

Gromacs-2021.3 (GPU)

4. Gromacs-2021.3 (GPU)

```
 Singularity /fs00/software/singularity-images/ngc_gromacs_2021.3.sif
Gromacs em nvt npt md 100%GPU
722080tiib 72rtxib 723090ib
```

		CPU	(GB)	(GB)	CPU (GHz)	GPU	GPU (GB)	(TFLOPS)
83a100ib	1	64	512	8	2.6	8	40	:82.92 :-----
723090ib	2	48	512	10.7	2.8	8	24	:4.30 :569.28
722080tiib	4	16	128	8.0	3.0	4	11	:3.07 :215.17
72rtxib	3	16	128	8.0	3.0	4	24	:2.30 :195.74

Gromacs-2021.3(GPU) GPU 102808
 464 residues, 9nt DNA, 31709 SOL, 94 NA, 94 CL 50 ns **83a100ib**
250 ns/day **723090ib** **220 ns/day** **722080tiib** **170 ns/day** **72rtxib**
180 ns/day 83a100ib 723090ib 80 NJOBS

/fs00/software/singularity-images/ngc_gromacs_2021.3.sif

em.lsf

```

#BSUB -q 72rtxib
#BSUB -gpu "num=1"
module load singularity/latest
export OMP_NUM_THREADS="$LSB_DJOB_NUMPROC"
SINGULARITY="singularity run --nv /fs00/software/singularity-images/ngc_gromacs_2021.3.sif"
${SINGULARITY} gmx grompp -f minim.mdp -c 1aki_solv_ions.gro -p topol.top -o em.tpr
${SINGULARITY} gmx mdrun -nb gpu -ntmpi 2 -deffnm em

```

nvt

```

#BSUB -q 72rtxib
#BSUB -gpu "num=1"
module load singularity/latest
export OMP_NUM_THREADS="$LSB_DJOB_NUMPROC"
SINGULARITY="singularity run --nv /fs00/software/singularity-images/ngc_gromacs_2021.3.sif"
${SINGULARITY} gmx grompp -f nvt.mdp -c em.gro -r em.gro -p topol.top -o nvt.tpr
${SINGULARITY} gmx mdrun -nb gpu -ntmpi 2 -deffnm nvt

```

npt

```

#BSUB -q 72rtxib
#BSUB -gpu "num=1"
module load singularity/latest

```

```
export OMP_NUM_THREADS="$LSB_DJOB_NUMPROC"
SINGULARITY="singularity run --nv /fs00/software/singularity-images/ngc_gromacs_2021.3.sif"
${SINGULARITY} gmx grompp -f npt.mdp -c nvt.gro -r nvt.gro -t nvt.cpt -p topol.top -o npt.tpr
${SINGULARITY} gmx mdrun -nb gpu -ntmpi 2 -deffnm npt
```



md

```
#BSUB -q 723090ib
#BSUB -gpu "num=1"
module load singularity/latest
export OMP_NUM_THREADS="$LSB_DJOB_NUMPROC"
SINGULARITY="singularity run --nv /fs00/software/singularity-images/ngc_gromacs_2021.3.sif"
${SINGULARITY} gmx grompp -f md.mdp -c npt.gro -t npt.cpt -p topol.top -o md_0_1.tpr
${SINGULARITY} gmx mdrun -nb gpu -bonded gpu -update gpu -pme gpu -pmefft gpu -deffnm md_0_1
```



md



```
#BSUB -q 723090ib
#BSUB -gpu "num=1"
module load singularity/latest
export OMP_NUM_THREADS="$LSB_DJOB_NUMPROC"
SINGULARITY="singularity run --nv /fs00/software/singularity-images/ngc_gromacs_2021.3.sif"
${SINGULARITY} gmx pdb2gmx -f protein.pdb -o protein_processed.gro -water tip3p -ignh -merge all <<< 4
${SINGULARITY} gmx editconf -f protein_processed.gro -o pro_newbox.gro -c -d 1.0 -bt cubic
${SINGULARITY} gmx solvate -cp pro_newbox.gro -cs spc216.gro -o pro_solv.gro -p topol.top
${SINGULARITY} gmx grompp -f ../MDP/ions.mdp -c pro_solv.gro -p topol.top -o ions.tpr
${SINGULARITY} gmx genion -s ions.tpr -o pro_solv_ions.gro -p topol.top -pname NA -nname CL -neutral <<< 13
${SINGULARITY} gmx grompp -f ../MDP/minim.mdp -c pro_solv_ions.gro -p topol.top -o em.tpr
${SINGULARITY} gmx mdrun -v -deffnm em
${SINGULARITY} gmx energy -f em.edr -o potential.xvg <<< "10 0"
${SINGULARITY} gmx grompp -f ../MDP/nvt.mdp -c em.gro -r em.gro -p topol.top -o nvt.tpr
${SINGULARITY} gmx mdrun -deffnm nvt
${SINGULARITY} gmx energy -f nvt.edr -o temperature.xvg <<< "16 0"
${SINGULARITY} gmx grompp -f ../MDP/npt.mdp -c nvt.gro -r nvt.gro -t nvt.cpt -p topol.top -o npt.tpr
${SINGULARITY} gmx mdrun -deffnm npt
${SINGULARITY} gmx energy -f npt.edr -o pressure.xvg <<< "18 0"
```

```

${SINGULARITY} gmx grompp -f ../MDP/md.mdp -c npt.gro -t npt.cpt -p topol.top -o md.tpr
${SINGULARITY} gmx mdrun -v -deffnm md
${SINGULARITY} gmx rms -f md.xtc -s md.tpr -o rmsd.xvg <<< "4 4"

