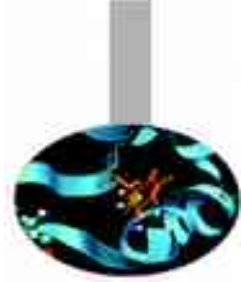


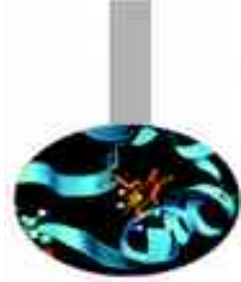
# Running MD code

# What for...?



1. Minimization
2. Molecular Dynamics (classic, brownian, Langevin)
3. Normal Mode Analysis
4. Essential Dynamics and Sampling
5. Free Energy calculations (FEP, Umbrella sampling, AFM)
6. Replica Exchange Molecular Dynamics
7. Coarse-Grained MD
8. Metadynamics
9. Much more...

# Generate topology (Gromacs)



Gromacs:

```
gmx pdb2gmx -f input_file.pdb -ignh -ter
```

input:

1. file\_in.pdb initial set of coordinates (either pdb or gro format)

output:

1. topol.top system topology
2. posre.itp position restraints file
3. conf.gro coordinate file (gro format by default)
4. topolA.itp, topolB.itp, etc topology of chain A, B, etc...
5. posreA.itp, posreB.itp, etc position restraints file for chain A, B, etc...

# How to generate the box (Gromacs)



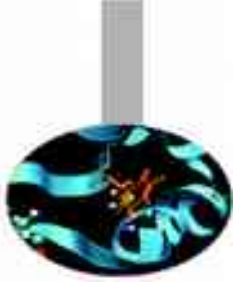
Structure generated file has to be immersed in a box of water molecules (or alternative solvent) prior to run an MD simulation. Different types of box are available in Gromacs (triclinic, cubic, dodecahedron or octahedron) and can be generated by the command:

```
gmx editconf -f conf.gro -bt triclinic -d 0.8 -o output.gro
```

Box type: triclinic in this case

Minimal solute-solvent distance along box axes.

# Box solvation (Gromacs)



Once defined, box has to be physically soaked with water (or alternative solvent). This can be easily performed by running the command:

```
gmx solvate -cp conf.gro -cs spc216.gro -o out.gro -p topol.top
```

# Ionic strength: genion



Grid based electrostatic treatment (Ewald sums, PME, etc.) are better performed with system net charge = 0. Namely, make sure that:

$$\text{solute charge} + \text{solvent charge} = 0$$

To set up box neutrality we can replace as many water molecule with corresponding positive or negative ions to generate a total charge = 0. To do so, we can run the genion command as follows:

```
gmx genion -s topol.tpr -seed XXX -o oution.gro  
-nn 20 -np 10 -p topol.top
```

This command replace randomly a total of 30 water molecules with 20 negative ions (chloride) and 10 positive ions (sodium) and updates the topol.top file with the new list of atoms.

# grompp: the GROMACS preprocessor



Command grompp generates a binary input file with all structural info and forcefield parameters needed to run an MD simulation.

```
gmx grompp -f param.mdp -c coord.gro -n index.ndx  
          -p topol.top -o topol.tpr
```

Grompp output is a binary file called topol.tpr that can be used as input for running the calculation. To visualize and check all info stored in the topol.tpr file we can use the following command:

```
gmx dump -s topol.tpr
```

## Output control

## Van der Waals and electrostatics

## Temperature and pressure coupling

```
title = Yo
cpp = cpp
Include = -I../top
define = -DPOSRES
integrator = md
dt = 0.002
nsteps = 500000
nstxout = 5000
nstvout = 5000
nstlog = 5000
nstenergy = 250
nstxtcou = 250
xtc_grps = Protein
energygrps = Protein SOL
nstlist = 10
ns_type = grid
rlist = 0.8
coulombtype = PME
rcoulomb = 1.4
rvdw = 0.8
tcoupl = V-Rescale
tc-grps = Protein SOL
tau_t = 0.1 0.1
ref_t = 300 300
pcoupl =
tau_p = 1.0
compressibility = 4.5e-5
ref_p = 1.0
gen_vel = yes
gen_temp = 300
gen_seed = 173529
constraints = all-bonds
```





# Output control parameters



nsteps	= 500000	Total number of steps
nstxout	= 5000	coords output frequency for traj.trr
nstvout	= 5000	velocity output frequency for traj.trr
nstlog	= 5000	output frequency for log file (md.log)
nstenergy	= 250	output frequency for energy file ener.edr
nstxtcout	= 250	coords output frequency for traj.xtc
xtc_grps	= Protein	content of file traj.xtc
energygrps	= Protein SOL	energy groups to store in file ener.edr

File [traj.xtc](#) contains coordinates of our simulated system. Atomic coordinates are saved in a compressed format so that to reduce file size. This file is the main trajectory file used for simulation analysis.

File [traj.trr](#) contains atomic coordinates, velocities and forces of our simulated system. These data are saved as 4 digits floating point numbers and are usefull to recover coordinates and velocities after a job crash or if we need velocities and forces for special analyses.

# NAMD input file



22

# standard NAMD benchmark 92k atoms

cellBasisVector1 108.8612 0.0 0.0

cellBasisVector2 0.0 108.8612 0.0

cellBasisVector3 0.0 0.0 77.758

cellOrigin 0.0 0.0 0.0

coordinates apoa1.pdb

temperature 300

seed 74269

switching on

switchdist 10

cutoff 12

pairlistdist 13.5

margin 0

stepspercycle 20

# electrostatics

PME on

PMEGridSizeX 108

PMEGridSizeY 108

PMEGridSizeZ 80

structure apoa1.psf

parameters par\_all22\_prot\_lipid.xplor

parameters par\_all22\_popc.xplor

exclude scaled1-4

1-4scaling 1.0

timestep 1.0

fullElectFrequency 4

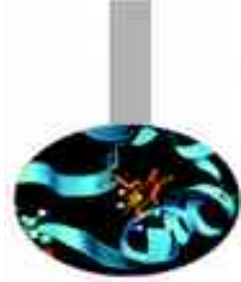
numsteps 10000

outputEnergies 200

outputtiming 200

outputname apoa1

# Amber input file



Typical Production MD NVT

&cntrl

ntx=5, irect=1,

ntc=2, ntf=2,

nstlim=2000,

ntpr=1000, ntwx=1000,

ntwr=2000,

dt=0.002, cut=8.,

ntt=1, tautp=10.0,

temp0=300.0,

ntb=2, ntp=1, barostat=2,

ioutfm=1,

/

# Run the simulation: Gromacs (serial)



```
gmx mdrun -s topol.tpr -dd dx dy dz -pd -npme N
```

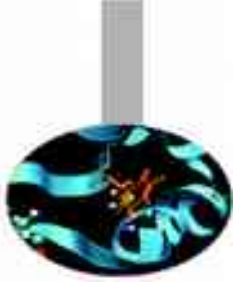
domain decomposition

particle decomposition

The command generates many output files at the end of the job. Among them:

