

Using Containers on Minerva and Accelerating Genome Analysis with Parabricks

Minerva Scientific Computing Environment

<https://labs.icahn.mssm.edu/minervalab>

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Outline

- ▶ Singularity/Apptainer access on Minerva
- ▶ Capabilities and Performance of Parabricks
- ▶ Parabricks for secondary analysis

Containers

- ▶ A standard unit of software that packages up code and all its dependencies, so the application runs quickly and reliably from one computing environment to another.

By-function containers provide:

- ▶ Software bundles for applications
- ▶ Self contained environment
- ▶ Platform/Host agnostic

<https://www.docker.com/resources/what-container>

<https://portal.biohpc.swmed.edu/content/guides/singularity-containers-biohpc/>

Host kernel
Filesystems /sc/arion, /home
Devices and drivers
Applications, Gnome Desktop, /usr/bin
Modules
...

OWN RootFS /
Applications
and dependencies

Container Namespace



Singularity on Minerva

Use singularity module on Minerva nodes

```
$ ml av singularity  
$ module load singularity/3.6.4
```

Pull image from Docker Hub [docker://](#), and Sylabs Cloud [library://](#)

```
$ singularity pull docker://gcc:10  
  
$ ls -l  
-rwxr-xr-x 1 yuj25 hpcstaff 396431360 Oct  4 17:04 gcc_10.sif  
  
$ singularity pull docker://rocker/rstudio:4.4.3
```

Convert Docker image to Singularity image

```
$ singularity build ollama.sif docker://ollama/ollama:latest
```

Singularity on Minerva

Images layers are cached in \$HOME/.singularity/cache/, may blow up your

\$HOME quota

```
$ singularity cache list -v
NAME                                DATE CREATED      SIZE              TYPE
0e3f4c426c9e5994ac625c            2021-04-23 16:46:57  440.43 MB        blob
0f46f97746e4df5959e8c8            2021-04-26 12:57:43  213.09 MB        blob

$ singularity cache clean
```

You can change the cache directory by specifying the SINGULARITY_CACHEDIR environment parameter:

```
$ SINGULARITY_CACHEDIR=/sc/arion/scratch/yuj25/containers/cache \
singularity pull docker://gcc:10

INFO:    Converting OCI blobs to SIF format
INFO:    Starting build...
Getting image source signatures
Copying blob 723254a2c089 done
Copying blob abe15a44e12f done
Copying blob 409a28e3cc3d done
```

Singularity on Minerva

Run interactively inside the image

```
$ singularity shell gcc_10.sif
Singularity> gcc -v
...
gcc version 10.4.0 (GCC)
Singularity> exit
exit
$ gcc -v
...
gcc version 4.8.5 20150623 (Red Hat 4.8.5-36) (GCC)
```

Run a custom command with exec

```
$ singularity exec gcc_10.sif gcc -v
...
gcc version 10.4.0 (GCC)
```

Singularity on Minerva

Run a container, with default runscript command

```
$ singularity pull library://sylabased/examples/lolcow
```

```
$ singularity run library://sylabased/examples/lolcow
```

```
/ Q: How many elephants can you fit in a \
| VW Bug? A: Four. Two in the front, two |
| in the back.                           |
```

```
| Q: How can you tell if four elephants |
| are in your refrigerator? A: There's a |
\ VW Bug in your driveway.              /
```

```
-----
 \   ^__^
  \  (oo)\_______
      (_____)\/    )\/\
          ||----w |
          ||     ||
```

Apptainer on Minerva

- ▶ Apptainer is the community-driven continuation of Singularity, maintained under the Linux Foundation
- ▶ It provides the same functionality and syntax as Singularity while offering better security, integration, and compatibility with HPC systems

```
$ml apptainer/1.3.6
$apptainer pull library://sylabstd/examples/lolcow
$apptainer run lolcow_latest.sif
```

```
/ It is easy to find fault, if one has \
| that disposition. There was once a man |
| who, not being able to find any other |
| fault with his coal, complained that |
| there were too many prehistoric toads |
| in it.                                |
|                                       |
| -- Mark Twain, "Pudd'nhead Wilson's  |
\ Calendar"                             /
```

```
-----
\      ^__^
\      (oo)\_______
          (_____)\/    )\ /\
              ||----w |
              ||     ||
```

```
$apptainer cache list -v
$apptainer shell
```

What is NVIDIA Parabricks?

