

Hiding large numbers of files in container overlays

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Zoom controls

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- To ask questions at any time, type in Chat, or Unmute to ask via audio
 - please address chat questions to "Everyone" (not direct chat!)
- Raise your hand in Participants



- Email training@westdri.ca
- Our winter/spring training schedule <https://bit.ly/wg2023a>
 - biweekly webinars
 - weekly online courses starting in February
 - in-person workshops @SFU @UBC
 - in-person spring/summer schools @UVic @SFU

Many files on a distributed filesystem (DFS) such as Lustre

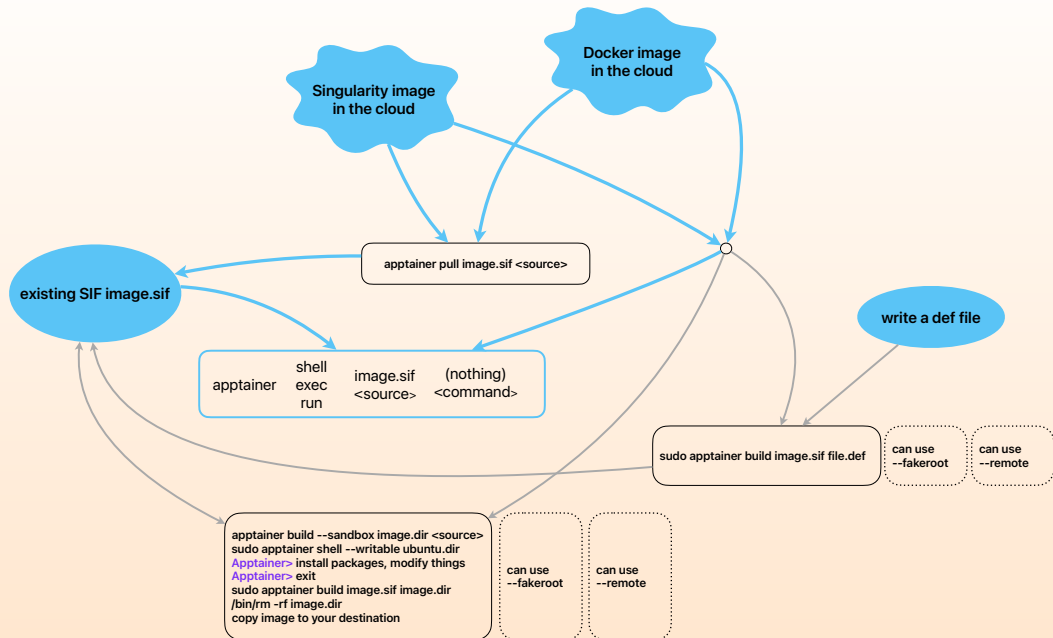
- Many of our cluster filesystems are Lustre-based
 - https://docs.alliancecan.ca/wiki/Storage_and_file_management#Filesystem_quotas_and_policies
 - highly scalable: up to hundreds of servers, tens of PBs, over a TB/s of aggregate throughput
 - employ multiple metadata + object storage servers that manage access to blocks stored on thousands of drives
 - multiple clients (individual cluster nodes) can access the DFS through the network
 - all clients see a unified namespace for all of the files and data in the filesystem
 - multiple levels of cache + data replication to improve I/O speeds and keep data highly available
 - ✓ One of the design goals: reach high bandwidth for large I/O operations (= reading/writing few large, unbroken data streams)
 - ✗ Having a large number of small files can lead to poor performance and strain the metadata server(s)
- ⇒ for best performance minimize the number of open/close operations on the filesystem
- ⇒ in other words, avoid high IOPS workflows (small reads/writes, writing many small files)

Possible solutions

- If you already have many small files:
 - use archival tools such as tar, DAR to pack them into few archive files
 - https://docs.alliancecan.ca/wiki/Handling_large_collections_of_files
 - <https://docs.alliancecan.ca/wiki/Dar>
 - only a partial solution ...
- Modify your code to streamline output to fewer, larger files
- Use local (to the node) temporary storage: \$SLURM_TMPDIR + copy out at the end
- Use an SQL database inside your code
 - does not apply to most workflows?
 - upcoming Feb-28 webinar
- ✓ Use container overlays
 - an overlay image is a file formatted as a filesystem
 - a container sees files inside, while the host OS sees only one large overlay image file
 - if your code does lots of small reads/writes, these are cached by the OS and Apptainer software before reaching the host's filesystem
- And few others ...