- |                |                                                         |
|----------------|---------------------------------------------------------|
| 1. confout.gro | final coordinates file (gro format)                     |
| 2. traj.xtc    | simulation trajectory file (compressed)                 |
| 3. traj.trr    | simulation trajectory file (coord+velocity, high prec.) |
| 4. ener.edr    | energy file                                             |
| 5. state.cpt   | checkpoint file for restarting runs.                    |
| 6. md.log      | log file with output control                            |

# Run the simulation: NAMD (parallel)



To run NAMD in a pure MPI job:

```
mpirun -np 16 namd2 config.namd > namd.log
```

To run NAMD in a Intel XeonPhi job:

```
mpirun -np 16 namd2.mic config.namd > namd.log
```

To run NAMD on a GPU

```
mpirun -np 16 namd2.cuda -idlepoll +devices 0,1 config.namd > namd.log
```

To run NAMD on a Xeon KNL

```
mpirun -np 16 namd2 -idlepoll +devices 0,1 config.namd > namd.log
```

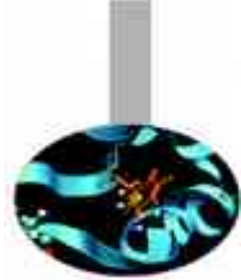
# Run the simulation: Amber



```
mpirun -np 16 pmemd.MPI -O -i run_md.CPU -o dimero_water_SCP_
```

```
mpirun -np 16 pmemd.cuda.MPI -O -i run_md.GPU -o dimero_water_SCP_
```

```
mpirun -np 16 pmemd.mic_offload.MPI -O -i run_md.MIC -o dimero_water_SCP_
```



# What if it all crashes...?

A .cpt file is produced by mdrun at specified intervals (mdrun -cpt n, where n is the interval in minutes), and contains information on all the state variables in a simulated system. In the case of a crash (hardware failure, power outage, etc), a checkpoint file can be used to resume the simulations exactly as it was before the failure. Simulations can also be extended using a checkpoint file ([www.gromacs.org](http://www.gromacs.org)).

```
mdrun -s topol.tpr -cpi state.cpt
```

```
mdrun -s topol.tpr -cpi state.cpt -append
```

Write down coordinates on previous generated files

# Gromacs 5.1.2, pure MPI

```
#!/bin/bash
```

```
#PBS -N gmx
```

```
#PBS -l select=1:ncpus=16:mpiprocs=16:mem=120GB
```

```
#PBS -l walltime=1:00:00
```

```
#PBS -W group_list=train_cmdR2016
```

```
#PBS -A train_cmdR2016
```

```
cd $PBS_O_WORKDIR ==> change to current dir
```

```
module load profile/advanced
```

```
module load autoload gromacs/5.1.2
```

```
export OMP_NUM_THREADS=1 ==> set nr. Of OpenMP threads to 1 per node
```

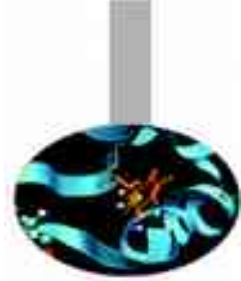
```
mdrun=$(which mdrun_mpi)
```

```
cmd="$mdrun -s topol.tpr -v -maxh 1.0 -nb cpu"
```

```
mpirun -np 16 $cmd
```



# Gromacs 5.1.2, pure MPI job

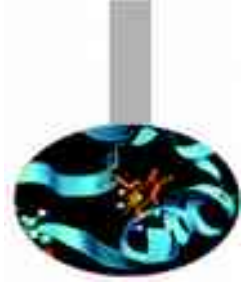


```

File Edit View Search Terminal Help
top - 18:59:29 up 191 days, 8:44, 1 user, load average: 6.03, 6.58, 9.76
Tasks: 521 total, 17 running, 504 sleeping, 0 stopped, 0 zombie
Cpu(s): 99.1%us, 0.9%sy, 0.0%ni, 0.0%id, 0.0%wa, 0.0%hi, 0.0%st, 0.0%st
Mem: 16294372k total, 1183416k used, 15110956k free, 49656k buffers
Swap: 8191992k total, 152304k used, 8039688k free, 184768k cached

  PID USER      PR  NI  VIRT  RES  SHR  S  %CPU  %MEM    TIME+  COMMAND
 25080 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.71  mdrun_mpi
 25082 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.71  mdrun_mpi
 25083 agrottes  20   0   192m  16m  13m  R  100.0  0.1   0:16.72  mdrun_mpi
 25088 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.72  mdrun_mpi
 25089 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.73  mdrun_mpi
 25091 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.72  mdrun_mpi
 25093 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.74  mdrun_mpi
 25094 agrottes  20   0   192m  15m  12m  R  100.0  0.1   0:16.72  mdrun_mpi
 25079 agrottes  20   0  52.2g  27m  23m  R  99.7  0.2   0:16.57  mdrun_mpi
 25081 agrottes  20   0   192m  15m  12m  R  99.7  0.1   0:16.72  mdrun_mpi
 25084 agrottes  20   0   192m  15m  12m  R  99.7  0.1   0:16.71  mdrun_mpi
 25085 agrottes  20   0   192m  15m  12m  R  99.7  0.1   0:16.71  mdrun_mpi
 25090 agrottes  20   0   192m  14m  11m  R  99.7  0.1   0:16.67  mdrun_mpi
 25092 agrottes  20   0   192m  17m  12m  R  99.7  0.1   0:16.71  mdrun_mpi
 25086 agrottes  20   0   192m  15m  12m  R  99.3  0.1   0:16.72  mdrun_mpi
 25087 agrottes  20   0   192m  15m  12m  R  99.3  0.1   0:16.69  mdrun_mpi
 3923  root      0  -20  8811m 320m  45m  S   0.3  2.0  1244:58  mdfsd
19541 abartoli  20   0   172m  5672 1048  S   0.3  0.0   0:00.87  daemon_mic_stat
25120 agrottes  20   0   15292 1576  940  R   0.3  0.0   0:00.04  top
  
