Parallel I/O

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Agenda

- Intro to I/O
- MPI-IO
- HDF5, NetCDF4
- Parallel HDF5/NetCDF4
- ADIOS







Data is getting bigger

- Increase in computing power makes simulations larger/more frequent
- Increase in sensor technology makes experiments/ observations larger
- Data sizes that used to be measured in MB/GB now measured in TB/PB.
- Easier to generate the data than to store it:



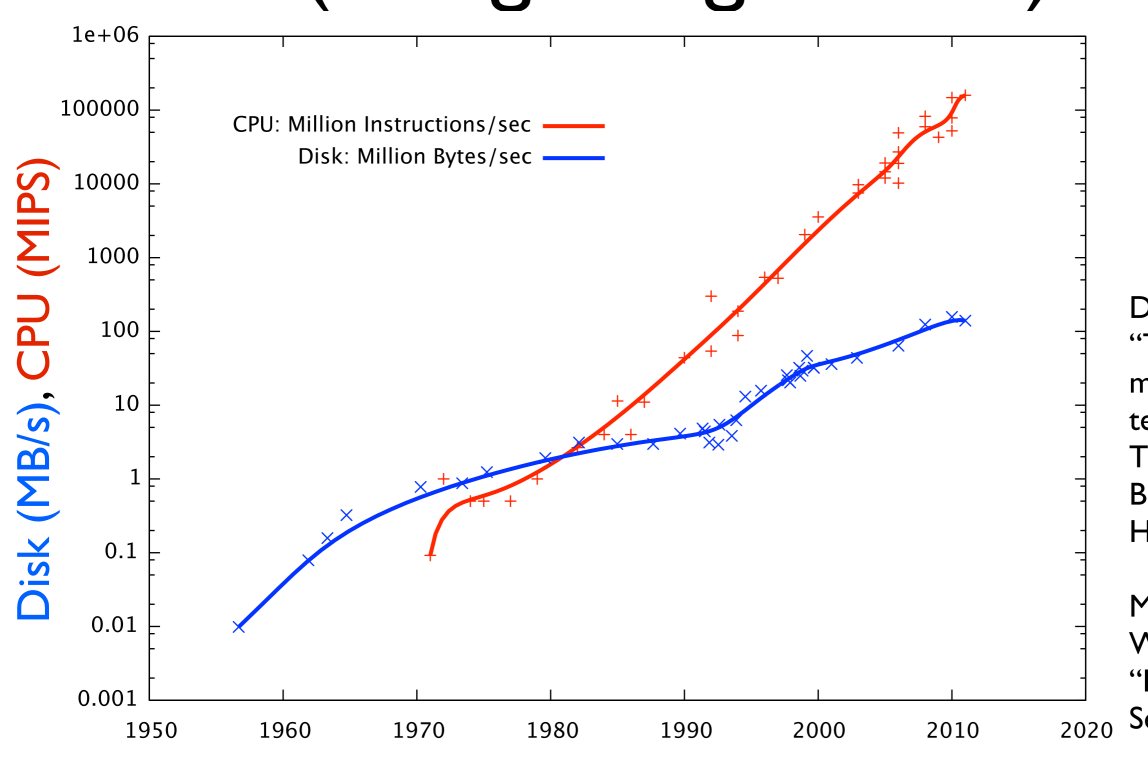
Economist, 27 Feb 2010







Disks are slower than CPUs (and getting slower)

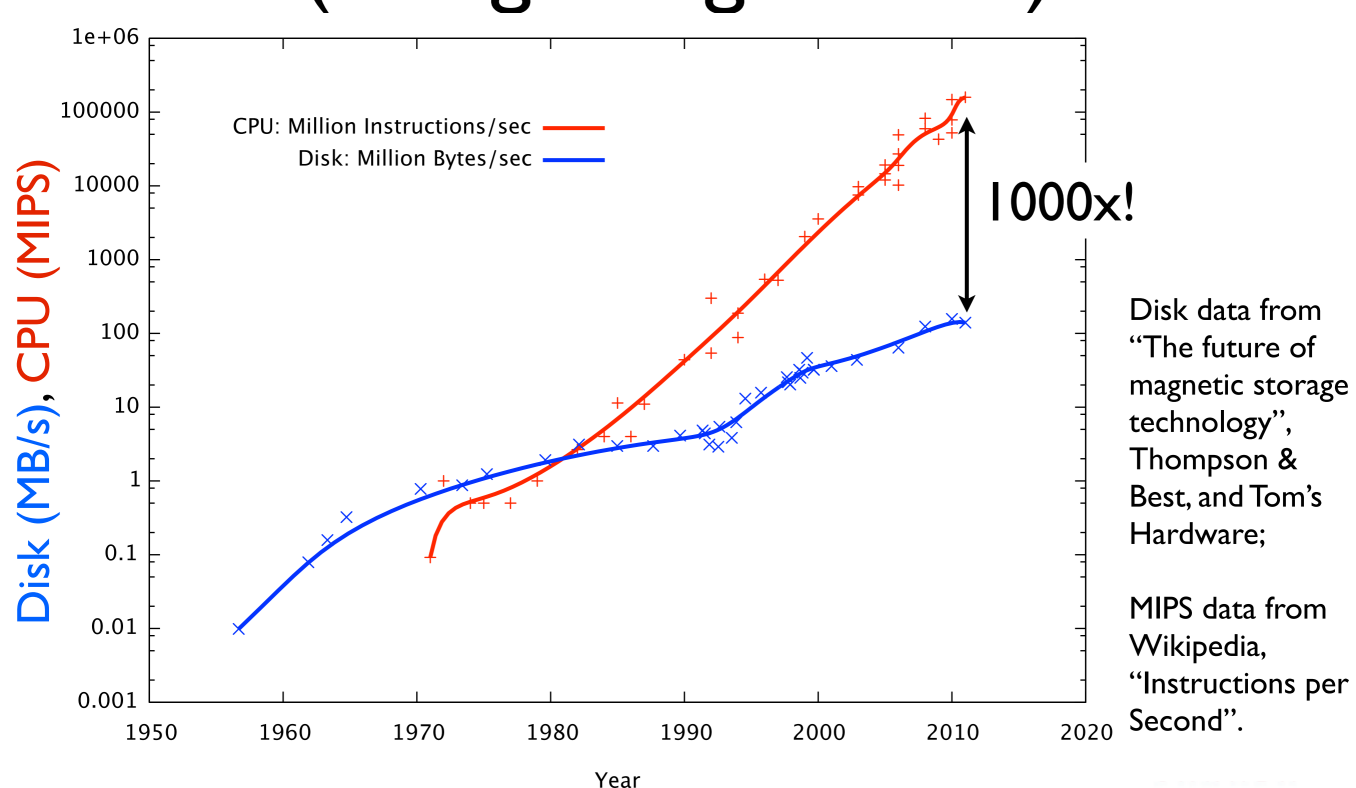


Year

Disk data from "The future of magnetic storage technology", Thompson & Best, and Tom's Hardware;

MIPS data from Wikipedia, "Instructions per Second".

Disks are slower than CPUs (and getting slower)



- Data rate: MB/s
 - Peak or sustained
 - Write faster than read
 - Network analogy: bandwidth
- IOPS: I/O Operations Per Second
 - open, close, seek, read, write
 - Network analogy: I/latency







Device	Bandwidth (MB/s)	IOPS
SATA HDD	100	100
SSD	250	10000

HD:

Open, Write, Close 1000x1kB files: 30.01s (eff: 0.033 MB/s)

Open, Write, Close Ix IMB file: 40ms (eff: 25 MB/s)







Device	Bandwidth (MB/s)	IOPS
SATA HDD	100	100
SSD	250	10000

SSD:

Open, Write, Close 1000x1kB files: 300ms (eff: 3.3 MB/s)

Open, Write, Close Ix IMB file: 4ms (eff: 232 MB/s)







- SSDs are much faster at IOPS:
 - No physical mechanisms that must move to file position
 - Read ~ as fast as write
- But still latency at controller, system calls, etc.
- SSDs will always be much more expensive than disk per unit storage - disk isn't going away.

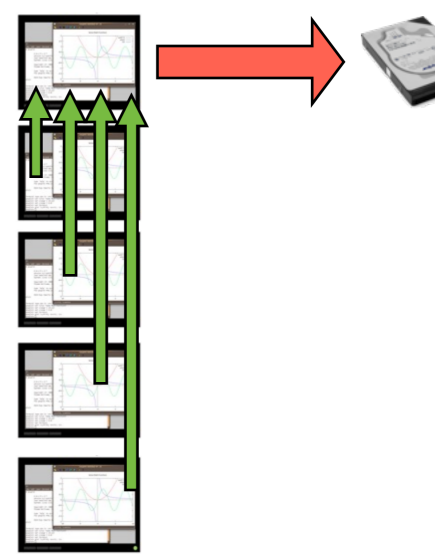






Planning your I/O

- Parallel computation, several options.
- Everyone sends data to process 0
- Process 0 writes.
- Serialize I/O huge bottleneck.



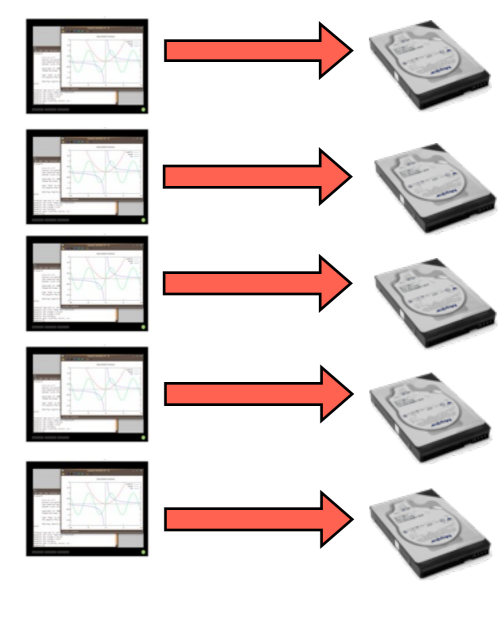






Planning your I/O

- Parallel computation, several options.
- Each process writes a file, possibly to local disk.
- Postpones the problem
 - how do you analyze,or restart withdifferent # of procs?







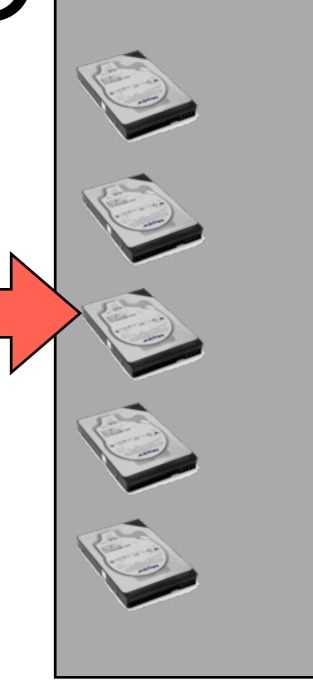


Parallel FS

Planning your I/O

- Parallel computation, several options.
- We're going to learn to avoid doing this by using Parallel I/O
- Coordinated single output of multiple processes.



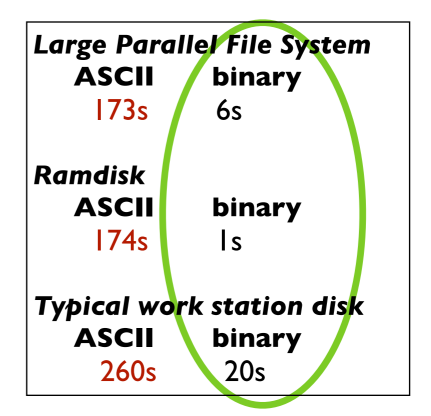








- Binary smaller files, much faster to read/write.
- You're not going to read GB/ TB of data yourself; don't bother trying.
- Write in I chunk, rather than a few #s at a time.



Timing data: writing 128M double-precision numbers







- All disk systems do best when reading/writing large, contiguous chunks
- I/O operations (IOPS) are themselves expensive
- moving around within a file
- opening/closing
- Seeks 3-15ms enough time to read 0.75 MB!

Typical work station disk binary - one large read

binary - 8k at a time 20s

binary - 8k chunks, lots of seeks | 150s

binary - seeky + open and closes 205s

Timing data: reading 128M double-precision numbers







- RAM is much better for random accesses
- Use right storage medium for the job!
- Where possible, read in contiguous large chunks, do random access in memory
- Much better if you use most of data read in



173s 6s

Ramdisk

ASCII binary

174s Is

Typical work station disk

ASCII binary

260s 20s

Ramdisk

binary - one large read

ls

binary - 8k at a time

ls

binary - 8k chunks, lots of seeks

ls

binary - seeky + open and closes

1.5s







Parallel I/O and large file systems

- Large disk systems featuring many servers, disks
- Can serve files to many clients concurrently
- Parallel File Systems -
- Lustre, Panasas, GlusterFS,
 Ceph, GPFS...



SciNet ~2k drives







SciNet's File Systems

- 2x DCS9900 couplets
- 1,790 ITB SATA disk drives
- I.4 PB of storage
- Single GPFS domain, accessed by all machines (TCS and GPC).
- Data to compute nodes via IB







SciNet's File Systems

- Designed for HPC workloads
- High bandwidth to large files big data.
- Does not do well with millions of little files:
 - wastes disk space (4MB block size)
 - lots of small scattered access is terrible for performance, even on desktop; multiply by hundreds of processors, can be disastrous







Device	Bandwidth (MB/s)	IOPS
SATA HDD	100	100
SSD	250	10000
SciNet GPFS	5000	30000

(SciNet GPFS used by ~3000 nodes.)







- Well built parallel file systems can greatly increase bandwidth
- Many pipes to network (servers), many spinning disks (bandwidth off of disks)
- But typically even worse penalties for seeky/IOPSy operations (coordinating all those disks.)
- Parallel FS can help with big data in two ways

Large Parallel File System

ASCII binary

173s 6s

Ramdisk

ASCII binary

174s Is

Typical work station disk

ASCII binary

260s 20s

Large Parallel File System binary - one large read 7.5s

binary - 8k at a time 62 s

binary - 8k chunks, lots of seeks 428 s

binary - seeky + open and closes 2137 s



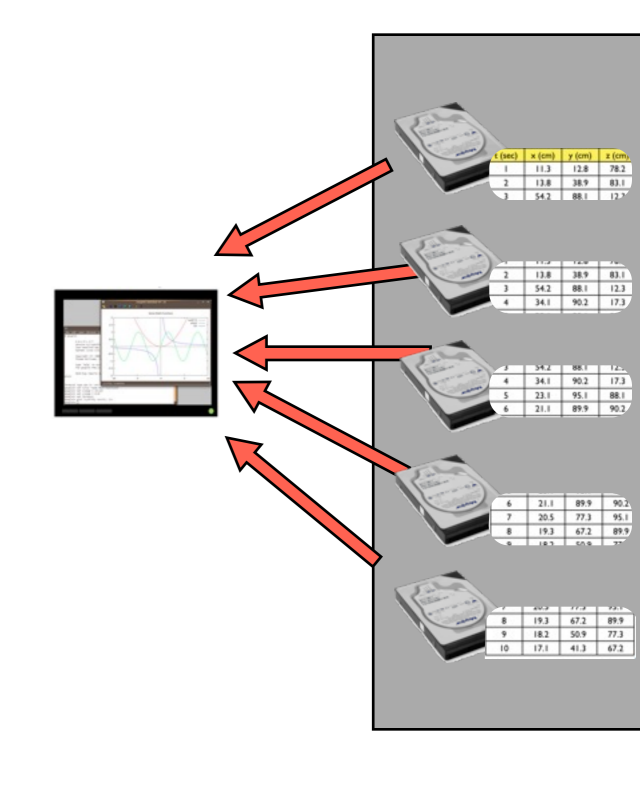




Parallel FS

Striping data across disks

- Single client can make use of multiple disk systems simultaneously
- "Stripe" file across many drives
- One drive can be finding next block while another is sending current block





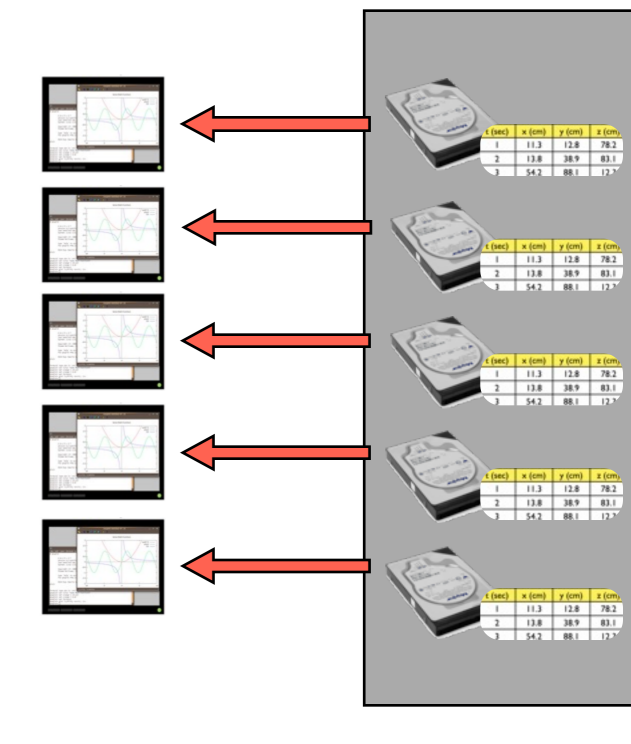




Parallel operations on separate data

- Or can do truly parallel operations
- multiple clients doing independent work
- Easy parallelism (good for lots of small data) - process many small files separately

Parallel FS





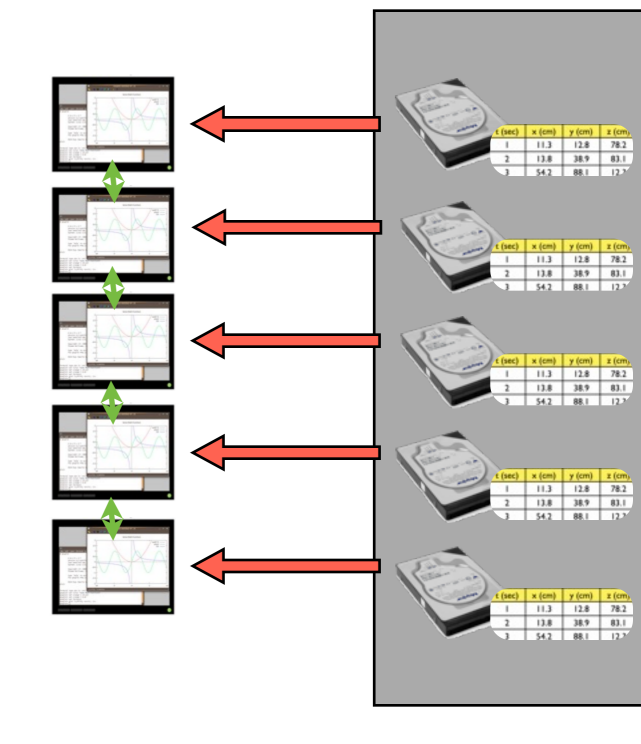




Parallel operations on separate data

- Or can do truly parallel operations
- multiple clients doing independent work
- Easy parallelism (good for lots of small data) - process many small files separately
- Harder parallelism each does part of a larger analysis job on a big file.

