

## Cannot open polymorphic potential file %s

The specified polymorphic potential file cannot be opened. Check that the path and name are correct.

## Cannot open print file %s

Self-explanatory.

## Cannot open processors output file

Self-explanatory.

## Cannot open restart file %s

Self-explanatory.

## Cannot open restart file for reading - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot open restart file for writing - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot open screen file

The screen file specified as a command-line argument cannot be opened. Check that the directory you are running in allows for files to be created.

## Cannot open temporary file for world counter.

Self-explanatory.

## Cannot open universe log file

For a multi-partition run, the master log file cannot be opened. Check that the directory you are running in allows for files to be created.

## Cannot open universe screen file

For a multi-partition run, the master screen file cannot be opened. Check that the directory you are running in allows for files to be created.

## Cannot read from restart file - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot read\_data without add keyword after simulation box is defined

Self-explanatory.

## Cannot read\_restart after simulation box is defined

The read\_restart command cannot be used after a read\_data, read\_restart, or create\_box command.

## Cannot redefine variable as a different style

An equal-style variable can be re-defined but only if it was originally an equal-style variable.

## Cannot replicate 2d simulation in z dimension

The replicate command cannot replicate a 2d simulation in the z dimension.

## Cannot replicate with fixes that store atom quantities

Either fixes are defined that create and store atom-based vectors or a restart file was read which included atom- based vectors for fixes. The replicate command cannot duplicate that information for new atoms. You should use the replicate command before fixes are applied to the system.

## Cannot reset timestep with a dynamic region defined

Dynamic regions (see the region command) have a time dependence. Thus you cannot change the timestep when one or more of these are defined.

## Cannot reset timestep with a time-dependent fix defined

You cannot reset the timestep when a fix that keeps track of elapsed time is in place.

## Cannot run 2d simulation with non-periodic Z dimension

Use the boundary command to make the z dimension periodic in order to run a 2d simulation.

## Cannot set bond topology types for atom style template

The bond, angle, etc types cannot be changed for this atom style since they are static settings in the molecule template files.

## Cannot set both respa pair and inner/middle/outer

In the rRESPA integrator, you must compute pairwise potentials either all together (pair), or in pieces (in- ner/middle/outer). You can’t do both.

## Cannot set cutoff/multi before simulation box is defined

Self-explanatory.

## Cannot set dpd/theta for this atom style

Self-explanatory.

## Cannot set dump\_modify flush for dump xtc

Self-explanatory.

## Cannot set mass for this atom style

This atom style does not support mass settings for each atom type. Instead they are defined on a per-atom basis in the data file.

## Cannot set meso/cv for this atom style

Self-explanatory.

## Cannot set meso/e for this atom style

Self-explanatory.

## Cannot set meso/rho for this atom style

Self-explanatory.

## Cannot set non-zero image flag for non-periodic dimension

Self-explanatory.

## Cannot set non-zero z velocity for 2d simulation

Self-explanatory.

## Cannot set quaternion for atom that has none

Self-explanatory.

## Cannot set quaternion with xy components for 2d system

Self-explanatory.

## Cannot set respa hybrid and any of pair/inner/middle/outer

In the rRESPA integrator, you must compute pairwise potentials either all together (pair), with different cutoff regions (inner/middle/outer), or per hybrid sub-style (hybrid). You cannot mix those.

## Cannot set respa middle without inner/outer

In the rRESPA integrator, you must define both a inner and outer setting in order to use a middle setting.

## Cannot set restart file size - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot set smd/contact/radius for this atom style

Self-explanatory.

## Cannot set smd/mass/density for this atom style

Self-explanatory.

## Cannot set temperature for fix rigid/nph

The temp keyword cannot be specified.

## Cannot set theta for atom that is not a line

Self-explanatory.

## Cannot set this attribute for this atom style

The attribute being set does not exist for the defined atom style.

## Cannot set variable z velocity for 2d simulation

Self-explanatory.

## Cannot skew triclinic box in z for 2d simulation

Self-explanatory.

## Cannot subtract groups using a dynamic group

This operation is not allowed.

## Cannot union groups using a dynamic group

This operation is not allowed.

## Cannot use -kokkos on without KOKKOS installed

Self-explanatory.

## Cannot use -reorder after -partition

Self-explanatory. See page discussion of command-line switches.

## Cannot use Ewald with 2d simulation

The kspace style ewald cannot be used in 2d simulations. You can use 2d Ewald in a 3d simulation; see the kspace\_modify command.

## Cannot use Ewald/disp solver on system with no charge, dipole, or LJ particles

No atoms in system have a non-zero charge or dipole, or are LJ particles. Change charges/dipoles or change options of the kspace solver/pair style.

## Cannot use EwaldDisp with 2d simulation

This is a current restriction of this command.

## Cannot use Kokkos pair style with rRESPA inner/middle

Self-explanatory.

## Cannot use NEB unless atom map exists

Use the atom\_modify command to create an atom map.

## Cannot use NEB with a single replica

Self-explanatory.

## Cannot use NEB with atom\_modify sort enabled

This is current restriction for NEB implemented in LAMMPS.

## Cannot use PPPM with 2d simulation

The kspace style pppm cannot be used in 2d simulations. You can use 2d PPPM in a 3d simulation; see the kspace\_modify command.

## Cannot use PPPMDisp with 2d simulation

The kspace style pppm/disp cannot be used in 2d simulations. You can use 2d pppm/disp in a 3d simulation; see the kspace\_modify command.

## Cannot use PRD with a changing box

The current box dimensions are not copied between replicas

## Cannot use PRD with a time-dependent fix defined

PRD alters the timestep in ways that will mess up these fixes.

## Cannot use PRD with a time-dependent region defined

PRD alters the timestep in ways that will mess up these regions.

## Cannot use PRD with atom\_modify sort enabled

This is a current restriction of PRD. You must turn off sorting, which is enabled by default, via the atom\_modify command.

## Cannot use PRD with multi-processor replicas unless atom map exists

Use the atom\_modify command to create an atom map.

## Cannot use TAD unless atom map exists for NEB

See atom\_modify map command to set this.

## Cannot use TAD with a single replica for NEB

NEB requires multiple replicas.

## Cannot use TAD with atom\_modify sort enabled for NEB

This is a current restriction of NEB.

## Cannot use a damped dynamics min style with fix box/relax

This is a current restriction in LAMMPS. Use another minimizer style.

## Cannot use a damped dynamics min style with per-atom DOF

This is a current restriction in LAMMPS. Use another minimizer style.

## Cannot use append/atoms in periodic dimension

The boundary style of the face where atoms are added can not be of type p (periodic).

## Cannot use atomfile-style variable unless atom map exists

Self-explanatory. See the atom\_modify command to create a map.

## Cannot use both com and bias with compute temp/chunk

Self-explanatory.

## Cannot use chosen neighbor list style with buck/coul/cut/kk

Self-explanatory.

## Cannot use chosen neighbor list style with buck/coul/long/kk

Self-explanatory.

## Cannot use chosen neighbor list style with buck/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with coul/cut/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with coul/debye/kk

Self-explanatory.

## Cannot use chosen neighbor list style with coul/dsf/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with coul/wolf/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with lj/charmm/coul/charmm/implicit/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/charmm/coul/charmm/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/charmm/coul/long/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/class2/coul/cut/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/class2/coul/long/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/class2/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/cut/coul/cut/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with lj/cut/coul/debye/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/cut/coul/long/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with lj/cut/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with lj/expand/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/gromacs/coul/gromacs/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/gromacs/kk

Self-explanatory.

## Cannot use chosen neighbor list style with lj/spica/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with pair eam/kk

That style is not supported by Kokkos.

## Cannot use chosen neighbor list style with pair eam/kk/alloy

Self-explanatory.

## Cannot use chosen neighbor list style with pair eam/kk/fs

Self-explanatory.

## Cannot use chosen neighbor list style with pair sw/kk

Self-explanatory.

## Cannot use chosen neighbor list style with tersoff/kk

Self-explanatory.

## Cannot use chosen neighbor list style with tersoff/zbl/kk

Self-explanatory.

## Cannot use compute chunk/atom bin z for 2d model

Self-explanatory.

## Cannot use compute cluster/atom unless atoms have IDs

Atom IDs are used to identify clusters.

## Cannot use create\_atoms rotate unless single style

Self-explanatory.

## Cannot use create\_bonds unless atoms have IDs

This command requires a mapping from global atom IDs to local atoms, but the atoms that have been defined have no IDs.

## Cannot use create\_bonds with non-molecular system

Self-explanatory.

## Cannot use cwiggle in variable formula between runs

This is a function of elapsed time.

## Cannot use delete\_atoms bond yes with atom\_style template

This is because the bonds for that atom style are hardwired in the molecule template.

## Cannot use delete\_atoms unless atoms have IDs

Your atoms do not have IDs, so the delete\_atoms command cannot be used.

## Cannot use delete\_bonds with non-molecular system

Your choice of atom style does not have bonds.

## Cannot use dump\_modify fileper without % in dump file name

Self-explanatory.

## Cannot use dump\_modify nfile without % in dump file name

Self-explanatory.

## Cannot use dynamic group with fix adapt atom

This is not yet supported.

## Cannot use fix TMD unless atom map exists

Using this fix requires the ability to lookup an atom index, which is provided by an atom map. An atom map does not exist (by default) for non-molecular problems. Using the atom\_modify map command will force an atom map to be created.

## Cannot use fix bond/break with non-molecular systems

Only systems with bonds that can be changed can be used. Atom\_style template does not qualify.

## Cannot use fix bond/create with non-molecular systems

Only systems with bonds that can be changed can be used. Atom\_style template does not qualify.

## Cannot use fix bond/swap with non-molecular systems

Only systems with bonds that can be changed can be used. Atom\_style template does not qualify.

## Cannot use fix box/relax on a 2nd non-periodic dimension

When specifying an off-diagonal pressure component, the second of the two dimensions must be periodic. E.g. if the xy component is specified, then the y dimension must be periodic.

## Cannot use fix box/relax on a non-periodic dimension

When specifying a diagonal pressure component, the dimension must be periodic.

## Cannot use fix box/relax with both relaxation and scaling on a tilt factor

When specifying scaling on a tilt factor component, that component can not also be controlled by the barostat.

E.g. if scalexy yes is specified and also keyword tri or xy, this is wrong.

## Cannot use fix box/relax with tilt factor scaling on a 2nd non-periodic dimension

When specifying scaling on a tilt factor component, the second of the two dimensions must be periodic. E.g. if the xy component is specified, then the y dimension must be periodic.

## Cannot use fix deform on a shrink-wrapped boundary

The x, y, z options cannot be applied to shrink-wrapped dimensions.

## Cannot use fix deform tilt on a shrink-wrapped 2nd dim

This is because the shrink-wrapping will change the value of the strain implied by the tilt factor.

## Cannot use fix deform trate on a box with zero tilt

The trate style alters the current strain.

## Cannot use fix deposit rigid and not molecule

Self-explanatory.

## Cannot use fix deposit rigid and shake

These two attributes are conflicting.

## Cannot use fix deposit shake and not molecule

Self-explanatory.

## Cannot use fix enforce2d with 3d simulation

Self-explanatory.

## Cannot use fix gcmc in a 2d simulation

Fix gcmc is set up to run in 3d only. No 2d simulations with fix gcmc are allowed.

## Cannot use fix gcmc shake and not molecule

Self-explanatory.

## Cannot use fix msst without per-type mass defined

Self-explanatory.

## Cannot use fix npt and fix deform on same component of stress tensor

This would be changing the same box dimension twice.

## Cannot use fix nvt/npt/nph on a 2nd non-periodic dimension

When specifying an off-diagonal pressure component, the second of the two dimensions must be periodic. E.g. if the xy component is specified, then the y dimension must be periodic.

## Cannot use fix nvt/npt/nph on a non-periodic dimension

When specifying a diagonal pressure component, the dimension must be periodic.

## Cannot use fix nvt/npt/nph with both xy dynamics and xy scaling

Self-explanatory.

## Cannot use fix nvt/npt/nph with both xz dynamics and xz scaling

Self-explanatory.

## Cannot use fix nvt/npt/nph with both yz dynamics and yz scaling

Self-explanatory.

## Cannot use fix nvt/npt/nph with xy scaling when y is non-periodic dimension

The second dimension in the barostatted tilt factor must be periodic.

## Cannot use fix nvt/npt/nph with xz scaling when z is non-periodic dimension

The second dimension in the barostatted tilt factor must be periodic.

## Cannot use fix nvt/npt/nph with yz scaling when z is non-periodic dimension

The second dimension in the barostatted tilt factor must be periodic.

## Cannot use fix pour rigid and not molecule

Self-explanatory.

## Cannot use fix pour rigid and shake

These two attributes are conflicting.

## Cannot use fix pour shake and not molecule

Self-explanatory.

## Cannot use fix pour with triclinic box

This option is not yet supported.

## Cannot use fix press/berendsen and fix deform on same component of stress tensor

These commands both change the box size/shape, so you cannot use both together.

## Cannot use fix press/berendsen on a non-periodic dimension

Self-explanatory.

## Cannot use fix press/berendsen with triclinic box

Self-explanatory.

## Cannot use fix reax/bonds without pair\_style reax

Self-explanatory.

## Cannot use fix rigid npt/nph and fix deform on same component of stress tensor

This would be changing the same box dimension twice.

## Cannot use fix rigid npt/nph on a non-periodic dimension

When specifying a diagonal pressure component, the dimension must be periodic.

## Cannot use fix rigid/small npt/nph on a non-periodic dimension

When specifying a diagonal pressure component, the dimension must be periodic.

## Cannot use fix shake with non-molecular system

Your choice of atom style does not have bonds.

## Cannot use fix ttm with 2d simulation

This is a current restriction of this fix due to the grid it creates.

## Cannot use fix ttm with triclinic box

This is a current restriction of this fix due to the grid it creates.

## Cannot use fix tune/kspace without a kspace style

Self-explanatory.

## Cannot use fix tune/kspace without a pair style

This fix (tune/kspace) can only be used when a pair style has been specified.

## Cannot use fix wall in periodic dimension

Self-explanatory.

## Cannot use fix wall zlo/zhi for a 2d simulation

Self-explanatory.

## Cannot use fix wall/reflect in periodic dimension

Self-explanatory.

## Cannot use fix wall/reflect zlo/zhi for a 2d simulation

Self-explanatory.

## Cannot use fix wall/srd in periodic dimension

Self-explanatory.

## Cannot use fix wall/srd more than once

Nor is their a need to since multiple walls can be specified in one command.

## Cannot use fix wall/srd without fix srd

Self-explanatory.

## Cannot use fix wall/srd zlo/zhi for a 2d simulation

Self-explanatory.

## Cannot use fix\_deposit unless atoms have IDs

Self-explanatory.

## Cannot use fix\_pour unless atoms have IDs

Self-explanatory.

## Cannot use include command within an if command

Self-explanatory.

## Cannot use lines with fix srd unless overlap is set

This is because line segments are connected to each other.

## Cannot use multiple fix wall commands with pair brownian

Self-explanatory.

## Cannot use multiple fix wall commands with pair lubricate

Self-explanatory.

## Cannot use multiple fix wall commands with pair lubricate/poly

Self-explanatory.

## Cannot use multiple fix wall commands with pair lubricateU

Self-explanatory.

## Cannot use neigh\_modify exclude with GPU neighbor builds

This is a current limitation of the GPU implementation in LAMMPS.

## Cannot use neighbor bins - box size << cutoff

Too many neighbor bins will be created. This typically happens when the simulation box is very small in some dimension, compared to the neighbor cutoff. Use the “nsq” style instead of “bin” style.

## Cannot use newton pair with beck/gpu pair style

Self-explanatory.

## Cannot use newton pair with born/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with born/coul/wolf/gpu pair style

Self-explanatory.

## Cannot use newton pair with born/gpu pair style

Self-explanatory.

## Cannot use newton pair with buck/coul/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with buck/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with buck/gpu pair style

Self-explanatory.

## Cannot use newton pair with colloid/gpu pair style

Self-explanatory.

## Cannot use newton pair with coul/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with coul/debye/gpu pair style

Self-explanatory.

## Cannot use newton pair with coul/dsf/gpu pair style

Self-explanatory.

## Cannot use newton pair with coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with dipole/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with dipole/sf/gpu pair style

Self-explanatory.

## Cannot use newton pair with dpd/gpu pair style

Self-explanatory.

## Cannot use newton pair with dpd/tstat/gpu pair style

Self-explanatory.

## Cannot use newton pair with eam/alloy/gpu pair style

Self-explanatory.

## Cannot use newton pair with eam/fs/gpu pair style

Self-explanatory.

## Cannot use newton pair with eam/gpu pair style

Self-explanatory.

## Cannot use newton pair with gauss/gpu pair style

Self-explanatory.

## Cannot use newton pair with gayberne/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/charmm/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/class2/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/class2/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cubic/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/coul/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/coul/debye/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/coul/dsf/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/coul/msm/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/expand/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/gromacs/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/spica/coul/long/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj/spica/gpu pair style

Self-explanatory.

## Cannot use newton pair with lj96/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with mie/cut/gpu pair style

Self-explanatory.

## Cannot use newton pair with morse/gpu pair style

Self-explanatory.

## Cannot use newton pair with resquared/gpu pair style

Self-explanatory.

## Cannot use newton pair with soft/gpu pair style

Self-explanatory.

## Cannot use newton pair with table/gpu pair style

Self-explanatory.

## Cannot use newton pair with yukawa/colloid/gpu pair style

Self-explanatory.

## Cannot use newton pair with yukawa/gpu pair style

Self-explanatory.

## Cannot use newton pair with zbl/gpu pair style

Self-explanatory.

## Cannot use non-zero forces in an energy minimization

Fix setforce cannot be used in this manner. Use fix addforce instead.

## Cannot use non-periodic boundares with fix ttm

This fix requires a fully periodic simulation box.

## Cannot use non-periodic boundaries with Ewald

For kspace style ewald, all 3 dimensions must have periodic boundaries unless you use the kspace\_modify com- mand to define a 2d slab with a non-periodic z dimension.

## Cannot use non-periodic boundaries with EwaldDisp

For kspace style ewald/disp, all 3 dimensions must have periodic boundaries unless you use the kspace\_modify command to define a 2d slab with a non-periodic z dimension.

## Cannot use non-periodic boundaries with PPPM

For kspace style pppm, all 3 dimensions must have periodic boundaries unless you use the kspace\_modify com- mand to define a 2d slab with a non-periodic z dimension.

## Cannot use non-periodic boundaries with PPPMDisp

For kspace style pppm/disp, all 3 dimensions must have periodic boundaries unless you use the kspace\_modify command to define a 2d slab with a non-periodic z dimension.

## Cannot use order greater than 8 with pppm/gpu.

Self-explanatory.

## Cannot use package gpu neigh yes with triclinic box

This is a current restriction in LAMMPS.

## Cannot use pair tail corrections with 2d simulations

The correction factors are only currently defined for 3d systems.

## Cannot use processors part command without using partitions

See the command-line -partition switch.

## Cannot use ramp in variable formula between runs

This is because the ramp() function is time dependent.

## Cannot use read\_data add before simulation box is defined

Self-explanatory.

## Cannot use read\_data extra with add flag

Self-explanatory.

## Cannot use read\_data offset without add flag

Self-explanatory.

## Cannot use read\_data shift without add flag

Self-explanatory.

## Cannot use region INF or EDGE when box does not exist

Regions that extend to the box boundaries can only be used after the create\_box command has been used.

## Cannot use set atom with no atom IDs defined

Atom IDs are not defined, so they cannot be used to identify an atom.

## Cannot use set mol with no molecule IDs defined

Self-explanatory.

## Cannot use swiggle in variable formula between runs

This is a function of elapsed time.

## Cannot use tris with fix srd unless overlap is set

This is because triangles are connected to each other.

## Cannot use variable energy with constant efield in fix efield

LAMMPS computes the energy itself when the E-field is constant.

## Cannot use variable energy with constant force in fix addforce

This is because for constant force, LAMMPS can compute the change in energy directly.

## Cannot use variable every setting for dump dcd

The format of DCD dump files requires snapshots be output at a constant frequency.

## Cannot use variable every setting for dump xtc

The format of this file requires snapshots at regular intervals.

## Cannot use vdisplace in variable formula between runs

This is a function of elapsed time.

## Cannot use velocity bias command without temp keyword

Self-explanatory.

## Cannot use velocity create loop all unless atoms have IDs

Atoms in the simulation to do not have IDs, so this style of velocity creation cannot be performed.

## Cannot use wall in periodic dimension

Self-explanatory.

## Cannot use write\_restart fileper without % in restart file name

Self-explanatory.

## Cannot use write\_restart nfile without % in restart file name

Self-explanatory.

## Cannot wiggle and shear fix wall/gran

Cannot specify both options at the same time.

## Cannot write to restart file - MPI error: %s

This error was generated by MPI when reading/writing an MPI-IO restart file.

## Cannot yet use KSpace solver with grid with comm style tiled

This is current restriction in LAMMPS.

## Cannot yet use comm\_style tiled with multi-mode comm

Self-explanatory.

## Cannot yet use comm\_style tiled with triclinic box

Self-explanatory.

## Cannot yet use compute tally with Kokkos

This feature is not yet supported.

## Cannot yet use fix bond/break with this improper style

This is a current restriction in LAMMPS.

## Cannot yet use fix bond/create with this improper style

This is a current restriction in LAMMPS.

## Cannot yet use minimize with Kokkos

This feature is not yet supported.

## Cannot yet use pair hybrid with Kokkos

This feature is not yet supported.

## Cannot zero Langevin force of 0 atoms

The group has zero atoms, so you cannot request its force be zeroed.

## Cannot zero gld force for zero atoms

There are no atoms currently in the group.

## Cannot zero momentum of no atoms

Self-explanatory.

## Change\_box command before simulation box is defined

Self-explanatory.

## Change\_box volume used incorrectly

The “dim volume” option must be used immediately following one or two settings for “dim1 . . . ” (and optionally “dim2 . . . ”) and must be for a different dimension, i.e. dim != dim1 and dim != dim2.

## Chunk/atom compute does not exist for compute angmom/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute com/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute gyration/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute inertia/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute msd/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute omega/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute property/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute temp/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute torque/chunk

Self-explanatory.

## Chunk/atom compute does not exist for compute vcm/chunk

Self-explanatory.

## Chunk/atom compute does not exist for fix ave/chunk

Self-explanatory.

## Comm tiled invalid index in box drop brick

Internal error check in comm\_style tiled which should not occur. Contact the developers.

## Comm tiled mis-match in box drop brick

Internal error check in comm\_style tiled which should not occur. Contact the developers.

## Comm\_modify group != atom\_modify first group

Self-explanatory.

## Communication cutoff for comm\_style tiled cannot exceed periodic box length

Self-explanatory.

## Communication cutoff too small for SNAP micro load balancing

This can happen if you change the neighbor skin after your pair\_style command or if your box dimensions grow during a run. You can set the cutoff explicitly via the comm\_modify cutoff command.

## Compute %s does not allow use of dynamic group

Dynamic groups have not yet been enabled for this compute.

## Compute for fix pafi does not calculate a local array

Self-explanatory.

## Compute for fix pafi must have 9 fields per atom

Self-explanatory.

## Compute ID for compute chunk /atom does not exist

Self-explanatory.

## Compute ID for compute chunk/atom does not exist

Self-explanatory.

## Compute gyration ID does not exist for compute gyration/shape

Self-explanatory. Provide a valid compute ID.

## Compute gyration/shape compute ID does not point to a gyration compute

Self-explanatory. Provide and ID of a compute gyration command.

## Compute ID for compute reduce does not exist

Self-explanatory.

## Compute ID for compute slice does not exist

Self-explanatory.

## Compute ID for fix ave/atom does not exist

Self-explanatory.

## Compute ID for fix ave/chunk does not exist

Self-explanatory.

## Compute ID for fix ave/correlate does not exist

Self-explanatory.

## Compute ID for fix ave/histo does not exist

Self-explanatory.

## Compute ID for fix ave/time does not exist

Self-explanatory.

## Compute ID for fix numdiff does not exist

Self-explanatory.

## Compute ID for fix numdiff/virial does not exist

Self-explanatory.

## Compute ID for fix store/state does not exist

Self-explanatory.

## Compute ID for fix vector does not exist

Self-explanatory.

## Compute ID must be alphanumeric or underscore characters

Self-explanatory.

## Compute angle/local used when angles are not allowed

The atom style does not support angles.

## Compute angmom/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute body/local requires atom style body

Self-explanatory.

## Compute bond/local used when bonds are not allowed

The atom style does not support bonds.

## Compute centro/atom requires a pair style be defined

This is because the computation of the centro-symmetry values uses a pairwise neighbor list.

## Compute chunk/atom bin/cylinder radius is too large for periodic box

Radius cannot be bigger than 1/2 of a non-axis periodic dimension.

## Compute chunk/atom bin/sphere radius is too large for periodic box

Radius cannot be bigger than 1/2 of any periodic dimension.

## Compute chunk/atom compute array is accessed out-of-range

The index for the array is out of bounds.

## Compute chunk/atom compute does not calculate a per-atom array

Self-explanatory.

## Compute chunk/atom compute does not calculate a per-atom vector

Self-explanatory.

## Compute chunk/atom compute does not calculate per-atom values

Self-explanatory.

## Compute chunk/atom cylinder axis must be z for 2d

Self-explanatory.

## Compute chunk/atom fix array is accessed out-of-range

The index for the array is out of bounds.

## Compute chunk/atom fix does not calculate a per-atom array

Self-explanatory.

## Compute chunk/atom fix does not calculate a per-atom vector

Self-explanatory.

## Compute chunk/atom fix does not calculate per-atom values

Self-explanatory.

## Compute chunk/atom for triclinic boxes requires units reduced

Self-explanatory.

## Compute chunk/atom ids once but nchunk is not once

You cannot assign chunks IDs to atom permanently if the number of chunks may change.

## Compute chunk/atom molecule for non-molecular system

Self-explanatory.

## Compute chunk/atom sphere z origin must be 0.0 for 2d

Self-explanatory.

## Compute chunk/atom stores no IDs for compute property/chunk

It will only store IDs if its compress option is enabled.

## Compute chunk/atom stores no coord1 for compute property/chunk

Only certain binning options for compute chunk/atom store coordinates.

## Compute chunk/atom stores no coord2 for compute property/chunk

Only certain binning options for compute chunk/atom store coordinates.

## Compute chunk/atom stores no coord3 for compute property/chunk

Only certain binning options for compute chunk/atom store coordinates.

## Compute chunk/atom variable is not atom-style variable

Self-explanatory.

## Compute chunk/atom without bins cannot use discard mixed

That discard option only applies to the binning styles.

## Compute cluster/atom cutoff is longer than pairwise cutoff

Cannot identify clusters beyond cutoff.

## Compute cluster/atom requires a pair style be defined

This is so that the pair style defines a cutoff distance which is used to find clusters.

## Compute cna/atom cutoff is longer than pairwise cutoff

Self-explanatory.

## Compute cna/atom requires a pair style be defined

Self-explanatory.

## Compute com/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute contact/atom requires a pair style be defined

Self-explanatory.

## Compute contact/atom requires atom style sphere

Self-explanatory.

## Compute coord/atom cutoff is longer than pairwise cutoff

Cannot compute coordination at distances longer than the pair cutoff, since those atoms are not in the neighbor list.

## Compute coord/atom requires a pair style be defined

Self-explanatory.

## Compute damage/atom requires peridynamic potential

Damage is a Peridynamic-specific metric. It requires you to be running a Peridynamics simulation.

## Compute dihedral/local used when dihedrals are not allowed

The atom style does not support dihedrals.

## Compute dilatation/atom cannot be used with this pair style

Self-explanatory.

## Compute dilatation/atom requires Peridynamic pair style

Self-explanatory.

## Compute does not allow an extra compute or fix to be reset

This is an internal LAMMPS error. Please report it to the developers.

## Compute erotate/asphere requires atom style ellipsoid or line or tri

Self-explanatory.

## Compute erotate/asphere requires extended particles

This compute cannot be used with point particles.