```

# Gromacs 5.1.2 MPI+CUDA job



```
#!/bin/bash
```

```
#PBS -N gmx
```

```
#PBS -l select=1:ncpus=2:mpiprocs=2:ngpus=2
```

```
#PBS -l walltime=1:00:00
```

```
#PBS -W group_list=train_cmdR2016
```

```
#PBS -A train_cmdR2016
```

```
cd $PBS_O_WORKDIR ==> change to current dir
```

```
module load profile/advanced
```

```
module load autoload gromacs/5.1.2
```

```
export OMP_NUM_THREADS=1 ==> set nr. Of OpenMP threads to 1
```

```
# ==> set total MPI tasks = 2 and bind to two GPUs
```

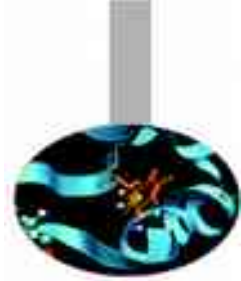
```
mdrun=$(which mdrun_mpi_cuda)
```

```
cmd="$mdrun -s topol.tpr -v -maxh 1.0 -gpu_id 01 "
```

```
mpirun -np 2 $cmd
```



# Gromacs - MPI+CUDA job

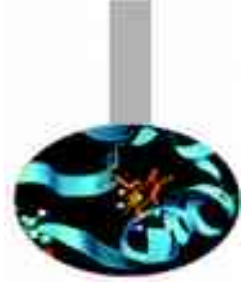


File Edit View Search Terminal Help

```
top - 18:55:58 up 191 days, 8:40, 1 user, load average: 2.63, 8.31, 11.05
Tasks: 509 total, 5 running, 504 sleeping, 0 stopped, 0 zombie
Cpu(s): 12.5%us, 0.1%sy, 0.0%ni, 87.4%id, 0.0%wa, 0.0%hi, 0.0%si, 0.0%st
Mem: 16294372k total, 1306364k used, 14988008k free, 49428k buffers
Swap: 8191992k total, 152304k used, 8039688k free, 185300k cached
```

PID	USER	PR	NI	VIRT	RES	SHR	S	%CPU	%MEM	TIME+	COMMAND
24907	agrottes	20	0	52.3g	106m	91m	R	99.7	0.7	0:20.38	mdrun_mpi
24908	agrottes	20	0	52.2g	103m	90m	R	99.4	0.7	0:20.51	mdrun_mpi
24915	agrottes	20	0	15292	1572	940	R	0.3	0.0	0:00.04	top
1	root	20	0	19352	660	432	S	0.0	0.0	0:17.84	init
2	root	20	0	0	0	0	S	0.0	0.0	0:00.64	kthreadd
3	root	RT	0	0	0	0	S	0.0	0.0	21:41.03	migration/0
4	root	20	0	0	0	0	S	0.0	0.0	34:50.30	ksoftirqd/0
5	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/0
6	root	RT	0	0	0	0	S	0.0	0.0	0:42.24	watchdog/0
7	root	RT	0	0	0	0	S	0.0	0.0	15:43.68	migration/1
8	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/1
9	root	20	0	0	0	0	S	0.0	0.0	17:55.57	ksoftirqd/1
10	root	RT	0	0	0	0	S	0.0	0.0	0:28.06	watchdog/1
11	root	RT	0	0	0	0	S	0.0	0.0	11:14.94	migration/2
12	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/2
13	root	20	0	0	0	0	S	0.0	0.0	12:02.44	ksoftirqd/2
14	root	RT	0	0	0	0	S	0.0	0.0	0:18.76	watchdog/2
15	root	RT	0	0	0	0	S	0.0	0.0	10:45.64	migration/3
16	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/3

# Gromacs MPI/OpenMP+CUDA job



```
#!/bin/bash
#PBS -N gmx
#PBS -l select=1:ncpus=16:mpiprocs=2:ngpus=2
#PBS -l walltime=1:00:00
#PBS -W group_list=train_cmdR2016
#PBS -A train_cmdR2016
```

```
cd $PBS_O_WORKDIR ==> change to current dir
```

```
module load profile/advanced
module load autoload gromacs/5.1.2
```

```
export OMP_NUM_THREADS=8 ==> set nr. Of OpenMP threads to 8
# ==> set 2 MPI tasks that bind to two GPUs
```

```
mdrun=$(which mdrun_mpi_cuda)
cmd="$mdrun -s topol.tpr -v -maxh 1.0 -gpu_id 01 "
mpirun -np 2 $cmd
```