Singularity / Apptainer containers

- Create a custom, secure virtual Linux environment (a container) that is different from the host Linux system, uses
 - uses kernel namespaces to virtualize and isolate OS resources (CPU, memory access, disk I/O, network access, user/group namespaces), so that processes inside the container see only a specific, virtualized set of resources
 - Linux control groups (cgroups) to control and limit the use of these resources
 - overlay filesystems to enable the appearance of writing to otherwise read-only filesystems
- Much more lightweight than a VM, as containers use many resources of the host's OS
- In a sense, give you control of your software environment without being root on the host system
 - with a catch: building containers from scratch usually requires root
- Ideal way to create a portable software environment: scientific software + all the dependencies



Creating and mounting an overlay

- A writable overlay sits on top of an immutable SIF container image, can be:
 - a sandbox directory (does not work for our purposes)
 - ✓ a standalone writable ext3 filesystem image
 - a writable ext3 image embedded into the SIF file (less flexible)

```
cd ~/scratch/containers
module load apptainer/1.1.3
apptainer pull ubuntu.sif docker://ubuntu:latest
apptainer overlay create --size 512 small.img # create a 0.5GB overlay image file

apptainer shell --overlay ./small.img ubuntu.sif
Apptainer> df -kh # the overlay should be mounted inside the container
Apptainer> mkdir -p /data # a new root-level dir will go into the overlay image
Apptainer> cd /data && df -kh . # using overlay; check for available space
Apptainer> for num in $(seq -w 00 19); do
    echo $num # next line generates a binary file (1-33)MB in size
    dd if=/dev/urandom of=test"$num" bs=1024 count=$(( RANDOM + 1024 ))
done
Apptainer> df -kh . # should take ~350 MB
Apptainer> exit

apptainer shell --overlay ./small.img ubuntu.sif
Apptainer> ls /data # here is your data
```

Creating and mounting an overlay (cont.)

- You can create an overlay image with a directory inside with something like this:

```
apptainer overlay create --create-dir /data --size 512 overlay.img
```

- You can mount the overlay in the read-only mode:

```
apptainer shell --overlay ./small.img:ro -B /home,/scratch ubuntu.sif
Apptainer> touch /data/test.txt # error: read-only file system
```

- To see the help page:

```
apptainer help overlay create
```

- Outside the container, when an overlay is *not* in use, you can resize it:

```
e2fsck -f small.img # good idea to check your overlay's filesystem first
resize2fs -p small.img 1G # resize your overlay
```


Sparse overlay images

- Empty blocks inside are not stored ⇒ use disk more efficiently
- Be careful using them: not all tools (e.g. backup/restore, scp, sftp, gunzip) recognize sparse files! ⇒ this can potentially lead to very bad things

1. Create a sparse overlay image:

```
apptainer overlay create --create-dir /data --size 512 --sparse sparse.img
ls -l sparse.img                # its apparent size is 512MB
du -h --apparent-size sparse.img # same
du -h sparse.img                # its actual size is much smaller (17MB)
```

2. Mount it and fill with some data:

```
apptainer shell --overlay sparse.img ubuntu.sif

Apptainer> cd /data
Apptainer> for num in $(seq -w 00 19); do
    echo $num      # next line generates a binary file (1-33)MB in size
    dd if=/dev/urandom of=test"$num" bs=1024 count=$(( RANDOM + 1024 ))
done
Apptainer> df -kh .          # should take just under 400MB, pay attention to "Used"

du -h sparse.img           # actual usage is now just under 400MB
```

Conda apptainer

- One possible use: install Conda into an overlay image
 - native Anaconda on our clusters is a bad idea <https://docs.alliancecan.ca/wiki/Anaconda/en>
 - installing into an overlay image takes a couple of minutes, results in over 17,000 files
 - **no need for root**, as you don't modify the container!
 - might still **not be the most efficient use** of resources (non-optimized binaries)

```
cd ~/scratch/containers
apptainer pull ubuntu.sif docker://ubuntu:latest
apptainer overlay create --size 550 --sparse conda.img # create a 550MB overlay image
wget https://repo.anaconda.com/miniconda/Miniconda3-py39_4.12.0-Linux-x86_64.sh # to host

apptainer shell --overlay ./conda.img -B /scratch ubuntu.sif
Apptainer> mkdir /conda && cd /conda && df -kh .
Apptainer> bash ~/scratch/containers/Miniconda3-py39_4.12.0-Linux-x86_64.sh
    agree to the license
    use /conda/miniconda3 for the installation path
    "no" to initialize Miniconda3, to avoid modifying your host's ~/.bashrc
Apptainer> find /conda/miniconda3/ -type f | wc -l # 17,722 files

du -h conda.img # uses 474M
apptainer shell --overlay ./conda.img -B /scratch ubuntu.sif
Apptainer> source /conda/miniconda3/bin/activate
Apptainer> python # works
```