Parallel FS



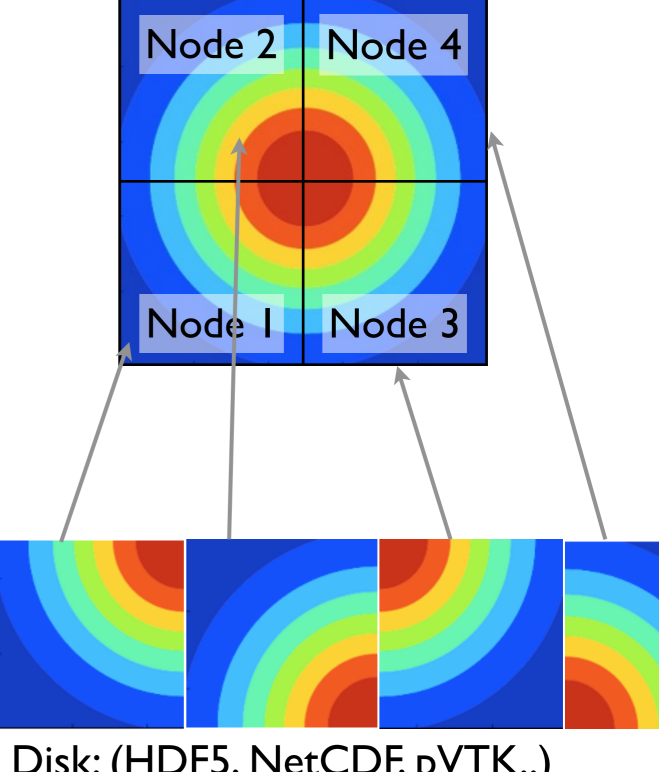






Data files must take advantage of parallel I/O

- For parallel operations on single big files, parallel filesystem isn't enough
- Data must be written in such a way that nodes can efficiently access relevant subregions
- HDF5, NetCDF formats typical examples for scientific data



Disk: (HDF5, NetCDF, pVTK..)

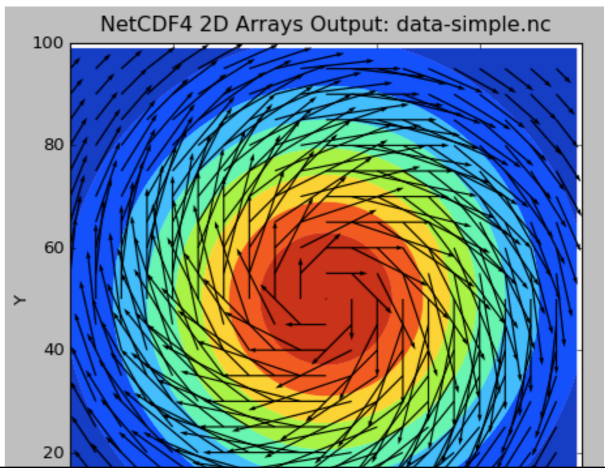






These formats are self-describing

- HDF5, NetCDF have other advantages anyway
- Binary
- Self describing contains not only data but names, descriptions of arrays, etc
- Many tools can read these formats
- Big data formats matterHPCS2012



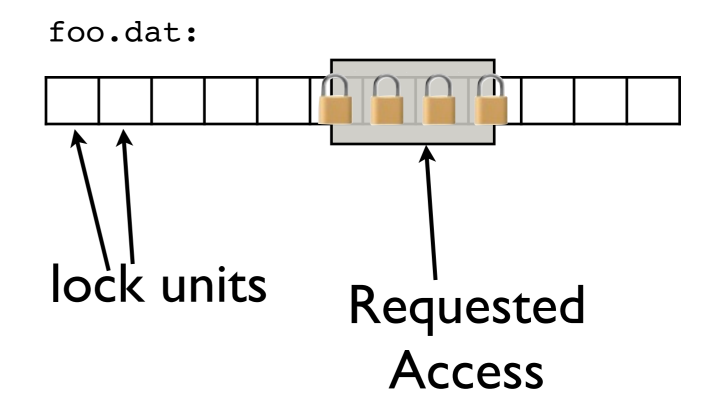
```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
  dimensions:
    X = 100 ;
    Y = 100 ;
    velocity components = 2 ;
  variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity components) ;
}
```





Coordinating I/O

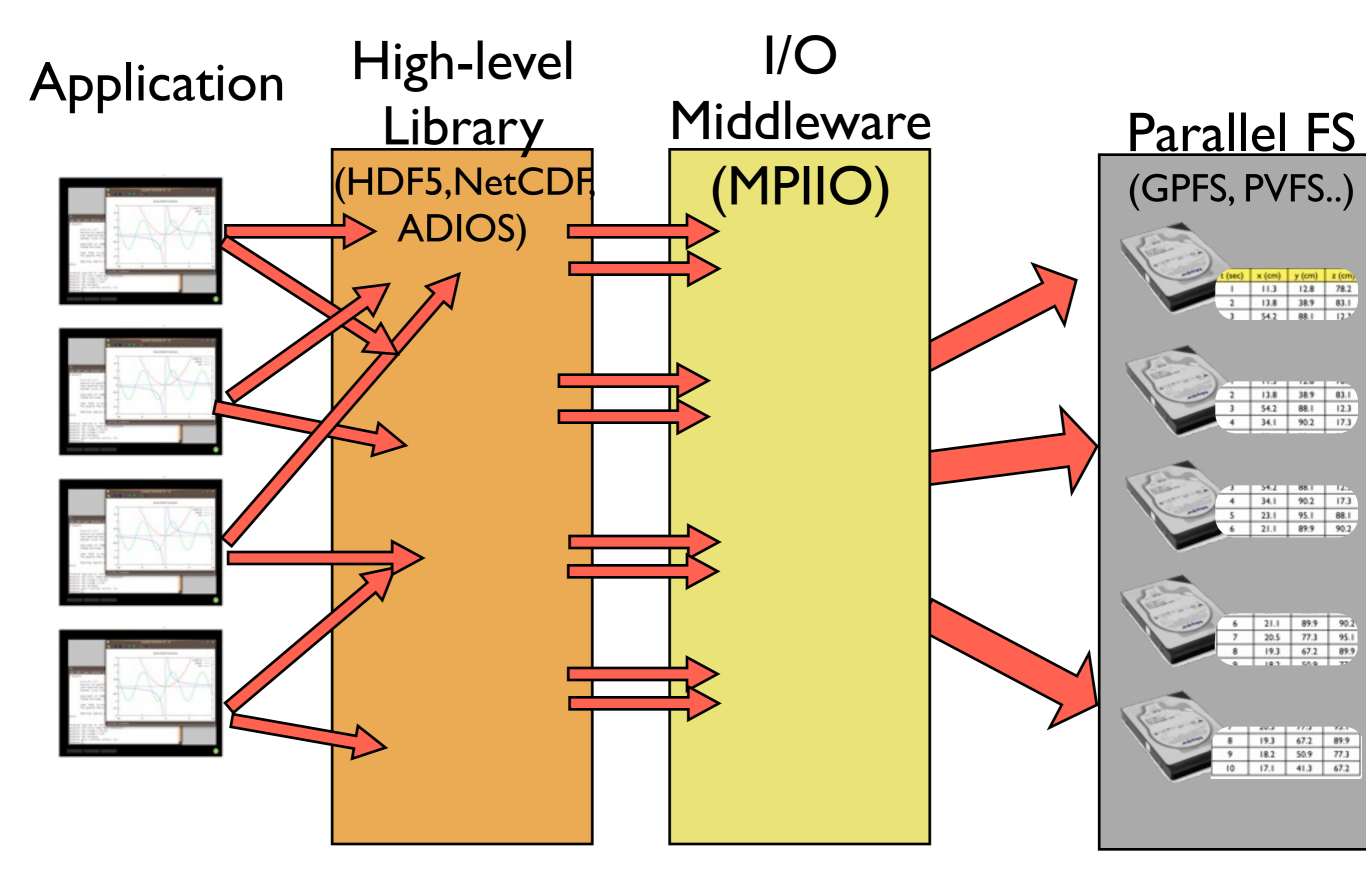
- Multiple nodes all accessing same filesystem.
- To prevent anarchy, locks for some requested accesses.
- File broken up into lock units, locks handed out upon request.
- "False sharing", etc, possible.
- Files and directories.
- Makes (some) IOPS even more expensive











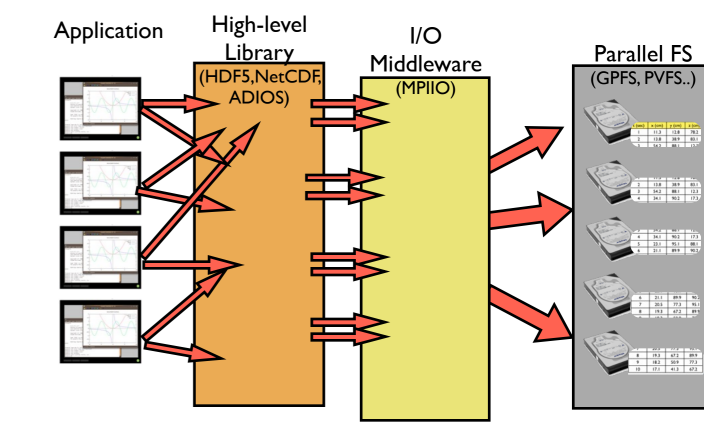






Abstraction Layers

- High Level libraries can simplify programmers tasks
 - Express IO in terms of the data structures of the code, not bytes and blocks
- I/O middleware can coordinate, improve performance
 - Data Sieving
 - 2-phase I/O



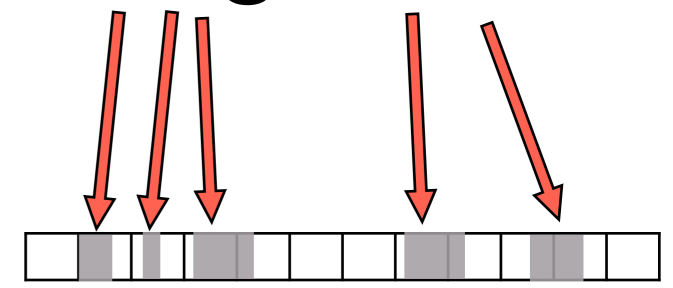


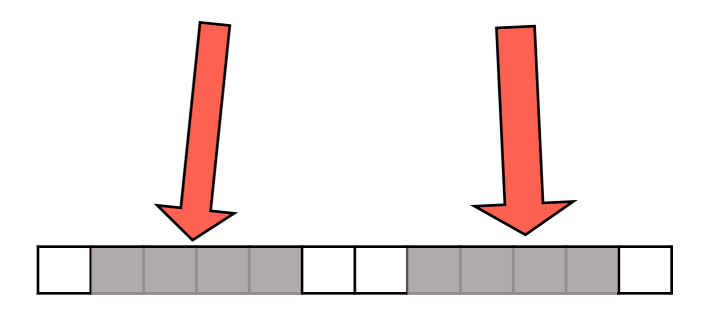




Data Sieving

- Combine many noncontiguous IO requests into fewer, bigger IO requests
- "Sieve" unwanted data out
- Reduces IOPS, makes use of high bandwidth for sequential IO





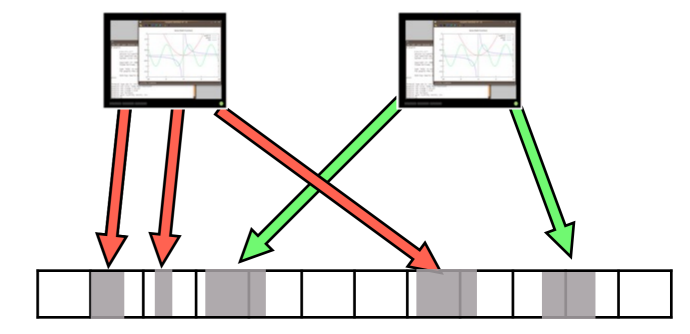


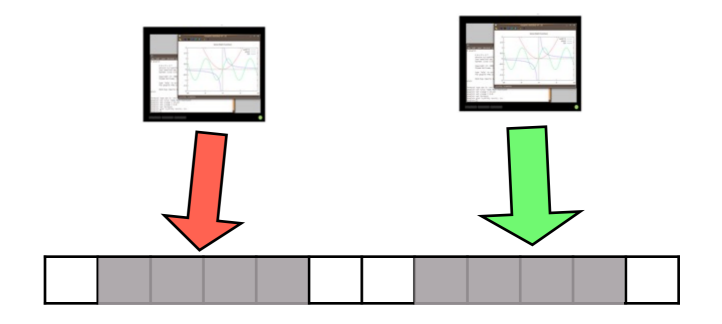


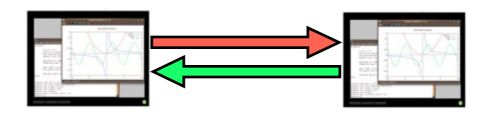


Two-Phase IO

- Collect requests into larger chunks
- Have individual nodes read big blocks
- Then use network communications to exchange pieces
- Fewer IOPS, faster IO
- Network communication usually faster













MPI-IO

- Part of MPI-2 standard
- Started at IBM Watson
- Maps I/O reads and writes to message passing
- ROMIO is the implementation found in MPICH2, OpenMPI
- Really only widelyavailable scientific computing parallel I/O middleware









```
$ cd parIO
$ source parallellibs
$ cd samples/mpiio
$ make
....

$ mpiexec -n 4 ./helloworldc
Rank 0 has message <Hello >
Rank 1 has message <World!>
Rank 2 has message <Hello >
Rank 3 has message <World!>
$ cat helloworld.txt
Hello World!Hello World! $
```







```
#include <stdio.h>
                                                            helloworldc.c
#include <string.h>
#include <mpi.h>
int main(int argc, char **argv) {
    int ierr, rank, size;
   MPI Offset offset;
   MPI File file;
   MPI Status status;
   const int msgsize=6;
    char message[msgsize+1];
    ierr = MPI Init(&argc, &argv);
    ierr = MPI Comm size(MPI COMM WORLD, &size);
    ierr |= MPI Comm rank(MPI COMM WORLD, &rank);
    if ((rank % 2) == 0) strcpy (message, "Hello "); else strcpy (message, "World!");
    offset = (msqsize*rank);
   MPI File open(MPI COMM WORLD, "helloworld.txt", MPI MODE CREATE MPI MODE WRONLY,
                  MPI INFO NULL, &file);
   MPI File seek(file, offset, MPI SEEK SET);
   MPI File write(file, message, msgsize, MPI CHAR, &status);
   MPI File close(&file);
   MPI Finalize();
   return 0;
```







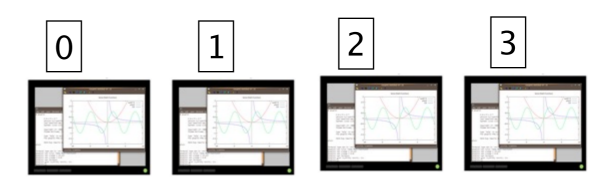
```
program MPIIO_helloworld
                                                              helloworldf.f90
    use mpi
    implicit none
    integer(mpi offset kind) :: offset
    integer, dimension(mpi_status_size) :: wstatus
    integer, parameter :: msgsize=6
    character(msgsize) :: message
    integer :: ierr, rank, comsize, fileno
    call MPI_Init(ierr)
    call MPI Comm size(MPI COMM WORLD, comsize, ierr)
    call MPI Comm rank(MPI COMM WORLD, rank, ierr)
    if (mod(rank, 2) == 0) then
       message = "Hello "
    else
       message = "World!"
    endif
    offset = rank*msqsize
    call MPI File open(MPI COMM WORLD, "helloworld.txt", ior(MPI MODE CREATE, MPI MODE WRONLY),&
                       MPI INFO NULL, fileno, ierr)
    call MPI File seek (fileno, offset, MPI SEEK SET, ierr)
    call MPI File write(fileno, message, msgsize, MPI CHARACTER, wstatus, ierr)
    call MPI_File_close(fileno, ierr)
    call MPI Finalize(ierr)
end program MPIIO helloworld
```







MPI-IO Hello World



mpiexec -n 4 ./helloworldc







MPI-IO Hello World

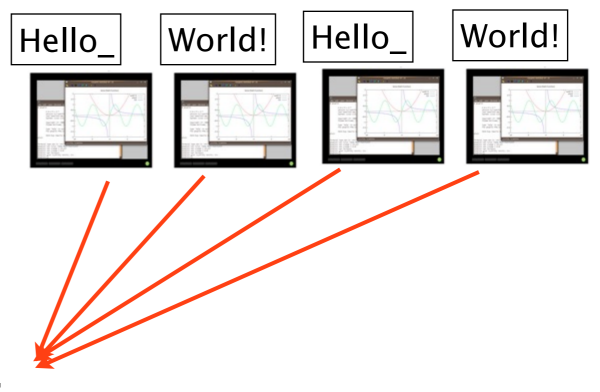


```
if ((rank % 2) == 0)
  strcpy (message, "Hello ");
else
  strcpy (message, "World!");
```







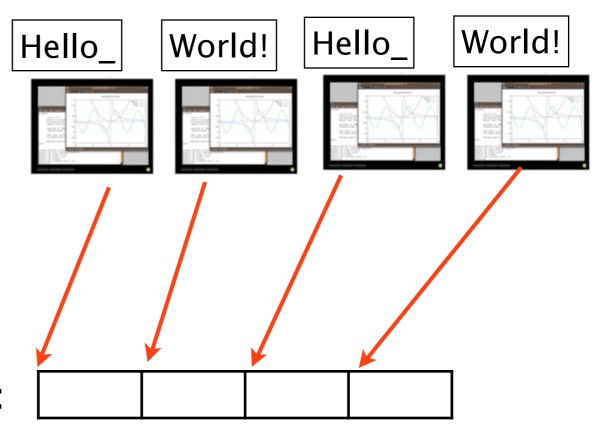


helloworld.txt:









helloworld.txt:

```
offset = (msgsize*rank);
MPI_File_seek(file, offset, MPI_SEEK_SET);
```







Hello_ World! Hello_ World!

Hello_World! Hello_World!

helloworld.txt:

MPI File write(file, message, msgsize, MPI CHAR, &status);









helloworld.txt: Hello_World! Hello_World!

MPI_File_close(&file);







Usual MPI startup/ teardown boilerplate

```
#include <stdio.h>
#include <string.h>
#include <mpi.h>
int main(int argc, char **argv) {
    int ierr, rank, size;
    MPI Offset offset;
    MPI File file;
    MPI Status status;
    const int msqsize=6;
    char message[msgsize+1];
    ierr = MPI Init(&argc, &argv);
    ierr = MPI Comm size(MPI COMM WORLD, &size);
    ierr = MPI Comm rank(MPI COMM WORLD, &rank);
    if ((rank % 2) == 0) strcpy (message, "Hello "); else strcpy (message)
    offset = (msqsize*rank);
    MPI File open(MPI COMM WORLD, "helloworld.txt", MPI MODE CREATE
                  MPI INFO NULL, &file);
    MPI File seek(file, offset, MPI SEEK SET);
    MPI File write(file, message, msgsize, MPI CHAR, &status);
    MPI File close(&file);
    MPI Finalize();
    return 0;
```







MPI_File_Open

Communicator; collective operation.







MPI_File_Open

Info allows us to send extra hints to MPI-IO layer about file(performance tuning, special case handling)

MPI_INFO_NULL: no extra info.

```
call MPI_File_Open( integer communicator, character(*) *filename, integer mode, integer info, integer handle, integer ierr);
```







Modes for MPI File Open():

MPI_MODE_RDONLY	read-only		
MPI_MODE_RDWR	read-write		
MPI_MODE_WRONLY	write-only		
MPI_MODE_CREATE	Create if doesn't exist.		
MPI_MODE_APPEND	On open, file pointers at end of file.		
MPI_MODE_EXCL	Fail if try to create, does exist.		
MPI_MODE_UNIQUE_OPEN	No one else is using this.		
MPI_MODE_SEQUENTIAL	Will be sequential access only.		
MPI_MODE_DELETE_ON_CLOSE	Delete when done. (OOC/scratch).		







MPI File Seek:

MPI_SEEK_SET	Set file pointer to position offset		
MPI_SEEK_CUR	pointer ← current position + offset		
MPI_SEEK_END	pointer ← end of file - offset		

Not collective; each adjusts its own local file pointer







MPI File write:

Not collective; each writes.