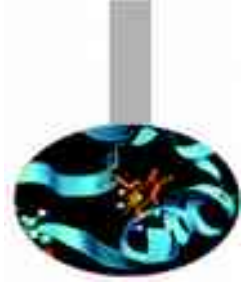




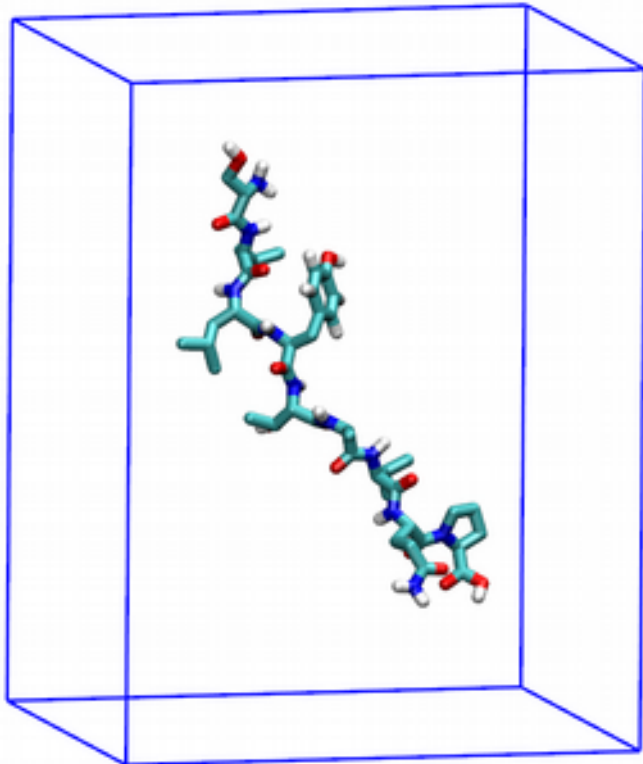
```
File Edit View Search Terminal Help
top - 18:53:13 up 191 days, 8:37, 1 user, load average: 25.69, 13.10, 12.81
Tasks: 509 total, 3 running, 506 sleeping, 0 stopped, 0 zombie
Cpu(s): 95.1%us, 4.9%sy, 0.0%ni, 0.0%id, 0.0%wa, 0.0%hi, 0.0%si, 0.0%st
Mem: 16294372k total, 1387360k used, 14907012k free, 49240k buffers
Swap: 8191992k total, 152304k used, 8039688k free, 187840k cached
```

PID	USER	PR	NI	VRT	RES	SHR	S	%CPU	%MEM	TIME+	COMMAND
24731	agrottes	20	0	53.4g	153m	92m	R	798.6	1.0	11:57.22	mdrun_mpi
24732	agrottes	20	0	53.3g	137m	92m	R	798.6	0.9	11:56.41	mdrun_mpi
67	root	20	0	0	0	0	S	0.3	0.0	128:39.59	events/0
75	root	20	0	0	0	0	S	0.3	0.0	75:53.93	events/8
19426	abartoli	20	0	178m	6692	1720	S	0.3	0.0	0:02.22	daemon_gpu_stat
1	root	20	0	19352	660	432	S	0.0	0.0	0:17.84	init
2	root	20	0	0	0	0	S	0.0	0.0	0:00.64	kthreadd
3	root	RT	0	0	0	0	S	0.0	0.0	21:41.03	migration/0
4	root	20	0	0	0	0	S	0.0	0.0	34:50.30	ksoftirqd/0
5	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/0
6	root	RT	0	0	0	0	S	0.0	0.0	0:42.24	watchdog/0
7	root	RT	0	0	0	0	S	0.0	0.0	15:43.68	migration/1
8	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/1
9	root	20	0	0	0	0	S	0.0	0.0	17:55.57	ksoftirqd/1
10	root	RT	0	0	0	0	S	0.0	0.0	0:28.06	watchdog/1
11	root	RT	0	0	0	0	S	0.0	0.0	11:14.94	migration/2
12	root	RT	0	0	0	0	S	0.0	0.0	0:00.01	migration/2
13	root	20	0	0	0	0	S	0.0	0.0	12:02.43	ksoftirqd/2
14	root	RT	0	0	0	0	S	0.0	0.0	0:18.76	watchdog/2

# MD Performance on hybrid CPU-GPU clusters



## Case study: small peptide in water



- Pure MPI (12 MPI procs) → 123.9 ns/day
- MPI-CUDA (2 MPI procs + 2 GPUs) → 148.0 ns/day
- MPI/OpenMP/CUDA (2 MPI procs + 6 threads + 2 GPUs) → 253.0 ns/day
- MPI + Intel-Phi (12 MPI procs + 34 threads) → 134.1 ns/day

Small peptide in a box of water, ~3300 atoms

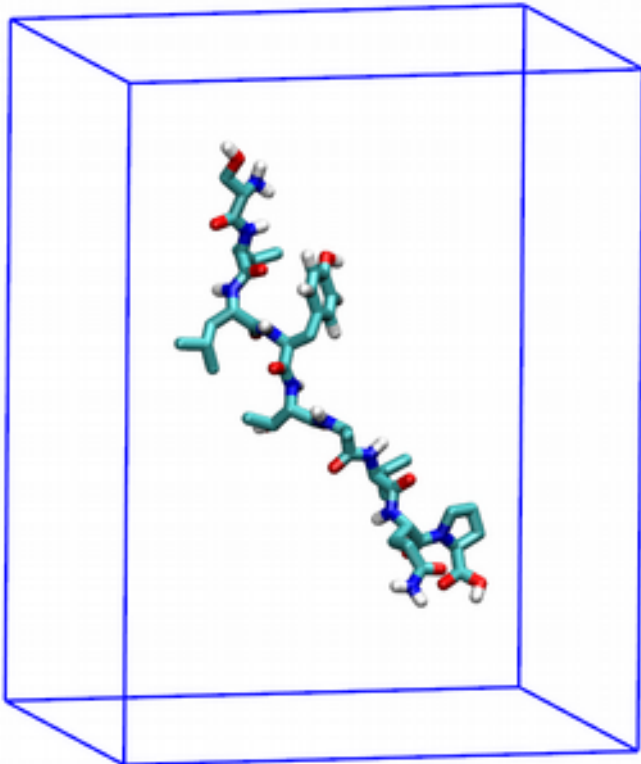
Gromacs 4.6.5 with GPU

PME for long electrostatics, 1 nm cut-off, T = 300 K

# MD Performance on hybrid CPU-GPU clusters



## Case study: small peptide in water



- Pure MPI (16 MPI procs) → 284.1 ns/day
- MPI-CUDA (2 MPI procs + 2 GPUs) → 258.3 ns/day
- MPI/OpenMP/CUDA (2 MPI procs + 8 threads + 2 GPUs) → 457.1 ns/day
- Intel-Phi (native mode) (16 MPI procs + 34 threads) → 68.6/day