```



GROMACS version: 2021.3-dev-20210818-11266ae-dirty-unknown

Precision: mixed

Memory model: 64 bit

MPI library: thread_mpi

OpenMP support: enabled (GMX_OPENMP_MAX_THREADS = 64)

GPU support: CUDA

SIMD instructions: AVX2_256

FFT library: fftw-3.3.9-sse2-avx-avx2-avx2_128-avx512

CUDA driver: 11.20

CUDA runtime: 11.40



ATOM 218234 (401 Protein residues, 68414 SOL, 9 Ion residues)

nsteps = 100000000 ; 200 ns

eScience GPU em nvt npt 1 GPU md

1 GPU

1	em	nvt	npt	md
---	72rtxib	722080tiib	722080tiib	723090ib
CPU time	1168.45	13960.33	42378.71	
Run time	79	1648	5586	117.428 ns/day 0.204 hour/ns
Turnaround time	197	1732	5661	
2	em	nvt	npt	md
---	72rtxib	722080tiib	72rtxib	722080tiib
CPU time	1399.30	15732.66	40568.04	
Run time	93	1905	5236	106.862 ns/day 0.225 hour/ns

<div><div></div><div>1</div></div>	em	nvt	npt	md
Turnaround time	181	1991	5479	
<div><div></div><div>3</div></div>	em	nvt	npt	md
---	72rtxib	72rtxib	72rtxib	72rtxib
CPU time	1368.11	5422.49	5613.74	
Run time	92	355	366	103.213 ns/day 0.233 hour/ns
Turnaround time	180	451	451	
<div><div></div><div>4</div></div>	em	nvt	npt	md
---	72rtxib	72rtxib	72rtxib	722080tiib
CPU time	1321.15	5441.60	5618.87	
Run time	89	356	369	111.807 ns/day 0.215 hour/ns
Turnaround time	266	440	435	
<div><div></div><div>5</div></div>	em	nvt	npt	md
---	72rtxib	72rtxib	72rtxib	72rtxib
CPU time	1044.17	5422.94	5768.44	
Run time	72	354	380	110.534 ns/day 0.217 hour/ns
Turnaround time	162	440	431	
<div><div></div><div>6</div></div>	em	nvt	npt	md
---	723090ib	723090ib	723090ib	723090ib
CPU time	1569.17	7133.74	6677.25	
Run time	81	326	325	114.362 ns/day 0.210 hour/ns
Turnaround time	75	320	300	
<div><div></div><div>7</div></div>	em	nvt	npt	md
---	723090ib	723090ib	723090ib	722080tiib
CPU time	1970.56	5665.71	6841.73	
Run time	91	253	327	111.409 ns/day 0.215 hour/ns
Turnaround time	123	251	328	

722080tiib 723090ib 83a100ib GPU 72rtxib

1.8 / / =0.45 / / 722080tiib 1.2 / / =0.3 / / 723090ib

1.8 / / =0.3 / / 83a100ib 4.8 / / =0.3 / / 4

6. Gromacs “num=4” 4

GPU http://bbs.keinsci.com/thread-13861-1-1.html

https://developer.nvidia.com/blog/creating-faster-molecular-dynamics-simulations-with-

gromacs-2020/ ATOM 500000 GPU

Gromacs Amber GPU

3090 tesla A100 GPU=4 gromacs

```
gmx mdrun -deffnm $file.pdb.md -ntmpi 4 -ntomp 7 -npme 1 -nb gpu -pme gpu -bonded gpu -pmefft gpu -v
```

Revision #11

Created 3 March 2022 15:46:38 by Yao Ge

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