# Batch Scripting for Parallel Systems

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## Purpose:

## **MINESMINESMINESM**

- To give you some ideas about what is possible
- To give you some examples to follow to help you write your own script
- Be a reference, where we can point people
- Document responses to questions

#### To Cover...

- Our test codes
- Bash useful concepts
- Basic Scripts
- Using Variables in Scripts
- Redirecting Output, getting output before a job finishes
- Getting Notifications
- Keeping a record of what you did
- Creating directories on the fly for each job
- Using local disk space

### To Cover...

## **MINESMINESMINESM**

- Multiple jobs on a node
  - Sequential
  - Multiple scripts one node
  - One Script different MPI jobs on different cores

#### To Cover...

- Mapping tasks to nodes
  - Less than N tasks per node
  - Different executables working together
  - Hybrid MPI/OpenMP jobs (MPI and Threading)
  - Running on heterogeneous nodes using all cores
- Job dependencies
  - Chaining jobs
  - Jobs submitting new jobs

## The Source and Scripts

These programs vary from a glorified "Hello World" to being very complex

We also include copies of all our scripts

<a href="http://geco.mines.edu/guide/scripts">http://geco.mines.edu/guide/scripts</a>

```
[joeuser@mio tests]$ [joeuser@mio tests]$ wget <a href="http://geco.mines.edu/scripts/morescripts.tgz">http://geco.mines.edu/scripts/morescripts.tgz</a>
[joeuser@mio tests]$ [joeuser@mio tests]$ tar -xzf <a href="morescripts.tgz">morescripts.tgz</a>
[joeuser@mio tests]$ [joeuser@mio tests]$ cd <a href="morescripts">morescripts</a>
```

```
[joeuser@mio somescripts]$ make 2> /dev/null
mpicc -o c ex00 c ex00.c
mpif90 -o f ex00 f ex00.f
rm -rf fmpi.mod
icc info.c -o info c
ifort info.f90 -o info f
cp info.py info p
chmod 700 info p
ifort -03 -mkl -openmp pointer.f90 -o fillmem
od -vAn -d -N1048576 < /dev/urandom > segment
tar -czf data.tgz segment
rm -rf segment*
mpicc -DDO LOCAL FILE TEST -c sinkfile.c
mpif90 sinkf.f90 sinkfile.o -o sinkf
mpicc -DDO LOCAL FILE TEST -DDO C TEST sinkfile.c -o sinkfile
rm *o *mod
chmod 700 nodes
```

#### What we have

- [c\_ex00.c, c\_ex00]
  - hello world in C and MPI
- [f\_ex00.f, f\_ex00]
  - hello world in Fortran and MPI
- [info.c, info\_c] [info.f90, info\_f] [info.py]
  - Serial programs in C, Fortran and Python that print the node name and process id. Creates a node name process id

#### info.c

```
#include <unistd.h>
#include <sys/types.h>
#include <stdio.h>
#include <stdlib.h>
main() {
    char name[128], fname[128];
    pid t mypid;
    FILE *f;
    char aline[128];
/* get the process id */
    mypid=getpid();
/* get the host name */
    gethostname(name, 128);
/* make a file name based on these two */
    sprintf(fname, "%s_%8.8d", name, (int) mypid);
/* open and write to the file */
    f=fopen(fname, "w");
    fprintf(f, "C says hello from %d on %s\n", (int)mypid, name);
```

#### info.f90

```
program info
  USE IFPOSIX ! needed by PXFGETPID
  implicit none
  integer ierr,mypid
  character(len=128) :: name, fname
! get the process id
  CALL PXFGETPID (mypid, ierr)
! get the node name
  call mynode(name)
! make a filename based on the two
  write(fname, '(a, "_", i8.8)')trim(name), mypid
! open and write to the file
  open(12,file=fname)
  write(12,*)"Fortran says hello from", mypid, " on ", trim(name)
end program
```

```
subroutine mynode(name)
! Intel Fortran subroutine to return
! the name of a node on which you are
! running
   USE IFPOSIX
   implicit none
   integer jhandle
   integer ierr,len
   character(len=128) :: name
   CALL PXFSTRUCTCREATE ("utsname", jhandle, ierr)
   CALL PXFUNAME (jhandle, ierr)
   call PXFSTRGET(jhandle, "nodename", name,len,ierr)
end subroutine
```

## info.py

```
#!/usr/bin/env python
import os
# get the process id
mypid=os.getpid()
# get the node name
name=os.uname()[1]
# make a filename based on the two
fname="%s_%8.8d" % (name,mypid)
# open and write to the file
f=open(fname,"w")
f.write("Python says hello from %d on %s\n" %(mypid,name))
```

## Example Output From the Serial Programs

```
[joeuser@mio cwp]$ ./info_c
[joeuser@mio cwp]$ ls -lt mio*
-rw-rw-r-- 1 joeuser joeuser 41 Jan 11 13:47
mio.mines.edu_00050205
[joeuser@mio cwp]$ cat mio.mines.edu_00050205
C says hello from 50205 on mio.mines.edu
[joeuser@mio cwp]$
```

## C MPI example

```
#include <stdio.h>
#include <stdlib.h>
#include <mpi.h>
#include <math.h>
This is a simple hello world program. Each processor prints out
it's rank and the size of the current MPI run (Total number of
processors).
int main(argc, argv)
int argc;
char *argv[];
   int myid, numprocs, mylen;
   char myname[MPI MAX PROCESSOR NAME];
   MPI Init(&argc,&argv);
   MPI Comm size(MPI COMM WORLD, &numprocs);
   MPI Comm rank(MPI COMM WORLD, &myid);
   MPI Get processor name(myname, &mylen);
/* print out my rank and this run's PE size*/
   printf("Hello from %d of %d on %s\n", myid, numprocs, myname);
   MPI Finalize();
```

## Fortran MPI example

```
This is a simple hello world program. Each processor
  prints out its rank and total number of processors
  in the current MPI run.
program hello
    include "mpif.h"
    character (len=MPI MAX PROCESSOR NAME):: myname
    call MPI INIT( ierr )
    call MPI COMM RANK( MPI COMM WORLD, myid, ierr )
    call MPI COMM SIZE( MPI COMM WORLD, numprocs, ierr )
    call MPI Get processor name(myname, mylen, ierr)
    write(*,*)"Hello from ",myid," of ",numprocs," on ",trim(myname)
    call MPI FINALIZE(ierr)
    stop
    end
```

## Fortran Matrix Inversion Example

- Fills a number of arrays with random data
- Does a matrix inversion
- Used to test the performance of individual cores of a processor
- Can also be used to test threading

```
include 'mkl_vsl.fi'

program testinvert
use numz

call my_clock(cnt1(i))
    CALL DGESV( N, NRHS, twod, LDA, IPIVs(:,i), Bs(:,i), LDB, INFOs(i) )
    call my_clock(cnt2(i))
    write(*,'(i5,i5,3(f12.3))')i,infos(i),cnt2(i),cnt1(i),real(cnt2(i)-cnt1(i),b8)
```

#### sinkfile.c

- A somewhat complicated example
- Does a parallel file copy
  - Copies a file seen by MPI task 0 to
  - Each nodes (not task) in an MPI program that does not share the node used by task 0
  - Used in a situation where MPI tasks might not share file systems

## Batch Scripts

- Batch scripts are just that scripts
  - Run with some "shell", bash, csh, tsh, python
  - Most of the things you can do in a normal script can be done in a batch script
- Some lines in the script are comments to the shell
- Comments can have meaning to the parallel environment
- The parallel environment can define/use variables

#### Bash

- Default shell on CSM machines
- Used to interact with the machine, run commands
- Bash commands can be run interactively or put in a script file
- A script file is really a "simple"
  - Program
  - List of commands
- We will use bash in our examples but other shells and scripting languages have similar capabilities
- First we discuss some features of bash

http://www.tldp.org/LDP/Bash-Beginners-Guide/html/

#### Notes on Commands

- > is used to sent output to a file (date > mylisting)
- >> append output to a file (ls >> mylisting)
- >& send output and error output to a file
- The; can be used to combine multiline commands on a single line. Thus the following are equivalent

date; echo "line 2"; uptime echo "line 2"

date

date

#### Notes on Commands

- Putting commands in ``returns the output of a command into a variable
- Can be use create a list with other commands such as "for loops"

```
myf90=`ls *f90`
echo $myf90
doint.f90 fourd.f90 tintel.f90 tp.f90 vect.f90
```

```
np=`expr 3 + 4`
np=`expr $PBS_NUM_NODES \* 4`
np=`expr $PBS_NUM_NODES / 4`
```