## Compute erotate/rigid with non-rigid fix-ID

Self-explanatory.

## Compute erotate/sphere requires atom style sphere

Self-explanatory.

## Compute erotate/sphere/atom requires atom style sphere

Self-explanatory.

## Compute event/displace has invalid fix event assigned

This is an internal LAMMPS error. Please report it to the developers.

## Compute group/group group ID does not exist

Self-explanatory.

## Compute gyration/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute heat/flux compute ID does not compute ke/atom

Self-explanatory.

## Compute heat/flux compute ID does not compute pe/atom

Self-explanatory.

## Compute heat/flux compute ID does not compute stress/atom

Self-explanatory.

## Compute hexorder/atom cutoff is longer than pairwise cutoff

Cannot compute order parameter beyond cutoff.

## Compute hexorder/atom requires a pair style be defined

Self-explanatory.

## Compute improper/local used when impropers are not allowed

The atom style does not support impropers.

## Compute inertia/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute ke/rigid with non-rigid fix-ID

Self-explanatory.

## Compute msd/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute msd/chunk nchunk is not static

This is required because the MSD cannot be computed consistently if the number of chunks is changing. Compute chunk/atom allows setting nchunk to be static.

## Compute nve/asphere requires atom style ellipsoid

Self-explanatory.

## Compute nvt/nph/npt asphere requires atom style ellipsoid

Self-explanatory.

## Compute nvt/nph/npt body requires atom style body

Self-explanatory.

## Compute omega/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute orientorder/atom cutoff is longer than pairwise cutoff

Cannot compute order parameter beyond cutoff.

## Compute orientorder/atom requires a pair style be defined

Self-explanatory.

## Compute pair must use group all

Pair styles accumulate energy on all atoms.

## Compute pe must use group all

Energies computed by potentials (pair, bond, etc) are computed on all atoms.

## Compute plasticity/atom cannot be used with this pair style

Self-explanatory.

## Compute plasticity/atom requires Peridynamic pair style

Self-explanatory.

## Compute pressure must use group all

Virial contributions computed by potentials (pair, bond, etc) are computed on all atoms.

## Compute pressure requires temperature ID to include kinetic energy

The keflag cannot be used unless a temperature compute is provided.

## Compute pressure temperature ID does not compute temperature

The compute ID assigned to a pressure computation must compute temperature.

## Compute property/atom floating point vector does not exist

The command is accessing a vector added by the fix property/atom command, that does not exist.

## Compute property/atom for atom property that is not allocated

Self-explanatory.

## Compute property/atom integer vector does not exist

The command is accessing a vector added by the fix property/atom command, that does not exist.

## Compute property/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute property/local cannot use these inputs together

Only inputs that generate the same number of datums can be used together. E.g. bond and angle quantities cannot be mixed.

## Compute property/local does not (yet) work with atom\_style template

Self-explanatory.

## Compute property/local for property that is not allocated

Self-explanatory.

## Compute rdf requires a pair style be defined

Self-explanatory.

## Compute reduce compute array is accessed out-of-range

An index for the array is out of bounds.

## Compute reduce compute calculates global values

A compute that calculates peratom or local values is required.

## Compute reduce compute does not calculate a local array

Self-explanatory.

## Compute reduce compute does not calculate a local vector

Self-explanatory.

## Compute reduce compute does not calculate a per-atom array

Self-explanatory.

## Compute reduce compute does not calculate a per-atom vector

Self-explanatory.

## Compute reduce fix array is accessed out-of-range

An index for the array is out of bounds.

## Compute reduce fix calculates global values

A fix that calculates peratom or local values is required.

## Compute reduce fix does not calculate a local array

Self-explanatory.

## Compute reduce fix does not calculate a local vector

Self-explanatory.

## Compute reduce fix does not calculate a per-atom array

Self-explanatory.

## Compute reduce fix does not calculate a per-atom vector

Self-explanatory.

## Compute reduce replace requires min or max mode

Self-explanatory.

## Compute reduce variable is not atom-style variable

Self-explanatory.

## Compute slice compute array is accessed out-of-range

An index for the array is out of bounds.

## Compute slice compute does not calculate a global array

Self-explanatory.

## Compute slice compute does not calculate a global vector

Self-explanatory.

## Compute slice compute does not calculate global vector or array

Self-explanatory.

## Compute slice compute vector is accessed out-of-range

The index for the vector is out of bounds.

## Compute slice fix array is accessed out-of-range

An index for the array is out of bounds.

## Compute slice fix does not calculate a global array

Self-explanatory.

## Compute slice fix does not calculate a global vector

Self-explanatory.

## Compute slice fix does not calculate global vector or array

Self-explanatory.

## Compute slice fix vector is accessed out-of-range

The index for the vector is out of bounds.

## Compute sna/atom cutoff is longer than pairwise cutoff

Self-explanatory.

## Compute sna/atom requires a pair style be defined

Self-explanatory.

## Compute snad/atom cutoff is longer than pairwise cutoff

Self-explanatory.

## Compute snad/atom requires a pair style be defined

Self-explanatory.

## Compute snav/atom cutoff is longer than pairwise cutoff

Self-explanatory.

## Compute snav/atom requires a pair style be defined

Self-explanatory.

## Compute stress/atom temperature ID does not compute temperature

The specified compute must compute temperature.

## Compute temp/asphere requires atom style ellipsoid

Self-explanatory.

## Compute temp/asphere requires extended particles

This compute cannot be used with point particles.

## Compute temp/body requires atom style body

Self-explanatory.

## Compute temp/body requires bodies

This compute can only be applied to body particles.

## Compute temp/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute temp/cs requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Compute temp/cs used when bonds are not allowed

This compute only works on pairs of bonded particles.

## Compute temp/partial cannot use vz for 2d systemx

Self-explanatory.

## Compute temp/profile cannot bin z for 2d systems

Self-explanatory.

## Compute temp/profile cannot use vz for 2d systemx

Self-explanatory.

## Compute temp/sphere requires atom style sphere

Self-explanatory.

## Compute ti kspace style does not exist

Self-explanatory.

## Compute ti pair style does not exist

Self-explanatory.

## Compute ti tail when pair style does not compute tail corrections

Self-explanatory.

## Compute torque/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Compute used in dump between runs is not current

The compute was not invoked on the current timestep, therefore it cannot be used in a dump between runs.

## Compute used in variable between runs is not current

Computes cannot be invoked by a variable in between runs. Thus they must have been evaluated on the last timestep of the previous run in order for their value(s) to be accessed. See the page for the variable command for more info.

## Compute used in variable thermo keyword between runs is not current

Some thermo keywords rely on a compute to calculate their value(s). Computes cannot be invoked by a variable in between runs. Thus they must have been evaluated on the last timestep of the previous run in order for their value(s) to be accessed. See the page for the variable command for more info.

## Compute vcm/chunk does not use chunk/atom compute

The style of the specified compute is not chunk/atom.

## Computed temperature for fix temp/berendsen cannot be 0.0

Self-explanatory.

## Computed temperature for fix temp/rescale cannot be 0.0

Cannot rescale the temperature to a new value if the current temperature is 0.0.

## Core/shell partner atom not found

Could not find one of the atoms in the bond pair.

## Core/shell partners were not all found

Could not find or more atoms in the bond pairs.

## Could not adjust g\_ewald\_6

The Newton-Raphson solver failed to converge to a good value for g\_ewald. This error should not occur for typical problems. Please send an email to the developers.

## Could not compute g\_ewald

The Newton-Raphson solver failed to converge to a good value for g\_ewald. This error should not occur for typical problems. Please send an email to the developers.

## Could not compute grid size

The code is unable to compute a grid size consistent with the desired accuracy. This error should not occur for typical problems. Please send an email to the developers.

## Could not compute grid size for Coulomb interaction

The code is unable to compute a grid size consistent with the desired accuracy. This error should not occur for typical problems. Please send an email to the developers.

## Could not compute grid size for Dispersion

The code is unable to compute a grid size consistent with the desired accuracy. This error should not occur for typical problems. Please send an email to the developers.

## Could not create 3d FFT plan

The FFT setup for the PPPM solver failed, typically due to lack of memory. This is an unusual error. Check the size of the FFT grid you are requesting.

## Could not create 3d grid of processors

The specified constraints did not allow a Px by Py by Pz grid to be created where Px \* Py \* Pz = P = total number of processors.

## Could not create 3d remap plan

The FFT setup in pppm failed.

## Could not create Python function arguments

This is an internal Python error, possibly because the number of inputs to the function is too large.

## Could not create numa grid of processors

The specified constraints did not allow this style of grid to be created. Usually this is because the total processor count is not a multiple of the cores/node or the user specified processor count is > 1 in one of the dimensions.

## Could not create twolevel 3d grid of processors

The specified constraints did not allow this style of grid to be created.

## Could not evaluate Python function input variable

Self-explanatory.

## Could not find Python function

The provided Python code was run successfully, but it not define a callable function with the required name.

## Could not find atom\_modify first group ID

Self-explanatory.

## Could not find change\_box group ID

Group ID used in the change\_box command does not exist.

## Could not find compute ID for PRD

Self-explanatory.

## Could not find compute ID for TAD

Self-explanatory.

## Could not find compute ID for temperature bias

Self-explanatory.

## Could not find compute ID to delete

Self-explanatory.

## Could not find compute displace/atom fix ID

Self-explanatory.

## Could not find compute event/displace fix ID

Self-explanatory.

## Could not find compute group ID

Self-explanatory.

## Could not find compute heat/flux compute ID

Self-explanatory.

## Could not find compute msd fix ID

Self-explanatory.

## Could not find compute msd/chunk fix ID

The compute creates an internal fix, which has been deleted.

## Could not find compute pressure temperature ID

The compute ID for calculating temperature does not exist.

## Could not find compute stress/atom temperature ID

Self-explanatory.

## Could not find compute vacf fix ID

Self-explanatory.

## Could not find compute/voronoi surface group ID

Self-explanatory.

## Could not find compute\_modify ID

Self-explanatory.

## Could not find custom per-atom property ID

Self-explanatory.

## Could not find delete\_atoms group ID

Group ID used in the delete\_atoms command does not exist.

## Could not find delete\_atoms region ID

Region ID used in the delete\_atoms command does not exist.

## Could not find displace\_atoms group ID

Group ID used in the displace\_atoms command does not exist.

## Could not find dump custom compute ID

Self-explanatory.

## Could not find dump custom fix ID

Self-explanatory.

## Could not find dump custom variable name

Self-explanatory.

## Could not find dump group ID

A group ID used in the dump command does not exist.

## Could not find dump local compute ID

Self-explanatory.

## Could not find dump local fix ID

Self-explanatory.

## Could not find dump modify compute ID

Self-explanatory.

## Could not find dump modify custom atom floating point property ID

Self-explanatory.

## Could not find dump modify custom atom integer property ID

Self-explanatory.

## Could not find dump modify fix ID

Self-explanatory.

## Could not find dump modify variable name

Self-explanatory.

## Could not find fix ID to delete

Self-explanatory.

## Could not find fix adapt storage fix ID

This should not happen unless you explicitly deleted a secondary fix that fix adapt created internally.

## Could not find fix halt variable name

Self-explanatory.

## Could not find fix gcmc exclusion group ID

Self-explanatory.

## Could not find fix gcmc rotation group ID

Self-explanatory.

## Could not find fix group ID

A group ID used in the fix command does not exist.

## Could not find fix msst compute ID

Self-explanatory.

## Could not find fix poems group ID

A group ID used in the fix poems command does not exist.

## Could not find fix recenter group ID

A group ID used in the fix recenter command does not exist.

## Could not find fix rigid group ID

A group ID used in the fix rigid command does not exist.

## Could not find fix srd group ID

Self-explanatory.

## Could not find fix\_modify ID

A fix ID used in the fix\_modify command does not exist.

## Could not find fix\_modify pressure ID

The compute ID for computing pressure does not exist.

## Could not find fix\_modify temperature ID

The compute ID for computing temperature does not exist.

## Could not find group clear group ID

Self-explanatory.

## Could not find group delete group ID

Self-explanatory.

## Could not find pair fix ID

A fix is created internally by the pair style to store shear history information. You cannot delete it.

## Could not find set group ID

Group ID specified in set command does not exist.

## Could not find specified fix gcmc group ID

Self-explanatory.

## Could not find thermo compute ID

Compute ID specified in thermo\_style command does not exist.

## Could not find thermo custom compute ID

The compute ID needed by thermo style custom to compute a requested quantity does not exist.

## Could not find thermo custom fix ID

The fix ID needed by thermo style custom to compute a requested quantity does not exist.

## Could not find thermo custom variable name

Self-explanatory.

## Could not find thermo fix ID

Fix ID specified in thermo\_style command does not exist.

## Could not find thermo variable name

Self-explanatory.

## Could not find thermo\_modify pressure ID

The compute ID needed by thermo style custom to compute pressure does not exist.

## Could not find thermo\_modify temperature ID

The compute ID needed by thermo style custom to compute temperature does not exist.

## Could not find undump ID

A dump ID used in the undump command does not exist.

## Could not find velocity group ID

A group ID used in the velocity command does not exist.

## Could not find velocity temperature ID

The compute ID needed by the velocity command to compute temperature does not exist.

## Could not find/initialize a specified accelerator device

Could not initialize at least one of the devices specified for the gpu package

## Could not grab element entry from EIM potential file

Self-explanatory

## Could not grab global entry from EIM potential file

Self-explanatory.

## Could not grab pair entry from EIM potential file

Self-explanatory.

## Could not initialize embedded Python

The main module in Python was not accessible.

## Could not open Python file

The specified file of Python code cannot be opened. Check that the path and name are correct.

## Could not process Python file

The Python code in the specified file was not run successfully by Python, probably due to errors in the Python code.

## Could not process Python string

The Python code in the here string was not run successfully by Python, probably due to errors in the Python code.

## Coulomb PPPMDisp order has been reduced below minorder

The default minimum order is 2. This can be reset by the kspace\_modify minorder command.

## Coulombic cutoff not supported in pair\_style buck/long/coul/coul

Must use long-range Coulombic interactions.

## Coulombic cutoff not supported in pair\_style lj/long/coul/long

Must use long-range Coulombic interactions.

## Coulombic cutoff not supported in pair\_style lj/long/tip4p/long

Must use long-range Coulombic interactions.

## Coulombic cutoffs of pair hybrid sub-styles do not match

If using a Kspace solver, all Coulombic cutoffs of long pair styles must be the same.

## Coulombic cut not supported in pair\_style lj/long/dipole/long

Must use long-range Coulombic interactions.

## Cound not find dump\_modify ID

Self-explanatory.

## Create\_atoms command before simulation box is defined

The create\_atoms command cannot be used before a read\_data, read\_restart, or create\_box command.

## Create\_atoms molecule has atom IDs, but system does not

The atom\_style id command can be used to force atom IDs to be stored.

## Create\_atoms molecule must have atom types

The defined molecule does not specify atom types.

## Create\_atoms molecule must have coordinates

The defined molecule does not specify coordinates.

## Create\_atoms region ID does not exist

A region ID used in the create\_atoms command does not exist.

## Create\_bonds command before simulation box is defined

Self-explanatory.

## Create\_bonds command requires no kspace\_style be defined

This is so that atom pairs that are already bonded to not appear in the neighbor list.

## Create\_bonds command requires special\_bonds 1-2 weights be 0.0

This is so that atom pairs that are already bonded to not appear in the neighbor list.

## Create\_bonds max distance > neighbor cutoff

Can only create bonds for atom pairs that will be in neighbor list.

## Create\_bonds requires a pair style be defined

Self-explanatory.

## Create\_box region ID does not exist

Self-explanatory.

## Create\_box region does not support a bounding box

Not all regions represent bounded volumes. You cannot use such a region with the create\_box command.

## Custom floating point vector for fix store/state does not exist

The command is accessing a vector added by the fix property/atom command, that does not exist.

## Custom integer vector for fix store/state does not exist

The command is accessing a vector added by the fix property/atom command, that does not exist.

## Custom per-atom property ID is not floating point

Self-explanatory.

## Custom per-atom property ID is not integer

Self-explanatory.

## Cut-offs missing in pair\_style lj/long/dipole/long

Self-explanatory.

## Cutoffs missing in pair\_style buck/long/coul/long

Self-explanatory.

## Cutoffs missing in pair\_style lj/long/coul/long

Self-explanatory.

## Cyclic loop in joint connections

Fix poems cannot (yet) work with coupled bodies whose joints connect the bodies in a ring (or cycle).

## Degenerate lattice primitive vectors

Invalid set of 3 lattice vectors for lattice command.

## Delete region ID does not exist

Self-explanatory.

## Delete\_atoms command before simulation box is defined

The delete\_atoms command cannot be used before a read\_data, read\_restart, or create\_box command.

## Delete\_atoms cutoff > max neighbor cutoff

Can only delete atoms in atom pairs that will be in neighbor list.

## Delete\_atoms mol yes requires atom attribute molecule

Cannot use this option with a non-molecular system.

## Delete\_atoms requires a pair style be defined

This is because atom deletion within a cutoff uses a pairwise neighbor list.

## Delete\_bonds command before simulation box is defined

The delete\_bonds command cannot be used before a read\_data, read\_restart, or create\_box command.

## Delete\_bonds command with no atoms existing

No atoms are yet defined so the delete\_bonds command cannot be used.

## Deposition region extends outside simulation box

Self-explanatory.

## Did not assign all atoms correctly

Atoms read in from a data file were not assigned correctly to processors. This is likely due to some atom coor- dinates being outside a non-periodic simulation box.

## Did not assign all restart atoms correctly

Atoms read in from the restart file were not assigned correctly to processors. This is likely due to some atom coordinates being outside a non-periodic simulation box. Normally this should not happen. You may wish to use the “remap” option on the read\_restart command to see if this helps.

## Did not find all elements in MEAM library file

Some requested elements were not found in the MEAM file. Check spelling etc.

## Did not find fix shake partner info

Could not find bond partners implied by fix shake command. This error can be triggered if the delete\_bonds command was used before fix shake, and it removed bonds without resetting the 1-2, 1-3, 1-4 weighting list via the special keyword.

## Did not find keyword in table file

Keyword used in pair\_coeff command was not found in table file.

## Did not set pressure for fix rigid/nph

The press keyword must be specified.

## Did not set temp for fix rigid/nvt/small

Self-explanatory.

## Did not set temp or press for fix rigid/npt/small

Self-explanatory.

## Did not set temperature for fix rigid/nvt

The temp keyword must be specified.

## Did not set temperature or pressure for fix rigid/npt

The temp and press keywords must be specified.

## Dihedral atom missing in delete\_bonds

The delete\_bonds command cannot find one or more atoms in a particular dihedral on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid dihedral.

## Dihedral atom missing in set command

The set command cannot find one or more atoms in a particular dihedral on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid dihedral.

## Dihedral atoms %d %d %d %d missing on proc %d at step %ld

One or more of 4 atoms needed to compute a particular dihedral are missing on this processor. Typically this is because the pairwise cutoff is set too short or the dihedral has blown apart and an atom is too far away.

## Dihedral atoms missing on proc %d at step %ld

One or more of 4 atoms needed to compute a particular dihedral are missing on this processor. Typically this is because the pairwise cutoff is set too short or the dihedral has blown apart and an atom is too far away.

## Dihedral charmm is incompatible with Pair style

Dihedral style charmm must be used with a pair style charmm in order for the 1-4 epsilon/sigma parameters to be defined.

## Dihedral coeff for hybrid has invalid style

Dihedral style hybrid uses another dihedral style as one of its coefficients. The dihedral style used in the dihe- dral\_coeff command or read from a restart file is not recognized.

## Dihedral coeffs are not set

No dihedral coefficients have been assigned in the data file or via the dihedral\_coeff command.

## Dihedral style hybrid cannot have hybrid as an argument

Self-explanatory.

## Dihedral style hybrid cannot have none as an argument

Self-explanatory.

## Dihedral style hybrid cannot use same dihedral style twice

Self-explanatory.

## Dihedral/improper extent > half of periodic box length

This error was detected by the neigh\_modify check yes setting. It is an error because the dihedral atoms are so far apart it is ambiguous how it should be defined.

## Dihedral\_coeff command before dihedral\_style is defined

Coefficients cannot be set in the data file or via the dihedral\_coeff command until an dihedral\_style has been assigned.

## Dihedral\_coeff command before simulation box is defined

The dihedral\_coeff command cannot be used before a read\_data, read\_restart, or create\_box command.

## Dihedral\_coeff command when no dihedrals allowed

The chosen atom style does not allow for dihedrals to be defined.

## Dihedral\_style command when no dihedrals allowed

The chosen atom style does not allow for dihedrals to be defined.

## Dihedrals assigned incorrectly

Dihedrals read in from the data file were not assigned correctly to atoms. This means there is something invalid about the topology definitions.

## Dihedrals defined but no dihedral types

The data file header lists dihedrals but no dihedral types.

## Dimension command after simulation box is defined

The dimension command cannot be used after a read\_data, read\_restart, or create\_box command.

## Disk limit not supported by OS or illegal path

Self-explanatory.

## Dispersion PPPMDisp order has been reduced below minorder

The default minimum order is 2. This can be reset by the kspace\_modify minorder command.

## Displace\_atoms command before simulation box is defined

The displace\_atoms command cannot be used before a read\_data, read\_restart, or create\_box command.

## Distance must be > 0 for compute event/displace

Self-explanatory.

## Divide by 0 in influence function

This should not normally occur. It is likely a problem with your model.

## Divide by 0 in influence function of pair peri/lps

This should not normally occur. It is likely a problem with your model.

## Divide by 0 in variable formula

Self-explanatory.

## Domain too large for neighbor bins

The domain has become extremely large so that neighbor bins cannot be used. Most likely, one or more atoms have been blown out of the simulation box to a great distance.

## Double precision is not supported on this accelerator

Self-explanatory

## Dump atom/gz only writes compressed files

The dump atom/gz output file name must have a .gz suffix.

## Dump cfg arguments can not mix xs|ys|zs with xsu|ysu|zsu

Self-explanatory.

## Dump cfg arguments must start with ‘mass type xs ys zs’ or ‘mass type xsu ysu zsu’

This is a requirement of the CFG output format. See the dump cfg doc page for more details.

## Dump cfg requires one snapshot per file

Use the wildcard “\*” character in the filename.

## Dump cfg/gz only writes compressed files

The dump cfg/gz output file name must have a .gz suffix.

## Dump custom and fix not computed at compatible times

The fix must produce per-atom quantities on timesteps that dump custom needs them.

## Dump custom compute does not calculate per-atom array

Self-explanatory.

## Dump custom compute does not calculate per-atom vector

Self-explanatory.

## Dump custom compute does not compute per-atom info

Self-explanatory.

## Dump custom compute vector is accessed out-of-range

Self-explanatory.

## Dump custom fix does not compute per-atom array

Self-explanatory.

## Dump custom fix does not compute per-atom info

Self-explanatory.

## Dump custom fix does not compute per-atom vector

Self-explanatory.

## Dump custom fix vector is accessed out-of-range

Self-explanatory.

## Dump custom variable is not atom-style variable

Only atom-style variables generate per-atom quantities, needed for dump output.

## Dump custom/gz only writes compressed files

The dump custom/gz output file name must have a .gz suffix.

## Dump dcd of non-matching # of atoms

Every snapshot written by dump dcd must contain the same # of atoms.

## Dump dcd requires sorting by atom ID

Use the dump\_modify sort command to enable this.

## Dump every variable returned a bad timestep

The variable must return a timestep greater than the current timestep.

## Dump file MPI-IO output not allowed with % in filename

This is because a % signifies one file per processor and MPI-IO creates one large file for all processors.

## Dump file does not contain requested snapshot

Self-explanatory.

## Dump file is incorrectly formatted

Self-explanatory.

## Dump image body yes requires atom style body

Self-explanatory.

## Dump image bond not allowed with no bond types

Self-explanatory.

## Dump image cannot perform sorting

Self-explanatory.

## Dump image line requires atom style line

Self-explanatory.

## Dump image requires one snapshot per file

Use a “\*” in the filename.

## Dump image tri requires atom style tri

Self-explanatory.

## Dump local and fix not computed at compatible times

The fix must produce per-atom quantities on timesteps that dump local needs them.

## Dump local attributes contain no compute or fix

Self-explanatory.

## Dump local cannot sort by atom ID

This is because dump local does not really dump per-atom info.

## Dump local compute does not calculate local array

Self-explanatory.

## Dump local compute does not calculate local vector

Self-explanatory.

## Dump local compute does not compute local info

Self-explanatory.

## Dump local compute vector is accessed out-of-range

Self-explanatory.

## Dump local count is not consistent across input fields

Every column of output must be the same length.

## Dump local fix does not compute local array

Self-explanatory.

## Dump local fix does not compute local info

Self-explanatory.

## Dump local fix does not compute local vector

Self-explanatory.

## Dump local fix vector is accessed out-of-range

Self-explanatory.

## Dump modify bcolor not allowed with no bond types

Self-explanatory.

## Dump modify bdiam not allowed with no bond types

Self-explanatory.

## Dump modify compute ID does not compute per-atom array

Self-explanatory.

## Dump modify compute ID does not compute per-atom info

Self-explanatory.

## Dump modify compute ID does not compute per-atom vector

Self-explanatory.

## Dump modify compute ID vector is not large enough

Self-explanatory.

## Dump modify element names do not match atom types

Number of element names must equal number of atom types.

## Dump modify fix ID does not compute per-atom array

Self-explanatory.

## Dump modify fix ID does not compute per-atom info

Self-explanatory.

## Dump modify fix ID does not compute per-atom vector

Self-explanatory.

## Dump modify fix ID vector is not large enough

Self-explanatory.

## Dump modify variable is not atom-style variable

Self-explanatory.

## Dump sort column is invalid

Self-explanatory.

## Dump xtc requires sorting by atom ID

Use the dump\_modify sort command to enable this.

## Dump xyz/gz only writes compressed files

The dump xyz/gz output file name must have a .gz suffix.

## Dump\_modify buffer yes not allowed for this style

Self-explanatory.

## Dump\_modify format string is too short

There are more fields to be dumped in a line of output than your format string specifies.

## Dump\_modify region ID does not exist

Self-explanatory.

## Dumping an atom property that is not allocated

The chosen atom style does not define the per-atom quantity being dumped.

## Duplicate atom IDs exist

Self-explanatory.

## Duplicate fields in read\_dump command

Self-explanatory.

## Duplicate particle in PeriDynamic bond - simulation box is too small

This is likely because your box length is shorter than 2 times the bond length.

## Electronic temperature dropped below zero

Something has gone wrong with the fix ttm electron temperature model.

## Element not defined in potential file

The specified element is not in the potential file.

## Empty brackets in variable

There is no variable syntax that uses empty brackets. Check the variable doc page.

## Energy was not tallied on needed timestep

You are using a thermo keyword that requires potentials to have tallied energy, but they did not on this timestep. See the variable page for ideas on how to make this work.

## Epsilon or sigma reference not set by pair style in PPPMDisp

Self-explanatory.

## Epsilon or sigma reference not set by pair style in ewald/n

The pair style is not providing the needed epsilon or sigma values.

## Error in MEAM parameter file: keyword %s (further information)

Self-explanatory. Check the parameter file.

## Error in vdw spline: inner radius > outer radius

A pre-tabulated spline is invalid. Likely a problem with the potential parameters.

## Error writing averaged chunk data

Something in the output to the file triggered an error.

## Error writing file header

Something in the output to the file triggered an error.

## Error writing out correlation data

Something in the output to the file triggered an error.

## Error writing out histogram data

Something in the output to the file triggered an error.

## Error writing out time averaged data

Something in the output to the file triggered an error.

## Failed to allocate %ld bytes for array %s

Your LAMMPS simulation has run out of memory. You need to run a smaller simulation or on more processors.

## Failed to open FFmpeg pipeline to file %s

The specified file cannot be opened. Check that the path and name are correct and writable and that the FFmpeg executable can be found and run.

## Failed to reallocate %ld bytes for array %s

Your LAMMPS simulation has run out of memory. You need to run a smaller simulation or on more processors.