Conda apptainer: installing Python packages into the overlay

```
# need more space if installing large packages
apptainer overlay create --size 2000 --sparse conda.img # need almost 2G for numpy
apptainer shell --overlay ./conda.img -B /scratch ubuntu.sif
... install Conda as described in the previous slide ...

# -C flag important to force writing config files locally, and not to the host
apptainer shell -C --overlay ./conda.img -B /scratch ubuntu.sif
Apptainer> cd /conda/miniconda3
Apptainer> source bin/activate
Apptainer> conda install numpy
Apptainer> du -skh . # used 1.5G
Apptainer> python
>>> import numpy as np
```

MPI codes in a container

There are fundamentally three different ways to run parallel MPI codes in a container:

1. Rely on **MPI inside the container**, e.g.

```
apptainer exec -B ... --pwd ... container.sif mpirun -np $SLURM_NTASKS ./mpicode
```

- single Apptainer process running on the node
- cgroup limitations from the Slurm job are passed into the container ⇒ set the number of available CPU cores
⇒ mpirun uses all available (in this job) CPU cores



limited to a single node



no need to adapt container's MPI to the host; just install SSH into the container

2. **Hybrid mode**: rely on mpirun on the host, and MPI inside the container to provide runtime MPI libraries, e.g.

```
mpirun -np $SLURM_NTASKS apptainer exec -B ... --pwd ... container.sif ./mpicode
```

- separate Apptainer process per each MPI rank



can span multiple nodes



container's MPI should be customized to the host (not that difficult)

3. **Bind mode** - not used in this presentation

Option 1: build a container with precompiled OpenFOAM

- Easy:

1. start with a stock `docker://ubuntu` image – already includes MPI
2. install `openfoam10` package from `http://dl.openfoam.org/ubuntu`

- Works well with MPI inside the container, can scale OpenFOAM up to an entire node

- In general, will not play nicely with host's MPI and Slurm, so will **not** be able to do

```
mpirun -np $SLURM_NTASKS apptainer exec -B ... --pwd ... container.sif ./mpicode
```

- Instructions on building the container, preparing and running a small parallel OpenFOAM simulation while writing output to an overlay

<https://github.com/razoumov/sharedSnippets/blob/main/openfoamPrecompiled.md>

Option 2: build a host-compatible OpenFOAM container (that can work in either mode)

1. Build an MPI container (that can talk to the host's MPI)

2. Run a test code with MPI **entirely inside the container**
3. Compile parallel OpenFOAM into this container
4. Run parallel OpenFOAM **with container's MPI** and with the case directory (and results) inside an overlay - limited to a single node

5. Run a test code with **hybrid approach**: host's MPI to launch the code, container's MPI for runtime libraries
6. Run parallel OpenFOAM in **hybrid mode**: organize output into multiple overlays
7. Post-process the results in the overlays

Step 1: build an MPI container

(that's compatible with host's Slurm and MPI)

- On the host, find out the supported MPI model:

```
srun --mpi=list # list the supported PMI implementations
scontrol show config | grep MpiDefault # show the default
```

e.g. on Arbutus Slurm supports pmi2 ⇒ inside the container build OpenMPI with pmi2 support

- Create a definition file that:

- bootstraps from `docker://ubuntu:22.04`
- installs the necessary fabric and PMI packages, Slurm client, few others requirements
- compiles the latest OpenMPI from source supporting hosts's PMI implementation
- instructions for building a container to run on Cedar, Magic Castle clusters on Arbutus
<https://github.com/razoumov/sharedSnippets/blob/main/mpiContainer.md>

- Build the container

```
sudo apptainer build mpi.sif parallelContainer.def
ls -lh mpi.sif # 294M
scp mpi.sif username@cluster:path
```

