Using the NIH HPC Storage Systems Effectively

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https://hpc.nih.gov
Motivation

• Often get e-mail complaining about slow access to directories
• File storage systems are often over-saturated, leading to system problems that take staff time to resolve
• Users’ work is often negatively impacted (sometimes without their knowledge).
Course overview/outline

• Overview of HPC systems storage
  • Different areas (/home, /data, /scratch, object store)
  • Quotas and quota increases
  • Snapshots

• Understanding input and output (I/O)
  • I/O patterns
  • Introduction to data and metadata
  • A brief look at HPC storage systems

• Putting it all together – using storage effectively
  • Profiling and benchmarking your application’s I/O usage
  • Understanding when you’re generating too much I/O on the system
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• **Putting it all together – using storage effectively**
  • Profiling and benchmarking your application’s I/O usage
  • Understanding when you’re generating too much I/O on the system
The NIH HPC filesystems

Summary of file storage options

<table>
<thead>
<tr>
<th>Location</th>
<th>Creation</th>
<th>Backups</th>
<th>Space</th>
<th>Available from</th>
</tr>
</thead>
<tbody>
<tr>
<td>/home</td>
<td>network (NFS)</td>
<td>yes</td>
<td>8 GB default quota</td>
<td>B,C,H</td>
</tr>
<tr>
<td>/scratch (nodes)</td>
<td>local</td>
<td>no</td>
<td>~850 GB shared</td>
<td>C</td>
</tr>
<tr>
<td>/scratch</td>
<td>network (NFS)</td>
<td>no</td>
<td>75 TB shared</td>
<td>B,H,C</td>
</tr>
<tr>
<td>/data</td>
<td>network (GPFS/NFS)</td>
<td>no</td>
<td>100 GB default quota</td>
<td>B,C,H</td>
</tr>
</tbody>
</table>

H = helix, B = biowulf login node, C = biowulf compute nodes

- /home is small – only non-buyin filesystem backed up to tape – **No shared areas!**
  - Snapshots and off-site tape back-ups
- /data - in practice you will keep most of your working files here
  - Quota increases available with **justified need**
  - Shared data directories available on request
  - Snapshots available (less frequent than /home)
- /scratch – shared area for **temporary data**
- /lscratch – compute node local scratch; allocated with batch jobs
What are local and network file systems?

• Network file systems are accessed by sending data to one or more servers
  • The server controls the disks that store the data
  • Multiple different client computers can access the filesystems simultaneously

• Local file systems do not send data over the network
  • Disks usually directly attached to the computer the accesses the file system
  • Only that single computer accesses the disks directly
General best practices

<table>
<thead>
<tr>
<th>BAD</th>
<th>GOOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Submitting a swarm without knowing how much data it will generate</td>
<td>Run a single job, sum up the output and tmp files, and figure out if you have enough space before submitting the swarm</td>
</tr>
<tr>
<td>Directory with 1 million files</td>
<td>Directories with &lt; 5,000 files</td>
</tr>
<tr>
<td>100 jobs all reading the same 50 GB file over and over from /data/$USER/</td>
<td>Use /scratch instead, copy the file there, and have each job access the file on local disk</td>
</tr>
<tr>
<td>100 jobs all writing and deleting large numbers of small temporary files</td>
<td>Use /scratch instead, have all tmp files written to local disk</td>
</tr>
<tr>
<td>Each collaborator having a copy of data on Biowulf</td>
<td>Ask for a shared area and keep shared files there to minimize duplication</td>
</tr>
<tr>
<td>Use Biowulf storage for archiving</td>
<td>Move unused or old data back to your local system</td>
</tr>
</tbody>
</table>

We’ll be talking a lot about the “whys” behind some of these rules!
The HPC Object Store

• “Web Scale” storage
  • Highly reliable (dispersed over multiple sites)
  • Easy to expand (just add more disks)
  • Accessed via simple list, put, get, delete semantics (examples forthcoming)