## Fewer SRD bins than processors in some dimension

This is not allowed. Make your SRD bin size smaller.

## File variable could not read value

Check the file assigned to the variable.

## Final box dimension due to fix deform is < 0.0

Self-explanatory.

## Fix %s does not allow use of dynamic group

Dynamic groups have not yet been enabled for this fix.

## Fix ID for compute chunk/atom does not exist

Self-explanatory.

## Fix ID for compute erotate/rigid does not exist

Self-explanatory.

## Fix ID for compute ke/rigid does not exist

Self-explanatory.

## Fix ID for compute reduce does not exist

Self-explanatory.

## Fix ID for compute slice does not exist

Self-explanatory.

## Fix ID for fix ave/atom does not exist

Self-explanatory.

## Fix ID for fix ave/chunk does not exist

Self-explanatory.

## Fix ID for fix ave/correlate does not exist

Self-explanatory.

## Fix ID for fix ave/histo does not exist

Self-explanatory.

## Fix ID for fix ave/time does not exist

Self-explanatory.

## Fix ID for fix store/state does not exist

Self-explanatory

## Fix ID for fix vector does not exist

Self-explanatory.

## Fix ID for read\_data does not exist

Self-explanatory.

## Fix ID for velocity does not exist

Self-explanatory.

## Fix ID must be alphanumeric or underscore characters

Self-explanatory.

## Fix SRD: bad bin assignment for SRD advection

Something has gone wrong in your SRD model; try using more conservative settings.

## Fix SRD: bad search bin assignment

Something has gone wrong in your SRD model; try using more conservative settings.

## Fix SRD: bad stencil bin for big particle

Something has gone wrong in your SRD model; try using more conservative settings.

## Fix SRD: too many big particles in bin

Reset the ATOMPERBIN parameter at the top of fix\_srd.cpp to a larger value, and re-compile the code.

## Fix SRD: too many walls in bin

This should not happen unless your system has been setup incorrectly.

## Fix adapt interface to this pair style not supported

New coding for the pair style would need to be done.

## Fix adapt kspace style does not exist

Self-explanatory.

## Fix adapt pair style does not exist

Self-explanatory

## Fix adapt pair style param not supported

The pair style does not know about the parameter you specified.

## Fix adapt requires atom attribute charge

The atom style being used does not specify an atom charge.

## Fix adapt requires atom attribute diameter

The atom style being used does not specify an atom diameter.

## Fix adapt type pair range is not valid for pair hybrid sub-style

Self-explanatory.

## Fix append/atoms requires a lattice be defined

Use the lattice command for this purpose.

## Fix ave/atom compute array is accessed out-of-range

Self-explanatory.

## Fix ave/atom compute does not calculate a per-atom array

Self-explanatory.

## Fix ave/atom compute does not calculate a per-atom vector

A compute used by fix ave/atom must generate per-atom values.

## Fix ave/atom compute does not calculate per-atom values

A compute used by fix ave/atom must generate per-atom values.

## Fix ave/atom fix array is accessed out-of-range

Self-explanatory.

## Fix ave/atom fix does not calculate a per-atom array

Self-explanatory.

## Fix ave/atom fix does not calculate a per-atom vector

A fix used by fix ave/atom must generate per-atom values.

## Fix ave/atom fix does not calculate per-atom values

A fix used by fix ave/atom must generate per-atom values.

## Fix ave/atom variable is not atom-style variable

A variable used by fix ave/atom must generate per-atom values.

## Fix ave/chunk compute does not calculate a per-atom array

Self-explanatory.

## Fix ave/chunk compute does not calculate a per-atom vector

Self-explanatory.

## Fix ave/chunk compute does not calculate per-atom values

Self-explanatory.

## Fix ave/chunk compute vector is accessed out-of-range

Self-explanatory.

## Fix ave/chunk does not use chunk/atom compute

The specified compute is not for a compute chunk/atom command.

## Fix ave/chunk fix does not calculate a per-atom array

Self-explanatory.

## Fix ave/chunk fix does not calculate a per-atom vector

Self-explanatory.

## Fix ave/chunk fix does not calculate per-atom values

Self-explanatory.

## Fix ave/chunk fix vector is accessed out-of-range

Self-explanatory.

## Fix ave/chunk variable is not atom-style variable

Self-explanatory.

## Fix ave/correlate compute does not calculate a scalar

Self-explanatory.

## Fix ave/correlate compute does not calculate a vector

Self-explanatory.

## Fix ave/correlate compute vector is accessed out-of-range

The index for the vector is out of bounds.

## Fix ave/correlate fix does not calculate a scalar

Self-explanatory.

## Fix ave/correlate fix does not calculate a vector

Self-explanatory.

## Fix ave/correlate fix vector is accessed out-of-range

The index for the vector is out of bounds.

## Fix ave/correlate variable is not equal-style variable

Self-explanatory.

## Fix ave/histo cannot input local values in scalar mode

Self-explanatory.

## Fix ave/histo cannot input per-atom values in scalar mode

Self-explanatory.

## Fix ave/histo compute array is accessed out-of-range

Self-explanatory.

## Fix ave/histo compute does not calculate a global array

Self-explanatory.

## Fix ave/histo compute does not calculate a global scalar

Self-explanatory.

## Fix ave/histo compute does not calculate a global vector

Self-explanatory.

## Fix ave/histo compute does not calculate a local array

Self-explanatory.

## Fix ave/histo compute does not calculate a local vector

Self-explanatory.

## Fix ave/histo compute does not calculate a per-atom array

Self-explanatory.

## Fix ave/histo compute does not calculate a per-atom vector

Self-explanatory.

## Fix ave/histo compute does not calculate local values

Self-explanatory.

## Fix ave/histo compute does not calculate per-atom values

Self-explanatory.

## Fix ave/histo compute vector is accessed out-of-range

Self-explanatory.

## Fix ave/histo fix array is accessed out-of-range

Self-explanatory.

## Fix ave/histo fix does not calculate a global array

Self-explanatory.

## Fix ave/histo fix does not calculate a global scalar

Self-explanatory.

## Fix ave/histo fix does not calculate a global vector

Self-explanatory.

## Fix ave/histo fix does not calculate a local array

Self-explanatory.

## Fix ave/histo fix does not calculate a local vector

Self-explanatory.

## Fix ave/histo fix does not calculate a per-atom array

Self-explanatory.

## Fix ave/histo fix does not calculate a per-atom vector

Self-explanatory.

## Fix ave/histo fix does not calculate local values

Self-explanatory.

## Fix ave/histo fix does not calculate per-atom values

Self-explanatory.

## Fix ave/histo fix vector is accessed out-of-range

Self-explanatory.

## Fix ave/histo input is invalid compute

Self-explanatory.

## Fix ave/histo input is invalid fix

Self-explanatory.

## Fix ave/histo input is invalid variable

Self-explanatory.

## Fix ave/histo inputs are not all global, peratom, or local

All inputs in a single fix ave/histo command must be of the same style.

## Fix ave/histo/weight value and weight vector lengths do not match

Self-explanatory.

## Fix ave/time cannot set output array intensive/extensive from these inputs

One of more of the vector inputs has individual elements which are flagged as intensive or extensive. Such an input cannot be flagged as all intensive/extensive when turned into an array by fix ave/time.

## Fix ave/time cannot use variable with vector mode

Variables produce scalar values.

## Fix ave/time columns are inconsistent lengths

Self-explanatory.

## Fix ave/time compute array is accessed out-of-range

An index for the array is out of bounds.

## Fix ave/time compute does not calculate a scalar

Self-explanatory.

## Fix ave/time compute does not calculate a vector

Self-explanatory.

## Fix ave/time compute does not calculate an array

Self-explanatory.

## Fix ave/time compute vector is accessed out-of-range

The index for the vector is out of bounds.

## Fix ave/time fix array cannot be variable length

Self-explanatory.

## Fix ave/time fix array is accessed out-of-range

An index for the array is out of bounds.

## Fix ave/time fix does not calculate a scalar

Self-explanatory.

## Fix ave/time fix does not calculate a vector

Self-explanatory.

## Fix ave/time fix does not calculate an array

Self-explanatory.

## Fix ave/time fix vector cannot be variable length

Self-explanatory.

## Fix ave/time fix vector is accessed out-of-range

The index for the vector is out of bounds.

## Fix ave/time variable is not equal-style variable

Self-explanatory.

## Fix balance rcb cannot be used with comm\_style brick

Comm\_style tiled must be used instead.

## Fix balance shift string is invalid

The string can only contain the characters “x”, “y”, or “z”.

## Fix bond/break needs ghost atoms from further away

This is because the fix needs to walk bonds to a certain distance to acquire needed info, The comm\_modify cutoff command can be used to extend the communication range.

## Fix bond/create angle type is invalid

Self-explanatory.

## Fix bond/create cutoff is longer than pairwise cutoff

This is not allowed because bond creation is done using the pairwise neighbor list.

## Fix bond/create dihedral type is invalid

Self-explanatory.

## Fix bond/create improper type is invalid

Self-explanatory.

## Fix bond/create induced too many angles/dihedrals/impropers per atom

See the read\_data command for info on using the “extra/angle/per/atom”, (or dihedral, improper) keywords to allow for additional angles, dihedrals, and impropers to be formed.

## Fix bond/create needs ghost atoms from further away

This is because the fix needs to walk bonds to a certain distance to acquire needed info, The comm\_modify cutoff command can be used to extend the communication range.

## Fix bond/react: Cannot use fix bond/react with non-molecular systems

Only systems with bonds that can be changed can be used. Atom\_style template does not qualify.

## Fix bond/react: Invalid template atom ID in map file

Atom IDs in molecule templates range from 1 to the number of atoms in the template.

## Fix bond/react: Rmax cutoff is longer than pairwise cutoff

This is not allowed because bond creation is done using the pairwise neighbor list.

## Fix bond/react: Molecule template ID for fix bond/react does not exist

A valid molecule template must have been created with the molecule command.

## Fix bond/react: Reaction templates must contain the same number of atoms

There should be a one-to-one correspondence between atoms in the pre-reacted and post-reacted templates, as specified by the map file.

## Fix bond/react: Unknown section in map file

Please ensure reaction map files are properly formatted.

## Fix bond/react: Atom/Bond type affected by reaction too close to template edge

This means an atom which changes type or connectivity during the reaction is too close to an ‘edge’ atom defined in the map file. This could cause incorrect assignment of bonds, angle, etc. Generally, this means you must include more atoms in your templates, such that there are at least two atoms between each atom involved in the reaction and an edge atom.

## Fix bond/react: Fix bond/react needs ghost atoms from farther away

This is because a processor needs to map the entire unreacted molecule template onto simulation atoms it knows about. The comm\_modify cutoff command can be used to extend the communication range.

## Fix bond/react: A deleted atom cannot remain bonded to an atom that is not deleted

Self-explanatory.

## Fix bond/react: First neighbors of chiral atoms must be of mutually different types

Self-explanatory.

## Fix bond/react: Chiral atoms must have exactly four first neighbors

Self-explanatory.

## Fix bond/react: Molecule template ‘Coords’ section required for chiralIDs keyword

The coordinates of atoms in the pre-reacted template are used to determine chirality.

## Fix bond/react special bond generation overflow

The number of special bonds per-atom created by a reaction exceeds the system setting. See the read\_data or create\_box command for how to specify this value.

## Fix bond/react topology/atom exceed system topology/atom

The number of bonds, angles etc per-atom created by a reaction exceeds the system setting. See the read\_data or create\_box command for how to specify this value.

## Fix bond/swap cannot use dihedral or improper styles

These styles cannot be defined when using this fix.

## Fix bond/swap requires pair and bond styles

Self-explanatory.

## Fix bond/swap requires special\_bonds = 0,1,1

Self-explanatory.

## Fix box/relax generated negative box length

The pressure being applied is likely too large. Try applying it incrementally, to build to the high pressure.

## Fix command before simulation box is defined

The fix command cannot be used before a read\_data, read\_restart, or create\_box command.

## Fix deform cannot use yz variable with xy

The yz setting cannot be a variable if xy deformation is also specified. This is because LAMMPS cannot deter- mine if the yz setting will induce a box flip which would be invalid if xy is also changing.

## Fix deform is changing yz too much with xy

When both yz and xy are changing, it induces changes in xz if the box must flip from one tilt extreme to another. Thus it is not allowed for yz to grow so much that a flip is induced.

## Fix deform tilt factors require triclinic box

Cannot deform the tilt factors of a simulation box unless it is a triclinic (non-orthogonal) box.

## Fix deform volume setting is invalid

Cannot use volume style unless other dimensions are being controlled.

## Fix deposit and fix rigid/small not using same molecule template ID

Self-explanatory.

## Fix deposit and fix shake not using same molecule template ID

Self-explanatory.

## Fix deposit molecule must have atom types

The defined molecule does not specify atom types.

## Fix deposit molecule must have coordinates

The defined molecule does not specify coordinates.

## Fix deposit molecule template ID must be same as atom\_style template ID

When using atom\_style template, you cannot deposit molecules that are not in that template.

## Fix deposit region cannot be dynamic

Only static regions can be used with fix deposit.

## Fix deposit region does not support a bounding box

Not all regions represent bounded volumes. You cannot use such a region with the fix deposit command.

## Fix deposit shake fix does not exist

Self-explanatory.

## Fix efield requires atom attribute q or mu

The atom style defined does not have this attribute.

## Fix efield with dipoles cannot use atom-style variables

This option is not supported.

## Fix evaporate molecule requires atom attribute molecule

The atom style being used does not define a molecule ID.

## Fix external callback function not set

This must be done by an external program in order to use this fix.

## Fix for fix ave/atom not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/atom is requesting a value on a non-allowed timestep.

## Fix for fix ave/chunk not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/chunk is requesting a value on a non-allowed timestep.

## Fix for fix ave/correlate not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/correlate is requesting a value on a non-allowed timestep.

## Fix for fix ave/histo not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/histo is requesting a value on a non-allowed timestep.

## Fix for fix ave/spatial not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/spatial is requesting a value on a non-allowed timestep.

## Fix for fix ave/time not computed at compatible time

Fixes generate their values on specific timesteps. Fix ave/time is requesting a value on a non-allowed timestep.

## Fix for fix store/state not computed at compatible time

Fixes generate their values on specific timesteps. Fix store/state is requesting a value on a non-allowed timestep.

## Fix for fix vector not computed at compatible time

Fixes generate their values on specific timesteps. Fix vector is requesting a value on a non-allowed timestep.

## Fix freeze requires atom attribute torque

The atom style defined does not have this attribute.

## Fix gcmc and fix shake not using same molecule template ID

Self-explanatory.

## Fix gcmc atom has charge, but atom style does not

Self-explanatory.

## Fix gcmc cannot exchange individual atoms belonging to a molecule

This is an error since you should not delete only one atom of a molecule. The user has specified atomic (non- molecular) gas exchanges, but an atom belonging to a molecule could be deleted.

## Fix gcmc does not (yet) work with atom\_style template

Self-explanatory.

## Fix gcmc molecule command requires that atoms have molecule attributes

Should not choose the gcmc molecule feature if no molecules are being simulated. The general molecule flag is off, but gcmc’s molecule flag is on.

## Fix gcmc molecule has charges, but atom style does not

Self-explanatory.

## Fix gcmc molecule must have atom types

The defined molecule does not specify atom types.

## Fix gcmc molecule must have coordinates

The defined molecule does not specify coordinates.

## Fix gcmc molecule template ID must be same as atom\_style template ID

When using atom\_style template, you cannot insert molecules that are not in that template.

## Fix gcmc put atom outside box

This should not normally happen. Contact the developers.

## Fix gcmc ran out of available atom IDs

See the setting for tagint in the src/lmptype.h file.

## Fix gcmc ran out of available molecule IDs

See the setting for tagint in the src/lmptype.h file.

## Fix gcmc region cannot be dynamic

Only static regions can be used with fix gcmc.

## Fix gcmc region does not support a bounding box

Not all regions represent bounded volumes. You cannot use such a region with the fix gcmc command.

## Fix gcmc region extends outside simulation box

Self-explanatory.

## Fix gcmc shake fix does not exist

Self-explanatory.

## Fix gld c coefficients must be >= 0

Self-explanatory.

## Fix gld needs more prony series coefficients

Self-explanatory.

## Fix gld prony terms must be > 0

Self-explanatory.

## Fix gld series type must be pprony for now

Self-explanatory.

## Fix gld start temperature must be >= 0

Self-explanatory.

## Fix gld stop temperature must be >= 0

Self-explanatory.

## Fix gld tau coefficients must be > 0

Self-explanatory.

## Fix halt variable is not equal-style variable

Self-explanatory.

## Fix heat group has no atoms

Self-explanatory.

## Fix heat kinetic energy of an atom went negative

This will cause the velocity rescaling about to be performed by fix heat to be invalid.

## Fix heat kinetic energy went negative

This will cause the velocity rescaling about to be performed by fix heat to be invalid.

## Fix in variable not computed at compatible time

Fixes generate their values on specific timesteps. The variable is requesting the values on a non-allowed timestep.

## Fix langevin angmom is not yet implemented with kokkos

This option is not yet available.

## Fix langevin angmom requires atom style ellipsoid

Self-explanatory.

## Fix langevin angmom requires extended particles

This fix option cannot be used with point particles.

## Fix langevin gjf and respa are not compatible

Self-explanatory.

## Fix langevin gjf cannot have period equal to dt/2

If the period is equal to dt/2 then division by zero will happen.

## Fix langevin gjf should come before fix nve

Self-explanatory.

## Fix langevin gjf with tbias is not yet implemented with kokkos

This option is not yet available.

## Fix langevin omega is not yet implemented with kokkos

This option is not yet available.

## Fix langevin omega requires atom style sphere

Self-explanatory.

## Fix langevin omega requires extended particles

One of the particles has radius 0.0.

## Fix langevin period must be > 0.0

The time window for temperature relaxation must be > 0

## Fix langevin variable returned negative temperature

Self-explanatory.

## Fix momentum group has no atoms

Self-explanatory.

## Fix move cannot define z or vz variable for 2d problem

Self-explanatory.

## Fix move cannot rotate aroung non z-axis for 2d problem

Self-explanatory.

## Fix move cannot set linear z motion for 2d problem

Self-explanatory.

## Fix move cannot set wiggle z motion for 2d problem

Self-explanatory.

## Fix msst compute ID does not compute potential energy

Self-explanatory.

## Fix msst compute ID does not compute pressure

Self-explanatory.

## Fix msst compute ID does not compute temperature

Self-explanatory.

## Fix msst requires a periodic box

Self-explanatory.

## Fix msst tscale must satisfy 0 <= tscale < 1

Self-explanatory.

## Fix npt/nph has tilted box too far in one step - periodic cell is too far from equilibrium state

Self-explanatory. The change in the box tilt is too extreme on a short timescale.

## Fix numdiff requires an atom map, see atom\_modify

Self-explanatory. Efficient loop over all atoms for numerical difference requires an atom map.

## Fix numdiff requires consecutive atom IDs

Self-explanatory. Efficient loop over all atoms for numerical difference requires consecutive atom IDs.

## Fix numdiff/virial must use group all

Virial contributions computed by this fix are computed on all atoms.

## Fix nve/asphere requires extended particles

This fix can only be used for particles with a shape setting.

## Fix nve/asphere/noforce requires atom style ellipsoid

Self-explanatory.

## Fix nve/asphere/noforce requires extended particles

One of the particles is not an ellipsoid.

## Fix nve/body requires atom style body

Self-explanatory.

## Fix nve/body requires bodies

This fix can only be used for particles that are bodies.

## Fix nve/line can only be used for 2d simulations

Self-explanatory.

## Fix nve/line requires atom style line

Self-explanatory.

## Fix nve/line requires line particles

Self-explanatory.

## Fix nve/sphere dipole requires atom attribute mu

An atom style with this attribute is needed.

## Fix nve/sphere requires atom style sphere

Self-explanatory.

## Fix nve/sphere requires extended particles

This fix can only be used for particles of a finite size.

## Fix nve/tri can only be used for 3d simulations

Self-explanatory.

## Fix nve/tri requires atom style tri

Self-explanatory.

## Fix nve/tri requires tri particles

Self-explanatory.

## Fix nvt/nph/npt asphere requires extended particles

The shape setting for a particle in the fix group has shape = 0.0, which means it is a point particle.

## Fix nvt/nph/npt body requires bodies

Self-explanatory.

## Fix nvt/nph/npt sphere requires atom style sphere

Self-explanatory.

## Fix nvt/npt/nph damping parameters must be > 0.0

Self-explanatory.

## Fix nvt/npt/nph dilate group ID does not exist

Self-explanatory.

## Fix nvt/sphere requires extended particles

This fix can only be used for particles of a finite size.

## Fix orient/fcc file open failed

The fix orient/fcc command could not open a specified file.

## Fix orient/fcc file read failed

The fix orient/fcc command could not read the needed parameters from a specified file.

## Fix orient/fcc found self twice

The neighbor lists used by fix orient/fcc are messed up. If this error occurs, it is likely a bug, so send an email to the [developers](https://www.lammps.org/authors.html).

## Fix peri neigh does not exist

Somehow a fix that the pair style defines has been deleted.

## Fix pour and fix rigid/small not using same molecule template ID

Self-explanatory.

## Fix pour and fix shake not using same molecule template ID

Self-explanatory.

## Fix pour insertion count per timestep is 0

Self-explanatory.

## Fix pour molecule must have atom types

The defined molecule does not specify atom types.

## Fix pour molecule must have coordinates

The defined molecule does not specify coordinates.

## Fix pour molecule template ID must be same as atom style template ID

When using atom\_style template, you cannot pour molecules that are not in that template.

## Fix pour polydisperse fractions do not sum to 1.0

Self-explanatory.

## Fix pour region ID does not exist

Self-explanatory.

## Fix pour region cannot be dynamic

Only static regions can be used with fix pour.

## Fix pour region does not support a bounding box

Not all regions represent bounded volumes. You cannot use such a region with the fix pour command.

## Fix pour requires atom attributes radius, rmass

The atom style defined does not have these attributes.

## Fix pour rigid fix does not exist

Self-explanatory.

## Fix pour shake fix does not exist

Self-explanatory.

## Fix press/berendsen damping parameters must be > 0.0

Self-explanatory.

## Fix property/atom cannot specify mol twice

Self-explanatory.

## Fix property/atom cannot specify q twice

Self-explanatory.

## Fix property/atom mol when atom\_style already has molecule attribute

Self-explanatory.

## Fix property/atom q when atom\_style already has charge attribute

Self-explanatory.

## Fix property/atom vector name already exists

The name for an integer or floating-point vector must be unique.

## Fix qeq has negative upper Taper radius cutoff

Self-explanatory.

## Fix qeq/comb group has no atoms

Self-explanatory.

## Fix qeq/comb requires atom attribute q

An atom style with charge must be used to perform charge equilibration.

## Fix qeq/dynamic group has no atoms

Self-explanatory.

## Fix qeq/dynamic requires atom attribute q

Self-explanatory.

## Fix qeq/fire group has no atoms

Self-explanatory.

## Fix qeq/fire requires atom attribute q

Self-explanatory.

## Fix qeq/point group has no atoms

Self-explanatory.

## Fix qeq/point has insufficient QEq matrix size

Occurs when number of neighbor atoms for an atom increased too much during a run. Increase SAFE\_ZONE and MIN\_CAP in fix\_qeq.h and re-compile.

## Fix qeq/point requires atom attribute q

Self-explanatory.

## Fix qeq/shielded group has no atoms

Self-explanatory.

## Fix qeq/shielded has insufficient QEq matrix size

Occurs when number of neighbor atoms for an atom increased too much during a run. Increase SAFE\_ZONE and MIN\_CAP in fix\_qeq.h and re-compile.

## Fix qeq/shielded requires atom attribute q

Self-explanatory.

## Fix qeq/slater could not extract params from pair coul/streitz

This should not happen unless pair coul/streitz has been altered.

## Fix qeq/slater group has no atoms

Self-explanatory.

## Fix qeq/slater has insufficient QEq matrix size

Occurs when number of neighbor atoms for an atom increased too much during a run. Increase SAFE\_ZONE and MIN\_CAP in fix\_qeq.h and re-compile.

## Fix qeq/slater requires atom attribute q

Self-explanatory.

## Fix reax/bonds numbonds > nsbmax\_most

The limit of the number of bonds expected by the ReaxFF force field was exceeded.

## Fix recenter group has no atoms

Self-explanatory.

## Fix restrain requires an atom map, see atom\_modify

Self-explanatory.

## Fix rigid atom has non-zero image flag in a non-periodic dimension

Image flags for non-periodic dimensions should not be set.

## Fix rigid file has no lines

Self-explanatory.

## Fix rigid langevin period must be > 0.0

Self-explanatory.

## Fix rigid molecule requires atom attribute molecule

Self-explanatory.

## Fix rigid npt/nph dilate group ID does not exist

Self-explanatory.

## Fix rigid npt/nph does not yet allow triclinic box

This is a current restriction in LAMMPS.

## Fix rigid npt/nph period must be > 0.0

Self-explanatory.

## Fix rigid npt/small t\_chain should not be less than 1

Self-explanatory.

## Fix rigid npt/small t\_order must be 3 or 5

Self-explanatory.

## Fix rigid nvt/npt/nph damping parameters must be > 0.0

Self-explanatory.

## Fix rigid nvt/small t\_chain should not be less than 1

Self-explanatory.

## Fix rigid nvt/small t\_iter should not be less than 1

Self-explanatory.

## Fix rigid nvt/small t\_order must be 3 or 5

Self-explanatory.

## Fix rigid xy torque cannot be on for 2d simulation

Self-explanatory.

## Fix rigid z force cannot be on for 2d simulation

Self-explanatory.

## Fix rigid/npt period must be > 0.0

Self-explanatory.

## Fix rigid/npt temperature order must be 3 or 5

Self-explanatory.

## Fix rigid/npt/small period must be > 0.0

Self-explanatory.

## Fix rigid/nvt period must be > 0.0

Self-explanatory.

## Fix rigid/nvt temperature order must be 3 or 5

Self-explanatory.

## Fix rigid/nvt/small period must be > 0.0

Self-explanatory.

## Fix rigid/small atom has non-zero image flag in a non-periodic dimension

Image flags for non-periodic dimensions should not be set.

## Fix rigid/small langevin period must be > 0.0

Self-explanatory.

## Fix rigid/small molecule must have atom types

The defined molecule does not specify atom types.

## Fix rigid/small molecule must have coordinates

The defined molecule does not specify coordinates.

## Fix rigid/small npt/nph period must be > 0.0

Self-explanatory.

## Fix rigid/small nvt/npt/nph damping parameters must be > 0.0

Self-explanatory.

## Fix rigid/small nvt/npt/nph dilate group ID does not exist

Self-explanatory.

## Fix rigid/small requires an atom map, see atom\_modify

Self-explanatory.

## Fix rigid/small requires atom attribute molecule

Self-explanatory.

## Fix rigid: Bad principal moments

The principal moments of inertia computed for a rigid body are not within the required tolerances.

## Fix shake cannot be used with minimization

Cannot use fix shake while doing an energy minimization since it turns off bonds that should contribute to the energy.

## Fix shake molecule template must have shake info

The defined molecule does not specify SHAKE information.

## Fix spring couple group ID does not exist

Self-explanatory.

## Fix srd can only currently be used with comm\_style brick

This is a current restriction in LAMMPS.

## Fix srd lamda must be >= 0.6 of SRD grid size

This is a requirement for accuracy reasons.

## Fix srd no-slip requires atom attribute torque

This is because the SRD collisions will impart torque to the solute particles.

## Fix srd requires SRD particles all have same mass

Self-explanatory.

## Fix srd requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Fix srd requires newton pair on

Self-explanatory.

## Fix store/state compute array is accessed out-of-range

Self-explanatory.

## Fix store/state compute does not calculate a per-atom array

The compute calculates a per-atom vector.

## Fix store/state compute does not calculate a per-atom vector

The compute calculates a per-atom vector.

## Fix store/state compute does not calculate per-atom values

Computes that calculate global or local quantities cannot be used with fix store/state.

## Fix store/state fix array is accessed out-of-range

Self-explanatory.

