

# Process MHC II DRB

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## Preface

All processing steps for analysing the Illumina MiSeq reads are outlined within this document. Most steps are based on several `python` scripts that may be executed using the command line (e.g. ‘Bash on Ubuntu in Windows’), while R functions can be called within this document.

```
library(magrittr)
source("R/clustering_functions.R")
```

## Prepare metafile

The first step of the processing uses the Qiime script `extract_barcode.py` to reorient forward and reverse reads and trimming barcodes from the sequences. In order to do so, a metafile is needed that gives the forward and reverse primer sequences respectively which allow to recognise the orientation of a read.

```
## R
## read barcodes
barcodes <-
  read.csv("data/mhc_barcode.csv", header = T, sep = ";", skip = 1)

## format left barcode
index_l <-
  barcodes[barcodes[["primer"]] == "Forward",c("Index","Sequence")]
names(index_l) <- c("index_L","BC_L")

## format right barcode
index_r <-
  barcodes[barcodes[["primer"]] == "Reverse",c("Index","Sequence")]
names(index_r) <- c("index_R","BC_R")

## read list of samples and join with barcodes
drb <-
  read.csv("documents/drb-samples-miseq.csv", header = T, sep = "\t") %>%
  dplyr::left_join(., index_l, by = "index_L" ) %>%
  dplyr::left_join(., index_r, by = "index_R")

## create metafile to extract barcodes using QIIME
drb_qiime <- data.frame(
  SampleID =
    paste0(drb[["sample"]], ".", drb[["position"]]),
  BarcodeSequence =
    paste0(drb[["BC_L"]], drb[["BC_R"]]),
  LinkerPrimerSequence =
    read.table("documents/mhc_primer_sequences.csv", header = T, sep = ",")[2,2] %>%
    as.character(),
  ReversePrimer =
    read.table("documents/mhc_primer_sequences.csv", header = T, sep = ",")[2,3] %>%
    as.character(),
```

```
  Description = NA)

drb_qiime$SampleID <-
  stringr::str_replace(drb_qiime$SampleID, "/", ".")

## Manually add '#' before each sample in the created file
write.table(x = drb_qiime,
            file = "miseq_reads/DRB-Pool/drb_barcodes.txt",
            sep = "\t",
            row.names = F,
            quote = F)
```

## Start processing MiSeq reads

```
# bash
#####
# Prerequisites: #
# #
# 'mhc_cluster' needs to be downloaded from https://github.com/mottensmann/mhc_cluster #
# and added to the 'PATH' variable. #
# Vsearch v.2.44, hmmer-3.1b2, usearch10, muscle3.8.31 are expected to be in 'PATH' #
# #
# Folder that are missing in the PATH can be added using the following notation #
# where '~' gives the path starting from the root. #
# Example: export PATH=~/.mhc_cluster:$PATH #
# Here, QIIME is assumed to be part of the conda environment #
# Raw reads may be downloaded as zip archives and saved in a subfolder raw_reads #
#####

## Start conda
## -----
source activate qiime191conda

## Set directory
## -----
cd miseq_reads/DQB-Pool/

## MUSCLE alignment of dqb sequences
## -----
muscle -in hmm/seal_dqb.fasta -out hmm/seal_dqb.afa ;

## create hidden markov model
## -----
hmmbuild hmm/seal_dqb.hmm hmm/seal_dqb.afa ;

## create auxiliary files for hmmscan
## -----
hmmcompress hmm/seal_dqb.hmm ;

## copy to folder lib in repository mhc_cluster
## -----
cd hmm/
find -name "seal_dqb*" -print -exec cp {} ~/.mhc_cluster/lib/ \;
cd ..

## Unzip raw reads
## -----
cd raw_reads
find . -iname "*.gz" -exec gunzip {} \;
cd ..

## Extract and strip barcodes from reads
## -----
extract_barcode.py --input_type barcode_paired_end
-f raw_reads/reads1.fastq -r raw_reads/reads2.fastq
```

```

--bc1_len 8 --bc2_len 8 -m dqb_barcode.txt
-a -o parsed_barcode/ ;

## Filter reads and truncate to 230 bp
## -----
vsearch --fastq_filter parsed_barcode/reads1.fastq
        -fastqout parsed_barcode/reads1_qual_filt.fastq
        -fastq_maxns 0 -fastq_maxee 2 -fastq_trunclen 230

## R
## Filter merged reads
## -----
source("R/call_filter_forward_reads_drb.R")

# bash
## Truncate filtered to get only exon sequences
## -----
fastx_trimmer -i parsed_barcode/reads1_qual_filt_cont_filt.fastq
              -o parsed_barcode/reads1_cluster_input.fastq -f 31 &

## Run MHC clustering pipeline
## -----
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/drb
               -hmm seal_drb.hmm
               -alpha '2.0'

## Explore sensitivity to parameter alpha
## -----
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '0.0' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '0.5' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '1.0' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '1.5' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '2.0' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '2.5' -hmm seal_drb.hmm ;
cluster_mhc2.py -f parsed_barcode/reads1_cluster_input.fastq
               -o clustered_reads/alpha_exploration/drb
               -alpha '3.0' -hmm seal_drb.hmm ;