Step 2: test a simple code inside the container

- Run a test code with container's `mpirun`

```
cd ~/scratch/containers/  
module load openmpi apptainer/1.1.3  
apptainer exec -B /scratch mpi.sif mpicc -O2 distributedPi.c -o distributedPi  
salloc --nodes=1 --ntasks=3 --time=0:5:0 --mem-per-cpu=3600  
apptainer exec -B /scratch mpi.sif mpirun -np $SLURM_NTASKS ./distributedPi
```


Step 3: compile parallel OpenFOAM into this container

OpenFOAM v2012 has a very subtle bug, resulting in an MPI deadlock in some of the runs with overlays ⇒ using the previous major release v1912

```
sudo apptainer build --sandbox 1912.dir mpi.sif # convert to a sandbox
sudo apptainer shell --writable 1912.dir
```

```
mkdir -p /opt/openfoam && cd /opt/openfoam
wget https://dl.openfoam.com/source/v1912/OpenFOAM-v1912.tgz
tar xvfz OpenFOAM-v1912.tgz && /bin/rm OpenFOAM-v1912.tgz
cd OpenFOAM-v1912
apt install -y flex libz-dev
WM_COMPILER_TYPE=ThirdParty
source etc/bashrc
./Allwmake -j 8 # took 1h35m
exit
```

```
sudo apptainer build openfoam1912.sif 1912.dir # rebuild the container
scp openfoam1912.sif username@cluster:path
```

Step 4: prepare and run parallel OpenFOAM with container's MPI

and with the case directory (and results) inside an overlay - limited to a single node

```
module load openmpi apptainer/1.1.3  &&  cd ~/scratch/containers
apptainer overlay create --size 100 --create-dir /results output.img
apptainer shell -B /scratch --overlay output.img openfoam1912.sif

. /opt/openfoam/OpenFOAM-v1912/etc/bashrc
cd /results
cp -r $FOAM_TUTORIALS/incompressible/simpleFoam/motorBike .  # parallel case
cd motorBike
mv system/decomposeParDict.6 system/decomposeParDict
>>> edit system/decomposeParDict
    numberOfSubdomains  3;
    n (3 1 1);
>>> edit system/controlDict
    endTime             500;
    writeInterval       10;
    writeFormat         binary;
mv 0.orig 0
blockMesh              # generate the mesh
decomposePar          # decompose the mesh and the ICs
exit

export OPENFOAM=/opt/openfoam/OpenFOAM-v1912/platforms/linux64GccDPInt32Opt
export APPTAINERENV_PREPEND_PATH=$OPENFOAM/bin
export APPTAINERENV_LD_LIBRARY_PATH=$OPENFOAM/lib:$OPENFOAM/lib/openmpi-system
export APPTAINERENV_WM_PROJECT_DIR=/opt/openfoam/OpenFOAM-v1912

salloc --nodes=1 --ntasks=3 --time=0:10:0 --mem-per-cpu=3600
apptainer exec -B /scratch --overlay output.img --pwd /results/motorBike openfoam1912.sif \
  mpirun -np $SLURM_NTASKS simpleFoam -parallel
exit the job
```

Step 4 (cont.): analyze results

```
salloc --ntasks=1 --time=0:10:0 --mem-per-cpu=3600
apptainer shell -B /scratch --overlay output.img openfoam1912.sif

cd /results/motorBike
ls processor*
find processor? -type f | wc -l          # 1542 decomposed files including the setup and ICs
reconstructPar                          # merge decomposed files
find . -type f | wc -l                  # 2476 files in total
rm -rf processor???? processor?/?      # delete the output decomposed files
find . -type f | wc -l                  # still 976 files including 50 outputs with 10 files each
exit the container

exit the job
```

- A typical production run would have ~ 1500 timesteps \times 128 cores \times ~ 10 files/core/step
⇒ 1,920,000 decomposed files ⇒ great to have these inside an overlay
- How about scaling to multiple nodes?

Complication

- To scale to multiple nodes, you will use host's MPI, launching a separate container per each MPI rank

```
mpirun -np $SLURM_NTASKS apptainer exec -B /scratch --pwd ... \  
openfoam1912.sif simpleFoam -parallel
```

- ... but you **cannot mount the same overlay file in multiple containers at the same time**
... especially if these containers are spread across multiple nodes

Complication

- To scale to multiple nodes, you will use host's MPI, launching a separate container per each MPI rank

```
mpirun -np $SLURM_NTASKS apptainer exec -B /scratch --pwd ... \  
openfoam1912.sif simpleFoam -parallel
```

... but you **cannot mount the same overlay file in multiple containers at the same time**
... especially if these containers are spread across multiple nodes