The command expr with "" can be used to do integer math

## For loops

```
myf90=`ls *f90`
          for f in $myf90; do file $f; done
                                                              myf90=`ls *f90`
          doint.f90:ASCII program text
                                                              for f in $myf90
          fourd.f90:ASCII program text
                                                                do file $f
          tintel.f90:ASCII program text
                                                              done
          tp.f90:ASCII program text
          vect.f90:ASCII program text
  for (( c=1; c<=5; c++ )); do echo "Welcome $c times..."; done
  Welcome I times...
  Welcome 2 times...
  Welcome 3 times...
  Welcome 4 times...
                                for c in 1 2 3 4 5; do echo "Welcome $c times..."; done
  Welcome 5 times...
                                Welcome I times...
                                Welcome 2 times...
                                Welcome 3 times...
                                Welcome 4 times...
                                Welcome 5 times...
for c in `seq 1 2 6`; do echo "Welcome $c times..."; date; done
Welcome I times...
                                                                                   for c in 'seq I 26'
Tue Jul 31 12:17:11 MDT 2012
                                                                                    do
Welcome 3 times...
                                                                                    echo "Welcome $c times..."
Tue Jul 31 12:17:11 MDT 2012
                                                                                    date
Welcome 5 times...
                                                                                   done
Tue Jul 31 12:17:11 MDT 2012
```

## Combing Operations

## **MINESMINESMINESM**

Operation	Effect
[!EXPR]	True if <b>EXPR</b> is false.
[(EXPR)]	Returns the value of <b>EXPR</b> . This may be used to override the normal precedence of operators.
[ EXPR1 -a EXPR2 ]	True if both <b>EXPR1</b> and <b>EXPR2</b> are true.
[ EXPR1 -o EXPR2 ]	True if either <b>EXPR1</b> or <b>EXPR2</b> is true.

## Test Variable Being Set and "if"

We do this loop 3 times.

- (I)"var" not set
- (2)"var" set but empty
- (3) var set and not empty

```
var is set but empty B
for i in 1 2 3; do
                                                        i = 3
    echo "i=" $i
                                                        var is set and not empty C
    if [ $i == 1 ] ; then unset var ; fi
    if [ $i == 2 ]; then var=""; fi
    if [ $i == 3 ]; then var="abcd"; fi
    if [ -z "$var" ] ; then echo "var is unset or empty A"; fi
    if [! -n "$var"];
                         then echo "var is unset or empty A2"; fi
    if [ -z "${var-x}" ] ; then echo "var is set but empty B"; fi
    if [ -n "$var" ];
                           then echo "var is set and not empty C"; fi
  echo
done
```

i=1

i=2

var is unset or empty A var is unset or empty A2

var is unset or empty A var is unset or empty A2

## String Tests

```
if test "abc" = "def" ;then echo "abc = def" ; else echo "nope 1" ; fi

if test "abc" != "def" ;then echo "abc != def" ; else echo "nope 2" ; fi

if [ "abc" \< "def" ];then echo "abc < def" ; else echo "nope 3" ; fi

if [ "abc" \> "def" ]; then echo "abc > def" ; else echo "nope 4" ; fi

if [ "abc" \> "abc" ]; then echo "abc > abc" ; else echo "nope 5" ; fi
```

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nope 1
abc != def
abc < def
nope 4
nope 5</pre>

## String Tests

```
if test "abc" = "def" ; then echo "abc = def" ; else echo "nope 1" ; fi
                                                               nope 1
if test "abc" != "def" ; then echo "abc != def" ; else echo "nope 2" ; fi
                                                               abc != def
if [ "abc" \< "def" ]; then echo "abc < def" ; else echo "nope 3" ; fi
                                                               abc < def
if [ "abc" \> "def" ]; then echo "abc > def"; else echo "nope 4"; fi
                                                               nope 4
if [ "abc" \> "abc" ]; then echo "abc > abc"; else echo "nope 5"; fi
                                                               nope 5
```

## File Tests

Test Meaning		
-a FILE ]	True if FILE exists.	
-b FILE]	True if FILE exists and is a block-special file.	
-c FILE]	True if FILE exists and is a character-special file.	
-d FILE ]	True if FILE exists and is a directory.	
e FILE	True if FILE exists.	
-f FILE ]	True if FILE exists and is a regular file.	
-g FILE ]	True if FILE exists and its SGID bit is set.	
-h FILE ]	True if FILE exists and is a symbolic link.	
-k FILE ]	True if FILE exists and its sticky bit is set.	
-p FILE ]	True if FILE exists and is a named pipe (FIFO).	
-r FILE ]	True if FILE exists and is readable.	
-s FILE ]	True if FILE exists and has a size greater than zero.	
-t FD]	True if file descriptor FD is open and refers to a terminal.	
-u FILE ]	True if FILE exists and its SUID (set user ID) bit is set.	
-w FILE ]	True if FILE exists and is writable.	
-x FILE ]	True if FILE exists and is executable.	
-O FILE ]	True if FILE exists and is owned by the effective user ID.	
-G FILE ]	True if FILE exists and is owned by the effective group ID.	
-L FILE ]	True if FILE exists and is a symbolic link.	
-N FILE ]	True if FILE exists and has been modified since it was last read.	
-s file ]	True if FILE exists and is a socket.	
FILE1 -nt FILE2]	True if FILE1 has been changed more recently than FILE2, or if FILE1 exists and FILE2 does not.	
FILE1 -ot FILE2 ]	True if FILE1 is older than FILE2, or is FILE2 exists and FILE1 does not.	
FILE1 -ef FILE2]	True if FILE1 and FILE2 refer to the same device and inode numbers.	

## Checking Terminal Input

```
echo "Do you want to proceed?"
echo -n "Y/N:
read yn
if [ \$yn = "y" ] || [ \$yn = "Y" ]; then
 echo "You said yes"
else
 echo "You said no"
```

Note spacing in the if statement. It is important!

fi

## Testing Return Code & /dev/null

- •Commands return an exit code
  - $\bullet 0 = success$
  - •not 0 = failure
- •The exit code from the previous command is stored in \$?
- •\$? can be echoed or tested
- •This is often used with piping output into /dev/null "the bit bucket" when you only want to know if a command was successful

```
ls a_dummy_file >& /dev/null

if [ $? -eq 0 ]; then
    echo "ls of a_dummy_file successful"
fi
```

#### While and with a Test and break

```
rm -f a dummy file
while true; do
  ls a dummy file >& /dev/null
  if [ $? -eq 0 ] ; then
    echo "ls of a dummy file successful"
  else
   echo "ls of a dummy file failed"
  fi
  if [ -a a dummy file ]; then
    echo "a dummy file exists, breaking"
    break
  else
    echo "a dummy file does not exist"
  fi
  touch a dummy file
  echo; echo "bottom of while loop"; echo
done
```

```
ls of a_dummy_file failed
a_dummy_file does not exist
```

bottom of while loop

ls of a\_dummy\_file successful
a\_dummy\_file exists, breaking

## Running Batch Scripts

- A batch script is submitted to a scheduler
  - pbs/torque/moab, sge, lsf, poe,slurm
  - Commands to submit scripts
    - qsub, msub, bsub, poe,sbatch
- The scheduler decides where and when to run your script
  - Wait for nodes to become available
  - Wait for other jobs to finish
  - Jobs are given a name so that you can track them in the system

## Related Commands SLURM

Command	Description - From http://slurm.schedmd.com/man_index.html	
sbatch	Submit a batch script to SLURM.	
srun	Run parallel jobs	
scancel	Used to signal (cancel) jobs or job steps that are under the control of Slurm.	
salloc	Obtain a SLURM job allocation (a set of nodes), Useful for interactive sessions.	
sacct	Displays accounting data for all jobs and job steps in the SLURM job accounting log or SLURM database	
sacctmgr	Used to view and modify Slurm account information.	
sattach	Attach to a SLURM job step.	
sdiag	scheduling diagnostic tool.	
sinfo	view information about SLURM nodes and partitions.	
smap	graphically view information about SLURM jobs, partitions, and set configurations parameters.	
sprio	view the factors that comprise a job's scheduling priority	
squeue	view information about jobs located in the SLURM scheduling queue.	
sreport	Generate reports from the slurm accounting data.	
sstat	Display various status information of a runங்கள் job/step.	

