

Miscellaneous MPI-IO topics

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### **MPI-IO Errors**



- Unlike the rest of MPI, MPI-IO errors are not fatal
  - probably don't want your program to crash if a file open fails
  - always need to check the error code!
- Many different error codes can be reported
  - I would suggest simply quitting if ierr != MPI\_SUCCESS
- Can change this behaviour for file operations
  - same functionality as MPI\_Errhandler\_create etc.
  - called MPI\_File\_create\_errhandler, ...
  - error handlers are attached to file handles rather than communicators
  - can set handler to be MPI\_ERRORS\_ARE\_FATAL

### Size of File on Disk

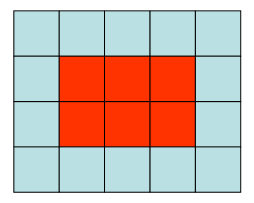


- Useful to check length of output file
  - ls -l <filename>
  - check that size (in bytes) is what you expect
- Can be confusing if file already exists
  - length will be increased if new file is longer than existing file
  - but may not be decreased if new file is shorter!
- Delete old files before running your test programs

# Datatype for MPI\_File\_read / write



- Usually pass the basic type of the array being processed
  - eg MPI\_FLOAT, MPI\_REAL
- Can pass derived types
  - useful for receiving the core of an array when local arrays have halos



```
MPI_File_read_all(fh, &x[1][1], 1, vector3x2, ...);
MPI_FILE_READ_ALL(fh, x(2,2) , 1, vector3x2, ...)
```

- or could use a 3x2 subarray and pass &x[0][0] or x(1,1)

# **General Decompositions**



- We have just considered block decompositions
  - where local array size is an exact multiple of global array size
- If the sizes don't match
  - define different sized subarrays on each process
  - eg processes at the edge of the grid have smaller subsections
- This does not generalize to block-cyclic decompositions
  - how do we specify discontinuous subarrays?

4	8	12	16
3	7	11	15
2	6	10	14
1	5	9	13

## **Distributed Arrays**



```
int MPI Type create darray(int size, int rank,
  int ndims, int array of qsizes[],
  int array_of_distribs[], int array_of_dargs[],
  int array of psizes[], int order,
  MPI Datatype oldtype, MPI Datatype *newtype);
MPI TYPE CREATE DARRAY(SIZE, RANK, NDIMS
  ARRAY OF GSIZES, ARRAY OF DISTRIBS, ARRAY_OF_DARGS,
  ARRAY OF PSIZES, ORDER, OLDTYPE, NEWTYPE, IERR)
INTEGER SIZE, RANK, NDIMS, ARRAY OF GSIZES(*),
  ARRAY OF DISTRIBS(*), ARRAY OF DARGS(*),
  ARRAY OF PSIZES(*), ORDER, OLDTYPE, NEWTYPE, IERR
```

- See the man page for full details!
  - uses HPF conventions for block-cyclic distributions

### **Unstructured Data**



### Imagine a particle simulation

- each particle is a compound object with a type and position (x,y,z)
  - eg a C struct or Fortran type
- each particle has unique global identifier 1, 2, 3, ..., N-1, N

#### Particles move around

- at the end of a simulation, each process will have:
  - a different numbers of particles
  - with a random mixture of global identifiers

#### Two choices

- write to file in the order they appear in the processes
- write to file with position based on global identifier

## Approach



- Define a derived type to match the particle object
  - eg MPI\_PARTICLE
  - use this as the etype
- Writing in process order
  - need to know where to start in the file
  - calculate the sum of the number of particles on previous ranks
    - using MPI\_Scan
- Writing in global order
  - call MPI\_Type\_indexed (or create\_indexed\_block)
  - use this as the filetype
  - write multiple instances of MPI\_PARTICLE

### **Unstructured Meshes**



- Similar to global ordering of particles
  - each element has both a local and global identifier
  - want the file to be ordered by the global id
- Define an MPI\_ELEMENT
  - use this as the etype
  - create an indexed filetype based on global id



This code spends a lot of time waiting while saving to disk

```
define big arrays: old and new
loop many times
  ! do a computationally expensive operation
  new = expensive_function(old)
  old = new
  every 10 iterations:
    save_to_disk(old)
 end loop
```

## Non-blocking IO



This code overlaps computation and IO

```
define big arrays: old and new
loop many times
  ! do a computationally expensive operation
  new = expensive_function(old)
  if (saving to disk):
    finish: isave_to_disk(old)
  old = new
  every 10 iterations:
    start: isave_to_disk(old)
 end loop
```

# Non-blocking IO in MPI-IO



#### Two forms

### General non-blocking

- MPI\_File\_iwrite(fh, buf, count, datatype, request)
- finish by waiting on request
- but no collective version

### Split collective

```
- MPI_File_write_all_begin(fh, buf, count, datatype)
```

```
- MPI_File_write_all_end(fh, buf, status)
```

- only a single outstanding IO operation at any one time
- allows for collective version