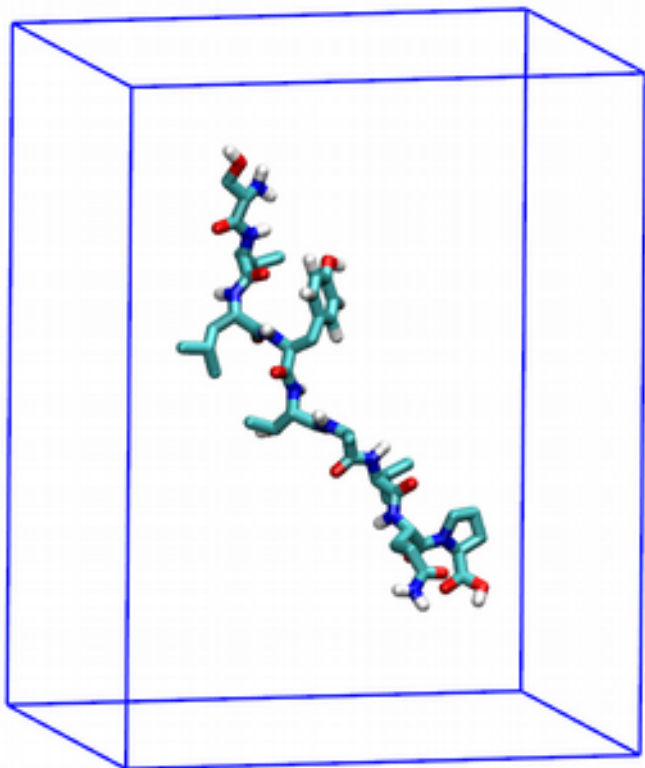
Small peptide in a box of water, ~3300 atoms

Gromacs 5.1.2 with GPU

PME for long electrostatics, 1 nm cut-off, T = 300 K

# MD Performance on hybrid CPU-GPU clusters

multiple MPI ranks with 2 GPUs



- MPI-CUDA (2 MPI procs + 2 GPUs) → 148.0 ns/day
- MPI-CUDA (4 MPI procs + 2 GPUs) → 304.1 ns/day
- MPI-CUDA (8 MPI procs + 2 GPUs) → 221.5 ns/day
- MPI-CUDA (8 MPI procs + 2 thrds + 2 GPUs) → not applicable
- MPI-CUDA (16 MPI + 2 GPUs) → not applicable

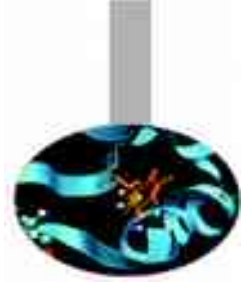
Small peptide in a box of water, ~3300 atoms

Gromacs 4.6.5 with GPU

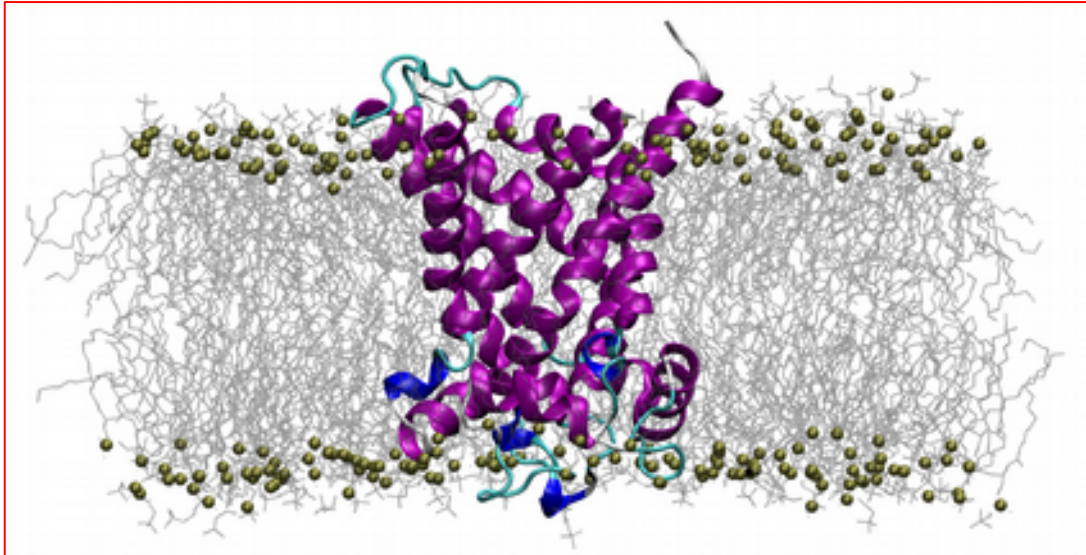
PME for long electrostatics, 1 nm cut-off, T = 300 K



# MD Performance on hybrid CPU-GPU clusters



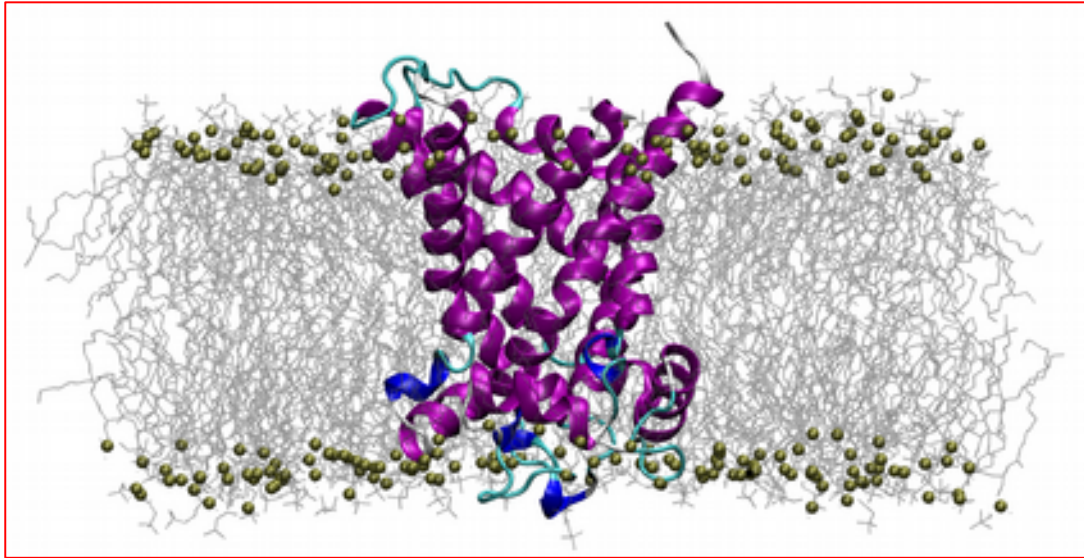
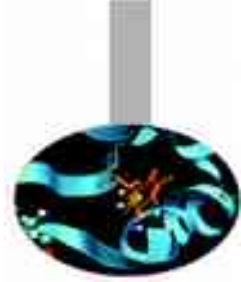
## Case study: membrane protein



