# Tinker-HP: Readme/Quickstart (v1.2)

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#### I. CITING TINKER-HP

If you use Tinker-HP please cite the following reference:

Tinker-HP: a Massively Parallel Molecular Dynamics Package for Multiscale Simulations of Large Complex Systems with Advanced Polarizable Force Fields. L. Lagardère, L.-H. Jolly, F. Lipparini, F. Aviat, B. Stamm, Z. F. Jing, M. Harger, H. Torabifard, G. A. Cisneros, M. J. Schnieders, N. Gresh, Y. Maday, P. Ren, J. W. Ponder, J.-P. Piquemal, Chem. Sci., 2018, 9, 956-972 (Open Access) DOI: 10.1039/C7SC04531J

If you use the AVX512 vectorized version of Tinker-HP 1.2, please also cite:

Raising the Performance of the Tinker-HP Molecular Modeling Package [Article v1.0]. L.-H. Jolly, A. Duran, L. Lagardère, J. W. Ponder, P. Y. Ren, J.-P. Piquemal, LiveCoMS, 2019, 1 (2), 10409 (Open Access) DOI: 10.33011/livecoms.1.2.10409

# II. PREREQUISITES

## A. Calculation Libraries

Tinker-HP requires the MKL library, a FFT library (such as FFTW) and a slightly modified 2DECOMP\_FFT library (shipped with Tinker-HP) in order to run. The 2DECOMP\_FFT library enables parallel 3D FFT computations based on 2d-pencils data distribution (see 2DECOMP\_FFT site) based on a sequential implementation of FFTs such as the one provided by the FFTW library.

# B. Parallel library

Tinker-HP also requires a recent enough MPI library supporting MPI 3.x standards such as non blocking collectives. The code has been extensively tested with recent IntelMPI versions (such as intel MPI 5.1) and better performances have been observed with this family of MPI implementation compared to other ones such as OpenMPI.

#### III. INSTALLATION

As Tinker-HP is shipped in source form, you need to compile it. This was not always an easy task in the previous releases. Tinker-HP now uses a configure script built with autotools packages from GNU to ease the compilation and installation process. The first thing you should do is to type:

```
autoconf ; automake
```

in the main directory. This will generate the configure file. Apart from the usual options available with all configure scripts, there are specific options for Tinker-HP.

Usage: ./configure [OPTION]... [VAR=VALUE]... Optional Features: --enable-debug Enable debug mode (check array bounds, implicit none, etc...). Should not be active in normal operations [default is no] Enable AVX512 Optimization for Skylake Processors --enable-skylake [default is no] Enable AVX512 Optimization for KNL (Xeon Phi) --enable-knl Processors [default is no] Enable generic FFT mode [default is yes] --enable-fft-generic --enable-fft-mkl Enable MKL FFT mode [default is no] --enable-fft-fftw3 Enable fftw3 FFT mode [default is no] --enable-fft-fftw3\_f03 Enable fftw3\_f03 FFT mode [default is no] --enable-plumed Enable plumed interface [default is no] --enable-colvars Enable Colvars interface [default is no]

```
Optional Packages:
--with-blaslib=<BLAS LIB> Specify BLAS library [mkl, lapack or /absolute/path/to/BLAS_library]
--with-fftlib=<FFT LIB> Specify a library for FFT called by 2decomp [mkl or fftw3 or /absolute/path/to/FFTW_library]
```

The ultimate goal of this script is to let you type

```
./configure; make; make install; cd example; ./ubiquitin2.run and have everything compiled, installed and running.
```

#### A. List of Options

As for all the configure scripts, you can choose the directory in which the binaries will be copied. So, configure has --prefix=<DIR>.