MPI_File_write:

- MPI File write is very much like a MPI_Send.
- "Sending" count of datatype from buf "to" the file.
- Here, writing 6 MPI_CHARs.
- Contiguous in memory starting in buffer.
- Status like a receive -- contains info about amount of data actually written, etc.







MPI File write:

count = 6

- To write out data that is noncontiguous in memory, same as MPI_Sending non-contig data:
 - Create type that describes data layout in memory
 - "Send" in units of that type.
- Noncontiguous data in memory is written out contiguously to the file, starting at the current (local) file pointer.

```
stride = 2

H@e # | * | ^ o & A

Hobline = I
```







```
integer, parameter :: msgsize=6, strsize=12
                                               helloworld-noncontigf.f90
character(strsize) :: message
integer :: everyother
! . . . .
if (mod(rank, 2) == 0) then
   message = "H@e#1*1^o* A"
else
   message = "WFoQr#1>d@! "
endif
! . . .
call MPI Type vector(msgsize, 1, 2, MPI CHARACTER, everyother, ierr)
call MPI_Type_commit(everyother, ierr)
call MPI File open(MPI COMM WORLD, "helloworld-nc.txt", ior(MPI MODE CREATE, MPI MODE WRONLY),&
                     MPI_INFO_NULL, fileno, ierr)
call MPI_File_seek (fileno, offset, MPI_SEEK_SET, ierr)
call MPI File write(fileno, message, 1, everyother, wstatus, ierr)
call MPI_File_close(fileno, ierr)
```

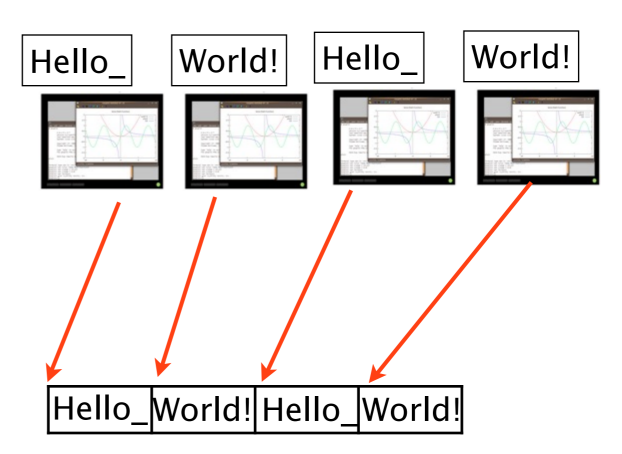






Seek + Write

- Works, but:
 - Very low level (gets complicated for less trivial data layouts)
 - Completely independent operations (seek, write)
 - Hard for any middleware to coordinate, improve anything.
 - Could do this with POSIX I/O.









MPI File write_at:







MPI_File_write_at:

Writes at a given offset

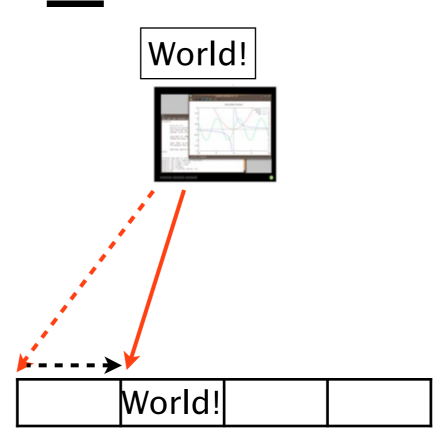






Write at

- Seek (relative to current, local, file pointer) + write in one operation.
- More explicit about what is going to happen some opt. possible.
- But actions of individual processors still independent - no collective optimization possible.









MPI File write at all:

writeatallc.c







MPI File write at all:

Writes at a given offset - Collective!

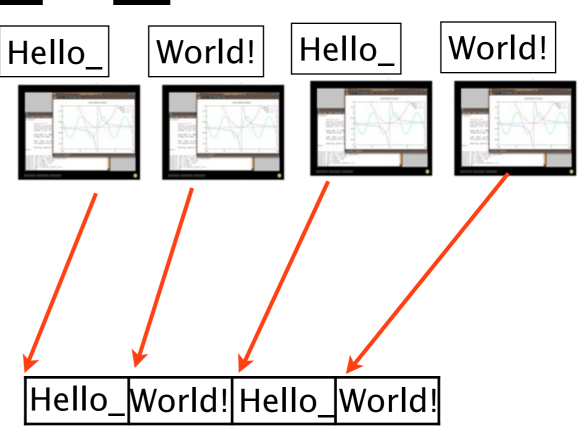






Write_at_all

- Much more explicit about what is going to happen globally.
- Collective operation.
- All processors participate.
- Higher order optimizations possible.









Non-contiguous in file

- Imagine having to write out a 2d file as to the right, with rank 0 "owning" the yellow data, etc.
- (eg, an image, or a complete checkpoint of a 2d domain).
- Would have to do repeated seeks, writing out one row at a time...

0	0	0	0	0	1	1	1	1	1
0	0	0	0	0	_	Τ	_	_	1
0	0	0	0	0	_	_	—	_	1
0	0	0	0	0	_	_	_	_	1
0	0	0	0	0	_	_	—	_	1
2	2	2	2	2	3	3	3	3	3
2	2	2	2	2	3	3	3	3	3
2	2	2	2	2	3	3	3	3	3
2	2	2	2	2	3	3	3	3	3







MPI-IO File View

```
    int MPI_File_set_view(
        MPI_File fh,
        MPI_Offset disp,
        MPI_Datatype etype,
        MPI_Datatype filetype,
        char *datarep,
        MPI_Info info)
    /* displacement in bytes from start */
        /* elementary type */
        /* file type; prob different for each proc */
        /* 'native' or 'internal' */
        /* MPI_INFO_NULL */
```

```
/ \\\/\/\/
etypes
disp
```



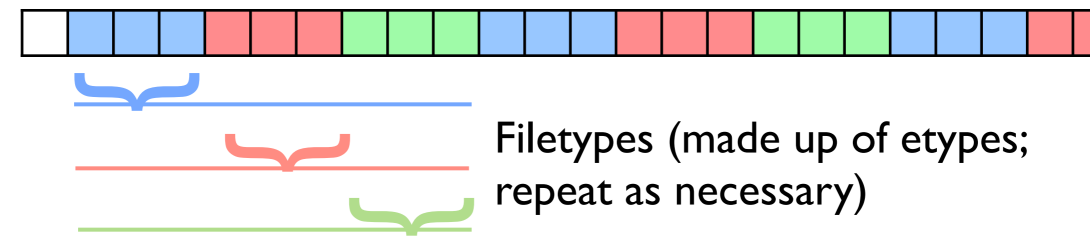




MPI-IO File View

```
    int MPI_File_set_view(
        MPI_File fh,
        MPI_Offset disp,
        MPI_Datatype etype,
        MPI_Datatype filetype,
        char *datarep,
        MPI_Info info)
```

```
/* displacement in bytes from start */
/* elementary type */
/* file type; prob different for each proc */
/* 'native' or 'internal' */
/* MPI INFO NULL */
```









MPI-IO File Write

```
    int MPI_File_write_all(
        MPI_File fh,
        void *buf,
        int count,
        MPI_Datatype datatype,
        MPI_Status *status)
```

Writes (_all: collectively) to part of file within view.







MPI_Type_create __subarray

- MPI_Type_create_subarray;
 piece of a multi-dimensional array.
- Much more convenient for higher-dimensional arrays
- (Otherwise, need vectors of vectors of vectors...)
- Here starts = [0,0],
 subsizes=[5,5], sizes=[10,10].

```
      0
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```

```
int MPI_Type_create_subarray(
    int ndims, int *array_of_sizes,
    int *array_of_subsizes,
    int *array_of_starts,
    int order,
    MPI_Datatype oldtype,
    MPI_Datatype &newtype);

call MPI_Type_create_subarray(
    integer ndims, [array_of_sizes],
    [array_of_subsizes],
    [array_of_starts],
    order, oldtype,
    newtype, ierr)
```







fileviewc.c

write locnrows*locncols contiguous MPI_CHARs to the (non-contiguous) view in file.







MPI-IO hands-on

- Fill in the blanks in sine.c or sinef.f90 to use MPI-IO to write out the I-d array of sin(x).
- Already written: decomposing the array, doing the calculation,
 MPI_File_open and MPI_file_close calls.
- Make sine (sinf) and make sineplot (sinefplot) to build the code, and run it and plot the results.
- Can use any of the approaches above
- ~15 minutes.







Formats for Scientific Data Management

NetCDF4, HDF5, and whatnot







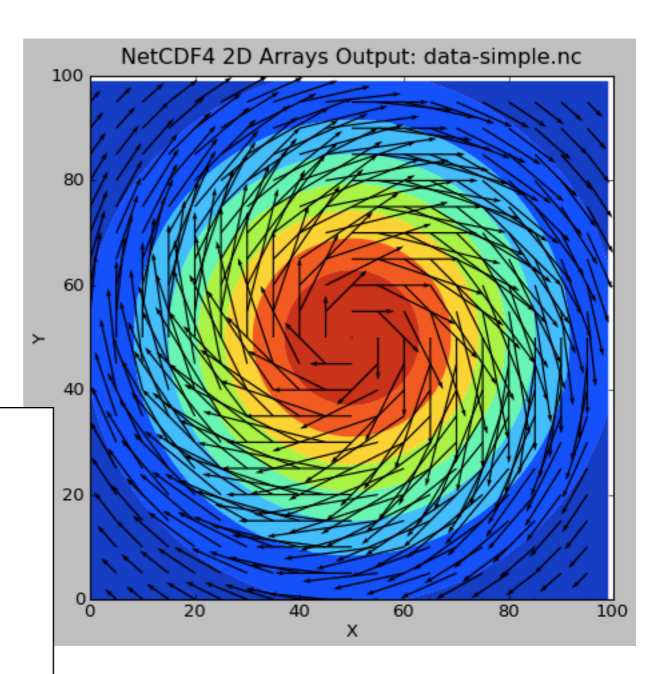
Sample Code

```
$ cd parIO/netcdf
```

```
$ make 2darray-simple (C), or
$ make f2darray-simple (F90)
```

```
$ ./{f,}2darray-simple
```

```
$ ls *.nc
$ ../plots.py *.nc
```

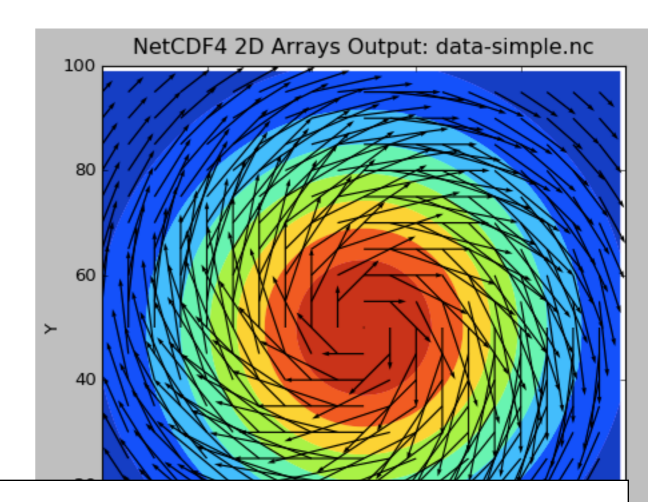






Sample Code

\$./2darray-simple --help



```
Options:
--nx=N (-x N): Set the number of grid cells in x direction.
--ny=N (-y N): Set the number of grid cells in y direction.
--filename=S (-f S): Set the output filename.

$ ./f2darray-simple --help
Usage: f2darray-simple [--help] [filename [nx [ny]]]
where filename is output filename, and
nx, ny are number of points in x and y directions.
```

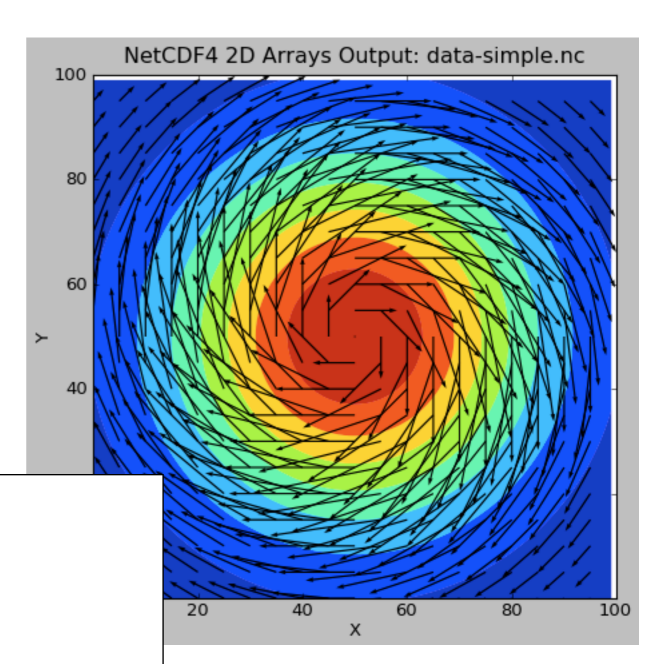






What is this .nc file?

```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
```



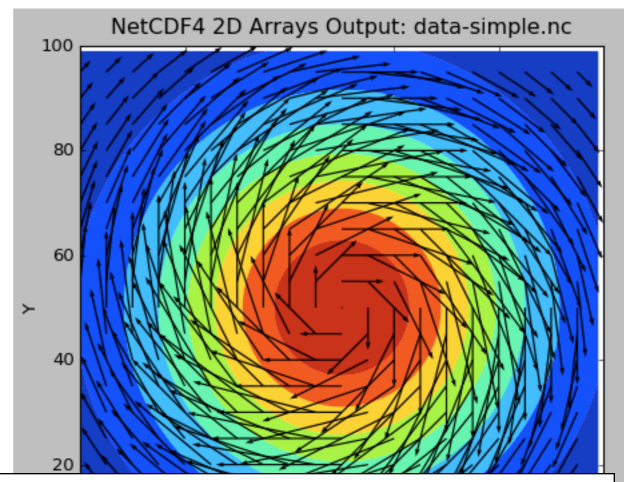






NetCDF

- NetCDF is a set of libraries and formats for:
 - portable,
 - efficient
 - "self-describing"
- way of storing and accessing large arrays (eg, for scientific data)
- Current version is NetCDF4



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
}
```

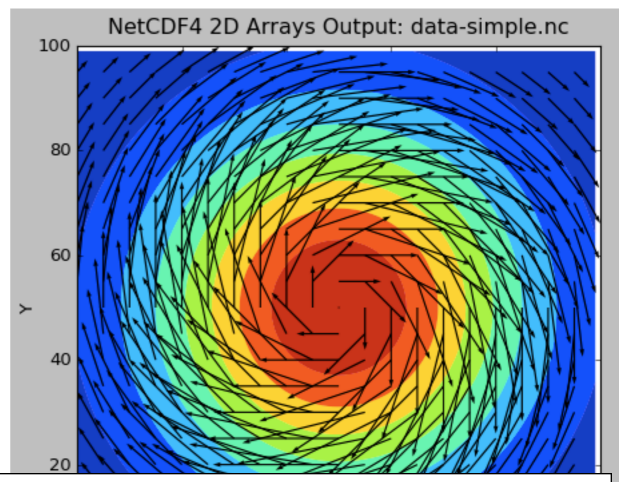






NetCDF: Portable

- Binary files, but common output format so that different sorts of machines can share files.
- Libraries accessible from C, C++, Fortran-77, Fortran 90/95/2003, python, etc.



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
  dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
  variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
}
```

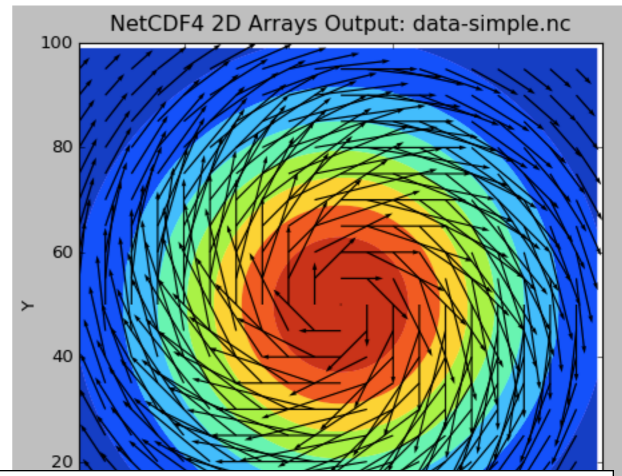






NetCDF: Self-Describing

- Header contains the metadata to describe the big data
- Lists:
 - Array names
 - Dimensions
 - shared dimensions information about how the arrays relate
 - Other, related information



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
  dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
    variables:
        double Density(Y, X) ;
        double Velocity(Y, X, velocity\ components) ;
}
```



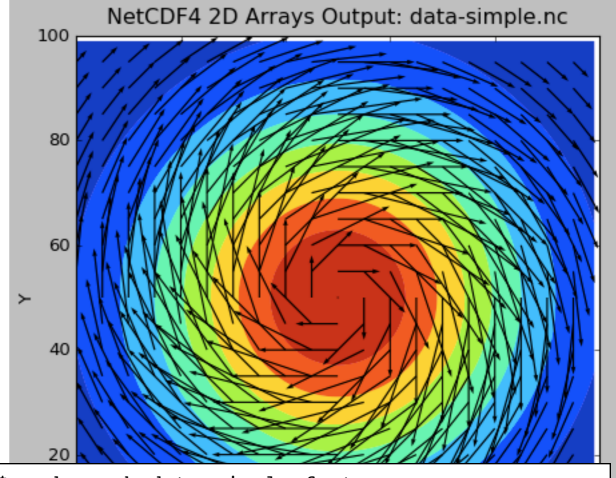




NetCDF: Efficient

- Binary, so less translation (as little is used as possible)
- IO libraries themselves are written for performance
- API, data format makes it easy to efficiently read, write subregions of arrays (slices, or 'hyperslabs')
- Still possible to make things slow
 - lots of metadata queries,

modifications HPCS2012



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
}
```





2darray-simple.c

```
#include "netcdf.h"
                                  Include function definitions
void writenetcdffile(rundata t rundata, double **dens,
                     double ***vel) {
    /* identifiers */
    int file id;
    /* return status */
    int status;
    /* Create a new file - clobber anything existing */
    status = nc_create(rundata.filename, NC_CLOBBER, &file_id);
    /* netCDF routines return NC_NOERR on success */
    if (status != NC_NOERR) {
        fprintf(stderr, "Could not open file %s\n", rundata.filename);
```







2darray-simple.c

```
#include "netcdf.h"
void writenetcdffile(rundata_t rundata, double **dens,
                    double ***vel) {
   /* identifiers */
    int file id;
                                 Create a new file, with name
                                          rundata.filename
   /* return status */
    int status;
   /* Create a new file clobber anything existing */
   status = nc_create(rundata.filename, NC_CLOBBER, &file_id);
   /* netCDF routines return NC_NOERR on success */
   if (status != NC_NOERR) {
       fprintf(stderr, "Could not open file %s\n", rundata.filename);
```







2darray-simple.c

```
#include "netcdf.h"
void writenetcdffile(rundata_t rundata, double **dens,
                    double ***vel) {
    /* identifiers */
    int file_id;
                                   Clobber anything already in
                                                 the file
    /* return status */
    int status;
    /* Create a new file - clobber anything existing */
    status = nc_create(rundata.filename, NC_CLOBBER, &file_id);
    /* netCDF routines return NC_NOERR on success */
    if (status != NC_NOERR) {
        fprintf(stderr, "Could not open file %s\n", rundata.filename);
```







2darray-simple.c

```
#include "netcdf.h"
void writenetcdffile(rundata_t rundata, double **dens,
                    double ***vel) {
    /* identifiers */
    int file id;
                                       Test the return codes
   /* return status */
    int status;
    /* Create a new file - clobber ___ything existing */
    status = nc_create(rundata_filename, NC_CLOBBER, &file_id);
    /* netCDF routines parn NC_NOERR on success */
    if (status != NC_NOERR) {
        fprintf(stderr, "Could not open file %s\n", rundata.filename);
```