A software suite designed to accelerate genomic data analysis by leveraging GPU (graphics processing unit) computing.

- High-speed genomic analysis
 - Integrates with widely used genomics analysis tools and pipelines (such as GATK)
 - Compatible with multiple bioinformatics formats
 - Preserves the accuracy of standard CPU-based genomics workflows
 - Compatible with common workflow managers WDL and NextFlow
- (<https://github.com/clara-parabricks-workflows>)

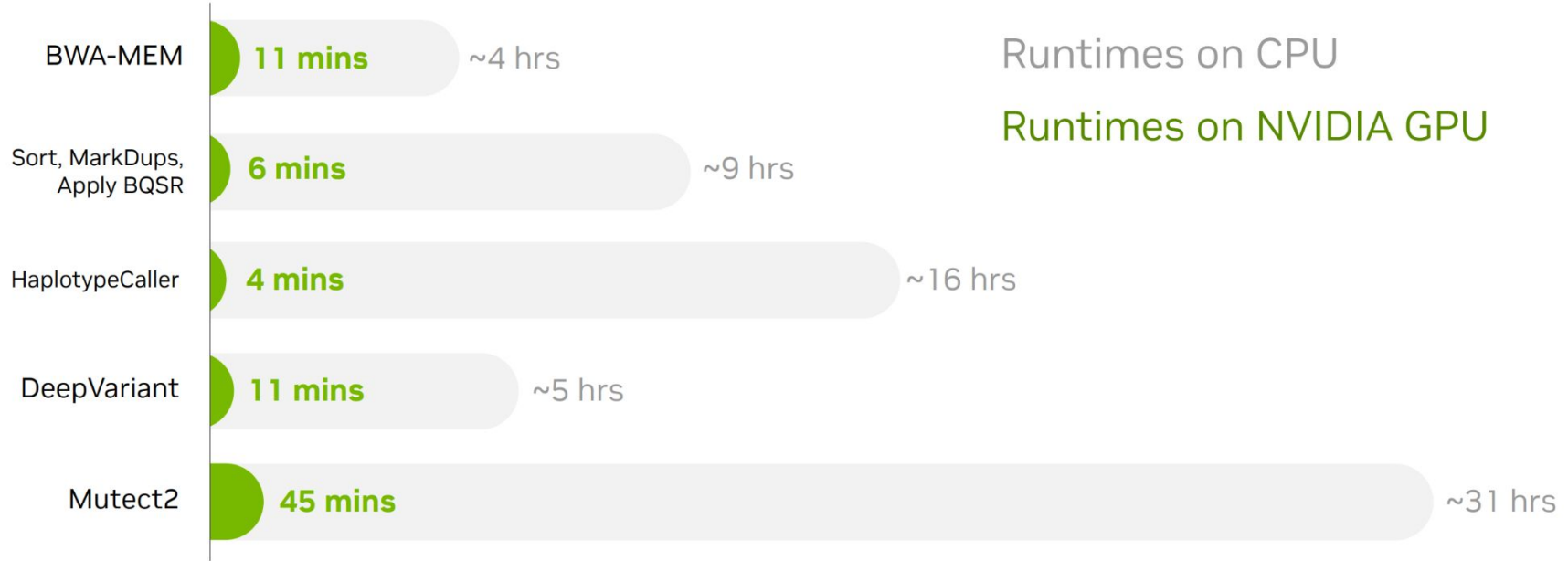
Tools Supported by Parabricks on Minerva

ALIGNMENT	PREPROCESSING	VARIANT CALLING	QUALITY CHECKS	GVCF PROCESSING
fq2bam (bwa-mem)	applybqsr	deepvariant	bammetrics	indexgvcf
STAR	bam2fq	haplotypecaller	collectmultiple metrics	dbsnp
MiniMap2	bqsr	mutectcaller		genotypegvcf
	bamsort	starfusion		preon
	markdup	somatic		postpon
		germline		
		deepvariant germline		
		PacBio germline		