## CSM Unique SLURM Commands

Command /opt/utility/*	Description	
sjobs	Summary of running and queued jobs	
slurmjobs	Show full information for all jobs -h for help	
slurmnodes	Show full information for all nodes (-h for help	
inuse	Node usage by group	
match	Creates an mpiexec "appfile" for MPMD runs and nonstandard mappings of tasks to nodes	
match_split	Creates an srun "multi-prog" for MPMD runs and nonstandard mappings of tasks to nodes	
phostname	Glorified MPI/OpenMP "hello world"  31	

## phostname "help"

```
[ioeuser@aun002 ~]$ /opt/utility/phostname -h
phostname arguments:
         -h : Print this help message
no arguments: Print a list of the nodes on which the command is run.
-f or -1
             : Same as no argument but print MPI task id and Thread id
               If run with OpenMP threading enabled OMP NUM THREADS > 1
               there will be a line per MPI task and Thread.
-F or -2
             : Add columns to tell first MPI task on a node and and the
               numbering of tasks on a node. (Hint: pipe this output in
               to sort -r
             : Print a listing of the environmental variables passed to
               MPI task. (Hint: use the -l option with SLURM to prepend MPI
               task #1)
[joeuser@aun002 ~]$
```

## A Simple Slurm Script for a MPI job

#!/bin/bash

```
#SBATCH --job-name="atest"
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=8
                                      Scripts contain comments
#SBATCH --time=00:02:00
                                     designated with a # that are
#SBATCH -o stdout
                                       interpreted by SLURM
#SBATCH -e stderr
#SBATCH --export=ALL
                                     and normal shell commands
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
                                        We go to this directory
cd ~/bins/example/mpi
                                        Run this MPI program on
srun - n 8 ./c ex00 \leftarrow
                                                8 cores
```

## A Simple Slurm Script for a MPI job

#!/bin/bash	This is a bash script
#SBATCHjob-name="atest"	Give our job a name in the
#SBATCHnodes=1	We want I node
#SBATCHntasks-per-node=8	We expect to run 8 tasks/node
#SBATCHtime=00:02:00	We want the node for 2 minutes
#SBATCHoutput=stdout	Output will go to a file "stdout"
#SBATCHerror=stderr	Errors will go to a file "stdout"
#SBATCHexport=ALL	Pass current environment to nodes
#SBATCHmail-type=ALL	Send email on abort, begin, end
#SBATCHmail-user=joeuser@mines.edu	Address for email
#	Just a normal "comment"
cd /home/joeuser/examples	Go to this directory first
srun -n 8 ./c_ex00	Run c_ex00 on 8 cores

## What happens when you run a script?

- You are given a collection of nodes
- You are logged on to one of the nodes, the primary compute node
- Any "normal" script command only run on the primary compute node
- Extra effort must be taken to run on all nodes
- srun or mpiexec (srun for Slurm, mpiexec is used for PBS)
  - Also Run only on the primary compute node
  - Makes the effort to launch MPI jobs on all nodes



### Slurm "script" Variables

Variable	Meaning	Typical Value
SLURM_SUBMIT_DIR	Directory for the script	/panfs/storage/scratch/joeuser
SLURM_JOB_USER	Who are you	joeuser
SLURM_EXPORT_ENV	Variables to export	ALL
SLURM_NNODES	# nodes for the job	2
SLURM_JOBID	Job ID	11160
SLURM_NODELIST	Compressed list of nodes	node[114-115]
SLURM_SUBMIT_HOST	Host used to launch job	aun002.mines.edu

You can also use variables you define before you submit your script and variables defined in your environment

### Example list of variables

```
SLURM CHECKPOINT IMAGE DIR=/bins/joeuser/examples/mpi
SLURM_NODELIST=node[114-115]
SLURM JOB NAME=atest
SLURMD NODENAME=node114
SLURM_TOPOLOGY_ADDR=node114
SLURM NTASKS PER NODE=8
SLURM PRIO PROCESS=0
SLURM NODE ALIASES=(null)
SLURM EXPORT ENV=ALL
SLURM TOPOLOGY ADDR PATTERN=node
SLURM NNODES=2
SLURM JOBID=11160
SLURM NTASKS=16
SLURM TASKS PER NODE=8(x2)
SLURM JOB ID=11160
SLURM_JOB_USER=joeuser
SLURM JOB UID=15049
SLURM_NODEID=0
SLURM_SUBMIT_DIR=/bins/joeuser/examples/mpi
SLURM TASK PID=14098
SLURM NPROCS=16
SLURM CPUS ON NODE=8
SLURM PROCID=0
SLURM_JOB_NODELIST=node[114-115]
SLURM LOCALID=0
SLURM_JOB_CPUS_PER_NODE=8(x2)
SLURM GTIDS=0
SLURM SUBMIT HOST=aun002.mines.edu
SLURM_JOB_PARTITION=aun
SLURM_JOB_NUM_NODES=2
```

```
#!/bin/bash
#SBATCH -- job-name="atest"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=8
#SBATCH --time=00:02:00
#SBATCH -o stdout
#SBATCH -e stderr
#SBATCH --export=ALL
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
                                        We go to "starting"
cd $SLURM SUBMIT DIR
                                            directory
srun -n 8 ./c_ex00
                                     Run this MPI program on
                                             8 cores
```

```
#!/bin/bash
#SBATCH --job-name="atest"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=8
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%j
#SBATCH --export=ALL
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
                                        We go to "starting"
cd $SLURM SUBMIT DIR
                                            directory
srun -n 8 ./c_ex00
                                     Run this MPI program on
                                             8 cores
```

```
cd $SLURM_SUBMIT_DIR
srun -n 8 ./c_ex00 >& myout.$SLURM_JOB_ID
```

Gives the program output as it runs / with each run having a unique output file

### Shorten JOBID

- \$PBS\_JOBID is of the form
  - 45682.mio.mines.edu
- How can we shorten this to just a number?
  - sed -e 's/\..\*//'
  - strips everything past the first period

```
$ echo $PBS_JOBID
201665.mio.mines.edu
$ MY_JOBID=`echo $PBS_JOBID | sed -e 's/\..*//'`
$ echo $MY_JOBID
201665
```

```
#!/bin/bash
#SBATCH --job-name="atest"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=8
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
                                    The output
#SBATCH -e stderr.%j
                                   from the script
#SBATCH --export=ALL
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
JOBID=`echo $SLURM JOB ID`
cd $SLURM SUBMIT DIR
srun -n 8 ./c_ex00 > my_out.$JOBID
              myout.#####
```



### We want...

- To keep records of what scripts we have run
- To be notified when a script runs
  - Under PBS
    - #PBS -M with -abe
    - Will send a notice at start and stop
  - Under Slurm
    - #SBATCH —mail-type=ALL
    - Produces information emails
  - We want more than the job number
  - We want everything

### Records Notifications

```
#!/bin/bash -x
#SBATCH --job-name="hybrid"
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=8
#SBATCH --ntasks=16
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%j
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
# Go to the directoy from which our job was launched
cd $SLURM_SUBMIT_DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
# Save a copy of our environment and script
echo $SLURM_JOB_NODELIST > nodes.$JOBID
cat $0 > script.$JOBID
printenv > env.$JOBID
#mail us the environment and other "stuff"
```

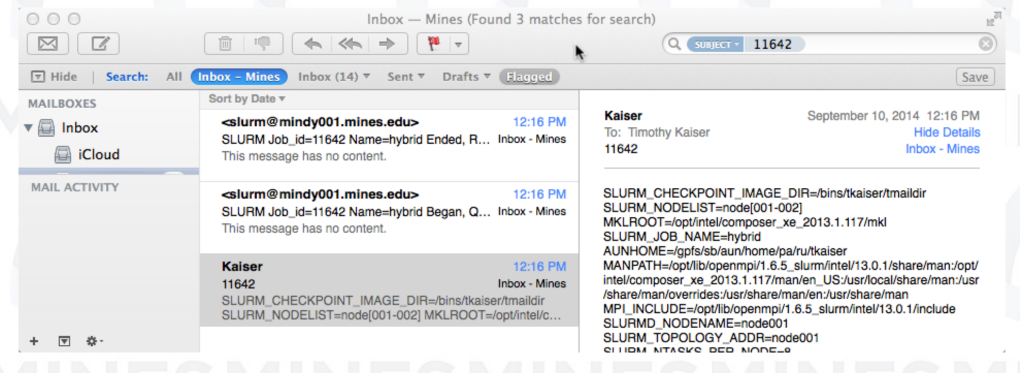
How can I record what I did and where?