- Solution: use a separate overlay image for each container (i.e., each MPI rank) (idea picked up from <https://pawseysc.github.io>)
 - single output (case) directory on the host's filesystem
 - inside it, use symbolic links to redirect output from each MPI rank to its own overlay
- ⇒ this will trick the simulation code into thinking that it's writing into a uniform filesystem

Step 5: run a test code with hybrid approach

(host's MPI to launch the code, container's MPI for runtime libraries)

```
mkdir -p ~/.openmpi
echo "btl_vader_single_copy_mechanism=none" >> ~/.openmpi/mca-params.conf
module load openmpi apptainer/1.1.3
export PMIX_MCA_psec=native # allow mpirun to use host's PMI
cd ~/scratch/containers
apptainer exec -B /scratch mpi.sif mpicc -O2 distributedPi.c -o distributedPi
salloc --ntasks=2 --time=0:5:0 --mem-per-cpu=3600
mpirun -np $SLURM_NTASKS apptainer exec -B /scratch mpi.sif ./distributedPi
```

Step 5: run a test code with hybrid approach

(host's MPI to launch the code, container's MPI for runtime libraries)

```
mkdir -p ~/.openmpi
echo "btl_vader_single_copy_mechanism=none" >> ~/.openmpi/mca-params.conf
module load openmpi apptainer/1.1.3
export PMIX_MCA_psec=native # allow mpirun to use host's PMI
cd ~/scratch/containers
apptainer exec -B /scratch mpi.sif mpicc -O2 distributedPi.c -o distributedPi
salloc --ntasks=2 --time=0:5:0 --mem-per-cpu=3600
mpirun -np $SLURM_NTASKS apptainer exec -B /scratch mpi.sif ./distributedPi
```

Later we'll use \$SLURM_PROCID variable from individual ranks

⇒ for that we'll use srun

```
# this breaks b/c of an environment bug
srun -n $SLURM_NTASKS apptainer exec -B /scratch mpi.sif ./distributedPi

# this works
echo "apptainer exec -B /scratch mpi.sif ./distributedPi" | srun -n $SLURM_NTASKS bash
# this prints each rank's ID
echo "apptainer exec mpi.sif echo '$$SLURM_PROCID' | srun -n $SLURM_NTASKS bash
```

Step 6a: run OpenFOAM in hybrid mode - prepare the simulation

```
cd ~/scratch/containers
module load openmpi apptainer/1.1.3
export PMIX_MCA_psec=native
/bin/rm -rf ~/scratch/OpenFOAM/*
apptainer shell -B /scratch --pwd ~/scratch/OpenFOAM openfoam1912.sif

. /opt/openfoam/OpenFOAM-v1912/etc/bashrc
cp -r $FOAM_TUTORIALS/incompressible/simpleFoam/motorBike .      # parallel case
cd motorBike                                                    # host's filesystem
mv system/decomposeParDict.6 system/decomposeParDict
>>> edit system/decomposeParDict
  numberOfSubdomains  3;
  n (3 1 1);
>>> edit system/controlDict
  endTime             500;
  writeInterval       10;
  writeFormat         binary;
  runtimeModifiable  false;  # disable loading of dictionaries at each timestep
mv 0.orig 0
blockMesh             # generate the mesh
decomposePar          # decompose the mesh (and the ICs)
exit
```


Step 6b: replace output dirs with links to equivalents in MPI overlays

```
cd ~/scratch/OpenFOAM/motorBike      # simulation directory on the host's filesystem
mkdir -p bak && mv processor* bak     # move them out of the way for now

cd ~/scratch/containers
# create a 100MB overlay
apptainer overlay create --size 100 --create-dir /results/motorBike results0.img
for rank in {1..2}; do
    cp results0.img results${rank}.img
done

for rank in {0..2}; do # for each processor copy the initial conditions and the mesh
    apptainer exec -B /scratch --overlay results${rank}.img openfoam1912.sif \
        cp -r ~/scratch/OpenFOAM/motorBike/bak/processor${rank} /results/motorBike
done

cd ~/scratch/OpenFOAM/motorBike
for rank in {0..2}; do
    # these links will be broken on the host
    # but they will work inside each rank's container
    ln -s /results/motorBike/processor${rank} processor${rank}
done
```