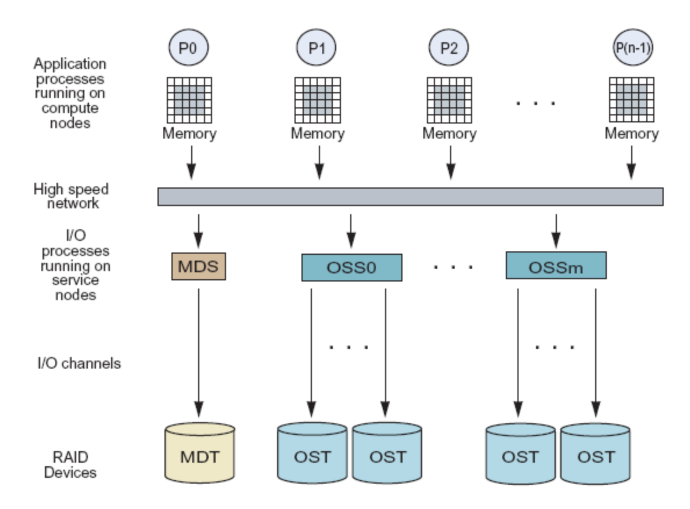
### Serial IO



- How can I read MPI-IO files in a serial program?
- Using native format
  - data is raw bytes
  - use fread in C or direct access unformatted IO in Fortran
  - see ioread.c and ioread.f90 for examples
  - Fortran approach is quite old-fashioned (direct access IO)
    - new access="stream" functionality makes this a bit simpler
- Other MPI-IO formats will require more work!
- Note that you can do single process IO in MPI-IO
  - pass MPI\_COMM\_SELF to MPI\_File\_open



Recall schematic overview of parallel file system Lustre



# Application-side parallel IO



- Implementing MPI-IO has achieved
  - all data going to a single file
  - minimal stress on Meta Data Server (MDS) a serial bottleneck
  - potential for many processes to write simultaneously
- But ...
  - performance requires multiple parallel writes to disk
  - in Lustre, requires multiple Object Storage Servers (OSS) writing to multiple Object Storage Targets (OST)
  - an OSS is like an IO server, an OST is like a physical disk
- User has control over assignment of files to OSTs
  - but default is only a few OSTs
  - MPI-IO performance not much better than naïve master IO

## **Lustre Striping**



- Can split a file across multiple OSTs
  - each block is called a "stripe"

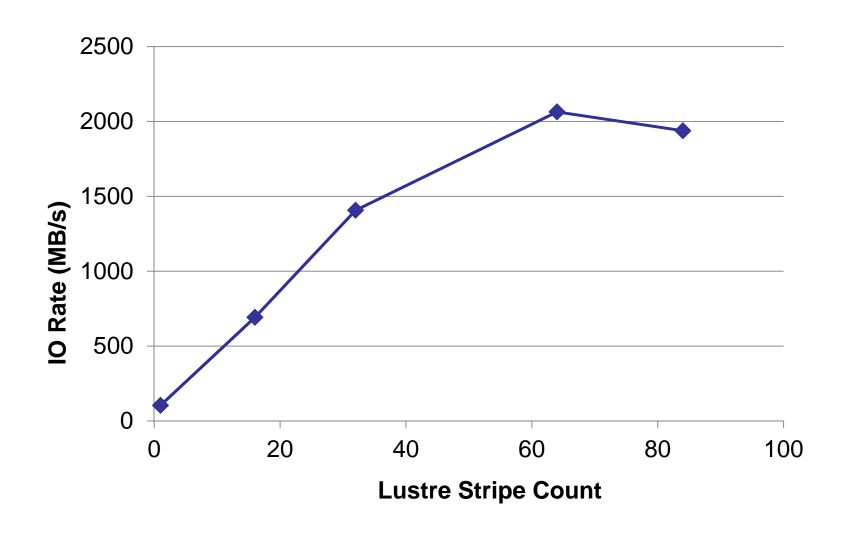
### • lfs setstripe -c 8 <filename>

- stripes across 8 OSTs
- has substantial benefits for performance

#### Test case

- 2048 x 2048 x 2048 array across 4096 processors (16 x 16 x 16)
- file size is 32 GB
- identical IO approach as used exercise
  - generalised to 3D
  - local halos automatically stripped off with derived type in MPI-IO write call



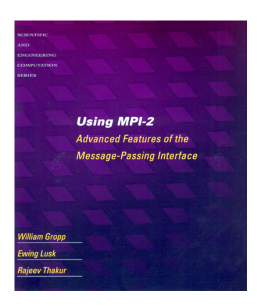


### Further information



- MPI-2 web pages
- Another tutorial
  - www.sdsc.edu/us/training/workshops/institute2005
    /docs/Thakur-MPI-IO.ppt
- MPI-2 book

http://www-unix.mcs.anl.gov/mpi/usingmpi2/



### **Further information**



ARCHER training: <a href="http://www.archer.ac.uk/training/">http://www.archer.ac.uk/training/</a>

Efficient Parallel I/O on ARCHER

- •Oxford, 10-11 Decmber 2015
- Registration is currently open

Finally:

Please fill in the feedback if you haven't already!

http://www.archer.ac.uk/training/feedback/