How can I know when a particular script starts and exactly what is running?

```
#### mail < env.$JOBID -s $JOBID $USER@mines.edu
#ssh $SLURM_SUBMIT_HOST "mail < $MYBASE/$JOBID/env.$JOBID -s $JOBID $SLURM_JOB_USER@mines.edu"
mkdir −p ~/tmail
cp env.$JOBID ~/tmail
export MAIL HOST=$SLURM_SUBMIT_HOST
export MAIL_HOST=mindy.mines.edu
ssh mindy.mines.edu "mail < ~/tmail/env.$JOBID -s $JOBID $SLURM JOB USER@mines.edu"
```

### Lots of records...

```
[joeuser@aun002 tmaildir]$ ls -l *11642*
-rw-rw-r-- 1 joeuser joeuser 6108 Sep 10 12:16 env.11642
-rw-rw-r-- 1 joeuser joeuser 14 Sep 10 12:16 nodes.11642
-rw-rw-r-- 1 joeuser joeuser 15934 Sep 10 12:16 output.11642
-rw-rw-r-- 1 joeuser joeuser 1014 Sep 10 12:16 script.11642
-rw-rw-r-- 1 joeuser joeuser 447 Sep 10 12:16 stderr.11642
-rw-rw-r-- 1 joeuser joeuser 0 Sep 10 12:16 stdout.11642
[joeuser@aun002 tmaildir]$
```



# More on variables and a

few other details

### A digression: /opt/utility/expands

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The CSM written utility /opt/utility/expands takes a Slurm style compressed node list and creates a full list similar to what is produces by PBS

```
/opt/utility/expands node[001-003,005-007,100] | sort -u
node001
node002
node003
node005
node006
node007
node100
```

```
#!/bin/bash -x
#SBATCH -- job-name="hybrid"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH --ntasks=1
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%i
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
echo $SLURM JOB NODELIST > nodes.$JOBID
export INPUT=sinput
export APP=fillmemc
/opt/utility/expands $SLURM JOB NODELIST > $APP.$INPUT.nodes.$JOBID
```

cat \$INPUT > \$APP.\$INPUT.input.\$JOBID

srun ./\$APP < \$INPUT >> \$APP.\$INPUT.output.\$JOBID

```
#!/bin/bash -x
#SBATCH -- job-name="hybrid"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH --ntasks=1
#SBATCH --exclusive
#SBATCH --export=ALL
                                                                  Create a short list of the nodes
#SBATCH --time=00:02:00
#SBATCH -o stdout.%i
                                                                  used in my job and give it a unique
#SBATCH -e stderr.%j
                                                                  name. We recommend people
                                                                  always do this
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
                                                                 The command "expands" takes a
                                                                 short list of nodes and expands it
echo $SLURM_JOB_NODELIST > nodes.$JOBID
                                                                 to a long list. Now we put our list in
                         We are going to use variables for
export INPUT=sinput
                                                                 a file that has application name, file
export APP=fillmemc
                         both our input file and application
                                                                 name, and input file as part of the
                         names
                                                                 file name
/opt/utility/expands $SLURM_JOB_NODELIST >
                                                   $APP.$INPUT.nodes.$JOBID
cat $INPUT > $APP.$INPUT.input.$JOBID
                                                                  Save a copy of our
srun ./$APP < $INPUT >> $APP.$INPUT.output.$JOBID
                                                                  input and put the
```

output in its own file

### What we get

```
[joeuser@aun001 memory]$ ls -l *11714*
                               11 Sep 10 13:31 fillmemc.sinput.input.11714
-rw-rw-r-- 1 joeuser joeuser
-rw-rw-r-- 1 joeuser joeuser 128 Sep 10 13:31 fillmemc.sinput.nodes.11714
-rw-rw-r-- 1 joeuser joeuser 3259 Sep 10 13:31 fillmemc.sinput.output.11714
-rw-rw-r-- 1 joeuser joeuser
                                8 Sep 10 13:31 nodes.11714
                              205 Sep 10 13:31 stderr.11714
-rw-rw-r-- 1 joeuser joeuser
-rw-rw-r-- 1 joeuser joeuser
                                0 Sep 10 13:31 stdout.11714
[joeuser@aun001 memory]$
[joeuser@aun001 memory]$ cat fillmemc.sinput.nodes.11714 | sort -u
node001
[joeuser@aun001 memory]$ cat fillmemc.sinput.nodes.11714 | wc
             16
                    128
[joeuser@aun001 memory]$
[joeuser@aun001 memory]$ cat fillmemc.sinput.input.11714
4096 64 1
[joeuser@aun001 memory]$ head fillmemc.sinput.output.11714
matrix size=
                     4096
copies=
                  64
 bytes=
            8589934592 gbytes=
                                   8.000
 using mkl for inverts
generating data for run
                                   1 of
                        2.114 threads= 16
generating time=
 starting inverts
             48675,423
                         48671.805
                                         3.618
   33
             48675,423
                                         3.618
                         48671.805
   57
             48675,423
                         48671.805
                                         3.618
```

### Multiple Executables Tricks

- Case I: Multiple jobs running on the same node at the same time
  - Independent
  - Launched from different scripts
- Case 2: Multiple executables running on the same node at the same time
  - Independent
  - Launched from a single script
- Either case could be serial or MPI
- Case 3: Using mpiexec or srun to launch several serial programs

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# Case I: Multiple Scripts

Or the same script several times



```
#!/bin/bash -x
#SBATCH --job-name="hybrid"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH --ntasks=1
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%i
#SBATCH -e stderr.%j
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
echo $SLURM JOB NODELIST > nodes.$JOBID
/opt/utility/expands $SLURM_JOB_NODELIST > $APP.$INPUT.nodes.$JOBID
export INPUT=sinput
                                      Save a list of nodes, first in just a nodes.* file
export APP=fillmemc
                                     and then a file that contains the input file name
cat $INPUT > $APP.$INPUT.input.$JOBID
srun ./$APP < $INPUT >> $APP.$INPUT.output.$JOBID
```

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We have our

application name

and input file set

as a variable .

```
#!/bin/bash -x
#SBATCH --job-name="hybrid"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH --ntasks=1
#SBATCH --share
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%j
# Go to the directoy from which our job was launched
cd $SLURM_SUBMIT_DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM_JOBID`
echo $SLURM_JOB_NODELIST > nodes.$JOBID
                                           We have commented out the line that sets
#export INPUT=sinput
                                           the input file name. We can (must) specify
export APP=fillmemc
                                           the input before running the script.
/opt/utility/expands $SLURM_JOB_NODELIST >
                                              $APP.$INPUT.nodes.$JOBID
cat $INPUT > $APP.$INPUT.input.$JOBID
export OMP_NUM_THREADS=2
      ./$APP < $INPUT >> $APP.$INPUT.output.$JOBID
srun
```

inputenv

# Assume we have 4 data sets and we are willing to run on any one node..

### MINESMINESMINESM

```
[joeuser@mio test] $ export INPUT=sinput1
Which node
             [joeuser@mio test]$ qsub from env
are we using?
             267335.mio.mines.edu
 It is in the
             [joeuser@mio test] $ cat mynodes*
mynodes* file n20
                                                                             We have specified the
             [joeuser@mio test] $ export INPUT=sinput2 <
                                                                             input file here. This is
             [joeuser@mio test] $ qsub from env -1 nodes=n20
                                                                             picked up by the script
             267336.mio.mines.edu
Force the rest
             [joeuser@mio test]$
of our jobs to
             [joeuser@mio test]$ export INPUT=sinput3
 the same
              jocuser@mio test]$ qsub from env>-1 nodes=n20
   node
             267337.mio.mines.edu
             [joeuser@mio test]$
             [joeuser@mio test]$ export INPUT=sinput4
             [joeuser@mio test]$ qsub from env -1 nodes=n20
             267338 mio mines edu
             [joeuser@mio test]$
```

(If you have a reserved node you can specify it for the first run also.)