ATP/ADP Mitochondrial Carrier,  
92K atoms  
Gromacs 5.0.4 with GPU/MIC  
PME for long electrostatics, 300 K,  
Cut-off = 1 nm  
Domain Decomposition

- Pure MPI (16 MPI procs) → 11.6 ns/day
- MPI-CUDA (2 MPI procs + 2 GPUs) → 9.4 ns/day
- MPI/OpenMP/CUDA (2 MPI procs + 8 threads + 2 GPUs) → 25.9 ns/day
- MPI + Intel Phi (8 MPI procs + 34 threads, Galileo) → 14.6 ns/day

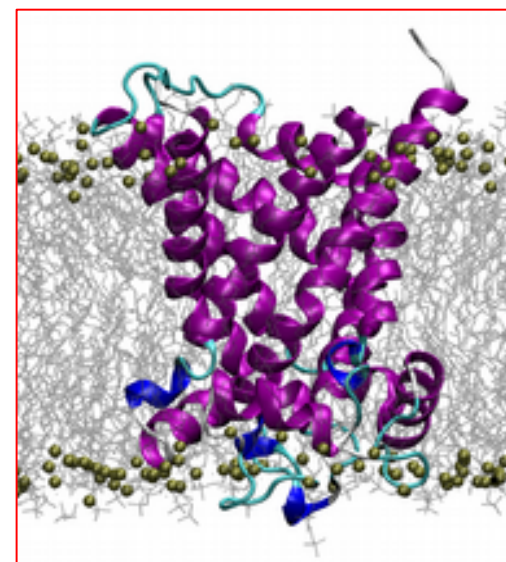
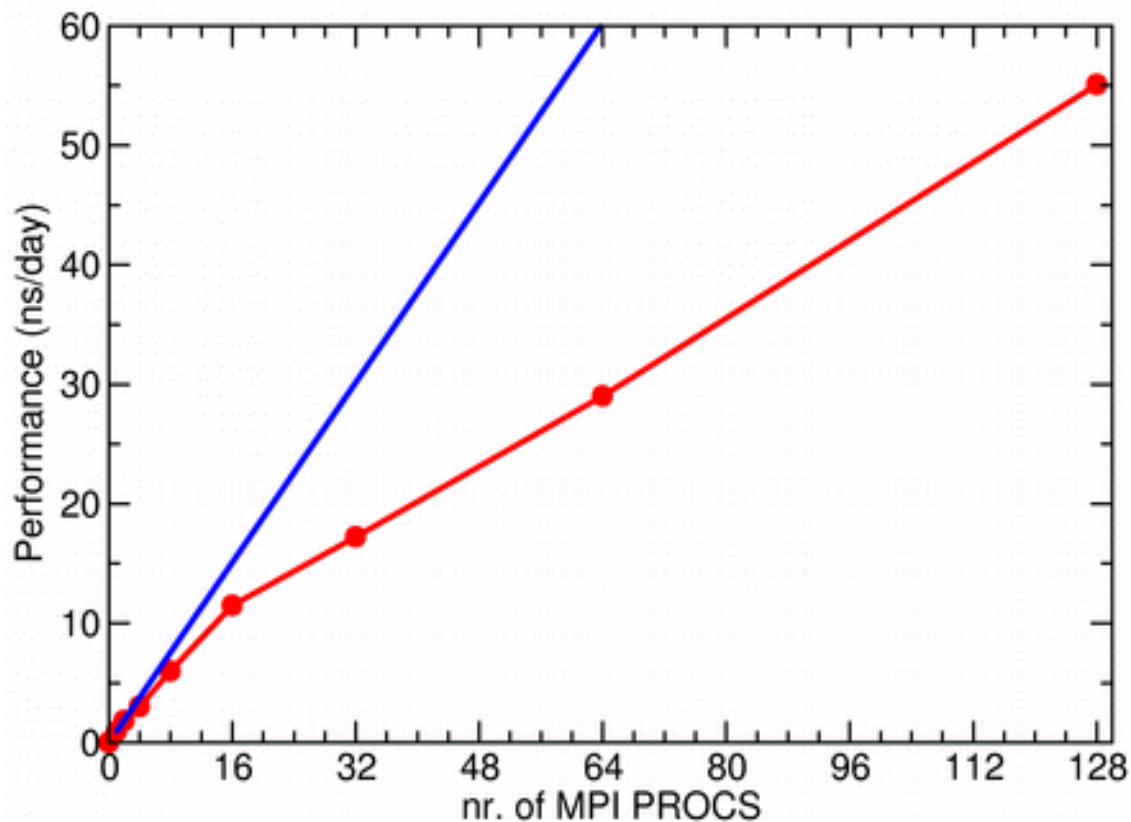
# MD Performance on hybrid CPU-GPU clusters (Eurora)



ATP/ADP Mitochondrial Carrier,  
92K atoms  
Gromacs 5.1.2 with GPU  
PME for long electrostatics, 300 K,  
Cut-off = 1 nm  
Domain Decomposition