```
subroutine writenetcdffile(rundata, dens, vel)
     use netcdf
                                                Import definitions
     implicit none
     type(rundata t), intent(IN) :: rundata
     double precision, intent(IN), dimension(:,:) :: dens
     double precision, intent(IN), dimension(:,:,:) :: vel
     integer :: file_id
     integer :: status
     ! create the file, check return code
     status = nf90 create(path=rundata%filename, cmode=NF90 CLOBBER,
ncid=file id)
     if (status /= NF90_NOERR) then
         print *,'Could not open file ', rundata%filename
     return
```







```
subroutine writenetcdffile(rundata, dens, vel)
     use netcdf
     implicit none
     type(rundata t), intent(IN) :: rundata
     double precision, intent(IN), dimension(:,:) :: dens
     double precision, intent(IN), dimension(:,:,:) :: vel
                                               Create file
     integer :: file_id
     integer :: status
     ! create the file, meck return code
     status = nf90 create(path=rundata%filename, cmode=NF90 CLOBBER,
ncid=file id)
     if (status /= NF90_NOERR) then
         print *,'Could not open file ', rundata%filename
     return
```







```
subroutine writenetcdffile(rundata, dens, vel)
    use netcdf
     implicit none
    type(rundata t), intent(IN) :: rundata
    double precision, intent(IN), dimension(:,:) :: dens
    double precision, intent(IN), dimension(:,:,:) :: vel
                                          C definitions are NC,
     integer :: file id
                                               F90 are NF90
     integer :: status
     ! create the file, chick return ode
    status = nf90_create(path=rundata%filename, cmode=NF90_CLOBBER,
ncid=file id)
     if (status /= NF90 NOERR) then
        print *,'Could not open file ', rundata%filename
    return
```

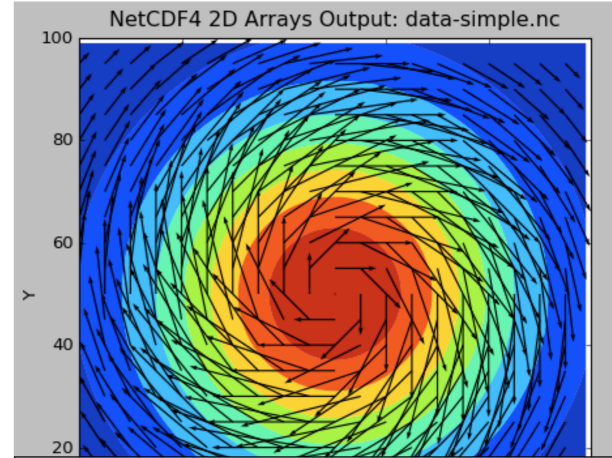






Writing a NetCDF File

- To write a NetCDF file, we go through the following steps:
 - Create the file (or open it for appending)
 - Define dimensions of the arrays we'll be writing
 - Define variables on those dimensions
 - End definition phase
 - Write variables
 - Close file



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
}
```

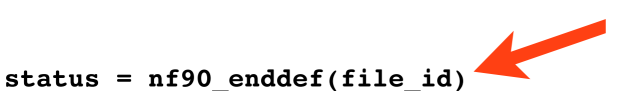






```
integer :: file id, xdim id, ydim id, vcomp id
integer :: dens id, vel id
integer, dimension(2) :: densdims
                                        Define the dimensions
integer, dimension(3) :: veldims
                                       in the file: name, size, id
status = nf90_def_dim(file_id, 'X', rundata%nx, xdim_id)
status = nf90_def_dim(file_id, 'Y', rundata%ny, ydim_id)
status = nf90 def dim(file id, 'velocity components', 2, vcomp id)
densdims = (/ xdim id, ydim id /)
veldims = (/ vcomp id, xdim id, ydim id /)
status = nf90 def var(file id, 'Density', NF90 DOUBLE, densdims, dens id)
if (status /= NF90_NOERR) print *, trim(nf90 strerror(status)), ' Dens'
status = nf90 def var(file id, 'Velocity', NF90 DOUBLE, veldims, vel id)
status = nf90 enddef(file id)
```

```
integer :: file id, xdim id, ydim id, vcomp id
integer :: dens id, vel id
integer, dimension(2) :: densdims
integer, dimension(3) :: veldims
status = nf90 def dim(file id, 'X', rundata%nx, xdim id)
status = nf90 def dim(file id, 'Y', rundata%ny, ydim id)
status = nf90 def dim(file id, 'velocity components', 2, vcomp id)
                                          Variables are defined in
densdims = (/ xdim_id, ydim_id /)
                                           terms of these dims
veldims = (/ vcomp_id, xdim_id, ydim_id /)
status = nf90_def_var(file_id, 'Density', NF90_DOUBLE, densdims, dens_id)
if (status /= NF90_NOERR) print *, trim(nf90 strerror(status)), ' Dens'
status = nf90_def_var(file_id, 'Velocity', NF90_DOUBLE, veldims, vel_id)
status = nf90 enddef(file id)
```



Once you're done defining things,

```
! Write out the values
status = nf90_put_var(file_id, dens_id, dens)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Dens'
status = nf90_put_var(file_id, vel_id, vel)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Vel'
status = nf90 close(file id)
```

cu

Writing data is easy.

```
! Write out the values
status = nf90_put_var(file_id, dens_id, dens)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Dens'
status = nf90_put_var(file_id, vel_id, vel)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Vel'
status = nf90_close(file_id)
```

cu

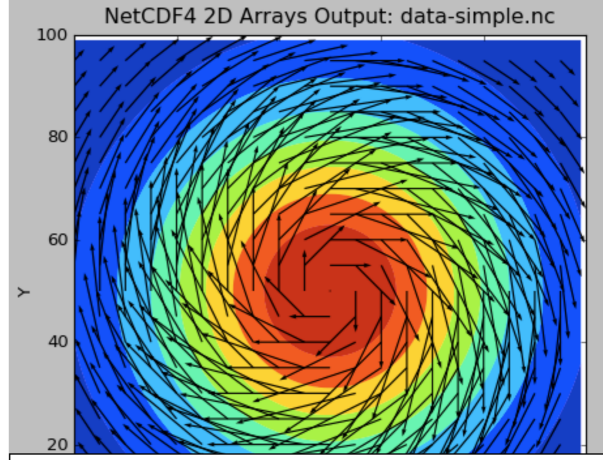
```
status = nf90_enddef(file_id)

! Write out the values
status = nf90_put_var(file_id, dens_id, dens)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Dens'
status = nf90_put_var(file_id, vel_id, vel)
if (status /= NF90_NOERR) print *, trim(nf90_strerror(status)), ' Vel'
status = nf90_close(file_id)
```

Closing the file is important!!

Reading a NetCDF File

- Flow is slightly different
 - Open the file for reading
 - Get dimension ids of the the dimensions in the files
 - Get dimension lengths so you can allocate the files
 - Get variable ids so you can access the data
 - Read variables
 - Close file



```
$ ncdump -h data-simple-fort.nc
netcdf data-simple-fort {
  dimensions:
    X = 100 ;
    Y = 100 ;
    velocity\ components = 2 ;
  variables:
    double Density(Y, X) ;
    double Velocity(Y, X, velocity\ components) ;
}
```







```
status = nf90_open(path=rundata%filename, mode=NF90_NOWRITE, ncid=file_id)
! find the dimensions
status = nf90 inq dimid(file id, 'X', xdim id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90 inq dimid(file id, 'velocity components', vcomp id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90 inquire dimension(file id, ydim id, len = rundata % ny )
status = nf90 inquire dimension(file id, vcomp id, len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90 inq varid(file id, 'Density', dens id)
status = nf90 inq varid(file id, 'Velocity', vel id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90 get var(file id, vel id, vel)
status = nf90 close(file id)
```

```
status = nf90 open(path=rundata%filename, mode=NF90 NOWRITE, ncid=file id)
! find the dimensions
status = nf90_inq_dimid(file_id, 'X', xdim_id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90_inq_dimid(file_id, 'velocity components', vcomp_id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90 inquire dimension(file id, ydim id, len = rundata % ny )
status = nf90 inquire dimension(file id, vcomp id, len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90 inq varid(file id, 'Density', dens id)
status = nf90 inq varid(file id, 'Velocity', vel id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90 get var(file id, vel id, vel)
status = nf90 close(file id)
```

```
status = nf90 open(path=rundata%filename, mode=NF90 NOWRITE, ncid=file id)
! find the dimensions
status = nf90 inq dimid(file id, 'X', xdim id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90 inq dimid(file id, 'velocity components', vcomp id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90_inquire_dimension(file_id, ydim_id, len = rundata % ny )
status = nf90_inquire_dimension(file_id, vcomp_id,len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90 inq varid(file id, 'Density', dens id)
status = nf90 inq varid(file id, 'Velocity', vel id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90 get var(file id, vel id, vel)
status = nf90 close(file id)
```

```
status = nf90 open(path=rundata%filename, mode=NF90 NOWRITE, ncid=file id)
! find the dimensions
status = nf90 inq dimid(file id, 'X', xdim id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90 inq dimid(file id, 'velocity components', vcomp id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90 inquire dimension(file id, ydim id, len = rundata % ny )
status = nf90 inquire dimension(file id, vcomp id, len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90_inq_varid(file_id, 'Density', dens_id)
status = nf90_inq_varid(file_id, 'Velocity', vel_id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90 get var(file id, vel id, vel)
status = nf90 close(file id)
```

```
status = nf90 open(path=rundata%filename, mode=NF90 NOWRITE, ncid=file id)
! find the dimensions
status = nf90 inq dimid(file id, 'X', xdim id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90 inq dimid(file id, 'velocity components', vcomp id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90 inquire dimension(file id, ydim id, len = rundata % ny )
status = nf90 inquire dimension(file id, vcomp id, len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90 inq varid(file id, 'Density', dens id)
status = nf90 inq varid(file id, 'Velocity', vel id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90_get_var(file_id, vel_id, vel)
status = nf90 close(file id)
```

```
status = nf90 open(path=rundata%filename, mode=NF90 NOWRITE, ncid=file id)
! find the dimensions
status = nf90 inq dimid(file id, 'X', xdim id)
status = nf90 inq dimid(file id, 'Y', ydim id)
status = nf90 inq dimid(file id, 'velocity components', vcomp id)
! find the dimension lengths
status = nf90_inquire_dimension(file_id, xdim_id, len = rundata % nx)
status = nf90 inquire dimension(file id, ydim id, len = rundata % ny )
status = nf90 inquire dimension(file id, vcomp id, len = rundata % nvelcomp)
! now we can allocate variable sizes
allocate(dens(rundata%nx, rundata%ny)) !...etc...
status = nf90 inq varid(file id, 'Density', dens id)
status = nf90 inq varid(file id, 'Velocity', vel id)
status = nf90_get_var(file_id, dens_id, dens)
status = nf90 get var(file id, vel id, vel)
status = nf90 close(file id)
```

```
status = nc open(rundata->filename, NC NOWRITE, &file id);
/* Get the dimensions */
status = nc inq dimid(file id, "X", &xdim id);
if (status != NC NOERR) fprintf(stderr, "Could not get X\n");
status = nc inq dimid(file id, "Y", &ydim id);
status = nc_inq_dimid(file_id, "velocity component", &vcomp_id);
status = nc inq dimlen(file id, xdim id, &(rundata->nx));
status = nc inq dimlen(file_id, ydim_id, &(rundata->ny));
status = nc inq dimlen(file id, vcomp id, &(rundata->nveldims));
nc_inq_varid(file_id, "Density", &dens_id);
nc inq varid(file id, "Velocity", &vel id);
nc_get_var_double(file_id, dens_id, &((*dens)[0][0]));
nc get var double(file id, vel id, &((*vel)[0][0][0]));
nc close(file id);
```

A Better example

- The above example is much more austere than a typical NetCDF file
- A more typical example is given in 2darray (or f2darray)
- make this, then run it
- ../plots.py data.nc
- (Same options as previous example)

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ component = 2 ;
variables:
 float X\ coordinate(X) ;
    X\ coordinate:units = "cm" ;
 float Y\ coordinate(Y) ;
    Y\ coordinate:units = "cm" ;
 double Density(X, Y) ;
    Density:units = "g/cm^3" ;
 double Velocity(velocity\
component, X, Y);
    Velocity:units = "cm/s" ;
```

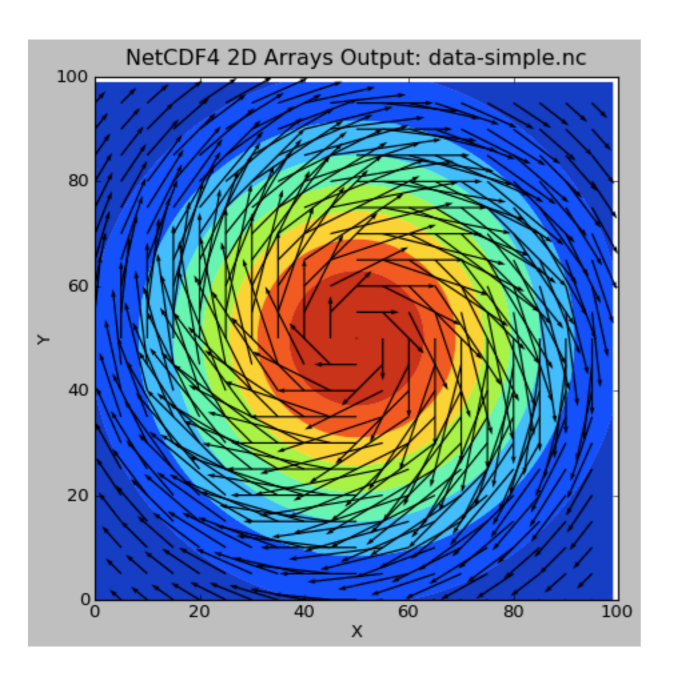


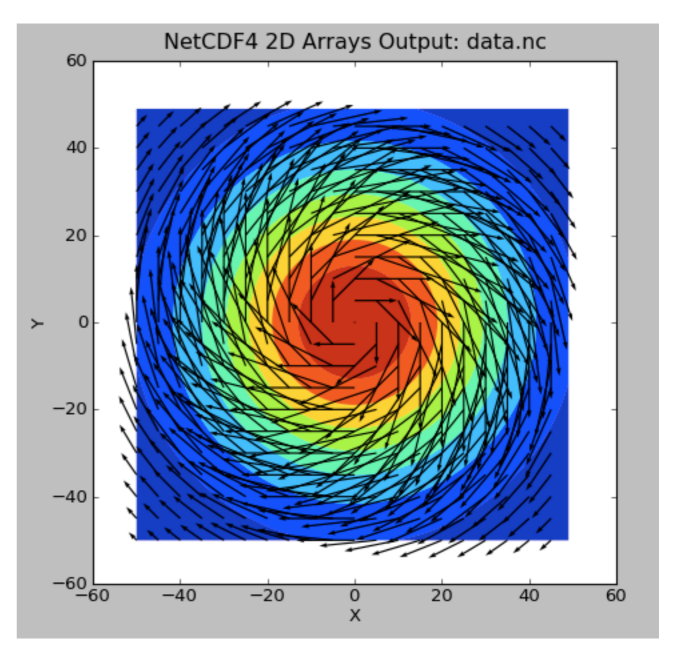




2darray.c

```
float *x, *y;
                                               Typically not only
const char *coordunit="cm";
                                            define dimensions but
for (i=0; i< rundata.nx; i++) x[i] = (1.*i-r)
                                            give coordinate values
for (i=0; i<rundata.ny; i++) y[i] = (1.*i-rt
/* define the dimensions */
nc_def_dim(file_id, "X", rundata.nx, &xdim_id);
nc def dim(file id, "Y", rundata.ny, &ydim id);
nc def dim(file id, "velocity component", 2, &vcomp id);
/* define the coordinate variables,... */
nc_def_var(file_id, "X coordinate", NC FLOAT, 1, &xdim id, &xcoord id);
nc def var(file id, "Y coordinate", NC_FLOAT, 1, &ydim_id, &ycoord_id);
/* ...and assign units to them as an attribute */
nc put att text(file id, xcoord id, "units", strlen(coordunit), coordunit);
nc_put_att_text(file_id, ycoord_id, "units", strlen(coordunit), coordunit);
```











2darray.c

```
float *x, *y;
                                             else) can have
const char *coordunit="cm";
                                               attributes:
for (i=0; i<rundata.nx; i++) x[i] = (1.*i-r)
                                         Name, and arbitrary
for (i=0; i< rundata.ny; i++) y[i] = (1.*i-r)
                                                   data
/* define the dimensions */
nc_def_dim(file_id, "X", rundata.nx, &xdim_id);
nc def dim(file id, "Y", rundata.ny, &ydim id);
nc def dim(file id, "velocity component", 2, &vcomp i/);
/* define the coordinate variables,... */
nc_def_var(file_id, "X coordinate", NC_FLOAT, 1, kdim_id, &xcoord_id);
/* ...and assign units to them as an attribute */
nc put att text(file id, xcoord id, "units", strlen(coordunit), coordunit);
nc_put_att_text(file_id, ycoord_id, "units", strlen(coordunit), coordunit);
```

Variables (or anything

NetCDF Attributes

- Any NetCDF object (data set, dimension) can have an arbitrary number of attributes associated with it
- Name, and any type or size...
- Like a variable! (But can't access only part of it).

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ component = 2 ;
variables:
 float X\ coordinate(X) ;
       coordinate:units
 float Y\ coordinate(Y)
       coordinate:units = "cm"
 double Density(X, Y) ;
    Density:units = "g/cm^3"
 double Velocity(velocity\
component, X, Y);
    Velocity:units = "cm/s"
```







NetCDF Attributes

- Attributes are assumed to be "small", though.
- Stored in header information (not with big data)
- Don't put large arrays in there

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ component = 2 ;
variables:
 float X\ coordinate(X)
    X\ coordinate:units = "cm"
 float Y\ coordinate(Y)
    Y\ coordinate:units = "cm"
 double Density(X, Y) ;
    Density:units = "g/cm^3"
 double Velocity(velocity\
component, X, Y);
    Velocity:units = "cm/s"
}
```







NetCDF Attributes

 Units are particularly useful attributes, as if a code needs data in some other units (MKS), can convert.

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ component = 2 ;
variables:
 float X\ coordinate(X)
    X\ coordinate:units = "cm"
 float Y\ coordinate(Y)
    Y\ coordinate:units = "cm"
 double Density(X, Y) ;
    Density:units = "g/cm^3"
 double Velocity(velocity\
component, X, Y);
    Velocity:units = "cm/s"
}
```







Limits to Self-Description

- But what if some codes expect "centimetre" and you use cm?
- Or their code uses "Dens" or "Rho" and yours uses "Density?" Or uses momentum rather than velocity?