Tinker-HP has a special interest to know if it will run on AVX-512 capable processors. So, configure has options for that:

```
--enable-slylake
--enable-knl
```

Recall that Tinker-HP needs to make a FFT decomposition with a modified version of the 2DECOMP\_FFT library, which in turn needs a working FFTW library. This is why you can find configure options about FFT interface and library:

```
--enable-fft-generic

--enable-fft-mkl

--enable-fft-fftw3

--enable-fft-fftw3_f03

--with-fftlib=<FFT_LIB>
```

Tinker-HP also needs some functions that resides in a working BLAS library. So, there is an option for that:

```
--with-blaslib=<BLAS LIB>
```

Tinker-HP is now able to be interfaced with PLUMED. So, there is an option for that:

```
--enable-plumed
```

Tinker-HP is now able to be interfaced with COLVARS. So, there is an option for that :

```
--enable-colvars
```

Finally, as there might be some execution problems, or compilation problems for the users who develop code, Tinker-HP has an --enable-debug option.

configure tries to find its path to reach a valid MPI compiler and a valid Fortran compiler by unsetting the environment variables \$FC and \$F77, and reading the environment variable \$PATH. It also tries to find valid FFT and BLAS libraries by reading \$FFTW and \$MKLROOT or \$LAPACK respectively. Most of the time, these environment variables are defined through the module framework. As a try, do a

```
module available 2>&1 | less
```

to see if you have the module framework installed on your machine, and to know what module you can load. configure then figures out how to build the correct Makefiles.

By default, configure chooses the MKL library from Intel as the BLAS and FFTW3 libraries, sets the --enable-fft-mkl option, does not make any processor optimization, and disables debugging. Thus, typing ./configure give the same result as if you have typed ./configure --enable-fft-mkl --with-blaslib=mkl --with-fftlib=mkl.

# B. Using configure

If you want to have different settings than those used by default, you'll have to give configure more information. Be aware that configure cannot magically guess anything. So, the information you give must be precise and complete.

1) Install Directory: By default, this is where you have unzipped and untarred the distribution. If you want another place, use --prefix=<DIR>. You can choose any directory you want, providing that you have permission to create this directory and/or write in it.

2) Processor optimization: The machine on which you compile is not always the one on which Tinker-HP will run. If you know that Tinker-HP is going to run on AVX-512 capable processors, you are strongly encouraged to use one of:

```
--enable-knl for KNL processors (also known as Xeon-Phi) --enable-slylake for Skylake processors.
```

as this will dramatically improve the execution speed. Otherwise, the optimization will be done using the capabilities of the compilation machine, as determined by the compiler.

3) FFT interface: You can choose the interface of FFTW you want to use. This has an effect on the 2DECOMP\_FFT library. So:

```
--enable-fft-generic gives the generic FFT, with no call to FFTW library
--enable-fft-mkl gives the MKL FFT. It also automatically selects the MKL library as the FFTW library.
--enable-fft-fftw3 gives the fftw3 interface, and is designed to work with an external FFTW library.
--enable-fft-fftw3_f03 gives the fftw3 Fortran2003 interface, and is designed to work with an external FFTW library
```

4) FFT Library: You can choose the FFTW library you want to use. It can come from the MKL suite, or some FFTW3 package (either system installed, or compiled by you). So:

```
--with-fftlib=mkl : selects the MKL library, but needs the variable $MKLROOT to be set to the absolute path of the MKL library : selects the MKL library by giving the absolute path of the MKL library : selects the FFTW3 library, but needs the variable $FFTW to be set to the absolute path of the FFTW3 library : selects the FFTW3 library by giving the absolute path of the FFTW3 library library by giving the absolute path of the FFTW3 library by giving the absolute path of the FFTW3 library
```

Here are typical commands you can type. If \$MKLROOT has been correctly set:

```
./configure --enable-fft-mkl --with-fftlib=mkl
```

If you wish to give the absolute path of the library:

```
./configure --enable-fft-mkl --with-fftlib=/path/to/mkl/library ./configure --enable-fft-fftw3 --with-fftlib=/path/to/fftw3/library
```

These last commands can also be written this way:

```
MKLROOT=/path/to/mkl/library ./configure --enable-fft-mkl --with-fftlib=mkl FFTW=/path/to/fftw3/library ./configure --enable-fft-fftw3 --with-fftlib=fftw3
```