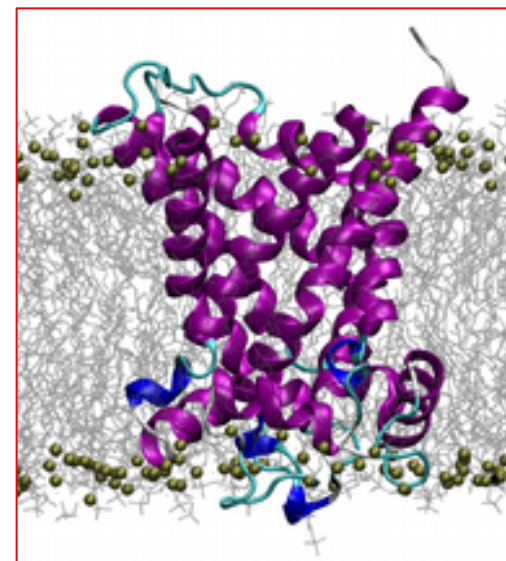
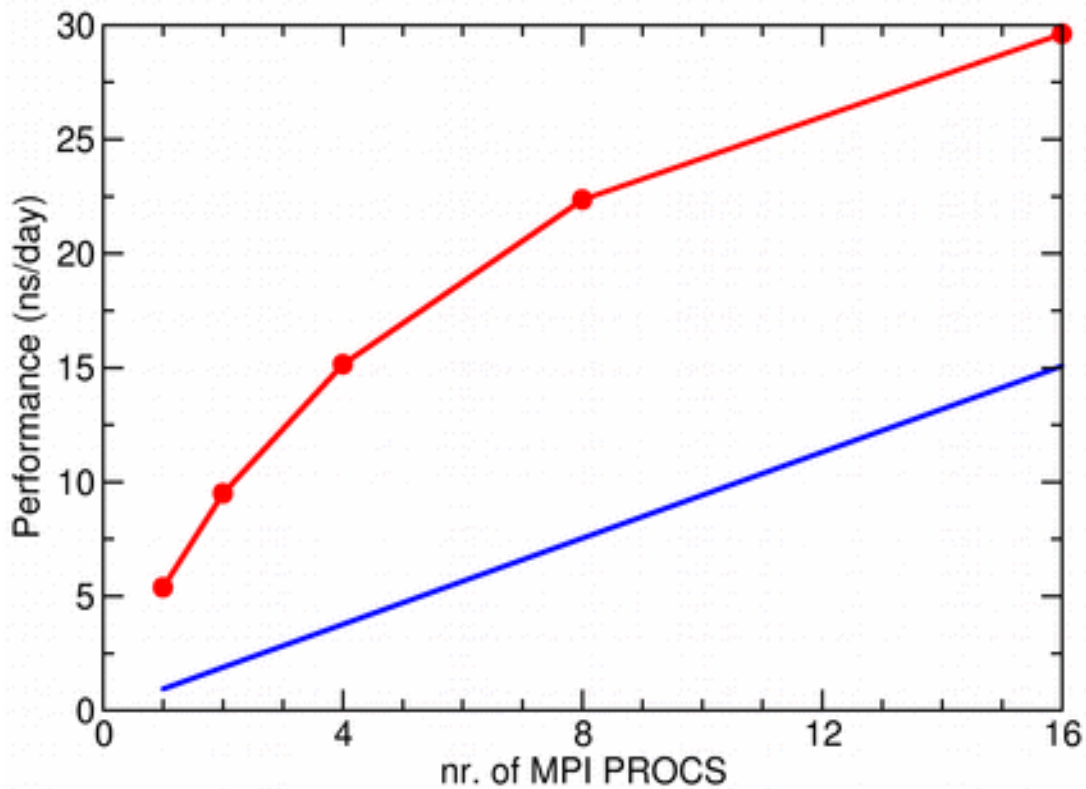
- Pure MPI (16 MPI) → 11.6 ns/day
- MPI-CUDA (2 MPI procs + 2 GPUs) → 9.4 ns/day
- MPI-CUDA (4 MPI procs + 2 GPUs) → 15.2 ns/day
- MPI-CUDA (8 MPI procs + 2 GPUs) → 23.0 ns/day
- MPI-CUDA (16 MPI + 2 GPUs) → 28.9 ns/day
- MPI-CUDA (8 MPI procs + 2 OpenMP + 2 GPUs) → 30.0 ns/day

# Speed up plot for pure MPI calculation



ATP/ADP Mitochondrial Carrier

# MD Performance on hybrid CPU-GPU clusters (Eurora)

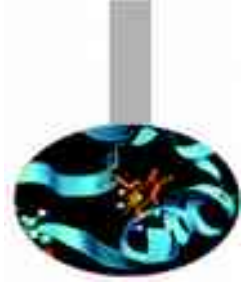


ATP/ADP Mitochondrial Carrier

# GPU acceleration in GROMACS



	Small peptide (3K atoms)	Membrane protein (92K atoms)
Pure MPI	1	1
MPI-CUDA	1.2x	0.8x
multiple MPI ranks/CUDA	2.5x	2.5x
MPI-OpenMP/CUDA	2.0x	2.1x
Intel Phi	1.1x	1.3x



```
#!/bin/bash
#PBS -N namd
#PBS -l select=1:ncpus=16:mpiprocs=16:nmics=2
#PBS -l walltime=00:30:00
#PBS -e job.err
#PBS -o job.out
#PBS -A train_cmdR2016_0
#PBS -W group_list=train_cmdR2016_0

start_time=$(date +%s)
cd ${PBS_O_WORKDIR}

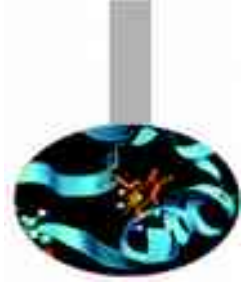
module load profile/advanced
module load autoload namd

source $INTEL_HOME/bin/compilervars.sh intel64
mpirun -n 16 namd2.mic md.namd
```

# Acceleration in NAMD (Galileo)



	Apoa1 (3000 atoms) NAMD 2.10	Speed up
Pure MPI	1.43 ns/day	1
MPI-CUDA	4.1 ns/day	2.8
Intel Xeon Phi	4.3 ns/day	3.0



```
#!/bin/bash
#PBS -N Amber
#PBS -l select=1:ncpus=16:mpiprocs=16:nmics=1:mem=14GB
#PBS -q parallel
#PBS -l walltime=1:00:00
#PBS -A train_cmdR2016
#PBS -W group_list=train_cmdR2016

cd $PBS_O_WORKDIR
module load autoload amber/14
exe=$AMBER_HOME/bin/pmemd.mic_offload.MPI
source $INTEL_HOME/bin/compilervars.sh intel64

mpirun -np 16 $exe -O -i mdin.CPU -o mdout-offload -p prmtop -c inpcrd
```