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ component = 2 ;
variables:
 float X\ coordinate(X) ;
    X\ coordinate:units = "cm"
 float Y\ coordinate(Y)
    Y\ coordinate:units = "cm"
 double Density(X, Y) ;
    Density:units = "g/cm^3";
 double Velocity(velocity\
component, X, Y);
    Velocity:units = "cm/s"
}
```







Conventions

- There are lists of conventions that you can follow for variable names, unit names, etc.
- If you are planning for interoperability with other codes, this is the way to go
- Codes expecting data following (say) CF conventions for geophys should recognize data in that



NetCDF Conventions

Unidata offers a repository and will maintain WWW links for sets of netCDF conventions, as suppo Conventions section of the netCDF User's Guide. The following sets of conventions are currently a

- CF Conventions (Recommended, if applicable)
- ACDD Conventions (Attribute Convention for Dataset Discovery)
- NCAR-RAF Conventions for Aircraft Data
- AMBER Trajectory Conventions for molecular dynamics simulations
- ARGO netCDF conventions for data centers
- National Oceanographic Data Center NetCDF Conventions
- Proposed CF Discrete Sampling Conventions (draft CF conventions for observational and potential)
- Developing Conventions for NetCDF-4
- COARDS Conventions (1995 standard that CF Conventions extends and generalizes)
- GDT Conventions (1999 standard that CF Conventions extends and generalizes)
- CDC Conventions (for gridded data, compatible with but more restrictive than COARDS)
- NUWG Conventions (1992-1995 effort to create some observational data conventions)



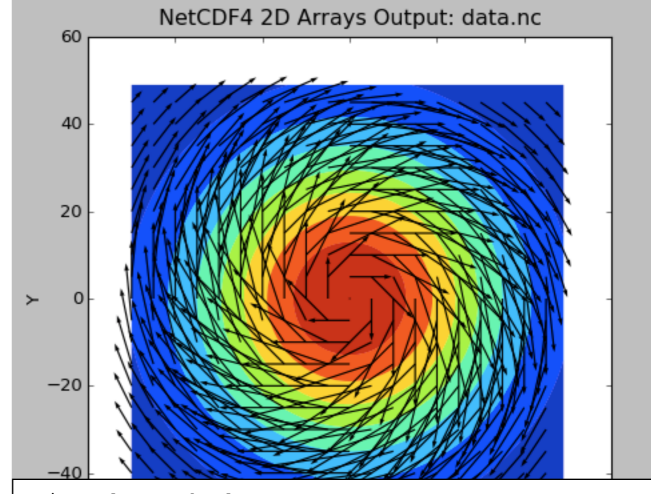




Big advantage of self-describing:

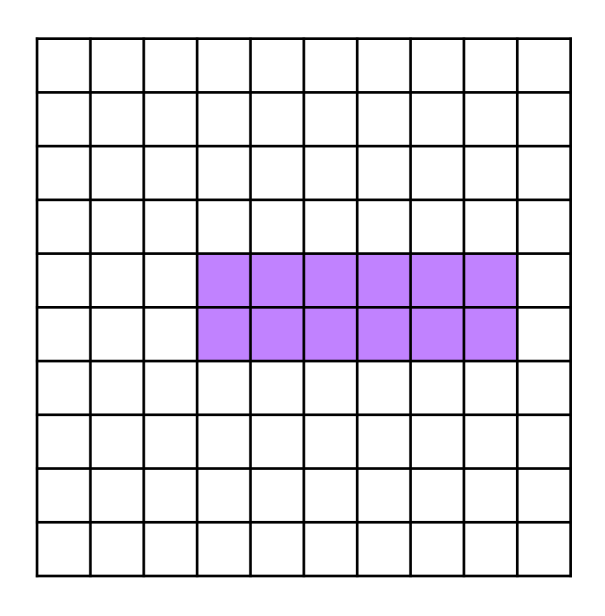
- Old program could easily read new file, even though data layout changed!
- Doesn't even need to know about attributes...
- New variables don't cause any problems - don't have to read them!
- Backwards compatibility
- But can look for them and use if





Accessing subregions in file

- nc_put_var_type or nf90_put_var puts whole array(by default)
- Subarrays can be specified with starts and counts







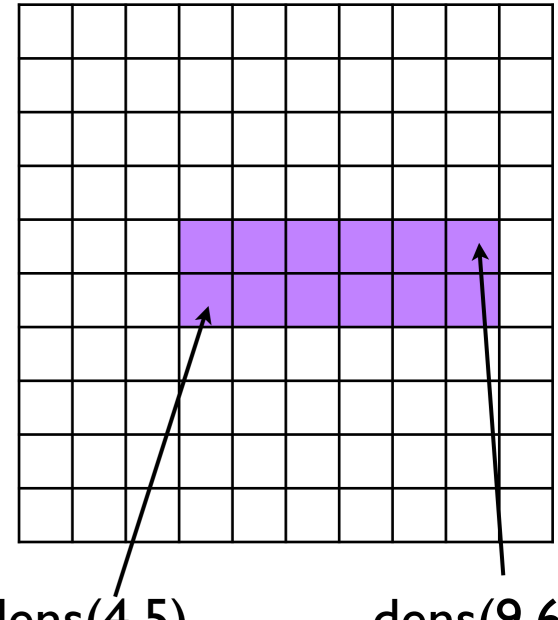


dens(10,10)

start(1) = 4
start(2) = 5

count(1) = 6
count(2) = 2

nf90_put_var(file_id, dens_id,
data, START=start, COUNT=count)



dens(4,5)

dens(9,6)





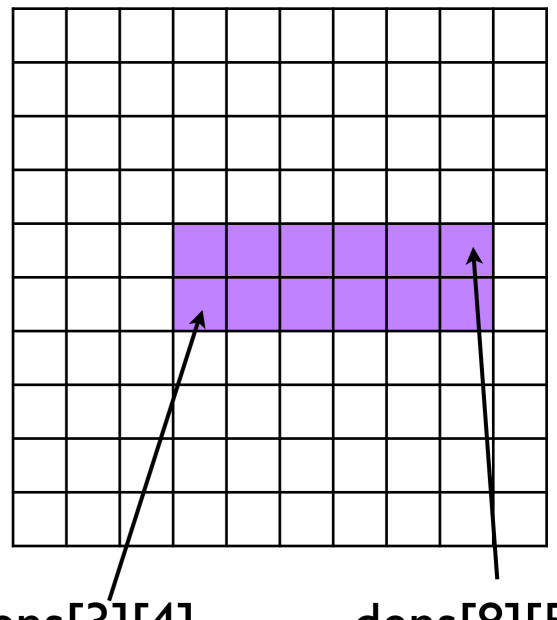


dens[10][10]

```
start[0] = 3;
start[1] = 4;

count[0] = 6;
count[1] = 2;

nc_put_vara_double(file_id,
dens_id, start, count, data);
```



dens[3][4]

dens[8][5]

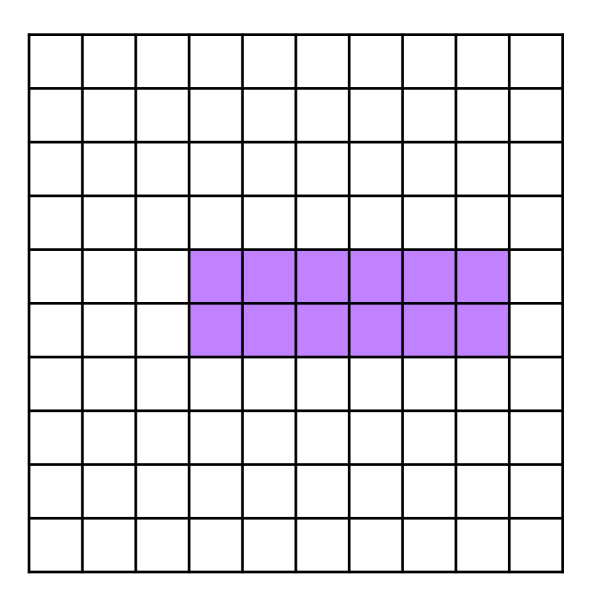






Accessing subregions in file

 Note that NetCDF libraries accepts starting conventions of C, Fortran as appropriate.



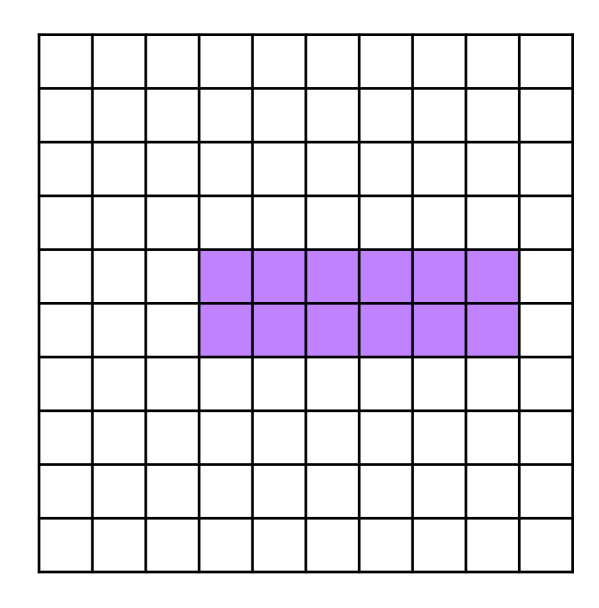






Accessing subregions in file

- Another thing this is good for; arrays in NetCDF can have a dimension of unlimited size (eg, can grow) - NetCDF3, only one dimension, NetCDF4, any
- Can use for timesteps, for instance.
- Any access to such a dataset is necessarily via subregions.









Fortran vs C array conventions

```
$ ncdump -h data.nc
netcdf data {
dimensions:
 X = 100 ;
 Y = 100;
 velocity\ component = 2 ;
variables:
  float X\ coordinate(X);
     X\ coordinate:units = "cm" ;
  float Y\ coordinate(Y) ;
     Y\ coordinate:units = "cm" ;
 double Density(X, Y) ;
     Density:units = "g/cm^3" ;
 double Velocity(velocity\
component, X, Y);
     Velocity:units = "cm/s";
```

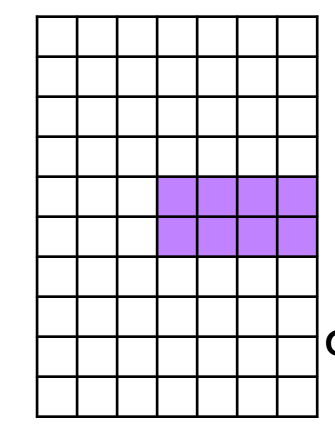
```
$ ncdump -h data-fort.nc
netcdf data-fort {
dimensions:
 X = 100;
 Y = 100 ;
 velocity\ components = 2 ;
variables:
  float X\ coordinate(X) ;
     X\ coordinate:units = "cm" ;
  float Y\ coordinate(Y) ;
     Y\ coordinate:units = "cm" ;
 double Density(Y, X);
     Density:units = "g/cm^3";
  double Velocity(Y, X, velocity\
components);
     Velocity:units = "cm/s" ;
```







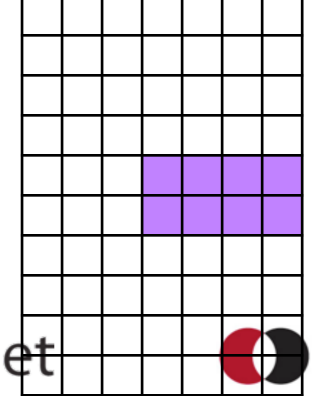
- Say in C you wanted to output in FORTRAN convention
- (i,j) in your array corresponds to (j,i) in data space in file
- nc_put_varm allows you to do this by mapping how indicies vary in memory compared to in file.



dens[7][10]

to

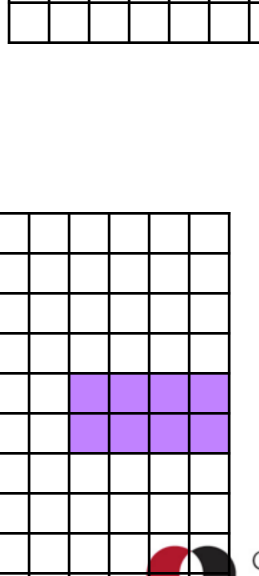
dens(10,7)

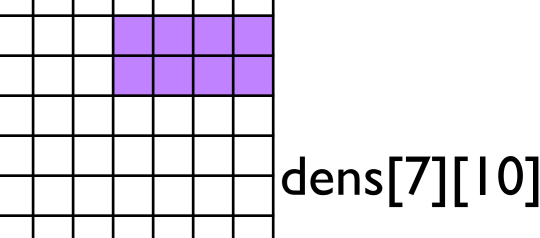






- Note this requires understanding how memory is laid out in your data structures, as with MPI & MPI-IO
- This is crucial for I/O, and for HPC in general
- C has more flexibility (== potential problems) in this regard.





to
dens(10,7)





- C: first array index most slowly varying.
- Eg, for a 3x4 array, each step in the 2nd index jumps you one position in memory,
- and in the first index, jumps you by 4.
- You could write this as (4,1)

Your picture of the array

[2][0]	[2][1]	[2][2]	[2][3]
[1][0]	[1][1]	[1][2]	[1][3]
[0][0]	[0][1]	[0][2]	[0][3]

[0][0]	[0][1]	[0][2]	[0][3]	[1][0]	[1][1]	[1][2]	[1][3]	[2][0]
--------	--------	--------	--------	--------	--------	--------	--------	--------







- But if you're writing to a fortran-convention file, you want this to go the other way
- In the file, one step in the Ist index should jump you by I, and the second by...

Your picture of the array

[2][0]	[2][1]	[2][2]	[2][3]
[1][0]	[1][1]	[1][2]	[1][3]
[0][0]	[0][1]	[0][2]	[0][3]

[0][0]	[0][1]	[0][2]	[0][3]	[1][0]	[י][י]	[1][2]	[1][3]	[2][0]
--------	--------	--------	--------	--------	--------	--------	--------	--------







- But if you're writing to a fortran-convention file, you want this to go the other way
- In the file, one step in the Ist index should jump you by I, and the second by 3.
- The map you want is (1,3)

Your picture of the array

[2][0]	[2][1]	[2][2]	[2][3]
[1][0]	[1][1]	[1][2]	[1][3]
[0][0]	[0][1]	[0][2]	[0][3]

[0][0]	[0][1]	[0][2]	[0][3]	[1][0]	[1][1]	[1][2]	[1][3]	[2][0]
--------	--------	--------	--------	--------	--------	--------	--------	--------







```
start = count = stride = NULL;
int imap[2] = {1,3};
```

nc_put_varm_double(file_id,
dens_id, start, count, stride,
imap, data);

nf90_put_var(file_id, dens_id,
data, MAP=(/4,1/))

Your picture of the array

[2][0]	[2][1]	[2][2]	[2][3]
[1][0]	[1][1]	[1][2]	[1][3]
[0][0]	[0][1]	[0][2]	[0][3]







More on NetCDF

- http://www.unidata.ucar.edu/ software/netcdf/
- Docs, mailing lists, tutorials, sample code, API, etc.









Sample Code

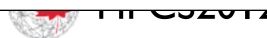


```
$ cd parIO/hdf5

$ source ../parallellibs
$ make serial or
$ make 2darray (C), or
$ make f2darray (F90)

$ ./{f,}2darray
$ ls *.h5

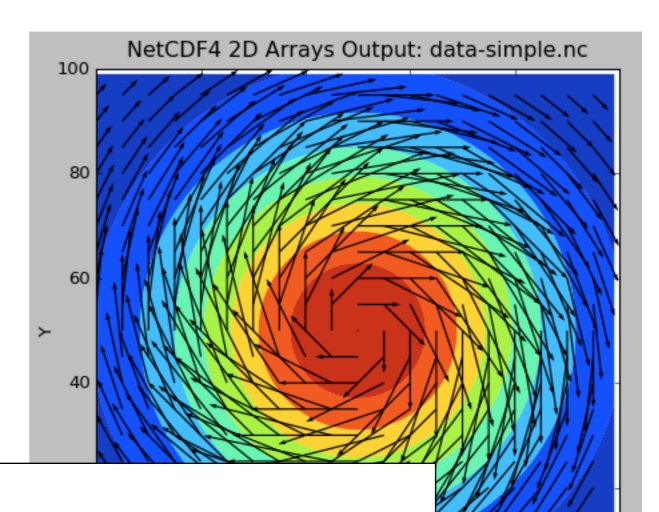
$ ../plots.py *.h5
```







What is this .h5 file?



\$ h5ls data-fort.h5

ArrayData

Group

OtherStuff

Group

\$ h5ls data-fort.h5/ArrayData

dens

Dataset {100, 100}

vel

Dataset {100, 100, 2}





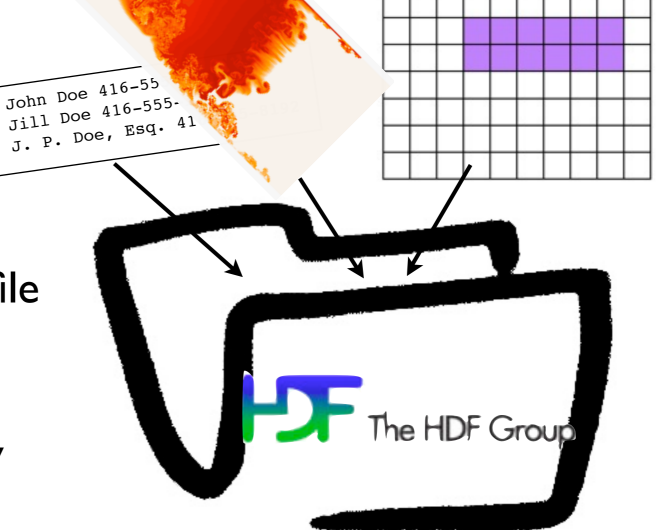


HDF5

- HDF5 is also self-describing file format and set of libraries
- Unlike NetCDF, much more general; can shove almost any type of data in there
- (We'll just be looking at large arrays, since that's our usual use case)
 HPCS2012







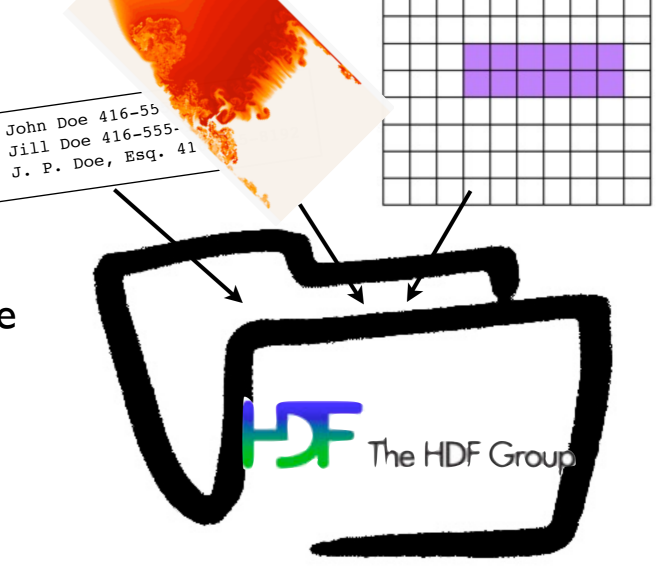
HDF5

- Much more general, and more low-level than NetCDF.
- (In fact, newest version of NetCDF implemented in HDF5).
- Pro: can do more!
- Con: have to do more.