5) BLAS library: You can choose the BLAS library you want to use. It can come from the MKL suite or some LAPACK package (either system installed, or compiled by you). So:

```
--with-blaslib=mkl : selects the MKL library, but needs the variable $MKLROOT to be set to the absolute path of the MKL library : selects the MKL library, by giving the absolute path of the MKL library : selects the LAPACK library, but needs the variable $LAPACK library objects to the absolute path of the LAPACK library is selects the LAPACK library, by giving the absolute path of the LAPACK library.
```

Here are typical commands you can type. If \$MKLROOT has been correctly set:

```
./configure --enable-fft-mkl --with-blas=mkl
```

If you wish to give the absolute path of the library:

```
./configure --enable-fft-mkl --with-blas=/path/to/mkl/library ./configure --enable-fft-mkl --with-blas=/path/to/lapack/library
```

These last commands can also be written this way:

```
MKLROOT=/path/to/mkl/library ./configure --enable-fft-mkl --with-blas=mkl LAPACK=/path/to/lapack/library ./configure --enable-fft-mkl --with-blas=lapack
```

6) DEBUG mode: This mode is primarily intended for developers, but can also be useful if you experience errors while running Tinker-HP. Adding --enable-debug to the configure command turns on **boundary checking**, forces **implicit none**, sets the optimization level to **0** (the lowest value) and enables **backtracing** and **all warnings**. The compilation produces all the binaries and gives them the .debug extension, so that you know that these binaries are not optimized.

#### C. Output of configure

configure produces a final log to resume what will be done. It displays using colors (if available) all the information you gave, and everything it has been able to catch from the environment.

Here is the result of a successful run of the configure command:

```
FFTW=/usr/local/fftw-3.3.7/Intel/2018/impi/
./configure --enable-debug --enable-fft-fftw3 --enable-plumed --with-fftlib=fftw3
--with-blaslib=lapack
```

where we give the absolute path of the FFTW3 library in the \$FFTW variable, ask for the DEBUG mode, enable the fftw3 interface, ask for the PLUMED interface, use the FFTW3 library and want the lapack library for BLAS, assuming that the \$LAPACK variable is already set.

```
configure:
configure: **
configure: ** Running Mode
                             : PLUMED DEBUG (binaries'extension is '.debug')
configure: ** MPI Fortran Wrapper : mpiifort
configure: ** Fortran Compiler
                           : ifort
configure: ** Fortran flags
                             : -fpp -DPLUMED -00 -q -u -warn all -check bounds
configure: **
                             : -no-ipo -no-prec-div -inline -heap-arrays
configure: **
                             : -traceback
configure: ** 2decomp Library
                            : -L ../2decomp_fft/src/ -12decomp_fft
configure: ** PLUMED Library
                             : -L ../plumed/Intel/lib/ -lplumed -lplumedKernel
configure: ** PLUMED Includes
                            : -I ../plumed/Intel/include
configure: ** FFTW3 Interface
                            : fftw3 of the FFTW3 library
                            : /usr/local/fftw-3.3.7/Intel/2018/impi//lib
configure: ** FFTW3 Path
configure: ** FFTW3 Includes
                            : -I /usr/local/fftw-3.3.7/Intel/2018/impi//include
configure: ** FFTW3 Library
                             : -lfftw3
configure: ** BLAS Type
                             : LAPACK
configure: ** BLAS Path
                             : /usr/local/Libraries/lapack-3.8.0/Intel/2018
configure: ** Prefix installation : /home/lhj/neutron/Tinker/REL/PME/v1.2
configure: ** Binaries location : /home/lhj/neutron/Tinker/REL/PME/v1.2/bin
configure: **
configure:
```

This log confirms that we are in DEBUG mode and that we use the Intel compiler ifort and the mpiifort wrapper from IntelMPI. As The PLUMED interface has been selected, the configure script shows the PLUMED settings. The installation directory where all binaries (with .debug extension) will be installed is shown as well.

In the case of the PLUMED or the COLVARS interface, the configure command will configure the plumed or the Colvars library as well.

# D. Making binaries

Once you are happy with the option you selected, it's time to run the make command, or even the make install command, which will compile and install all at once.

As the compilation process takes care of the dependencies between subroutines and modules, you can safely use the -j flag of the make command to do parallel compilation. This would dramatically speedup the compilation process.