### Our output files:

```
numbers
-rw-rw-r-- 1 joeuser joeuser 3395 Feb 15 11:31 fillmemc.sinput1.267335.mio.mines.edu
-rw-rw-r-- 1 joeuser joeuser 5035 Feb 15 11:32 fillmemc.sinput2.267336.mio.mines.edu
-rw-rw-r-- 1 joeuser joeuser 6675 Feb 15 11:32 fillmemc.sinput3.267337.mio.mines.edu
-rw-rw-r-- 1 joeuser joeuser 1541 Feb 15 11:29 fillmemc.sinput4.267334.mio.mines.edu
-rw-rw-r-- 1 joeuser joeuser 1755 Feb 15 11:31 fillmemc.sinput4.267338.mio.mines.edu
[joeuser@mio test]$
```

The input file name becomes part of the output file name

Note different job

Our execution line

[joeuser@mio test]\$ ls -1 fillmemc.sinput\*

< \$INPUT >> \$APP.\$INPUT.\$PBS JOBID ./\$APP

# Assume we have 4 data sets and we are willing to run on any one node..

```
[joeuser@aun001 memory]$ export INPUT=sinput1
            [joeuser@aun001 memory]$ sbatch env2
Which node
            Submitted batch job 11834
are we using?

√[joeuser@aun001 memory]$ cat nodes.11834

 It is in the
            node001
                                                                         We have specified the
 nodes* file
                                                                         input file here. This is
            [joeuser@aun001 memory] $ export INPUT=sinput2
                                                                         picked up by the script
            [joeuser@aun001 memory] $ sbatch --nodelist=node001 env2
Force the rest
            Submitted batch job 11835
of our jobs to
            [joeuser@aun001 memory]$ export INPUT=sinput3
 the same
             joeuser@aun001 memory]$ sbate --nodelist=node001 env2
   node
            Submitted batch job 11836
            [joeuser@aun001 memory] $ sbatch -- nodelist=node001 env2
            Submitted batch job 11837
```

(If you have a reserved node you can specify it for the first run also.)

### Our output files:

```
[joeuser@aun001 memory]$ squeue -u joeuser
             JOBID PARTITION
                                  NAME
                                           USER ST
                                                          TIME
                                                                NODES NODELIST (REASON)
             11836
                                hybrid
                                        joeuser R
                                                          0:10
                                                                    1 node001
                       debug
             11837
                       debug
                               hybrid
                                        joeuser R
                                                          0:10
                                                                    1 node001
             11835
                       debug
                                hybrid
                                        joeuser R
                                                          0:24
                                                                    1 node001
```

joeuser R

[joeuser@mio test]\$ ls -l fillmemc.sinput\*

debug

11834

Note different job numbers

1 node001

```
-rw-rw-r-- 1 joeuser joeuser 2763 Sep 10 20:35 fillmemc.sinput3.output.11836
-rw-rw-r-- 1 joeuser joeuser 2716 Sep 10 20:35 fillmemc.sinput4.output.11837
-rw-rw-r-- 1 joeuser joeuser 2481 Sep 10 20:35 fillmemc.sinput2.output.11835
-rw-rw-r-- 1 joeuser joeuser 2481 Sep 10 20:34 fillmemc.sinput1.output.11834
[joeuser@mio test]$
```

hybrid

The input file name becomes part of the output file name

1:24

Our execution line

./\$APP < \$INPUT >> \$APP.\$INPUT.\$PBS\_JOBID

# Case 2: Multiple Executables Same Script

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```
#!/bin/bash -x
#SBATCH -- job-name="hybrid"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH --ntasks=1
#SBATCH --share
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%j
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
echo $SLURM JOB NODELIST > nodes.$JOBID
export APP=fillmemc
                                                We launch the application
                                                 over a list of input files
export OMP NUM THREADS=2
for INPOT in sinput1 sinput2 sinput3 sinput4; do
        ./$APP $INPUT >> $APP.$INPUT.output.$JOBID &
  srun
done
wait
          The wait command "holds" the node until
                                                Forces the job into the background so we
             all of your applications are done
                                                         can launch the next
```

multiwait

We have our

application name

and input file set

as a variable

### Our output files:

Note common job numbers with different input files

```
[joeuser@aun001 memory]$ ls -lt fillmemc* head -4
-rw-rw-r-- 1 joeuser joeuser 695 Sep 10 20:52 fillmemc.sinput3.output.11839
-rw-rw-r-- 1 joeuser joeuser 695 Sep 10 20:52 fillmemc.sinput1.output.11839
-rw-rw-r-- 1 joeuser joeuser 695 Sep 10 20:52 fillmemc.sinput2.output.11839
-rw-rw-r-- 1 joeuser joeuser 695 Sep 10 20:52 fillmemc.sinput4.output.11839
```

srun ./\$APP < \$INPUT >> \$APP.\$INPUT.\$JOB &

### mpiexec/srun and serial applications

- Some versions of mpiexec and srun will work with nonMPI programs
- Creates specified number of copies of the program, all independent

### Our Batch file, batch I

```
#!/bin/bash -x
#SBATCH -- job-name="hybrid"
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=8
#SBATCH --ntasks=16
#SBATCH --share
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH -o stdout.%j
#SBATCH -e stderr.%j
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
                                               info_p is a python program that
echo $SLURM JOB NODELIST > nodes.$JOBID
                                               creates a file based on node name
export APP=info.py
                                               and process id
        ./$APP
  srun
wait
```

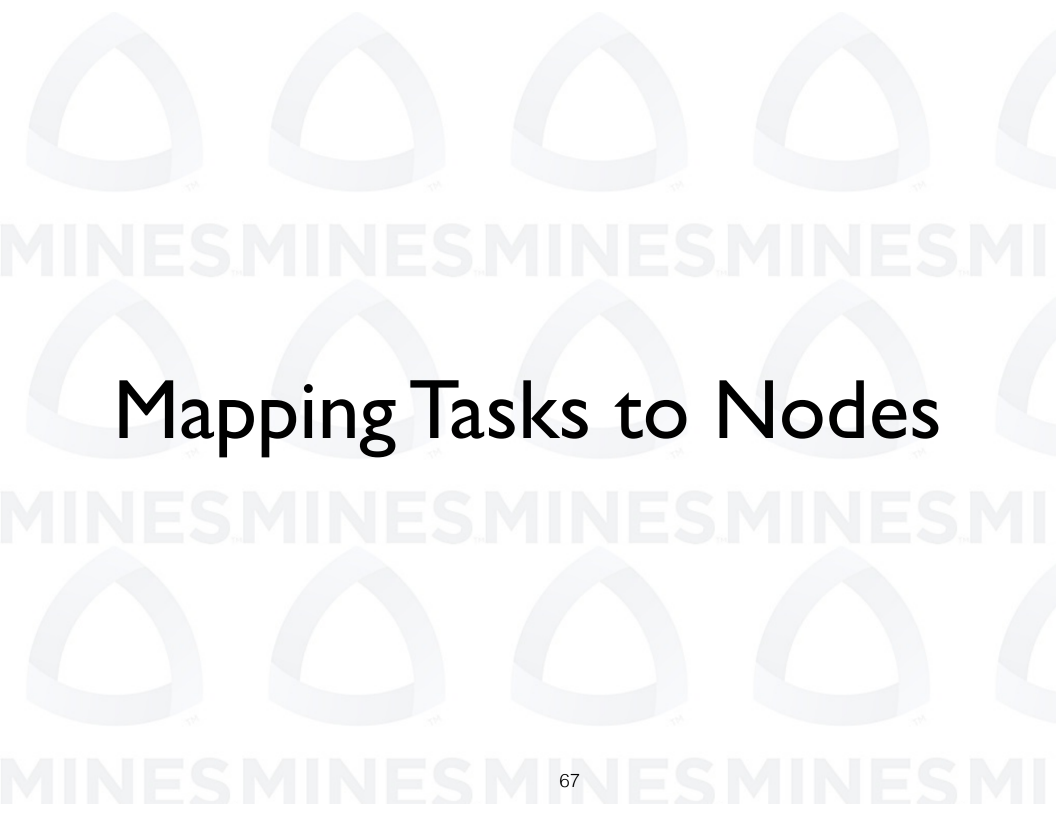
# Running a serial program with mpiexec

export MYPROGRAM=info\_p

[joeuser@aun001 memory]\$ sbatch -p debug serial Submitted batch job 11841