## Fix store/state fix does not calculate a per-atom array

The fix calculates a per-atom vector.

## Fix store/state fix does not calculate a per-atom vector

The fix calculates a per-atom array.

## Fix store/state fix does not calculate per-atom values

Fixes that calculate global or local quantities cannot be used with fix store/state.

## Fix store/state for atom property that is not allocated

Self-explanatory.

## Fix store/state variable is not atom-style variable

Only atom-style variables calculate per-atom quantities.

## Fix temp/berendsen period must be > 0.0

Self-explanatory.

## Fix temp/berendsen variable returned negative temperature

Self-explanatory.

## Fix temp/csld is not compatible with fix rattle or fix shake

These two commands cannot currently be used together with fix temp/csld.

## Fix temp/csld variable returned negative temperature

Self-explanatory.

## Fix temp/csvr variable returned negative temperature

Self-explanatory.

## Fix temp/rescale variable returned negative temperature

Self-explanatory.

## Fix tfmc displacement length must be > 0

Self-explanatory.

## Fix tfmc is not compatible with fix shake

These two commands cannot currently be used together.

## Fix tfmc temperature must be > 0

Self-explanatory.

## Fix thermal/conductivity swap value must be positive

Self-explanatory.

## Fix tmd must come after integration fixes

Any fix tmd command must appear in the input script after all time integration fixes (nve, nvt, npt). See the fix tmd documentation for details.

## Fix ttm electron temperatures must be > 0.0

Self-explanatory.

## Fix ttm electronic\_density must be > 0.0

Self-explanatory.

## Fix ttm electronic\_specific\_heat must be > 0.0

Self-explanatory.

## Fix ttm electronic\_thermal\_conductivity must be >= 0.0

Self-explanatory.

## Fix ttm gamma\_p must be > 0.0

Self-explanatory.

## Fix ttm gamma\_s must be >= 0.0

Self-explanatory.

## Fix ttm number of nodes must be > 0

Self-explanatory.

## Fix ttm v\_0 must be >= 0.0

Self-explanatory.

## Fix used in compute chunk/atom not computed at compatible time

The chunk/atom compute cannot query the output of the fix on a timestep it is needed.

## Fix used in compute reduce not computed at compatible time

Fixes generate their values on specific timesteps. Compute reduce is requesting a value on a non-allowed timestep.

## Fix used in compute slice not computed at compatible time

Fixes generate their values on specific timesteps. Compute slice is requesting a value on a non-allowed timestep.

## Fix vector cannot set output array intensive/extensive from these inputs

The inputs to the command have conflicting intensive/extensive attributes. You need to use more than one fix vector command.

## Fix vector compute does not calculate a scalar

Self-explanatory.

## Fix vector compute does not calculate a vector

Self-explanatory.

## Fix vector compute vector is accessed out-of-range

Self-explanatory.

## Fix vector fix does not calculate a scalar

Self-explanatory.

## Fix vector fix does not calculate a vector

Self-explanatory.

## Fix vector fix vector is accessed out-of-range

Self-explanatory.

## Fix vector variable is not equal-style variable

Self-explanatory.

## Fix viscosity swap value must be positive

Self-explanatory.

## Fix viscosity vtarget value must be positive

Self-explanatory.

## Fix wall cutoff <= 0.0

Self-explanatory.

## Fix wall/colloid requires atom style sphere

Self-explanatory.

## Fix wall/colloid requires extended particles

One of the particles has radius 0.0.

## Fix wall/gran is incompatible with Pair style

Must use a granular pair style to define the parameters needed for this fix.

## Fix wall/gran requires atom style sphere

Self-explanatory.

## Fix wall/piston command only available at zlo

The face keyword must be zlo.

## Fix wall/region colloid requires atom style sphere

Self-explanatory.

## Fix wall/region colloid requires extended particles

One of the particles has radius 0.0.

## Fix wall/region cutoff <= 0.0

Self-explanatory.

## Fix\_modify pressure ID does not compute pressure

The compute ID assigned to the fix must compute pressure.

## Fix\_modify temperature ID does not compute temperature

The compute ID assigned to the fix must compute temperature.

## For triclinic deformation, specified target stress must be hydrostatic

Triclinic pressure control is allowed using the tri keyword, but non-hydrostatic pressure control can not be used in this case.

## Found no restart file matching pattern

When using a “\*” in the restart file name, no matching file was found.

## GPU library not compiled for this accelerator

Self-explanatory.

## GPU package does not (yet) work with atom\_style template

Self-explanatory.

## GPU particle split must be set to 1 for this pair style.

For this pair style, you cannot run part of the force calculation on the host. See the package command.

## GPUs are requested but Kokkos has not been compiled for CUDA

Re-compile Kokkos with CUDA support to use GPUs.

## Ghost velocity forward comm not yet implemented with Kokkos

This is a current restriction.

## Gmask function in equal-style variable formula

Gmask is per-atom operation.

## Gravity changed since fix pour was created

The gravity vector defined by fix gravity must be static.

## Gravity must point in -y to use with fix pour in 2d

Self-explanatory.

## Gravity must point in -z to use with fix pour in 3d

Self-explanatory.

## Grmask function in equal-style variable formula

Grmask is per-atom operation.

## Group ID does not exist

A group ID used in the group command does not exist.

## Group ID in variable formula does not exist

Self-explanatory.

## Group all cannot be made dynamic

This operation is not allowed.

## Group command before simulation box is defined

The group command cannot be used before a read\_data, read\_restart, or create\_box command.

## Group dynamic cannot reference itself

Self-explanatory.

## Group dynamic parent group cannot be dynamic

Self-explanatory.

## Group dynamic parent group does not exist

Self-explanatory.

## Group region ID does not exist

A region ID used in the group command does not exist.

## If read\_dump purges it cannot replace or trim

These operations are not compatible. See the read\_dump doc page for details.

## Illegal . . . command

Self-explanatory. Check the input script syntax and compare to the documentation for the command. You can use -echo screen as a command-line option when running LAMMPS to see the offending line.

## Illegal COMB parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal COMB3 parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal Stillinger-Weber parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal Tersoff parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal Vashishta parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal compute voronoi/atom command (occupation and (surface or edges))

Self-explanatory.

## Illegal coul/streitz parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal dump\_modify sfactor value (must be > 0.0)

Self-explanatory.

## Illegal dump\_modify tfactor value (must be > 0.0)

Self-explanatory.

## Illegal fix gcmc gas mass <= 0

The computed mass of the designated gas molecule or atom type was less than or equal to zero.

## Illegal fix tfmc random seed

Seeds can only be nonzero positive integers.

## Illegal fix wall/piston velocity

The piston velocity must be positive.

## Illegal integrate style

Self-explanatory.

## Illegal nb3b/harmonic parameter

One or more of the coefficients defined in the potential file is invalid.

## Illegal number of angle table entries

There must be at least 2 table entries.

## Illegal number of bond table entries

There must be at least 2 table entries.

## Illegal number of pair table entries

There must be at least 2 table entries.

## Illegal or unset periodicity in restart

This error should not normally occur unless the restart file is invalid.

## Illegal range increment value

The increment must be >= 1.

## Illegal simulation box

The lower bound of the simulation box is greater than the upper bound.

## Illegal size double vector read requested

This error should not normally occur unless the restart file is invalid.

## Illegal size integer vector read requested

This error should not normally occur unless the restart file is invalid.

## Illegal size string or corrupt restart

This error should not normally occur unless the restart file is invalid.

## Imageint setting in lmptype.h is invalid

Imageint must be as large or larger than smallint.

## Imageint setting in lmptype.h is not compatible

Format of imageint stored in restart file is not consistent with LAMMPS version you are running. See the settings in src/lmptype.h

## Improper atom missing in delete\_bonds

The delete\_bonds command cannot find one or more atoms in a particular improper on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid improper.

## Improper atom missing in set command

The set command cannot find one or more atoms in a particular improper on a particular processor. The pairwise cutoff is too short or the atoms are too far apart to make a valid improper.

## Improper atoms %d %d %d %d missing on proc %d at step %ld

One or more of 4 atoms needed to compute a particular improper are missing on this processor. Typically this is because the pairwise cutoff is set too short or the improper has blown apart and an atom is too far away.

## Improper atoms missing on proc %d at step %ld

One or more of 4 atoms needed to compute a particular improper are missing on this processor. Typically this is because the pairwise cutoff is set too short or the improper has blown apart and an atom is too far away.

## Improper coeff for hybrid has invalid style

Improper style hybrid uses another improper style as one of its coefficients. The improper style used in the improper\_coeff command or read from a restart file is not recognized.

## Improper coeffs are not set

No improper coefficients have been assigned in the data file or via the improper\_coeff command.

## Improper style hybrid cannot have hybrid as an argument

Self-explanatory.

## Improper style hybrid cannot have none as an argument

Self-explanatory.

## Improper style hybrid cannot use same improper style twice

Self-explanatory.

## Improper\_coeff command before improper\_style is defined

Coefficients cannot be set in the data file or via the improper\_coeff command until an improper\_style has been assigned.

## Improper\_coeff command before simulation box is defined

The improper\_coeff command cannot be used before a read\_data, read\_restart, or create\_box command.

## Improper\_coeff command when no impropers allowed

The chosen atom style does not allow for impropers to be defined.

## Improper\_style command when no impropers allowed

The chosen atom style does not allow for impropers to be defined.

## Impropers assigned incorrectly

Impropers read in from the data file were not assigned correctly to atoms. This means there is something invalid about the topology definitions.

## Impropers defined but no improper types

The data file header lists improper but no improper types.

## Incompatible KIM Simulator Model

The requested KIM Simulator Model was defined for a different MD code and thus is not compatible with LAMMPS.

## Incompatible units for KIM Simulator Model

The selected unit style is not compatible with the requested KIM Simulator Model.

## Incomplete use of variables in create\_atoms command

The var and set options must be used together.

## Inconsistent iparam/jparam values in fix bond/create command

If itype and jtype are the same, then their maxbond and newtype settings must also be the same.

## Inconsistent line segment in data file

The end points of the line segment are not equal distances from the center point which is the atom coordinate.

## Inconsistent triangle in data file

The centroid of the triangle as defined by the corner points is not the atom coordinate.

## Inconsistent use of finite-size particles by molecule template molecules

Not all of the molecules define a radius for their constituent particles.

## Incorrect # of floating-point values in Bodies section of data file

See page for body style.

## Incorrect # of integer values in Bodies section of data file

See page for body style.

## Incorrect %s format in data file

A section of the data file being read by fix property/atom does not have the correct number of values per line.

## Incorrect SNAP parameter file

The file cannot be parsed correctly, check its internal syntax.

## Incorrect args for angle coefficients

Self-explanatory. Check the input script or data file.

## Incorrect args for bond coefficients

Self-explanatory. Check the input script or data file.

## Incorrect args for dihedral coefficients

Self-explanatory. Check the input script or data file.

## Incorrect args for improper coefficients

Self-explanatory. Check the input script or data file.

## Incorrect args for pair coefficients

Self-explanatory. Check the input script or data file.

## Incorrect args in pair\_style command

Self-explanatory.

## Incorrect atom format in data file

Number of values per atom line in the data file is not consistent with the atom style.

## Incorrect atom format in neb file

The number of fields per line is not what expected.

## Incorrect bonus data format in data file

See the read\_data page for a description of how various kinds of bonus data must be formatted for certain atom styles.

## Incorrect boundaries with slab Ewald

Must have periodic x,y dimensions and non-periodic z dimension to use 2d slab option with Ewald.

## Incorrect boundaries with slab EwaldDisp

Must have periodic x,y dimensions and non-periodic z dimension to use 2d slab option with Ewald.

## Incorrect boundaries with slab PPPM

Must have periodic x,y dimensions and non-periodic z dimension to use 2d slab option with PPPM.

## Incorrect boundaries with slab PPPMDisp

Must have periodic x,y dimensions and non-periodic z dimension to use 2d slab option with pppm/disp.

## Incorrect conversion in format string

A format style variable was not using either a %f, a %g, or a %e conversion. Or an immediate variable with format suffix was not using either a %f, a %g or a %e conversion in the format suffix.

## Incorrect element names in ADP potential file

The element names in the ADP file do not match those requested.

## Incorrect element names in EAM potential file

The element names in the EAM file do not match those requested.

## Incorrect format of . . . section in data file

Number or type of values per line in the given section of the data file is not consistent with the requirements for this section.

## Incorrect format in COMB potential file

Incorrect number of words per line in the potential file.

## Incorrect format in COMB3 potential file

Incorrect number of words per line in the potential file.

## Incorrect format in MEAM library file

Incorrect number of words per line in the potential file.

## Incorrect format in SNAP coefficient file

Incorrect number of words per line in the coefficient file.

## Incorrect format in SNAP parameter file

Incorrect number of words per line in the parameter file.

## Incorrect format in Stillinger-Weber potential file

Incorrect number of words per line in the potential file.

## Incorrect format in TMD target file

Format of file read by fix tmd command is incorrect.

## Incorrect format in Tersoff potential file

Incorrect number of words per line in the potential file.

## Incorrect format in Vashishta potential file

Incorrect number of words per line in the potential file.

## Incorrect format in coul/streitz potential file

Incorrect number of words per line in the potential file.

## Incorrect format in nb3b/harmonic potential file

Incorrect number of words per line in the potential file.

## Incorrect integer value in Bodies section of data file

See page for body style.

## Incorrect multiplicity arg for dihedral coefficients

Self-explanatory. Check the input script or data file.

## Incorrect number of elements in potential file

Self-explanatory.

## Incorrect rigid body format in fix rigid file

The number of fields per line is not what expected.

## Incorrect rigid body format in fix rigid/small file

The number of fields per line is not what expected.

## Incorrect sign arg for dihedral coefficients

Self-explanatory. Check the input script or data file.

## Incorrect table format check for element types

Self-explanatory.

## Incorrect velocity format in data file

Each atom style defines a format for the Velocity section of the data file. The read-in lines do not match.

## Incorrect weight arg for dihedral coefficients

Self-explanatory. Check the input script or data file.

## Index between variable brackets must be positive

Self-explanatory.

## Indexed per-atom vector in variable formula without atom map

Accessing a value from an atom vector requires the ability to lookup an atom index, which is provided by an atom map. An atom map does not exist (by default) for non-molecular problems. Using the atom\_modify map command will force an atom map to be created.

## Initial temperatures not all set in fix ttm

Self-explanatory.

## Input line quote not followed by white-space

An end quote must be followed by white-space.

## Insertion region extends outside simulation box

Self-explanatory.

## Insufficient Jacobi rotations for POEMS body

Eigensolve for rigid body was not sufficiently accurate.

## Insufficient Jacobi rotations for body nparticle

Eigensolve for rigid body was not sufficiently accurate.

## Insufficient Jacobi rotations for rigid body

Eigensolve for rigid body was not sufficiently accurate.

## Insufficient Jacobi rotations for rigid molecule

Eigensolve for rigid body was not sufficiently accurate.

## Insufficient Jacobi rotations for triangle

The calculation of the inertia tensor of the triangle failed. This should not happen if it is a reasonably shaped triangle.

## Insufficient memory on accelerator

There is insufficient memory on one of the devices specified for the gpu package

## Internal error in atom\_style body

This error should not occur. Contact the developers.

## Invalid -reorder N value

Self-explanatory.

## Invalid Angles section in molecule file

Self-explanatory.

## Invalid Bonds section in molecule file

Self-explanatory.

## Invalid Boolean syntax in if command

Self-explanatory.

## Invalid Charges section in molecule file

Self-explanatory.

## Invalid Coords section in molecule file

Self-explanatory.

## Invalid Diameters section in molecule file

Self-explanatory.

## Invalid Dihedrals section in molecule file

Self-explanatory.

## Invalid Impropers section in molecule file

Self-explanatory.

## Invalid Kokkos command-line args

Self-explanatory. See Section 2.7 of the manual for details.

## Invalid LAMMPS restart file

The file does not appear to be a LAMMPS restart file since it does not contain the correct magic string at the beginning.

## Invalid Masses section in molecule file

Self-explanatory.

## Invalid molecule ID in molecule file

Molecule ID must be a non-zero positive integer.

## Invalid Molecules section in molecule file

Self-explanatory.

## Invalid REAX atom type

There is a mis-match between LAMMPS atom types and the elements listed in the ReaxFF force field file.

## Invalid Special Bond Counts section in molecule file

Self-explanatory.

## Invalid Types section in molecule file

Self-explanatory.

## Invalid angle count in molecule file

Self-explanatory.

## Invalid angle table length

Length must be 2 or greater.

## Invalid angle type in Angles section of data file

Angle type must be positive integer and within range of specified angle types.

## Invalid angle type in Angles section of molecule file

Self-explanatory.

## Invalid angle type index for fix shake

Self-explanatory.

## Invalid args for non-hybrid pair coefficients

“NULL” is only supported in pair\_coeff calls when using pair hybrid

## Invalid argument to factorial %d

N must be >= 0 and <= 167, otherwise the factorial result is too large.

## Invalid atom ID in %s section of data file

An atom in a section of the data file being read by fix property/atom has an invalid atom ID that is <= 0 or > the maximum existing atom ID.

## Invalid atom ID in Angles section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Angles section of molecule file

Self-explanatory.

## Invalid atom ID in Atoms section of data file

Atom IDs must be positive integers.

## Invalid atom ID in Bodies section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Bonds section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Bonds section of molecule file

Self-explanatory.

## Invalid atom ID in Bonus section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Dihedrals section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Fragments section of molecule file

Self-explanatory.

## Invalid atom ID in Impropers section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in Velocities section of data file

Atom IDs must be positive integers and within range of defined atoms.

## Invalid atom ID in dihedrals section of molecule file

Self-explanatory.

## Invalid atom ID in impropers section of molecule file

Self-explanatory.

## Invalid atom ID in variable file

Self-explanatory.

## Invalid atom IDs in neb file

An ID in the file was not found in the system.

## Invalid atom diameter in molecule file

Diameters must be >= 0.0.

## Invalid atom mass for fix shake

Mass specified in fix shake command must be > 0.0.

## Invalid atom mass in molecule file

Masses must be > 0.0.

## Invalid atom type in Atoms section of data file

Atom types must range from 1 to specified # of types.

## Invalid atom type in create\_atoms command

The create\_box command specified the range of valid atom types. An invalid type is being requested.

## Invalid atom type in create\_atoms mol command

The atom types in the defined molecule are added to the value specified in the create\_atoms command, as an offset. The final value for each atom must be between 1 to N, where N is the number of atom types.

## Invalid atom type in fix atom/swap command

The atom type specified in the atom/swap command does not exist.

## Invalid atom type in fix bond/create command

Self-explanatory.

## Invalid atom type in fix deposit command

Self-explanatory.

## Invalid atom type in fix deposit mol command

The atom types in the defined molecule are added to the value specified in the create\_atoms command, as an offset. The final value for each atom must be between 1 to N, where N is the number of atom types.

## Invalid atom type in fix gcmc command

The atom type specified in the gcmc command does not exist.

## Invalid atom type in fix pour command

Self-explanatory.

## Invalid atom type in fix pour mol command

The atom types in the defined molecule are added to the value specified in the create\_atoms command, as an offset. The final value for each atom must be between 1 to N, where N is the number of atom types.

## Invalid atom type in molecule file

Atom types must range from 1 to specified # of types.

## Invalid atom type in neighbor exclusion list

Atom types must range from 1 to Ntypes inclusive.

## Invalid atom type index for fix shake

Atom types must range from 1 to Ntypes inclusive.

## Invalid atom types in pair\_write command

Atom types must range from 1 to Ntypes inclusive.

## Invalid atom vector in variable formula

The atom vector is not recognized.

## Invalid atom\_style body command

No body style argument was provided.

## Invalid atom\_style command

Self-explanatory.

## Invalid attribute in dump custom command

Self-explanatory.

## Invalid attribute in dump local command

Self-explanatory.

## Invalid attribute in dump modify command

Self-explanatory.

## Invalid basis setting in create\_atoms command

The basis index must be between 1 to N where N is the number of basis atoms in the lattice. The type index must be between 1 to N where N is the number of atom types.

## Invalid basis setting in fix append/atoms command

The basis index must be between 1 to N where N is the number of basis atoms in the lattice. The type index must be between 1 to N where N is the number of atom types.

## Invalid bin bounds in compute chunk/atom

The lo/hi values are inconsistent.

## Invalid bin bounds in fix ave/spatial

The lo/hi values are inconsistent.

## Invalid body nparticle command

Arguments in atom-style command are not correct.

## Invalid bond count in molecule file

Self-explanatory.

## Invalid bond table length

Length must be 2 or greater.

## Invalid bond type in Bonds section of data file

Bond type must be positive integer and within range of specified bond types.

## Invalid bond type in Bonds section of molecule file

Self-explanatory.

## Invalid bond type in create\_bonds command

Self-explanatory.

## Invalid bond type in fix bond/break command

Self-explanatory.

## Invalid bond type in fix bond/create command

Self-explanatory.

## Invalid bond type index for fix shake

Self-explanatory. Check the fix shake command in the input script.

## Invalid coeffs for this dihedral style

Cannot set class 2 coeffs in data file for this dihedral style.

## Invalid color in dump\_modify command

The specified color name was not in the list of recognized colors. See the dump\_modify doc page.

## Invalid color map min/max values

The min/max values are not consistent with either each other or with values in the color map.

## Invalid command-line argument

One or more command-line arguments is invalid. Check the syntax of the command you are using to launch LAMMPS.

## Invalid compute ID in variable formula

The compute is not recognized.

## Invalid create\_atoms rotation vector for 2d model

The rotation vector can only have a z component.

## Invalid custom OpenCL parameter string.

There are not enough or too many parameters in the custom string for package GPU.

## Invalid cutoff in comm\_modify command

Specified cutoff must be >= 0.0.

## Invalid cutoffs in pair\_write command

Inner cutoff must be larger than 0.0 and less than outer cutoff.

## Invalid d1 or d2 value for pair colloid coeff

Neither d1 or d2 can be < 0.

## Invalid data file section: Angle Coeffs

Atom style does not allow angles.

## Invalid data file section: AngleAngle Coeffs

Atom style does not allow impropers.

## Invalid data file section: AngleAngleTorsion Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: AngleTorsion Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: Angles

Atom style does not allow angles.

## Invalid data file section: Bodies

Atom style does not allow bodies.

## Invalid data file section: Bond Coeffs

Atom style does not allow bonds.

## Invalid data file section: BondAngle Coeffs

Atom style does not allow angles.

## Invalid data file section: BondBond Coeffs

Atom style does not allow angles.

## Invalid data file section: BondBond13 Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: Bonds

Atom style does not allow bonds.

## Invalid data file section: Dihedral Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: Dihedrals

Atom style does not allow dihedrals.

## Invalid data file section: Ellipsoids

Atom style does not allow ellipsoids.

## Invalid data file section: EndBondTorsion Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: Improper Coeffs

Atom style does not allow impropers.

## Invalid data file section: Impropers

Atom style does not allow impropers.

## Invalid data file section: Lines

Atom style does not allow lines.

## Invalid data file section: MiddleBondTorsion Coeffs

Atom style does not allow dihedrals.

## Invalid data file section: Triangles

Atom style does not allow triangles.

## Invalid delta\_conf in tad command

The value must be between 0 and 1 inclusive.

## Invalid density in Atoms section of data file

Density value cannot be <= 0.0.

## Invalid density in set command

Density must be > 0.0.

## Invalid diameter in set command

Self-explanatory.

## Invalid dihedral count in molecule file

Self-explanatory.

## Invalid dihedral type in Dihedrals section of data file

Dihedral type must be positive integer and within range of specified dihedral types.

## Invalid dihedral type in dihedrals section of molecule file

Self-explanatory.

## Invalid dipole length in set command

Self-explanatory.

## Invalid displace\_atoms rotate axis for 2d

Axis must be in z direction.

## Invalid dump dcd filename

Filenames used with the dump dcd style cannot be binary or compressed or cause multiple files to be written.

## Invalid dump frequency

Dump frequency must be 1 or greater.

## Invalid dump image element name

The specified element name was not in the standard list of elements. See the dump\_modify doc page.

## Invalid dump image filename

The file produced by dump image cannot be binary and must be for a single processor.

## Invalid dump image theta value

Theta must be between 0.0 and 180.0 inclusive.

## Invalid dump image zoom value

Zoom value must be > 0.0.

## Invalid dump movie filename

The file produced by dump movie cannot be binary or compressed and must be a single file for a single processor.

## Invalid dump xtc filename

Filenames used with the dump xtc style cannot be binary or compressed or cause multiple files to be written.

## Invalid dump xyz filename

Filenames used with the dump xyz style cannot be binary or cause files to be written by each processor.

## Invalid dump\_modify threshold operator

Operator keyword used for threshold specification in not recognized.

## Invalid entry in -reorder file

Self-explanatory.

## Invalid fix ID in variable formula

The fix is not recognized.

## Invalid fix ave/time off column

Self-explanatory.

## Invalid fix box/relax command for a 2d simulation

Fix box/relax styles involving the z dimension cannot be used in a 2d simulation.

## Invalid fix box/relax command pressure settings

If multiple dimensions are coupled, those dimensions must be specified.

## Invalid fix box/relax pressure settings

Settings for coupled dimensions must be the same.

## Invalid fix halt attribute

Self-explanatory.

## Invalid fix halt operator

Self-explanatory.

## Invalid fix nvt/npt/nph command for a 2d simulation

Cannot control z dimension in a 2d model.

## Invalid fix nvt/npt/nph command pressure settings

If multiple dimensions are coupled, those dimensions must be specified.

## Invalid fix nvt/npt/nph pressure settings

Settings for coupled dimensions must be the same.

## Invalid fix press/berendsen for a 2d simulation

The z component of pressure cannot be controlled for a 2d model.

## Invalid fix press/berendsen pressure settings

Settings for coupled dimensions must be the same.

## Invalid fix qeq parameter file

Element index > number of atom types.

## Invalid fix rigid npt/nph command for a 2d simulation

Cannot control z dimension in a 2d model.

## Invalid fix rigid npt/nph command pressure settings

If multiple dimensions are coupled, those dimensions must be specified.

## Invalid fix rigid/small npt/nph command for a 2d simulation

Cannot control z dimension in a 2d model.

## Invalid fix rigid/small npt/nph command pressure settings

If multiple dimensions are coupled, those dimensions must be specified.

## Invalid flag in force field section of restart file

Unrecognized entry in restart file.

## Invalid flag in header section of restart file

Unrecognized entry in restart file.

## Invalid flag in peratom section of restart file

The format of this section of the file is not correct.

## Invalid flag in type arrays section of restart file

Unrecognized entry in restart file.

## Invalid frequency in temper command

Nevery must be > 0.

## Invalid group ID in neigh\_modify command

A group ID used in the neigh\_modify command does not exist.

## Invalid group function in variable formula

Group function is not recognized.

## Invalid group in comm\_modify command

Self-explanatory.

## Invalid image up vector

Up vector cannot be (0,0,0).

## Invalid immediate variable

Syntax of immediate value is incorrect.

## Invalid improper count in molecule file

Self-explanatory.

## Invalid improper type in Impropers section of data file

Improper type must be positive integer and within range of specified improper types.

## Invalid improper type in impropers section of molecule file

Self-explanatory.

## Invalid index for non-body particles in compute body/local command

Only indices 1,2,3 can be used for non-body particles.

## Invalid index in compute body/local command

Self-explanatory.

## Invalid is\_active() function in variable formula

Self-explanatory.

## Invalid is\_available() function in variable formula

Self-explanatory.

## Invalid is\_defined() function in variable formula

Self-explanatory.

## Invalid keyword in angle table parameters

Self-explanatory.

## Invalid keyword in bond table parameters

Self-explanatory.

## Invalid keyword in compute angle/local command

Self-explanatory.

## Invalid keyword in compute bond/local command

Self-explanatory.

## Invalid keyword in compute dihedral/local command

Self-explanatory.

## Invalid keyword in compute improper/local command

Self-explanatory.

## Invalid keyword in compute pair/local command

Self-explanatory.

## Invalid keyword in compute property/atom command

Self-explanatory.