Anyway, on modern machines, the compilation is not very long, except for 2 or 3 subroutines that can take up to 5(!) minutes to compile, depending on the compiler you use and even on the fastest machines. Everything should compile and link gracefully.

Using make install will copy the binaries into the directory you selected with the --prefix=<DIR> option. The binaries produced will have the following extension:

Mode	Extension
NORMAL	.prod
PLUMED NORMAL	_plumed.prod
COLVARS NORMAL	_colvars.prod
DEBUG	.debug
PLUMED DEBUG	_plumed.debug
COLVARS DEBUG	_colvars.debug

the make install command will also create 5 shell scripts, named after the binaries, to setup the proper running environment. Don't forget to install the binaries you created, or you will not be able to run the examples.

#### IV. NOTE FOR DEVELOPERS

#### A. Writing new sources

We don't want to impose you a unique style of writing. Indeed, we don't have one. But we just want to give you some rules we believe are important for the consistency of Tinker-HP's code.

- a) File format: We use FIXED FORM format throughout all the code, even though the code is written in FORTRAN90. This is mandatory. The compilation process would not work otherwise.
- b) Editing: We always use lowercase letters for code (except for printing purposes). We indent all lines embedded in do....enddo, do....while, etc... statements, or in if...else...endif constructs.
- c) Variable declarations: You are required to use implicit none. If you compile in debug mode, that will be enforced by the compiler.

The order we use to declare variables is:

- 1) integer (4 bytes sized)
- 2) real (8 bytes sized)
- 3) logical
- 4) array (in the same order)
- 5) character (single string or array)

We always try not to mix different types of variables in the same declaration line. This is not just because it is easier to read. That is also because it is more memory efficient, particularly for vectorization, where alignment in memory is crucial. character variables should be put at the very end, since they can have arbitrary lengths and almost never align to a memory boundary. We also try to choose significant names for the variables.

d) Comments: We always begin a comment line by the c character. The ! character should only appear in the middle of a line. This is because the ! character at the beginning is reserved to introduce compiler directives.

If ever you create modules, please comment all the new variables you create, like:

С	maxvalue	atoms directly bonded to an atom
С	maxgrp	user-defined groups of atoms
С	maxtyp	force field atom type definitions
С	maxclass	force field atom class definitions

In subroutines or functions, give as many comments as you believe is needed to understand what your code is doing. That would be precious for you, and for us as well.

# B. Compiling new sources

If you make development on Tinker-HP, it is likely that you would need to add subroutines and modules in the source directory.

All modules should be put in files named MOD\_xxxxx.f, even though they are written in FORTRAN90. All functions and routines should be put in files with names beginning by a lowercase letter and with .f extension. Please, try to find significant names (epolar1tcg2shortreal.f is far better than ep1tc2shre.f). You are required to follow this scheme as much as possible.

To compile your new sources, you should add them in the Makefile.am file of the source directory. We've put some comments in this file, to help you know where to put things. Search for the string Add in the file.

There are 3 different cases<sup>1</sup>:

<sup>&</sup>lt;sup>1</sup>Of course, you can match all three at the same time!

```
yourmain_SOURCES = yourmain.f
```

and

yourmain\_DEPENDENCIES = libtinkermod.a libtinkercalc.a

after the similar lines concerning testgrad.

In the Makefile.am file of the scripts directory, you should also add the line:

```
-(cd $bindir; $(LN_S) -f $(LINK_TO) yourmain)
```

in the install-exec-hook: section, and add yourmain at the end of the uninstall-binSCRIPTS: section.

2) You created new module(s). Add lines

```
libtinkermod_a_SOURCES += MOD_xxxxx.f
just below<sup>2</sup> libtinkermod_a_SOURCES += MOD_virial.f
```

3) You created functions and subroutines. Add lines like

```
libtinkercalc_a_SOURCES += yourexplicitfilename.f
just below<sup>3</sup> the line libtinkercalc a SOURCES += version.f
```

You should now go in the main directory, where configure ac resides, and type autoconf and automake autoconf should not generate any message. automake will probably do, mainly because of a different version than the one we used to create the distribution. In this case, just type aclocal before running automake again. These 2 (or 3) commands will generate a new configure script that takes care of your new sources. You should then run ./configure<sup>4</sup>, compile, install and enjoy debugging your code.