• Different from file based storage systems
  • Objects are accessed by NAME, not PATH
  • Completely flat name space
  • No concept of directories, but “/” is a valid character in object names
  • Data and metadata are stored together with the object (sometimes true in file storage systems as well)

• More info: https://hpc.nih.gov/storage/object.html

• Unless otherwise noted, the rest of this class deals with file (not object) storage. Separate object storage class!
Use cases for object storage

• Read-intensive workloads
  • Object storage is much more efficient at reading than writing.
  • An *entire* object has to be re-written for each change
    • Computationally expensive to process and disperse the data
    • Lots of over-writing

• Static data
  • Related to the above
  • Data that doesn’t change often, but still used
  • E.g. reference genomics files
Understanding your disk quota

Use the `checkquota` command
- Shows all directories you have access to.
- Mailbox is only shown on helix - mail service discontinued Nov. 30.

You can also get this information from your user dashboard

Some storage systems have limits on the number of files – please keep these in mind (more on this later).
Requesting quota increases

• Default data directory quota: 100 GB

• Quota increases must be justified.
  • Explain number of files, size of files, and scientific use.
  • Don’t use “prior experience” or “my lab mate said” or “I need more”

• **No** quota increases on home directories
Quota increases: now on the user dashboard!
Quota increases: now on the user dashboard!

Can only request quota increases on data directories you own!
Quota increases: now on the user dashboard!

- Only two things to fill in now!
  - Amount of space
  - Justification
- Also need to accept storage policies
  - NO PHI/PII
  - NO BACKUPS!

HPC Disk Storage Request

This form should be completed by NIH HPC users who require additional storage space in their /data area.

There are no time limits or other restrictions placed on the use of data directory space on the Helix and Biowulf systems, but please use the space responsibly; even hundreds of terabytes won’t last forever if files are never deleted. Disk space on Helix and Biowulf should never be used as archival storage.

Location and User Information

- Location: /data/btmiller (gs2)
- Current Quota: 500 GB
- Name: Benjamin Timothy Miller
- Helix/Biowulf username: btmiller
- IC: NHLBI
- Telephone: +1 301 827 5281
- Email: btmiller@nihbi.nih.gov
- Principal Investigator: Bailey, Steven A

Space and Justification

- Additional space required: example: 250 GB
- How this space requirement was estimated: The input data for a single run of xx program is 100 MB. The output data is about 200 MB, and each run requires 100 MB of temp space. I expect to have about 500 such runs, so we’ll need 400MB*500 = 240 GB.

Policies

Data directories are not backed up to tape. Irreplaceable data should be backed up to your local storage (More info). Personally identifiable information (PII) or Protected Health Information (PHI) data cannot be stored anywhere on the NIH HPC systems (More info).

Please confirm that you are aware of these policies: [ ] Agree

Submit Reset Test (staff only)
Using space efficiently

• Instead of running to the quota request form, think about whether you can be using space more efficiently.
  • Move files back to your local computing infrastructure when you’re done processing
  • Delete any raw data or intermediate files that you’re sure you won’t need again (or that are backed up elsewhere)
  • Compress (gzip, bzip2, etc.) files when not in active use (note: not all files compress well).
    • FASTQ files should always be compressed
    • Molecular dynamics binary trajectories generally don’t compress much

• Storage space on the NIH HPC file systems is NOT to be used for archiving. However, we will be introducing a time-limited archive using the object store.
Shared data directories

- Requested from https://hpc.nih.gov/nih/shared_data_request.html
  - Now also redirects to the user dashboard.

- A new group will be created (group name must begin with a capital letter).
  - Group members must be specified

- Justification for the storage request must be given as with a personal data directory quota increase.

- Don’t open your personal data directory to world access!
  - Shared group directories should only be accessible by the group that owns them.