## Invalid keyword in compute property/chunk command

Self-explanatory.

## Invalid keyword in compute property/local command

Self-explanatory.

## Invalid keyword in dump cfg command

Self-explanatory.

## Invalid keyword in pair table parameters

Keyword used in list of table parameters is not recognized.

## Invalid length in set command

Self-explanatory.

## Invalid mass in set command

Self-explanatory.

## Invalid mass line in data file

Self-explanatory.

## Invalid mass value

Self-explanatory.

## Invalid math function in variable formula

Self-explanatory.

## Invalid math/group/special function in variable formula

Self-explanatory.

## Invalid option in lattice command for non-custom style

Certain lattice keywords are not supported unless the lattice style is “custom”.

## Invalid order of forces within respa levels

For respa, ordering of force computations within respa levels must obey certain rules. E.g. bonds cannot be compute less frequently than angles, pairwise forces cannot be computed less frequently than kspace, etc.

## Invalid pair table cutoff

Cutoffs in pair\_coeff command are not valid with read-in pair table.

## Invalid pair table length

Length of read-in pair table is invalid

## Invalid param file for fix qeq/shielded

Invalid value of gamma.

## Invalid param file for fix qeq/slater

Zeta value is 0.0.

## Invalid partitions in processors part command

Valid partitions are numbered 1 to N and the sender and receiver cannot be the same partition.

## Invalid python command

Self-explanatory. Check the input script syntax and compare to the documentation for the command. You can use -echo screen as a command-line option when running LAMMPS to see the offending line.

## Invalid radius in Atoms section of data file

Radius must be >= 0.0.

## Invalid random number seed in fix ttm command

Random number seed must be > 0.

## Invalid random number seed in set command

Random number seed must be > 0.

## Invalid replace values in compute reduce

Self-explanatory.

## Invalid rigid body ID in fix rigid file

The ID does not match the number of an existing ID of rigid bodies that are defined by the fix rigid command.

## Invalid rigid body ID in fix rigid/small file

The ID does not match the number of an existing ID of rigid bodies that are defined by the fix rigid/small command.

## Invalid run command N value

The number of timesteps must fit in a 32-bit integer. If you want to run for more steps than this, perform multiple shorter runs.

## Invalid run command start/stop value

Self-explanatory.

## Invalid run command upto value

Self-explanatory.

## Invalid seed for Marsaglia random # generator

The initial seed for this random number generator must be a positive integer less than or equal to 900 million.

## Invalid seed for Park random # generator

The initial seed for this random number generator must be a positive integer.

## Invalid shake angle type in molecule file

Self-explanatory.

## Invalid shake atom in molecule file

Self-explanatory.

## Invalid shake bond type in molecule file

Self-explanatory.

## Invalid shake flag in molecule file

Self-explanatory.

## Invalid shape in Ellipsoids section of data file

Self-explanatory.

## Invalid shape in Triangles section of data file

Two or more of the triangle corners are duplicate points.

## Invalid shape in set command

Self-explanatory.

## Invalid shear direction for fix wall/gran

Self-explanatory.

## Invalid special atom index in molecule file

Self-explanatory.

## Invalid special function in variable formula

Self-explanatory.

## Invalid style in pair\_write command

Self-explanatory. Check the input script.

## Invalid syntax in variable formula

Self-explanatory.

## Invalid t\_event in prd command

Self-explanatory.

## Invalid t\_event in tad command

The value must be greater than 0.

## Invalid template atom in Atoms section of data file

The atom indices must be between 1 to N, where N is the number of atoms in the template molecule the atom belongs to.

## Invalid template index in Atoms section of data file

The template indices must be between 1 to N, where N is the number of molecules in the template.

## Invalid thermo keyword in variable formula

The keyword is not recognized.

## Invalid threads\_per\_atom specified.

For 3-body potentials on the GPU, the threads\_per\_atom setting cannot be greater than 4 for NVIDIA GPUs.

## Invalid timestep reset for fix ave/atom

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid timestep reset for fix ave/chunk

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid timestep reset for fix ave/correlate

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid timestep reset for fix ave/histo

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid timestep reset for fix ave/spatial

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid timestep reset for fix ave/time

Resetting the timestep has invalidated the sequence of timesteps this fix needs to process.

## Invalid tmax in tad command

The value must be greater than 0.0.

## Invalid type for mass set

Mass command must set a type from 1-N where N is the number of atom types.

## Invalid label2type() function syntax in variable formula

The first argument must be a label map kind (atom, bond, angle, dihedral, or improper) and the second argument must be a valid type label that has been assigned to a numeric type.

## Invalid use of library file() function

This function is called through the library interface. This error should not occur. Contact the developers if it does.

## Invalid value in set command

The value specified for the setting is invalid, likely because it is too small or too large.

## Invalid variable evaluation in variable formula

A variable used in a formula could not be evaluated.

## Invalid variable in next command

Self-explanatory.

## Invalid variable name

Variable name used in an input script line is invalid.

## Invalid variable name in variable formula

Variable name is not recognized.

## Invalid variable style in special function next

Only file-style or atomfile-style variables can be used with next().

## Invalid variable style with next command

Variable styles *equal* and *world* cannot be used in a next command.

## Invalid volume in set command

Volume must be > 0.0.

## Invalid wiggle direction for fix wall/gran

Self-explanatory.

## Invoked angle equil angle on angle style none

Self-explanatory.

## Invoked angle single on angle style none

Self-explanatory.

## Invoked bond equil distance on bond style none

Self-explanatory.

## Invoked bond single on bond style none

Self-explanatory.

## Invoked pair single on pair style none

A command (e.g. a dump) attempted to invoke the single() function on a pair style none, which is illegal. You are probably attempting to compute per-atom quantities with an undefined pair style.

## Invoking coulombic in pair style lj/coul requires atom attribute q

The atom style defined does not have this attribute.

## Invoking coulombic in pair style lj/long/dipole/long requires atom attribute q

The atom style defined does not have these attributes.

## KIM Simulator Model has no Model definition

There is no model definition (key: model-defn) in the KIM Simulator Model. Please contact the OpenKIM database maintainers to verify and potentially correct this.

## KOKKOS package does not yet support comm\_style tiled

Self-explanatory.

## KOKKOS package requires a kokkos enabled atom\_style

Self-explanatory.

## KSpace accuracy must be > 0

The kspace accuracy designated in the input must be greater than zero.

## KSpace accuracy too large to estimate G vector

Reduce the accuracy request or specify gewald explicitly via the kspace\_modify command.

## KSpace accuracy too low

Requested accuracy must be less than 1.0.

## KSpace solver requires a pair style

No pair style is defined.

## KSpace style does not yet support triclinic geometries

The specified kspace style does not allow for non-orthogonal simulation boxes.

## KSpace style has not yet been set

Cannot use kspace\_modify command until a kspace style is set.

## KSpace style is incompatible with Pair style

Setting a kspace style requires that a pair style with matching long-range Coulombic or dispersion components be used.

## Keyword %s in MEAM parameter file not recognized

Self-explanatory.

## Kokkos has been compiled for CUDA but no GPUs are requested

One or more GPUs must be used when Kokkos is compiled for CUDA.

## Kspace\_modify mesh parameter must be all zero or all positive

Valid kspace mesh parameters are >0. The code will try to auto-detect suitable values when all three mesh sizes are set to zero (the default).

## Kspace\_modify mesh/disp parameter must be all zero or all positive

Valid kspace mesh/disp parameters are >0. The code will try to auto-detect suitable values when all three mesh sizes are set to zero **and** the required accuracy via *force/disp/real* as well as *force/disp/kspace* is set.

## Kspace style does not support compute group/group

Self-explanatory.

## Kspace style pppm/disp/tip4p requires newton on

Self-explanatory.

## Kspace style pppm/tip4p requires newton on

Self-explanatory.

## Kspace style requires atom attribute q

The atom style defined does not have these attributes.

## Kspace\_modify eigtol must be smaller than one

Self-explanatory.

## LAMMPS is not built with Python embedded

This is done by including the PYTHON package before LAMMPS is built. This is required to use python-style variables.

## LAMMPS unit\_style lj not supported by KIM models

Self-explanatory. Check the input script or data file.

## LJ6 off not supported in pair\_style buck/long/coul/long

Self-explanatory.

## Label map is incomplete: all types must be assigned a unique type label

For a given type-kind (atom types, bond types, etc.) to be written to the data file, all associated types must be assigned a type label, and each type label can be assigned to only one numeric type.

## Label wasn’t found in input script

Self-explanatory.

## Labelmap command before simulation box is defined

The labelmap command cannot be used before a read\_data, read\_restart, or create\_box command.

## Lattice orient vectors are not orthogonal

The three specified lattice orientation vectors must be mutually orthogonal.

## Lattice orient vectors are not right-handed

The three specified lattice orientation vectors must create a right-handed coordinate system such that a1 cross a2

= a3.

## Lattice primitive vectors are collinear

The specified lattice primitive vectors do not for a unit cell with non-zero volume.

## Lattice settings are not compatible with 2d simulation

One or more of the specified lattice vectors has a non-zero z component.

## Lattice spacings are invalid

Each x,y,z spacing must be > 0.

## Lattice style incompatible with simulation dimension

2d simulation can use sq, sq2, or hex lattice. 3d simulation can use sc, bcc, or fcc lattice.

## Log of zero/negative value in variable formula

Self-explanatory.

## Lost atoms via balance: original %ld current %ld

This should not occur. Report the problem to the developers.

## Lost atoms: original %ld current %ld

Lost atoms are checked for each time thermo output is done. See the thermo\_modify lost command for options. Lost atoms usually indicate bad dynamics, e.g. atoms have been blown far out of the simulation box, or moved further than one processor’s subdomain away before reneighboring.

## MEAM library error %d

A call to the MEAM Fortran library returned an error.

## MPI\_LMP\_BIGINT and bigint in lmptype.h are not compatible

The size of the MPI datatype does not match the size of a bigint.

## MPI\_LMP\_TAGINT and tagint in lmptype.h are not compatible

The size of the MPI datatype does not match the size of a tagint.

## MSM can only currently be used with comm\_style brick

This is a current restriction in LAMMPS.

## MSM grid is too large

The global MSM grid is larger than OFFSET in one or more dimensions. OFFSET is currently set to 16384. You likely need to decrease the requested accuracy.

## MSM order must be 4, 6, 8, or 10

This is a limitation of the MSM implementation in LAMMPS: the MSM order can only be 4, 6, 8, or 10.

## Mass command before simulation box is defined

The mass command cannot be used before a read\_data, read\_restart, or create\_box command.

## Matrix factorization to split dispersion coefficients failed

This should not normally happen. Contact the developers.

## Min\_style command before simulation box is defined

The min\_style command cannot be used before a read\_data, read\_restart, or create\_box command.

## Minimization could not find thermo\_pe compute

This compute is created by the thermo command. It must have been explicitly deleted by a uncompute command.

## Minimize command before simulation box is defined

The minimize command cannot be used before a read\_data, read\_restart, or create\_box command.

## Mismatched brackets in variable

Self-explanatory.

## Mismatched compute in variable formula

A compute is referenced incorrectly or a compute that produces per-atom values is used in an equal-style variable formula.

## Mismatched fix in variable formula

A fix is referenced incorrectly or a fix that produces per-atom values is used in an equal-style variable formula.

## Mismatched parameter in MEAM library file: z!=lat

The coordination number and lattice do not match, check that consistent values are given.

## Mismatched variable in variable formula

A variable is referenced incorrectly or an atom-style variable that produces per-atom values is used in an equal- style variable formula.

## Modulo 0 in variable formula

Self-explanatory.

## Molecule IDs too large for compute chunk/atom

The IDs must not be larger than can be stored in a 32-bit integer since chunk IDs are 32-bit integers.

## Molecule auto special bond generation overflow

Counts exceed maxspecial setting for other atoms in system.

## Molecule file has angles but no nangles setting

Self-explanatory.

## Molecule file has body params but no setting for them

Self-explanatory.

## Molecule file has bonds but no nbonds setting

Self-explanatory.

## Molecule file has dihedrals but no ndihedrals setting

Self-explanatory.

## Molecule file has fragments but no nfragments setting

Self-explanatory.

## Molecule file has impropers but no nimpropers setting

Self-explanatory.

## Molecule file has no Body Doubles section

Self-explanatory.

## Molecule file has no Body Integers section

Self-explanatory.

## Molecule file has no Fragments section

Self-explanatory.

## Molecule file has special flags but no bonds

Self-explanatory.

## Molecule file needs both Special Bond sections

Self-explanatory.

## Molecule file requires atom style body

Self-explanatory.

## Molecule file shake flags not before shake atoms

The order of the two sections is important.

## Molecule file shake flags not before shake bonds

The order of the two sections is important.

## Molecule file shake info is incomplete

All 3 SHAKE sections are needed.

## Molecule file special list does not match special count

The number of values in an atom’s special list does not match count.

## Molecule file z center-of-mass must be 0.0 for 2d

Self-explanatory.

## Molecule file z coord must be 0.0 for 2d

Self-explanatory.

## Molecule natoms must be 1 for body particle

Self-explanatory.

## Molecule sizescale must be 1.0 for body particle

Self-explanatory.

## Molecule template ID for atom\_style template does not exist

Self-explanatory.

## Molecule template ID for create\_atoms does not exist

Self-explanatory.

## Molecule template ID for fix deposit does not exist

Self-explanatory.

## Molecule template ID for fix gcmc does not exist

Self-explanatory.

## Molecule template ID for fix pour does not exist

Self-explanatory.

## Molecule template ID for fix rigid/small does not exist

Self-explanatory.

## Molecule template ID for fix shake does not exist

Self-explanatory.

## Molecule template ID must be alphanumeric or underscore characters

Self-explanatory.

## Molecule topology/atom exceeds system topology/atom

The number of bonds, angles, etc per-atom in the molecule exceeds the system setting. See the create\_box command for how to specify these values.

## Molecule topology type exceeds system topology type

The number of bond, angle, etc types in the molecule exceeds the system setting. See the create\_box command for how to specify these values.

## More than one fix deform

Only one fix deform can be defined at a time.

## More than one fix freeze

Only one of these fixes can be defined, since the granular pair potentials access it.

## More than one fix shake

Only one fix shake can be defined.

## Mu not allowed when not using semi-grand in fix atom/swap command

Self-explanatory.

## Must define angle\_style before Angle Coeffs

Must use an angle\_style command before reading a data file that defines Angle Coeffs.

## Must define angle\_style before BondAngle Coeffs

Must use an angle\_style command before reading a data file that defines Angle Coeffs.

## Must define angle\_style before BondBond Coeffs

Must use an angle\_style command before reading a data file that defines Angle Coeffs.

## Must define bond\_style before Bond Coeffs

Must use a bond\_style command before reading a data file that defines Bond Coeffs.

## Must define dihedral\_style before AngleAngleTorsion Coeffs

Must use a dihedral\_style command before reading a data file that defines AngleAngleTorsion Coeffs.

## Must define dihedral\_style before AngleTorsion Coeffs

Must use a dihedral\_style command before reading a data file that defines AngleTorsion Coeffs.

## Must define dihedral\_style before BondBond13 Coeffs

Must use a dihedral\_style command before reading a data file that defines BondBond13 Coeffs.

## Must define dihedral\_style before Dihedral Coeffs

Must use a dihedral\_style command before reading a data file that defines Dihedral Coeffs.

## Must define dihedral\_style before EndBondTorsion Coeffs

Must use a dihedral\_style command before reading a data file that defines EndBondTorsion Coeffs.

## Must define dihedral\_style before MiddleBondTorsion Coeffs

Must use a dihedral\_style command before reading a data file that defines MiddleBondTorsion Coeffs.

## Must define improper\_style before AngleAngle Coeffs

Must use an improper\_style command before reading a data file that defines AngleAngle Coeffs.

## Must define improper\_style before Improper Coeffs

Must use an improper\_style command before reading a data file that defines Improper Coeffs.

## Must define pair\_style before Pair Coeffs

Must use a pair\_style command before reading a data file that defines Pair Coeffs.

## Must define pair\_style before PairIJ Coeffs

Must use a pair\_style command before reading a data file that defines PairIJ Coeffs.

## Must have more than one processor partition to temper

Cannot use the temper command with only one processor partition. Use the -partition command-line option.

## Must not have multiple fixes change box parameter . . .

Self-explanatory.

## Must read Angle Type Labels before Angles

An Angle Type Labels section of a data file must come before the Angles section.

## Must read Atom Type Labels before Atoms

An Atom Type Labels section of a data file must come before the Atoms section.

## Must read Atoms before Angles

The Atoms section of a data file must come before an Angles section.

## Must read Atoms before Bodies

The Atoms section of a data file must come before a Bodies section.

## Must read Atoms before Bonds

The Atoms section of a data file must come before a Bonds section.

## Must read Atoms before Dihedrals

The Atoms section of a data file must come before a Dihedrals section.

## Must read Atoms before Ellipsoids

The Atoms section of a data file must come before a Ellipsoids section.

## Must read Atoms before Impropers

The Atoms section of a data file must come before an Impropers section.

## Must read Atoms before Lines

The Atoms section of a data file must come before a Lines section.

## Must read Atoms before Triangles

The Atoms section of a data file must come before a Triangles section.

## Must read Atoms before Velocities

The Atoms section of a data file must come before a Velocities section.

## Must read Bond Type Labels before Bonds

A Bond Type Labels section of a data file must come before the Bonds section.

## Must read Dihedral Type Labels before Dihedrals

An Dihedral Type Labels section of a data file must come before the Dihedrals section.

## Must read Improper Type Labels before Impropers

An Improper Type Labels section of a data file must come before the Impropers section.

## Must re-specify non-restarted pair style (xxx) after read\_restart

For pair styles, that do not store their settings in a restart file, it must be defined with a new ‘pair\_style’ command after read\_restart.

## Must set both respa inner and outer

Cannot use just the inner or outer option with respa without using the other.

## Must set number of threads via package omp command

Because you are using the OPENMP package, set the number of threads via its settings, not by the pair\_style snap nthreads setting.

## Must shrink-wrap piston boundary

The boundary style of the face where the piston is applied must be of type s (shrink-wrapped).

## Must specify a region in fix deposit

The region keyword must be specified with this fix.

## Must specify a region in fix pour

Self-explanatory.

## Must specify at least 2 types in fix atom/swap command

Self-explanatory.

## Must use ‘kim\_style init’ command before simulation box is defined

Self-explanatory.

## Must use ‘kim\_style define’ command after simulation box is defined

Self-explanatory.

## Must use ‘kim\_style init’ command before ‘kim\_style define’

Self-explanatory.

## Must use ‘kspace\_modify pressure/scalar no’ for rRESPA with kspace\_style MSM

The kspace scalar pressure option cannot (yet) be used with rRESPA.

## Must use ‘kspace\_modify pressure/scalar no’ for tensor components with kspace\_style msm

Otherwise MSM will compute only a scalar pressure. See the kspace\_modify command for details on this setting.

## Must use ‘kspace\_modify pressure/scalar no’ to obtain per-atom virial with kspace\_style MSM

The kspace scalar pressure option cannot be used to obtain per-atom virial.

## Must use ‘kspace\_modify pressure/scalar no’ with GPU MSM Pair styles

The kspace scalar pressure option is not (yet) compatible with GPU MSM Pair styles.

## Must use ‘kspace\_modify pressure/scalar no’ with kspace\_style msm/cg

The kspace scalar pressure option is not compatible with kspace\_style msm/cg.

## Must use -in switch with multiple partitions

A multi-partition simulation cannot read the input script from stdin. The -in command-line option must be used to specify a file.

## Must use Kokkos half/thread or full neighbor list with threads or GPUs

Using Kokkos half-neighbor lists with threading is not allowed.

## Must use a block or cylinder region with fix pour

Self-explanatory.

## Must use a block region with fix pour for 2d simulations

Self-explanatory.

## Must use a bond style with TIP4P potential

TIP4P potentials assume bond lengths in water are constrained by a fix shake command.

## Must use a molecular atom style with fix poems molecule

Self-explanatory.

## Must use a z-axis cylinder region with fix pour

Self-explanatory.

## Must use an angle style with TIP4P potential

TIP4P potentials assume angles in water are constrained by a fix shake command.

## Must use atom map style array with Kokkos

See the atom\_modify map command.

## Must use atom style with molecule IDs with fix bond/swap

Self-explanatory.

## Must use pair\_style comb or comb3 with fix qeq/comb

Self-explanatory.

## Must use variable energy with fix addforce

Must define an energy variable when applying a dynamic force during minimization.

## Must use variable energy with fix efield

You must define an energy when performing a minimization with a variable E-field.

## NEB command before simulation box is defined

Self-explanatory.

## NEB requires damped dynamics minimizer

Use a different minimization style.

## NEB requires use of fix neb

Self-explanatory.

## NL ramp in wall/piston only implemented in zlo for now

The ramp keyword can only be used for piston applied to face zlo.

## Need nswaptypes mu values in fix atom/swap command

Self-explanatory.

## Needed bonus data not in data file

Some atom styles require bonus data. See the read\_data page for details.

## Needed molecular topology not in data file

The header of the data file indicated bonds, angles, etc would be included, but they are not present.

## Neigh\_modify exclude molecule requires atom attribute molecule

Self-explanatory.

## Neigh\_modify include group != atom\_modify first group

Self-explanatory.

## Neighbor delay must be 0 or multiple of every setting

The delay and every parameters set via the neigh\_modify command are inconsistent. If the delay setting is non-zero, then it must be a multiple of the every setting.

## Neighbor include group not allowed with ghost neighbors

This is a current restriction within LAMMPS.

## Neighbor list overflow, boost neigh\_modify one

There are too many neighbors of a single atom. Use the neigh\_modify command to increase the max number of neighbors allowed for one atom. You may also want to boost the page size.

## Neighbor multi not yet enabled for ghost neighbors

This is a current restriction within LAMMPS.

## Neighbor multi not yet enabled for granular

Self-explanatory.

## Neighbor multi not yet enabled for rRESPA

Self-explanatory.

## Neighbor page size must be >= 10x the one atom setting

This is required to prevent wasting too much memory.

## New atom IDs exceed maximum allowed ID

See the setting for tagint in the src/lmptype.h file.

## New bond exceeded bonds per atom in create\_bonds

See the read\_data command for info on using the “extra/bond/per/atom” keyword to allow for additional bonds to be formed

## New bond exceeded bonds per atom in fix bond/create

See the read\_data command for info on using the “extra/bond/per/atom” keyword to allow for additional bonds to be formed

## New bond exceeded special list size in fix bond/create

See the “read\_data extra/special/per/atom” command (or the “create\_box extra/special/per/atom” command) for info on how to leave space in the special bonds list to allow for additional bonds to be formed.

## Newton bond change after simulation box is defined

The newton command cannot be used to change the newton bond value after a read\_data, read\_restart, or cre- ate\_box command.

## Next command must list all universe and uloop variables

This is to ensure they stay in sync.

## No Kspace style defined for compute group/group

Self-explanatory.

## No OpenMP support compiled in

An OpenMP flag is set, but LAMMPS was not built with OpenMP support.

## No angle style is defined for compute angle/local

Self-explanatory.

## No angles allowed with this atom style

Self-explanatory.

## No atoms in data file

The header of the data file indicated that atoms would be included, but they are not present.

## No basis atoms in lattice

Basis atoms must be defined for lattice style user.

## No bodies allowed with this atom style

Self-explanatory. Check data file.

## No bond style is defined for compute bond/local

Self-explanatory.

## No bonds allowed with this atom style

Self-explanatory.

## No box information in dump. You have to use ‘box no’

Self-explanatory.

## No count or invalid atom count in molecule file

The number of atoms must be specified.

## No dihedral style is defined for compute dihedral/local

Self-explanatory.

## No dihedrals allowed with this atom style

Self-explanatory.

## No dump custom arguments specified

The dump custom command requires that atom quantities be specified to output to dump file.

## No dump local arguments specified

Self-explanatory.

## No ellipsoids allowed with this atom style

Self-explanatory. Check data file.

## No fix gravity defined for fix pour

Gravity is required to use fix pour.

## No improper style is defined for compute improper/local

Self-explanatory.

## No impropers allowed with this atom style

Self-explanatory.

## No input values for fix ave/spatial

Self-explanatory.

## No lines allowed with this atom style

Self-explanatory. Check data file.

## No matching element in ADP potential file

The ADP potential file does not contain elements that match the requested elements.

## No matching element in EAM potential file

The EAM potential file does not contain elements that match the requested elements.

## No molecule topology allowed with atom style template

The data file cannot specify the number of bonds, angles, etc, because this info if inferred from the molecule templates.

## No overlap of box and region for create\_atoms

Self-explanatory.

## No pair coul/streitz for fix qeq/slater

These commands must be used together.

## No pair hbond/dreiding coefficients set

Self-explanatory.

## No pair style defined for compute group/group

Cannot calculate group interactions without a pair style defined.

## No pair style is defined for compute pair/local

Self-explanatory.

## No pair style is defined for compute property/local

Self-explanatory.

## No rigid bodies defined

The fix specification did not end up defining any rigid bodies.

## No triangles allowed with this atom style

Self-explanatory. Check data file.

## No values in fix ave/chunk command

Self-explanatory.

## No values in fix ave/time command

Self-explanatory.

## Non digit character between brackets in variable

Self-explanatory.

## Non integer # of swaps in temper command

Swap frequency in temper command must evenly divide the total # of timesteps.

## Non-numeric box dimensions - simulation unstable

The box size has apparently blown up.

## Non-zero atom IDs with atom\_modify id = no

Self-explanatory.

## Non-zero read\_data shift z value for 2d simulation

Self-explanatory.

## Nprocs not a multiple of N for -reorder

Self-explanatory.

## Number of core atoms != number of shell atoms

There must be a one-to-one pairing of core and shell atoms.

## Numeric index is out of bounds

A command with an argument that specifies an integer or range of integers is using a value that is less than 1 or greater than the maximum allowed limit.

## One or more Atom IDs is negative

Atom IDs must be positive integers.

## One or more atom IDs is too big

The limit on atom IDs is set by the SMALLBIG, BIGBIG, SMALLSMALL setting in your LAMMPS build. See the *Build settings* page for more info.

## One or more atom IDs is zero

Either all atoms IDs must be zero or none of them.

## One or more atoms belong to multiple rigid bodies

Two or more rigid bodies defined by the fix rigid command cannot contain the same atom.

## One or more rigid bodies are a single particle

Self-explanatory.

## One or zero atoms in rigid body

Any rigid body defined by the fix rigid command must contain 2 or more atoms.

## Only 2 types allowed when not using semi-grand in fix atom/swap command

Self-explanatory.

## Only one cut-off allowed when requesting all long

Self-explanatory.

## Only one cutoff allowed when requesting all long

Self-explanatory.

## Only zhi currently implemented for fix append/atoms

Self-explanatory.

## Out of range atoms - cannot compute MSM

One or more atoms are attempting to map their charge to a MSM grid point that is not owned by a processor. This is likely for one of two reasons, both of them bad. First, it may mean that an atom near the boundary of a processor’s subdomain has moved more than 1/2 the *neighbor skin distance* without neighbor lists being rebuilt and atoms being migrated to new processors. This also means you may be missing pairwise interactions that need to be computed. The solution is to change the re-neighboring criteria via the *neigh\_modify* command. The safest settings are “delay 0 every 1 check yes”. Second, it may mean that an atom has moved far outside a processor’s subdomain or even the entire simulation box. This indicates bad physics, e.g. due to highly overlapping atoms, too large a timestep, etc.

## Out of range atoms - cannot compute PPPM

One or more atoms are attempting to map their charge to a PPPM grid point that is not owned by a processor. This is likely for one of two reasons, both of them bad. First, it may mean that an atom near the boundary of a processor’s subdomain has moved more than 1/2 the *neighbor skin distance* without neighbor lists being rebuilt and atoms being migrated to new processors. This also means you may be missing pairwise interactions that need to be computed. The solution is to change the re-neighboring criteria via the *neigh\_modify* command. The safest settings are “delay 0 every 1 check yes”. Second, it may mean that an atom has moved far outside a processor’s subdomain or even the entire simulation box. This indicates bad physics, e.g. due to highly overlapping atoms, too large a timestep, etc.