How to see content inside output directories?

```
cd ~/scratch/containers
apptainer shell --overlay results0.img -B /scratch \
  --pwd ~/scratch/OpenFOAM/motorBike openfoam1912.sif
ls processor*    # only processor0 will show files
exit
>>> the same when mounting results1.img - only processor1 will show files, and so on ...

cd ~/scratch/containers
salloc --ntasks=3 --time=0:60:0 --mem-per-cpu=3600
# look inside each rank's directory
echo 'apptainer exec --overlay results${SLURM_PROCID}.img -B /scratch \
  --pwd ~/scratch/OpenFOAM/motorBike openfoam1912.sif \
  ls processor${SLURM_PROCID}/' | srun -n $SLURM_NTASKS bash
```

Step 6c: run the simulation

```
export OPENFOAM=/opt/openfoam/OpenFOAM-v1912/platforms/linux64GccDPInt32Opt
export APPTAINERENV_PREPEND_PATH=$OPENFOAM/bin
export APPTAINERENV_LD_LIBRARY_PATH=$OPENFOAM/lib:$OPENFOAM/lib/openmpi-system
export APPTAINERENV_WM_PROJECT_DIR=/opt/openfoam/OpenFOAM-v1912
```

```
salloc --ntasks=3 --time=0:30:0 --mem-per-cpu=3600
```

```
echo 'apptainer exec --overlay results${SLURM_PROCID}.img -B /scratch \
--pwd ~/scratch/OpenFOAM/motorBike openfoam1912.sif simpleFoam \
--parallel' | srun -n $SLURM_NTASKS bash
```

```
echo 'apptainer exec --overlay results${SLURM_PROCID}.img -B /scratch \
--pwd ~/scratch/OpenFOAM/motorBike openfoam1912.sif \
ls processor${SLURM_PROCID}/' | srun -n $SLURM_NTASKS bash # check the results
```

Step 7: postprocessing – reconstruct ...

We can't mount more than one writable overlay per container ... but we can mount
(1) multiple overlays as read-only + (2) one writable overlay:

```
cd ~/scratch/containers
apptainer overlay create --size 300 --create-dir /analyze reconstructed.img
salloc --time=0:30:0 --mem-per-cpu=3600
apptainer shell \
  --overlay $(echo results{0..2}.img:ro | tr ' ' ,) \      # per-rank results
  --overlay reconstructed.img \                          # reconstructed results
  -B /scratch --pwd ~/scratch/OpenFOAM/motorBike openfoam1912.sif

ls processor*      # see output from all ranks
mv * /analyze/     # fast: not moving the results, just the initial setup + links
cd /analyze

reconstructPar
exit
apptainer shell --overlay reconstructed.img -B /scratch --pwd analyze openfoam1912.sif

ls                # here are your reconstructed results
ls processor*/    # these links are now are broken
```

Step 7: postprocessing – ... or not

1. Install/compile parallel ParaView server into another MPI container, call it `paraview.sif`
2. Run parallel ParaView server

```
cd ~/scratch/containers
salloc --ntasks=3 --time=0:60:0 --mem-per-cpu=3600
echo 'apptainer exec --overlay results${SLURM_PROCID}.img -B /scratch \
  --pwd ~/scratch/OpenFOAM/motorBike paraview.sif \
  pvserver --force-offscreen-rendering' | srun -n $SLURM_NTASKS bash
```

3. Take note of the listening node's name and port number

```
ssh <username>@<cluster>.computecanada.ca -L 11111:<node>:<port>
```

4. Connect from the ParaView client on your computer to `localhost:11111`
5. Create an empty `case.foam` file in the case directory
6. Load `case.foam`, set Case Type = Decomposed

Summary

- Overlays give us a great way to add writable layers to a container, no need for root to compile software into an overlay
- Can mix and match containers and overlays
- Only one limitation: you can mount only one writable overlay per container at a time
 - ⇒ can mount many as read-only
 - ⇒ can mount one writable overlay per MPI rank (if running one container per MPI rank)
- The examples in these slides shown with `salloc`, but of course all production runs should be scheduled with `sbatch` and proper Slurm scripts
- In a parallel run, what if all output is happening on a single MPI rank?
 - ⇒ mount the overlay on this rank, need only one overlay
- In a parallel run, what if all your processors write their output into the same directory?
 - well, could they be doing parallel I/O, writing into a few large files?
 - if not, and you have the many-files problem ⇒ probably there is a custom solution for each code, talk to us
 - or modify your code's output, so that it does not create the problem in the first place, or make it write output files into per-processor directories + apply the solution from this presentation

Questions?