- Requestor becomes “group owner”
  - The only one who can request a quota increase
  - The only one who can request users be added to or removed from the group
Shared data directories and permissions

- New directories created under the top level will generally **NOT** have the SGID bit set.
- Users can override group ownership and permissions of directories that they create within the shared area.
  - Some applications do this without informing the user.
- Coordination and care among group members is needed.
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  - Some applications do this without informing the user.

Coordination and care among group members is needed.

```
helix:/spin1/users 6$ ls -ld /data/SomeLab/
drwxrws--- 3 btmiller SomeLab 512 Sep 23 11:43 /data/SomeLab/
```

Group name – same as the shared data directory name.
Shared data directories and permissions

Permissions – all group members can write at the top level. The SGID bit (“s” in the group execute permissions) indicates all files created in this directory will have a group ownership of SomeLab.

• New directories created under the top level will generally NOT have the SGID bit set.
• Users can override group ownership and permissions of directories that they create within the shared area.
  • Some applications do this without informing the user.
• Coordination and care among group members is needed.
  • Set umask to 007 so that all files/directories created become group writeable
  • Some users request that their primary group be changed to the shared group
A few notes about /scratch

• /scratch is a **network accessed, global** /scratch directory
  • Prior to mid-2015 (before Biowulf 2), /scratch was a **local** disk file system
  • The same scratch directory is available on helix, biowulf, and all of the compute nodes.
  • Files deleted if not accessed for 10 days!

• Good use-cases for /scratch
  • A temporary means of sharing data with a colleague who has a HPC account
  • Storing lightly accessed temporary files that must be accessed from multiple nodes.

• Bad use-cases for /scratch
  • Storing application temporary files that are only accessed from a single host (use /lscratch instead).
  • Storing anything that must be read and written frequently (use /data or /lscratch instead).
  • Storing valuable, hard to reproduce data (no back-ups; use /home or /data as appropriate).
Using /lscratch

• /lscratch = local scratch space on a compute node
• Allocated as a generic resource:
  • sbatch ... --gres=lscratch:100 ← allocates 100 GB
  • swarm -f swarmfile --gres=lscratch:100 ...
  • I bid for sinteractive
  • At /lscratch/$SLURM_JOBID
• Benefits
  • Local to node, no network traffic
  • Fewer users sharing it
  • New nodes have faster solid-state disks
When to use /lscratch (and when not)

• Use /lscratch when...
  • Many jobs will be independently reading in the same data file.
  • Many jobs will be independently writing out output files.
  • Many jobs will be writing out a lot of independent temporary files
  • Jobs will be doing large amounts of random I/O (more on this later)
    • Swarms that read/write a lot of I/O should almost always use lscratch!!!

• Don’t use /lscratch (or why bother) when...
  • Your job needs to write out a file visible to multiple nodes (this is rare?)
  • Your job is only reading/writing a few files and not very much data.

• If in doubt, e-mail staff@hpc.nih.gov
Workflow for /Iscratch

• Copy high-intensity input data from network directory to Iscratch
  • Don’t have 50 independent processes all reading the same input
  • Copy the input into each process’s Iscratch directory
  • Multiple copies per host? Can avoid this with clever scripting...
  • Remember to copy results back when done – Iscratch goes away after your job finishes!

• Use Iscratch for temporary files/scratch space
  • Set environment variables – e.g. $TMPDIR, $SCRATCH
  • Scratch/temp space variables are often program specific – need to know your application

• Use Iscratch as a local cache for objects from the object store.
Snapshots

• No back-ups of HPC data directories, but there are snapshots.
• A snapshot is a copy of the directory as it existed at a given point in time.