<https://docs.nvidia.com/clara/parabricks/4.3.0/toolreference.html>

Up to 80x Acceleration

Gold-standard results, faster



v3.8 Benchmarks

Dataset: HG002 30x WGS, except Mutect2 on SEQC2 50x WGS

CPU: m5.24xlarge; GPU: 8xA100, except DeepVariant & Mutect2 on 8xV100

Parabricks Testing in the Command Line

```
[choh07@li04e03 ~]$ bsub -P acc_hpcstaff -q gpu -gpu num=1 -R h100nvl -n 8 -R span[hosts=1] -R rusage[mem=8GB] -W 1:00 -XF -Is /bin/bash
Job <205034142> is submitted to queue <gpu>.
<<ssh X11 forwarding job>>
<<Waiting for dispatch ...>>
Warning: Permanently added '172.28.3.144' (ED25519) to the list of known hosts.
<<Starting on lg05e15>>
[choh07@lg05e15 ~]$
```

```
[choh07@lg05e15 ~]$ ml parabricks
The singularity image is /hpc/packages/minerva-centos7/parabricks/4.3.0-1/clara-parabricks_4.3.0-1.sif.
[choh07@lg05e15 ~]$
[choh07@lg05e15 ~]$ PARABRICKS=/hpc/packages/minerva-centos7/parabricks/4.3.0-1/clara-parabricks_4.3.0-1.sif
[choh07@lg05e15 ~]$
[choh07@lg05e15 ~]$ ml
```

Currently Loaded Modules:

1) singularity/3.6.4 2) cuda/12.1.1 3) parabricks/4.3.0-1

Parabricks Testing in the Command Line

```
[choh07@lg05e15 ~]$ singularity exec $PARABRICKS pbrun --help
```

```
Please visit https://docs.nvidia.com/clara/#parabricks for detailed documentation
```

```
usage: pbrun <command> [<args>]
```

```
Help: pbrun -h
```

```
command can be a TOOL or FULL PIPELINE. Example:
```

```
pbrun fq2bam --ref genome.fa --in-fq sample_1.fq.gz sample_2.fq.gz --out-bam sample.bam
```

```
pbrun germline --ref genome.fa --in-fq sample_1.fq.gz sample_2.fq.gz --out-bam sample.bam --out-variants sample.vcf
```

```
command options for standalone TOOL
```

applybqsr	- Apply BQSR report to a BAM file and generate a new BAM file
bam2fq	- Convert a BAM file to FASTQ
bammetrics	- Collect WGS Metrics on a BAM file
bamsort	- Sort a BAM file
bqsr	- Collect BQSR report on a BAM file
collectmultiplemetrics	- Collect multiple classes of metrics on a BAM file
dbsnp	- Annotate variants based on a dbsnp
deepvariant	- Run GPU-DeepVariant for calling germline variants
fq2bam	- Run bwa mem, co-ordinate sorting, marking duplicates, and Base Quality Score Recalibration
fq2bam_meth	- Run GPU-accelerated bwa-meth compatible alignment, co-ordinate sorting, marking duplicates, and Base Quality Score Recalibration
fq2bamfast	- Run newly optimized version of bwa mem, co-ordinate sorting, marking duplicates, and Base Quality Score Recalibration
genotypegvcf	- Convert a GVCf to VCF
haplotypcaller	- Run GPU-HaplotypeCaller for calling germline variants
indexgvcf	- Index a GVCf file
markdup	- Identifies duplicate reads
minimap2	- Align long read sequences against a large reference database to convert FASTQ to BAM/CRAM
mutectcaller	- Run GPU-Mutect2 for tumor-normal analysis
postpon	- Generate the final VCF output of doing mutect pon
prepon	- Build an index for PON file, which is the prerequisite to performing mutect pon
rna_fq2bam	- Run RNA-seq data through the fq2bam pipeline
starfusion	- Identify candidate fusion transcripts supported by Illumina reads

```
command options for commonly used FULL PIPELINES
```

deepvariant_germline	- Run the germline pipeline from FASTQ to VCF using a deep neural network analysis
pacbio_germline	- Run the germline pipeline from FASTQ to VCF by aligning long read sequences with minimap2 and using a deep neural network analysis
germline	- Run the germline pipeline from FASTQ to VCF
somatic	- Run the somatic pipeline from FASTQ to VCF

```
Information about the software
```