## Out of range atoms - cannot compute PPPMDisp

One or more atoms are attempting to map their charge to a PPPM grid point that is not owned by a processor. This is likely for one of two reasons, both of them bad. First, it may mean that an atom near the boundary of a processor’s subdomain has moved more than 1/2 the *neighbor skin distance* without neighbor lists being rebuilt and atoms being migrated to new processors. This also means you may be missing pairwise interactions that need to be computed. The solution is to change the re-neighboring criteria via the *neigh\_modify* command. The safest settings are “delay 0 every 1 check yes”. Second, it may mean that an atom has moved far outside a processor’s subdomain or even the entire simulation box. This indicates bad physics, e.g. due to highly overlapping atoms, too large a timestep, etc.

## Overflow of allocated fix vector storage

This should not normally happen if the fix correctly calculated how long the vector will grow to. Contact the developers.

## Overlapping large/large in pair colloid

This potential is infinite when there is an overlap.

## Overlapping small/large in pair colloid

This potential is infinite when there is an overlap.

## POEMS fix must come before NPT/NPH fix

NPT/NPH fix must be defined in input script after all poems fixes, else the fix contribution to the pressure virial is incorrect.

## PPPM can only currently be used with comm\_style brick

This is a current restriction in LAMMPS.

## PPPM grid is too large

The global PPPM grid is larger than OFFSET in one or more dimensions. OFFSET is currently set to 4096. You likely need to decrease the requested accuracy.

## PPPM grid stencil extends beyond nearest neighbor processor

This is not allowed if the kspace\_modify overlap setting is no.

## PPPM order < minimum allowed order

The default minimum order is 2. This can be reset by the kspace\_modify minorder command.

## PPPM order cannot be < 2 or > than %d

This is a limitation of the PPPM implementation in LAMMPS.

## PPPMDisp Coulomb grid is too large

The global PPPM grid is larger than OFFSET in one or more dimensions. OFFSET is currently set to 4096. You likely need to decrease the requested accuracy.

## PPPMDisp Dispersion grid is too large

The global PPPM grid is larger than OFFSET in one or more dimensions. OFFSET is currently set to 4096. You likely need to decrease the requested accuracy.

## PPPMDisp can only currently be used with comm\_style brick

This is a current restriction in LAMMPS.

## PPPMDisp coulomb order cannot be greater than %d

This is a limitation of the PPPM implementation in LAMMPS.

## PPPMDisp used but no parameters set, for further information please see the pppm/disp documentation

An efficient and accurate usage of the pppm/disp requires settings via the kspace\_modify command. Please see the pppm/disp documentation for further instructions.

## PRD command before simulation box is defined

The prd command cannot be used before a read\_data, read\_restart, or create\_box command.

## PRD nsteps must be multiple of t\_event

Self-explanatory.

## PRD t\_corr must be multiple of t\_event

Self-explanatory.

## Package command after simulation box is defined

The package command cannot be used after a read\_data, read\_restart, or create\_box command.

## Package gpu command without GPU package installed

The GPU package must be installed via “make yes-gpu” before LAMMPS is built.

## Package intel command without INTEL package installed

The INTEL package must be installed via “make yes-intel” before LAMMPS is built.

## Package kokkos command without KOKKOS package enabled

The KOKKOS package must be installed via “make yes-kokkos” before LAMMPS is built, and the “-k on” must be used to enable the package.

## Package omp command without OPENMP package installed

The OPENMP package must be installed via “make yes-openmp” before LAMMPS is built.

## Pair body requires atom style body

Self-explanatory.

## Pair body requires body style nparticle

This pair style is specific to the nparticle body style.

## Pair brownian requires atom style sphere

Self-explanatory.

## Pair brownian requires extended particles

One of the particles has radius 0.0.

## Pair brownian requires monodisperse particles

All particles must be the same finite size.

## Pair brownian/poly requires atom style sphere

Self-explanatory.

## Pair brownian/poly requires extended particles

One of the particles has radius 0.0.

## Pair brownian/poly requires newton pair off

Self-explanatory.

## Pair coeff for hybrid has invalid style

Style in pair coeff must have been listed in pair\_style command.

## Pair coul/wolf requires atom attribute q

The atom style defined does not have this attribute.

## Pair cutoff < Respa interior cutoff

One or more pairwise cutoffs are too short to use with the specified rRESPA cutoffs.

## Pair dipole/cut requires atom attributes q, mu, torque

The atom style defined does not have these attributes.

## Pair dipole/cut/gpu requires atom attributes q, mu, torque

The atom style defined does not have this attribute.

## Pair dipole/long requires atom attributes q, mu, torque

The atom style defined does not have these attributes.

## Pair dipole/sf/gpu requires atom attributes q, mu, torque

The atom style defined does not one or more of these attributes.

## Pair distance < table inner cutoff

Two atoms are closer together than the pairwise table allows.

## Pair distance > table outer cutoff

Two atoms are further apart than the pairwise table allows.

## Pair dpd requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair gayberne epsilon a,b,c coeffs are not all set

Each atom type involved in pair\_style gayberne must have these 3 coefficients set at least once.

## Pair gayberne requires atom style ellipsoid

Self-explanatory.

## Pair gayberne requires atoms with same type have same shape

Self-explanatory.

## Pair gayberne/gpu requires atom style ellipsoid

Self-explanatory.

## Pair gayberne/gpu requires atoms with same type have same shape

Self-explanatory.

## Pair granular requires atom attributes radius, rmass

The atom style defined does not have these attributes.

## Pair granular requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair granular with shear history requires newton pair off

This is a current restriction of the implementation of pair granular styles with history.

## Pair hybrid single calls do not support per sub-style special bond values

Self-explanatory.

## Pair hybrid sub-style does not support single call

You are attempting to invoke a single() call on a pair style that does not support it.

## Pair hybrid sub-style is not used

No pair\_coeff command used a sub-style specified in the pair\_style command.

## Pair inner cutoff < Respa interior cutoff

One or more pairwise cutoffs are too short to use with the specified rRESPA cutoffs.

## Pair inner cutoff >= Pair outer cutoff

The specified cutoffs for the pair style are inconsistent.

## Pair line/lj requires atom style line

Self-explanatory.

## Pair lj/long/dipole/long requires atom attributes mu, torque

The atom style defined does not have these attributes.

## Pair lubricate requires atom style sphere

Self-explanatory.

## Pair lubricate requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair lubricate requires monodisperse particles

All particles must be the same finite size.

## Pair lubricate/poly requires atom style sphere

Self-explanatory.

## Pair lubricate/poly requires extended particles

One of the particles has radius 0.0.

## Pair lubricate/poly requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair lubricate/poly requires newton pair off

Self-explanatory.

## Pair lubricateU requires atom style sphere

Self-explanatory.

## Pair lubricateU requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair lubricateU requires monodisperse particles

All particles must be the same finite size.

## Pair lubricateU/poly requires ghost atoms store velocity

Use the comm\_modify vel yes command to enable this.

## Pair lubricateU/poly requires newton pair off

Self-explanatory.

## Pair peri lattice is not identical in x, y, and z

The lattice defined by the lattice command must be cubic.

## Pair peri requires a lattice be defined

Use the lattice command for this purpose.

## Pair peri requires an atom map, see atom\_modify

Even for atomic systems, an atom map is required to find Peridynamic bonds. Use the atom\_modify command to define one.

## Pair resquared epsilon a,b,c coeffs are not all set

Self-explanatory.

## Pair resquared epsilon and sigma coeffs are not all set

Self-explanatory.

## Pair resquared requires atom style ellipsoid

Self-explanatory.

## Pair resquared requires atoms with same type have same shape

Self-explanatory.

## Pair resquared/gpu requires atom style ellipsoid

Self-explanatory.

## Pair resquared/gpu requires atoms with same type have same shape

Self-explanatory.

## Pair style AIREBO requires atom IDs

This is a requirement to use the AIREBO potential.

## Pair style AIREBO requires newton pair on

See the newton command. This is a restriction to use the AIREBO potential.

## Pair style BOP requires atom IDs

This is a requirement to use the BOP potential.

## Pair style BOP requires newton pair on

See the newton command. This is a restriction to use the BOP potential.

## Pair style COMB requires atom IDs

This is a requirement to use the AIREBO potential.

## Pair style COMB requires atom attribute q

Self-explanatory.

## Pair style COMB requires newton pair on

See the newton command. This is a restriction to use the COMB potential.

## Pair style COMB3 requires atom IDs

This is a requirement to use the COMB3 potential.

## Pair style COMB3 requires atom attribute q

Self-explanatory.

## Pair style COMB3 requires newton pair on

See the newton command. This is a restriction to use the COMB3 potential.

## Pair style LCBOP requires atom IDs

This is a requirement to use the LCBOP potential.

## Pair style LCBOP requires newton pair on

See the newton command. This is a restriction to use the Tersoff potential.

## Pair style MEAM requires newton pair on

See the newton command. This is a restriction to use the MEAM potential.

## Pair style SNAP requires newton pair on

See the newton command. This is a restriction to use the SNAP potential.

## Pair style Stillinger-Weber requires atom IDs

This is a requirement to use the SW potential.

## Pair style Stillinger-Weber requires newton pair on

See the newton command. This is a restriction to use the SW potential.

## Pair style Tersoff requires atom IDs

This is a requirement to use the Tersoff potential.

## Pair style Tersoff requires newton pair on

See the newton command. This is a restriction to use the Tersoff potential.

## Pair style Vashishta requires atom IDs

This is a requirement to use the Vashishta potential.

## Pair style Vashishta requires newton pair on

See the newton command. This is a restriction to use the Vashishta potential.

## Pair style bop requires comm ghost cutoff at least 3x larger than %g

Use the communicate ghost command to set this. See the pair bop page for more details.

## Pair style born/coul/long requires atom attribute q

An atom style that defines this attribute must be used.

## Pair style born/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style born/coul/wolf requires atom attribute q

The atom style defined does not have this attribute.

## Pair style buck/coul/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style buck/coul/long requires atom attribute q

The atom style defined does not have these attributes.

## Pair style buck/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style buck/long/coul/long requires atom attribute q

The atom style defined does not have this attribute.

## Pair style coul/cut requires atom attribute q

The atom style defined does not have these attributes.

## Pair style coul/cut/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style coul/debye/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style coul/dsf requires atom attribute q

The atom style defined does not have this attribute.

## Pair style coul/dsf/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style coul/long/gpu requires atom attribute q

The atom style defined does not have these attributes.

## Pair style coul/streitz requires atom attribute q

Self-explanatory.

## Pair style does not have extra field requested by compute pair/local

The pair style does not support the pN value requested by the compute pair/local command.

## Pair style does not support bond\_style quartic

The pair style does not have a single() function, so it can not be invoked by bond\_style quartic.

## Pair style does not support compute group/group

The pair\_style does not have a single() function, so it cannot be invoked by the compute group/group command.

## Pair style does not support compute pair/local

The pair style does not have a single() function, so it can not be invoked by compute pair/local.

## Pair style does not support compute property/local

The pair style does not have a single() function, so it can not be invoked by fix bond/swap.

## Pair style does not support fix bond/swap

The pair style does not have a single() function, so it can not be invoked by fix bond/swap.

## Pair style does not support pair\_write

The pair style does not have a single() function, so it can not be invoked by pair write.

## Pair style does not support rRESPA inner/middle/outer

You are attempting to use rRESPA options with a pair style that does not support them.

## Pair style granular with history requires atoms have IDs

Atoms in the simulation do not have IDs, so history effects cannot be tracked by the granular pair potential.

## Pair style hbond/dreiding requires an atom map, see atom\_modify

Self-explanatory.

## Pair style hbond/dreiding requires atom IDs

Self-explanatory.

## Pair style hbond/dreiding requires molecular system

Self-explanatory.

## Pair style hbond/dreiding requires newton pair on

See the newton command for details.

## Pair style hybrid cannot have hybrid as an argument

Self-explanatory.

## Pair style hybrid cannot have none as an argument

Self-explanatory.

## Pair style is incompatible with KSpace style

If a pair style with a long-range Coulombic component is selected, then a kspace style must also be used.

## Pair style is incompatible with TIP4P KSpace style

The pair style does not have the requires TIP4P settings.

## Pair style lj/charmm/coul/charmm requires atom attribute q

The atom style defined does not have these attributes.

## Pair style lj/charmm/coul/long requires atom attribute q

The atom style defined does not have these attributes.

## Pair style lj/charmm/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/class2/coul/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/class2/coul/long requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/class2/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/cut/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/debye/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/dsf requires atom attribute q

The atom style defined does not have these attributes.

## Pair style lj/cut/coul/dsf/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/long requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/tip4p/cut requires atom IDs

This is a requirement to use this potential.

## Pair style lj/cut/tip4p/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style lj/cut/tip4p/cut requires newton pair on

See the newton command. This is a restriction to use this potential.

## Pair style lj/cut/tip4p/long requires atom IDs

There are no atom IDs defined in the system and the TIP4P potential requires them to find O,H atoms with a water molecule.

## Pair style lj/cut/tip4p/long requires atom attribute q

The atom style defined does not have these attributes.

## Pair style lj/cut/tip4p/long requires newton pair on

This is because the computation of constraint forces within a water molecule adds forces to atoms owned by other processors.

## Pair style lj/gromacs/coul/gromacs requires atom attribute q

An atom\_style with this attribute is needed.

## Pair style lj/long/dipole/long does not currently support respa

This feature is not yet supported.

## Pair style lj/long/tip4p/long requires atom IDs

There are no atom IDs defined in the system and the TIP4P potential requires them to find O,H atoms with a water molecule.

## Pair style lj/long/tip4p/long requires atom attribute q

The atom style defined does not have these attributes.

## Pair style lj/long/tip4p/long requires newton pair on

This is because the computation of constraint forces within a water molecule adds forces to atoms owned by other processors.

## Pair style lj/spica/coul/long/gpu requires atom attribute q

The atom style defined does not have this attribute.

## Pair style nb3b/harmonic requires atom IDs

This is a requirement to use this potential.

## Pair style nb3b/harmonic requires newton pair on

See the newton command. This is a restriction to use this potential.

## Pair style nm/cut/coul/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style nm/cut/coul/long requires atom attribute q

The atom style defined does not have this attribute.

## Pair style peri requires atom style peri

Self-explanatory.

## Pair style polymorphic requires atom IDs

This is a requirement to use the polymorphic potential.

## Pair style polymorphic requires newton pair on

See the newton command. This is a restriction to use the polymorphic potential.

## Pair style reax requires atom IDs

This is a requirement to use the ReaxFF potential.

## Pair style reax requires atom attribute q

The atom style defined does not have this attribute.

## Pair style reax requires newton pair on

This is a requirement to use the ReaxFF potential.

## Pair style requires a KSpace style

No kspace style is defined.

## Pair style requires use of kspace\_style ewald/disp

Self-explanatory.

## Pair style sw/gpu requires atom IDs

This is a requirement to use this potential.

## Pair style sw/gpu requires newton pair off

See the newton command. This is a restriction to use this potential.

## Pair style vashishta/gpu requires atom IDs

This is a requirement to use this potential.

## Pair style vashishta/gpu requires newton pair off

See the newton command. This is a restriction to use this potential.

## Pair style tersoff/gpu requires atom IDs

This is a requirement to use the tersoff/gpu potential.

## Pair style tersoff/gpu requires newton pair off

See the newton command. This is a restriction to use this pair style.

## Pair style tip4p/cut requires atom IDs

This is a requirement to use this potential.

## Pair style tip4p/cut requires atom attribute q

The atom style defined does not have this attribute.

## Pair style tip4p/cut requires newton pair on

See the newton command. This is a restriction to use this potential.

## Pair style tip4p/long requires atom IDs

There are no atom IDs defined in the system and the TIP4P potential requires them to find O,H atoms with a water molecule.

## Pair style tip4p/long requires atom attribute q

The atom style defined does not have these attributes.

## Pair style tip4p/long requires newton pair on

This is because the computation of constraint forces within a water molecule adds forces to atoms owned by other processors.

## Pair table cutoffs must all be equal to use with KSpace

When using pair style table with a long-range KSpace solver, the cutoffs for all atom type pairs must all be the same, since the long-range solver starts at that cutoff.

## Pair table parameters did not set N

List of pair table parameters must include N setting.

## Pair tersoff/zbl requires metal or real units

This is a current restriction of this pair potential.

## Pair tersoff/zbl/kk requires metal or real units

This is a current restriction of this pair potential.

## Pair tri/lj requires atom style tri

Self-explanatory.

## Pair yukawa/colloid requires atom style sphere

Self-explanatory.

## Pair yukawa/colloid requires atoms with same type have same radius

Self-explanatory.

## Pair yukawa/colloid/gpu requires atom style sphere

Self-explanatory.

## PairKIM only works with 3D problems

This is a current limitation.

## Pair\_coeff command before pair\_style is defined

Self-explanatory.

## Pair\_coeff command before simulation box is defined

The pair\_coeff command cannot be used before a read\_data, read\_restart, or create\_box command.

## Pair\_modify command before pair\_style is defined

Self-explanatory.

## Pair\_modify special setting for pair hybrid incompatible with global special\_bonds setting

Cannot override a setting of 0.0 or 1.0 or change a setting between 0.0 and 1.0.

## Pair\_write command before pair\_style is defined

Self-explanatory.

## Particle on or inside fix wall surface

Particles must be “exterior” to the wall in order for energy/force to be calculated.

## Particle outside surface of region used in fix wall/region

Particles must be inside the region for energy/force to be calculated. A particle outside the region generates an error.

## Per-atom compute in equal-style variable formula

Equal-style variables cannot use per-atom quantities.

## Per-atom energy was not tallied on needed timestep

You are using a thermo keyword that requires potentials to have tallied energy, but they did not on this timestep. See the variable page for ideas on how to make this work.

## Per-atom fix in equal-style variable formula

Equal-style variables cannot use per-atom quantities.

## Per-atom virial was not tallied on needed timestep

You are using a thermo keyword that requires potentials to have tallied the virial, but they did not on this timestep. See the variable page for ideas on how to make this work.

## Per-processor system is too big

The number of owned atoms plus ghost atoms on a single processor must fit in 32-bit integer.

## Potential energy ID for fix neb does not exist

Self-explanatory.

## Potential energy ID for fix nvt/nph/npt does not exist

A compute for potential energy must be defined.

## Potential file has duplicate entry

The potential file has more than one entry for the same element.

## Potential file is missing an entry

The potential file does not have a needed entry.

## Power by 0 in variable formula

Self-explanatory.

## Pressure ID for fix box/relax does not exist

The compute ID needed to compute pressure for the fix does not exist.

## Pressure ID for fix modify does not exist

Self-explanatory.

## Pressure ID for fix npt/nph does not exist

Self-explanatory.

## Pressure ID for fix press/berendsen does not exist

The compute ID needed to compute pressure for the fix does not exist.

## Pressure ID for fix rigid npt/nph does not exist

Self-explanatory.

## Pressure ID for thermo does not exist

The compute ID needed to compute pressure for thermodynamics does not exist.

## Pressure control can not be used with fix nvt

Self-explanatory.

## Pressure control can not be used with fix nvt/asphere

Self-explanatory.

## Pressure control can not be used with fix nvt/body

Self-explanatory.

## Pressure control can not be used with fix nvt/sllod

Self-explanatory.

## Pressure control can not be used with fix nvt/sphere

Self-explanatory.

## Pressure control must be used with fix nph

Self-explanatory.

## Pressure control must be used with fix nph/asphere

Self-explanatory.

## Pressure control must be used with fix nph/body

Self-explanatory.

## Pressure control must be used with fix nph/small

Self-explanatory.

## Pressure control must be used with fix nph/sphere

Self-explanatory.

## Pressure control must be used with fix nphug

A pressure control keyword (iso, aniso, tri, x, y, or z) must be provided.

## Pressure control must be used with fix npt

Self-explanatory.

## Pressure control must be used with fix npt/asphere

Self-explanatory.

## Pressure control must be used with fix npt/body

Self-explanatory.

## Pressure control must be used with fix npt/sphere

Self-explanatory.

## Processor count in z must be 1 for 2d simulation

Self-explanatory.

## Processor partitions do not match number of allocated processors

The total number of processors in all partitions must match the number of processors LAMMPS is running on.

## Processors command after simulation box is defined

The processors command cannot be used after a read\_data, read\_restart, or create\_box command.

## Processors custom grid file is inconsistent

The vales in the custom file are not consistent with the number of processors you are running on or the Px,Py,Pz settings of the processors command. Or there was not a setting for every processor.

## Processors grid numa and map style are incompatible

Using numa for gstyle in the processors command requires using cart for the map option.

## Processors part option and grid style are incompatible

Cannot use gstyle numa or custom with the part option.

## Processors twogrid requires proc count be a multiple of core count

Self-explanatory.

## Pstart and Pstop must have the same value

Self-explanatory.

## Python function evaluation failed

The Python function did not run successfully and/or did not return a value (if it is supposed to return a value). This is probably due to some error condition in the function.

## Python function is not callable

The provided Python code was run successfully, but it not define a callable function with the required name.

## Python invoke of undefined function

Cannot invoke a function that has not been previously defined.

## Python variable does not match Python function

This matching is defined by the python-style variable and the python command.

## Python variable has no function

No python command was used to define the function associated with the python-style variable.

## QEQ with ‘newton pair off’ not supported

See the newton command. This is a restriction to use the QEQ fixes.

## R0 < 0 for fix spring command

Equilibrium spring length is invalid.

## RATTLE coordinate constraints are not satisfied up to desired tolerance

Self-explanatory.

## RATTLE determinant = 0.0

The determinant of the matrix being solved for a single cluster specified by the fix rattle command is numerically invalid.

## RATTLE failed

Certain constraints were not satisfied.

## RATTLE velocity constraints are not satisfied up to desired tolerance

Self-explanatory.

## Read data add offset is too big

It cannot be larger than the size of atom IDs, e.g. the maximum 32-bit integer.

## Read dump of atom property that is not allocated

Self-explanatory.

## Read rerun dump file timestep > specified stop

Self-explanatory.

## Read restart MPI-IO input not allowed with % in filename

This is because a % signifies one file per processor and MPI-IO creates one large file for all processors.

## Read\_data shrink wrap did not assign all atoms correctly

This is typically because the box-size specified in the data file is large compared to the actual extent of atoms in a shrink-wrapped dimension. When LAMMPS shrink-wraps the box atoms will be lost if the processor they are re-assigned to is too far away. Choose a box size closer to the actual extent of the atoms.

## Read\_dump command before simulation box is defined

The read\_dump command cannot be used before a read\_data, read\_restart, or create\_box command.

## Read\_dump field not found in dump file

Self-explanatory.

## Read\_dump triclinic status does not match simulation

Both the dump snapshot and the current LAMMPS simulation must be using either an orthogonal or triclinic box.

## Read\_dump xyz fields do not have consistent scaling/wrapping

Self-explanatory.

## Reax\_defs.h setting for NATDEF is too small

Edit the setting in the ReaxFF library and re-compile the library and re-build LAMMPS.

## Reax\_defs.h setting for NNEIGHMAXDEF is too small

Edit the setting in the ReaxFF library and re-compile the library and re-build LAMMPS.

## Receiving partition in processors part command is already a receiver

Cannot specify a partition to be a receiver twice.

## Region ID for compute chunk/atom does not exist

Self-explanatory.

## Region ID for compute reduce/region does not exist

Self-explanatory.

## Region ID for compute temp/region does not exist

Self-explanatory.

## Region ID for dump custom does not exist

Self-explanatory.

## Region ID for fix addforce does not exist

Self-explanatory.

## Region ID for fix atom/swap does not exist

Self-explanatory.

## Region ID for fix ave/spatial does not exist

Self-explanatory.

## Region ID for fix aveforce does not exist

Self-explanatory.

## Region ID for fix deposit does not exist

Self-explanatory.

## Region ID for fix efield does not exist

Self-explanatory.

## Region ID for fix evaporate does not exist

Self-explanatory.

## Region ID for fix gcmc does not exist

Self-explanatory.

## Region ID for fix heat does not exist

Self-explanatory.

## Region ID for fix setforce does not exist

Self-explanatory.

## Region ID for fix wall/region does not exist

Self-explanatory.

## Region ID for group dynamic does not exist

Self-explanatory.

## Region ID in variable formula does not exist

Self-explanatory.

## Region cannot have 0 length rotation vector

Self-explanatory.

## Region for fix oneway does not exist

Self-explanatory.

## Region intersect region ID does not exist

Self-explanatory.

## Region union or intersect cannot be dynamic

The sub-regions can be dynamic, but not the combined region.

## Region union region ID does not exist

One or more of the region IDs specified by the region union command does not exist.

## Replacing a fix, but new style != old style

A fix ID can be used a second time, but only if the style matches the previous fix. In this case it is assumed you want to reset a fix’s parameters. This error may mean you are mistakenly re-using a fix ID when you do not intend to.

## Replicate command before simulation box is defined

The replicate command cannot be used before a read\_data, read\_restart, or create\_box command.

## Replicate did not assign all atoms correctly

Atoms replicated by the replicate command were not assigned correctly to processors. This is likely due to some atom coordinates being outside a non-periodic simulation box.

## Replicated system atom IDs are too big

See the setting for tagint in the src/lmptype.h file.

## Replicated system is too big

See the setting for bigint in the src/lmptype.h file.

## Required border comm not yet implemented with Kokkos

There are various limitations in the communication options supported by Kokkos.

## Rerun command before simulation box is defined

The rerun command cannot be used before a read\_data, read\_restart, or create\_box command.

## Rerun dump file does not contain requested snapshot

Self-explanatory.

## Resetting timestep size is not allowed with fix move

This is because fix move is moving atoms based on elapsed time.

## Respa inner cutoffs are invalid

The first cutoff must be <= the second cutoff.

## Respa levels must be >= 1

Self-explanatory.

## Respa middle cutoffs are invalid

The first cutoff must be <= the second cutoff.

## Restart file MPI-IO output not allowed with % in filename

This is because a % signifies one file per processor and MPI-IO creates one large file for all processors.

## Restart file byte ordering is not recognized

The file does not appear to be a LAMMPS restart file since it does not contain a recognized byte-ordering flag at the beginning.

## Restart file byte ordering is swapped

The file was written on a machine with different byte-ordering than the machine you are reading it on. Convert it to a text data file instead, on the machine you wrote it on.

## Restart file incompatible with current version

This is probably because you are trying to read a file created with a version of LAMMPS that is too old compared to the current version. Use your older version of LAMMPS and convert the restart file to a data file.

## Restart file is a MPI-IO file

The file is inconsistent with the filename you specified for it.

## Restart file is a multi-proc file

The file is inconsistent with the filename you specified for it.

## Restart file is not a MPI-IO file

The file is inconsistent with the filename you specified for it.

## Restart file is not a multi-proc file

The file is inconsistent with the filename you specified for it.

## Restart variable returned a bad timestep

The variable must return a timestep greater than the current timestep.

## Restrain atoms %d %d %d %d missing on proc %d at step %ld

The 4 atoms in a restrain dihedral specified by the fix restrain command are not all accessible to a processor. This probably means an atom has moved too far.

## Restrain atoms %d %d %d missing on proc %d at step %ld

The three atoms in a restrain angle specified by the fix restrain command are not all accessible to a processor. This probably means an atom has moved too far.

## Restrain atoms %d %d missing on proc %d at step %ld

The two atoms in a restrain bond specified by the fix restrain command are not all accessible to a processor. This probably means an atom has moved too far.

## Reuse of compute ID

A compute ID cannot be used twice.

## Reuse of dump ID

A dump ID cannot be used twice.

## Reuse of molecule template ID

The template IDs must be unique.

## Reuse of region ID

A region ID cannot be used twice.

## Rigid body atoms %d %d missing on proc %d at step %ld

This means that an atom cannot find the atom that owns the rigid body it is part of, or vice versa. The solution is to use the communicate cutoff command to ensure ghost atoms are acquired from far enough away to encompass the max distance printed when the fix rigid/small command was invoked.