```
/* identifiers */
hid_t file_id, dens_dataset_id, vel_dataset_id;
hid t dens dataspace id, vel dataspace id;
/* sizes */
hsize t densdims[2], veldims[3];
/* status */
herr t status;
/* Create a new file - truncate anything existing, use default properties
* /
file id = H5Fcreate(rundata.filename, H5F ACC TRUNC, H5P DEFAULT,
H5P DEFAULT);
/* HDF5 routines generally return a negative number on failure.
 * Should check return values! */
if (file_id < 0) {</pre>
    fprintf(stderr, "Could not open file %s\n", rundata.filename);
    return; }
```

```
/* identifiers */
hid_t file_id, dens_dataset_id, vel_dataset_id;
hid t dens dataspace id, vel dataspace id;
                                        NetCDF used ints for
/* sizes */
                                          everything - HDF5
hsize t depsdims[2], veldims[3];
                                        distinguishes between
/* status */
                                        ids, sizes, errors, uses
herr t status;
                                             its own types.
/* Create a new file - truncate anything
* /
file id = H5Fcreate(rundata.filename, H5F ACC TRUNC, H5P DEFAULT,
H5P DEFAULT);
/* HDF5 routines generally return a negative number on failure.
 * Should check return values! */
if (file id < 0) {
   fprintf(stderr, "Could not open file %s\n", rundata.filename);
   return; }
```

```
/* identifiers */
hid t file_id, dens_dataset_id, vel_dataset_id;
hid t dens dataspace id, vel dataspace id;
/* sizes */
hsize t densdims[2], veldims[3];
                                                  H5F, H5P...?
/* status */
herr t status;
/* Create a new file - trustate anything existing, use efault properties
* /
file id = H5Fcreate(rundata.filename, H5F_ACC_TRUNC, H5P_DEFAULT,
H5P DEFAULT);
/* HDF5 routines generally return a negative number on failure.
 * Should check return values! */
if (file_id < 0) {</pre>
    fprintf(stderr, "Could not open file %s\n", rundata.filename);
    return; }
```

Decomposing the HDF5 API

- HDF5 API is large
- Constants, function calls start with H5x; x tells you what part of the library
- Table tells you (some) of those parts...
- Fortran the same, but usually

end	with	F
HP	CS20	1 2



H5A	Attributes		
H5D	Datasets		
H5E	Errors		
H5F	Files		
H5G	Groups		
H5P	Properties		
H5S	Data Spaces		
H5T	Data Types		



```
/* Create the data space for the two datasets. */
densdims[0] = rundata.nx; densdims[1] = rundata.ny;
veldims[0] = 2; veldims[1] = rundata.nx; veldims[2] = rundata.ny;
dens dataspace id = H5Screate simple(2, densdims, NULL);
vel dataspace id = H5Screate simple(3, veldims, NULL);
/* Create the datasets within the file.
 * H5T IEEE F64LE is a standard (IEEE) double precision (64 bit)
 * floating (F) data type and will work on any machine.
 * H5T NATIVE DOUBLE would work too */
dens dataset id = H5Dcreate(file id, "dens", H5T IEEE F64LE,
                                dens dataspace id, H5P DEFAULT,
                                H5P DEFAULT, H5P DEFAULT);
vel_dataset_id = H5Dcreate(file_id, "vel", H5T_IEEE F64LE,
                                vel dataspace id, H5P DEFAULT,
                                H5P DEFAULT, H5P DEFAULT);
```

```
/* Create the data space for the two datasets. */
densdims[0] = rundata.nx; densdims[1] = rundata.ny;
veldims[0] = 2; veldims[1] = rundata.nx; veldims[2] = rundata.ny;
dens dataspace id = H5Screate simple(2, densdims, NULL);
vel dataspace id = H5Screate simple(3, veldims, NULL);
/* Create the datasets within the file
 * H5T IEEE F64LE is a stanlard (IEEE)
 * floating (F) data type and will work
 * H5T NATIVE DOUBLE would work too
dens_dataset_id = H5Dcreate(file id, "
                                dens_da
                                H5P DEI
vel_dataset_id = H5Dcreate(file_id,
                                vel dat
                                H5P DEI
```

All data (in file or in mem) in HDF5 has a dataspace it lives in.

In NetCDF, just cartesian product of dimensions; here more general

```
/* Create the data space for the two datasets. */
densdims[0] = rundata.nx; densdims[1] =
                                      Creating a data set like
veldims[0] = 2; veldims[1] = rundata.nx
                                        defining a variable in
dens dataspace id = H5Screate simple(2,
                                                NetCDF.
vel dataspace id = H5Screate simple(3)
                                       Also declare the type
/* Create the datasets within the file.
                                       you want it to be on
 * H5T IEEE F64LE is a standard (IEEE)
                                                   disk.
 * floating (F) data type and will work
 * H5T NATIVE DOUBLE rould work too */
dens dataset id = H5Dcreate(file id, "dens", H5T IEEE F64LE,
                              dens dataspace id, H5P DEFAULT,
                              H5P DEFAULT, H5P DEFAULT);
vel_dataset_id = H5Dcreate(file_id, "vel", H5T_IEEE F64LE,
                              vel dataspace id, H5P DEFAULT,
                              H5P DEFAULT, H5P DEFAULT);
```

status = H5Fclose(file id);

```
/* Write the data. We're writing it from memory, where it is saved
 * in NATIVE DOUBLE format */
status = H5Dwrite(dens dataset id, H5T NATIVE DOUBLE, H5S ALL, H5S ALL,
H5P DEFAULT, & (dens[0][0]));
status = H5Dwrite(vel dataset id, H5T NATIVE_DOUBLE, H5S_ALL, H5S_ALL,
H5P DEFAULT, &(vel[0][0][0]));
/* End access to groups data sets and release resources used by them */
status = H5Sclose(dens_data_pace_id);
status = H5Dclose(dens dataset id);
status = H5Sclose(vel dataspace N);
status = H5Dclose(vel_dataset_id);
                                      Write memory from all
/* Close the file */
```

Write memory from all of memory to all of the dataset on the file.

Values in mem are in the native double precision format.

```
/* Write the data. We're writing it from memory, where it is saved
 * in NATIVE DOUBLE format */
status = H5Dwrite(dens dataset id, H5T NATIVE DOUBLE, H5S ALL, H5S ALL,
H5P DEFAULT, & (dens[0][0]));
status = H5Dwrite(vel dataset id, H5T NATIVE_DOUBLE, H5S_ALL, H5S_ALL,
H5P DEFAULT, &(vel[0][0][0]));
/* End access to groups & data sets and release resources used by them */
status = H5Sclose(dens_dataspace_id);
status = H5Dclose(dens dataset id);
status = H5Sclose(vel dataspace id);
status = H5Dclose(vel_dataset_id);
                                          Close everything
/* Close the file */
status = H5Fclose(file id);
```

```
integer(hid t) :: file id
integer(hid_t) :: dens_space_id, vel_space_id
integer(hid t) :: dens id, vel id
integer(hsize t), dimension(2) :: densdims
integer(hsize t)__dimension(3) :: veldims
                                            Fortran: values are
integer :: status
                                             integer(hid_t) or
! first we have to open the FORTRAN inter
                                              integer(hsize_t)
call h5open f(status)
! create the file, check return code
call h5fcreate f(rundata%filename, H5F ACC TRUNC F, file id, status)
if (status /= 0) then
   print *, 'Could not open file ', rundata%filename
   return
endif
```

```
integer(hid t) :: file id
integer(hid_t) :: dens_space_id, vel_space_id
integer(hid t) :: dens id, vel id
integer(hsize t), dimension(2) :: densdims
integer(hsize t), dimension(3) :: veldims
integer :: status
! first we have to open the FORTRAN interface.
call h5open_f(status)
                                            Have to start the
                                          FORTRAN interface
! create the file, check return code
call h5fcreate f(rundata%filename, H5F ACC TRUNC F, file id, status)
if (status /= 0) then
   print *, 'Could not open file ', rundata%filename
   return
endif
```

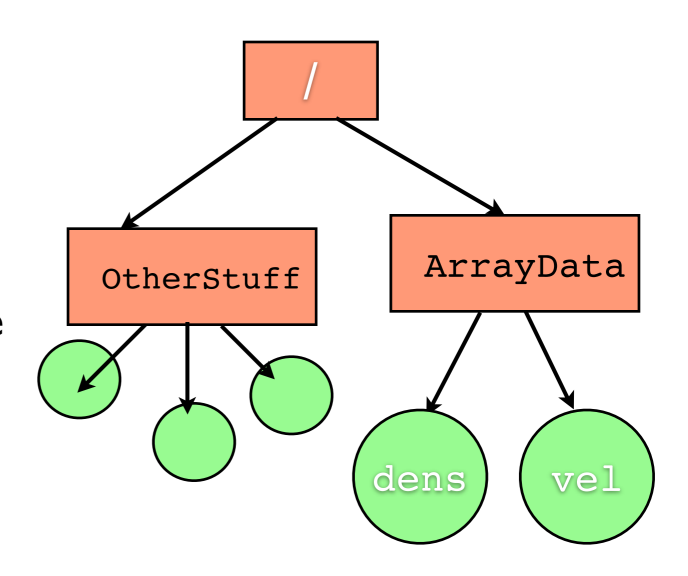
```
integer(hid t) :: file id
integer(hid_t) :: dens_space_id, vel_space_id
integer(hid t) :: dens id, vel id
integer(hsize t), dimension(2) :: densdims
integer(hsize t), dimension(3) :: veldims
integer :: status
! first we have to open the FORTRAN interface.
call h5open_f( tus)
                                         See what I mean about
! create the file, check return
call h5fcreate_f(: ___aata%filename, H5F_ACC_TRUNC_F, file_id, status)
if (status /= 0) then
   print *, 'Could not open file ', rundata%filename
   return
endif
```

```
! create the dataspaces corresponding to our variables
densdims = (/ rundata % nx, rundata % ny /)
call h5screate simple f(2, densdims, dens space id, status)
veldims = (/ 2, rundata % nx, rundata % ny /)
call h5screate simple f(3, veldims, vel space id, status)
! now that the dataspaces are defined, we can define variables on them
call h5dcreate f(file id, "dens", H5T IEEE F64LE, dens space id, dens id,
status)
call h5dcreate_f(file_id, "vel" , H5T_IEEE_F64LE, vel_space_id, vel_id,
status)
```

In F90 interface, a lot of less-common arguments are optional; fewer H5P_DEFAULTs kicking around

HDF5 Groups

- HDF5 has a structure a bit like a unix filesystem:
- "Groups" directories
- "Datasets" files
- NetCDF4 now has these, but breaks compatibility with NetCDF3 files









2darray.c

```
/* Create a new group within the new file */
arr group id = H5Gcreate(file id, "/ArrayData", H5P DEFAULT, H5P DEFAULT,
H5P DEFAULT);
dens dataset id = H5Dcreate(file id, "/ArrayData/dens", H5T IEEE F64LE,
                                dens dataspace id, H5P DEFAULT,
                                H5P DEFAULT, H5P DEFAULT);
vel dataset id = H5Dcreate(file id, "/ArrayData/vel", H5T IEEE F64LE,
                                vel_dataspa__id, H5P_DEFAULT,
                                H5P DEFAULT H5P DEFAULT);
```

Can specify that a dataset goes in a group by giving it an "absolute path"...

```
/* Create a new group within the new file */
arr group id = H5Gcreate(file id, "/ArrayData", H5P DEFAULT, H5P DEFAULT,
H5P DEFAULT);
dens dataset id = H5Dcreate(arr group id, "dens", H5T IEEE F64LE,
                                dens dataspace id, H5P DEFAULT,
                                H5P DEFAULT, H5P DEFAULT);
vel dataset id = H5Dcreate(arr group id, "vel", H5T IEEE F64LE,
                                v dataspace_id, H5P_DEFAULT,
                                H5P DEFAULT, H5P DEFAULT);
```

...or just by creating it *in* the group, rather than the file.

What NetCDF, HDF aren't

- Databases
- Seem like lots of information, in key value pairs.
- Relational databases interrelated tables of small
 pieces of data
- Very easy/fast to query

Books

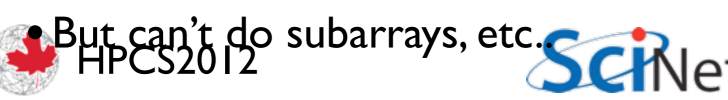
bid	title	isbn	author	date	volume
1	Big Cats	24589673-0	Cat, Simon	2003	2
2	Plants	24316759-1	Smith, Rose	1967	1
3	Sailing	34817645-0	Jones, Tom	1868	1
_					

Transactions

tid	date	bid	pid	duedate	
1	02/11/08	3	2	16/11/08	
2	04/11/08	1	3	18/11/08	

Borrowers

pid	firstname	lastname	address	phone	fines
1	Fred	Thompson	2 Reach Rd.	827-9867	2.25
2	Sam	Trunker	23 stone St.	243-0955	0
3	Tony	Sanchas	4 two Rd.	123-6453	0





Databases for science

INSERT INTO benchmarkruns
values (newrunnum, datestr,
timestr, juliannum)

• • •

SELECT nprocs, test, size,
transport, mpitype, runtime,
mopsperproc, run FROM
mpirundata WHERE (success=1)

run#	success	size	transport	•••
93	no	I2k	eth	
	yes	512	eth	
87	yes	64	ib	
13	no	32	eth	

• • •







Parallel I/O using NetCDF4, HDF5



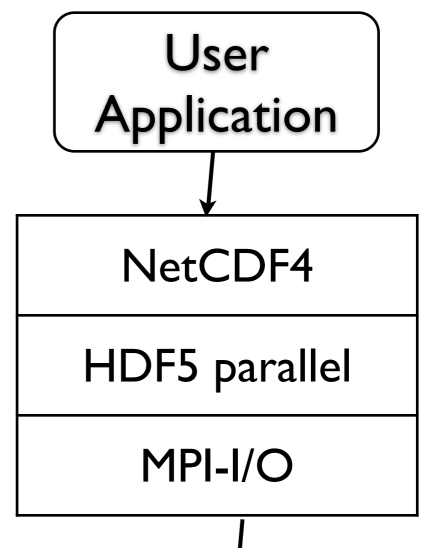




Parallel I/O libraries

- Can use the same NetCDF(4),
 HDF5 libraries to do Parallel
 IO on top of the MPI-I/O
 library
- Reading file afterwards, can't tell the difference.
- Fairly minor differences in function calls to do parallel I/O
- Hard part is figuring out what/ where to write







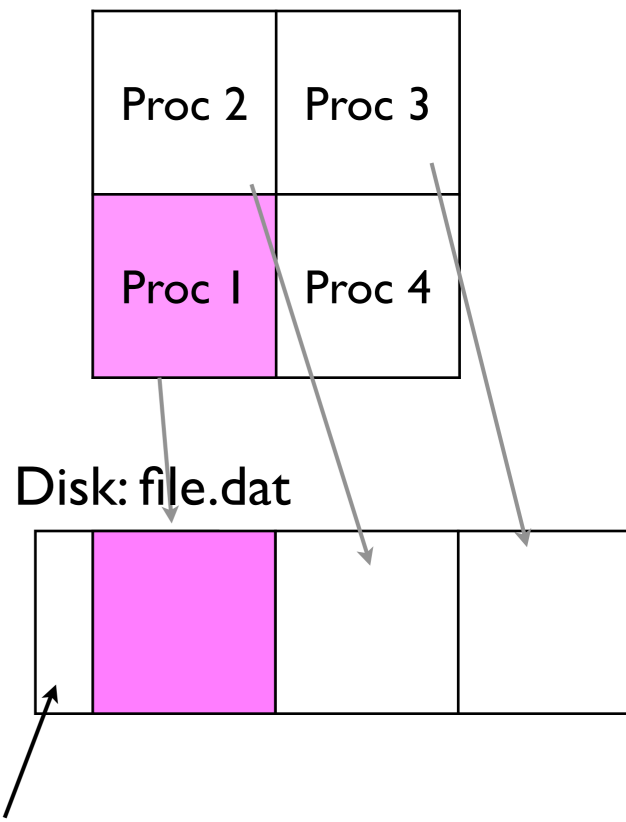
npute • calcul

CANADA



Parallel IO to One file

- Can be made to work efficiently, but must write to disjoint chunks of file
- Should write big disjoint chunks of file.





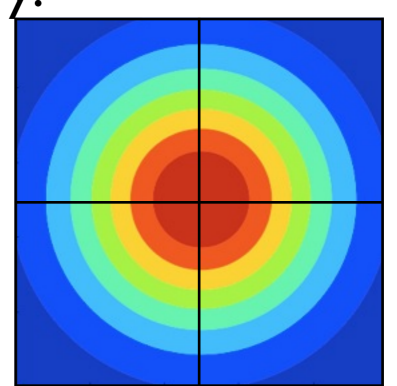




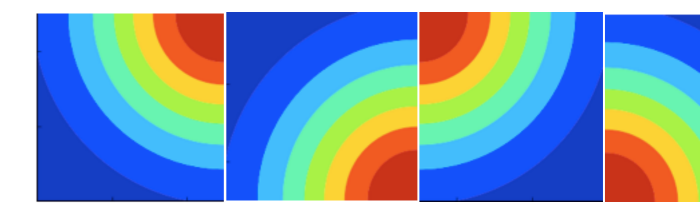
Memory:

How do you decide where to write?

- One possibility: each processor writes out its part of problem, in order.
- Pros can be super fast.
- Cons Output depends on number of processors run on. Analysis routines, restarts...



Disk:







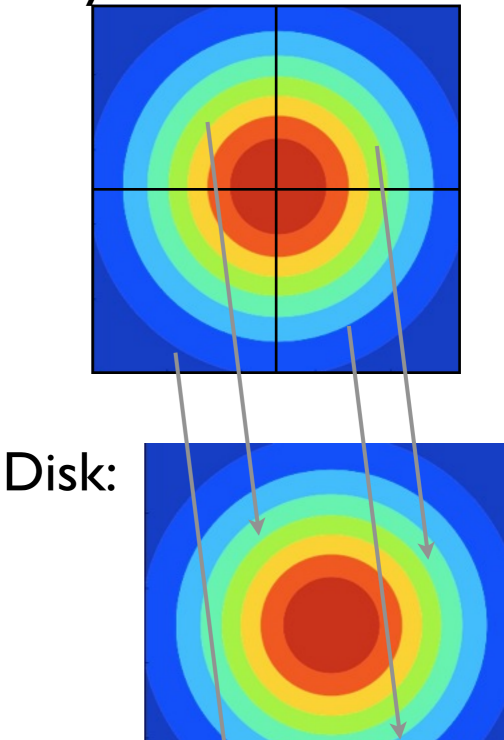


How do you decide where to write?

- Other possibility: Write out chunks as they would be in memory on serial machine
- Pros: File looks the same no matter how many processes were used to write.
- Cons: Noncontig access; may be slower, but MPI-IO collective + good parallel FS should make competitive.

HPCS2012

Memory:





Sample Code

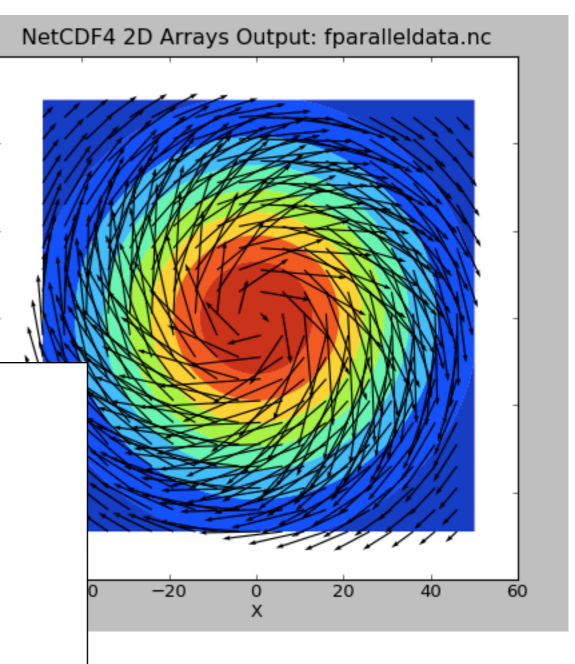
```
$ cd
$ cd parIO/netcdf

$ make parallel2darray (C), or
$ make fparallel2darray (F90)