```
[joeuser@aun001 memory]$ cat nodes.11841
node[001-002]
[joeuser@aun001 memory]$ ls -lt node00*
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007602
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007604
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007607
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007606
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007605
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007608
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007609
-rw-rw-r-- 1 joeuser joeuser 39 Sep 10 21:05 node002_00007603
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026256
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026257
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026253
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026255
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026254
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026259
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026260
-rw-rw-r-- 1 joeuser joeuser 40 Sep 10 21:05 node001_00026258
```

[joeuser@aun001 memory]\$ cat node001\_00026256
Python says hello from 26256 on node001



### Need better than default mappings...

- Want to use less than all of the nodes on a node
  - Large memory/task
  - Hybrid MPI/OpenMPI
- Different executables on various cores (MPMD)
- Heterogeneous environment with different core counts

# Method for OpenMPI and MVAPICH2

- Same method works for both versions of MPI
- Create a description of your job on the fly from inside your script
- The description is a mapping of programs to cores
- Tell mpiexec/mpirun to use your description to launch your job
- We created a utility script to make it easy

### Alternate syntax for mpiexec

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- Normally you specify the number of MPI tasks on the mpiexec line
- The alternate syntax is to provide an "appfile"
  - mpiexec -app appfile
  - The appfile is a mapping of executables to nodes

Note: The normal "parallel run" command under slrum is srun. CSM machines AuN and Mio also support mpiexec under slurm. The method discussed here will work on these machines. However, there is also a slurm specific method for doing mappings that will be discussed below. It splits the node list and the application list into two separate files. An easy way to use the slurm specific method is to first create the appfile as discussed here. There is a CSM written utility to split the appfile into the two separate files.

### Appfile format

- Collection of lines of the form
  - -host <host name> -np <number of copies to run on host><program name>
- Specify different application names in your appfile for MPMD
- You can specify a node or program more than once

### Examples

#### These two are equivalent

#### **Appfile Example 1**

```
-host compute-1-1 -np 1 myprogram
```

### **Appfile Example 2**

```
-host compute-1-1 -np 4 myprogram
```

Note: You should specify the full path to your program

#### These two are not equivalent

### **Appfile Example 3**

```
-host compute-1-1 -np 2 aya.out
-host compute-2-3 -np 2 bee.out
```

### **Appfile Example 3**

```
-host compute-1-1 -np 1 aya.out
-host compute-2-3 -np 1 aya.out
-host compute-1-1 -np 2 bee.out
-host compute-2-3 -np 2 bee.out
```

# Difficulty and Solution

# **MINES MINES MINES M**

- Problem:
  - Names of the nodes that you are using are not known until after the job is submitted
  - You need to create the appfile on the fly from within your PBS script

# Difficulty and Solution

- Solution:
  - Under PBS the variable \$PBS\_NODEFILE contains the name of a file that has the list of nodes on which your job will run.
  - Under Slrum the variable SLURM\_JOB\_NODELIST has a compressed list of nodes.
  - We have created a script "match" which takes a list of nodes and a list of applications to run on those nodes and creates an applie
  - Located on Mio and AuN at /opt/utility/match

### Solution

Given your \$PBS\_NODEFILE and a list of programs in a file app\_list the simplest usage of match is:

match \$PBS\_NODEFILE app\_list > appfile
mpiexec --app appfile

For myapich2 replace -- app with -- configfile

### Match notes

- Number of applications that get launched
  - Is equal to the length of the longer of the two lists, the node file list, or the application list
  - If the lists are not the same length then multiple copies will be launched
  - Match also takes an optional replication count, the number copies of an application to run on a node
- Feel free to copy and modify the match script for your own needs

# Examples

```
#get a copy of all of our nodes, each node will be
#listed 8 times
cat $PBS_NODEFILE > fulllist
```

#save a nicely sorted short list of nodes, each node only
#listed one time
sort-u \$PBS NODEFILE > shortlist

#### fulllist.

compute-8-15.local compute-8-15.local compute-8-15.local compute-8-15.local compute-8-15.local compute-8-15.local compute-8-15.local compute-8-15.local compute-8-13.local compute-8-13.local

compute-8-13.local

### shortlist

compute-8-15.local compute-8-13.local

/lustre/home/apps/utility/nsort \$PBS\_NODEFILE
Also works to give a sorted list

# Examples

- We have two programs we are going to play with f\_ex00 and c\_ex00
- We have two program lists that we are going to use
  - oneprogram
    - c\_ex00
  - twoprograms
    - c\_ex00
    - f\_ex00

# match fulllist twoprograms > appfile l

```
-host
       compute-8-15.local
                                  1
                                     c ex00
                             -np
-host
       compute-8-15.local
                                  1
                                      f ex00
                             -np
-host
       compute-8-15.local
                                     c ex00
                             -np
                                  1
       compute-8-15.local
                                  1
                                     f ex00
-host
                             -np
-host
       compute-8-15.local
                             -np
                                  1
                                     c ex00
-host
       compute-8-15.local
                                  1
                                      f ex00
                             -np
       compute-8-15.local
                                     c ex00
-host
                                  1
                             -np
       compute-8-15.local
                                      f ex00
-host
                                  1
                             -np
-host
       compute-8-13.local
                                  1
                                     c ex00
                             -np
-host
       compute-8-13.local
                                      f ex00
                             -np
                                  1
                                     c ex00
-host
       compute-8-13.local
                             -np
                                  1
       compute-8-13.local
                                     f ex00
-host
                                  1
                             -np
-host
       compute-8-13.local
                             -np
                                     c ex00
-host
       compute-8-13.local
                                  1
                                      f ex00
                             -np
-host
       compute-8-13.local
                                  1
                                     c ex00
                             -np
-host
       compute-8-13.local
                                  1
                                      f ex00
                             -np
```

# match shortlist twoprograms > appfile2

# **MINESMINESMINESMI**

```
-host compute-8-13.local -np 1 c_ex00
-host compute-8-15.local -np 1 f_ex00
```

# match shortlist twoprograms 2 > appfile3

# **MINESMINESMINESMI**

```
-host compute-8-13.local -np 2 c_ex00
-host compute-8-15.local -np 2 f_ex00
```

# match shortlist oneprogram 2 > appfile4

# MINES, MINES, MINES, MINES, M

```
-host compute-8-13.local -np 2 c_ex00
-host compute-8-15.local -np 2 c_ex00
```

This will be useful for hybrid MPI OpenMP

### Can take names from command line

match <node list file> -p"list of programs" [<number of copies per node>]

Run I copy of c\_ex01 on the first node in shortlist and 8 copies of f\_ex01 on the second node

If you don't specify the number of copies then do I per core

# Running on Heterogeneous Nodes

- Mixed numbers of cores (8,12, 16)
- Want to use all of the cores
- The number of cores expected on a node is ppn=N
- Could use match with a fixed core count but this might leave some open or over subscribed

```
match shortlist -p"c_ex01" 8 8
```

 If you don't specify the number of copies then you will be given 1 per core

```
match shortlist -p"c_ex01"
```

# Slurm Specific mapping

- Slurm has several ways to specify application to node mapping
- mpiexec works on Mio and AuN
- Another way:
  - Node list and application list go in separate files, say hostlist and applist
  - To Run on 12 cores:

