# 1 Navigate to the data folder

```
cd /opt/data/datasets
ls
ls -lh
```

# 2 FASTQC

# 2.1 demo\_miseq\_lpa5104:

- 1. cd demo miseq lpa5104
- 2. mkdir fastqc report
- 3. /opt/tools/FastQC/fastqc \*.fastq -outdir=./fastqc\_report/
- 4. ls fastqc-report (use autocomplete)
- 5. Download the html file locally and look into both reads of 5296 and of 5118
- 6. merge all R1 and R2 reads separately:

```
cat 5*R1*.fastq > merged_R1.fastq
cat 5*R2*.fastq > merged_R2.fastq
```

7. /opt/tools/FastQC/fastqc merged R1.fastq --

outdir=./fastqc\_report/
8. /opt/tools/FastQC/fastqc merged\_R2.fastq -outdir=./fastqc report/

- 9. download the reports locally and look at it in firefox
- 10. run multigc in top folder of the project (no need to enter in fastqc\_report):

```
multiqc.
```

11. download the multiqc report html file and look at it in firefox

# 2.2 demo\_qc

- ⇒ cd /opt/data/datasets/demo qc (use Tab autocomplete)
- ⇒ mkdir fastqc report
- ⇒ merge R1 and R2 reads separately directly from .gz file:

```
zcat *R1*.fastq > demo_qc_merged_R1.fastq
zcat_*R2*.fastq > demo_qc_merged_R2.fastq
```

# 2.3 demo\_np\_incseq

1. cd /opt/data/datasets/demo np incseq

- 2. ls ./fastq
- 3. too many files
- 4. show first 10 files to learn the file name structure

```
ls ./fastq -lh | head
```

5. Several thousand file cannot be concatenated with cat; use:

```
cd /opt/data/datasets/demo np incseq small/
```

6. finds all files and concatenates them

```
find ./fastq -maxdepth 1 -type f -exec cat {} + > incseq.fastq
```

finds only files with the name structure  $\ensuremath{\mathsf{SP}^{\star}}$ .fastq and concatenates them

```
find ./fastq -maxdepth 1 -name SP*.fastq -exec cat {} + >
fastq/incseq.fastq
```

- 7. mkdir fastqc report
- 8. /opt/tools/FastQC/fastqc fastq/incseq.fastq -outdir=./fastqc report

## 2.4 demo np lpa5104

# 3 Annotation

- 1) cd /opt/data/vcf
- 2) ls -lh
- 3) Use the file /opt/data/vcf/chr6 lpa enhancer plg.vcf
- 4) count number of lines with wc-l

```
chr6 lpa enhancer plg.vcf | wc -1
```

5) look at the file

```
head chr6 lpa enhancer plg.vcf -n 40
```

6) count the number of variant-Lines with grep | wc

```
grep "PASS" chr6_lpa_enhancer_plg.vcf | wc -l
```

7) run vep on chr6\_lpa\_enhancer\_plg.vcf with

```
/opt/tools/ensembl-vep/vep --everything --database -i
/opt/data/vcf/chr6_lpa_enhancer_plg.vcf -o ~/vep_report.txt
```

8) Look into file and show that here is no annotation to LPA, do

```
grep "SYMBOL=LPA" | wc -l
```

- 9) => this is a 1000G file; is on hg19
- 10) Switch to hg19 annotation by

```
/opt/tools/ensembl-vep/vep --everything --database --assembly
GRCh37 --port 3337 -i /opt/data/vcf/chr6_lpa_enhancer_plg.vcf -o
~/vep_report_hg19.txt
```

11) Count the number of lines.

```
wc -l vep_report_hg19.txt
Take note
```

- 12) What is the number of SNPs? The number of SNPs in the vcf (from the grep | wc –l command on "PASS" done previously) returned a lower number of SNPs than the number of hits. Additionally mind that the VCF file is supposed to contain a much larger region of the genome. What is happening?
- 13) Look into the file

```
head vep_report_hg19.txt -n 20
```

=> not enough lines displayed, we are still in the header. Show hundred lines:

```
head vep_report_hg19.txt -n 100
```

- 14) Check the reported transcript IDs => multiple transcripts!
- 15) Filter only form the canonical LPA transcript ENST00000316300 by doing a grep for "ENST00000316300":

```
grep "ENST00000316300" vep_report_hg19.txt >
vep_report_hg19_filtered.txt
```

16) count lines again:

```
wc-I vep_report_hg19_filtered.txt
```

Number of lines is now much lower. We have filtered only for the relevant transcript.