## Rigid body has degenerate moment of inertia

Fix poems will only work with bodies (collections of atoms) that have non-zero principal moments of inertia. This means they must be 3 or more non-collinear atoms, even with joint atoms removed.

## Rigid fix must come before NPT/NPH fix

NPT/NPH fix must be defined in input script after all rigid fixes, else the rigid fix contribution to the pressure virial is incorrect.

## Rmask function in equal-style variable formula

Rmask is per-atom operation.

## Run command before simulation box is defined

The run command cannot be used before a read\_data, read\_restart, or create\_box command.

## Run command start value is after start of run

Self-explanatory.

## Run command stop value is before end of run

Self-explanatory.

## Run\_style command before simulation box is defined

The run\_style command cannot be used before a read\_data, read\_restart, or create\_box command.

## SRD bin size for fix srd differs from user request

Fix SRD had to adjust the bin size to fit the simulation box. See the cubic keyword if you want this message to be an error vs warning.

## SRD bins for fix srd are not cubic enough

The bin shape is not within tolerance of cubic. See the cubic keyword if you want this message to be an error vs warning.

## SRD particle %d started inside big particle %d on step %ld bounce %d

See the inside keyword if you want this message to be an error vs warning.

## SRD particle %d started inside wall %d on step %ld bounce %d

See the inside keyword if you want this message to be an error vs warning.

## Same dimension twice in fix ave/spatial

Self-explanatory.

## Sending partition in processors part command is already a sender

Cannot specify a partition to be a sender twice.

## Set command before simulation box is defined

The set command cannot be used before a read\_data, read\_restart, or create\_box command.

## Set command floating point vector does not exist

Self-explanatory.

## Set command integer vector does not exist

Self-explanatory.

## Set command with no atoms existing

No atoms are yet defined so the set command cannot be used.

## Set region ID does not exist

Region ID specified in set command does not exist.

## Shake angles have different bond types

All 3-atom angle-constrained SHAKE clusters specified by the fix shake command that are the same angle type, must also have the same bond types for the two bonds in the angle.

## Shake atoms %d %d %d %d missing on proc %d at step %ld

The 4 atoms in a single shake cluster specified by the fix shake command are not all accessible to a processor. This probably means an atom has moved too far.

## Shake atoms %d %d %d missing on proc %d at step %ld

The three atoms in a single shake cluster specified by the fix shake command are not all accessible to a processor. This probably means an atom has moved too far.

## Shake atoms %d %d missing on proc %d at step %ld

The two atoms in a single shake cluster specified by the fix shake command are not all accessible to a processor. This probably means an atom has moved too far.

## Shake cluster of more than 4 atoms

A single cluster specified by the fix shake command can have no more than 4 atoms.

## Shake clusters are connected

A single cluster specified by the fix shake command must have a single central atom with up to 3 other atoms bonded to it.

## Shake determinant = 0.0

The determinant of the matrix being solved for a single cluster specified by the fix shake command is numerically invalid.

## Shake fix must come before NPT/NPH fix

NPT fix must be defined in input script after SHAKE fix, else the SHAKE fix contribution to the pressure virial is incorrect.

## Shear history overflow, boost neigh\_modify one

There are too many neighbors of a single atom. Use the neigh\_modify command to increase the max number of neighbors allowed for one atom. You may also want to boost the page size.

## Small to big integers are not sized correctly

This error occurs when the sizes of smallint, imageint, tagint, bigint, as defined in src/lmptype.h are not what is expected. Contact the developers if this occurs.

## Smallint setting in lmptype.h is invalid

It has to be the size of an integer.

## Smallint setting in lmptype.h is not compatible

Smallint stored in restart file is not consistent with LAMMPS version you are running.

## Special list size exceeded in fix bond/create

See the “read\_data extra/special/per/atom” command (or the “create\_box extra/special/per/atom” command) for info on how to leave space in the special bonds list to allow for additional bonds to be formed.

## Species XXX is not supported by this KIM Simulator Model

The kim\_style define command was referencing a species that is not present in the requested KIM Simulator Model.

## Specified processors != physical processors

The 3d grid of processors defined by the processors command does not match the number of processors LAMMPS is being run on.

## Specified target stress must be uniaxial or hydrostatic

Self-explanatory.

## Sqrt of negative value in variable formula

Self-explanatory.

## Subsequent read data induced too many angles per atom

See the extra/angle/per/atom keyword for the create\_box or the read\_data command to set this limit larger

## Subsequent read data induced too many bonds per atom

See the extra/bond/per/atom keyword for the create\_box or the read\_data command to set this limit larger

## Subsequent read data induced too many dihedrals per atom

See the extra/dihedral/per/atom keyword for the create\_box or the read\_data command to set this limit larger

## Subsequent read data induced too many impropers per atom

See the extra/improper/per/atom keyword for the create\_box or the read\_data command to set this limit larger

## Substitution for illegal variable

Input script line contained a variable that could not be substituted for.

## Support for writing images in JPEG format not included

LAMMPS was not built with the -DLAMMPS\_JPEG switch in the Makefile.

## Support for writing images in PNG format not included

LAMMPS was not built with the -DLAMMPS\_PNG switch in the Makefile.

## Support for writing movies not included

LAMMPS was not built with the -DLAMMPS\_FFMPEG switch in the Makefile

## System in data file is too big

See the setting for bigint in the src/lmptype.h file.

## System is not charge neutral, net charge = %g

The total charge on all atoms on the system is not 0.0. For some KSpace solvers this is an error.

## TAD nsteps must be multiple of t\_event

Self-explanatory.

## TIP4P hydrogen has incorrect atom type

The TIP4P pairwise computation found an H atom whose type does not agree with the specified H type.

## TIP4P hydrogen is missing

The TIP4P pairwise computation failed to find the correct H atom within a water molecule.

## TMD target file did not list all group atoms

The target file for the fix tmd command did not list all atoms in the fix group.

## Tad command before simulation box is defined

Self-explanatory.

## Tagint setting in lmptype.h is invalid

Tagint must be as large or larger than smallint.

## Tagint setting in lmptype.h is not compatible

Format of tagint stored in restart file is not consistent with LAMMPS version you are running. See the settings in src/lmptype.h

## Target pressure for fix rigid/nph cannot be < 0.0

Self-explanatory.

## Target pressure for fix rigid/npt/small cannot be < 0.0

Self-explanatory.

## Target temperature for fix nvt/npt/nph cannot be 0.0

Self-explanatory.

## Target temperature for fix rigid/npt cannot be 0.0

Self-explanatory.

## Target temperature for fix rigid/npt/small cannot be 0.0

Self-explanatory.

## Target temperature for fix rigid/nvt cannot be 0.0

Self-explanatory.

## Target temperature for fix rigid/nvt/small cannot be 0.0

Self-explanatory.

## Temper command before simulation box is defined

The temper command cannot be used before a read\_data, read\_restart, or create\_box command.

## Temperature ID for fix bond/swap does not exist

Self-explanatory.

## Temperature ID for fix box/relax does not exist

Self-explanatory.

## Temperature ID for fix nvt/npt does not exist

Self-explanatory.

## Temperature ID for fix press/berendsen does not exist

Self-explanatory.

## Temperature ID for fix rigid nvt/npt/nph does not exist

Self-explanatory.

## Temperature ID for fix temp/berendsen does not exist

Self-explanatory.

## Temperature ID for fix temp/csld does not exist

Self-explanatory.

## Temperature ID for fix temp/csvr does not exist

Self-explanatory.

## Temperature ID for fix temp/rescale does not exist

Self-explanatory.

## Temperature compute degrees of freedom < 0

This should not happen if you are calculating the temperature on a valid set of atoms.

## Temperature control can not be used with fix nph

Self-explanatory.

## Temperature control can not be used with fix nph/asphere

Self-explanatory.

## Temperature control can not be used with fix nph/body

Self-explanatory.

## Temperature control can not be used with fix nph/sphere

Self-explanatory.

## Temperature control must be used with fix nphug

The temp keyword must be provided.

## Temperature control must be used with fix npt

Self-explanatory.

## Temperature control must be used with fix npt/asphere

Self-explanatory.

## Temperature control must be used with fix npt/body

Self-explanatory.

## Temperature control must be used with fix npt/sphere

Self-explanatory.

## Temperature control must be used with fix nvt

Self-explanatory.

## Temperature control must be used with fix nvt/asphere

Self-explanatory.

## Temperature control must be used with fix nvt/body

Self-explanatory.

## Temperature control must be used with fix nvt/sllod

Self-explanatory.

## Temperature control must be used with fix nvt/sphere

Self-explanatory.

## Temperature control must not be used with fix nph/small

Self-explanatory.

## Temperature for fix nvt/sllod does not have a bias

The specified compute must compute temperature with a bias.

## Tempering could not find thermo\_pe compute

This compute is created by the thermo command. It must have been explicitly deleted by a uncompute command.

## Tempering fix ID is not defined

The fix ID specified by the temper command does not exist.

## Tempering temperature fix is not valid

The fix specified by the temper command is not one that controls temperature (nvt or langevin).

## Test\_descriptor\_string already allocated

This is an internal error. Contact the developers.

## The package gpu command is required for gpu styles

Self-explanatory.

## Thermo and fix not computed at compatible times

Fixes generate values on specific timesteps. The thermo output does not match these timesteps.

## Thermo compute array is accessed out-of-range

Self-explanatory.

## Thermo compute does not compute array

Self-explanatory.

## Thermo compute does not compute scalar

Self-explanatory.

## Thermo compute does not compute vector

Self-explanatory.

## Thermo compute vector is accessed out-of-range

Self-explanatory.

## Thermo custom variable cannot be indexed

Self-explanatory.

## Thermo custom variable is not equal-style variable

Only equal-style variables can be output with thermodynamics, not atom-style variables.

## Thermo every variable returned a bad timestep

The variable must return a timestep greater than the current timestep.

## Thermo fix array is accessed out-of-range

Self-explanatory.

## Thermo fix does not compute array

Self-explanatory.

## Thermo fix does not compute scalar

Self-explanatory.

## Thermo fix does not compute vector

Self-explanatory.

## Thermo fix vector is accessed out-of-range

Self-explanatory.

## Thermo keyword in variable requires thermo to use/init pe

You are using a thermo keyword in a variable that requires potential energy to be calculated, but your thermo output does not use it. Add it to your thermo output.

## Thermo keyword in variable requires thermo to use/init press

You are using a thermo keyword in a variable that requires pressure to be calculated, but your thermo output does not use it. Add it to your thermo output.

## Thermo keyword in variable requires thermo to use/init temp

You are using a thermo keyword in a variable that requires temperature to be calculated, but your thermo output does not use it. Add it to your thermo output.

## Thermo style does not use press

Cannot use thermo\_modify to set this parameter since the thermo\_style is not computing this quantity.

## Thermo style does not use temp

Cannot use thermo\_modify to set this parameter since the thermo\_style is not computing this quantity.

## Thermo\_modify every variable returned a bad timestep

The returned timestep is less than or equal to the current timestep.

## Thermo\_modify int format does not contain d character

Self-explanatory.

## Thermo\_modify pressure ID does not compute pressure

The specified compute ID does not compute pressure.

## Thermo\_modify temperature ID does not compute temperature

The specified compute ID does not compute temperature.

## Thermo\_style command before simulation box is defined

The thermo\_style command cannot be used before a read\_data, read\_restart, or create\_box command.

## This variable thermo keyword cannot be used between runs

Keywords that refer to time (such as cpu, elapsed) do not make sense in between runs.

## Threshold for an atom property that is not allocated

A dump threshold has been requested on a quantity that is not defined by the atom style used in this simulation.

## Timestep must be >= 0

Specified timestep is invalid.

## Too big a problem to use velocity create loop all

The system size must fit in a 32-bit integer to use this option.

## Too big a timestep for dump dcd

The timestep must fit in a 32-bit integer to use this dump style.

## Too big a timestep for dump xtc

The timestep must fit in a 32-bit integer to use this dump style.

## Too few bits for lookup table

Table size specified via pair\_modify command does not work with your machine’s floating point representation.

## Too few lines in %s section of data file

Self-explanatory.

## Too few values in body lines in data file

Self-explanatory.

## Too few values in body section of molecule file

Self-explanatory.

## Too many -pk arguments in command line

The string formed by concatenating the arguments is too long. Use a package command in the input script instead.

## Too many MSM grid levels

The max number of MSM grid levels is hardwired to 10.

## Too many args in variable function

More args are used than any variable function allows.

## Too many atom pairs for pair bop

The number of atomic pairs exceeds the expected number. Check your atomic structure to ensure that it is realistic.

## Too many atom sorting bins

This is likely due to an immense simulation box that has blown up to a large size.

## Too many atom triplets for pair bop

The number of three atom groups for angle determinations exceeds the expected number. Check your atomic structure to ensure that it is realistic.

## Too many atoms for dump dcd

The system size must fit in a 32-bit integer to use this dump style.

## Too many atoms for dump xtc

The system size must fit in a 32-bit integer to use this dump style.

## Too many elements extracted from MEAM library.

Increase ‘maxelt’ in meam.h and recompile.

## Too many exponent bits for lookup table

Table size specified via pair\_modify command does not work with your machine’s floating point representation.

## Too many groups

The maximum number of atom groups (including the “all” group) is given by MAX\_GROUP in group.cpp and is 32.

## Too many iterations

You must use a number of iterations that fit in a 32-bit integer for minimization.

## Too many lines in one body in data file - boost MAXBODY

MAXBODY is a setting at the top of the src/read\_data.cpp file. Set it larger and re-compile the code.

## Too many local+ghost atoms for neighbor list

The number of nlocal + nghost atoms on a processor is limited by the size of a 32-bit integer with 2 bits removed for masking 1-2, 1-3, 1-4 neighbors.

## Too many mantissa bits for lookup table

Table size specified via pair\_modify command does not work with your machine’s floating point representation.

## Too many masses for fix shake

The fix shake command cannot list more masses than there are atom types.

## Too many molecules for fix poems

The limit is 2^31 = ~2 billion molecules.

## Too many molecules for fix rigid

The limit is 2^31 = ~2 billion molecules.

## Too many neighbor bins

This is likely due to an immense simulation box that has blown up to a large size.

## Too many timesteps

The cumulative timesteps must fit in a 64-bit integer.

## Too many timesteps for NEB

You must use a number of timesteps that fit in a 32-bit integer for NEB.

## Too many total atoms

See the setting for bigint in the src/lmptype.h file.

## Too many total bits for bitmapped lookup table

Table size specified via pair\_modify command is too large. Note that a value of N generates a 2^N size table.

## Too many values in body lines in data file

Self-explanatory.

## Too many values in body section of molecule file

Self-explanatory.

## Too much buffered per-proc info for dump

The size of the buffered string must fit in a 32-bit integer for a dump.

## Too much per-proc info for dump

Number of local atoms times number of columns must fit in a 32-bit integer for dump.

## Topology type exceeds system topology type

The number of bond, angle, etc types exceeds the system setting. See the create\_box or read\_data command for how to specify these values.

## Tree structure in joint connections

Fix poems cannot (yet) work with coupled bodies whose joints connect the bodies in a tree structure.

## Tried to convert a double to int, but input\_double > INT\_MAX

Self-explanatory.

## Trying to build an occasional neighbor list before initialization completed

This is not allowed. Source code caller needs to be modified.

## Two fix ave commands using same compute chunk/atom command in incompatible ways

They are both attempting to “lock” the chunk/atom command so that the chunk assignments persist for some number of timesteps, but are doing it in different ways.

## Two groups cannot be the same in fix spring couple

Self-explanatory.

## The %s type label %s is already in use for type %s

For a given type-kind (atom types, bond types, etc.), a given type label can be assigned to only one numeric type.

## Type label string %s for %s type %s is invalid

See the labelmap command documentation for valid type labels.

## Unable to initialize accelerator for use

There was a problem initializing an accelerator for the gpu package

## Unbalanced quotes in input line

No matching end double quote was found following a leading double quote.

## Unexpected end of -reorder file

Self-explanatory.

## Unexpected empty line in Angle Coeffs section

Read a blank line where there should be coefficient data.

## Unexpected empty line in Bond Coeffs section

Read a blank line where there should be coefficient data.

## Unexpected empty line in Dihedral Coeffs section

Read a blank line where there should be coefficient data.

## Unexpected empty line in Improper Coeffs section

Read a blank line where there should be coefficient data.

## Unexpected empty line in Pair Coeffs section

Read a blank line where there should be coefficient data.

## Unexpected end of custom file

Self-explanatory.

## Unexpected end of data file

LAMMPS hit the end of the data file while attempting to read a section. Something is wrong with the format of the data file.

## Unexpected end of dump file

A read operation from the file failed.

## Unexpected end of fix rigid file

A read operation from the file failed.

## Unexpected end of fix rigid/small file

A read operation from the file failed.

## Unexpected end of molecule file

Self-explanatory.

## Unexpected end of neb file

A read operation from the file failed.

## Units command after simulation box is defined

The units command cannot be used after a read\_data, read\_restart, or create\_box command.

## Universe/uloop variable count < # of partitions

A universe or uloop style variable must specify a number of values >= to the number of processor partitions.

## Unrecognized angle style

The choice of angle style is unknown.

## Unrecognized atom style

The choice of atom style is unknown.

## Unrecognized body style

The choice of body style is unknown.

## Unrecognized bond style

The choice of bond style is unknown.

## Unknown category for info is\_active()

Self-explanatory.

## Unknown category for info is\_available()

Self-explanatory.

## Unknown category for info is\_defined()

Self-explanatory.

## Unrecognized command: %s

The command is not known to LAMMPS. Check the input script.

## Unrecognized compute style

The choice of compute style is unknown.

## Unrecognized dihedral style

The choice of dihedral style is unknown.

## Unrecognized dump reader style

The choice of dump reader style via the format keyword is unknown.

## Unrecognized dump style

The choice of dump style is unknown.

## Unknown error in GPU library

Self-explanatory.

## Unrecognized fix style

The choice of fix style is unknown.

## Unknown identifier in data file: %s

A section of the data file cannot be read by LAMMPS.

## Unrecognized improper style

The choice of improper style is unknown.

## Unknown keyword in thermo\_style custom command

One or more specified keywords are not recognized.

## Unrecognized kspace style

The choice of kspace style is unknown.

## Unknown name for info newton category

Self-explanatory.

## Unknown name for info package category

Self-explanatory.

## Unknown name for info pair category

Self-explanatory.

## Unrecognized pair style

The choice of pair style is unknown.

## Unknown pair\_modify hybrid sub-style

The choice of sub-style is unknown.

## Unrecognized region style

The choice of region style is unknown.

## Unknown section in molecule file

Self-explanatory.

## Unknown table style in angle style table

Self-explanatory.

## Unknown table style in bond style table

Self-explanatory.

## Unknown table style in pair\_style command

Style of table is invalid for use with pair\_style table command.

## Unknown unit\_style

Self-explanatory. Check the input script or data file.

## Unrecognized lattice type in MEAM library file

The lattice type in an entry of the MEAM library file is not valid.

## Unrecognized lattice type in MEAM parameter file

The lattice type in an entry of the MEAM parameter file is not valid.

## Unrecognized pair style in compute pair command

Self-explanatory.

## Unsupported mixing rule in kspace\_style ewald/disp

Only geometric mixing is supported.

## Unsupported order in kspace\_style ewald/disp

Only 1/r^6 dispersion or dipole terms are supported.

## Unsupported order in kspace\_style pppm/disp, pair\_style %s

Only pair styles with 1/r and 1/r^6 dependence are currently supported.

## Unsupported parameter in MEAM library file

Self-explanatory.

## Use cutoff keyword to set cutoff in single mode

Mode is single so cutoff/multi keyword cannot be used.

## Use cutoff/multi keyword to set cutoff in multi mode

Mode is multi so cutoff keyword cannot be used.

## Using fix nvt/sllod with inconsistent fix deform remap option

Fix nvt/sllod requires that deforming atoms have a velocity profile provided by “remap v” as a fix deform option.

## Using fix nvt/sllod with no fix deform defined

Self-explanatory.

## Using fix srd with inconsistent fix deform remap option

When shearing the box in an SRD simulation, the remap v option for fix deform needs to be used.

## Using pair lubricate with inconsistent fix deform remap option

Must use remap v option with fix deform with this pair style.

## Using pair lubricate/poly with inconsistent fix deform remap option

If fix deform is used, the remap v option is required.

## Using suffix gpu without GPU package installed

Self-explanatory.

## Using suffix intel without INTEL package installed

Self-explanatory.

## Using suffix kk without KOKKOS package enabled

Self-explanatory.

## Using suffix omp without OPENMP package installed

Self-explanatory.

## Using update dipole flag requires atom attribute mu

Self-explanatory.

## Using update dipole flag requires atom style sphere

Self-explanatory.

## Variable ID in variable formula does not exist

Self-explanatory.

## Variable atom ID is too large

Specified ID is larger than the maximum allowed atom ID.

## Variable evaluation before simulation box is defined

Cannot evaluate a compute or fix or atom-based value in a variable before the simulation has been setup.

## Variable evaluation in fix wall gave bad value

The returned value for epsilon or sigma < 0.0.

## Variable evaluation in region gave bad value

Variable returned a radius < 0.0.

## Variable for compute ti is invalid style

Self-explanatory.

## Variable for create\_atoms is invalid style

The variables must be equal-style variables.

## Variable for displace\_atoms is invalid style

It must be an equal-style or atom-style variable.

## Variable for dump every is invalid style

Only equal-style variables can be used.

## Variable for dump image center is invalid style

Must be an equal-style variable.

## Variable for dump image phi is invalid style

Must be an equal-style variable.

## Variable for dump image theta is invalid style

Must be an equal-style variable.

## Variable for dump image zoom is invalid style

Must be an equal-style variable.

## Variable for fix adapt is invalid style

Only equal-style variables can be used.

## Variable for fix addforce is invalid style

Self-explanatory.

## Variable for fix aveforce is invalid style

Only equal-style variables can be used.

## Variable for fix deform is invalid style

The variable must be an equal-style variable.

## Variable for fix efield is invalid style

The variable must be an equal- or atom-style variable.

## Variable for fix gravity is invalid style

Only equal-style variables can be used.

## Variable for fix heat is invalid style

Only equal-style or atom-style variables can be used.

## Variable for fix indent is invalid style

Only equal-style variables can be used.

## Variable for fix indent is not equal style

Only equal-style variables can be used.

## Variable for fix langevin is invalid style

It must be an equal-style variable.

## Variable for fix move is invalid style

Only equal-style variables can be used.

## Variable for fix setforce is invalid style

Only equal-style variables can be used.

## Variable for fix temp/berendsen is invalid style

Only equal-style variables can be used.

## Variable for fix temp/csld is invalid style

Only equal-style variables can be used.

## Variable for fix temp/csvr is invalid style

Only equal-style variables can be used.

## Variable for fix temp/rescale is invalid style

Only equal-style variables can be used.

## Variable for fix wall is invalid style

Only equal-style variables can be used.

## Variable for fix wall/reflect is invalid style

Only equal-style variables can be used.

## Variable for fix wall/srd is invalid style

Only equal-style variables can be used.

## Variable for group dynamic is invalid style

The variable must be an atom-style variable.

## Variable for group is invalid style

Only atom-style variables can be used.

## Variable for region cylinder is invalid style

Only equal-style variables are allowed.

## Variable for region is invalid style

Only equal-style variables can be used.

## Variable for region is not equal style

Self-explanatory.

## Variable for region sphere is invalid style

Only equal-style variables are allowed.

## Variable for restart is invalid style

Only equal-style variables can be used.

## Variable for set command is invalid style

Only atom-style variables can be used.

## Variable for thermo every is invalid style

Only equal-style variables can be used.

## Variable for velocity set is invalid style

Only atom-style variables can be used.

## Variable for voronoi radius is not atom style

Self-explanatory.

## Variable formula compute array is accessed out-of-range

Self-explanatory.

## Variable formula compute vector is accessed out-of-range

Self-explanatory.

## Variable formula fix array is accessed out-of-range

Self-explanatory.

## Variable formula fix vector is accessed out-of-range

Self-explanatory.

## Variable has circular dependency

A circular dependency is when variable “a” in used by variable “b” and variable “b” is also used by variable “a”. Circular dependencies with longer chains of dependence are also not allowed.

## Variable name between brackets must be alphanumeric or underscore characters

Self-explanatory.

## Variable name for compute chunk/atom does not exist

Self-explanatory.

## Variable name for compute reduce does not exist

Self-explanatory.

## Variable name for compute ti does not exist

Self-explanatory.

## Variable name for create\_atoms does not exist

Self-explanatory.

## Variable name for displace\_atoms does not exist

Self-explanatory.

## Variable name for dump every does not exist

Self-explanatory.

## Variable name for dump image center does not exist

Self-explanatory.

## Variable name for dump image phi does not exist

Self-explanatory.

## Variable name for dump image theta does not exist

Self-explanatory.

## Variable name for dump image zoom does not exist

Self-explanatory.

## Variable name for fix adapt does not exist

Self-explanatory.

## Variable name for fix addforce does not exist

Self-explanatory.

## Variable name for fix ave/atom does not exist

Self-explanatory.

## Variable name for fix ave/chunk does not exist

Self-explanatory.

## Variable name for fix ave/correlate does not exist

Self-explanatory.

## Variable name for fix ave/histo does not exist

Self-explanatory.

## Variable name for fix ave/spatial does not exist

Self-explanatory.

## Variable name for fix ave/time does not exist

Self-explanatory.

## Variable name for fix aveforce does not exist

Self-explanatory.

## Variable name for fix deform does not exist

Self-explanatory.

## Variable name for fix efield does not exist

Self-explanatory.

## Variable name for fix gravity does not exist

Self-explanatory.

## Variable name for fix heat does not exist

Self-explanatory.

## Variable name for fix indent does not exist

Self-explanatory.

## Variable name for fix langevin does not exist

Self-explanatory.

## Variable name for fix move does not exist

Self-explanatory.

## Variable name for fix setforce does not exist

Self-explanatory.

## Variable name for fix store/state does not exist

Self-explanatory.

## Variable name for fix temp/berendsen does not exist

Self-explanatory.

## Variable name for fix temp/csld does not exist

Self-explanatory.

## Variable name for fix temp/csvr does not exist

Self-explanatory.

## Variable name for fix temp/rescale does not exist

Self-explanatory.

## Variable name for fix vector does not exist

Self-explanatory.

## Variable name for fix wall does not exist

Self-explanatory.

## Variable name for fix wall/reflect does not exist

Self-explanatory.

## Variable name for fix wall/srd does not exist

Self-explanatory.

## Variable name for group does not exist

Self-explanatory.

## Variable name for group dynamic does not exist

Self-explanatory.

## Variable name for region cylinder does not exist

Self-explanatory.

## Variable name for region does not exist

Self-explanatory.

## Variable name for region sphere does not exist

Self-explanatory.

## Variable name for restart does not exist

Self-explanatory.

## Variable name for set command does not exist

Self-explanatory.

## Variable name for thermo every does not exist

Self-explanatory.

## Variable name for velocity set does not exist

Self-explanatory.

## Variable name for voronoi radius does not exist

Self-explanatory.

## Variable name must be alphanumeric or underscore characters

Self-explanatory.

## Variable uses atom property that is not allocated

Self-explanatory.

## Velocity command before simulation box is defined

The velocity command cannot be used before a read\_data, read\_restart, or create\_box command.

## Velocity command with no atoms existing

A velocity command has been used, but no atoms yet exist.

## Velocity ramp in z for a 2d problem

Self-explanatory.

## Velocity rigid used with non-rigid fix-ID

Self-explanatory.

## Velocity temperature ID does calculate a velocity bias

The specified compute must compute a bias for temperature.

## Velocity temperature ID does not compute temperature

The compute ID given to the velocity command must compute temperature.

## Verlet/split can only currently be used with comm\_style brick

This is a current restriction in LAMMPS.

## Verlet/split does not yet support TIP4P

This is a current limitation.

## Verlet/split requires 2 partitions

See the -partition command-line switch.

## Verlet/split requires Rspace partition layout be multiple of Kspace partition layout in each dim

This is controlled by the processors command.

## Verlet/split requires Rspace partition size be multiple of Kspace partition size

This is so there is an equal number of Rspace processors for every Kspace processor.