Parabricks Testing in the Command Line

```
[choh07@lg05e15 ~]$ singularity exec $PARABRICKS pbrun bam2fq --help
Please visit https://docs.nvidia.com/clara/#parabricks for detailed documentation
```

```
usage: pbrun bam2fq [<args>]
Help: pbrun bam2fq -h
```

Run bam2fq to convert BAM/CRAM to FASTQ.

optional arguments:

-h, --help show this help message and exit

Input Output file options:

Options for Input and Output files for this tool.

--ref REF Path to the reference file. This argument is only required for CRAM input. (default: **None**)
--in-bam IN_BAM Path to the input BAM/CRAM file to convert to fastq.gz. (default: **None**)
--out-prefix OUT_PREFIX Prefix filename for output fastq files. (default: **None**)

Tool Options:

Options specific to the tool.

--out-suffixF OUT_SUFFIXF Output suffix used for paired reads that are first in pair. The suffix must end with ".gz". (default: **_1.fastq.gz**)
--out-suffixF2 OUT_SUFFIXF2 Output suffix used for paired reads that are second in pair. The suffix must end with ".gz". (default: **_2.fastq.gz**)
--out-suffixO OUT_SUFFIXO Output suffix used for orphan/unmatched reads that are first in pair. The suffix must end with ".gz". If no suffix is provided, these reads will be ignored. (default: **None**)
--out-suffixO2 OUT_SUFFIXO2 Output suffix used for orphan/unmatched reads that are second in pair. The suffix must end with ".gz". If no suffix is provided, these reads will be ignored. (default: **None**)
--out-suffixS OUT_SUFFIXS Output suffix used for single-end/unpaired reads. The suffix must end with ".gz". If no suffix is provided, these reads will be ignored. (default: **None**)
--rg-tag RG_TAG Split reads into different fastq files based on the read group tag. Must be either PU or ID. (default: **None**)
--remove-qc-failure Remove reads from the output that have abstract QC **failure**. (default: **None**)

Parabricks Testing in the Command Line

```
[choh07@lg05e15 ~]$ singularity exec --nv $PARABRICKS pbrun fq2bam --help
Please visit https://docs.nvidia.com/clara/#parabricks for detailed documentation
```

```
usage: pbrun fq2bam [<args>]
Help: pbrun fq2bam -h
```

Run GPU-bwa mem, co-ordinate sorting, marking duplicates, and Base Quality Score Recalibration to convert FASTQ to BAM/CRAM.

optional arguments:

-h, --help show this help message and exit

Input Output file options:

Options for Input and Output files for this tool.

--ref REF Path to the reference file. (default: **None**)

--in-fq [IN_FQ [IN_FQ ...]]

Path to the pair-ended FASTQ files followed by optional read groups with quotes (Example:

"@RG\tID:foo\tLB:lib1\tPL:bar\tSM:sample\tPU:foo"). The files must be in fastq or fastq.gz format. All sets of inputs should have a **read** group; otherwise, **none** should have a read group, and it will be automatically added by the pipeline. This option can be repeated multiple times. Example 1: **--in-fq** sampleX_1_1.fastq.gz sampleX_1_2.fastq.gz **--in-fq** sampleX_2_1.fastq.gz sampleX_2_2.fastq.gz.

Example 2: **--in-fq** sampleX_1_1.fastq.gz sampleX_1_2.fastq.gz "@RG\tID:foo\tLB:lib1\tPL:bar\tSM:sample\tPU:unit1" **--in-fq** sampleX_2_1.fastq.gz sampleX_2_2.fastq.gz "@RG\tID:foo2\tLB:lib1\tPL:bar\tSM:sample\tPU:unit2". For the same sample, Read Groups should have the same sample name (SM) and a different ID and PU. (default: **None**)

--in-se-fq [IN_SE_FQ [IN_SE_FQ ...]]