$ mpirun -np 4 parallel2darray

$ ls *.nc
$ source ../seriallibs
```

../plots.py paralleldata.nc





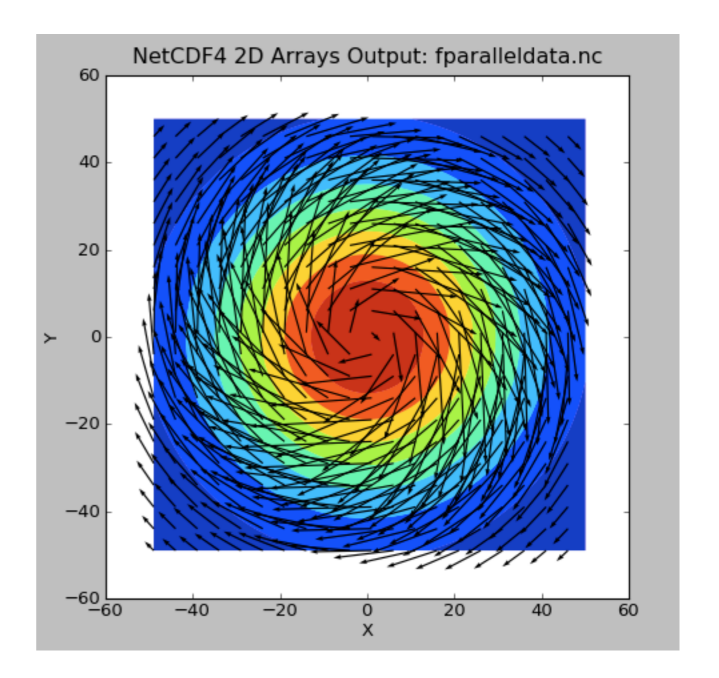




Sample Code

- Can do an ncdump -h...
- No trace of being written by different files
- Looks the same; code to read in is identical
- And not that much harder to code!
- By far the trickiest part is figuring out where in the file to write.

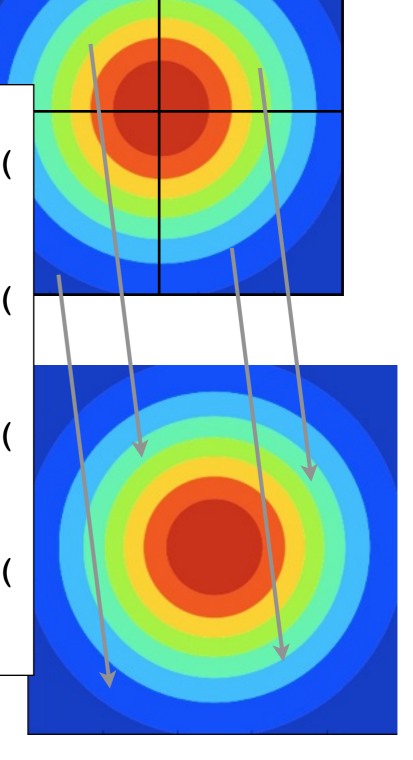
HPCS2012





Memory:

```
$ mpirun -np 4 ./fparallel2darray
[ 0] gets (0, 0): local points = (
50, 50); global points = (100, 100).
[ 1] gets (1, 0): local points = (
50, 50); global points = (100, 100).
[ 2] gets (0, 1): local points = (
50, 50); global points = (100, 100).
[ 3] gets (1, 1): local points = (
50, 50); global points = (100, 100).
```

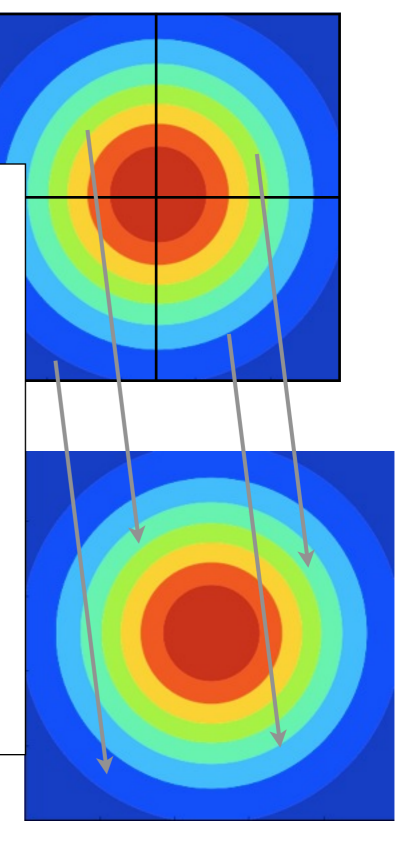








Memory:









```
call MPI Info create(info, status)
call MPI Info set(info, "IBM largeblock io", "true", status)
mode flag = IOR(NF90 MPIIO, NF90 CLOBBER)
mode flag = IOR(mode flag, NF90 NETCDF4)
status = nf90_create_par(rundata%filename, mode_flag,
MPI_COMM_WORLD, info, file id)
if (status /= NF90 NOERR) then
      print *,'Could not open file ', rundata%filename
      return
endif
```

Icul

```
call MPI Info create(info, status)
call MPI Info set(info, "IBM largeblock io", "true", status)
mode flag = IOR(NF90 MPIIO, NF90 CLOBBER)
mode flag = IOR(mode flag, NF90 NETCDF4)
status = nf90_create_par(rundata%filename, mode_flag,
MPI_COMM_WORLD, info, file id)
if (status /= NF90 NOERR) hen
      print *, 'Could not oben file ', rundata%filename
      return
endif
```

create_par rather than create

```
call MPI Info create(info, status)
call MPI Info set(info, "IBM largeblock io", "true", status)
mode flag = IOR(NF90 MPIIO, NF90 CLOBBER)
mode_flag = IOR(mode_flag, NF90_NETCDF4)
status = nf90 create par(rundata%f ename, mode_flag,
MPI COMM WORLD, info, file id)
if (status /= NF90 NOERR) then
      print *, 'Could not open file ', rundata%filename
      return
endif
```

mode_flag = CLOBBER | MPIIO | NETCDF4

```
call MPI Info create(info, status)
call MPI Info set(info,"IBM largeblock io","true", status)
mode flag = IOR(NF90 MPIIO, NF90 CLOBBER)
mode flag = IOR(mode flag, NF90 NETCDF4)
status = nf90 create par(rundata%filename, mode flag,
MPI_COMM_WORLD, info, file id)
if (status _ NF90_NOERR) then
      print *,'Could not open file ', rundata%filename
      return
endif
```

Extra arguments: communicator that will do the I/O

```
call MPI_Info_create(info, status)
call MPI_Info_set(info,"IBM_largeblock_io","true", status)
mode flag = IOR(NF90 MPIIO, NF90 CLOBBER)
mode flag = IOR(mode flag, NF90 NETCDF4)
status = nf90 create par(rundata%filename, mode flag,
MPI_COMM_WORLD, info, file id)
if (status /= NF9 NOERR) then
      print *,'Could not open file ', rundata%filename
      return
endif
```

Extra arguments: MPI Info; can pass MPI-I/O "hints"

```
status = nf90 def dim(file id, 'X', rundata%globalnx, xdim id)
status = nf90 def dim(file id, 'Y', rundata%globalny, ydim id)
status = nf90 def dim(file id, 'velocity components', 2,
vcomp id)
! now that the dimensions are defined, define variables
densdims = (/ xdim id, ydim id /)
veldims = (/ vcomp id, xdim id, ydim id /)
status = nf90_def_var(file_id, 'Density', NF90 DOUBLE, densdims,
dens id)
status = nf90 def var(file id, 'Velocity', NF90 DOUBLE, veldims,
vel id)
```

Defining variables identical (but global v local)







```
status = nf90_var_par_access(file_id, dens_id, NF90_COLLECTIVE)
status = nf90_var_par_access(file_id, vel_id, NF90_COLLECTIVE)

status = nf90_put_var(file_id, dens_id, dens, start=densstarts, count=denscounts)
status = nf90_put_var(file_id, vel_id, vel, start=velstarts, count=velcounts)

status = nf90_close(file_id)
```

Define how we'll be accessing variables - COLLECTIVE vs INDEPENDANT. (eg, Write_all vs. Write).







```
status = nf90 var par access(file id, dens id, NF90 COLLECTIVE)
status = nf90 var par access(file id, vel id, NF90 COLLECTIVE)
status = nf90_put_var(file_id, dens_id, dens, start=densstarts,
count=denscounts)
status = nf90_put_var(file_id, vel_id, vel, start=velstarts,
count=velcounts)
status = nf90 close(file id)
```

put_var is exactly like serial with subsections - starts, counts







```
status = nf90_var_par_access(file_id, dens_id, NF90_COLLECTIVE)
status = nf90_var_par_access(file_id, vel_id, NF90_COLLECTIVE)

status = nf90_put_var(file_id, dens_id, dens, start=densstarts, count=denscounts)
status = nf90_put_var(file_id, vel_id, vel, start=velstarts, count=velcounts)

status = nf90_close(file_id)
```



close is the same as ever.







```
serial.c
/* name of units for dens, vel */
const char *densunit="q/cm^3";
const char *velunit="cm/s";
/* return status */
int status;
/* set up x, y coordinates */
x = (float *)malloc(rundata.nx * sizeof(float));
y = (float *)malloc(rundata.my * sizeof(float));
for (i=0; i<rundata.nx; i++)
   x[i] = (1. *i-rundata. nx/2.);
for (i=0; i<rundata.ny; i++)
    y[i] = (1, *i-rundata, ny/2.);
/* Create a new file - clobber anything existing */
status = nc create(rundata.filename, NC_CLOBBER, &file_id);
/* netCDF routines return NC NOERR on success */
if (status != NC NOERR) {
    fprintf(stderr, "Could not open file %s\n", rundata.filename);
    return;
/* define the dimensions */
nc_def_dim(file_id, "X", rundata.nx, &xdim_id);
nc_def_dim(file_id, "Y", rundata.ny, &ydim_id);
nc_def_dim(file_id, "velocity component", 2, &vcomp_id);
/* define the coordinate variables,... */
```

```
parallel.c
/* name of units for dens, vel */
const char *densunit="q/cm^3";
const char *velunit="cm/s";
/* offsets for sub-regions of arrays */
size_t starts[3];
size_t counts[3];
/* return status */
int status;
/* MPI-IO hints for performance */
MPI_Info info;
/* set up x, y coordinates */
x = (float *)malloc(rundata.qlobalnx * sizeof(float));
y = (float *)malloc(rundata. qlobalny * sizeof(float));
for (i=0; i<rundata qlobalnx; i++)
    x[i] = (1. *i-rundata. qlobalnx/2.);
for (i=0; i<rundata. qlobalny; i++)
    v[i] = (1.*i-rundata.qlobalny/2.);
/* set the MPI-IO hints for better performance on GPFS */
MPI Info create(&info);
MPI_Info_set(info,"IBM_largeblock_io","true");
/* Create a new file - clobber anything existing */
status = nc create_par(rundata.filename, NC_MPIIO|NC_CLOBBER|N
                   MPI COMM WORLD, info, &file id);
/* netCDF routines return NC NOERR on success */
if (status != NC NOERR) {
    fprintf(stderr, "Could not open file %s\n", rundata.filenam
    return;
/* define the dimensions */
nc_def_dim(file_id, "X", rundata.globalnx, &xdim_id);
nc_def_dim(file_id, "Y", rundata.globalny, &ydim_id);
nc_def_dim(file_id, "velocity component", 2, &vcomp_id);
/* define the coordinate variables,... */
                              COTTIPALE + Calcal
```

CANADA

```
serial.c
nc_def_var(file_id, "Density", NC_DOUBLE, 2, densdims, &dens_id);
nc def var(file id, "Velocity", NC DOUBLE, 3, veldims, &vel id);
/* assign units to the variables */
nc put att text(file id, dens id, "units", strlen(densunit), densunit);
nc put att text(file id, vel id, "units", strlen(velunit), velunit);
/* we are now done defining variables and their attributes */
nc enddef(file id);
/* Write out the data to the variables we've defined */
nc_put_var_float(file_id, xcoord_id, x);
nc_put_var_float(file_id, ycoord_id, y);
nc_put_var_double(file_id, dens_id, &(dens[0][0]));
nc_put_var_double(file_id, vel_id, &(vel[0][0][0]));
```

```
nc_def_var(file_id, "Density", NC_DOUBLE, 2, densdims, &dens_id)
nc def var (file id, "Velocity", NC DOUBLE, 3, veldims, &vel id);
/* assign units to the variables */
nc put att text(file id, dens id, "units", strlen(densunit), dens
nc put att text(file id, vel id, "units", strlen(velunit), velu
/* we are now done defining variables and their attributes */
nc enddef(file id);
/* Write out the data to the variables we've defined */
nc_put_var_float(file_id, xcoord_id, x);
nc_put_var_float(file_id, ycoord_id, y);
/* The big data will be written to collectively;
 * the alternative is NC_INDEPENDENT */
nc var par access(file id, dens id, NC COLLECTIVE);
nc var par access(file id, vel id, NC COLLECTIVE);
/* densities */
starts[0] = (rundata.globalnx/rundata.npx)*rundata.myx;
starts[1] = (rundata.qlobalny/rundata.npy)*rundata.myy;
counts[0] = rundata.localnx;
counts[1] = rundata.localny;
nc put vara double(file id, dens id, starts, counts, &(dens[0][0]
/* velocities */
starts[0] = 0;
starts[1] = (rundata.globalnx/rundata.npx)*rundata.myx;
starts[2] = (rundata.qlobalny/rundata.npy)*rundata.myy;
counts[0] = 2;
counts[1] = rundata.localnx;
counts[2] = rundata.localny;
nc_put_vara_double(file_id, vel_id, starts, counts, &(vel[0][0][0]
nc_close(file_id);
return;
```

parallel.c

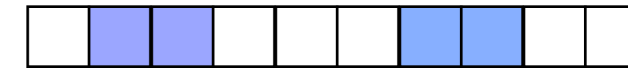
nc_close(file_id);
return;







- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride

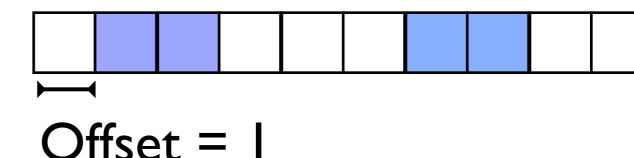








- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride



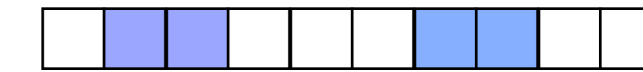






- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride

blocksize = 2

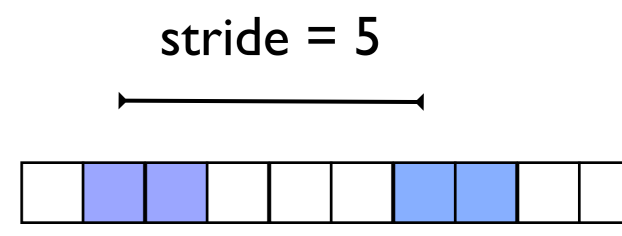








- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride



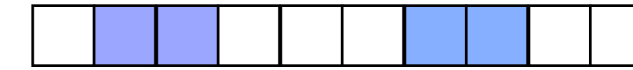






- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride
- (MPI_Type_vector)

count = 2







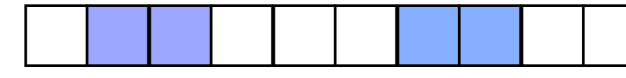


- Parallel HDF5 similar to parallel NetCDF - fairly modest changes to structure of code
- Different (more low-level, natch) way of dealing with sub-regions
- Offset, block, count, stride
- Hyperslab one of these per dimensions.
- (offset,block) just like (start, counts) in netcdf.











```
/* set the MPI-IO hints for better performance on GPFS */
MPI Info create(&info);
MPI Info set(info, "IBM largeblock io", "true");
/* Set up the parallel environment for file access*/
fap id = H5Pcreate(H5P FILE ACCESS);
/* Include the file access property with IBM hint */
H5Pset_fapl_mpio(fap_id, MPI_COMM_WORLD, info);
/* Set up the parallel environment */
dist id = H5Pcreate(H5P DATASET XFER);
/* we'll be writing collectively */
H5Pset dxpl mpio(dist id, H5FD MPIO COLLECTIVE);
```

lcul

```
/* set the MPI-IO hints for better performance on GPFS */
MPI Info create(&info);
MPI Info set(info, "IBM largeblock io", "true");
/* Set up the parallel environment for file access*/
fap id = H5Pcreate(H5P FILE ACCESS);
/* Include the file access property with IBM hint */
H5Pset_fapl_mpio(fap_id, MPI_COMM_WORLD, info);
/* Set up the parallel enveronment */
dist id = H5Pcreate(H5P DFTASET XFER);
/* we'll be writing collectively */
H5Pset dxpl mpio(dist id H5FD MPIO COLLECTIVE);
```

Same as NetCDF; this is a property of the file

```
/* set the MPI-IO hints for better performance on GPFS */
MPI Info create(&info);
MPI Info set(info, "IBM largeblock io", "true");
/* Set up the parallel environment for file access*/
fap id = H5Pcreate(H5P FILE ACCESS);
/* Include the file access property with IBM hint */
H5Pset fapl mpio(fap id, MPI COMM WORLD, info);
/* Set up the parallel environment */
dist id = H5Pcreate(H5P_DATASET_XFER);
/* we'll be writing collectively */
H5Pset_dxpl_mpio(dist_id, H5FD_MPIO_COLLECTIVE);
```

Collective/independant: this is a property of accessing a variable

lcul

```
offsets[0] = (rundata.globalnx/rundata.npx)*rundata.myx;
offsets[1] = (rundata.globalny/rundata.npy)*rundata.myy;
blocks[0] = rundata.localnx;
strides[0] = strides[1] = 1;
counts[0] = counts[1] = 1;
globaldensspace = H5Dget space(dens dataset id);
H5Sselect hyperslab(globaldensspace, H5S SELECT SET, offsets,
strides, counts, blocks);
status = H5Dwrite(dens dataset id, H5T NATIVE DOUBLE,
loc dens dataspace id, globaldensspace, dist id, &(dens[0]
[0]));
```

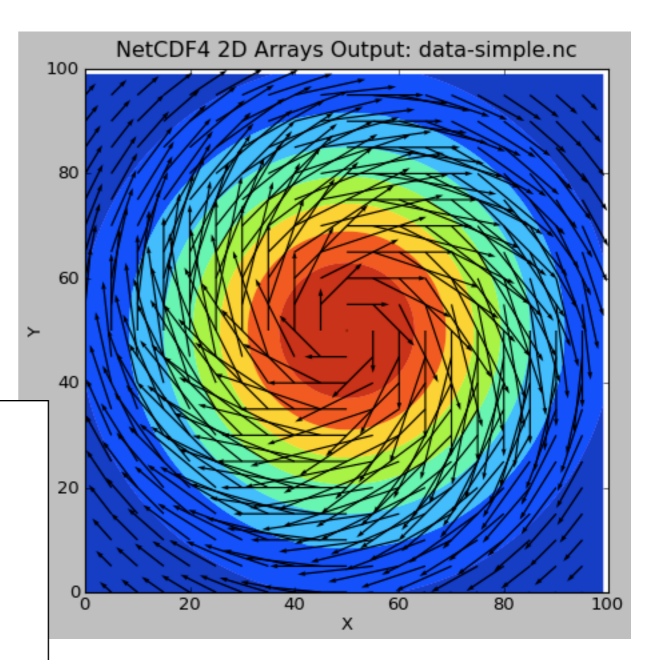
Select hyperslab, and write; parallelism is in distribution_id

Projects

\$ cd parIO/hydro{c,f}
Write hdf5, netcdf outputs

\$ cd parIO/hydro{c,f}-mpi Write ppm output in MPI-IO, (started) and output in parallel hdf5, netcdf

\$ cd parIO/nbody Write parallel hdf5, netcdf, MPI-IO outputs for gravitational particles (FORTRAN)

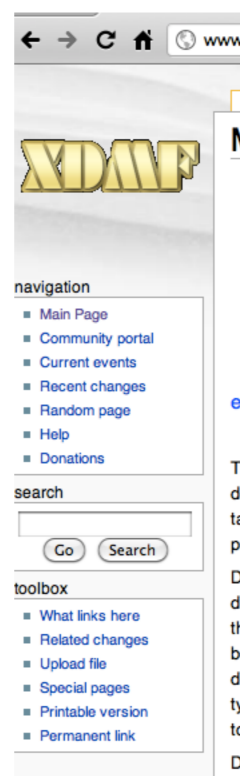






Conventions for HDF5

- XDMF
- An XML description of your HDF5 files
- A way of encoding "conventions" for HDF5
- Important for interoperatbility (eg, w/ viz packages)







eXtensible Data Model and Format

The need for a standardized method to exchange scient development of the eXtensible Data Model and Format (take advantage of widely used visualization programs like previously stand alone codes.

Data format refers to the raw data to be manipulated. Infidimensions completely describe the any dataset regardle themselves. We refer to the description of the data as **L** be passed between modules easily. Heavy data may be different nature of heavy and light data, they are stored typically stored using HDF5. While we could have chose tool to have access to the compiled HDF5 libraries in or

Data model refers to the intended use of the data. For ex geometry for a grid or calculated vector values. Without describes the data, it is purely light data and thus stored

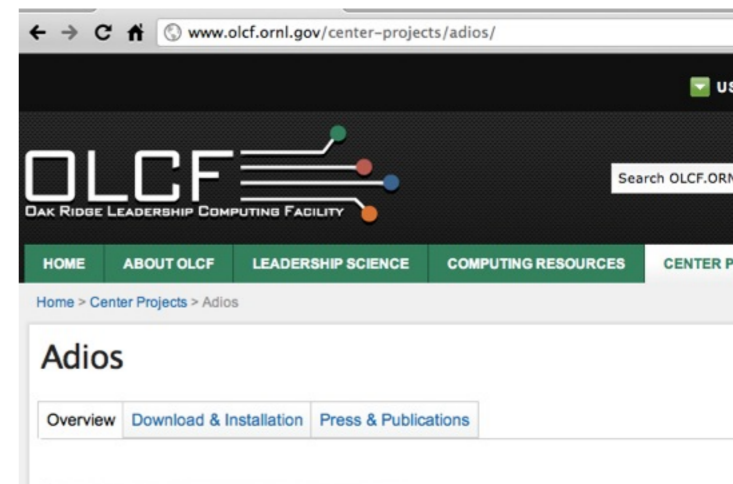






Adaptable IO System

- ADIOS
- A library for IO for scientific code
- Uses MPIIO, HDF5, etc... under the hood
- Allows changing of IO strategy, method; no rewriting code and maybe not even a recompile.





The Adaptable IO System (ADIOS) provides a simple, flexible way for scientists to desribe the data in their code that may need to be written, read, or processed outside of the running simulation. By provid an external to the code XML file describing the various elements, their types, and how you wish to process them this run, the routines in the host code (either Fortran or C) can transparently change how they process the data.

The in code IO routines were modeled after standard Fortran POSIX IO routines for simplicity and clar The additional complexity including organization into hierarchies, data type specifications, process







parlO/adios/parallel2darray.{c,f90}