## Virial was not tallied on needed timestep

You are using a thermo keyword that requires potentials to have tallied the virial, but they did not on this timestep. See the variable page for ideas on how to make this work.

## Voro++ error: narea and neigh have a different size

This error is returned by the Voro++ library.

## Wall defined twice in fix wall command

Self-explanatory.

## Wall defined twice in fix wall/reflect command

Self-explanatory.

## Wall defined twice in fix wall/srd command

Self-explanatory.

## Water H epsilon must be 0.0 for pair style lj/cut/tip4p/cut

This is because LAMMPS does not compute the Lennard-Jones interactions with these particles for efficiency reasons.

## Water H epsilon must be 0.0 for pair style lj/cut/tip4p/long

This is because LAMMPS does not compute the Lennard-Jones interactions with these particles for efficiency reasons.

## Water H epsilon must be 0.0 for pair style lj/long/tip4p/long

This is because LAMMPS does not compute the Lennard-Jones interactions with these particles for efficiency reasons.

## World variable count does not match # of partitions

A world-style variable must specify a number of values equal to the number of processor partitions.

## Write\_data command before simulation box is defined

Self-explanatory.

## Write\_restart command before simulation box is defined

The write\_restart command cannot be used before a read\_data, read\_restart, or create\_box command.

## Zero length rotation vector with displace\_atoms

Self-explanatory.

## Zero length rotation vector with fix move

Self-explanatory.

## Zero-length lattice orient vector

Self-explanatory.

# Warning messages

This is an alphabetic list of the WARNING messages LAMMPS prints out and the reason why. If the explanation here is not sufficient, the documentation for the offending command may help. Warning messages also list the source file and line number where the warning was generated. For example, a message like this:

WARNING: Bond atom missing in box size check (domain.cpp:187)

means that line #187 in the file src/domain.cpp generated the error. Looking in the source code may help you figure out what went wrong.

Doc page with [*ERROR messages*](#_bookmark2)

## Adjusting Coulombic cutoff for MSM, new cutoff = %g

The adjust/cutoff command is turned on and the Coulombic cutoff has been adjusted to match the user-specified accuracy.

## Angle atoms missing at step %ld

One or more of three atoms needed to compute a particular angle are missing on this processor. Typically this is because the pairwise cutoff is set too short or the angle has blown apart and an atom is too far away.

## Angle style in data file differs from currently defined angle style

Self-explanatory.

## Angles are defined but no angle style is set

The topology contains angles, but there are no angle forces computed since there was no angle\_style command.

## Atom style in data file differs from currently defined atom style

Self-explanatory.

## Bond atom missing in box size check

The second atom needed to compute a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond atom missing in image check

The second atom in a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond atoms missing at step %ld

The second atom needed to compute a particular bond is missing on this processor. Typically this is because the pairwise cutoff is set too short or the bond has blown apart and an atom is too far away.

## Bond style in data file differs from currently defined bond style

Self-explanatory.

## Bonds are defined but no bond style is set

The topology contains bonds, but there are no bond forces computed since there was no bond\_style command.

## Bond/angle/dihedral extent > half of periodic box length

This is a restriction because LAMMPS can be confused about which image of an atom in the bonded interaction is the correct one to use. “Extent” in this context means the maximum end-to-end length of the bond/angle/dihedral. LAMMPS computes this by taking the maximum bond length, multiplying by the number of bonds in the inter- action (e.g. 3 for a dihedral) and adding a small amount of stretch.

## Both groups in compute group/group have a net charge; the Kspace boundary correction to energy will be non-zero

Self-explanatory.

## Calling write\_dump before a full system init.

The write\_dump command is used before the system has been fully initialized as part of a ‘run’ or ‘minimize’ command. Not all dump styles and features are fully supported at this point and thus the command may fail or produce incomplete or incorrect output. Insert a “run 0” command, if a full system init is required.

## Cannot count rigid body degrees-of-freedom before bodies are fully initialized

This means the temperature associated with the rigid bodies may be incorrect on this timestep.

## Cannot count rigid body degrees-of-freedom before bodies are initialized

This means the temperature associated with the rigid bodies may be incorrect on this timestep.

## Cannot include log terms without 1/r terms; setting flagHI to 1

Self-explanatory.

## Cannot include log terms without 1/r terms; setting flagHI to 1.

Self-explanatory.

## Charges are set, but coulombic solver is not used

Self-explanatory.

## Charges did not converge at step %ld: %lg

Self-explanatory.

## Communication cutoff is 0.0. No ghost atoms will be generated. Atoms may get lost

The communication cutoff defaults to the maximum of what is inferred from pair and bond styles (will be zero, if none are defined) and what is specified via *comm\_modify cutoff* (defaults to 0.0). If this results to 0.0, no ghost atoms will be generated and LAMMPS may lose atoms or use incorrect periodic images of atoms in interaction lists. To avoid, either use *pair style zero* with a suitable cutoff or use *comm\_modify cutoff* .

## Communication cutoff is shorter than a bond length based estimate. This may lead to errors.

Since LAMMPS stores topology data with individual atoms, all atoms comprising a bond, angle, dihedral or improper must be present on any subdomain that “owns” the atom with the information, either as a local or a ghost atom. The communication cutoff is what determines up to what distance from a subdomain boundary ghost atoms are created. The communication cutoff is by default the largest non-bonded cutoff plus the neighbor skin distance, but for short or non-bonded cutoffs and/or long bonds, this may not be sufficient. This warning indicates that there is an increased risk of a simulation stopping unexpectedly because of Bond/Angle/Dihedral/Improper atoms missing. It can be silenced by manually setting the communication cutoff via *comm\_modify cutoff* . However, since the heuristic used to determine the estimate is not always accurate, it is not changed automatically and the warning may be ignored depending on the specific system being simulated.

## Communication cutoff is too small for SNAP micro load balancing, increased to %lf

Self-explanatory.

## Compute cna/atom cutoff may be too large to find ghost atom neighbors

The neighbor cutoff used may not encompass enough ghost atoms to perform this operation correctly.

## Computing temperature of portions of rigid bodies

The group defined by the temperature compute does not encompass all the atoms in one or more rigid bodies, so the change in degrees-of-freedom for the atoms in those partial rigid bodies will not be accounted for.

## Create\_bonds max distance > minimum neighbor cutoff

This means atom pairs for some atom types may not be in the neighbor list and thus no bond can be created between them.

## Delete\_atoms cutoff > minimum neighbor cutoff

This means atom pairs for some atom types may not be in the neighbor list and thus an atom in that pair cannot be deleted.

## Dihedral atoms missing at step %ld

One or more of 4 atoms needed to compute a particular dihedral are missing on this processor. Typically this is because the pairwise cutoff is set too short or the dihedral has blown apart and an atom is too far away.

## Dihedral problem

Conformation of the 4 listed dihedral atoms is extreme; you may want to check your simulation geometry.

## Dihedral problem: %d %ld %d %d %d %d

Conformation of the 4 listed dihedral atoms is extreme; you may want to check your simulation geometry.

## Dihedral style in data file differs from currently defined dihedral style

Self-explanatory.

## Dihedrals are defined but no dihedral style is set

The topology contains dihedrals, but there are no dihedral forces computed since there was no dihedral\_style command.

## Dump dcd/xtc timestamp may be wrong with fix dt/reset

If the fix changes the timestep, the dump dcd file will not reflect the change.

## Energy due to X extra global DOFs will be included in minimizer energies

When using fixes like box/relax, the potential energy used by the minimizer is augmented by an additional energy provided by the fix. Thus the printed converged energy may be different from the total potential energy.

## Estimated error in splitting of dispersion coeffs is %g

Error is greater than 0.0001 percent.

## Ewald/disp Newton solver failed, using old method to estimate g\_ewald

Self-explanatory. Choosing a different cutoff value may help.

## FENE bond too long

A FENE bond has stretched dangerously far. It’s interaction strength will be truncated to attempt to prevent the bond from blowing up.

## FENE bond too long: %ld %d %d %g

A FENE bond has stretched dangerously far. It’s interaction strength will be truncated to attempt to prevent the bond from blowing up.

## FENE bond too long: %ld %g

A FENE bond has stretched dangerously far. It’s interaction strength will be truncated to attempt to prevent the bond from blowing up.

## Fix halt condition for fix-id %s met on step %ld with value %g

Self explanatory.

## Fix SRD walls overlap but fix srd overlap not set

You likely want to set this in your input script.

## Fix bond/create is used multiple times or with fix bond/break - may not work as expected

When using fix bond/create multiple times or in combination with fix bond/break, the individual fix instances do not share information about changes they made at the same time step and thus it may result in unexpected behavior.

## Fix bond/react: Atom affected by reaction too close to template edge

This means an atom which changes type or connectivity during the reaction is too close to an ‘edge’ atom defined in the superimpose file. This could cause incorrect assignment of bonds, angle, etc. Generally, this means you must include more atoms in your templates, such that there are at least two atoms between each atom involved in the reaction and an edge atom.

## Fix bond/swap will ignore defined angles

See the page for fix bond/swap for more info on this restriction.

## Fix deposit near setting < possible overlap separation %g

This test is performed for finite size particles with a diameter, not for point particles. The near setting is smaller than the particle diameter which can lead to overlaps.

## Fix evaporate may delete atom with non-zero molecule ID

This is probably an error, since you should not delete only one atom of a molecule.

## Fix gcmc using full\_energy option

Fix gcmc has automatically turned on the full\_energy option since it is required for systems like the one specified by the user. User input included one or more of the following: kspace, triclinic, a hybrid pair style, an eam pair style, or no “single” function for the pair style.

## Fix langevin gjf using random gaussians is not implemented with kokkos

This will most likely cause errors in kinetic fluctuations.

## Fix property/atom mol or charge w/out ghost communication

A model typically needs these properties defined for ghost atoms.

## Fix qeq CG convergence failed (%g) after %d iterations at %ld step

Self-explanatory.

## Fix qeq has non-zero lower Taper radius cutoff

Absolute value must be <= 0.01.

## Fix qeq has very low Taper radius cutoff

Value should typically be >= 5.0.

## Fix qeq/dynamic tolerance may be too small for damped dynamics

Self-explanatory.

## Fix qeq/fire tolerance may be too small for damped fires

Self-explanatory.

## Fix rattle should come after all other integration fixes

This fix is designed to work after all other integration fixes change atom positions. Thus it should be the last integration fix specified. If not, it will not satisfy the desired constraints as well as it otherwise would.

## Fix recenter should come after all other integration fixes

Other fixes may change the position of the center-of-mass, so fix recenter should come last.

## Fix srd SRD moves may trigger frequent reneighboring

This is because the SRD particles may move long distances.

## Fix srd grid size > 1/4 of big particle diameter

This may cause accuracy problems.

## Fix srd particle moved outside valid domain

This may indicate a problem with your simulation parameters.

## Fix srd particles may move > big particle diameter

This may cause accuracy problems.

## Fix srd viscosity < 0.0 due to low SRD density

This may cause accuracy problems.

## Fixes cannot send data in Kokkos communication, switching to classic communication

This is current restriction with Kokkos.

## For better accuracy use ‘pair\_modify table 0’

The user-specified force accuracy cannot be achieved unless the table feature is disabled by using ‘pair\_modify table 0’.

## Geometric mixing assumed for 1/r^6 coefficients

Self-explanatory.

## Group for fix\_modify temp != fix group

The fix\_modify command is specifying a temperature computation that computes a temperature on a different group of atoms than the fix itself operates on. This is probably not what you want to do.

## H matrix size has been exceeded: m\_fill=%d H.m=%dn

This is the size of the matrix.

## Ignoring unknown or incorrect info command flag

Self-explanatory. An unknown argument was given to the info command. Compare your input with the docu- mentation.

## Improper atoms missing at step %ld

One or more of 4 atoms needed to compute a particular improper are missing on this processor. Typically this is because the pairwise cutoff is set too short or the improper has blown apart and an atom is too far away.

## Improper problem: %d %ld %d %d %d %d

Conformation of the 4 listed improper atoms is extreme; you may want to check your simulation geometry.

## Improper style in data file differs from currently defined improper style

Self-explanatory.

## Impropers are defined but no improper style is set

The topology contains impropers, but there are no improper forces computed since there was no improper\_style command.

## Inconsistent image flags

The image flags for a pair on bonded atoms appear to be inconsistent. Inconsistent means that when the coor- dinates of the two atoms are unwrapped using the image flags, the two atoms are far apart. Specifically they are further apart than half a periodic box length. Or they are more than a box length apart in a non-periodic dimension. This is usually due to the initial data file not having correct image flags for the two atoms in a bond that straddles a periodic boundary. They should be different by 1 in that case. This is a warning because in- consistent image flags will not cause problems for dynamics or most LAMMPS simulations. However they can cause problems when such atoms are used with the fix rigid or replicate commands. Note that if you have an infinite periodic crystal with bonds then it is impossible to have fully consistent image flags, since some bonds will cross periodic boundaries and connect two atoms with the same image flag.

## Increasing communication cutoff for GPU style

The pair style has increased the communication cutoff to be consistent with the communication cutoff require- ments for this pair style when run on the GPU.

## KIM Model does not provide ‘energy’; Potential energy will be zero

Self-explanatory.

## KIM Model does not provide ‘forces’; Forces will be zero

Self-explanatory.

## KIM Model does not provide ‘particleEnergy’; energy per atom will be zero

Self-explanatory.

## KIM Model does not provide ‘particleVirial’; virial per atom will be zero

Self-explanatory.

## Kspace\_modify slab param < 2.0 may cause unphysical behavior

The kspace\_modify slab parameter should be larger to ensure periodic grids padded with empty space do not overlap.

## Less insertions than requested

The fix pour command was unsuccessful at finding open space for as many particles as it tried to insert.

## Library error in lammps\_gather\_atoms

This library function cannot be used if atom IDs are not defined or are not consecutively numbered.

## Library error in lammps\_scatter\_atoms

This library function cannot be used if atom IDs are not defined or are not consecutively numbered, or if no atom map is defined. See the atom\_modify command for details about atom maps.

## Likewise 1-2 special neighbor interactions != 1.0

The topology contains bonds, but there is no bond style defined and a 1-2 special neighbor scaling factor was not

1.0. This means that pair style interactions may have scaled or missing pairs in the neighbor list in expectation of interactions for those pairs being computed from the bond style.

## Likewise 1-3 special neighbor interactions != 1.0

The topology contains angles, but there is no angle style defined and a 1-3 special neighbor scaling factor was not

1.0. This means that pair style interactions may have scaled or missing pairs in the neighbor list in expectation of interactions for those pairs being computed from the angle style.

## Likewise 1-4 special neighbor interactions != 1.0

The topology contains dihedrals, but there is no dihedral style defined and a 1-4 special neighbor scaling factor was not 1.0. This means that pair style interactions may have scaled or missing pairs in the neighbor list in expectation of interactions for those pairs being computed from the dihedral style.

## Lost atoms via change\_box: original %ld current %ld

The command options you have used caused atoms to be lost.

## Lost atoms via displace\_atoms: original %ld current %ld

The command options you have used caused atoms to be lost.

## Lost atoms: original %ld current %ld

Lost atoms are checked for each time thermo output is done. See the thermo\_modify lost command for options. Lost atoms usually indicate bad dynamics, e.g. atoms have been blown far out of the simulation box, or moved further than one processor’s subdomain away before reneighboring.

## MSM mesh too small, increasing to 2 points in each direction

Self-explanatory.

## Mismatch between velocity and compute groups

The temperature computation used by the velocity command will not be on the same group of atoms that velocities are being set for.

## Mixing forced for lj coefficients

Self-explanatory.

## Molecule attributes do not match system attributes

An attribute is specified (e.g. diameter, charge) that is not defined for the specified atom style.

## Molecule has bond topology but no special bond settings

This means the bonded atoms will not be excluded in pairwise interactions.

## Molecule template for create\_atoms has multiple molecules

The create\_atoms command will only create molecules of a single type, i.e. the first molecule in the template.

## Molecule template for fix gcmc has multiple molecules

The fix gcmc command will only create molecules of a single type, i.e. the first molecule in the template.

## Molecule template for fix shake has multiple molecules

The fix shake command will only recognize molecules of a single type, i.e. the first molecule in the template.

## More than one compute centro/atom

It is not efficient to use compute centro/atom more than once.

## More than one compute cluster/atom

It is not efficient to use compute cluster/atom more than once.

## More than one compute cna/atom defined

It is not efficient to use compute cna/atom more than once.

## More than one compute contact/atom

It is not efficient to use compute contact/atom more than once.

## More than one compute coord/atom

It is not efficient to use compute coord/atom more than once.

## More than one compute damage/atom

It is not efficient to use compute ke/atom more than once.

## More than one compute dilatation/atom

Self-explanatory.

## More than one compute erotate/sphere/atom

It is not efficient to use compute erorate/sphere/atom more than once.

## More than one compute hexorder/atom

It is not efficient to use compute hexorder/atom more than once.

## More than one compute ke/atom

It is not efficient to use compute ke/atom more than once.

## More than one compute orientorder/atom

It is not efficient to use compute orientorder/atom more than once.

## More than one compute plasticity/atom

Self-explanatory.

## More than one compute sna/atom

Self-explanatory.

## More than one compute sna/grid

Self-explanatory.

## More than one compute sna/grid/local

Self-explanatory.

## More than one compute snad/atom

Self-explanatory.

## More than one compute snav/atom

Self-explanatory.

## More than one fix poems

It is not efficient to use fix poems more than once.

## More than one fix rigid

It is not efficient to use fix rigid more than once.

## Neighbor exclusions used with KSpace solver may give inconsistent Coulombic energies

This is because excluding specific pair interactions also excludes them from long-range interactions which may not be the desired effect. The special\_bonds command handles this consistently by ensuring excluded (or weighted) 1-2, 1-3, 1-4 interactions are treated consistently by both the short-range pair style and the long-range solver. This is not done for exclusions of charged atom pairs via the neigh\_modify exclude command.

## New thermo\_style command, previous thermo\_modify settings will be lost

If a thermo\_style command is used after a thermo\_modify command, the settings changed by the thermo\_modify command will be reset to their default values. This is because the thermo\_modify command acts on the currently defined thermo style, and a thermo\_style command creates a new style.

## No Kspace calculation with verlet/split

The second partition performs a kspace calculation so the kspace\_style command must be used.

## No automatic unit conversion to XTC file format conventions possible for units lj

This means no scaling will be performed.

## No fixes defined, atoms won’t move

If you are not using a fix like nve, nvt, npt then atom velocities and coordinates will not be updated during timestepping.

## No joints between rigid bodies, use fix rigid instead

The bodies defined by fix poems are not connected by joints. POEMS will integrate the body motion, but it would be more efficient to use fix rigid.

## Not using real units with pair reaxff

This is most likely an error, unless you have created your own ReaxFF parameter file in a different set of units.

## Number of MSM mesh points changed to be a multiple of 2

MSM requires that the number of grid points in each direction be a multiple of two and the number of grid points in one or more directions have been adjusted to meet this requirement.

## OMP\_NUM\_THREADS environment is not set.

This environment variable must be set appropriately to use the OPENMP package.

## One or more atoms are time integrated more than once

This is probably an error since you typically do not want to advance the positions or velocities of an atom more than once per timestep.

## One or more chunks do not contain all atoms in molecule

This may not be what you intended.

## One or more dynamic groups may not be updated at correct point in timestep

If there are other fixes that act immediately after the initial stage of time integration within a timestep (i.e. after atoms move), then the command that sets up the dynamic group should appear after those fixes. This will ensure that dynamic group assignments are made after all atoms have moved.

## One or more respa levels compute no forces

This is computationally inefficient.

## Pair COMB charge %.10f with force %.10f hit max barrier

Something is possibly wrong with your model.

## Pair COMB charge %.10f with force %.10f hit min barrier

Something is possibly wrong with your model.

## Pair brownian needs newton pair on for momentum conservation

Self-explanatory.

## Pair dpd needs newton pair on for momentum conservation

Self-explanatory.

## Pair dsmc: num\_of\_collisions > number\_of\_A

Collision model in DSMC is breaking down.

## Pair dsmc: num\_of\_collisions > number\_of\_B

Collision model in DSMC is breaking down.

## Pair style in data file differs from currently defined pair style

Self-explanatory.

## Pair style restartinfo set but has no restart support

This pair style has a bug, where it does not support reading and writing information to a restart file, but does not set the member variable “restartinfo” to 0 as required in that case.

## Particle deposition was unsuccessful

The fix deposit command was not able to insert as many atoms as needed. The requested volume fraction may be too high, or other atoms may be in the insertion region.

## Proc subdomain size < neighbor skin, could lead to lost atoms

The decomposition of the physical domain (likely due to load balancing) has led to a processor’s subdomain being smaller than the neighbor skin in one or more dimensions. Since reneighboring is triggered by atoms moving the skin distance, this may lead to lost atoms, if an atom moves all the way across a neighboring processor’s subdomain before reneighboring is triggered.

## Reducing PPPM order b/c stencil extends beyond nearest neighbor processor

This may lead to a larger grid than desired. See the kspace\_modify overlap command to prevent changing of the PPPM order.

## Reducing PPPMDisp Coulomb order b/c stencil extends beyond neighbor processor

This may lead to a larger grid than desired. See the kspace\_modify overlap command to prevent changing of the PPPM order.

## Reducing PPPMDisp dispersion order b/c stencil extends beyond neighbor processor

This may lead to a larger grid than desired. See the kspace\_modify overlap command to prevent changing of the PPPM order.

## Replacing a fix, but new group != old group

The ID and style of a fix match for a fix you are changing with a fix command, but the new group you are specifying does not match the old group.

## Replicating in a non-periodic dimension

The parameters for a replicate command will cause a non-periodic dimension to be replicated; this may cause unwanted behavior.

## Resetting reneighboring criteria during PRD

A PRD simulation requires that neigh\_modify settings be delay = 0, every = 1, check = yes. Since these settings were not in place, LAMMPS changed them and will restore them to their original values after the PRD simulation.

## Resetting reneighboring criteria during TAD

A TAD simulation requires that neigh\_modify settings be delay = 0, every = 1, check = yes. Since these settings were not in place, LAMMPS changed them and will restore them to their original values after the PRD simulation.

## Resetting reneighboring criteria during minimization

Minimization requires that neigh\_modify settings be delay = 0, every = 1, check = yes. Since these settings were not in place, LAMMPS changed them and will restore them to their original values after the minimization.

## Restart file used different # of processors

The restart file was written out by a LAMMPS simulation running on a different number of processors. Due to round-off, the trajectories of your restarted simulation may diverge a little more quickly than if you ran on the same # of processors.

## Restart file used different 3d processor grid

The restart file was written out by a LAMMPS simulation running on a different 3d grid of processors. Due to round-off, the trajectories of your restarted simulation may diverge a little more quickly than if you ran on the same # of processors.

## Restart file used different boundary settings, using restart file values

Your input script cannot change these restart file settings.

## Restart file used different newton bond setting, using restart file value

The restart file value will override the setting in the input script.

## Restart file used different newton pair setting, using input script value

The input script value will override the setting in the restart file.

## Restrain problem: %d %ld %d %d %d %d

Conformation of the 4 listed dihedral atoms is extreme; you may want to check your simulation geometry.

## Running PRD with only one replica

This is allowed, but you will get no parallel speed-up.

## SRD bin shifting turned on due to small lamda

This is done to try to preserve accuracy.

## SRD bin size for fix srd differs from user request

Fix SRD had to adjust the bin size to fit the simulation box. See the cubic keyword if you want this message to be an error vs warning.

## SRD bins for fix srd are not cubic enough

The bin shape is not within tolerance of cubic. See the cubic keyword if you want this message to be an error vs warning.

## SRD particle %d started inside big particle %d on step %ld bounce %d

See the inside keyword if you want this message to be an error vs warning.

## SRD particle %d started inside wall %d on step %ld bounce %d

See the inside keyword if you want this message to be an error vs warning.

## Shake determinant < 0.0

The determinant of the quadratic equation being solved for a single cluster specified by the fix shake command is numerically suspect. LAMMPS will set it to 0.0 and continue.

## Shell command ‘%s’ failed with error ‘%s’

Self-explanatory.

## Shell command returned with non-zero status

This may indicate the shell command did not operate as expected.

## Should not allow rigid bodies to bounce off reflecting walls

LAMMPS allows this, but their dynamics are not computed correctly.

## Should not use fix nve/limit with fix shake or fix rattle

This will lead to invalid constraint forces in the SHAKE/RATTLE computation.

## Simulations might be very slow because of large number of structure factors

Self-explanatory.

## Slab correction not needed for MSM

Slab correction is intended to be used with Ewald or PPPM and is not needed by MSM.

## Specifying an ‘subset’ value of ‘0’ is equivalent to no ‘subset’ keyword

Self-explanatory.

## System is not charge neutral, net charge = %g

The total charge on all atoms on the system is not 0.0. For some KSpace solvers this is only a warning.

## Table inner cutoff >= outer cutoff

You specified an inner cutoff for a Coulombic table that is longer than the global cutoff. Probably not what you wanted.

## Temperature for MSST is not for group all

User-assigned temperature to MSST fix does not compute temperature for all atoms. Since MSST computes a global pressure, the kinetic energy contribution from the temperature is assumed to also be for all atoms. Thus the pressure used by MSST could be inaccurate.

## Temperature for NPT is not for group all

User-assigned temperature to NPT fix does not compute temperature for all atoms. Since NPT computes a

global pressure, the kinetic energy contribution from the temperature is assumed to also be for all atoms. Thus the pressure used by NPT could be inaccurate.

## Temperature for fix modify is not for group all

The temperature compute is being used with a pressure calculation which does operate on group all, so this may be inconsistent.

## Temperature for thermo pressure is not for group all

User-assigned temperature to thermo via the thermo\_modify command does not compute temperature for all atoms. Since thermo computes a global pressure, the kinetic energy contribution from the temperature is assumed to also be for all atoms. Thus the pressure printed by thermo could be inaccurate.

## The fix ave/spatial command has been replaced by the more flexible fix ave/chunk and compute chunk/atom commands – fix ave/spatial will be removed in the summer of 2015

Self-explanatory.

## The minimizer does not re-orient dipoles when using fix efield

This means that only the atom coordinates will be minimized, not the orientation of the dipoles.

## Too many common neighbors in CNA %d times

More than the maximum # of neighbors was found multiple times. This was unexpected.

## Too many inner timesteps in fix ttm

Self-explanatory.

## Too many neighbors in CNA for %d atoms

More than the maximum # of neighbors was found multiple times. This was unexpected.

## Use special bonds = 0,1,1 with bond style fene

Most FENE models need this setting for the special\_bonds command.

## Use special bonds = 0,1,1 with bond style fene/expand

Most FENE models need this setting for the special\_bonds command.

## Using a many-body potential with bonds/angles/dihedrals and special\_bond exclusions

This is likely not what you want to do. The exclusion settings will eliminate neighbors in the neighbor list, which the many-body potential needs to calculated its terms correctly.

## Using compute temp/deform with inconsistent fix deform remap option

Fix nvt/sllod assumes deforming atoms have a velocity profile provided by “remap v” or “remap none” as a fix deform option.

## Using compute temp/deform with no fix deform defined

This is probably an error, since it makes little sense to use compute temp/deform in this case.

## Using fix srd with box deformation but no SRD thermostat

The deformation will heat the SRD particles so this can be dangerous.

## Using kspace solver on system with no charge

Self-explanatory.

## Using largest cut-off for lj/long/dipole/long long long

Self-explanatory.

## Using largest cutoff for buck/long/coul/long

Self-explanatory.

## Using largest cutoff for lj/long/coul/long

Self-explanatory.

## Using largest cutoff for pair\_style lj/long/tip4p/long

Self-explanatory.

## Using package gpu without any pair style defined

Self-explanatory.

## Using pair potential shift with pair\_modify compute no

The shift effects will thus not be computed.

## Using pair tail corrections with nonperiodic system

This is probably a bogus thing to do, since tail corrections are computed by integrating the density of a periodic system out to infinity.

## Using pair tail corrections with pair\_modify compute no

The tail corrections will thus not be computed.