# 17) head vep report hg19 filtered.txt

The column names are missing. How to extract them? Do a grep on a word, which is specific for the column name row, write this into a file and then concatenate the files

⇒ Search for a word, which denotes the column header line

```
head test_vep_hg19.txt -n 100
```

⇒ The Word "Location" is typical for the header. Extract the respective line

```
grep "Location" test_vep_hg19.txt > header.txt
```

 $\Rightarrow$  look into the file:

```
cat header.txt
```

⇒ Concatenate the two files and write a new file with header

```
cat header.txt vep_report_hg19_filtered.txt >
vep_report_hg19_filtered_header.txt
```

18) Do a grep for a list of items

We build quickly a list by writing two SNP names into a file (">>" means append to an existing file)

```
echo "rs10455872" > snplist.txt
echo "rs3798220" >> snplist.txt
```

Now we grep for the contents of the file (option –f)

```
grep -w -f snplist.txt vep_report_hg19_filtered.txt >
listgrep.txt
```

- -w search only for whole words
- -f use the given file as input

# 4 Nanopore data

# 4.1 demo\_np\_lpa5104

1. PORETOOLS STATS & HIST on fast5 files

```
poretools stats fast5/
poretools hist fast5/
```

the hist command cannot be executed on the remote terminal

```
poretools hist --min-length 1000 fast5/
```

use the min-length parameter to specify a lower bound of the read lengths to be displayed in the histogram (to get rid of all very short fragments)

## 2. PAUVRE on fastq

```
pauvre marginplot -n --fastq <fastq_file>

write pauvre report in a file
pauvre marginplot -n --fastq <fastq_file> > pauvre_report.txt

look at report with (several options)
cat pauvre_report.txt
less pauvre_report.txt (hit "q" to exit)
vi pauvre report.txt (write ":q" to exit)
```

## 4.2 demo\_np\_incseq\_small

Several thousand file cannot be concatenated with cat; use:

```
cd /opt/data/datasets/demo_np_incseq_small/
finds all files and concatenates them
find ./fastq -maxdepth 1 -type f -exec cat {} + > incseq.fastq

finds only files with the name structure SP*.fastq and concatenates them
find ./fastq -maxdepth 1 -name SP*.fastq -exec cat {} + > fastq/incseq.fastq

poretools stats fast5/
poretools hist fast5/
pauvre marginplot -n --fastq <fastq_file>
```

#### 4.3 Examples of commands for the last day

This commands are not copy/paste ready but need to be adapted according to your data. Use manual and help page

### **Basecalling**

for help

```
read fast5 basecaller.py -h
```

go to run folder, which contains subfolder with raw FAST5 data

```
cd /opt/data/datasets/demo_np_incseq_small/
mkdir fast5_alba
```

#### 1. Do basecalling of called FAST5 with ALBACORE

```
read_fast5_basecaller.py \
-f FLO-MIN106 \
-k SQK-RBK001 \
--barcoding -r -t 8 \
-i /media/qf0297/Volume/data_np/meth_validation2/fast5_all/ \
-s /media/qf0297/Volume/data_np/meth_validation2/fast5_alba/ \
-n 0 -o fastq,fast5 -q 320000
```

-q: use a number which is greater than the number of FAST5 files

### 2. Kraken (metagenomics analysis)

```
/opt/tools/kraken-tool/kraken --db
/opt/tools/kraken/minikraken_20171013_4GB/ --threads 1 --fastq-
input --preload --output out-folder <input>
/opt/tools/kraken-tool/kraken-report --db minikraken_20171013_4GB/
out-folder > here.report
```

### 3. Methylation analysis (in-vitro methylated PCR product)

### The steps are

- create FASTA files with poretools (MUST be poretools)
- check the number of reads in the fasta. Does it match the number of fast5 reads?
- IMPORTANT: once fasta has bee ncreated do not change the relative position of the files with respect to the fast5 files!
- Align to reference with bwa
- create a sorted BAM with samtools
- do methylation calling with nanopolish
- use the nanopolish script calculate\_methylation\_frequency.py to generate a tabular methylation level summary per position

```
poretools fasta fast5/ > fasta/reads.fasta
grep ">" reads.fasta | wc -l

bwa index meth_ref.fasta
bwa mem -x ont2d meth_ref.fasta fasta/reads.fasta > meth_01.sam
samtools view -bS meth_01.sam > meth_01.bam
samtools sort meth_01.bam sorted_meth_01
samtools index sorted_meth_01.bam

/opt/tools/nanopolish/nanopolish call-methylation --progress -t 8 -r
reads.fasta -g meth_ref.fasta -b sorted_meth_01.bam >
methylation_01.tsv

python
/opt/tools/nanopolish/nanopolish/scripts/calculate_methylation_frequency.py -c 2.5 -i methylation_01.tsv > methylation_01_freq.tsv
```

Use the values from the .tsv file to generate e.g. boxplots of the methylation level using R or online tools like http://shiny.chemgrid.org/boxplotr/