#### V. EXECUTABLES

After having successfully compiled the code, five executable files should be present in the install directory: analyze, bar, dynamic, testgrad and minimize<sup>5</sup>, which are the analogous of the binaries of the Tinker-8 release and require similarly a geometry (given by a \*.xyz file), a simulation setup (given by a \*.key file) and possibly a restart (given by a \*.dyn file) for the dynamic program.

All these executables must run in the same environment you had during the compilation phase. That means the same set of modules, or the correct LIBRARY\_PATH. They should be launched with the mpirun -np x prefix in order to run in parallel with x MPI processes.

#### A. General remarks

The only boundary conditions that are available in this release are periodic boundary conditions treated with Particle Mesh Ewald. Furthermore, only orthorhombic unit cells can be used.

Classical force fields such as AMBER, CHARMM and OPLS are available in Tinker-HP as well as polarizable force fields such as AMOEBA.

# B. analyze

The analyze executable allows potential energy analysis. Compared to the Tinker-8 software, the only option compatible with this binary is "e".

For example the command line:

```
mpirun -np 16 ../bin/analyze dhfr2 e
```

will give you as an output the potential energy terms of the geometry given by a dhfr2.xyz file and with the simulation setup given by the dhfr2.key file. Furthermore, this computation will run on 16 MPI processes.

# C. dynamic

The dynamic executable allows to run molecular dynamics simulation. As for Tinker-8, the command line used to run the MD should give first the number of MD steps to make, then the size of each time step (in femtoseconds), then the time between each writing of geometry (in picoseconds), then the statistical ensemble to sample: 1 is NVE, 2 is NVT, 4 is NPT. For NVT and NPT simulations, this number should be followed by the temperature (in Kelvin) of the simulation, and for NPT simulation by the pressure (in Atmosphere) of the simulation.

For example the command lines:

```
mpirun -np 16 ../bin/dynamic dhfr2 1000 1 1 1
```

will give you as an output 1000 MD steps in NVE for the dhfr2 system, with a 1 fs time step and a 1 ps frequency output.

<sup>&</sup>lt;sup>2</sup>As the compilation process takes care of all the dependencies, the positions of the lines you add are not really significant. But putting the new lines at the end is just a way of remembering they are – well – new.

<sup>&</sup>lt;sup>3</sup>Same remark as above.

<sup>&</sup>lt;sup>4</sup>Presumably with the --enable-debug option flag.

<sup>&</sup>lt;sup>5</sup>If you have ever compiled with --enable-debug before, you should have 5 more binaries.

```
mpirun -np 16 ./dynamic dhfr2 1000 1 1 2 300
```

will give you as an output 1000 MD steps in NVT at 300K for the dhfr2 system, with a 1 fs time step and a 1 ps frequency output.

```
mpirun -np 16 ./dynamic dhfr2 1000 1 1 4 300 1
```

will give you as an output 1000 MD steps in NPT at 300K and 1atm for the dhfr2 system, with a 1 fs time step and a 1 ps frequency output.

# D. testgrad

The testgrad program is absolutely equivalent to the one of the Tinker-8 release: it allows the output of the components of the analytical and/or numerical gradients of the different energy terms.

For example, the command line:

```
mpirun -np 16 ../bin/testgrad dhfr2 Y Y 0.0001 Y
```

will give you as an output all the analytical and numerical gradients (computed with an increment of 0.0001 Angstroms for the positions of the atoms) of all the energy terms of the dhfr2 system.

## E. minimize

The minimize program computes energy minimization starting from a given structure, using a low memory quasi-newton BFGS algorithm as in Tinker-8. The command line used should give the numerical threshold for the convergence of the algorithm.

For example, the command line:

```
mpirun -np 16 ../bin/minimize dhfr2 0.1
```

will compute energy minimization on the dhfr2 structure until the RMS on the gradient is inferior to 0.1. The new geometry will be written at each iteration of the algorithm in the file dhfr2.xyz\_2.