Path to the single-ended FASTQ file followed by optional read group with quotes (Example:

"@RG\tID:foo\tLB:lib1\tPL:bar\tSM:sample\tPU:foo"). The file must be in fastq or fastq.gz format. Either all sets of inputs have a **read** group, or **none** should have one, and it will be automatically added by the pipeline. This option can be repeated multiple times. Example 1: **--in-se-fq** sampleX_1.fastq.gz **--in-se-fq** sampleX_2.fastq.gz . Example 2: **--in-se-fq** sampleX_1.fastq.gz

"@RG\tID:foo\tLB:lib1\tPL:bar\tSM:sample\tPU:unit1" **--in-se-fq** sampleX_2.fastq.gz "@RG\tID:foo2\tLB:lib1\tPL:bar\tSM:sample\tPU:unit2". For the same sample, Read Groups should have the same sample name (SM) and a different ID and PU. (default: **None**)

--in-fq-list IN_FQ_LIST

Path to a file that contains the locations of pair-ended FASTQ files. Each line must contain the location of two FASTQ files followed

Parabricks Testing in the Command Line

```
[choh07@lg05e15 ~]$ singularity exec --nv --bind $DATA_DIR:$DATA_DIR $PARABRICKS pbrun fq2bam --ref $REF --in-fq $INP_fq1 $INP_fq2 --out-bam $OUT_BAM --num-gpus 1
Please visit https://docs.nvidia.com/clara/#parabricks for detailed documentation

[Parabricks Options Msg]: Checking argument compatibility
[Parabricks Options Msg]: Automatically generating ID prefix
[Parabricks Options Msg]: Read group created for
/sc/arion/projects/hpcstaff/jobmix_lsf/parabricks/parabricks_sample/Data/sample_1.fq.gz and
/sc/arion/projects/hpcstaff/jobmix_lsf/parabricks/parabricks_sample/Data/sample_2.fq.gz
[Parabricks Options Msg]: @RG\tID:HK3TJBCX2.1\tLB:lib1\tPL:bar\tSM:sample\tPU:HK3TJBCX2.1
[PB Info 2025-Oct-13 18:23:32] -----
[PB Info 2025-Oct-13 18:23:32] ||                      Parabricks accelerated Genomics Pipeline                      ||
[PB Info 2025-Oct-13 18:23:32] ||                      Version 4.3.0-1                      ||
[PB Info 2025-Oct-13 18:23:32] ||                      GPU-BWA mem, Sorting Phase-I                      ||
[PB Info 2025-Oct-13 18:23:32] -----
[M::bwa_idx_load_from_disk] read 0 ALT contigs
[PB Info 2025-Oct-13 18:23:36] GPU-BWA mem
[PB Info 2025-Oct-13 18:23:36] ProgressMeter      Reads      Base Pairs Aligned
[PB Info 2025-Oct-13 18:23:57] 5043564      580000000
[PB Info 2025-Oct-13 18:24:10] 10087128  1160000000
[PB Info 2025-Oct-13 18:24:22] 15130692  1740000000
[PB Info 2025-Oct-13 18:24:35] 20174256  2320000000
[PB Info 2025-Oct-13 18:24:48] 25217820  2900000000
[PB Info 2025-Oct-13 18:25:00] 30261384  3480000000
[PB Info 2025-Oct-13 18:25:12] 35304948  4060000000
[PB Info 2025-Oct-13 18:25:25] 40348512  4640000000
[PB Info 2025-Oct-13 18:25:38] 45392076  5220000000
[PB Info 2025-Oct-13 18:25:52] 50435640  5800000000
[PB Info 2025-Oct-13 18:26:07]
GPU-BWA Mem time: 151.395909 seconds
[PB Info 2025-Oct-13 18:26:07] GPU-BWA Mem is finished.

[main] CMD: /usr/local/parabricks/binaries/bin/bwa mem -Z ./pbOpts.txt -F 0 /sc/arion/projects/hpcstaff/jobmix_lsf/parabricks/parabricks_sample/Ref/Homo_sapiens_assembl
y38.fasta /sc/arion/projects/hpcstaff/jobmix_lsf/parabricks/parabricks_sample/Data/sample_1.fq.gz /sc/arion/projects/hpcstaff/jobmix_lsf/parabricks/parabricks_sampl
e/Data/sample_2.fq.gz @RG\tID:HK3TJBCX2.1\tLB:lib1\tPL:bar\tSM:sample\tPU:HK3TJBCX2.1
[main] Real time: 154.845 sec; CPU: 1174.795 sec
[PB Info 2025-Oct-13 18:26:07] -----
[PB Info 2025-Oct-13 18:26:07] ||                      Program:                      GPU-BWA mem, Sorting Phase-I                      ||
[PB Info 2025-Oct-13 18:26:07] ||                      Version:                      4.3.0-1                      ||
[PB Info 2025-Oct-13 18:26:07] ||                      Start Time:                  Mon Oct 13 18:23:32 2025                      ||
[PB Info 2025-Oct-13 18:26:07] ||                      End Time:                    Mon Oct 13 18:26:07 2025                      ||
[PB Info 2025-Oct-13 18:26:07] ||                      Total Time:                  2 minutes 35 seconds                      ||
[PB Info 2025-Oct-13 18:26:07] -----
```