```

```

# R
## split reads in amplicons
demultiplex_fastq(
reads = "miseq_reads/DRB-Pool/parsed_barcodes/reads1_cluster_input.fastq",
out = "miseq_reads/DRB-Pool/demultiplexed",
outname = "read1.fastq")

# bash
## Apply to samples individually
## -----
cd /demultiplexed/
find . -type d | while read d; do
    (cd $d/
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '0.0' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '0.5' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '1.0' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '1.5' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '2.0' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '2.5' ;
    cluster_mhc2.py -f read1.fastq -o temp -hmm seal_drb.hmm -alpha '3.0' ;
    )
done
cd ../

## R
## Remove temporary created files
remove_mhc_cluster_files(
    parentfolder = "miseq_reads/DRB-Pool/demultiplexed/",
    fastq = "read1.fastq")

## Pool alleles of all individually processed amplicons
filen <- c("temp_pct_1.0_a_0.0_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_0.5_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_1.0_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_1.5_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_2.0_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_2.5_ee_1.0.fixed.otus.fa",
           "temp_pct_1.0_a_3.0_ee_1.0.fixed.otus.fa")
lapply(filen, function(x) {
    pool_zotus(parentfolder = "miseq_reads/DRB-Pool/demultiplexed/",
               filen = x)})

# bash
## Cluster using based on individually identified Zotus
## -----
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
    -ref demultiplexed/temp_pct_1.0_a_0.0_ee_1.0.fixed.otus.fa_pooled_zotus.fa
    -o reference_based/a_0.0 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
    -ref demultiplexed/temp_pct_1.0_a_0.5_ee_1.0.fixed.otus.fa_pooled_zotus.fa
    -o reference_based/a_0.5 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
    -ref demultiplexed/temp_pct_1.0_a_1.0_ee_1.0.fixed.otus.fa_pooled_zotus.fa
    -o reference_based/a_1.0 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq

```

```
-ref demultiplexed/temp_pct_1.0_a_1.5_ee_1.0.fixed.otus.fa_pooled_zotus.fa
-o reference_based/a_1.5 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
-ref demultiplexed/temp_pct_1.0_a_2.0_ee_1.0.fixed.otus.fa_pooled_zotus.fa
-o reference_based/a_2.0 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
-ref demultiplexed/temp_pct_1.0_a_2.5_ee_1.0.fixed.otus.fa_pooled_zotus.fa
-o reference_based/a_2.5 ;
cluster_against_db.py -f parsed_barcodes/reads1_cluster_input.fastq
-ref demultiplexed/temp_pct_1.0_a_3.0_ee_1.0.fixed.otus.fa_pooled_zotus.fa
-o reference_based/a_3.0 ;
```

---

```
sessionInfo()
> R version 3.4.3 (2017-11-30)
> Platform: x86_64-w64-mingw32/x64 (64-bit)
> Running under: Windows 10 x64 (build 16299)
>
> Matrix products: default
>
> locale:
> [1] LC_COLLATE=English_United Kingdom.1252
> [2] LC_CTYPE=English_United Kingdom.1252
> [3] LC_MONETARY=English_United Kingdom.1252
> [4] LC_NUMERIC=C
> [5] LC_TIME=English_United Kingdom.1252
>
> attached base packages:
> [1] stats      graphics  grDevices  utils
> [5] datasets  methods   base
>
> other attached packages:
> [1] magrittr_1.5 knitr_1.17
>
> loaded via a namespace (and not attached):
> [1] compiler_3.4.3 backports_1.1.2
> [3] rprojroot_1.3-1 tools_3.4.3
> [5] htmltools_0.3.6 yaml_2.1.16
> [7] Rcpp_0.12.14    stringi_1.1.6
> [9] rmarkdown_1.8  stringr_1.2.0
> [11] digest_0.6.13  evaluate_0.10.1
```

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