#### VI. KEYWORDS

The main keywords of Tinker-8 are available in Tinker-HP. So, a description of these keywords can be found in the Tinker user guide. Let us review a few of these and some new ones which are specific to Tinker-HP.

# A. Keywords specific to the dynamic program

• Integrators: As in Tinker, the integrator of a dynamic is imposed by the keyword "integrator x", x being one of the available integrator:

**BEEMAN** : The default one

VERLET : Verlet

**BBK** : Langevin Dynamics for constant temperature simulations **BAOAB** : Langevin Dynamics for constant temperature simulations

**RESPA**: Bonded/non bonded respa-split with a velocity-verlet inner loop and with a 0.25 fs default timestep

for the inner loop

BAOABRESPA: Bonded/non bonded respa-split for Langevin dynamics with a BAOAB inner loop, the default time

step for the inner loop is also 0.25 fs

RESPA1 : (Bonded)/(short range non bonded)/(long range non bonded) three level respa1-split with a velocity

verlet inner loop. The default timesteps are 0.25 fs for the inner loop and 2 fs for the intermediate one

BAOABRESPA1: (Bonded)/(short range non bonded)/(long range non bonded) three level respa1-split for Langevin

dynamics with a BAOAB inner loop. The default timesteps are 0.25 fs for the inner loop and 2 fs for

the intermediate one

For all the Langevin integrators (BBK, BAOAB, BAOABRESPA, BAOABRESPA1 and BAOABPISTON), the friction (in  $ps^{-1}$ ) can be controlled by the keyword friction x, the default being  $1 ps^{-1}$ .

For RESPA, BAOABRESPA, RESPA1 and BAOABRESPA1, the inner timestep can be imposed by the keyword dshort x, x being its desired value in ps.

For **RESPA1** and **BAOABRESPA1**, the intermediate timestep can be imposed by the keyword dinter x, x being its desired value in ps.

• Thermostats and barostats: Aside from the Langevin integrators, the thermostats available in Tinker-HP are Berendsen, Bussi (which is the default one) and Andersen. The barostats available in Tinker-HP are the Berendsen (which is the default one), the Monte-Carlo one and the Langevin Piston one. These option can be set by putting the keywords: thermostat berendsen, thermostat bussi, thermostat andersen, barostat berendsen

,barostat montecarlo and barostat langevin in the key file. Note that the Langevin piston barostat is only compatible with BAOAB integrators.

Tinker-HP deals with restart files for dynamic trajectories the same way as Tinker-8 does by creating a \*.dyn file encompassing current positions, velocities and accelerations of the system.

# B. New keywords specific to Tinker-HP

Some new keywords have been introduced in Tinker-HP. The first one concern the algorithm used to converge the polarization equations.

polar-alg x: choose the algorithm used to compute the dipoles solution of the polarization equations. x can be:

1 : Conjugate Gradient with a diagonal preconditioner

2: Jacobi/DIIS

of them being prefixed by tcg:

3 : Truncated Conjugate Gradient (TCG)5 : Divide and Conquer Jacobi/DIIS (default)

TCG is a systematically improvable method with 4 tunable parameters that can be controlled by different keywords, each

**tcgorder**  $\times$  : order of the TCG truncation,  $\times$  can take the value 1 (TCG1) or 2 (TCG2), default is 2 **tcgprec**  $\times$  : use of a diagonal preconditioner.  $\times$  can take the value 1 (YES) or 0 (No), default is 1

teggues x : use of a "direct guess" ( $polarizability \times permanent_electric_field$ ) as a guess. x can take the value 1 (YES)

or 0 (No), default is 0.

tcgpeekx : use of a peek step. x can take the value 1 (YES) or 0 (No), default is 1. When a peek step is used, a Jacobi

Over Relaxation (JOR) is applied to the TCG values of the dipoles with a parameter  $\omega$ . By default, this value

is  $\omega = 1$ . (regular Jacobi step) but three keywords can modify this:

**tcgomega**  $\times$  : change the value of the  $\omega$  parameter to  $\times$ .

**tcgomegafit** : impose a regular fitting of the  $\omega$  parameter to match at regular intervals the fully converged

polarization energy.