Running Parabricks (fq2bam)

Run Parabricks on Minerva

```
singularity exec --nv --bind $WORK_DIR:$WORK_DIR \  
  $PARABRICKS pbrun fq2bam \  
  --ref $WORK_DIR/${REFERENCE_FILE} \  
  --in-fq $WORK_DIR/${INPUT_FASTQ_1} $WORK_DIR/${INPUT_FASTQ_2} \  
  --knownSites $WORK_DIR/${KNOWN_SITES_FILE} \  
  --out-bam $WORK_DIR/${OUTPUT_BAM} \  
  --out-recal-file $WORK_DIR/${OUTPUT_RECAL_FILE}
```

Compatible CPU-based BWA-MEM, GATK4 Commands

```
# Run bwa-mem and pipe the output to create a sorted BAM.  
$ bwa mem \  
  -t 32 \  
  -K 10000000 \  
  -R '@RG\tID:sample_rg1\tLB:lib1\tPL:bar\tSM:sample\tPU:sample_rg1' \  
  <INPUT_DIR>/${REFERENCE_FILE} <INPUT_DIR>/${INPUT_FASTQ_1} <INPUT_DIR>/${INPUT_FASTQ_2} | \  
gatk SortSam \  
  --java-options -Xmx30g \  
  --MAX_RECORDS_IN_RAM 5000000 \  
  -I /dev/stdin \  
  -O cpu.bam \  
  --SORT_ORDER coordinate  
  
# Mark duplicates.  
$ gatk MarkDuplicates \  
  --java-options -Xmx30g \  
  -I cpu.bam \  
  -O mark_dups_cpu.bam \  
  -M metrics.txt  
  
# Generate a BQSR report.  
$ gatk BaseRecalibrator \  
  --java-options -Xmx30g \  
  --input mark_dups_cpu.bam \  
  --output <OUTPUT_DIR>/${OUTPUT_RECAL_FILE} \  
  --known-sites <INPUT_DIR>/${KNOWN_SITES_FILE} \  
  --reference <INPUT_DIR>/${REFERENCE_FILE}
```

Running Parabricks (rna_fq2bam)

Run Parabricks on Minerva

```
singularity exec --nv --bind $WORK_DIR:$WORK_DIR \  
  $PARABRICKS pbrun rna_fq2bam \  
  --in-fq $WORK_DIR/${INPUT_FASTQ_1} $WORK_DIR/${INPUT_FASTQ_2} \  
  --genome-lib-dir $WORK_DIR/${PATH_TO_GENOME_LIBRARY}/ \  
  --output-dir $WORK_DIR/${PATH_TO_OUTPUT_DIRECTORY} \  
  --ref $WORK_DIR/${REFERENCE_FILE} \  
  --out-bam $WORK_DIR/${OUTPUT_BAM} \  
  --read-files-command zcat
```