```
void writeadiosfile(rundata t *rundata, double **dens, double ***vel) {
               adios err=0;
    int
               adios groupsize, adios totalsize;
   uint64 t
   int64 t adios handle;
   MPI Comm = MPI COMM WORLD;
   int size;
   MPI Comm size(comm, &size);
   adios init ("adios global.xml");
   adios open (&adios handle, "ArrayData", rundata->filename, "w", &comm);
   #include "gwrite ArrayData.ch"
    if (adios err)
        fprintf(stderr, "Error doing adios write.\n");
   adios close (adios handle);
```







parlO/adios/{,f}array_global.xml

```
<?xml version="1.0"?>
<adios-config host-language="C">
    <adios-group name="ArrayData" coordination-communicator="comm">
        <var name="rundata->localnx"
                                      type="integer" />
                                      type="integer" />
        <var name="rundata->localny"
        <var name="rundata->globalnx" type="integer" />
        <var name="rundata->globalny" type="integer" />
        <var name="rundata->startx"
                                      type="integer" />
        <var name="rundata->starty"
                                      type="integer" />
        <var name="size" type="integer" />
        <qlobal-bounds dimensions="2,rundata->qlobalnx,rundata->qlobalny"
                       offsets="0, rundata->startx, rundata->starty">
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                       dimensions="2,rundata->localnx,rundata->localny" />
        </global-bounds>
        <global-bounds dimensions="rundata->globalnx,rundata->globalny"
                       offsets="rundata->startx,rundata->starty">
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                       dimensions="rundata->localnx,rundata->localny" />
        </global-bounds>
    </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
```







ADIOS workflow

- Write XML file describing data, layout
- gpp.py [file].xml generates C or Fortran code: adios calls, size calculation
- Build code
- Separates data layout, code.





```
offsets="0, rundata->startx, rundata->st
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                        dimensions="2, rundata->localnx, rundata
        </qlobal-bounds>
        <qlobal-bounds dimensions="rundata->qlobalnx,rundata-
                        offsets="rundata->startx,rundata->star
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                        dimensions="rundata->localnx,rundata->
        </global-bounds>
    </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
void writeadiosfile(rundata t *rundata, double **dens, double
                adios err=0;
    int
                adios_groupsize, adios_totalsize;
    uint64 t
    int64 t
                adios handle;
                comm = MPI COMM WORLD;
   MPI_Comm
    int size;
   MPI_Comm_size(comm, &size);
    adios init ("adios global.xml");
    adios_open (&adios_handle, "ArrayData", rundata->filename
    #include "gwrite ArrayData.ch"
    if (adios err)
        fprintf(stderr, "Error doing adios write.\n");
    adios_close (adios_handle);
```

<adios-group name="ArrayData" coordination-communicator=</pre>

<var name="rundata->globalnx" type="integer" />
<var name="rundata->globalny" type="integer" />

<global-bounds dimensions="2,rundata->globalnx,rundat

type="integer" />

type="integer" />

type="integer" />

type="integer" />

<?xml version="1.0"?>

<adios-config host-language="C">

<var name="rundata->localnx"

<var name="rundata->startx"
<var name="rundata->starty"

<var name="size" type="integer" />

<var name="rundata->localny"



ADIOS workflow

- Separation isn't perfect; xml file references code variables, etc.
- But allows "componentization" of I/O.
- Changes that don't result in changes to grwrite_Array.ch don't require recompilation (eg, only changing number, size of variables in group).

```
<?xml version="1.0"?>
<adios-config host-language="C">
    <adios-group name="ArrayData" coordination-communicator=</pre>
        <var name="rundata->localnx"
                                       type="integer" />
                                       type="integer" />
        <var name="rundata->localny"
        <var name="rundata->globalnx" type="integer" />
        <var name="rundata->globalny" type="integer" />
                                       type="integer" />
        <var name="rundata->startx"
        <var name="rundata->starty"
                                       type="integer" />
        <var name="size" type="integer" />
        <global-bounds dimensions="2,rundata->globalnx,rundat
                        offsets="0, rundata->startx, rundata->st
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                        dimensions="2, rundata->localnx, rundata
        </global-bounds>
        <qlobal-bounds dimensions="rundata->qlobalnx,rundata-
                        offsets="rundata->startx,rundata->star
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                        dimensions="rundata->localnx,rundata->
        </global-bounds>
    </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
```

```
void writeadiosfile(rundata t *rundata, double **dens, double
                adios err=0;
   int
                adios_groupsize, adios_totalsize;
   uint64 t
   int64 t
                adios handle;
   MPI_Comm
                comm = MPI_COMM_WORLD;
   int size;
   MPI_Comm_size(comm, &size);
   adios init ("adios global.xml");
   adios_open (&adios_handle, "ArrayData", rundata->filename
   #include "gwrite ArrayData.ch"
   if (adios err)
       fprintf(stderr, "Error doing adios write.\n");
   adios_close (adios_handle);
```







Variable Groups

- Multiple groups of variables possible: (eg) restart files vs. files for analysis
- Variables can appear in mutiple groups
- Each group can be handled with different methods

```
<?xml version="1.0"?>
<adios-config host-language="C">
    <adios-group name="ArrayData" coordination-communicator=</pre>
        <var name="rundata->localnx"
                                       type="integer" />
        <var name="rundata->localny"
                                       type="integer" />
        <var name="rundata->globalnx" type="integer" />
        <var name="rundata->globalny" type="integer" />
                                       type="integer" />
        <var name="rundata->startx"
        <var name="rundata->starty"
                                       type="integer" />
        <var name="size" type="integer" />
        <global-bounds dimensions="2,rundata->globalnx,rundat
                       offsets="0, rundata->startx, rundata->st
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                       dimensions="2, rundata->localnx, rundata
        </global-bounds>
        <qlobal-bounds dimensions="rundata->qlobalnx,rundata-
                       offsets="rundata->startx,rundata->star
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                       dimensions="rundata->localnx,rundata->
        </global-bounds>
    </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
void writeadiosfile(rundata t *rundata, double **dens, double
    int
                adios err=0;
                adios_groupsize, adios_totalsize;
    uint64 t
    int64 t
                adios handle;
                comm = MPI_COMM_WORLD;
   MPI_Comm
    int size;
   MPI Comm size(comm, &size);
    adios init ("adios global.xml");
    adios_open (&adios_handle, "ArrayData", rundata->filename
    #include "gwrite ArrayData.ch"
    if (adios err)
        fprintf(stderr, "Error doing adios write.\n");
```

adios_close (adios_handle);







I/O methods

- Possible methods: parallel HDF5 (PHDF5), NetCDF (NC4), one-per-process posix files (POSIX), it's own native format (BP) using MPI-IO (MPI)
- Change between methods: edit xml file, that's it.
- P-I (PHDF5, NC4,MPI), P-P (POSIX), or even P-M possible (PHDF5, etc with multiple communicators)

```
<?xml version="1.0"?>
<adios-config host-language="C">
    <adios-group name="ArrayData" coordination-communicator=</pre>
        <var name="rundata->localnx"
                                       type="integer" />
                                       type="integer" />
        <var name="rundata->localny"
        <var name="rundata->globalnx" type="integer" />
        <var name="rundata->globalny" type="integer" />
                                       type="integer" />
        <var name="rundata->startx"
                                       type="integer" />
        <var name="rundata->starty"
        <var name="size" type="integer" />
        <global-bounds dimensions="2,rundata->globalnx,rundata
                       offsets="0,rundata->startx,rundata->st
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                       dimensions="2, rundata->localnx, rundata
        </qlobal-bounds>
        <global-bounds dimensions="rundata->globalnx,rundata-
                       offsets="rundata->startx,rundata->star
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                       dimensions="rundata->localnx,rundata->
        </global-bounds>
    </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
void writeadiosfile(rundata t *rundata, double **dens, double
                adios err=0;
    int
                adios_groupsize, adios_totalsize;
    uint64 t
    int64 t
                adios handle;
                comm = MPI_COMM_WORLD;
   MPI_Comm
    int size;
   MPI Comm size(comm, &size);
    adios init ("adios global.xml");
    adios_open (&adios_handle, "ArrayData", rundata->filename
   #include "gwrite ArrayData.ch"
    if (adios err)
        fprintf(stderr, "Error doing adios write.\n");
```

adios_close (adios_handle);







Simplifies 10 code

- Even if you aren't planning to switch between IO strategies, can greatly simplify code
- Many mechanical steps (eg, pasting together rectangular multi-dimentional arrays) done for you.
- Eliminates tedious, error-prone boilerplate code

```
<adios-group name="ArrayData" coordination-communicator=</pre>
        <var name="rundata->localnx"
                                       type="integer" />
        <var name="rundata->localny"
                                       type="integer" />
        <var name="rundata->globalnx" type="integer" />
        <var name="rundata->globalny" type="integer" />
        <var name="rundata->startx"
                                       type="integer" />
                                       type="integer" />
        <var name="rundata->starty"
        <var name="size" type="integer" />
        <qlobal-bounds dimensions="2,rundata->qlobalnx,rundat
                       offsets="0,rundata->startx,rundata->st
           <var name="vel" gwrite="vel[0][0]" type="double"</pre>
                       dimensions="2, rundata->localnx, rundata
        </global-bounds>
        <qlobal-bounds dimensions="rundata->globalnx,rundata-
                       offsets="rundata->startx,rundata->star
           <var name="dens" gwrite="dens[0]" type="double"</pre>
                       dimensions="rundata->localnx,rundata->
        </qlobal-bounds>
   </adios-group>
<method group="ArrayData" method="PHDF5" />
<buffer size-MB="2" allocate-time="now"/>
</adios-config>
void writeadiosfile(rundata t *rundata, double **dens, double
    int
                adios err=0;
                adios_groupsize, adios_totalsize;
   uint64 t
                adios handle;
   int64 t
   MPI Comm
                comm = MPI COMM WORLD;
   int size;
   MPI Comm size(comm, &size);
    adios init ("adios global.xml");
   adios open (&adios handle, "ArrayData", rundata->filename
   #include "gwrite ArrayData.ch"
   if (adios err)
        fprintf(stderr, "Error doing adios write.\n");
    adios close (adios handle);
                                compute • calcul
```

<?xml version="1.0"?>

<adios-config host-language="C">







```
void writeadiosfile(rundata t *rundata, double **dens, double ***vel) {
               adios err=0;
    int
   uint64_t adios_groupsize, adios_totalsize;
   int64_t adios handle;
   MPI Comm = MPI COMM WORLD;
   int size;
   MPI Comm size(comm, &size);
   adios init ("adios global.xml");
   adios open (&adios handle, "ArrayData", rundata->filename, "w", &comm);
   #include "gwrite ArrayData.ch"
    if (adios err)
       fprintf(stderr, "Error doing adios write. \n");
   adios close (adios handle);
```







```
void writehdf5file(rundata t rundata, double **dens, double ***vel) {
    /* identifiers */
   hid_t file_id, arr_group_id, dens_dataset_id, vel_dataset_id;
   hid t dens dataspace_id, vel_dataspace_id;
   hid t loc dens dataspace id, loc vel dataspace id;
   hid t globaldensspace, globalvelspace;
   hid t dist id;
   hid t fap id;
   /* sizes */
   hsize t densdims[2], veldims[3];
   hsize t locdensdims[2], locveldims[3];
   /* status */
   herr t status;
    /* MPI-IO hints for performance */
   MPI Info info;
    /* parameters of the hyperslab */
   hsize t counts[3];
   hsize t strides[3];
   hsize t offsets[3];
   hsize t blocks[3];
```









```
/* set the MPI-IO hints for better performance on GPFS */
MPI Info create(&info);
MPI Info set(info, "IBM largeblock_io", "true");
/* Set up the parallel environment for file access*/
fap id = H5Pcreate(H5P FILE ACCESS);
/* Include the file access property with IBM hint */
H5Pset fapl mpio(fap id, MPI COMM WORLD, info);
/* Set up the parallel environment */
dist id = H5Pcreate(H5P DATASET XFER);
/* we'll be writing collectively */
H5Pset dxpl mpio(dist id, H5FD MPIO COLLECTIVE);
/* Create a new file - truncate anything existing, use default properties */
file id = H5Fcreate(rundata.filename, H5F ACC TRUNC, H5P DEFAULT, fap id);
/* HDF5 routines generally return a negative number on failure.
 * Should check return values! */
if (file id < 0) {
    fprintf(stderr, "Could not open file %s\n", rundata.filename);
    return;
```







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```
/* Create a new group within the new file */
arr group id = H5Gcreate(file id, "/ArrayData", H5P DEFAULT, H5P DEFAULT,
                         H5P DEFAULT);
/* Give this group an attribute listing the time of calculation */
    hid t attr id, attr sp id;
    struct tm *t;
    time t now;
    int yyyymm;
    now = time(NULL);
    t = localtime(&now);
    yyyymm = (1900+t->tm year)*100+t->tm mon;
    attr sp id = H5Screate(H5S SCALAR);
    attr id = H5Acreate(arr group id, "Calculated on (YYYYMM)", H5T STD U32LE,
                        attr sp id, H5P DEFAULT, H5P DEFAULT);
    printf("yymm = %d\n",yyyymm);
    H5Awrite(attr id, H5T NATIVE INT, &yyyymm);
    H5Aclose(attr id);
    H5Sclose(attr sp id);
}
/* Create the data space for the two global datasets. */
densdims[0] = rundata.globalnx; densdims[1] = rundata.globalny;
veldims[0] = 2; veldims[1] = rundata.globalnx; veldims[2] = rundata.globalny;
```







```
dens dataspace id = H5Screate simple(2, densdims, NULL);
vel dataspace id = H5Screate simple(3, veldims, NULL);
/* Create the datasets within the file.
 * H5T IEEE F64LE is a standard (IEEE) double precision (64 bit) floating (F) data
 * and will work on any machine. H5T NATIVE DOUBLE would work too, but would give
 * different results on GPC and TCS */
dens dataset id = H5Dcreate(file id, "/ArrayData/dens", H5T IEEE F64LE,
                            dens_dataspace_id, H5P_DEFAULT, H5P DEFAULT, H5P DEFAUL
vel dataset id = H5Dcreate(file id, "/ArrayData/vel", H5T IEEE F64LE,
                            vel dataspace id, H5P DEFAULT, H5P DEFAULT, H5P DEFAUI
/* Now create the data space for our sub-regions. These are the data spaces
 * of our actual local data in memory. */
locdensdims[0] = rundata.localnx; locdensdims[1] = rundata.localny;
locveldims[0] = 2; locveldims[1] = rundata.localnx; locveldims[2] = rundata.localny
loc dens dataspace id = H5Screate simple(2, locdensdims, NULL);
loc vel dataspace id = H5Screate simple(3, locveldims, NULL);
```







```
/*
  * Now we have to figure out the `hyperslab' within the global
  * data that corresponds to our local data.
  *
  * Hyperslabs are described by an array of counts, strides, offsets,
  * and block sizes.
  *
           -offx--
  *
                                   offy
  *
                                   localny
                  localnx
     In this case the blocksizes are (localnx, localny) and the offsets are
     (offx,offy) = ((myx)/nxp*globalnx, (myy/nyp)*globalny)
  */
```







```
offsets[0] = (rundata.globalnx/rundata.npx)*rundata.myx;
offsets[1] = (rundata.globalny/rundata.npy)*rundata.myy;
blocks[0] = rundata.localnx;
blocks[1] = rundata.localny;
strides[0] = strides[1] = 1;
counts[0] = counts[1] = 1;
/* select this subset of the density variable's space in the file */
globaldensspace = H5Dget space(dens dataset id);
H5Sselect hyperslab(globaldensspace, H5S SELECT SET, offsets, strides, counts, block
/* For the velocities, it's the same thing but there's a count of two,
 * (one for each velocity component) */
offsets[1] = (rundata.globalnx/rundata.npx)*rundata.myx;
offsets[2] = (rundata.globalny/rundata.npy)*rundata.myy;
blocks[1] = rundata.localnx;
blocks[2] = rundata.localny;
strides[0] = strides[1] = strides[2] = 1;
counts[0] = 2; counts[1] = counts[2] = 1;
offsets[0] = 0;
blocks[0] = 1;
```







```
globalvelspace = H5Dget space(vel dataset id);
    H5Sselect hyperslab(globalvelspace, H5S SELECT SET, offsets, strides, counts, blocks
    /* Write the data. We're writing it from memory, where it is saved
     * in NATIVE DOUBLE format */
    status = H5Dwrite(dens dataset id, H5T NATIVE DOUBLE, loc dens dataspace id, global
dist id, &(dens[0][0]));
    status = H5Dwrite(vel_dataset_id, H5T_NATIVE_DOUBLE, loc_vel_dataspace_id, globalv
dist id, &(vel[0][0][0]));
    /* End access to groups & data sets and release resources used by them */
    status = H5Sclose(dens dataspace id);
    status = H5Dclose(dens_dataset_id);
    status = H5Sclose(vel dataspace id);
    status = H5Dclose(vel dataset id);
    status = H5Gclose(arr group id);
    status = H5Pclose(fap id);
    status = H5Pclose(dist id);
    /* Close the file */
    status = H5Fclose(file_id);
    return;
```







ADIOS hands-on:

- Modify XML file, try outputting with method of MPI (use bpls or bp2hdf on resulting file), POSIX (Netcdf won't work at this point)
- Try a different IO strategy; do contiguous parallel IO by having single file but with each process' data contiguous in file, one after another. With large file size (--nx=10000 --ny=10000) and 8/16 processes, what are the timings between "straight" PHDF5, MPI, POSIX, and this approach? (And how long would it have taken you to do this without ADIOS?)
- Advanced: Break up MPI_COMM_WORLD into 2 communicators using MPI_Comm_split, call the new communicator comm, and output 2 files from the 8/16 processes using PHDF5 or MPI.