• Current snapshot retention policies:
  • /home – 7 hourly, 6 daily, 8 weekly
  • /data – 2 daily, 2 weekly
  • Weekly snapshots are taken on Sundays for NFS home directories and on Tuesdays for GPFS data directories.
Accessing snapshots

- Navigate to `.snapshot/` in your `/data` directory
  - Will NOT tab complete; need to type out the full directory name.
- Will see several subdirectories that contain read-only copies of your directory at a point in time.
- Can copy data from snapshots into your “main” home directory if, for example, you accidentally delete a file!

```
[helix:/data/btmiller 6$ ls .snapshot/
  helix:/data/btmiller 7$ ls .snapshot/weekly.2017-10-22_0015/
  biowulf-class  fscrwaler  linpack  SAPT2016.1  svn
  bwa_indices  gl_runner  mpsis  SAPT2016.1.tar.gz  swarm
  charm  gromacs  namd  sastemp  tmp
  clev-3.10.0.146 gs6_tim_iozone.tgz netapp_sunrpc_iozone sbin  trace
  clev_put.patch hpe  objtest  slurm  Trilinos
  ddn  ibdump-4.0.0-2.x86_64.rpm ofed_bug  src  zaki_test
  debian_jessie.img ibutils  prog  svn
  dist  infiniband  projects  swarm
  fio  intel  stress-ng-0.07.11  tmp
  freebayes  lammps  stress-ng-0.07.11.tar.gz
```
Snapshots are NOT back-ups

Snapshots are stored on the same physical hardware as home/data directories. Therefore, in the event of a hardware or facilities failure, the snapshots will also be lost!

You must back up irretrievable data to a local system (or put it in your /home directory if it will fit).
A guide to choosing a storage system

• /home
  • Small files that are very important, low I/O intensity
  • Not for files that need to be shared with other users

• /data
  • Bulk data files and scripts
  • Consider using shared data areas for sharing
  • Shared, parallel filesystems – intensive, single node I/O -> /lscratch.

• /scratch
  • Low I/O intensity job data that needs to be accessed from multiple nodes.
  • Good for short-term sharing.

• /lscratch
  • High I/O intensity job data that only needs to be accessed from a single node.

• Object
  • Primarily read-only data; low to medium intensity, but large capacity.
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  • I/O patterns
  • Introduction to data and metadata
  • A brief look at HPC storage systems

• Putting it all together – using storage effectively
  • Profiling and benchmarking your application’s I/O usage
  • Understanding when you’re generating too much I/O on the system
A block about blocks

• A filesystem organizes files and directories into **blocks** of data
  • This is because hard disks can only read and write information in discrete-sized chunks.

• Each non-zero length file you store takes up at least one full block on the filesystem.
  • The exact size of a block differs from filesystem to filesystem and may be chosen by the administrator who creates the filesystem.

<table>
<thead>
<tr>
<th>Name: myfile.txt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Owner: 917 (btmiller)</td>
</tr>
<tr>
<td>Group: 49 (staff)</td>
</tr>
<tr>
<td>Permissions: rw-r-r-</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Block 1</th>
<th>Block 2</th>
<th>Block 3</th>
<th>Block 4</th>
<th>Block 5</th>
<th>Block 6</th>
<th>Block 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hello H</td>
<td>PC stor</td>
<td>age exp</td>
<td>erts. H</td>
<td>you doi</td>
<td>ng? \0</td>
<td></td>
</tr>
</tbody>
</table>
Sequential vs. random access

• Sequential access: blocks in a file are read one after the other
  • Example: reading in a series of sequences from a file, one after another.
  • Sometimes referred to as “streaming”.
  • E.g. reading a FASTQ file into memory

• Random access: blocks in the file are read in a random order.
  • Common in database applications
  • Much harder for I/O systems to optimize
  • Generally MUCH slower!
  • E.g. Pulling non-adjacent sequences from a BAM file.
I/O comes in all sizes

• In addition to the pattern of I/O, we also have to be concerned with the size of I/O.

