# Benchmarks Gromacs

# 9 juillet 2019

## $1 \quad avx2$

## 1.1 Number of MPI rank: 1

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#### PERFORMANCE RESULTS

gmx tune\_pme for GROMACS 2020-dev-20190708-e6a250d12-dirty-unknown Number of ranks : 1 The mpirun command is : mpirun Passing # of ranks via : -npThe mdrun command is : gmx\_mpi mdrun mdrun args benchmarks : -resetstep 1500 -o bench.trr -x bench.xtc -cpo bench.c Benchmark steps : 1000 dlb equilibration steps: 1500 mdrun args at launchtime: Repeats for each test Input file : ../topol.tpr PME/PP load estimate: 0.999977 Number of particles : 6 : PME Coulomb type Grid spacing x y z : 0.115742 0.115742 0.115742Van der Waals type : Cut-off

Will try these real/reciprocal workload settings: No. scaling rcoulomb nkx nky nkz spacing rvdw tpr file 1.000000 1.000000 108 108 108 0.120000 1.000000 ../topol\_bench00.t

Individual timings for input file 0 (../topol\_bench00.tpr): PME ranks Gcycles ns/day PME/f Remark 0 271.609 9.171 OK. 0 268.941 9.262 OK. -1(-1)269.297 9.249 OK. -1(-1)272.294 9.147 OK.

Tuning took 1.6 minutes.

Summary of successful runs:

Line	tpr	PME ranks	Gcycles Av.	Std.dev.	ns/day	PME/ f
0	0	0	270.275	1.887	9.216	_
1	0	-1(-1)	270.796	2.119	9.198	_

Best performance was achieved with 0 PME ranks (see line 0) Please use this command line to launch the simulation:

mpirun -np 1 gmx\_mpi mdrun -npme 0 -s ../topol.tpr

#### 1.2 Number of MPI rank: 2

#### PERFORMANCE RESULTS

gmx tune\_pme for GROMACS 2020-dev-20190708-e6a250d12-dirty-unknown

Number of ranks : 2

The mpirun command is : mpirun Passing # of ranks via : -np

The mdrun command is : gmx\_mpi mdrun

mdrun args benchmarks : -resetstep 1500 -o bench.trr -x bench.xtc -cpo bench.c

Benchmark steps : 1000 dlb equilibration steps: 1500

mdrun args at launchtime:

Repeats for each test : 2

: ../topol.tpr Input file

PME/PP load estimate : 0.999977

Number of particles : 6 : PME Coulomb type

Grid spacing x y z : 0.115742 0.115742 0.115742

Van der Waals type : Cut-off

Will try these real/reciprocal workload settings:

scaling No. rcoulomb rvdw tpr file nkx nky nkz spacing

1.000000 1.000000 108 108 0.120000 1.000000 ../topol\_bench00.t 108

Individual timings for input file 0 (../topol\_bench00.tpr):

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PME ranks	Gcycles	ns/day	PME/ f	Remark
0	349.458	7.128	_	OK.
0	434.702	5.730	_	OK.

Tuning took 2.3 minutes.

Summary of successful runs:									
Line tpr PME rank	cs Gcycles Av.	Std.dev.	ns/day	PME/ f					
DD grid	·		, -	,					
0 0 0	392.080	60.277	6.429	_					
2 1 1									
1 0 -1( 0	381.979	6.190	6.522	_					
2 1 1									

Best performance was achieved with the automatic number of PME ranks (see line 1 Please use this command line to launch the simulation:

mpirun -np 2 gm $x_-mpi$  mdrun -npme -1 -s ../topol.tpr