**tcgomegafitfreq** x: number of timesteps between two updates of the fitted  $\omega$  parameter. The default of x is 1000.

When the RESPA1 and BAOABRESPA1 integrators are used, one has to solve the short range real space polarization equations at the intermediate time steps. By default, this is done using the same algorithm as the one used to solve the complete polarization equations at the outer time steps. But one can chose a different algorithm to solve the short range polarization by using the keyword **polar-algshort** with the same possible values as for **polar-alg**. If TCG is chosen as a short range polarization solver, one can define all the related option for this solver by taking the same keywords defined above and adding the suffix **short**:

tcgordershort : order of the short range TCG truncation tcgprecshort : use of diagonal preconditioner or not

tcgguessshort : use of a "direct guess"

tcgpeekshort : use of a peek

 ${\it tcgomegashort}\ \omega$  : choice of the peek step parameter

With the introduction of Steered Molecular Dynamics come two keywords:

CVSMD for Constant Velocity Steered Molecular Dynamics

**CFSMD** for Constant Force Steered Molecular Dynamics.

The use of those two forms of steered molecular dynamics (SMD) is described in a dedicated tutorial which can be found in the tutorials/SMD/ directory, along with two subdirectories (CFSMD and CVSMD) containing input files, running scripts and output files. The 2 Tinker Archives corresponding to **CVSMD** and **CFSMD** (178MB each) can be downloaded at http://tinker-hp.ip2ct.upmc.fr/?Download-Process.

During equilibration of a solvated protein it can be useful to impose positional restrains on its alpha carbon, this is why we introduced the keyword: **restrain-backbone** x that puts such restrains with a spring constant of x in kcal/mol/Å on all the atoms that have the tag 'CA' in the xyz file. If x is not defined, a default spring constant of 100 kcal/mol/Å is used.

**I/O** of trajectories works as in Tinker 8: if no keyword is specified then frames will printed individually at the selected frequency. By adding the keyword archive then the frames will be directly appended in an \*.arc file.

Moreover, Tinker-HP now supports **CHARMM dcd format**: adding the line **dcdio** to the keyfile will print a trajectory with the dcd format. When running the analyze or the bar postprocessing programs, if **dcdio** is specified in the key file, then the programs will look for the associated \*dcd file and process the frames it contains.

Unlike Tinker-8, the neighbor-lists for non bonded interactions are computed every x steps (x=20 being the default for a 2 fs time step) and not adjusted dynamically at each time step. This frequency of update can be modified by using the keywords **nlupdate x**, 1 corresponding to a neighbor-list update at each time step (which can be useful during equilibration for example).

Other keywords have been introduced to specify parallel options when running Tinker-HP. A 3D domain decomposition is used to divide the unit cell in as many subdomains as MPI tasks (nproc). If the user does not define it, 3 integers Nx, Ny and Nz so that  $Nx \times Ny \times Nz = nproc$  will be chosen to divide each axis of the simulation box and thus define the 3d domain decomposition. It is possible to manually impose this choice by using the keyword: **DECOMP3D Nx Ny Nz** with Nx, Ny and Nz integers such that  $Nx \times Ny \times Nz = nproc$ .

The current distributed release is only available with the PME algorithm which involves direct and reciprocal space computations in the electrostatic and polarization interactions. As the reciprocal space interactions are known to have a less efficient parallel scaling (because of FFTs) it is possible to specify a lower number of MPI processes that will be dedicated to these computations for both the computation of electrostatic and polarization interactions. This can be done by using the keyword **pme-procs** x, corresponding to x MPI processes dedicated to reciprocal space computations.

To find the ideal x value of **pme-procs**, a good starting point, when a large number of cores is used, is usually to dedicate about  $\frac{1}{4}$  of the total cores to reciprocal space computations. But this parameter depends greatly on the machine used and on the setup of your simulation so it should be adjusted manually by comparing the timings obtained with different values for **pme-procs**. During a dynamic, detailed timings are written when the **verbose** keyword is in the \*.key file. In the future, the parameter **pme-procs** will be adapted heuristically by the program as it is done in popular MD packages.

