

# Outline of Assignment 4, Part I

## Deadlock-free communication

pmd.c

```
if even
    MPI_Send()
    MPI_Recv()
else if odd
    MPI_Recv()
    MPI_Send()
else (self)
    memory copy
```

No ifs & buts!  
cleaner

pmd\_irecv.c

```
MPI_Irecv()
MPI_Send() // OK to send to self
MPI_Wait()
```

## Where?

```
2 in atom_copy()
2 in atom_move()
```

4 code segments in total

## Computation (ns)/communication ( $\mu$ s-ms) overlap

```
MPI_Irecv()
?  
MPI_Send()  
MPI_Wait()
```

[discovery ~]\$ ping hpc-transfer.usc.edu  
time=0.074 ms

...

# Bash Programming

## pmd\_irecv.sl

```
mpicc -O -o pmd_irecv pmd_irecv.c -lm
counter=0
while [ $counter -lt 3 ]; do
    echo "***** Asynchronous *****"
    mpirun -n $SLURM_NTASKS ./pmd_irecv
    echo "***** Synchronous *****"
    mpirun -n $SLURM_NTASKS ./pmd
    let counter+=1
done
```

Value of a variable

Print to terminal

Input-parameter file pmd.in should be in the same directory

Evaluate a mathematical expression & stores its result into a variable

mpicc -O -o pmd pmd.c -lm

See “Bash scripting tutorial for beginners”

<https://linuxconfig.org/bash-scripting-tutorial-for-beginners>

# Runtime Fluctuation

- Due to (1) network interference & (2) shared access to computing nodes, measured runtimes will fluctuate
- The latter could be avoided by exclusive access (`#SBATCH --exclusive`), but **please do not use** this since it will cause very low utilization of computing resources & slow down other users' work

```
***** Asynchronous *****
CPU & COMT = 5.190056e-01 1.406116e-01
***** Synchronous *****
CPU & COMT = 5.163595e-01 1.690525e-01

***** Asynchronous *****
CPU & COMT = 5.119587e-01 1.358150e-01
***** Synchronous *****
CPU & COMT = 5.163731e-01 1.707310e-01

***** Asynchronous *****
CPU & COMT = 5.117363e-01 1.349801e-01
***** Synchronous *****
CPU & COMT = 5.155800e-01 1.672257e-01
```

*pmd\_irecv.c*

*pmd.c*

CPU & COMT reports total run time & communication time, respectively

# Resource Usage (1)

- Start interactive job on discovery & start a MPI program on one of the allocated computing nodes

```
[anakano@discovery cs596]$ salloc --nodes=4 --ntasks-per-node=4 -t 30
salloc: Nodes d05-[33-36] are ready for job
[anakano@d05-33 cs596]$ mpirun -n 16 ./pmd_irecv
...
```

- In another terminal, log in to another allocated node & type 'top' to see running processes

```
[anakano@discovery cs596]$ ssh d05-34
[anakano@d05-34 ~]$ top
top - 07:42:03 up 47 days, 18:34, 2 users, load average: 4.37, 3.33, 3.15
Tasks: 315 total, 8 running, 307 sleeping, 0 stopped, 0 zombie

PID USER      PR  NI    VIRT    RES    SHR S  %CPU  %MEM     TIME+ COMMAND
 3262 rvandamm  20   0 1168000    1.0g  25228 R   100.0   0.5   1090:38 rna_denovo.stat
 3263 rvandamm  20   0 1344840    1.2g  25228 R    99.7   0.6   1090:38 rna_denovo.stat
23608 anakano   20   0 432324 110840  8660 R    99.7   0.1    0:26.48 pmd_irecv
23609 anakano   20   0 432332 108808  8672 R    99.7   0.1    0:26.41 pmd_irecv
23610 anakano   20   0 432324 110856  8676 R    99.7   0.1    0:26.51 pmd_irecv
23607 anakano   20   0 432328 108732  8604 R    99.3   0.1    0:26.43 pmd_irecv
15225 sgopalan  20   0   11.4g  11.2g   7576 R    99.0   5.9   2072:59 R
19675 telegraf  20   0 1507240  49764 18380 S     0.3   0.0   14:08.81 telegraf
23588 anakano   20   0  164372    2508  1612 R     0.3   0.0    0:00.12 top
     1 root      20   0   43572    3956  2528 S     0.0   0.0    2:02.10 systemd
```

4 instances (ranks) of pmd\_irecv are running per node

# Resource Usage (2)

- Type '1' (toggle to show detailed core usage): two users (including myself) are not making full use of cores; let others utilize the unused resources by avoiding exclusive access

```
%Cpu0 : 0.0 us, 0.0 sy, 0.0 ni, 99.7 id, 0.0 wa, 0.0 hi, 0.3 si, 0.0 st
%Cpu1 :100.0 us, 0.0 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu2 :100.0 us, 0.0 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu3 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu4 : 76.3 us, 23.0 sy, 0.0 ni, 0.7 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu5 :100.0 us, 0.0 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu6 :100.0 us, 0.0 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu7 : 99.7 us, 0.3 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu8 : 99.7 us, 0.3 sy, 0.0 ni, 0.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu9 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu10 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu11 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu12 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu13 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu14 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu15 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu16 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu17 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu18 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu19 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu20 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu21 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu22 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
%Cpu23 : 0.0 us, 0.0 sy, 0.0 ni,100.0 id, 0.0 wa, 0.0 hi, 0.0 si, 0.0 st
```

17 out of 24 cores unused

# Note on Assignment 4, Part II

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- Hands-on experience in a common situation of adding new analysis functionality to an existing MPI simulation code, *via* minimally invasive surgery of the code
- Note the header, `pmd.h`, in the homework package, `csci596-as04`, was set for **Part I**:

```
int vproc[3] = {2,2,4}, nproc = 16;
```

The number of MPI ranks should match `nproc` in `pmd.h`:

```
mpirun -n 16 ./pmd (also ./pmd_irecv)
```

- Due to ‘shadow’ analysis ranks, the total number of ranks to be spawned by `mpirun` in **Part II** should instead be twice the number of spatial subsystems, `nproc`, in `pmd_split.c`:

In `pmd_split.h`:

```
int vproc[3] = {2,2,2}, nproc = 8;
```

Run:

```
#SBATCH --nodes=2
```

```
#SBATCH --ntasks-per-node=8
```

```
mpirun -n $SLURM_NTASKS ./pmd_split // $SLURM_NTASKS = 16
```

# Message Composition

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- **Multidimensional arrays are sent as one-dimensional arrays**

```
double rv[NMAX][3];  
double dbuf[NDBUF];
```