Compatible CPU-based STAR, GATK4 Commands

```
# STAR Alignment  
$ ./STAR \  
  --genomeDir <INPUT_DIR>/${PATH_TO_GENOME_LIBRARY} \  
  --readFilesIn <INPUT_DIR>/${INPUT_FASTQ_1} <INPUT_DIR>/${INPUT_FASTQ_2} \  
  --outFileNamePrefix <OUTPUT_DIR>/${PATH_TO_OUTPUT_DIRECTORY}/ \  
  --outSAMtype BAM SortedByCoordinate \  
  --readFilesCommand zcat  
  
# Mark Duplicates  
$ gatk MarkDuplicates \  
  --java-options -Xmx30g \  
  -I Aligned.sortedByCoord.out.bam \# This filename is determined by STAR.  
  -O <OUTPUT_DIR>/${NAME_OF_OUTPUT_BAM_FILE} \  
  -M metrics.txt
```

Running Parabricks (haplotypcaller)

Run Parabricks on Minerva

```
singularity exec --nv --bind $WORK_DIR:$WORK_DIR \  
  $PARABRICKS pbrun haplotypcaller \  
  --ref $WORK_DIR/${REFERENCE_FILE} \  
  --in-bam $WORK_DIR/${INPUT_BAM} \  
  --in-recal-file $WORK_DIR/${INPUT_RECAL_FILE} \  
  --out-variants $WORKDIR/${OUTPUT_VCF}
```

Compatible CPU-based GATK4 Commands

```
# Run ApplyBQSR Step  
$ gatk ApplyBQSR \  
  --java-options -Xmx30g \  
  -R Ref/Homo_sapiens_assembly38.fasta \  
  -I mark_dups_cpu.bam \  
  --bqsr-recal-file recal_file.txt \  
  -O cpu_nodups_BQSR.bam  
  
#Run Haplotype Caller  
$ gatk HaplotypeCaller \  
  --java-options -Xmx30g \  
  --input cpu_nodups_BQSR.bam \  
  --output result_cpu.vcf \  
  --reference Ref/Homo_sapiens_assembly38.fasta \  
  --native-pair-hmm-threads 16
```

Batch Job Submission Example (fq2bam)

```
[choh07@li03c02 pbtest]$ cat fq2bam.lsf
#!/bin/bash
#BSUB -J fq2bam                # Job name
#BSUB -P acc_hpcstaff          # allocation account
#BSUB -q gpuexpress            # queue
#BSUB -n 64                    # number of compute cores
#BSUB -W 10                    # walltime in HH:MM
#BSUB -R h10080g
#BSUB -gpu num=4
#BSUB -R rusage[mem=4000]      # 256 GB of memory (4 GB/core * 64 cores)
#BSUB -R span[hosts=1]        # all cores from the same node
#BSUB -o %J.stdout             # output log (%J : JobID)
#BSUB -eo %J.stderr            # error log
#BSUB -L /bin/bash            # Initialize the execution environment

ml parabricks

# Set Input Parameters
PARABRICKS=/hpc/packages/minerva-centos7/parabricks/4.3.0-1/clara-parabricks_4.3.0-1.sif
DATA_DIR=/sc/arion/scratch/choh07/pbtest
INP_fq1=$DATA_DIR/Data/sample_1.fq.gz
INP_fq2=$DATA_DIR/Data/sample_2.fq.gz
REF=$DATA_DIR/Ref/Homo_sapiens_assembly38.fasta
OUT_BAM=$DATA_DIR/output.bam

# Run FQ2BAM (FASTA + FASTQ ==> BAM)
singularity exec \
    --nv --bind $DATA_DIR:$DATA_DIR $PARABRICKS \
    pbrun fq2bam --ref $REF --in-fq $INP_fq1 $INP_fq2 --out-bam $OUT_BAM --num-gpus 4
[choh07@li03c02 pbtest]$
[choh07@li03c02 pbtest]$
[choh07@li03c02 pbtest]$ bsub < fq2bam.lsf
Job <144140780> is submitted to queue <gpuexpress>.
[choh07@li03c02 pbtest]$
```