• Which do you think will be faster?
  • An application that reads 100 MB sequentially by issuing 50 read requests of 2 MB each
  • An application that reads 100 MB sequentially by issuing 5000 read requests of 20 KB each

• The first case is (usually) a lot faster
  • There’s a certain amount of overhead in performing requests.
  • Some systems will try to coalesce multiple small requests into bigger ones
  • With random I/O, this can be difficult to impossible.
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Unless you are writing your own application, you don’t have any direct control over this. However, it’s something to be aware of!
Data vs. metadata

• When we think of a file, we usually think in terms of the **data** it contains.
  • A BAM file contains sequence alignments
  • A molecular dynamics trajectory contains atomic coordinates/velocities
  • An MRI output contains an image of someone’s brain.
  • An EM image contains a picture of a cell in the body.

• **Metadata** literally means “data about data”
Data vs. metadata

• Examples of metadata
  • When the file was created/accessed
  • The sample ID from which the file was generated
  • The access permissions of the file
  • Where the data is located physically on the underlying disk
  • Can you think of more examples?
A conceptual diagram

• Remember our simple schematic...

Name: myfile.txt
Owner: 917 (btmiller)
Group: 49 (staff)
Permissions: rw-r-r-

This is (some of) the METADATA

This is the DATA in the file.
Some notes on directories

• Directories are special files that hold pointers (links) to other files.
• The more files there are in a directory, the larger amount of space the directory blocks will take up on disk
• Listing directories, resolving path names, moving files, etc. all require operations on the directory blocks.
  • The file system has to iterate through files in the directory individually
  • The bigger the directory is, the longer these operations will take.
  • This is why the HPC staff recommends having < 5000 files per directory
  • Especially true with directories that will have lots of operations happening simultaneously!
File and directory parallel access

• Reading and writing the same file from multiple parallel jobs can cause contention.
  • Especially with writing – due to the need for locks to avoid corruption

• Likewise, lots of different processes listing, creating files, etc. in the same directory is a bottleneck.

• Constant creation and deletion of files can create performance issues, particularly when multiple processes are doing it in the same directory.
Exercise: Good practice or bad practice?

• Which of the following are not good practice? Why?
  • Having 1,000,000 data files in a single directory.
  • Having separate runs of a program write output to separate files.
  • Reading a data file in once at program initiation, and then keeping the data cached in memory.
  • Using the name of a file to encode program results.
  • Having a swarm of 1,000 jobs each use the same temporary directory in /scratch.
Basic storage system architecture

• /home, /scratch, and some /data directories are on a large storage system that uses NFS.
Basic storage system architecture

• Other data directories are on systems running IBM’s General Parallel File System (GPFS)
• Use “checkquota --gpfs” to determine if any of your data directories are on GPFS.
NFS and GPFS

• NFS and GPFS have different back-end implementations, but from a user’s perspective, they work the same way.

• The systems perform similarly, though file system performance is variable dependent on how many users are accessing a given filesystem at any one time.
  • There is no significant performance advantage to using one system vs. the other.

• Main difference from a feature perspective: GPFS has access control lists (ACLs) whereas the NFS implementation we use does not.
  • ACLs are an advanced way of setting granular file/directory permissions – see https://hpc.nih.gov/storage/acls.html for more details
  • We will not discuss ACLs further (unless someone really wants to).
Figuring out where your data is stored

• You can use the “-a” flag on checkquota to see what filesystems the directories you have access to are on.
• spin1 = NFS, /gs[2-8] = GPFS
• Can also use “checkquota --gpfs"
• NEVER refer to any data directory by its absolute path (i.e. use /data/username NOT /spin1/users/username
  • The storage admins move directories for a variety of reasons, so the absolute paths can and do change.

```
[teacher@biowulf ~]$ checkquota -a
Mount Used Quota Percent Files  Limit
/gs3/users/cpo: 37.1 GB 100.0 GB 37.14% 118498 n/a
/home/teacher: 3.2 GB 8.0 GB 40.12% 52786  n/a
/spin1/users/teacher: 118.8 GB 500.0 GB 23.76% 16226 31129581
```
What you can do to avoid bottlenecks!

• Use /Iscratch whenever possible!!!
  • Since Iscratch is local to the node – avoid all network operations
  • Also, Iscratch on newer nodes is provisioned with SSDs!