```
export SLURM_HOSTFILE=hostlist
srun -n12 --multi-prog applist
```

# Slurm Specific mapping

The hostlist and app list are of the form:

```
HOSTLIST:APPLIST:node0010 hellocnode0021 hellofnode0022 hellofnode0023 hellof
```

- We have the issues creating these files as we do for the mpiexec appfile
- We have a script that converts a mpiexec app file to these separate files

# match\_split

### Usage:

/opt/utility/match\_split [MATCHFILE applist hostlist]

A post processing script for the CSM utility match.

It takes the mpiexec "appfile" output from match and creates srun style application list and hostfile list files for use with the srun option "--multi-prog".

You can pipe "|" match into this program in which case /opt/utility/match\_split will create the files applist and hostlist.

Of you can specify the file created by match on the command line in which case the files created will be of the form MATCHFILE applist and MATCHFILE hostlist.

Finally, you can specify all three files on the command line

/opt/utility/match split MATCHFILE applist hostlist.

To run a slurm job using these files you do two things:

export SLURM\_HOSTFILE=hostlist
srun -n12 --multi-prog applist

where -n12 specifies the total number of MPI tasks to start.

# match\_split

### Examples:

```
[joeuser@aun001 mpi]$ cat matchfile
-host node001 -np 1 helloc
-host node002 -np 3 hellof
[joeuser@aun001 mpi]$
[joeuser@aun001 mpi]$ /opt/utility/match split matchfile
[joeuser@aun001 mpi]$ cat matchfile applist
0 helloc
1 hellof
2 hellof
3 hellof
[joeuser@aun001 mpi]$ cat matchfile_hostlist
node001
node002
node002
node002
export SLURM HOSTFILE=matchfile hostlist
srun -n4 --multi-prog matchfile applist
```

# match\_split

```
[joeuser@aun001 mpi]$ match shortlist -p"c01 f01" 3 2 | /opt/utility/match_split
[joeuser@aun001 mpi]$
[joeuser@aun001 mpi]$ cat applist
0 c01
1 c01
2 c01
3 f01
4 f01
[joeuser@aun001 mpi]$ cat hostlist
node001
node001
node001
node002
node002
export SLURM HOSTFILE=hostlist
srun -n5 --multi-prog applist
```

# Slurm script for match and match\_split

```
#!/bin/bash -x
#SBATCH -- job-name="match"
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=16
#SBATCH --ntasks=32
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH -time=00:02:00
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
echo $SLURM JOB NODELIST > nodes.$JOBID
#create a shortlist of nodes
/opt/utility/expands $SLURM JOB NODELIST |
                                           sort -u > shortlist
#run match to create a mpiexec appfile
/opt/utility/match shortlist -p"helloc hellof" 4 8 > appfile
#run the job using mpiexec
mpiexec --app appfile > outone.$JOBID
#run match split to create a srun applist and hostlist
/opt/utility/match split appfile applist hostlist
#run the job using srun
export SLURM HOSTFILE=hostlist
srun -n12 --multi-prog applist > out2.$JOBID
```

First run we use match to create and appfile and run using mpiexec

Then we create separate app list and hostlist files and run using srun split

# [joeuser@aun001 mpi]\$ cat appfile

```
-host
     node001
                         helloc
                 -np
                      4
-host node002
                 -np
                         hellof
[joeuser@aun001 mpi]$ cat applist
                                              [joeuser@aun001 mpi]$ cat hostlist
0 helloc
                                             node001
1 helloc
                                             node001
2 helloc
                                             node001
3 helloc
                                             node001
4 hellof
                                             node002
5 hellof
                                             node002
6 hellof
                                             node002
7 hellof
                                             node002
8 hellof
                                             node002
9 hellof
                                             node002
10 hellof
                                             node002
11 hellof
                                             node002
```

```
[joeuser@aun001 mpi]$ cat out2.11895
                                         sort
C-> Hello from node001 # 0 of 12
C-> Hello from node001 # 1 of 12
C-> Hello from node001 # 2 of 12
C-> Hello from node001 # 3 of 12
F-> Hello from
                             node002 #
                                           10 of
                                                    12
F-> Hello from
                             node002 #
                                           11 of
                                                    12
F-> Hello from
                             node002 #
                                            4 of
                                                    12
F-> Hello from
                             node002 #
                                            5 of
                                                    12
F-> Hello from
                             node002 #
                                            6 of
                                                    12
F-> Hello from
                             node002 #
                                                    12
                                            7 of
F-> Hello from
                             node002 #
                                              of
                                                    12
F-> Hello from
                                            9 of
                                                    12
                             node002 #
                      91
```

# Creating Directories on the fly Using Local Disk

## A Directory for Each Run

```
#!/bin/bash -x
#SBATCH -- job-name="hybrid"
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=8
#SBATCH --ntasks=16
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
                                                                 We could create NEW DIR based on a
# Create a "base name" for a directory
                                                                time stamp, the year, month, day, hour,
# in which our job will run
# For production runs this should be in $SCRATCH
                                                                               minute, and second.
MYBASE=$SLURM SUBMIT DIR
#MYBASE=$SCRATCH/mc2 tests
# We could create a directoy for our run based on the date/time
export NEW DIR=`date +%y%m%d%H%M%S`
                                                                                            or
# But here we create a directoy for our run based on the $JOBID and go there
mkdir -p $MYBASE/$JOBID
cd $MYBASE/$JOBID
odir=`pwd`
                                                                                 Based on JOBID
export ODIR=`pwd`
                                                       "Copy" our data to
# Create a link back to our starting directory
ln -s $SLURM SUBMIT DIR submit
                                                       the new directory
cp $SLURM SUBMIT DIR/data.tgz .
                                                       from our staring
tar -xzf data.tgz
                                                       directory
```

# Local Disk Space

- Most parallel machines have some disk space that is local
  - Can only be seen by tasks on the same nodes
  - Can't be seen from the primary compute node
  - Might be faster than shared space
  - Size? Location?
    - Usually a bad idea to use /tmp
    - On "Rocks" it's /state/partition I
- Usage is up to local policy

# Using Local disk

- Figure out where it is
- Create directories
- Copy to the new directory
- Compute
- Copy what you want to shared storage
- Clean up after yourself

### Here it is...

- Figure out where local disk is
- Create a shared directory where all of the results will be copied in the end
- Get a list of nodes
- Use ssh to create a directory on each node
- Do all the "normal" saves done in other examples
- Go to the new directory (This only happens on master node.)
- Use "match" to create an appfile
- Run the application (All tasks will get launched in the same named directory)
- Use scp to copy the files to shared storage
- Clean up

```
#!/bin/bash -x
                                      Set Up
#SBATCH --job-name="hybrid"
#SBATCH --nodes=4
#SBATCH --ntasks-per-node=4
#SBATCH --ntasks=16
#SBATCH --exclusive
#SBATCH --export=ALL
#SBATCH --time=00:02:00
#SBATCH --mail-type=ALL
#SBATCH --mail-user=joeuser@mines.edu
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short |OBID base on the one provided by the scheduler
|OBID=`echo $SLURM |OBID`
# Create a "base name" for a directory
# in which our job will run
# For production runs this should be in $SCRATCH
MYBASE=$SLURM SUBMIT DIR
#MYBASE=$SCRATCH/mc2 tests
#We could create a directoy for our run based on the date/time
export NEW DIR='date +%y%m%d%H%M%S'
# But here we create a directoy for our run based on the $JOBID and go there
mkdir -p $MYBASE/$|OBID
cd $MYBASE/$IOBID
odir='pwd'
export ODIR='pwd'
```

local

# Set Up 2