Batch Job Submission Example (deepvariant)

```
[choh07@li03c02 pbtest]$ cat deepvariant.lsf
#!/bin/bash
#BSUB -J deepvariant          # Job name
#BSUB -P acc_hpcstaff         # allocation account
#BSUB -q gpuexpress           # queue
#BSUB -n 64                   # number of compute cores
#BSUB -W 10                   # walltime in HH:MM
#BSUB -R h10080g              #
#BSUB -gpu num=4              #
#BSUB -R rusage[mem=4000]     # 256 GB of memory (4 GB/core * 64 cores)
#BSUB -R span[hosts=1]       # all cores from the same node
#BSUB -o %J.stdout            # output log (%J : JobID)
#BSUB -eo %J.stderr          # error log
#BSUB -L /bin/bash           # Initialize the execution environment

ml parabricks

# Set Input Parameters
PARABRICKS=/hpc/packages/minerva-centos7/parabricks/4.3.0-1/clara-parabricks_4.3.0-1.sif
DATA_DIR=/sc/arion/scratch/choh07/pbtest
INP_fq1=$DATA_DIR/Data/sample_1.fq.gz
INP_fq2=$DATA_DIR/Data/sample_2.fq.gz
REF=$DATA_DIR/Ref/Homo_sapiens_assembly38.fasta
IN_BAM=$DATA_DIR/output.bam
OUT_VARIANTS=$DATA_DIR/haplotype.vcf

# Run DeepVariant (BAM ==> VCF)
singularity exec \
  --nv --bind $DATA_DIR:$DATA_DIR $PARABRICKS \
  pbrun deepvariant \
    --ref $REF \
    --in-bam $IN_BAM \
    --out-variants $OUT_VARIANTS \
    --num-cpu-threads-per-stream 6 \
    --num-streams-per-gpu 2 \
    --num-gpus 4
```

Monitoring Resource Usage

```
[choh07@li03c02 pbtest]$ bjobs -gpu -l 144140780
```

```
Job <144140780>, Job Name <fq2bam>, User <choh07>, Project <acc_hpcstaff>, Appl  
ication <default>, Status <DONE>, Queue <gpuepress>, Job  
Priority <50>, Command <#!/bin/bash;#BSUB -J fq2bam
```

```
;
```

```
Mon Nov  4 23:44:46: Done successfully. The CPU time used is 2052.0 seconds.
```

```
HOST: lg05g29; CPU_TIME: 2052 seconds
```

```
GPU ID: 0
```

```
Total Execution Time: 132 seconds
```

```
Energy Consumed: 8459 Joules
```

```
SM Utilization (%): Avg 15, Max 93, Min 0
```

```
Memory Utilization (%): Avg 1, Max 9, Min 0
```

```
Max GPU Memory Used: 17448304640 bytes
```

```
GPU ID: 1
```

```
Total Execution Time: 132 seconds
```

```
Energy Consumed: 8017 Joules
```

```
SM Utilization (%): Avg 14, Max 94, Min 0
```

```
Memory Utilization (%): Avg 1, Max 9, Min 0
```

```
Max GPU Memory Used: 17448304640 bytes
```

```
GPU ID: 2
```

```
Total Execution Time: 132 seconds
```

```
Energy Consumed: 8185 Joules
```

```
SM Utilization (%): Avg 13, Max 96, Min 0
```

```
Memory Utilization (%): Avg 1, Max 9, Min 0
```

```
Max GPU Memory Used: 17448304640 bytes
```

```
GPU ID: 3
```

```
Total Execution Time: 132 seconds
```

```
Energy Consumed: 9335 Joules
```

```
SM Utilization (%): Avg 12, Max 94, Min 0
```

```
Memory Utilization (%): Avg 1, Max 9, Min 0
```

```
Max GPU Memory Used: 17448304640 bytes
```

```
GPU Energy Consumed: 33996.000000 Joules
```

```
RUNLIMIT
```

```
10.0 min
```

```
MEMLIMIT
```

```
3.9 G
```

```
MEMORY USAGE:
```

```
MAX MEM: 36.6 Gbytes; AVG MEM: 12.4 Gbytes; MEM Efficiency: 14.66%
```

```
CPU USAGE:
```

```
CPU PEAK: 21.63 ; CPU PEAK DURATION: 60 second(s)
```

```
CPU AVERAGE EFFICIENCY: 20.52% ; CPU PEAK EFFICIENCY: 33.80%
```

Acknowledgements

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Last but not Least

Got a problem? Need a program installed? Send an email to:

hpchelp@hpc.mssm.edu