• Avoid many parallel I/O operations.
  • They tend to oversaturate the disks

• Try to do I/O on large chunks
  • i.e. read and write large amounts of data
  • Less network overhead, and easier for the disk systems to optimize
  • If you’re using someone else’s code, this is difficult/impossible

• Avoid excessive metadata operations
  • Tend to be filtered to a small amount of disks/controllers.
  • This includes directory operations!
Exercise – where is the bottleneck?

• For the ”bad practices” we identified earlier (marked in red), what bottlenecks are likely to be relevant?
  • Having 1,000,000 data files in a single directory.
  • Having separate runs of a program write output to separate files.
  • Reading a data file in once at program initiation, and then keeping the data cached in memory.
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How do you know if you’re abusing the storage systems?

• This can be difficult to know, but there are a few clues
  • Very slow access to working directories involved with the job. (ls etc. take a long time to return)
  • For swarms, job completion speed was acceptable for small numbers of jobs, but gets dramatically slower as the size or number of jobs increases.
    • You did test with small-scale jobs first, right?!
  • The problem may be with another user’s jobs, but if you started seeing problems right after you started a bunch of jobs, you are the prime suspect.

• HPC staff will notify you if we notice your jobs having an impact
  • However, please be proactive and don’t wait for us to notice the problem
  • If we send you mail, it means you’re having a significant negative impact on system performance.
Refactoring a workload – general principles

• Look for places where lots of parallel processes are doing I/O
  • Think about if only one process could do I/O and communicate with other processes (probably not possible with swarm).
  • Can some or all of that I/O use /lscratch instead of /scratch or /data?

• Think about bottlenecks in the workflow
  • E.g. the whole workload has to wait until one file is updated
  • Does the usage on a shared filesystem cause delays in this process?

• Does the workflow behavior change over time?
  • Do jobs have different I/O patterns in the beginning, middle, or end of their runs.
  • Would staggering this I/O be possible?
Workload refactoring: an example

- Start of a real-world genomics pipeline
  - Aligns sequences, creates index
  - RSeQC performs QC steps before continuing the pipeline
- Pipeline ran much more quickly after workload was refactored to use I/O more efficiently!
Original Pipeline

- Lots of read and writes back to network storage.
- Many parallel processes reading the same data from the storage system.
- Depending on the exact analysis done by RSeQC, random I/O is heavy.
  - Remember, lots of RSeQC processes in parallel.
- How would you fix this?
Refactored pipeline

- Only writes to networked storage when needed.
  - E.g. not after the initial read, only when index is built.
- Instead of each RSeQC reading the data from network storage, use local scratch
- Only write out final result files.
IOPS comparison

- **Bottom line 1:** User was able to do more science in less time.
- **Bottom line 2:** HPC storage admins did not have to troubleshoot performance problems.

![IOPS graph](image)

- **Run 1** [10 samples]
- **Run 2** [20 samples]
staff@hpc.nih.gov
Wrap-up; Q&A

• Thank you for coming!
  • We hope you are able to apply the lessons learned to your own particular storage issues.
  • PLEASE reach out to staff@hpc.nih.gov for assistance; we’d love to work with you **proactively** instead of **reactively**.

• Please provide feedback on this presentation!
  • E-mail Tim btmiller@helix.nih.gov
  • E-mail Mark patkus@helix.nih.gov
  • General questions staff@hpc.nih.gov
Upcoming Seminars

• November 30, 1 - 3 pm  
  **Python in HPC**  
  Overview of python tools used in high performance computing, and how to improve the performance of your python jobs on Biowulf

• Jan 16, 1 - 3 pm  
  **Relion tips and tricks, and Parallel jobs and benchmarking**  
  Mechanics and best practices for submitting RELION jobs to the batch system from both the command line and via the RELION GUI, as well as methods for monitoring and evaluating the results. Scaling of parallel jobs, how to benchmark to make effective use of your allocated resources