As explained in the beginning of the document, Tinker-HP distributes the data on a grid used to run 3D FFT in 2d pencils which can be associated to a 2d processor grid as explained in  $2DECOMP\_FFT$  web page. By default, the library looks for an optimal 2d processor grid given the number of cores available but this decomposition can be imposed by the user by setting the keyword: **decompfft-grid n1 n2**, corresponding to a  $n1 \times n2$  processor grid.

Note that the user does not have to specify that he wants neighbor-lists to be used as it is the only available option in Tinker-HP.

Tinker-HP is now compatible with the **PLUMED** library that provides enhanced sampling algorithms, free-energy methods and, more generally, tools to analyze data coming from MD simulations. The user only has to add the line:

# plumed input.dat output.dat

to the Tinker-HP \*.key file in order to run an MD coupled to **PLUMED** with the file **input.dat** as a **PLUMED** input. See https://www.plumed.org/doc for a detailed description of the **PLUMED** capabilities. Input files to compute the PMF associated to the distance between a Na+ and a Cl- ion with the amber99 force field and metadynamics are given: **nacl\_amber.xyz**, **nacl\_amber.key** and **plumed.dat**.

Tinker-HP is now also compatible with the **COLVARS** library that provides similar enhanced sampling algorithms, free-energy methods and, more generally, tools to analyze data coming from MD simulations. To run such a simulation, a COLVARS input with the same name as the Tinker-HP one and the \*.colvars extension has to be added in the working directory. See https://colvars.github.io/master/colvars-refman-tinkerhp.pdf for a detailed description of the **COLVARS** capabilities in combination with Tinker-HP. Input files to compute the PMF associated to the distance between a Na+ and a Cl- ion with the amber99 force field and Adaptive Biasing Force (ABF) are given: nacl\_amber.xyz, nacl\_amber.key and nacl\_amber.colvars.

# VII. EXAMPLES

- 4 examples of systems with associated \*.key files are given in the distribution: ubiquitin2, dhfr2, puddle and pond. The sizes of these systems are respectively: 9737, 23558, 96000 and 288000 atoms, making them good various benchmarks for the program.
  - 4 different setups are given for the ubiquitin system with 4 different key files:

1) **ubiquitin2.key** : regular (langevin with BAOAB integration based) 2 fs respa (bonded/non-bonded split)

computations with DC-JI/DIIS as a polarization solver

2) ubiquitin2tcg.key : 2 fs respa computations with TCG2 (with a diagonal preconditioner, no guess and a peek

step with  $\omega = 1$  as a polarization solver

3) ubiquitin2respa1.key : 6 fs respa1 (bonded/short range non-bonded/long range non-bonded split) langevin with

BAOAB integration computations with DC-JI/DIIS as a short and total polarization solver

4) ubiquitin2respa1tcg.key: 10 fs respa1 langevin with BAOAB integration computations with heavy hydrogen, TCG1

(with a diagonal preconditioner, no guess and no peek step) as a short range polarization

solver and DC-JI/DIIS as a total polarization solver

A fifth example is reserved for debug purposes. It's exactly the same as the first one. ./ubiquitin2.debug.run runs the dynamic.debug binary.

#### VIII. TUTORIALS

Several tutorials can be found in the directory **tutorials** of the release:

A general tutorial to prepare systems for Tinker/Tinker-HP

A tutorial to use Umbrella Sampling with Tinker/Tinker-HP

A tutorial to use Steered MD with Tinker-HP

A tutorial to run simulations including nuclear quantum effects (with path integral techniques or approximate generalized langevin equality of the control o

A tutorial to run alchemical free energy simulations with the Lambda-ABF method: Lambda-ABF.md

## IX. SUPPORT

Tinker-HP is maintained by few people. That means we cannot promise you to answer in a minute to your requests. Anyway, if you have any question or need any support for Tinker-HP, feel free to send a mail to our team at TinkerHP\_Support@ip2ct.upmc.fr. We will answer as soon as we can, providing that we can!