```
In -s $SLURM SUBMIT DIR submit
cp $SLURM SUBMIT DIR/data.tgz.
tar -xzf data.tgz
if [ -n "$JOBTMP" ]; then
 echo using $JOBTMP from environment
else
 export JOBTMP=~/scratch/local_sim
fi
#get a list of nodes...
export nlist=\'/opt/utility/expands\'
# For each node...
for i in $nlist
do
# Create my temporary directory in /scratch on each node
 ssh $i mkdir -p $JOBTMP/$NEW DIR
# Copy my data
 echo $USER@$i:$JOBTMP/$NEW_DIR
 scp * $USER@$i:$JOBTMP/$NEW_DIR
done
# save a copy of our nodes
```

/opt/utility/expands > nlist.\$JOBID

# Create a link back to our starting directory

### Run

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export APP=\$SLURM\_SUBMIT\_DIR/sinkfile match \$ODIR/flist.\$JOBID -p"\$APP" > appfile mpiexec -app appfile >& screen.\$JOBID

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# Clean Up

# Remove the temporary directory ssh \$i rm -r \$JOBTMP/\$NEW\_DIR done



# Running jobs in sequence

- In theory a batch script can submit another script
- One script creates a second then runs it
- Most systems don't support submission from compute nodes
- Must run from primary compute node
  - ssh mio.mines.edu "cd rundir; qsub next\_run"
- In most cases it is better to use the batch dependency option

# Depend section of the srun man page

### -d, --dependency=<dependency list>

Defer the start of this job until the specified dependencies have been satisfied completed. < dependency list > is of the form < type:job id[:job id][,type:job id[:job\_id]] >. Many jobs can share the same dependency and these jobs may even belong to different users. The value may be changed after job submission using the scontrol command.

#### after:job\_id[:jobid...]

This job can begin execution after the specified jobs have begun execution.

#### afterany:job\_id[:jobid...]

This job can begin execution after the specified jobs have terminated.

### afternotok:job\_id[:jobid...]

This job can begin execution after the specified jobs have terminated in some failed state (non-zero exit code, node failure, timed out, etc).

#### afterok:job id[:jobid...]

This job can begin execution after the specified jobs have successfully executed (ran to completion with an exit code of zero).

#### expand:job\_id

Resources allocated to this job should be used to expand the specified job. The job to expand must share the same QOS (Quality of Service) and partition. Gang scheduling of resources in the partition is also not supported.

#### singleton

This job can begin execution after any previously launched jobs sharing the same job name and user have terminated.

srun --dependency=after:123 /tmp/script
srun --dependency=before:234 /tmp/script

# srun - requesting specific nodes

- srun normally gives you any node
- Can select nodes based on queues
- Can select nodes base on name

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```
sbatch -p debug
sbatch -p group_name
sbatch -nodelist=node[001-006]
```

# A Digression, Not a Batch Script...

fi

This script prints
your nodes if
running in batch or
all nodes if running
on the front end
nodes

# Slurm array jobs

### Slurm allows array jobs:

Job arrays offer a mechanism for submitting and managing collections of similar jobs quickly and easily. All jobs must have the same initial options (e.g. size, time limit, etc.), however it is possible to change some of these options after the job has begun execution using the command specifying the *JobID* of the array or individual *ArrayJobID*.

Job arrays will have two additional environment variable set. **SLURM\_ARRAY\_JOB\_ID** will be set to the first job ID of the array. **SLURM\_ARRAY\_TASK\_ID** will be set to the job array index value. For example a job submission of this sort:

```
sbatch --array=1-3 -N1 some_script will generate a job array containing three jobs. If the sbatch command responds Submitted batch job 36 then the environment variables will be set as follows:
```

```
SLURM_JOBID=36
SLURM_ARRAY_JOB_ID=36
SLURM_ARRAY_TASK_ID=1
```

```
SLURM_JOBID=37
SLURM_ARRAY_JOB_ID=36
SLURM_ARRAY_TASK_ID=2
```

```
SLURM_JOBID=38
SLURM_ARRAY_JOB_ID=36
SLURM_ARRAY_TASK_ID=3
```

```
Redirection Revisited
#!/bin/bash
#PBS -1 nodes=1:ppn=8
#PBS -1 walltime=00:02:00
#PBS -N testIO
#PBS -o out.$PBS JOBID
#PBS -e err.$PBS JOBID
#PBS -V
#PBS -m abe
#PBS -M joeuser@mines.edu
cd $PBS O WORKDIR
################
# http://compgroups.net/comp.unix.shell/bash-changing-stdout/497180
# set up our redirects of stdout and stderr
                           # 1 and 2 are file descriptors for
                           # stdout and stderr
                           # 3 and 4 are descriptors to logfile
                           # we will use 3 for stdout 4 for stderr
exec 3>>logfile. date + "%y%m%d%H%M%S"
                           # anything that goes to 4 will go to 3
                           # which is our file we have created
exec 4>&3
                           # save "pointers" to stdin and stdout
exec 5>&1 6>&2
                           # redirect stdin and stdout to file
exec 1>&3 2>&4
###############
# normal commands
                           # this line goes to stdout
echo this is a test from stdout
                           # this line goes to stderr
echo this is a test from stderr >&2
                           # error message goes to stderr
ls file that does not exist
1s
mpiexec -n 8 ./c ex00 > myout.$PBS JOBID
mpiexec -n 8 ./c ex00
###############
                           # restore original stdin and stdout
exec 1>&5 2>&6
                           # close logfile descriptors
3>&- 4>&-
                           # close saved stdin and stdout
5>&- 6>&-
```

# Slurm array jobs

```
#!/bin/bash -x
#SBATCH -- job-name="array"
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
                                                      This script is similar to the last but we
#SBATCH --ntasks=1
                                                                       use
#SBATCH --share
                                                            SLURM ARRAY TASK ID
#SBATCH --export=ALL
#SBATCH --time=00:02:00
                                                                       and
#SBATCH -o stdout.%j
                                                             SLURM ARRAY JOB ID
#SBATCH -e stderr.%j
                                                          to define input and output files
# Go to the directoy from which our job was launched
cd $SLURM SUBMIT DIR
# Create a short JOBID base on the one provided by the scheduler
JOBID=`echo $SLURM JOBID`
echo $SLURM JOB NODELIST > nodes.$JOBID
export INPUT=sinput${SLURM ARRAY TASK ID}
export APP=fillmemc
/opt/utility/expands $SLURM JOB NODELIST >
                                            $APP.$INPUT.nodes.$JOBID
cat $INPUT > $APP.$INPUT.input.$JOBID
export OMP NUM THREADS=2
    ./$APP < $INPUT >> $APP.$INPUT.output.$JOBID.$SLURM ARRAY JOB ID.$SLURM ARRAY TASK ID
srun
```

# Slurm array jobs

[joeuser@aun001 memory]\$ sbatch -p debug --nodelist=node002 --array=1-4 array Submitted batch job 11968

```
[joeuser@aun001 memory]$ squeue -u joeuser
             JOBID PARTITION
                                                                NODES NODELIST (REASON)
                                  NAME
                                           USER ST
                                                          TIME
           11968 1
                               hybrid
                                                          0:05
                                                                    1 node002
                       debug
                                        joeuser
           11968 2
                       debug hybrid
                                        joeuser
                                                          0:05
                                                                    1 node002
           11968 3
                       debug
                               hybrid
                                        joeuser
                                                                    1 node002
                                                          0:05
           11968 4
                       debug
                               hybrid
                                        joeuser
                                                          0:05
                                                                    1 node002
```

Here we run 4 jobs on the same node producing:

```
[joeuser@aun001 memory]$ ls -lt | head
total 12768
-rw-rw-r-- 1 joeuser joeuser
                               1791 Sep 11 14:50 stderr.11968
-rw-rw-r-- 1 joeuser joeuser
                                803 Sep 11 14:50 stderr.11970
                                803 Sep 11 14:50 stderr.11971
-rw-rw-r-- 1 joeuser joeuser
                                803 Sep 11 14:50 stderr.11969
-rw-rw-r-- 1 joeuser joeuser
-rw-rw-r-- 1 joeuser joeuser
                               2763 Sep 11 14:50 fillmemc.sinput1.output.11968.11968.1
                               2763 Sep 11 14:50 fillmemc.sinput4.output.11971.11968.4
-rw-rw-r-- 1 joeuser joeuser
                               2763 Sep 11 14:50 fillmemc.sinput2.output.11969.11968.2
-rw-rw-r-- 1 joeuser joeuser
                               2763 Sep 11 14:50 fillmemc.sinput3.output.11970.11968.3
-rw-rw-r-- 1 joeuser joeuser
```