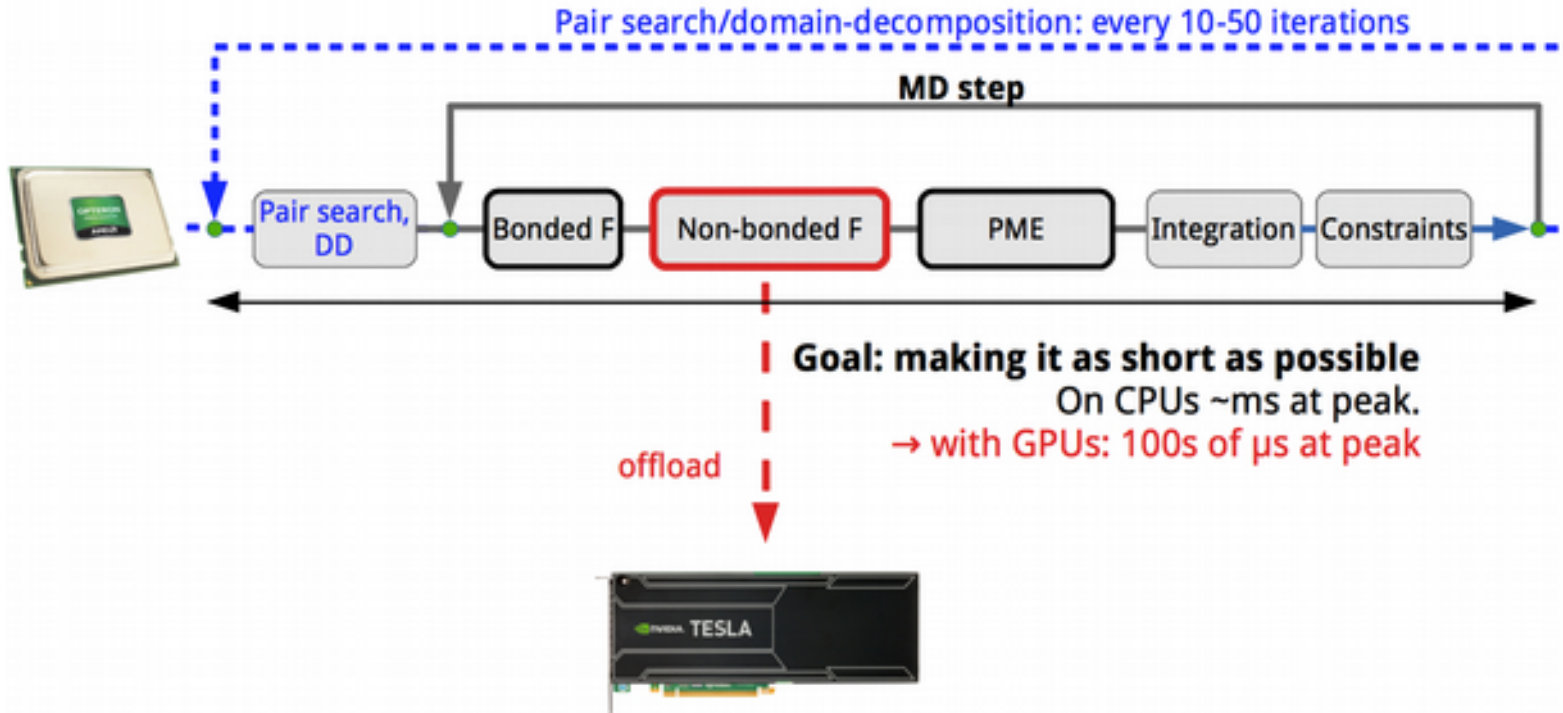
# MD Optimizzation on hybrid CPU-GPU Clusters



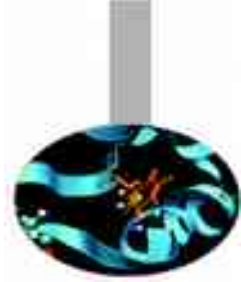
- System size and composition:
- Choice of PME vs. Cut-off based electrostatics
- Larger cut-off radius means a larger Verlet list ==> GPU is better than CPU
- Pure MPI jobs are suitable for small-sized systems

# GPU acceleration in GROMACS

[www.gromacs.org](http://www.gromacs.org)



The idea behind the native GPU acceleration in GROMACS is that **we offload the heavy nonbonded force calculation to an accelerator** (either a GPU or Xeon Phi), while the CPU does bonded forces and lattice summation (PME) in the mean time.



To address the bottleneck caused by multi-threading inefficiencies, it can be advantageous to reduce the number of OpenMP threads per rank. However, to not leave cores empty, this requires using more MPI ranks, hence more PP ranks, and therefore ranks will have to **share GPUs**. GPU sharing is possible by passing a GPU ID to mdrun multiple times, e.g. `-gpu_id 0011` will allow the first two PP ranks in a compute node to use GPU0 and the third and fourth GPU1.

### Example #1:

```
#PBS -l select=1:ncpus=8:mpiprocs=8:ngpus=2:mem=14GB
```

```
...
```

```
OMP_NUM_THREADS=1
```

```
...
```

```
mpirun -np 8 mdrun_mpi_cuda -s topol.tpr -maxh 1.0 -deffnm test -gpu_id 00001111
```

### Example #2:

```
#PBS -l select=1:ncpus=16:mpiprocs=16:ngpus=2:mem=14GB
```

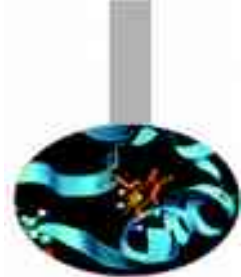
```
...
```

```
OMP_NUM_THREADS=1
```

```
...
```

```
mpirun -np 16 mdrun_mpi_cuda -s topol.tpr -maxh 1.0 -deffnm test -gpu_id 0000000011111111
```





### Example #3:

```
#PBS -l select=1:ncpus=8:mpiprocs=8:ngpus=2:mem=14GB
...
OMP_NUM_THREADS=1
...
mpirun -np 8 mdrun_mpi_cuda -s topol.tpr -maxh 1.0 -deffnm test -gpu_id 01010101
```

### Example #4 (Galileo):

```
#PBS -l select=1:ncpus=16:mpiprocs=8:ngpus=2:mem=14GB
...
OMP_NUM_THREADS=2
...
mpirun -np 8 mdrun_mpi_cuda -s topol.tpr -maxh 1.0 -deffnm test -gpu_id 00001111
```

